Atmos. Chem. Phys., 16, 5467–5479, 2016 www.atmos-chem-phys.net/16/5467/2016/ doi:10.5194/acp-16-5467-2016 © Author(s) 2016. CC Attribution 3.0 License.





The global tropospheric ammonia distribution as seen in the 13-year AIRS measurement record

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Received: 13 October 2015 – Published in Atmos. Chem. Phys. Discuss.: 21 December 2015 Revised: 8 April 2016 – Accepted: 10 April 2016 – Published: 2 May 2016

Abstract. Ammonia (NH₃) plays an increasingly important role in the global biogeochemical cycle of reactive nitrogen as well as in aerosol formation and climate. We present extensive and nearly continuous global ammonia measurements made by the Atmospheric Infrared Sounder (AIRS) from the Aqua satellite to identify and quantify major persistent and episodic sources as well as to characterize seasonality. We examine the 13-year period from September 2002 through August 2015 with a retrieval algorithm using an optimal estimation technique with a set of three, spatially and temporally uniform a priori profiles. Vertical profiles show good agreement (\sim 5–15 %) between AIRS NH₃ and the in situ profiles from the winter 2013 DISCOVER-AQ (DISCOVER-Air Quality) field campaign in central California, despite the likely biases due to spatial resolution differences between the two instruments. The AIRS instrument captures the strongest consistent NH₃ concentrations due to emissions from the anthropogenic (agricultural) source regions, such as South Asia (India/Pakistan), China, the United States (US), parts of Europe, Southeast (SE) Asia (Thailand/Myanmar/Laos), the central portion of South America, as well as Western and Northern Africa. These correspond primarily to irrigated croplands, as well as regions with heavy precipitation, with extensive animal feeding operations and fertilizer applications where a summer maximum and a secondary spring maximum are reliably observable. In the Southern Hemisphere (SH) regular agricultural fires contribute to a spring maximum. Regions of strong episodic emissions include Russia and Alaska as well as parts of South

America, Africa, and Indonesia. Biomass burning, especially wildfires, dominate these episodic NH₃ high concentrations.

1 Introduction

Global ammonia (NH₃) emissions are increasing due to the increased agricultural livestock numbers coupled with the increasing use of nitrogen fertilization. Atmospheric ammonia has impacts upon local scales, acidification and eutrophication of the ecosystems, and international (transboundary), as well as local, scales through formation of fine ammonium containing aerosols (Sutton et al., 2007, 2008; Galloway et al., 2008; Erisman et al., 2013). Ammonia reacts rapidly with sulfuric (H_2SO_4), nitric (HNO_3), and hydrochloric (HCl) acids to form a large fraction of secondary aerosols, i.e., fine particulate matter (PM_{2.5}) (particles less than 2.5 micrometers in diameter) (Malm et al., 2004). These ammonium containing aerosols affect Earth's radiative balance, both directly by scattering incoming radiation and indirectly by acting as cloud condensation nuclei (e.g., Adams et al., 2001; Martin et al., 2004; Abbatt et al., 2006; Wang et al., 2008; Henze et al., 2012). A large percentage of PM2.5 can penetrate human respiratory systems and deposit in the lungs and alveolar regions, thus endangering public health (e.g., Pope et al., 2002). Ammonia deposition modifies the transport lifetimes, and deposition patterns of sulfur dioxide (SO₂) and nitrogen dioxide (NO_x) (Wang et al., 2008; Henze et al., 2012). Additionally, ammonia increases the concentrations of the greenhouse gas nitrous oxide (N_2O) (EPA, 2011) and, together with NH_4^+ content in soils, NH_3 is involved in CH_4 production and release (Fowler et al., 2009). NH_3 can also contribute to increases in radiative forcing through conversion of organic carbon (OC) into brown carbon (BrC) (Up-dyke et al., 2012). Therefore, monitoring NH_3 global distribution of sources is important to human health, with respect to air and water quality, and climate change.

Atmospheric ammonia concentrations have been modeled from a three-dimensional coupled oxidant-aerosol model (GEOS-Chem) (Bey et al., 2001) to estimate natural and transboundary pollution influences on sulfate-nitrateammonium aerosol concentrations in the United States (US) (Park et al., 2004). We used the simulated NH₃ fields from GEOS-Chem as the retrieval a priori for this study. A number of ammonia-related science studies and top-down inventory studies are based on GEOS-Chem and its adjoint (Henze et al., 2007; Heald et al., 2012; Paulot et al., 2013, 2014; Zhu et al., 2013; Paulot and Jacob, 2014). The model's ammonia emissions were based on annual data from the 1990 $1^{\circ} \times 1^{\circ}$ GEIA inventory of Bouwman et al. (1997). Table 1b from Park et al. (2004) shows a summary of global and contiguous US ammonia emissions for 2001. The inventory's categories include anthropogenic sources: domesticated animals, fertilizers, human bodies, industry, fossil fuels, and natural sources: oceans, crops, soils, and wild animals. Additional emissions from biomass burning and biofuel used were computed using the global inventories of Duncan et al. (2003) and Yevich and Logan (2003), with an emission factor of 1.3 g NH₃ per kilogram dry mass burned (Andreae and Merlet, 2001). For the emissions from domesticated animals and soils, the GEOS-Chem model used the exponential dependencies on temperature reported by Aneja et al. (2000) and Roelle and Aneja (2002), respectively. Ammonia emissions from crops and fertilizers were assumed to vary seasonally with the number of daylight hours (Adams et al., 1999). Seasonal variations in biomass burning and biofuel emissions in the model were specified based on satellite observations (Duncan et al., 2003) and the heating degree-days approach (Park et al., 2004). The GEOS-Chem model can be used to generate 3-D global monthly mean fields of NH₃ concentrations, or higher temporal resolutions (e.g., daily or hourly), for various years.

Satellite remote sensing offers unique opportunities to monitor environmental variables with relatively high temporal and spatial coverages. Ammonia measurements with large, daily global coverage are challenging and lacking, partly due to the relatively short (hours to a day) lifetime of NH₃ near the Earth's surface, and partly because its retrievals require high sensitivity that can be only obtained from areas with high thermal contrast (TC, the temperature difference between that of the surface temperature and of the first discernable atmospheric layer) near the surface (Clarisse et al., 2010). First measurements of ammonia from space were reported over Beijing and San Diego, CA areas, as examples, with the Tropospheric Emission Spectrometer (TES, Beer et al., 2008) and in biomass burning plumes with the Infrared Atmospheric Sounding Interferometer (IASI, Coheur et al., 2009) satellite. Shephard et al. (2011) documented the TES ammonia retrieval methodology. TES NH₃ data have been utilized jointly with GEOS-Chem in various emission source studies (e.g. Alvarado et al., 2011; Pinder et al., 2011; Walker et al., 2012; Zhu et al., 2013). Luo et al. (2015) compared TES NH₃ versus carbon monoxide (CO) ratios, using data from the year 2007, to those of the GEOS-Chem model with a focus on biomass burning emissions using TES representative volume mixing ratio values (Shephard et al., 2011).

The first global map of ammonia was created from IASI measurements by correlating observed brightness temperature differences between strong NH₃ absorbing channels and weak ones to NH₃ total columns using averaged data sets from 2008 (Clarisse et al., 2009). It was later concluded that this method tends to underestimate the global emission inventories at a number of global NH₃ hotspots using IASI radiances. Clarisse et al. (2010) examined the ammonia amounts in the San Joaquin Valley of California in the US using an optimal estimation (OE) retrieval method (Rodgers, 2000) with a global uniform a priori and IASI radiances and compared them with TES measurements. They studied the factors influencing the ability to use satellite infrared (IR) instruments to retrieve accurate NH3 columns and concentrations, finding that the main factors were NH₃ concentrations and thermal contrast. They concluded that through retrieval and forward radiative transfer model runs, if both of the NH₃ concentrations and thermal contrast are large enough, it is possible to quantify ammonia near the lowest level of the atmosphere. R'Honi et al. (2013) discussed the elevated concentrations of NH3 and HCOOH emitted by the 2010 Russian wildfires. Additional IASI NH₃ algorithm approaches and validations are also discussed by Van Damme et al. (2014, 2015) and Whitburn et al. (2015). Heald et al. (2012) used IASI ammonia products jointly with the GEOS-Chem output to study inorganic aerosol loading and atmospheric ammonia concentrations over the US.

Global ammonia sources and variability based on continuous monitoring with longer than a decade record (13 years) have not been hitherto available. This study introduces a newly developed daily and global ammonia product from the Atmospheric Infrared Sounder (AIRS) on the NASA EOS Aqua satellite, spanning September 2002 through August 2015. The AIRS orbit covers nearly the entire globe twice daily, and due to cloud clearing, recovers up to 70 % of cloudy coverage (Susskind et al., 2003; Warner et al., 2013). Additionally, AIRS is in the afternoon Equator crossing time; and therefore, it offers high sensitivity due to higher surface temperature and provides higher thermal contrast to NH₃ measurements.

In the next section, we detail the methodology used to develop the global products of NH_3 and present the discussions for data quality. In Sect. 3, we show examples of validation cases using in situ data from a recent NASA

2 Methodology

The AIRS instrument is a grating spectrometer with 2378 separate spectral channels between 650 and 2670 cm⁻¹ (15.3–3.8 µm) with a spectral resolving power on the order of 1200. Twelve channels of the AIRS radiances in the window regions (860–875, 928–932, and 965–967 cm⁻¹) are currently used to retrieve NH₃. These channels are carefully selected so that the retrievals are based on the NH₃ sensitivity, while the effects of the surface and overlapping gases are minimized. AIRS cloud clearing, described by Susskind et al. (2003), increases the data coverage significantly to nearly 50-70% of the total measurements, instead of the pure clear coverage of approximately 10–15% at a 13.5 km² singleview pixel size (Warner et al., 2013). AIRS NH₃ retrievals are based on the cloud-cleared radiances (CCRs) from AIRS L2 products. The averaging kernel (AK) peaks at about 918 hPa giving AIRS good sensitivity to lower tropospheric NH₃ because the planetary boundary layer generally extends above this altitude at the overpass local time of 1:30 p.m.

The algorithm used in this AIRS NH₃ study was based on a retrieval module developed for AIRS carbon monoxide (CO) products (Warner et al., 2010). This module was built upon and added to the current AIRS operational system or team algorithm (Susskind et al., 2003), but used a different minimization method. The NH3 module uses AIRS Version 6 (V6) Level 2 (L2) profiles and errors from the previous retrieval steps (i.e., surface, clouds, water vapor, ozone, methane, CO) as input to the AIRS forward model the stand-alone AIRS radiative transfer algorithm (SARTA) (Strow et al., 2003). We used SARTA with the addition of NH₃ as a variable gas, which was carried out by co-author Strow and co-workers, since the official AIRS forward model does not include NH3 absorption as a variable. AIRS NH3 retrievals use an OE method following the formulations given by Rodgers (2000), and also described by Pan et al. (1998). The OE retrieval output quantities not only include the NH3 concentrations, but also provide the AKs, the error covariance, and the degrees of freedom for signal (DOFS), which benefit model verifications and data assimilation by using well-quantified errors.

Given a model of the instrument's signals, in the OE method, the forward equation for the NH_3 profile retrieval problem can be written as

$$\mathbf{y} = f(\mathbf{x}, \mathbf{b}) + \mathbf{n}_{\varepsilon},\tag{1}$$

where y is the vector of measured radiances, x is the state vector (variables to be retrieved from the measurements), b represents all other parameters used by the forward model, f(x, b) is the forward model function, and n_{ε} is the instru-

ment noise. For the variables that obey a Gaussian distribution, this inverse problem is equivalent to the maximum likelihood solution. By using a Newtonian iteration; the solution to Eq. (1) can be written as (Rodgers, 2000)

$$\mathbf{x}_{n+1} = \mathbf{x}_{a} + \mathbf{C}_{a}\mathbf{K}_{n}^{T} \left(\mathbf{K}_{n}\mathbf{C}_{a}\mathbf{K}_{n}^{T} + \mathbf{C}_{e}\right)^{-1} [\mathbf{y} - \mathbf{y}_{n} - \mathbf{K}_{n}(\mathbf{x}_{a} - \mathbf{x}_{n})]$$
(2)

where n is the order of iteration and C_e is the measurement error covariance matrix.

 $\mathbf{K}_n = \partial f(\mathbf{x}, \mathbf{b}) / \partial \mathbf{x}$ is the jacobian matrix for iteration \mathbf{n} , which is the sensitivity matrix of the forward model to the state vector \mathbf{x} . \mathbf{x}_a is the mean of the a priori distribution and \mathbf{C}_a is the a priori error covariance matrix for \mathbf{x}_a .

As defined by the retrieval formulations, the AKs are computed using the following equation:

$$\mathbf{A} = \mathbf{C}_{\mathbf{a}}\mathbf{K}^{T}(\mathbf{K}\mathbf{C}_{\mathbf{a}}\mathbf{K}^{T} + \mathbf{C}_{\mathbf{e}})^{-1}\mathbf{K}$$
(3)

and,

$$\mathbf{x}' \approx \mathbf{A}\mathbf{x} + (\mathbf{I} - \mathbf{A})\mathbf{x}_{\mathbf{a}},\tag{4}$$

where **I** represents the identity matrix and x is the true state. Equation (4) states that in the absence of other error sources the retrieved state is a weighted mean of the true state and the a priori state, with the weight **A** for the true state and **I** – **A** for the a priori. This shows the importance of AKs as diagnostics of the retrieval. The closer the matrix **A** is to the identity matrix the more the retrieved state resembles the true state.

The optimal estimation method requires an a priori mean profile and a corresponding error covariance matrix that represent the current knowledge of the geophysical property, i.e., NH₃, prior to the retrieval. Due to the high spatial variability and short lifetime of NH₃, a simple fixed a priori for all emission scenarios is not appropriate. We developed a global mean, multi-year averaged (2003-2012), three-tier a priori from GEOS-Chem model (v9-02) simulations for high, moderate, and low pollution. We used GEOS-5 MERRA data sets from the NASA Global Modeling and Assimilation Office (Rienecker et al., 2011) to drive the meteorological fields in the GEOS-Chem simulations. Figure 1 shows the a priori mean profiles (solid curve with squares) and the error covariance matrices (horizontal bars) for the low (left panel), the moderate (middle panel), and the high pollution (right panel), respectively. The high pollution range was defined by profiles with volume mixing ratios (VMRs) greater than or equal to 5 parts-per-billion-volume (ppbv) at the surface. The moderate pollution range includes the profiles with surface VMRs greater than or equal to 1 ppbv but less than 5 ppbv,



Figure 1. The a priori profiles and the square root of the diagonal terms of the error covariance matrices for the low pollution (left panel), the moderate pollution (middle panel), and the high pollution scenarios (right panel), respectively.

or greater than 1 ppbv at any level between the surface and 500 hPa. The low pollution is then defined as being lower than the lower bounds of the moderate pollution range. The profiles were adjusted to match AIRS forward model levels. The modeled profiles are extrapolated near the surface with additional constraints to eliminate high values in the model near the surface, which are likely seen by satellite sensors.

The same set of the three-tier a priori profiles is used globally and throughout the AIRS data record. Thus, any spatial and temporal NH₃ variations detected using this algorithm are from AIRS measurements. To select one of the three a priori profiles for each AIRS pixel, we examine the brightness temperature difference between a strong and a weak channel, divided by the measurement noise of the strong channel, defined as a "difference of brightness temperature index" (DBTI). This is similar to the method used by TES NH₃ and described by Shephard et al. (2011). The DBTIs vary with meteorological conditions and, most importantly, the thermal contrast at the surface. To take these effects into account, we simulate the relationship between the brightness temperature differences and TC under various meteorological conditions using SARTA. We randomly picked 13790 profiles from AIRS L2 products over land from the months of January, April, July, and October in years 2003, 2008, and 2011. We then perturbed the NH₃ values spanning the three a priori mean profiles using the range of 0-100 ppbv multiplied by a random number for each atmospheric profile. The observed brightness temperatures are compared with the simulated values at a given TC to determine the level of a priori for the full retrievals. Figure 2 depicts a relationship between the DBTI and DOFS for the three emission levels with low emissions in blue, moderate emissions in green, and high emissions in red. The higher DBTIs are correlated with higher DOFS, which represent higher surface thermal contrast (Deeter et al., 2007).

The NH₃ retrieval quality assurance levels are determined based on the retrieval sensitivities under various meteorological and surface conditions using the AKs and the DOFS.



Figure 2. Correlation between the DBTI (difference of brightness temperature index) and DOFS (degrees of freedom for signal) for the three emission scenarios with low pollution in blue, moderate pollution in green, and high pollution in red.

We also take into account the performance of the retrievals against surface thermal contrasts from AIRS products. Additionally, we examine the retrieval residuals, χ^2 , and the number of iterations to set proper quality assurance flags. The retrieval residuals in Kelvin (K) are defined by the square root of the mean variance of the observed brightness temperatures minus calculated. The NH₃ retrieval quality is affected by the meteorological properties, such as the vertical temperature and water vapor profiles, surface temperatures, and emissivity, which are used to model the atmosphere. We also adapt the error information provided by the AIRS CCR for the relevant channels, which includes meteorological quantities that are used in deriving the AIRS CCR (http://disc.sci.gsfc.nasa.gov/AIRS/documentation/ v6 docs/v6releasedocs-1/V6 Level 2 Cloud Cleared Radiances.pdf). This error information is flagged by Q0, Q1, and Q2 with Q0 having the highest quality and Q2 being unusable. In the remaining discussions of this study, we used χ^2 between 0.9 and 27, considering that the channels used



Figure 3. AIRS NH₃ validation against CRDS (the cavity ring down spectrometer) spiral profiles collected during the DISCOVER-AQ CA (16 January–6 February 2013). The red curves represent AIRS retrieved profiles, gray curves are the a priori profiles, green solid lines are in situ spiral profiles, and the blue dashed lines are the convolved profiles using AIRS NH₃ AKs. The *x* axis is linear from 0 to 25 ppbv and logarithmic from 25 to 150 ppbv.

are not all spectrally independent. The number of iterations limit was set at 10, meanwhile, only the cases with retrieval residuals less than 1 K are used. We also excluded cases with the surface thermal contrast between -4 and +4 K, to avoid ambiguous a priori levels; however, this primarily affects areas over the global oceans. Any additional screening of the data for higher quality requirements, e.g., the use of DOFS, will be discussed case by case. Although we have developed AIRS NH₃ products for all available data sets, only the daytime and land cases are discussed in this study. Additionally, only radiances with quality flag as Q0 are selected for the discussions in the following sections to ensure the best accuracy.

3 Validation with in situ measurements

Validations of retrievals using in situ measurements are vital to quantifying uncertainties in the concentrations, sources, transport patterns, and trends using satellite data. Direct measurements of tropospheric NH₃ are relatively sparse and in situ measurements above the ground level, necessary to validate satellite retrievals, are available for only limited locations and time periods (e.g. Nowak et al., 2007, 2010, 2012). Validation of AIRS NH₃ data sets with available in situ measurements is a continuous effort as more in situ measurements become available. As an example of our validation effort, we use the DISCOVER-AQ NH₃ measurements over California (https://www-air.larc.nasa.gov/cgi-bin/ArcView/discover-aq.ca-2013). The sampling inlet and NH₃ calibration set-up used during DISCOVER AQ

with the cavity ring down spectrometer (CRDS) (G2103, Picarro Inc.) is the same as used with the Chemical Ionization Mass Spectrometry (CIMS) and described in Nowak et al. (2007). The CRDS, aboard the NASA P-3B aircraft during DISCOVER-AQ CA, data period covers 16 January to 6 February 2013. The in situ NH₃ vertical profiles were made in the Southern San Joaquin Valley of California. This region inside the central valley of California, between the coastal mountains in the west and the Sierra Nevada Mountains in the east, consists largely of farmland with scattered dairy farms. Although most of the area is rural, the profiles were made near the small cities of Hanford and Corcoran. We only select spiral profiles from the flights within 45 km of the center of the retrieved AIRS profiles, for the closest match, and within 3 h of the measurement window, similar to the method used for AIRS CO validation (Warner et al., 2006).

Figure 3 shows four retrieval profiles that show high NH_3 concentrations and meet the matching criteria, where the red curves represent AIRS retrieved profiles, gray curves are the a priori profiles, green solid lines are in situ spiral profiles, and the blue dashed lines are the convolved in situ profiles by AIRS NH_3 AKs. The in situ spiral profiles are taken by flying an aircraft in the spiral shape in descending or ascending order near a central location, hence are the closest to being the true vertical profiles. Note that in Fig. 3, the *x* axis is linear from 0 to 25 ppbv and logarithmic from 25 to 150 ppbv. The convolved in situ profiles take into account satellite retrieval sensitivities, making them appropriate to compare against satellite retrievals (Rodgers and Connor, 2003). The convolution calculations follow Eqs. (3) and (4) in Sect. 2. The top



Figure 4. (upper panel) AIRS global NH₃ VMRs at 918 hPa, averaged from September 2002 through August 2015. The colorbar is linear from 0 to 5 ppbv and 5 to 10 ppbv, but with different increments. (lower panel) The total occurrences (number of days) of high concentrations (VMRs > 1.0 ppbv at 918 hPa) in the 13-year period. Red/blue colors indicate relatively high/low occurrences of high concentrations, respectively.

left panel shows a case measured on 16 January 2013 with the retrieval quality at 0, DOFS at 0.64, χ^2 at 1.91, the retrieval residual at 0.07 K, and the measurement time differences at 1.31 h. The distance between the in situ profile and the center of the AIRS profile is approximately 13.5 km. The top right panel shows four in situ profiles from 21 January 2013 with AIRS retrieved profile quality at 0, DOFS at 0.66, χ^2 at 1.26, the retrieval residual at 0.07 K, the time differences ranging from 0.58 to 1.68 h, and the distance differences at approximately 56 km for all four profiles. The two profiles in the bottom left panel are also from 21 January 2013, with quality at 0, DOFS at 0.83, χ^2 at 0.31, the retrieval residual at 0.06 K. The time differences to the AIRS retrieved profile are 1.02 and -1.25 h, and the distances are 38.3 and 38.7 km, respectively. In the bottom right panel, there are four profiles taken from 4 February 2013, with the retrieval profile quality at 0, DOFS at 0.84, χ^2 at 1.1, and the retrieval residual at 0.05 K. The time differences between the in situ and the retrieved profiles are 1.63, 1.40, -0.47, and -0.71, and the distances are 5.1, 45.2, 4.9, and 45.2 km, respectively. Some



Figure 5. AIRS NH₃ DOFS values averaged over September 2002– August 2015 period. Red/blue colors indicate relatively high/low DOFS, respectively.

of the AIRS retrievals collocate with several in situ profiles, and these show substantial spatial variability.

Over regions with high NH₃ in situ concentrations, the convolved in situ profiles agree with the retrievals within > 1to $\sim 3 \text{ ppbv}$ ($\sim 5-15 \%$) near the top of the boundary layer, as seen in the top two panels in Fig. 3. These two AIRS NH₃ profiles show good retrieval sensitivities with DOFS at approximately 0.64 and 0.66, χ^2 at 1.91 and 1.26, and the residual at 0.07 K, respectively. The top left in situ profile is relatively close (13.5 km) to the center of the AIRS pixel, whereas the top right in situ profiles are further away $(\sim 46 \text{ km})$ from the center of the AIRS pixel. When the NH₃ amount is low and there is very little sensitivity in AIRS measurements, the convolved profiles converge to the a priori profiles, as seen in the profiles with low NH3 concentrations in the top right panel and in the bottom left panel. In the bottom right panel, there are four in situ profiles close to the AIRS profile - the AIRS pixel measures the average effect of the area represented by the four in situ profiles. Below 925–950 hPa in height, the in situ NH₃ mixing ratios are significantly higher than the retrieved profiles, indicating a limitation of satellite remote sensing in capturing near surface composition properties. Note again that each AIRS profile covers a surface area of 45 km² where in situ observed NH₃ amounts can vary by a factor of ten. The aircraft in situ flights sometimes are biased by their proximity to strong local point sources. Therefore, the differences between the retrievals and in situ measurements are likely due to sampling issues, although the retrieved profile matches the average of the in situ profiles as discussed above. Nonetheless, the vertical profiles show good agreement ($\sim 5-15$ %) between AIRS NH₃ and the in situ profiles in the examples given above.

4 Global ammonia concentrations

The AIRS global NH₃ VMRs at 918 hPa, averaged from September 2002 through August 2015, are shown in the upper panel of Fig. 4. The lower panel in Fig. 4 shows the total occurrences of elevated concentrations (VMRs \geq 1.0 ppbv at 918 hPa) for the same data set. The occurrences, in numbers of days, are good indicators of the types of emission sources either due to recurring agricultural practices or episodic forest fires. It is important to analyze the NH₃ VMRs together with the occurrences to identify major emission sources. Another important quantity used in the NH₃ source analysis is the retrieval DOFS. Figure 5 shows the AIRS NH₃ DOFS values being in a range of 0.1 to slightly above 1.0. The regions with DOFS greater than 0.4 are generally associated with high NH₃ concentrations and strong signal to noise ratios. We used a threshold level of DOFS of 0.1 to screen the retrievals in the Fig. 4 top panel to eliminate noise and to focus on where AIRS sensitivity is high. Areas with DOFS < 0.1 in the whole data record are indicated in white. The AIRS retrievals are sensitive to NH₃ concentrations in the lowest layer of the atmosphere between 850 hPa and the surface, with sensitivity peaking at approximately 918 hPa based on the retrieved AKs (not shown). Therefore, we use NH₃ VMRs at this level for all discussions in this study. There are diurnal variations in the data sets (not shown) that may be due to a number of factors including the day-night differences of emissions and chemical reactions and possibly measurement sensitivities, which are beyond the scope of this paper and will be studied at a later time. Also note that the missing data over land in certain regions are either due to high elevation (above the 918 hPa altitude level), and therefore not shown, or persistent cloudy days.

Globally, AIRS shows strong NH₃ hotspots from biogenic and anthropogenic sources including South Asia (India/Pakistan), East Asia (China), the central US, parts of Europe, Southeast Asia (Thailand/Myanmar/Laos), the central portion of South America, and Western and Northern Africa, where both the NH₃ VMRs and the frequent occurrences are high. The primary sources for these regions are from human activities, e.g., livestock waste management and other agricultural activities. The NH₃ concentrations over these hot spots vary from ~ 2.5 to above 10 ppbv, averaged over 13 years covering both strong and weak emission periods. Also seen are large regions of high NH3 concentrations due to biomass burning events over Russia, Alaska, South America, Africa, and Indonesia, represented by high VMRs and low occurrences. High concentrations of NH₃ are persistent over South America and reflect emissions from biomass burning that are trapped by the Andes http://earthobservatory.nasa.gov/IOTD/view.php? id=8033&eocn=image&eoci=related image. The hot spot over South Asia corresponds to the heavily populated Indo-Gangetic Plain with plentiful, fertile croplands and extensive livestock, and bounded on the north by the Himalayas (Ya-



Figure 6. (top panel) The NH₃ VMRs from the persistent sources filtered with the collocated occurrences of elevated concentrations (\geq 1.4 ppbv) using a threshold of greater than 40 days; (middle panel) pasture and Cropland Map (http://OurWorldInData.org); and (bottom panel) irrigated agricultural land areas (http://data. worldbank.org). Footnote: The World Bank provided the statement that the maps displayed on the World Bank web site are for reference only and do not imply any judgment on the legal status of any territory, or any endorsement or acceptance of such boundaries.

maji et al., 2004). The absolute maximum on Fig. 4 is found over the Punjab which has the highest population density in Pakistan.

To understand the persistent emission sources, we filtered the NH₃ VMRs with the collocated occurrences of elevated concentrations (\geq 1.4 ppbv) greater than 40 days; and the results are shown in Fig. 6 top panel. Although a sufficient concentration (\geq 1.4 ppbv) threshold is used to calculate occurrences of the persistent sources, we used all VMR values, with DOFS greater than 0.1, for the VMR maps. The persistent NH₃ sources not only include those large regions listed above, but also include small geographical areas such as in the San Joaquin Valley of central California in the US (with low sulfur emissions and where livestock are plentiful); the Po Valley, Italy; Fergana Valley, Uzbekistan; Azerbaijan; the Nile Delta and along the banks of the Nile River in Egypt; and the Sichuan Basin in China. Some of these source locations are consistent with those previously reported by Clarisse et al. (2009). These emission hotspots are compared with the "Pasture and Cropland Map" (see middle panel in



Figure 7. AIRS NH₃ VMRs at 918 hPa averaged between September 2002 and August 2015 for December–January–February (DJF, upper left panel), March–April–May (MAM, upper right panel), June–July–August (JJA, lower left panel), and September–October–November (SON, lower right panel), with DOFS greater than 0.1 and no cutoff limit for the VMRs. Red/purple colors indicate relatively high/low NH₃ VMRs.



Figure 8. As in Fig. 7 except for the occurrences of high concentrations (VMRs \geq 1 ppbv). Red/blue colors indicate relatively high/low occurrences of high concentrations.

Fig. 6), posted by http://OurWorldInData.org, located at the Institute for New Economic Thinking at the Oxford Martin School. AIRS NH₃ source regions are strongly correlated with cropland areas, e.g., over India, China, the middle US, Western Africa, eastern South America, and Europe. Note that four of the strongest emission regions correspond to high percentage irrigated agricultural areas (see bottom panel in Fig. 6), i.e., over Pakistan, India, northern Italy, and Azerbaijan adjacent to the Caspian Sea. The irrigated agricultural land includes that irrigated by controlled flooding. These data are provided by the World Bank (http://data.worldbank.org) where the color values are the percent agricultural irrigated land of total agricultural land. These irrigation activities are associated with periods of fertilization and ammonia release. Sommer et al. (2004) studied the relationship between the fertilizing time and the ammonia release time and indicated that the fertilizers applied in March can be released in the June to August time frame depending on the amount of precipitation. The irrigation practices may have the same effect as high amounts of precipitation.

Over China, the AIRS retrieval can match high-resolution inventories distinguishing the two major animal husbandry areas in east-central China (Henan, Shandong, and Hebei provinces) as well as Sichuan to the southwest (Huang et al., 2012). Additional weaker, but persistent, NH₃ sources are also seen in the Fig. 6 top panel that are likely related to livestock and agriculture practices. These source regions include areas in eastern North Carolina (consistent with Wu et al., 2007), Arizona near Phoenix, in the east coast of Spain near Barcelona and Águilas, and over large areas in the Netherlands, in Mozambique in Africa, and the Gambela National Park region between Ethiopia and South Sudan.

5 Seasonal variability

Seasonal variations are shown in Fig. 7 in the four NH₃ VMR maps, averaged between September 2002 and August 2015, for December-January-February (DJF, upper left panel), March-April-May (MAM, upper right panel), June-July-August (JJA, lower left panel), and September-October-November (SON, lower right panel), respectively, with DOFS greater than 0.1 and no cutoff for the VMRs. Globally, the highest concentrations are in the NH summer and spring seasons, with the exception from strong biomass burning (BB) sources, i.e., over South America, Southeast Asia, and Russia in the NH fall season. The highest NH₃ concentrations over non-BB dominant regions occur over India, China, the Mid-West US, and parts of Europe in the summer months. The longest high concentration seasons are over northern India, collocated with the measurement of high NH_4^+ in the precipitation over India reported by Kulshrestha et al. (2005). The seasonal NH₃ VMR distributions in China, Europe, and the US are also consistent, to a large extent, with the Paulot et al. (2014) study of agricultural emissions in-



Figure 9. The NH₃ monthly mean variations (solid line) in the NH (upper panel) and the SH (lower panel), respectively. The long-dash lines show the 1σ standard deviation (SD); and the shaded areas represent the maximum and minimum range of each data set.

ventory derived by high-resolution inversion of ammonium wet deposition data. This is especially true for the spring season, as seen in Fig. A1 of Paulot et al. (2014), showing MASAGE_NH3 (Magnitude and Seasonality of Agricultural Emissions for NH₃) emissions of NH₃ from fertilizers.

High average concentrations (Fig. 7) with low frequencies of occurrences (Fig. 8) generally indicate NH₃ from biomass burning (BB). The greatest emissions from BB in the NH appear in the summer months over Siberia and eastern Russia as well as over Alaska, US. The highest concentrations due to BB in the SH appear over South America in September to November (spring for the SH) when precipitation is minimal and burning extensive (Oliveras et al., 2014). Over SE Asia where the dry season and most BB occur in March to May, we find another local maximum (Lin et al., 2013). Over Africa high concentrations from BB occur in the Western and Central regions, although both high concentrations and frequencies of occurrences appear in the Sahel just south of the Sahara in the NH winter. In that region persistent burning of agricultural waste has been reported (Haywood et al., 2008); see also http://rapidfire.sci.gsfc.nasa.gov/cgi-bin/ imagery/firemaps.cgi.

Ammonia seasonal variations are presented (Fig. 9) using the monthly mean VMRs averaged over the 13-year period. Simple hemispheric averages of NH₃ concentrations for all cases do not accurately reflect the seasonality of the important agricultural activities in the NH, due to the mixing with BB cases and low NH₃ regions, as well as regions with missing values due to weeks of persistent cloud cover. To understand how NH₃ emissions vary seasonally due to human activities, we focus on the NH3 concentrations from the continuous emission sources. As in the case of Fig. 6, where we showed continuous sources using screening by the occurrences of elevated concentrations, we select the occurrence thresholds at concentration levels higher than 1.4 ppbv on at least 40 days of the 13-year record. Figure 9 shows the monthly mean variations of NH₃ (solid line) in both the NH (upper panel) and the SH (lower panel); the dashed lines show the $\pm 1\sigma$ (standard deviation, SD) and the shaded areas represent the maximum and minimum range of each data set. In the NH, the high emission period starts in April and the NH₃ concentrations peak in June. The NH average of the VMR concentrations from April through July is in the range of 3.7-4.0 ppbv; and it gradually decreases to the minimum of below 2 ppbv in November-December-January. The range of monthly mean variability between different years is also larger from April to September (at ~ 1 ppbv) than in the winter months (at ~ 0.4 ppbv). The SD decreases from the summer values of 0.6 to 0.3 ppbv in the winter.

Seasonal variation in the SH (lower panel in Fig. 9) shows that the primary sources of NH₃ emission are from BB, as was seen in the NH₃ seasonal maps (e.g., Fig. 7). Although the filtering for the continuous emission sources eliminated some large occasional fires (i.e., over Indonesia), there are still regularly occurring fires, such as those over the central part of South America. The NH₃ concentrations in the SH peaks in September with an average value near 3.5 ppbv and decreases sharply after the SH spring season. The season of high concentrations in the SH is much shorter than in the NH, as demonstrated by the widths of the seasonal distribution curves. The largest SD occurs in September with a magnitude of 2 ppbv, but the variation between different years in the winter is very small (~ 0.25 ppbv).

6 Summary

The AIRS ammonia (NH₃) measurements with a 13-year data record provide global daily maps, identify major source regions, and show seasonal cycles. This enables studies for detailed locations of the sources and their spatial and temporal variations. The AIRS NH₃ products using the optimal estimation (OE) retrievals provide retrieval sensitivity properties, in addition to NH₃ concentrations, such as the averaging kernels (AKs), error covariance matrices, and the degrees of freedom for signal (DOFS). This will facilitate sensor inter-comparisons, model verifications, and data assimilation of satellite retrievals. AIRS measurements can not only capture high biomass burning emissions (e.g., over Russia, Alaska, South America, Africa, and Indonesia) and/or accumulated concentrations such as in various valleys (e.g., San Joaquin Valley, California in the US, the Po Valley, Italy, Fer-

gana Valley, Uzbekistan, and the Sichuan Basin in China), but also emissions due to routine animal feeding and agriculture activities (e.g., Azerbaijan, Nile Delta and along the banks of the Nile River in Egypt, the Mid-West US, North Carolina, US, the east coast of Spain, in the Netherlands, in Mozambique and Ethiopia, Africa, and especially the Indo-Gangetic Plain of South Asia). Over China, the AIRS retrieval can match high-resolution inventories distinguishing the two major animal husbandry areas in east-central China and the Sichuan Basin. Preliminary validation results show excellent agreement with in situ airborne measurements (to within 5-15 % of the retrieved profiles). Note that since each AIRS profile covers a surface area of 45 km² where the NH₃ amounts can vary largely, the simple numerical differences may not be the optimal way to validate satellite ammonia products.

We use frequent occurrences of NH₃ elevated concentrations to select persistent sources. This distinguishes the NH₃ emissions due to human activities versus occasional fires or retrieval noise. We show the persistent ammonia sources correlate well with cropland usage, particularly in regions where irrigation is a routine practice. We show that the hemispheric seasonal variation using sources screened by the high NH₃ frequent occurrences. The NH high NH₃ concentrations occur in the spring and summer with highest from April to July and lowest in November through January. In the SH, the NH₃ concentration is highest in September, this is most likely due to BB emissions shown by the high VRMs and relatively low frequent occurrences.

Detailed examinations of specific regions are needed and will be included in future studies to improve our understanding of the processes that control the NH₃ distribution and variability. The recent NH3 trends from AIRS 13-year measurements will also be a subject of future studies since the scope of this paper is to focus on the algorithm details and the global distributions. Results in this study are focused on land and daytime only. Future studies will include more complicated surface types, i.e., ocean surfaces and regions with lower thermal contrast. The diurnal variations will also be an important topic in the future studies. We have used the pixels with the highest quality cloud-cleared radiances (at 45 km² spatial resolution) defined by the earlier steps of AIRS retrievals, while a future direction will be to also use the higher spatial resolution single-view pixels (at 13.5 km^2) under clear-sky conditions (Warner et al., 2013).

Data availability

Data used for this study are included in the first author Warner's webpage (http://www.atmos.umd.edu/~juying/ ACP_2015_819/) under Data Links.

Acknowledgements. This study was funded by NASA's The Science of Terra and Aqua program under grant numbers NNX11AG39G and NNX12AJ05G. We wish to acknowledge the

GEOS-Chem team, AIRS science team, DISCOVER-AQ team. MERRA data used in this study/project have been provided by GMAO at NASA Goddard Space Flight Center through the NASA GES DISC online archive. Resources supporting this work were provided by the NASA High-End Computing (HEC) Program through the NASA Center for Climate Simulation (NCCS) at Goddard Space Flight Center.

Edited by: W. Lahoz

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