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Retrieval of ammonia from ground-based FTIR solar spectra

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Abstract

We present a retrieval method for ammonia (NH$_3$) total columns from ground-based Fourier Transform InfraRed (FTIR) observations. Observations from Bremen (53.10° N, 8.85° E), Lauder (45.04° S, 169.68° E), Reunion (20.9° S, 55.50° E) and Jungfraujoch (46.55° N, 7.98° E) were used to illustrate the capabilities of the method. NH$_3$ mean total columns ranging three orders of magnitude were obtained with higher values at Bremen (mean of 13.47 × 10$^{15}$ molecules cm$^{-2}$) to the lower values at Jungfraujoch (mean of 0.18 × 10$^{15}$ molecules cm$^{-2}$). In conditions with high surface concentrations of ammonia, as in Bremen, it is possible to retrieve information on the vertical gradient as two layers can be discriminated. The retrieval there is most sensitive to ammonia in the planetary boundary layer, where the trace gas concentration is highest. For conditions with low concentrations only the total column can be retrieved. Combining the systematic and random errors we have a mean total error of 26% for all spectra measured at Bremen (Number of spectra (N) = 554), 30% for all spectra from Lauder (N = 2412), 25% for spectra from Reunion (N = 1262) and 34% for spectra measured at Jungfraujoch (N = 2702). The error is dominated by the systematic uncertainties in the spectroscopy parameters. Station specific seasonal cycles were found to be consistent with known seasonal cycles of the dominant ammonia sources in the station surroundings. The developed retrieval methodology from FTIR-instruments provides a new way to obtain highly time-resolved measurements of ammonia burdens. FTIR-NH$_3$ observations will be useful for understanding the dynamics of ammonia concentrations in the atmosphere and for satellite and model validation. It will also provide additional information to constrain the global ammonia budget.
1 Introduction

Nitrogen emissions in the form of ammonia (NH$_3$), which largely derive from agriculture, have been associated with acidification and eutrophication of soils and surface waters (Krupa, 2003; Vitousek et al., 1997), which may reduce biodiversity in vulnerable ecosystems (Bobbink et al., 1998, 2010). Ammonia also reacts with nitric acid and sulphuric acid to form ammonium salts, which account for a large fraction of particulate matter concentrations (Schaap et al., 2004). Particulate matter is major contributor to smog and is related to negative health impacts (Pope et al., 2009). Moreover ammonium salts play an important role in the radiance balance of the Earth, thus having an impact on climate change (Charlson et al., 1991; Erisman et al., 2007). Recently it was shown that reduced nitrogen also plays a role in the fixation of carbon dioxide (CO$_2$) (Reay et al., 2008). Human activities have increased the global emissions of reactive nitrogen (N$_r$) to the atmosphere (Holland et al., 1999). Current global N$_r$ emissions have been estimated to be almost four times larger compared to pre-industrial levels (Fowler et al., 2013) with NH$_3$ emissions amounting to 49.3 Tg in 2008 (EDGAR-Emission Database for Global Atmospheric Research, 2011). Consequently this has led to large increases in atmospheric nitrogen deposition (Rodhe et al., 2002; Dentener et al., 2006). Biomass burning was found to account for 11% of the global emission budget of NH$_3$ (Bouwman et al., 1997). While the agricultural emissions dominate in the Northern Hemisphere, biomass burning is one of the main sources of the NH$_3$ concentrations in the Southern Hemisphere.

Despite its central role in many environmental threats, little is known about the ammonia budget and its distribution across the globe. Uncertainties in global and regional emission rates are large with errors of more than 50% (Erisman et al., 2007; Sutton et al., 2013). Ammonia concentrations have a large variability in time and space, a short lifetime in the order of hours, and the lack of globally distributed observations hamper our understanding. Surface observations are available but these are not homogenously distributed over the globe and most observation sites are located in the Northern Hemi-
sphere. Most sites provide data with a poor temporal resolution e.g. many observation networks use passive samplers with a sampling time of 2 or 4 weeks (Thijsse et al., 1998; Puchalski et al., 2011) whereas emission and deposition dynamics affect concentrations on the scale of hours to days. Systems with higher sampling frequency such as the AMANDA, MARGA and (denuder) filter packs are available but the number of measurement networks using these instruments is limited as they are often costly to operate (Erisman et al., 2001; Thomas et al., 2009; Mount et al., 2002; Hansen et al., 2003). Moreover, measuring NH₃ is challenging and existing in-situ measurement techniques are often prone to sampling artefacts (von Bobrutzki et al., 2010). Recent advances in open path remote sensing techniques, like (mini-) DOAS systems and open path Quantum Cascaded Laser instruments show large potential to overcome part of these sampling issues (Volten et al., 2012; Miller et al., 2014), but are still in the development stage and not widely applied yet. Another aspect is the lack of vertical information, as most instruments only measure surface concentrations (Erisman et al., 1998, 2007; Van Damme et al., 2015a). Some recent airborne measurements have been made (Nowak et al., 2007, 2010; Leen et al., 2013) but only during dedicated campaigns with limited temporal and spatial coverage. In short, it is very difficult to obtain detailed knowledge on the global ammonia budget using currently available field observations.

Remote sensing products from atmospheric satellite sounders such as the Infrared Atmospheric Sounding Interferometer (IASI), the Tropospheric Emission Spectrometer (TES) and the Cross-track Infrared Sounder (CR-IS) (Van Damme et al., 2014a; Shephard et al., 2011) have become available and show good promise to improve NH₃ concentration monitoring (Van Damme et al., 2014b; Luo et al., 2015; Whitburn et al., 2015). However, these data sets are constrained by the overpass time of the satellite and the atmospheric conditions (cloud coverage, thermal contrast, etc.). Moreover, the uncertainties associated to the data are relatively large, which calls for a detailed evaluation of the data. A recent study (Van Damme et al., 2015a) showed a number of challenges related to the validation. First, reliable hourly in-situ data is sparse.
Second, when not using optimal estimation satellite product as is the case for the IASI-NH$_3$ retrieval, one has to assume a vertical profile to link surface concentrations to a column value. Third, the ground-based observations are often influenced by local sources, whereas the IASI satellite data are averaged observations over a circular footprint with at best a diameter of 12 km. Hence, a measurement methodology that would provide columnar and vertical profiles of ammonia concentrations at a high temporal resolution would be highly beneficial for evaluating the merits of the novel satellite products. Fourier Transform infrared spectrometry (FTIR) provides this methodology. Atmospheric sounders have a long history for validation of satellite products. FTIR observations are already commonly used for the validation of satellite products of among others, Carbon monoxide (CO), Methane (CH$_4$) and Nitrous Oxide (N$_2$O) (Kerzenmacher et al., 2012; Dils et al., 2006).

FTIR spectrometry is a well-established remote sensing technique for the observation of atmospheric trace gases (Rao and Weber, 1992). FTIR has so far been used to estimate ammonia emissions from fires by Paton-Walsh et al. (2005) but only on a campaign basis, not long-term monitoring. There are several monitoring stations with FTIR instruments operated on a regular basis, providing long-term time series for a suite of key tropospheric and stratospheric species, including Carbon Dioxide (CO$_2$), Carbon Monoxide (CO) and Ozone (O$_3$). So far nobody has systematically analysed the FTIR measurements for NH$_3$. We have developed a NH$_3$ retrieval strategies for four Network for detection of Atmospheric Composition Change (NDACC) FTIR stations, spanning very different concentration conditions (polluted and remote sites), in order to obtain time-series of NH$_3$ total columns and show its value for describing temporal variations.

First we present the measurement sites and the retrieval strategies in Sect. 2. We describe the characteristics of the retrieval in Sect. 3.1.1 and the uncertainty budget in Sect. 3.1.2. Section 3.2 constitutes of an interpretation of the results in combination with a comparison with existing datasets of CO total columns and temperature to distinguish between emission sources. We summarize the results in Sect. 4.
2 Measurement sites and retrieval strategies

2.1 Sites description

Ground-based FTIR instruments measure the solar absorption spectra under cloud-free conditions by using a Fourier Transform Spectrometer. These spectra can be analysed by using a line by line model (Pougatchev et al., 1995; Hase et al., 2004, 2006), which models the spectroscopic absorption lines by using known parameters from a spectroscopic database (e.g. HITRAN, Rothman et al., 2013) in combination with the radiative state of the atmosphere, and an optimal estimation inversion scheme (Rodgers, 2000). Information on vertical concentration profiles can be retrieved using the pressure broadening of the absorption lines. For the NDACC network the spectral region measured is the near- to mid-infrared domain (740 to 4250 cm\(^{-1}\), i.e. 13.5 to 2.4 µm) and a HgCdTe or InSb cooled detector (Zander et al., 2008) and a suite of optical filters are used to optimize the signal-to-noise ratio in the complementary spectral regions. Instruments in the network are routinely checked and characterized using laboratory measurements of HBr lines and the linefit software (Hase et al., 1999) to assess the instrument line shape, alignment and measurement noise levels. Four NDACC stations are used in our study, two in each hemisphere:

- The site of Bremen (53.10° N, 8.85° E) is especially suitable to measure variations in ammonia concentrations as the surrounding state, Lower Saxony, which is a region with intensive agricultural activities with high and temporal variable emissions (Dämmgen et al., 2005). In short, the ammonia total columns (molecules cm\(^{-2}\)) at Bremen are expected to reach high values compared to background stations. The Universität Bremen operates a Bruker 125HR spectrometer and a solar tracker by Bruker GmbH, directly on the university campus.

- The Jungfraujoch station (46.55° N, 7.98° E) is a high altitude station (3580 m a.s.l.) located in Switzerland (Zander et al., 2008). There are no large emissions sources surrounding the station itself as it is mostly located in the free

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troposphere. At Jungfraujoch, a Bruker 120HR instrument is in operation since the early 1990s. For the current study, specific for the Jungfraujoch site, we used a subset of spectra recorded during the 2004–2013 time period with apparent solar zenith angles (SZA) between 70 and 85° to increase the capability to retrieve the very low ammonia concentrations.

– The Lauder (45.04° S, 169.68° E) National Institute of Water and Atmospheric Research (NIWA) atmospheric research station in Central Otago, New Zealand at an altitude of 370 (m a.s.l.). Long-term operations started in 1991 with a Bruker 120M (Griffith et al., 2003). This instrument was replaced with a Bruker 120HR in October 2001. Ammonia emissions in the surrounding valley are mostly due to livestock grazing on the pastures and a by-product of seasonal fertilizer application. In recent years there has been an increase in cattle grazing and crop cultivation (EDGAR-Emission Database for Global Atmospheric Research, 2011).

– Reunion Island (20.9° S, 55.50° E) is located in the Indian Ocean to the east of Madagascar. The station is located at the University campus of St.-Denis on the north side of the island. Agricultural activities are mostly related to sugar cane production. The island is prone to some local biomass burning and wild fire events, which are known to emit ammonia. It is also very close to Madagascar, a region with frequent and intense biomass burning events, and it has been found using backward trajectory that the emissions in Madagascar can be transported to Reunion Island within one day (Vigouroux et al., 2009). The measurements used in this study are performed with a Bruker 120M spectrometer. Details on the measurements can be found in Senten et al. (2008) and Vigouroux et al. (2012).

These stations are expected to provide significant differences in variability and levels of ammonia, making them suitable to demonstrate the strength of our retrieval scheme for application across the whole network. A summary of the station descriptions is given in Table 1. CO columns were obtained from the NDACC database to be used for comparison in Sect. 3.
2.2 NH₃ retrieval strategies

The ammonia absorption lines from its ν₂ vibrational band can be observed in the 700–1350 cm⁻¹ wavenumber range, which are also used in the retrieval of satellite products of ammonia (e.g. Clarisse et al., 2009; Van Damme et al., 2014a). In this spectral range the FTIR spectra can be measured using a potassium bromide (KBr) beam-splitter in combination with a mercury cadmium telluride (MCT) nitrogen cooled detector (Zander et al., 2008). The retrieval scheme of trace gas concentrations from FTIR spectra is built on the use of a set of spectral micro windows containing absorption lines of the targeted species, with minimum interference by other atmospheric species or solar lines. Two slightly different set of spectral micro-windows were used at the four stations, but they both use the same main NH₃ absorption lines. The target and interfering species are summarized in Table 2, with the profile retrieved species indicated in bold. To properly estimate ammonia, interfering species like O₃ and water vapour (H₂O) that overlap NH₃ lines in the ν₂ vibrational band have to be accounted for. Two micro windows were chosen that contain as little interfering species as possible. In both sets, the first micro window (MW1) covers the NH₃ absorption line at 930.75 cm⁻¹. At Bremen/Lauder, the choice was to use only isolated NH₃ absorption features to avoid possible problems due to line mixing, therefore the spectral window MW1 is only 1 cm⁻¹ wide [930.32–931.32, MW1]. Figure 1 shows an example of a synthetic spectrum calculated to fit an observation that was measured with the 125HR in Bremen on the 19 April 2010 at 09:59 (UTC) (Solar Zenith Angle of 45°). The NH₃ concentrations on this day were slightly higher than average resulting in slightly stronger NH₃ absorption features in the spectra. The top two figures show the absorption contributions of the absorbing species in both micro windows. The bottom two panels show an enlarged version of the figure to distinguish the interfering species with smaller absorption features. At Reunion Island/Jungfraujoch, MW1 was extended [929.4–931.4, MW1] to cover another NH₃ line at 929.9 cm⁻¹. This improved the retrieval for Reunion Island because at this location the NH₃ concentration levels are much lower than at Bremen and the water vapour
concentrations are much higher. In this high humidity condition, the 930.75 cm\(^{-1}\) line is not isolated from H\(_2\)O, and it improved the retrieval to add the more isolated one at 929.9 cm\(^{-1}\) (see Fig. 2). The main interfering species in MW1 are CO\(_2\), N\(_2\)O, and H\(_2\)O. Minor interfering species are SF\(_6\) and CFC-12. The second window is spanning the NH\(_3\) line at 967.35 cm\(^{-1}\). Again, different widths are used for Bremen/Lauder [966.97–967.68, MW2] and Reunion Island/Jungfraujoch [962.7–970, MW2]. The very weak absorption signatures at Reunion Island and Jungfraujoch are close to the noise level and therefore the whole NH\(_3\) absorption shape is retrieved (about 964–968 cm\(^{-1}\), see Fig. 2) rather than a single line. The main interfering species in MW2 are O\(_3\), CO\(_2\) and H\(_2\)O for all sites. At Reunion Island HDO is also interfering in MW2 as well as the isotopologue \(^{686}\)O\(_3\) (i.e. \(^{16}\)O-\(^{18}\)O-\(^{16}\)O), which has been fitted in addition to the main \(^{666}\)O\(_3\). At Jungfraujoch apart from CO\(_2\), two O\(_3\) isotopologues (the most abundant and \(^{686}\)O\(_3\)) and water vapour which are the main interferences, N\(_2\)O, CFC-12, SF\(_6\) and HDO absorptions are also retrieved. Typical NH\(_3\) absorptions are weak, on the order of a few tenths of a percent. The typical measurement noise (SNR) differs per spectra and site but ranges between \(\sim 250\) at Lauder to \(\sim 450\) at Bremen. Channelling was not an issue in any of the spectra and did not need to be fitted.

Except at Jungfraujoch where S FIT2 is used, the retrieval is performed using the more recent S FIT4.0.9.4 algorithm (Pougatchev et al., 1995; Hase et al., 2004, 2006). Both versions use a form of the optimal estimation method (Rodgers et al., 2000) to retrieve the volume mixing ratios and total columns of NH\(_3\) and makes use of a-priori information (profile and covariance matrix). For Bremen, Lauder and Jungfraujoch the used NH\(_3\) a-priori volume mixing ratios are based on balloon observations (Toon et al., 1999, NH\(_3\) available in dataset but not reported). The shape of the balloon measurements profile was kept constant but extended and scaled to expected surface concentrations. The a-priori surface volume mixing ratio is estimated to be 10 ppb for Bremen (Dammgen et al., 2005). Although the shape of NH\(_3\) profiles do change through time, the largest share of NH\(_3\) is expected to be in the mixing layer, which is represented by the lowest layers in the calculation (Van Damme et al., 2015a; Nowak et al., 2010). At Re-
union Island, the a priori profile was taken from the MOZART model (Louisa Emmons, private communication). The a-priori profile peaks at a higher altitude (4–5 km) instead of the boundary layer as in Bremen, as NH$_3$ is expected to originate mainly from transport of biomass burning emissions at this location. At all stations, the a-priori profiles of the interfering species were taken from the Whole Atmosphere Community Climate Model (Chang et al., 2008).

At Bremen and Lauder, the a priori covariance matrices only have diagonal values corresponding to standard deviations of 100 % for all layers with no interlayer correlation, chosen in relation to the large range of possible concentrations and variations between layers. At Jungfraujoch and Reunion Island, we did not use the a priori covariance matrix as in optimal estimation but the Tikhonov type $L_1$ regularization (e.g. Sussmann et al., 2009) was adopted for the Jungfraujoch retrievals. After several tests, values of 50 and 250 were adopted for the alpha parameter and the signal to noise for inversion, respectively. A Tikhonov regularization with an alpha parameter value of 50 was also adopted for the Reunion retrievals. The signal to noise ratio is calculated for each spectra, the mean value being 365.

Daily temperature and pressure profiles for the meteorological variables were taken from NCEP (National Center for Environment Prediction). For the radiative transfer calculations the profiles were split into about 50 levels, depending slightly on the station, from ground up to 80 km (100 km in the case of Jungfraujoch and Reunion Island). The layers have a typical thickness of 500 m in the troposphere up to 2 km for the higher layers. For the line spectroscopy we use the HITRAN 2012 database (Rothman et al., 2013) in combination with a number of corrections for CO$_2$ (ATMOS, Brown et al., 1996) (except for Jungfraujoch for which the HITRAN lines are used) and sets of pseudo lines generated by G.C. Toon (NASA-JPL) to account for broad unresolved absorptions by heavy molecules (e.g. CFC-12, SF6).

Figure 3 shows an example of the fit in both micro windows for the same measured spectra as used in Fig. 1. The top two and bottom two panels show the calculated
(Green line) and measured spectrum (Blue line) and the residual of both micro windows. The simultaneous fits are good with a standard deviation of 0.15 % in both cases.

3 Results of the FTIR retrievals

3.1 Characteristics of the NH$_3$ retrievals

3.1.1 Vertical information

The retrieved vertical information differs from station to station. The top of Fig. 4 shows for the 4 stations the average NH$_3$ volume mixing ratios (VMR) for each of the retrieved layers (blue line) and the a priori profile that was used as input in the retrieval (green line). The bottom of Fig. 4 shows the averaging kernels for each of the 4 stations averaged over all available observations. As mentioned earlier most of the NH$_3$ at Bremen is in the lowest layers. In Fig. 4 this is also observed as the averaging kernel shows the most sensitivity in the lowest layers (red and green lines for the layers 0.03–0.5 and 0.5–1 km). The combination of the two spectral micro windows on average contain 1.9$^\circ$ of freedom for signal (DOFS) for the Bremen spectra, which means around two independent vertical layers can be retrieved. The two separate layers consist of a layer covering ground–1 km and one that covers 1–6 km height, which can be observed in Fig. 4. It must be taken into account however that the shown averaging kernels are a mean for all observations and thus the retrievable number of layers and combined layer depths vary from spectra to spectra. On average, the Lauder spectra have a DOFS of 1.4. There is only vertical information for multiple layers during periods with increased NH$_3$ total columns, which mostly occur during summer. Similar to Bremen averaging kernels peak near the surface. At Reunion Island only 1.0 DOFS is achieved, which means that only total columns can be retrieved. All the averaging kernels are peaking at the same altitude (about 5 km), which is also the peak of the
a priori profile (Fig. 4). Similar to the Reunion spectra the Jungfraujoch spectra do not have vertical information with a DOF of 1.

### 3.1.2 Uncertainties budget

For the error analysis the posteriori error calculation included in the SFIT4 package is used. The error calculation is based on the error estimation approach by Rodgers (2000). It allows the calculation of the error by attributing errors to each of the parameters used in the retrieval. The error budget can be divided into three contributions, the error due to the forward model parameters, the measurement noise and the error due to the vertical resolution of the retrieval (smoothing error). The assumed uncertainties for the used parameters in the retrieval are listed in Table 3 for the parameters used in the calculation for Bremen, Lauder and Reunion. For Jungfraujoch, the error computation was performed using the perturbation method, the spectra of 2009 to 2011 and the Rodger formalism as explained e.g. in Franco et al. (2015). For Reunion Island, the covariance matrix used for the smoothing error has diagonal elements representing 150% of variability from the a priori profile. To reflect the error in the NCEP temperature profiles we assume an uncertainty of about 2 K in the troposphere and a 5 K uncertainty in the stratosphere. For the uncertainty in the NH$_3$ line parameters we assume values as stated in the HITRAN 2012 database. We assume a conservative 20% uncertainty for the intensity and 10% for both the temperature and pressure broadening coefficients.

The results of the error calculation are listed in in Table 4. Combining the systematic and random errors we have a mean total error of 25.8% for all the spectra measured at Bremen ($N = 554$), 30.2% for the spectra from Lauder ($N = 2412$), 25.2% for the Reunion spectra ($N = 1262$) and 34.23% for the Jungfraujoch spectra ($N = 2702$). The errors are dominated by uncertainties in the spectroscopy. In detail, the random error sources amount to a mean error of 9.1% for the Bremen spectra, which is mostly due to uncertainty in temperature, measurement noise and the zero level of the sensor (i.e. an instrument property). In the case of the systematic error, with a mean error of
23.5\%, the error is for the largest part due to the spectroscopy (i.e. line parameters)
with smaller contributions of the temperature, zero level, phase and the smoothing
error. The results are similar for the Lauder, Reunion and Jungfraujoch spectra with
most of the uncertainty coming from the line parameters. Hence, the pressure and
temperature broadening parameters of the ammonia absorption lines are critical for
the NH$_3$ concentrations.

3.2 Time series

Figure 5 shows the NH$_3$ total columns retrieved from all available spectra from 2004–
2013. Table 5 gives a summary of statistics of the retrieved NH$_3$ columns. Individual
measurements at Bremen (blue) show high concentrations, especially in spring
with an overall mean column total of $13.7 \times 10^{15}$ molecules NH$_3$ cm$^{-2}$ and a root mean
square (RMS) of 20.22 indicating a large variability in the observations. The amplitude
of the spring peaks vary throughout the years, with maxima in 2010 and 2013 reaching
$\sim 93 \times 10^{15}$ and $85 \times 10^{15}$ molecules NH$_3$ cm$^{-2}$. The variability through the years is
carried by changes in meteorology, emissions and timing of the measurements. Gaps
in the data are due to days with overcast and instrument downtimes. The individual
observed columns are sorted into monthly averages to analyse the seasonal variability
and to understand the processes driving the NH$_3$ concentrations. This is shown in Fig. 6
together with monthly averages of surface temperature and CO total columns. NH$_3$
column total concentrations at Bremen (Blue line) have a seasonal cycle with highest
levels during spring, the summer months and autumn. The maximum concentrations
occur around April which is consistent with temporal emission patterns for manure appli-
cation reported for this region (Friedrich and Reis, 2004; Martin et al., 2015; Paulot
et al., 2014). The baseline variability with higher concentrations in summer can be ex-
plained by an increase in volatilization rates of NH$_3$, emitted from livestock housing,
which is driven by animal activity and temperature (Gyldenkaerne et al., 2005). A com-
parison with CO is made to distinguish between agricultural and fire emissions sources.
A correlation between NH$_3$ and CO columns is not observed, which is consistent with agriculture as the dominant source of ammonia.

On average the measurements at Lauder (Fig. 5, red line, top panel) yield a column total of $4.17 \times 10^{15}$ molecules NH$_3$ cm$^{-2}$. These levels are about 1/3 of the concentrations measured at Bremen (blue, top panel). Spectra from Lauder are available for most days in the retrieved time series, which makes it easier to discern peaks and variability. Distinctive peaks are only visible in the summers. Maxima during spring times are not often observed. The peak values are similar in between years, with maxima typically around $30 \times 10^{15}$ molecules NH$_3$ cm$^{-2}$. As for Bremen the RMS of 5.95 reflects a large variability in the observations between individual retrievals. The average error is $1.34 \times 10^{15}$ which is around a quarter of the mean. Figure 6 shows the seasonal cycle of Lauder (red line, top left panel). The seasonal variation of NH$_3$ coincides with that of the atmospheric temperature (red line, bottom right panel) and with the livestock emissions in the surrounding region, which are strongly correlated with temperature.

The bottom panel of Fig. 5 shows the observations from Reunion (green symbols, bottom panel). The mean column total observed at Reunion is $0.82 \times 10^{15}$ molecules NH$_3$ cm$^{-2}$. The concentrations are low during most of the year. However, peaks reaching densities of $\sim 6 \times 10^{15}$ molecules NH$_3$ cm$^{-2}$ can be observed during the end of each year. The peaks in September–November coincide with the dry season indicating that emissions are mostly due to biomass burning and large fire events (Vigouroux et al., 2012). This is supported by the increased CO concentrations, which are also observed in October and November (see, bottom left panel, Fig. 6). NH$_3$ surface concentration measurements are not available for this region but a recent paper by Van Damme et al. (2015b), which uses IASI-NH$_3$ observations, shows similar seasonal cycles for the south eastern parts of Africa (Madagascar). Temperature is almost constant throughout the year and not a major factor in the seasonality of Reunion.

Observations from Jungfraujoch have the lowest mean concentration of all four stations (Fig. 5, orange line), with a mean of $0.18 \times 10^{15}$ molecules NH$_3$ cm$^{-2}$. The low concentrations at Jungfraujoch are expected, as the station is located in the free tro-
posphere high above the surrounding valleys. Transport of NH$_3$ from the valleys only occurs sporadically during days with intense vertical mixing. This was also observed in an earlier study of CO concentrations (Barret et al., 2003). The Jungfraujoch observations show almost no seasonal effects with only a minimal increase during the summer months. The low concentrations measured at Jungfraujoch support our assumption on the vertical distribution of the ammonia concentrations with low values in the troposphere that were used in our a-priori.

4 Conclusions and perspectives

In this study we presented a new method to retrieve ammonia total columns from ground-based FTIR solar spectra. Observations from four complementary stations were used to illustrate the capabilities of the retrieval method. NH$_3$ total columns ranging three orders of magnitude were obtained with high abundances at Bremen (mean of $13.47 \times 10^{15}$ molecules cm$^{-2}$) to low columns at Jungfraujoch (mean of $0.18 \times 10^{15}$ molecules cm$^{-2}$). The very low levels obtained at the Jungfraujoch demonstrate the sensitivity of the retrieval method we developed. A separate error calculation shows random errors in the order of 10% and systematic errors of 25% for individual observations. The errors are dominated by uncertainties in spectroscopy, atmospheric temperature and deviations in instrumental parameters. For conditions with high surface concentrations of ammonia, as in Bremen, it is possible to retrieve information on the vertical gradient as two layers can be discriminated. At Bremen, the retrieval there is most sensitive to ammonia in the planetary boundary layer, where most of the ammonia is expected. For conditions with lower concentrations only the total columns can be retrieved. Station specific seasonal cycles were found to be consistent with known seasonal cycles of the dominant ammonia sources in the station surroundings. For example, highest levels in Bremen were observed during spring time when manure is applied to the fields with column total concentrations reaching up to $93 \times 10^{15}$ molecules cm$^{-2}$.
Remote sensing techniques avoid sampling artefacts common to other techniques such as filter packs (Puchalski et al., 2011; von Bobrutzki et al., 2010). For in-situ observations open path remote sensing techniques, e.g. DOAS and QCL instruments, are starting to be used (Volten et al., 2010; Miller et al., 2014). The FTIR-NH₃ observations would be an excellent addition to these approaches as it provides the NH₃ total column and profiles, including vertical information for sites sampling high ammonia levels. With a mean error of ~25% for all observations in high ammonia source areas the accuracy of the FTIR retrievals is comparable to that reported for satellite products (TES, IASI, CrIS). Compared to the in-situ open path remote sensing methods the FTIR method has a higher uncertainty, but this is a trade-off for the ability to retrieve vertical information. To improve the accuracy of the FTIR-NH₃ retrieval a reassessment of the spectral line parameters is necessary.

Observations from existing networks commonly represent daily or even monthly averaged concentration values, which severely complicates any attempt to validate satellite observations. The novel FTIR-NH₃ observations enable a direct validation of satellite products. As the FTIR-NH₃ product provides averaging kernels a direct comparison can be made with optimal estimation satellite retrievals while taking account of the a-priori information and vertical sensitivity of both instruments (Rogers and Connor, 2003). A dedicated field campaign was executed at the Cabauw Experimental Site for Atmospheric Remote Sensing (CESAR) in the Netherlands (spring and summer 2014) to validate the IASI-NH₃ using a range of instruments including mini-DOAS instruments and a Bruker IFS-NH₃ using a Bruker IFS-66 instrument (Dammers et al. in prep).

The uncertainty in the emission distributions hampers the performance and prediction capabilities of air quality and climate models (Heald et al., 2012). Emissions are usually based on nationally reported yearly emission inventories (Pouliot et al., 2012) and gridded by distributing the emissions following animal numbers and agricultural land use (Bouwman et al., 2002; Keunen et al., 2011). To improve on static emission time profiles, a new direction is to include the impact of meteorological variability of ammonia emissions in modelling systems (Sutton et al., 2013). Recently, such
an improvement was shown to greatly enhance the performance of air quality models (Skjøth et al., 2011). Similar to satellite observations, FTIR total columns in combination with surface observations could provide the means to evaluate the emission modelling through comparing trends and concentration anomalies within and between years. For this purpose continuous time series are necessary. Due to the lack of continuous data (i.e. more than one observation per hour) we could not derive a typical diurnal cycle in this study, whereas this would be highly useful for model evaluation. Improved knowledge on the diurnal cycles may also greatly help to interpret model evaluation results against satellite data as they provide snapshots, e.g. daily IASI’s observations at 9.30. Also, the model-measurement comparison would be less sensitive to modelling errors in the turbulent vertical exchange as the ammonia is integrated over vertical.

The developed retrieval methodology from FTIR-instruments provides a new way to obtain vertically and temporally resolved measurements on ammonia concentrations. FTIR-NH$_3$ observations may prove very valuable for satellite and model validation and may provide a complementary source of information to constrain the global ammonia budget.

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many colleagues who have contributed to FTIR data acquisition at the various sites.

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Table 1. FTIR stations used in the analysis. The location, longitude, latitude and altitude are given for each station as well as the instrument used for the measurements. Some station specifics are given in the last column.

<table>
<thead>
<tr>
<th>Station</th>
<th>Location</th>
<th>Longitude</th>
<th>Latitude</th>
<th>Altitude (m a.s.l.)</th>
<th>Instrument</th>
<th>Station specifics</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bremen</td>
<td>Germany</td>
<td>8.85° E</td>
<td>53.10° N</td>
<td>27</td>
<td>Bruker 125 HR</td>
<td>City, fertilizers, livestock</td>
</tr>
<tr>
<td>Lauder</td>
<td>New Zealand</td>
<td>169.68° E</td>
<td>45.04° S</td>
<td>370</td>
<td>Bruker 120 HR</td>
<td>Fertilizers, livestock</td>
</tr>
<tr>
<td>Reunion</td>
<td>Indian Ocean</td>
<td>55.5° E</td>
<td>20.90° S</td>
<td>85</td>
<td>Bruker 120M</td>
<td>Fertilizers, fires</td>
</tr>
<tr>
<td>Jungfraujoch</td>
<td>Switzerland</td>
<td>7.98° E</td>
<td>46.55° N</td>
<td>3580</td>
<td>Bruker 120 HR</td>
<td>High altitude, no large sources</td>
</tr>
</tbody>
</table>
Table 2. Micro windows used in the NH$_3$ retrieval at the four stations.

<table>
<thead>
<tr>
<th>Stations</th>
<th>Micro window</th>
<th>Spectral range (cm$^{-1}$)</th>
<th>Interfering species (Profile retrieved species in bold)</th>
<th>SNR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bremen and Lauder</td>
<td>MW1</td>
<td>930.32–931.32</td>
<td>NH$_3$, H$_2$O, O$_3$, CO$_2$, N$_2$O, HNO$_3$, SF$_6$, CFC-12, solar lines</td>
<td>Bremen – Real SNR mean value of 450</td>
</tr>
<tr>
<td></td>
<td>MW2</td>
<td>966.97–967.68</td>
<td>NH$_3$, H$_2$O, O$_3$, CO$_2$, N$_2$O, HNO$_3$, solar lines</td>
<td>Lauder – Real SNR mean value of 250</td>
</tr>
<tr>
<td>Reunion</td>
<td>MW1</td>
<td>929.4–931.4</td>
<td>NH$_3$, H$_2$O, O$_3$, CO$_2$, N$_2$O, HNO$_3$, SF$_6$, CFC-12</td>
<td>Reunion – Real SNR mean value of 365</td>
</tr>
<tr>
<td></td>
<td>MW2</td>
<td>962.1–970.0</td>
<td>NH$_3$, H$_2$O, O$_3$, CO$_2$, N$_2$O, HNO$_3$, HDO, $^{68}$O$_3$, solar lines</td>
<td></td>
</tr>
<tr>
<td>Jungfraujoch</td>
<td>MW1</td>
<td>929.4–931.4</td>
<td>NH$_3$, H$_2$O, O$_3$, CO$_2$, N$_2$O, HNO$_3$, SF$_6$, CFC-12</td>
<td></td>
</tr>
<tr>
<td></td>
<td>MW2</td>
<td>962.1–970.0</td>
<td>NH$_3$, H$_2$O, O$_3$, CO$_2$, N$_2$O, HDO, $^{68}$O$_3$, solar lines</td>
<td>Jungfraujoch – Fixed at 250</td>
</tr>
</tbody>
</table>
Table 3. Random and Systematic uncertainties used in the error calculation.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>SFIT 4 (Bremen, Lauder, Reunion)</th>
<th>SFIT 2 (Jungfraujoch)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Random uncertainty</td>
<td>Systematic uncertainty</td>
</tr>
<tr>
<td>Temperature</td>
<td>2 K troposphere</td>
<td>2 K troposphere</td>
</tr>
<tr>
<td></td>
<td>5 K stratosphere</td>
<td>5 K stratosphere</td>
</tr>
<tr>
<td>Solar line shift</td>
<td>0.005 cm(^{-1})</td>
<td>0.005 cm(^{-1})</td>
</tr>
<tr>
<td>Solar line strength</td>
<td>0.1 %</td>
<td>0.1 %</td>
</tr>
<tr>
<td>Solar zenith angle</td>
<td>0.01 °</td>
<td>0.01 °</td>
</tr>
<tr>
<td>Phase</td>
<td>0.001 Rad</td>
<td>0.001 Rad</td>
</tr>
<tr>
<td>Zero level</td>
<td>0.01</td>
<td>0.01</td>
</tr>
<tr>
<td>Background curvature</td>
<td>0.001 cm(^{-2})</td>
<td>Influence a priori profiles</td>
</tr>
<tr>
<td>Field of view</td>
<td>0.001</td>
<td></td>
</tr>
<tr>
<td>Line intensity</td>
<td>20.0 %</td>
<td></td>
</tr>
<tr>
<td>Line T broadening</td>
<td>10.0 %</td>
<td></td>
</tr>
<tr>
<td>Line P broadening</td>
<td>10.0 %</td>
<td></td>
</tr>
<tr>
<td>Interfering species</td>
<td>HITRAN2012: varies</td>
<td></td>
</tr>
</tbody>
</table>
Table 4. Mean random and systematic errors for each of the individual NH$_3$ retrieval parameters. The table is split into two sections to cover both the error calculation using SFIT4 (Bremen, Lauder, Reunion) and SFIT2 (Jungfraujoch). At the bottom the errors are summarized into total mean errors for each of the stations.

<table>
<thead>
<tr>
<th>Station Parameter</th>
<th>Bremen Mean Random Error (%)</th>
<th>Bremen Mean Systematic Error (%)</th>
<th>Lauder Mean Random Error (%)</th>
<th>Lauder Mean Systematic Error (%)</th>
<th>Reunion Mean Random Error (%)</th>
<th>Reunion Mean Systematic Error (%)</th>
<th>Jungfraujoch Mean Random Error (%)</th>
<th>Jungfraujoch Mean Systematic Error (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temperature</td>
<td>4.9</td>
<td>4.9</td>
<td>3.6</td>
<td>3.6</td>
<td>2.7</td>
<td>2.9</td>
<td>15.2</td>
<td>1.9</td>
</tr>
<tr>
<td>Solar zenith angle</td>
<td>1.6</td>
<td>1.6</td>
<td>1.1</td>
<td>1.1</td>
<td></td>
<td></td>
<td>1.9</td>
<td>1.4</td>
</tr>
<tr>
<td>Phase</td>
<td>1.0</td>
<td>1.0</td>
<td>1.1</td>
<td>1.1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Zero level</td>
<td>5.0</td>
<td>5.0</td>
<td>6.8</td>
<td>6.8</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Measurement noise</td>
<td>4.5</td>
<td>4.5</td>
<td>8.4</td>
<td>8.4</td>
<td>10.9</td>
<td></td>
<td>18.2</td>
<td></td>
</tr>
<tr>
<td>Interfering Species</td>
<td>1.3</td>
<td>2.4</td>
<td>0.9</td>
<td>8.7 (H$_2$O line pressure broadening)</td>
<td></td>
<td>1.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Retrieval parameters</td>
<td>0.1</td>
<td>0.1</td>
<td></td>
<td></td>
<td></td>
<td>1.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Background curvature</td>
<td></td>
<td>1.1</td>
<td>1.2</td>
<td>0.3</td>
<td></td>
<td>1.0</td>
<td>1.4</td>
<td></td>
</tr>
<tr>
<td>Smoothing error</td>
<td>2.8</td>
<td>8.1</td>
<td>10.3</td>
<td>17.8</td>
<td></td>
<td>Model parameters</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Spectroscopy</td>
<td>21.0</td>
<td>22.7</td>
<td>30.2</td>
<td>27.0</td>
<td>15.3</td>
<td>20.0</td>
<td>5.4</td>
<td>20.1</td>
</tr>
<tr>
<td>Influence a priori</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>6.6</td>
<td>6.1</td>
</tr>
<tr>
<td>Subtotal error</td>
<td>9.1</td>
<td>25.3</td>
<td>12.0</td>
<td>30.2</td>
<td>15.3</td>
<td>20.0</td>
<td>25.3</td>
<td>23.1</td>
</tr>
<tr>
<td>Total error</td>
<td>23.5</td>
<td>30.2</td>
<td>15.3</td>
<td>20.0</td>
<td>25.2</td>
<td>34.2</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Table 5. Statistics of the NH$_3$ columns. (Nr: number of data points, DOFS: Degree of Freedom for Signal, Mean ± the error of the mean, RMS: Root Mean Square). Total columns are given in $1 \times 10^{15}$ molecules NH$_3$ cm$^{-2}$.

<table>
<thead>
<tr>
<th>Station</th>
<th>Nr</th>
<th>Mean DOFS</th>
<th>Mean (molecules × $1 \times 10^{15}$)</th>
<th>Median (molecules × $1 \times 10^{15}$)</th>
<th>RMS (molecules × $1 \times 10^{15}$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Bremen</td>
<td>554</td>
<td>1.9</td>
<td>13.75 ± 4.24</td>
<td>9.51</td>
<td>20.22</td>
</tr>
<tr>
<td>Lauder</td>
<td>2412</td>
<td>1.4</td>
<td>4.17 ± 1.40</td>
<td>2.85</td>
<td>5.95</td>
</tr>
<tr>
<td>Reunion</td>
<td>1262</td>
<td>1.0</td>
<td>0.80 ± 0.54</td>
<td>0.56</td>
<td>1.14</td>
</tr>
<tr>
<td>Jungfraujoch</td>
<td>2702</td>
<td>1.0</td>
<td>0.18 ± 0.07</td>
<td>0.15</td>
<td>0.22</td>
</tr>
</tbody>
</table>
Figure 1. Calculated spectrum for both spectral windows measured with the 125HR in Bremen on the 19 April 2010 at 09:59 (UTC) corresponding to a total column of $18.83 \times 10^{15}$ molecules cm$^{-2}$. The top two panels show the individual contributions of the different species in the first (MW1) and second (MW2) spectral windows. The second row show the same calculated spectra but now with the $y$ axis scaled to show the minor interfering species.
**Figure 2.** Example of a synthetic atmospheric spectrum for both spectral windows at Reunion Island, computed for the 5 June 2011 and a total column of $1.07 \times 10^{15}$ molecules cm$^{-2}$. The top panel shows the individual contributions of the main species in the first spectral window. The bottom panel shows the second spectral window.
Figure 3. Measured and calculated spectrum for both spectral windows measured with the 125HR in Bremen on the 19 April 2010 at 09:59 (UTC) corresponding to a total column of $18.83 \times 10^{15}$ molecules NH$_3$ cm$^{-2}$. The top two panels show the observed (blue line) and calculated (green line) spectra for MW1 (left) and MW2 (right). The bottom two figures show the residuals of the fits in both spectral windows.
Figure 4. Top panels: the retrieved NH₃ profile (blue) and the a-priori profile (green) in order from left to right: Bremen (Left), Lauder (Left middle), Reunion island (right middle) and Jungfraujoch (right). Horizontal lines indicate the standard deviation in all observations for each layer. Bottom panels: the normalized averaging kernel for each of the stations.
Figure 5. Time series of retrieved NH$_3$ columns (in molecules NH$_3$ cm$^{-2}$). From top to bottom the figure shows the Bremen (blue), Lauder (red), Reunion (green) and Jungfraujoch (yellow) total columns. The bars reflect the errors on the individual observations.
Figure 6. 2004–2013 monthly averaged columns for NH$_3$, CO and temperature. The top two panels show the monthly NH$_3$ column concentrations (molecules NH$_3$ cm$^{-2}$) for each of the four stations. Vertical lines indicate the mean monthly error. The bottom two panels show additional column concentrations of CO (bottom, left) and temperature (bottom, right).