

Data assimilation of CrIS-NH₃ satellite observations for improving spatiotemporal NH₃ distributions in LOTOS-EUROS.

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Abstract. Atmospheric levels of ammonia (NH₃) have substantially increased during the last century, posing a hazard to both human health and environmental quality. The atmospheric budget of NH₃, however, is still highly uncertain due to an overall lack of observations. Satellite observations of atmospheric NH₃ may help us in the current observational and knowledge gaps. Recent observations of the Cross-track Infrared Sounder (CrIS) provide us with daily, global distributions of NH₃. In this study, the CrIS-NH₃ product is assimilated into the LOTOS-EUROS chemistry transport model using two different methods aimed at improving the modelled spatio-temporal NH₃ distributions. In the first method NH₃ surface concentrations from CrIS are used to fit spatially varying NH₃ emission time factors to redistribute model input NH₃ emissions over the year. The second method uses the CrIS-NH₃ column data profile to adjust the NH₃ emissions using a Local Ensemble Transform Kalman Filter (LETKF) in a top-down approach. The two methods are tested separately and combined, focusing on a region in western Europe (Germany, Belgium, and the Netherlands). In this region, the mean CrIS-NH₃ total columns were up to a factor 2 higher than the simulated NH₃ columns between 2014 and 2018, which, after assimilating the CrIS-NH₃ columns using the LETKF algorithm, led to an increase of the total NH₃ emissions of up to approximately 30%. Our results illustrate that CrIS-NH₃ observations can be used successfully to estimate spatially variable NH₃ time factors, and improve NH₃ emission distributions temporally, especially in spring (March to May). Moreover, the use of the CrIS-based NH₃ time factors resulted in an improved comparison with the onset and duration of the NH₃ spring peak observed at observation sites at hourly resolution in the Netherlands. Assimilation of the CrIS-NH₃ columns with the LETKF algorithm is mainly advantageous for improving the spatial concentration distribution of the modelled NH₃ fields. Compared to in-situ observations, a combination of both methods led to the most significant improvements in modelled monthly NH₃ surface concentration and NH₄⁺ wet deposition fields, illustrating the usefulness of the CrIS-NH₃ products to improve the temporal representativity of the model and better constrain the budget in agricultural areas.

1. Introduction

Ammonia (NH_3) is an alkaline gas in the Earth's atmosphere. NH_3 is highly reactive and readily reacts with available acids, forming aerosol components harmful to human health (Pope et al., 2009, Lelieveld et al., 2015, Giannakis et al., 2019) and, directly and indirectly, impacting global climate change (Erisman et al., 2011, Myhre et al., 2013). NH_3 is emitted from a large number of sources, including agriculture, natural nitrogen fixation in oceans and plants, volcanic eruptions, and biomass-, industrial- and fossil fuel burning (Erisman et al., 2015). Globally, agriculture is the largest source of NH_3 . Agricultural emissions of NH_3 consist of, among others, volatilized NH_3 after manure and chemical fertilizer application, livestock housing and grazing and harvesting of crops. About 40% of the total global NH_3 emissions follow directly from volatilization of animal manure and chemical fertilizer, a spatially variable process highly controlled by the temperature and acidity of soils (Sutton et al., 2013). In western Europe, for instance, agriculture is an even more dominant source of NH_3 and contributes to 85-100% of all NH_3 emissions (Hertel et al., 2011). After the emitted NH_3 is transported through the atmosphere, it is deposited back to the Earth's surface through the processes of wet and dry deposition. Excess amounts of reactive nitrogen deposition can cause several adverse effects, such as eutrophication in aquatic ecosystems and soil acidification (Erisman et al., 2007) and biodiversity loss in terrestrial ecosystems (Bobbink et al., 2010).

Even though NH_3 at its current levels is an important threat to human health and environmental quality, its atmospheric budget is still very uncertain. NH_3 concentrations are highly variable in space and time and are difficult to be reliably measured in-situ due to the sticky nature of NH_3 leading to potential adsorption to parts of the measurement devices (von Bobruzki et al., 2010). Globally, only a few NH_3 measurement networks exist, most of which contain only a small number of locations. Moreover, most measurements are performed at a coarse temporal resolution (weeks to months), while most atmospheric processes occur on much shorter time scales. Due to the lack of dense and precise measurement networks, measures for NH_3 emission controls currently rely mostly on estimates from models, for instance from chemical transport models (CTMs). CTMs simulate atmospheric processes such as emissions, transport, deposition and chemical conversion to estimate the spatial and temporal distribution of atmospheric NH_3 . However, these models involve large uncertainties. On the one hand, model assumptions and parameterizations are uncertain due to insufficient or lack of knowledge of some of the processes, for instance, the limited understanding of bi-directional fluxes of NH_3 (Schrader and Brümmner, 2014, Schrader et al., 2018) or the direct effect of meteorology on NH_3 emissions (Sutton et al., 2013). On the other hand, uncertainties stem from the underlying input data and the spatial and temporal variation in emissions. Compared to other air pollutants, NH_3 emission inputs are especially uncertain due to their large spatiotemporal variability resulting from the diverse nature of agricultural sources (Behera et al., 2013). In Europe, the uncertainty of the total annual reported NH_3 emissions on a country level is for instance already estimated to be at least around ~30% (EEA, 2019). Naturally, NH_3 emissions from individual sources have a much higher uncertainty due to errors related to spatial and temporal **re**distribution. So as to reduce the uncertainty in modelled NH_3 fields from CTMs, it is vital to better understand both the spatiotemporal distribution of NH_3 emissions.

65 With the scarceness of in-situ measurements and uncertainties in existing models, the atmospheric NH₃ budget remains among the least known parts of the nitrogen cycle (Erismann et al., 2007). Recent satellite observations of NH₃ in the lower troposphere can help us to fill in both observational and knowledge gaps. Satellite instruments, such as the NASA Tropospheric Emission Spectrometer (TES) (Beer et al., 2008), ESA's Infrared Atmospheric Sounding Interferometers (IASI) (Clarisse et al., 2009), the NASA Atmospheric Infrared Sounder (AIRS) (Warner et al., 2016), the Thermal And Near-infrared Spectrometer for
70 Observation-Fourier Transform Spectrometer (TANSO-FTS) (Someya et al., 2020) and the NASA/NOAA Cross-track Infrared Sounder (CrIS) (Shephard and Cady-Pereira, 2015) provide global observations of atmospheric NH₃. Out of the operational satellites that observe NH₃ with twice daily global coverage, CrIS is the newest instrument and has the lowest radiometric noise in the spectral region used for NH₃ (Zavalyov et al., 2013). Moreover, CrIS has **increased** **greater** vertical sensitivity **for** to near-surface NH₃, and provides retrievals of the vertical distribution of NH₃ (Shephard et al., 2020).

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Measurements of These atmospheric trace gases **measurements** with satellites have opened up new ways to study the atmospheric budget. Recently, satellite observations have successfully been used for direct estimates of emissions and lifetimes of various other atmospheric species (e.g., SO₂, NO₂, CO₂) of single anthropogenic or natural point sources (e.g., Fioletov et al., 2015, Nassar et al., 2017) or even multiple sources at a time (Fioletov et al., 2017, Beirle et al., 2019). For NH₃ specifically,
80 multiple studies have reported emissions and atmospheric lifetime estimates either based on satellite data (e.g., Zhu et al., 2013, Whitburn et al., 2015, Van Damme et al., 2018, Zhang et al., 2018, Cao et al., 2020, Evangelio et al., 2021) or directly estimated from satellite data (e.g., Van Damme et al., 2018, Adams et al., 2019, Dammers et al., 2019). Here, also different forms of model inversions have been used. Overall, these studies indicate an underestimation of both anthropogenic and natural NH₃ emissions in the current emission inventories. In addition to estimating NH₃ emissions, various studies used satellite
85 observations to estimate dry deposition fluxes of NH₃ (Kharol et al., 2018, Van der Graaf et al., 2018, Lui et al., 2020).

In this manuscript, we describe two methods to improve both the temporal and spatial variation of NH₃ emissions in the LOTOS-EUROS chemistry transport model with CrIS-NH₃ observations. The first method aims at deriving an improved set of a-priori, observation-based NH₃ time factors to be used for the **temporal** **daily** distribution of agricultural emission sources within LOTOS-EUROS. In this method, the temporal variation of NH₃ surface concentrations from CrIS is used. The second
90 method is used to assimilate the CrIS-NH₃ observations into the LOTOS-EUROS model. **For this, a** **using a** Local Ensemble Transform Kalman Filter (LETKF) approach **is used** as data-assimilation system. **which strength lies in enhancing existing** **spatial patterns**. The impact of the two methods, both individually and combined, on the simulated NH₃ emissions, concentration and deposition fields is then evaluated. The focus region of the study is a low-to-high NH₃ emission area within
95 western Europe (The Netherlands, Germany, Belgium), which is representative for other intense agricultural regions in the world. Moreover, the NH₃ emissions within this region are relatively well known and in-situ observations are sufficiently available.

2. Methodology

2.1. LOTOS EUROS

100 LOTOS-EUROS is an Eulerian chemistry transport model (Manders et al., 2017) that could be used to simulate trace gas and
aerosol concentrations in the lower troposphere. The model has an intermediate complexity with limited run time, allowing
ensemble-based simulations and assimilation studies. LOTOS-EUROS uses meteorological data as input, which in this study
is taken from the using European Centre for Medium-Range Weather Forecasts (ECMWF). The gas-phase chemistry follows
a carbon-bond mechanism (Schaap et al., 2008). The dry deposition fluxes are calculated with the Deposition of Acidifying
105 Compounds (DEPAC) 3.11 module, following the resistance approach and it includes a calculation of bi-directional NH_3 fluxes
(Van Zanten et al., 2010, Wichink Kruit et al., 2012). The wet deposition fluxes are computed using the CAMx (Comprehensive
Air quality Model with Xtensions) approach, which includes both in-cloud and below-cloud scavenging (Banzhaf et al., 2012).
The anthropogenic emissions are taken from CAMS-REG-AP (Copernicus Atmospheric Monitoring Services Regional Air
Pollutants) emissions dataset v2.2 (Granier et al., 2019). For Germany, high resolution gridded NH_3 emission inputs (GRETA)
110 are used (Schaap et al., 2018). In this study, a region in Western Europe (47°N-56°N, 2°E-16°E) is modelled, which includes
all of Germany, the Netherlands and Belgium (Fig. 2). A spatial resolution of 0.20° longitude by 0.10° latitude is used,
corresponding to approximately 12 by 12 square kilometers, which is also roughly the footprint size of CrIS (14 by 14 km² at
nadir). The vertical grid extends up to 200 hPa and is split up into 13 vertical layers. This captures the largest part of
atmospheric NH_3 , as it is a relatively short-lived species mainly located in the boundary layer. The interfaces of the vertical
115 layers are based on the pressure layers used in the ECMWF meteorological input data. LOTOS-EUROS is part of the
operational Copernicus Atmosphere Monitoring Service (CAMS) ensemble forecasts and analysis for Europe (Marécal et al.,
2015). The model has participated in multiple model intercomparison studies (e.g., Bessagnet et al., 2016, Colette et al., 2017,
Vivanco et al., 2018), showing overall good performance.

2.2. Datasets

120 2.2.1. CrIS NH_3

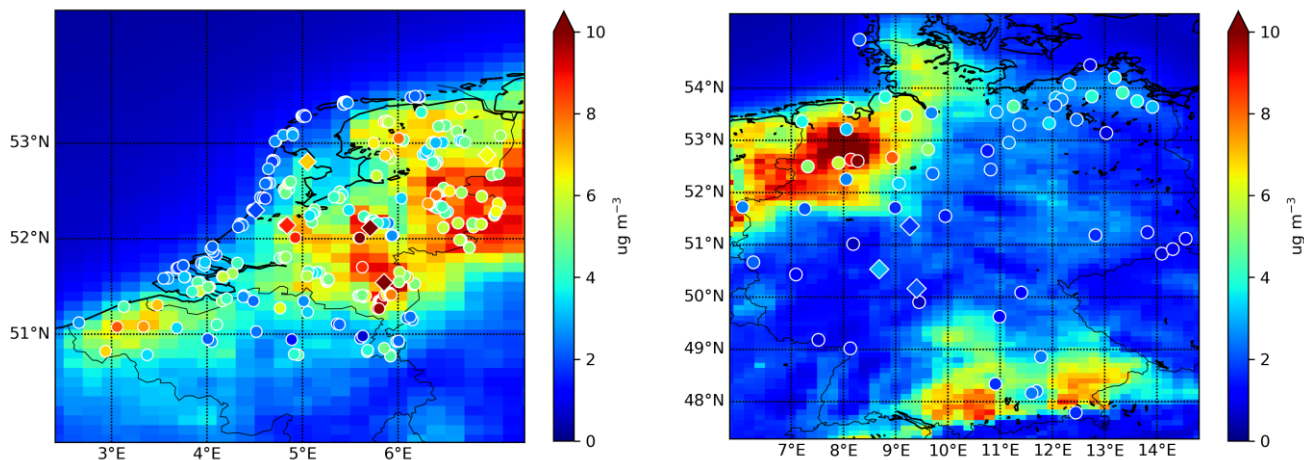
The Cross-Track Infrared Sounder (CrIS) is an instrument aboard NASA's sun-synchronous, Earth orbiting Suomi NPP
satellite with an equatorial overpass at 13:30 and 1:30 LST. The CrIS sensor has a spectral resolution of 0.625 cm⁻¹ (Shephard
et al., 2015) and a detection limit of 0.3-0.5 ppbv under favorable conditions (Shephard et al., 2020). The instrument has a
wide swath of up to 2200 km with pixels of approximately 14 km in size at nadir. Compared to other NH_3 satellite sounders
125 (e.g., AIRS, IASI), CrIS has **an improved greater vertical sensitivity of NH_3 close to the surface due to its low spectral noise
of approximately 0.04K at 280K in the NH_3 spectral region (Zavyalov et al., 2013).** Moreover, CrIS has a relatively high near-
surface sensitivity **and an overpass time around 1:30 LST, which coincides with the time of the day with the highest thermal
contrast.** The peak sensitivity of the instrument is typically between 900 and 700 hPa, which corresponds to approximately 1
to 3 km (Shephard et al., 2020). The CrIS NH_3 total columns have an estimated total random measurement error of around 10-

130 15%, and an estimated random total error of ~30%. Due to the limited vertical resolution, the NH₃ concentrations at individual retrieval levels have a higher random measurement error of about 10-30% and a total error of ~60-100% (Shephard et al., 2020). Version 1.3 of the CrIS-NH₃ product has been evaluated against in-situ Fourier Transform Infrared (FTIR) measurements (Dammers et al., 2017) showing an overall good performance and high correlations of $r \sim 0.8$. In this study, we used version 1.5 of the CrIS fast physical retrieval (FPR)-NH₃ product, which is based on the optimal estimation method
135 (Rodgers, 2000). More details about the CrIS FPR-NH₃ product can be found in (Shephard et al., 2020). Here, we used daytime observations of NH₃ (partial) column concentrations and surface concentrations made between January 2014 and December 2018 from the first CrIS sensor, which has the longest observing period. During this 5-year period, a virtually continuous timeseries of CrIS observations was available. More recent observations were not used due to the technical issues of the CrIS instrument during the summertime in 2019, and the potentially anomalous situation resulting from the COVID-19 outbreak in
140 2020.

2.2.2. In-situ observations

Several measurement networks were used to evaluate the simulated concentration and deposition fields. The NH₃ surface concentrations are evaluated against observations from the Dutch Meetnet Ammoniak in Natuurgebieden (MAN) network (Lolkema et al., 2015), the Dutch Landelijk Meetnet Luchtkwaliteit (LML) network (van Zanten et al., 2017), the Belgium
145 Flanders Environment Agency (VMM) network (den Bril et al., 2011) and the German Environment Agency (UBA) network (Schaap et al., 2018). The locations of these sites are shown in Fig. 1. The MAN network provides monthly mean NH₃ surface concentrations since 2005, spread over 80 mostly low NH₃ emission nature areas in the Netherlands. The measurements are performed with low-cost passive samplers from Gradko and have an estimated uncertainty of ~20% for high concentrations and ~41% for low concentrations (Lolkema et al., 2015). The NH₃ concentrations in Flanders are measured with passive
150 samplers from Radiello and IVL samplers (den Bril et al., 2011). The LML network observes hourly NH₃ concentrations at six different locations in the Netherlands with different emission regimes (high, moderate, low). Initially, continuous flow denuders from AMOR were used, which have a reported uncertainty of at least 9% for hourly concentrations (Blank et al., 2001). Around 2016, the AMOR instruments were replaced by miniDOAS instruments (Berkhout et al., 2017), which are active differential optical absorption spectrometers. For evaluation of the wet deposition fields, observations from wet-only
155 samplers from the Dutch Landelijk Meetnet Regenwatersamenstelling (LMRe) network (van Zanten et al., 2017), whose locations largely overlap with the LML network, and the UBA network (Schaap et al., 2018) are used. The locations of the wet-only samplers are shown in Fig. S1.

Mean NH₃ surface concentration (2014-2017)



160 **Figure 1: Locations of stations that measure NH₃ surface concentrations. The circles depict passive samplers and the diamonds hourly observations stations.**

2.3. Fitting method for deriving CrIS-based NH₃ time factors

A non-linear least squares method is used to fit a trimodal gaussian curve to the scaled NH₃ surface concentrations (see section 2.3.3) from CrIS in each grid cell. The Trust Region Reflective algorithm is used to perform the minimization (Conn et al., 2000). The minimalization algorithm is restrained with initial parameter guesses and bounds for three fitted gaussians. The three gaussians represent the spring (μ_1 , σ_1 , and A_1), autumn (μ_2 , σ_2 , and A_2) and summer peak (μ_3 , σ_3 , and A_3) in NH₃ emissions, respectively. The initial parameter guesses are based on the standard MACC-III (Kuenen et al., 2014) NH₃ emission time profiles. The bounds are defined as follows:

- 170 • the mean values ($\mu_{1,2,3}$) are permitted to shift by one month (30 days) to cover the most probable emission peaks
- the standard deviations ($\sigma_{1,2,3}$) are permitted to vary by half their initial value guess (i.e., $\pm 0.5\sigma$)
- the fitted amplitude of the spring peak (A_1) is allowed to be between 0.1 and 1.0 and amplitudes of the autumn and summer gaussians ($A_{2,3}$) between 0.1 and 0.8

175 An overview of the permitted parameter bounds is given in Table 1. The range in permitted $A_{1,2,3}$ values is quite large, allowing the minimization algorithm to fit meaningful trimodal curves for different types of time variant NH₃ emission sources simultaneously (e.g., flatter peaks for emissions that are mainly dependent of temperature and specific periods, such as open stables, a sharper more distinct spring and autumn peaks for emissions following fertilizer or manure application).

	Spring peak			Autumn peak			Summer peak		
	μ_1 (doty)	σ_1 (days)	A_1 (-)	μ_2 (doty)	σ_2 (days)	A_2 (-)	μ_3 (doty)	σ_3 (days)	A_3 (-)
Lower bound	47.4	13.1	0.1	222.8	11.6	0.1	148.9	26.9	0.1
First guess (MACC-III)	77.4	26.1	0.96	252.8	23.2	0.26	178.9	53.7	0.21
Upper bound	107.4	39.1	1.0	282.8	34.8	0.8	208.9	107.4	0.8

180 Table 1: Initial parameter guesses and parameter bounds used in the trimodal fit algorithm.

After the daily NH₃ time factors are fitted, the diurnal variation from the MACC-III NH₃ time factors is added to obtain hourly time factors. The resulting hourly CrIS-based time factors are used as input for all time-variant NH₃ sources from agriculture subcategories in LOTOS-EUROS, i.e., continuous NH₃ point sources emissions remain time-invariant.

185 2.3.2. Data selection

The CrIS NH₃ concentrations in the lowest retrieval level, i.e., closest to the surface, are used to adjust the daily variability in the hourly time profiles spatially on a regular 0.1° by 0.05° grid. First, ~~to collect enough observations,~~ to collect a sufficient number of observations for the fitting algorithm, the CrIS NH₃ surface concentrations with a quality flag of at least 3 and within a selection radius of 1° around the center points of each grid cell are selected. The daily average NH₃ concentrations throughout the year are computed after application of a simple outlier filter (>99th percentile excluded given more than 3 observations). Due to the lower number of observations during winter, and to avoid a bias towards higher values due to lower thermal contrast, observations in January, November and December are ignored. During these months it is anyway prohibited to apply fertilizer or spread manure in parts of the regions (for the Netherlands, see RVO, 2021), and in combination with the colder temperatures, NH₃ concentrations are expected to be low due to low volatilization rates (e.g., Søggaard et al., 2002).

195 2.3.3. Correction for local emission to concentration ratio

The relationship between NH₃ emissions and surface concentrations differs per by region and changes throughout the year due to changes in the meteorological and chemical circumstances conditions. To correct for this, the following adjustment factor is applied to the daily CrIS NH₃ surface concentrations. The factor is based on is derived from the NH₃ emission and simulated surface concentration fields from LOTOS-EUROS, which are used to compute the local ratio of the smoothed daily total NH₃ emissions to the NH₃ surface concentrations at the CrIS overpass time per grid cell. These are used as a first order approximation for the relation between the emission and concentration. The ratios are rescaled by the mean annual values for each grid cell to obtain a unitless daily scaling factor (Fig. S2). After multiplying the daily averaged CrIS NH₃ surface concentrations with this scaling factor, a $\pm 1\sigma$ filter is used to smoothen out the daily time series. To avoid too much flattening of the spring emission peak, a separate filter is applied for the spring period. NH₃ emissions originating from the application

205 of synthetic or manure fertilizers are mainly found during this period, at the beginning of the growing season. This may lead to an increase in observed NH₃ concentrations, that would be filtered out when the same filter is applied for the entire year. Finally, the scaled NH₃ surface concentrations are normalized for each grid cell.

2.4. Data assimilation system

2.4.1. Local Ensemble Transform Kalman Filter

210 The Ensemble Kalman Filter (Evensen, 2003) is a sequential data assimilation method that could be used to combine model simulations with observation. In this study, the Local Ensemble Transform Kalman Filter (LETKF) formulation is used (Hunt et al, 2007) following the implementation by (Shin et al., 2016). The LETKF performs an analysis per grid cell based on nearby observations only and it therefore computationally advantageous compared to the regular implementation of the Ensemble Transform Kalman Filter. The basic idea behind an Ensemble Kalman Filter is to express the probability function of the state
 215 in terms of an ensemble with N possible states x_1, x_2, \dots, x_N , each considered to be a possible sample out of the distribution of the true state. In this study, the state contains the NH₃ concentrations in a three-dimensional grid and two-dimensional NH₃ emission perturbation factors β . The perturbation factors describe the uncertainty in the emissions, and are modelled as samples out of normal distribution with zero mean and standard deviation σ . Spatial variations are initially not defined, but are introduced by a localization length scale that is described below. The temporal variation in the emission factors is described
 220 by temporal correlation coefficient α , that depends on temporal length scale τ (Lopez-Restrepo et al., 2020, Barbu et al., 2009):

$$\alpha_k = e^{-|t_k - t_{k-1}|/\tau} \quad (\text{Eq. 1})$$

An initial ensemble is created by generating random samples of the perturbation factors. The ensemble is then propagated in time in what is called the *forecast* step between consecutive *analysis* times for which observations are available. In the *forecast*
 225 step, the model propagates the *analysed* ensemble members from time t_{k-1} to time t_k following:

$$\mathbf{x}_i(k) = \mathbf{M}_{k-1}(\mathbf{x}_i^a(k-1)) \quad (\text{Eq. 2})$$

where operator \mathbf{M}_{k-1} describes the model simulation, including application of the perturbation factors that are present in \mathbf{x} . The ensemble mean \bar{x} and forecast error covariance \mathbf{P} at time k are expressed as:

$$\bar{x} = \frac{1}{N} \sum_{i=1}^N x_i \quad (\text{Eq. 3})$$

$$\mathbf{P} = \frac{1}{N-1} \sum_{i=1}^N (x_i - \bar{x})(x_i - \bar{x})^T \quad (\text{Eq. 4})$$

When CrIS observations (\mathbf{y}^{CrIS}) are available (at time t_k), the LETKF algorithm *analyses* the ensemble by incorporating the observations to reduce the ensemble spread. The *analysed* ensemble members are computed from:

$$\mathbf{x}_i^a = \mathbf{x}_i + \mathbf{P}^a \mathbf{H}^T \mathbf{R}^{-1} (\mathbf{y}^{\text{CrIS}} - \mathbf{h}(x_i) + \mathbf{v}_i) \quad (\text{Eq. 5})$$

In here, $\mathbf{h}(x_i)$ represents the simulated retrieval from the state \mathbf{x}_i , or in particular from the concentration array in \mathbf{x}_i . Operator \mathbf{H} is a linearization of $\mathbf{h}(x)$ to x (see section 2.4.4). The matrix \mathbf{R} is the *observation representation error covariance*, which describes the difference between the simulation and the observation due to measurement and representation errors:

$$240 \quad \mathbf{y}^{\text{CrIS}} - \mathbf{h}(x_i) \sim \mathcal{N}(0, \mathbf{R}) \quad (\text{Eq. 6})$$

The actual implementation of \mathbf{h} , \mathbf{H} , and \mathbf{R} are described below. The *analysis covariance* \mathbf{P}^a is computed from:

$$\mathbf{P}^a = [\mathbf{P}\mathbf{H}^T\mathbf{R}^{-1}\mathbf{H} + \mathbf{I}]^{-1} \mathbf{P} \quad (\text{Eq. 7})$$

2.4.2. Observation simulation

245 The simulated observation vector $\mathbf{h}(x_i)$ represents the simulated retrieval, which is what the satellite observes from concentrations described in 3-dimensional grid cell x_i , and is computed from:

$$\mathbf{h}(x_i) = \mathbf{y}_a - \mathbf{A}\mathbf{y}_a + \mathbf{A}\mathbf{G}\mathbf{x}_i \quad (\text{Eq. 8})$$

Here, matrix \mathbf{G} , the *gridding operator*, is applied to horizontally and vertically match the simulated partial NH_3 columns in LOTOS-EUROS with the retrieval CrIS pressure levels. Here, *air-mass weighted averaging is used to average the model layers to the retrieval levels*. The relationship between the true and the retrieved atmospheric NH_3 profiles, i.e., the vertical sensitivity of the CrIS measurements, is described by averaging kernel \mathbf{A} . The full relationship between the true and the observed state is given by $\mathbf{y}^{\text{true}} = \mathbf{h}(x^{\text{true}}) + \mathbf{v}$, which can be rewritten to (Eq. 9) (Rodgers and Connor, 2003):

$$250 \quad \mathbf{y}^{\text{true}} = \mathbf{y}_a + \mathbf{A}(\mathbf{G}\mathbf{x}^{\text{true}} - \mathbf{y}_a) + \mathbf{v} \quad (\text{Eq. 9})$$

255

with \mathbf{y}_a the a-priori profile that is part of the CrIS retrieval product. The error \mathbf{v} is a sample of the *observation representation error taken from a normal distribution* that describes the possible differences between simulation and retrieval:

$$\mathbf{v} \sim \mathcal{N}(0, \mathbf{R}) \quad (\text{Eq. 10})$$

260 In this study, \mathbf{R} is set to the retrieval error covariance that is part of the CrIS product. The linearized observation operator becomes:

$$\mathbf{H} = \mathbf{A}\mathbf{G} \quad (\text{Eq. 11})$$

2.4.3. Analysis per grid cell

265 The analysis described above is applied per model grid cell; for the exact implementation we refer to Shin et al. (2016). First, the simulated observation vectors $\mathbf{h}(x_i)$ are computed for all ensemble members. For the grid cell to be analyzed, all simulations are collected that are within 3.5ρ distance, where ρ is called the localization length scale as well as the matching

actual observations \mathbf{y}^{CrIS} . The state elements corresponding to the grid cell are then analyzed using the collected observations and simulations, where the weight of observations further away is limited using Gaussian correlation that decays with distance and that uses the same correlation length scale ρ that is used for collection.

270 **2.4.4. Observation selection**

CrIS observations with the highest quality flag, QF = 5, were used. **These observations have a relatively higher impact because of their low uncertainty.** As the assumed vertical NH_3 profile shape in background areas used in the CrIS retrieval and in LOTOS-EUROS differ, CrIS retrievals with “unpolluted” a-priori profiles were filtered out. The original CrIS retrieval is performed in the log domain and therefore either the averaging kernels A from CrIS need to be linearized or the LOTOS-
275 EUROS profiles transformed to the log-domain. Linearization of the kernel is only accurate for higher concentrations, and since this is the case for the selected “polluted” retrievals, this option was found to be suitable.

2.4.5. Parameter calibration

In this study, we used a localization radius of $\rho = 15$ km, a standard deviation of $\sigma = 0.5$ and a temporal correlation length of $\tau = 3$ days. Two experiments were performed to study the effect of ρ , σ and τ in more detail. A description of the experiments
280 can be found in section S1 of the supplementary materials. A limited ensemble size of $N=12$ was found to be sufficient to describe the imposed model uncertainty, which is not too complicated due to short life-time of NH_3 and therefore strong relation between concentrations and nearby emissions.

3. Results

3.1. Direct comparison of NH_3 concentrations from CrIS and LOTOS-EUROS

285 Before looking at the effects of assimilating the CrIS observations, a direct comparison of the modelled and observed NH_3 column densities was made. The simulated NH_3 concentrations from the default run in LOTOS-EUROS were sampled at the locations of the CrIS observations, and after application of the averaging kernels compared with the retrievals. The observed and simulated NH_3 total columns averaged over all years are shown in Fig. 2. Similar maps per year are available in Fig. S3 of the supplementary materials. The general pattern of the NH_3 total column densities matches quite well. For instance, the
290 observed and simulated NH_3 columns are very similar in southwestern Germany, and close to the Dutch border. The CrIS NH_3 total columns are generally higher than the simulated NH_3 total columns. This is for instance found in large parts of northeastern Germany, along the Belgium coast and in the south of the Netherlands. Here, the observed NH_3 columns were on average approximately a factor 2 higher than the simulated NH_3 columns. Moreover, the observed NH_3 total columns are consistently higher than the simulated NH_3 columns in background areas, with a bias between the observed and modelled concentrations
295 of approximately $\sim 0.5 \times 10^{16}$ molecules/cm².

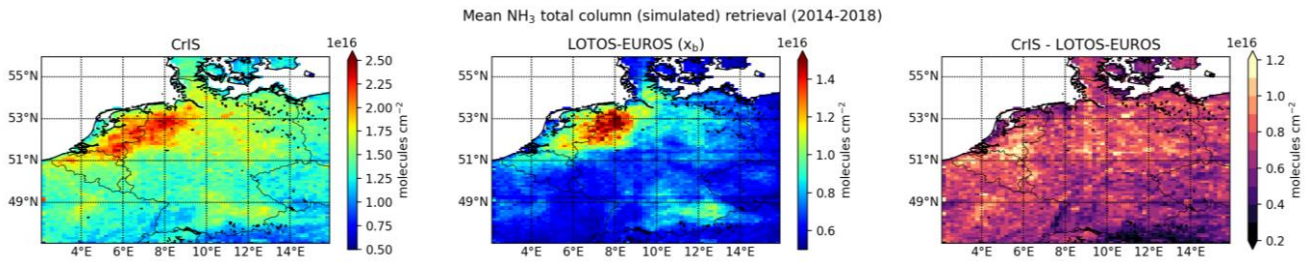


Figure 2: Mean retrieved (left) and simulated (middle) NH_3 total column from 2014-2018, and their absolute difference (right).

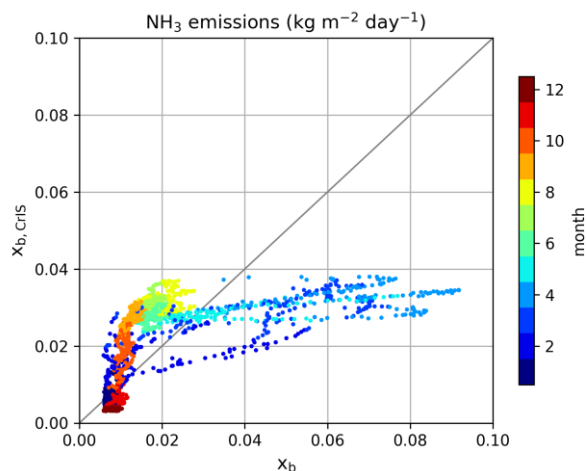
3.2. CrIS-based NH_3 time factors

3.2.1. Effect on NH_3 emissions in LOTOS-EUROS

300 Following the method described in section 2.3, temporal profiles for the NH_3 have been obtained per grid cell. Compared to the original model, the new time profiles vary spatially. Fig. 3 shows a comparison of the daily grid-averaged NH_3 emissions between the default background model run (x_b) and the background run with the CrIS-based NH_3 time factors ($x_{b,\text{CrIS}}$), using a different color for each month. The default NH_3 time factors from MACC-III provide more intra-annual variation than the CrIS-based NH_3 time factors. The default time factors include a very high peak in spring and much lower peaks during summer and autumn (i.e., $A_1/A_3 = 4.57$, $A_1/A_2 = 3.70$). Fig. S4 shows the fitted spring parameters (μ_1 , σ_1 and A_1). The NH_3 spring peak present in the CrIS- NH_3 surface concentrations is generally lower than the default NH_3 spring peak. In large parts of the model region, the CrIS-observed NH_3 spring peak is subsequently lower and less sharp. Compared to the default NH_3 time factors, the amplitude of the spring peak in the CrIS-based NH_3 time factors is now generally much lower. The amplitude of the spring peak differs almost by a factor 2 on average. As a result, there is a decrease in springtime NH_3 emissions, especially in March and April. The CrIS-based NH_3 time factors, and consequently the NH_3 emissions, are, on the other hand, generally higher later in the year. The NH_3 emissions are on average approximately 50% higher in summer and the beginning of autumn (June to September), and approximately twice as high in October.

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315 **Figure 3: Daily grid-average NH₃ emission, colored per month. Here, x_b represents the default background run and $x_{b,CrIS}$ the background run with CrIS-based NH₃ time factors.**

3.2.2. Effect on NH₃ concentrations and deposition fields in LOTOS-EUROS

The changes in modelled NH₃ surface concentration, total column concentrations and NH_x total deposition from 2014 to 2018 related to the use of the CrIS-based NH₃ time factors **alone** are shown in Fig. 4, Fig. S5 in the supplementary materials and Fig. 5. Here, x_b represents the default background run and $x_{b,CrIS}$ the background run with the CrIS-based NH₃ time factors.

320 The use of the CrIS-based emission time profiles led to an overall increase in mean NH₃ surface concentrations. The absolute change is largest in areas with already relatively high NH₃ surface concentrations, for instance in northwestern Germany, where the mean NH₃ surface concentrations increased with up to 2 µg/m³. The mean NH₃ surface concentrations increased with up to ~25% due to the change in NH₃ time factors. The effect of the CrIS-based NH₃ time factors on the NH₃ total column concentrations is smaller, with minor changes from minus ~5% up to 5%. The mean NH₃ total column concentrations generally

325 increase in areas with already high NH₃ concentrations, such as large parts of the Netherlands, and decrease in background areas with little NH₃ emissions, for instance in central Germany. The use of the CrIS-based NH₃ time factors led to ~10% less total NH_x deposition along the northwestern coast, including agricultural hotspots such as the Netherlands and northwestern Germany, and an increase of up to ~10% in background areas.

Fig. S6 compares the daily, grid averaged, NH₃ surface concentrations, total column concentrations and NH_x wet and dry

330 deposition, with different colors per month. Here, a similar redistribution is observed for the NH₃ concentration and deposition fields as seen earlier for the NH₃ emission fields. Compared to the default background run (x_b), the NH₃ concentration fields were up to a factor 2 lower during March and April. The NH₃ total columns decreased in spring, the largest decrease occurring in April (up to ~60%). The NH₃ surface concentrations increased during the summer and the beginning of autumn, up to ~50% in September. During these months, a similar but slightly lower increase in the NH₃ total column concentrations is observed.

335 Because the CrIS-based NH_3 time factors vary per year, the interannual variation in the modelled NH_3 fields is much larger. Fig. S7 shows the relative changes in NH_3 surface concentration, total column concentration and NH_x deposition fields per year. Overall, the mean NH_3 surface concentration increases by up to $\sim 30\%$ per year. The largest increases occurred in 2016 and 2018, years with relatively high summer temperatures (Copernicus Climate Change Service, 2021). The variation in the annual mean NH_3 total column concentrations is much smaller (-15 to $+15\%$). The relative change in NH_x budget shows much more variation, with the most prominent increase occurring in 2014 ($+25\%$) and decreases occurring in 2018 (-25%).

The temporal redistribution of the NH_3 emissions thus significantly impacts the modelled NH_3 concentration and deposition fields, too. Generally, a part of the initial spring NH_3 emissions is now attributed to the summer and autumn months. Depending on the degree of redistribution, there are distinct changes in the NH_x budget. Looking at the fitted spring peak parameters (Fig. S4) and the matching CrIS-based NH_3 factors at hourly measurement sites (Fig. S8), clear interannual differences are observed.

345 For instance, a relatively sharp spring peak was observed over the Netherlands in 2014. In 2018, on the other hand, the fitted spring peak had a distinctly lower amplitude and started later in the year. Moreover, a relatively large rise in NH_3 time factors was observed again in late summer and autumn (July to September). Compared to 2014, this resulted in a relatively larger redistribution of the NH_3 emissions towards warmer months. The higher temperatures resulted in lower dry deposition velocities and more vertical mixing and transport of NH_3 , leading to an overall decrease in NH_x deposition over the

350 Netherlands. Moreover, the summer of 2018 was relatively dry, also leading to higher NH_3 total column concentrations and a decrease in wet NH_x deposition.

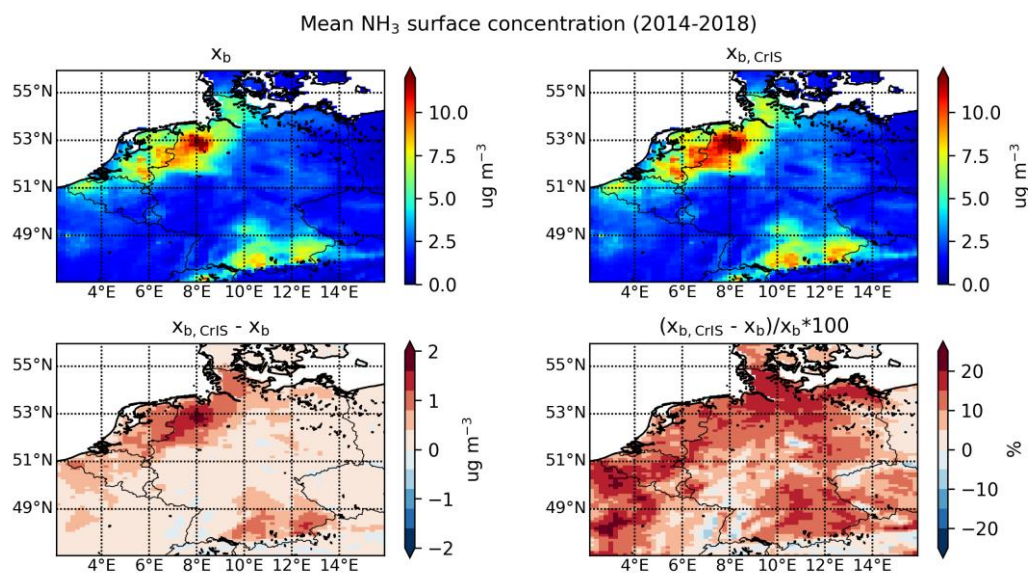
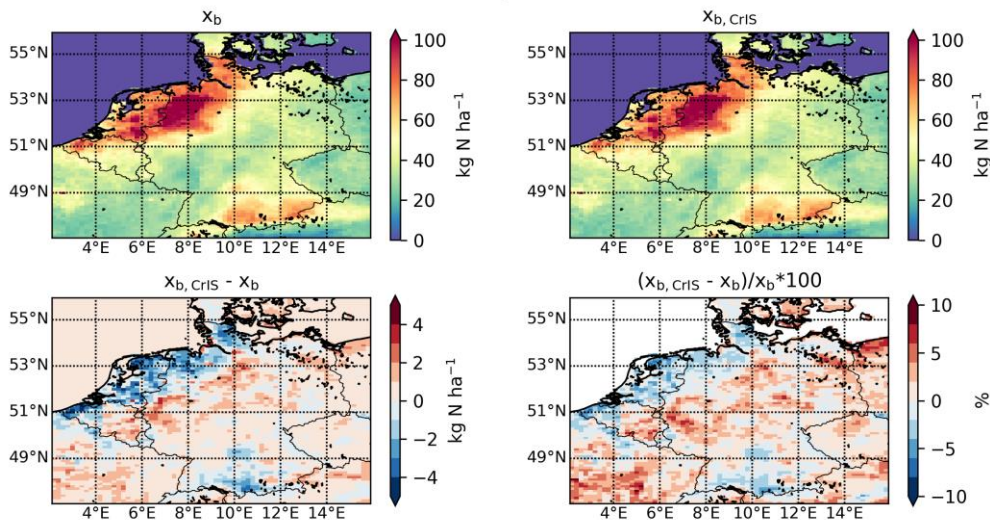


Figure 4: The mean NH_3 surface concentration over 2014 to 2018 from the (top left) default background run (x_b) and the (top right) background run with CrIS-based NH_3 time factors ($x_{b, \text{CrIS}}$) and their (bottom left) absolute and (bottom right) relative difference.

Total NH_x budget (2014-2018)



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Figure 5: The total NH_x deposition from 2014 to 2018 from the (top left) default background run (x_b) and the (top right) background run with CrIS-based NH₃ time factors ($x_{b,CrIS}$) and their (bottom left) absolute and (bottom right) relative difference.

3.3. Local Ensemble Transform Kalman Filter

3.3.1. Effect on NH₃ emissions and concentrations in LOTOS-EUROS

360 The CrIS-NH₃ columns were assimilated using the Local Ensemble Transform Kalman Filter (LETKF) described in section 2.4. Assimilations were performed using either the default emission time profiles (x_a), or using the CrIS-based profiles ($x_{a,CrIS}$). The total NH₃ emissions from 2014 to 2018 and the relative and absolute changes compared to background simulations x_b and $x_{b,CrIS}$ are shown in Fig. 6. The corresponding mean NH₃ surface and total column concentrations are shown in Fig. S9 and Fig. S10 of the supplementary materials. The absolute NH₃ emission updates by the LETKF are, as expected, largest in regions with already high NH₃ emissions. There is a maximum increase of ~30% in total NH₃ emission by the LETKF over the entire period for some grid cells. Relatively, the largest changes are found in the southern parts of the Netherlands (province of Noord-Brabant), the west coast of Belgium (province of West-Vlaanderen), the northeastern parts of Germany and France. Compared to the analysis run using default emission time profiles (x_a), the analysis runs with the CrIS-based NH₃ profiles ($x_{a,CrIS}$) generally have more NH₃ emission and consequently higher NH₃ surface and total column concentrations. The long-term spatial patterns of the emission updates by the LETKF, however, remained very similar.

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Total NH₃ emissions (2014-2018)

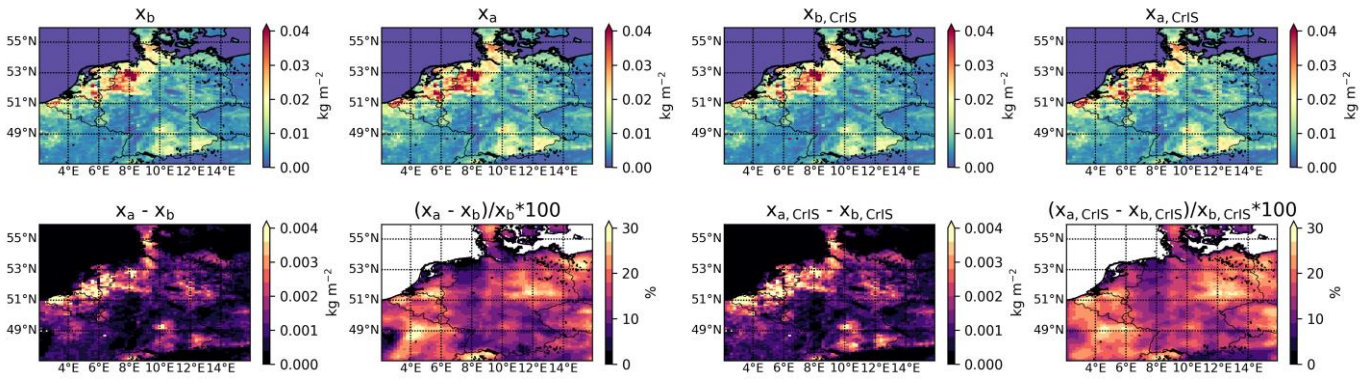
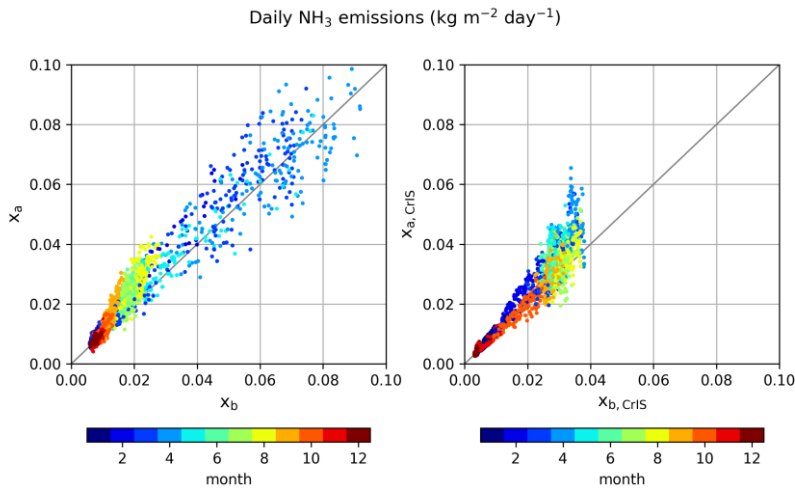


Figure 6: The total NH₃ emissions in 2014-2018 in the background runs x_b and $x_{b,CrIS}$ and in analysis runs x_a and $x_{a,CrIS}$ (top panels), as well as their absolute and relative difference (bottom panels).

375 To study the effect of the LETKF in more detail, the daily grid average NH₃ emissions of the background runs (x_b and $x_{b,CrIS}$) are plotted against analysis runs (x_a and $x_{a,CrIS}$) in Fig. 7. **Similar** Similarly, for the NH₃ surface and total column concentrations are plotted in Fig. S11 of the supplement. In the runs with the default NH₃ time factors (x_b and x_a), data assimilation of the CrIS-NH₃ columns led to both positive and negative emission updates in spring. In the summer, on the contrary, it mostly resulted in an increase in NH₃ emissions. In the runs with the CrIS-based NH₃ time factors ($x_{b,CrIS}$ and $x_{a,CrIS}$), the pattern is

380 distinctly different. Compared to the default runs, the NH₃ emission updates in spring are now smaller and largely positive, with the largest updates occurring in April. Moreover, the NH₃ emission updates were generally smaller during summer, too. This is related to the fact that the CrIS-NH₃ surface concentrations were used to fit the NH₃ time factors, which resulted in the model being closer to the CrIS observations already.



385 **Figure 7: Daily grid average NH₃ emissions in 2014-2018 from the (left) default background run (x_b) versus analysis run (x_a), and from the (right) background run with the CrIS-based NH₃ time factors ($x_{b,CrIS}$) versus analysis run $x_{a,CrIS}$, colored per month.**

Perturbation factor β is the computed multiplication factor by which the initial input NH_3 emissions are updated in the LETKF. The mean perturbation factors β per year are shown in Fig. S12 of the supplementary materials. The pattern of the NH_3 emission updates does not change drastically between years, which points to a consistent, spatial misdistribution of the emissions in the current inventory. By far the largest mean NH_3 emission updates took place in 2018, followed by 2015.

Fig. 8 shows timeseries of the daily grid average NH_3 emissions in both background runs x_b and $x_{b,\text{CrIS}}$ and analysis runs x_a and $x_{a,\text{CrIS}}$. Fig. 9 and S13 show the corresponding timeseries and changes in NH_3 surface and total column concentrations. The NH_3 emissions in the default background run (x_b) have a strong, annually reoccurring spring peak. After this peak, the NH_3 emissions decrease steeply and then slightly increase again in late summer and autumn (August and September). In analysis run x_a , the spring NH_3 emissions are both positively and negatively adjusted. Later in the year, almost only positive emission updates are found. The largest positive NH_3 emission updates occurred around August and September, which suggests an underestimation of the autumn NH_3 peak in the default runs.

In the background runs with the CrIS-based NH_3 time factors ($x_{b,\text{CrIS}}$), the NH_3 emissions are much more evenly distributed over the year. In contrast to the default runs, practically only positive NH_3 emission updates occurred in the analysis run ($x_{a,\text{CrIS}}$). The largest NH_3 updates took place during spring (March to May). The flattening of the NH_3 emissions led to a flattening in NH_3 concentration fields, too. Compared to default runs (x_b and x_a), there is much less interannual variation in the NH_3 surface and total column concentrations. As a result, the NH_3 concentrations during summer and autumn could be at the same level or even higher than the springtime concentrations. During the warm summer of 2018 (Copernicus Climate Change Service, 2021), for instance, the NH_3 concentrations in August even clearly exceed the spring NH_3 concentrations.

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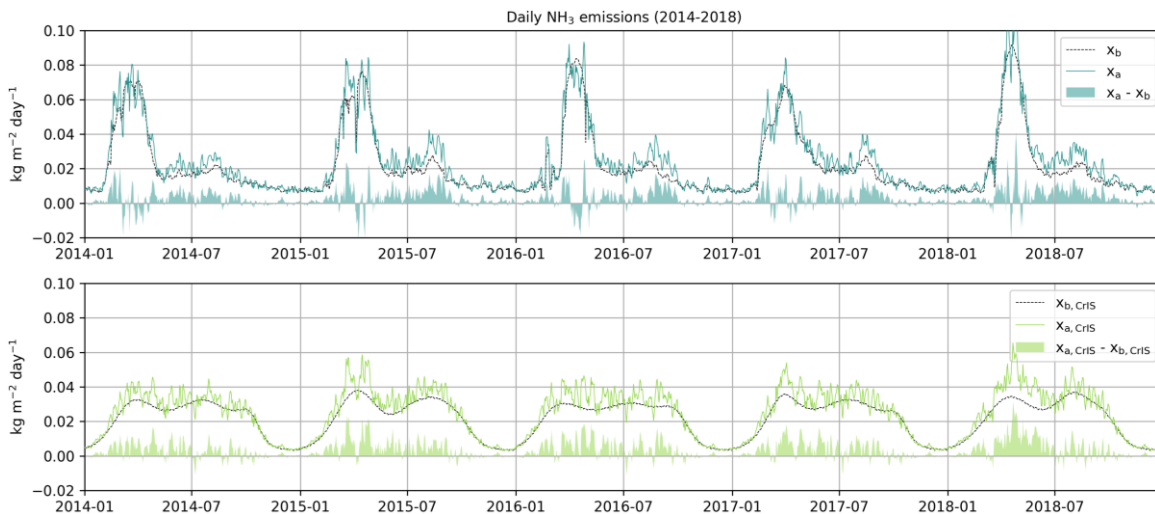


Figure 8: Timeseries of the daily grid-averaged NH_3 emissions in the background and analysis runs, and their absolute difference. The top figure (blue) represents the default background (x_b) and analysis run (x_a). The bottom figure (green) the background ($x_{b,\text{CrIS}}$) and analysis run ($x_{a,\text{CrIS}}$) with the CrIS-based NH_3 time factors.

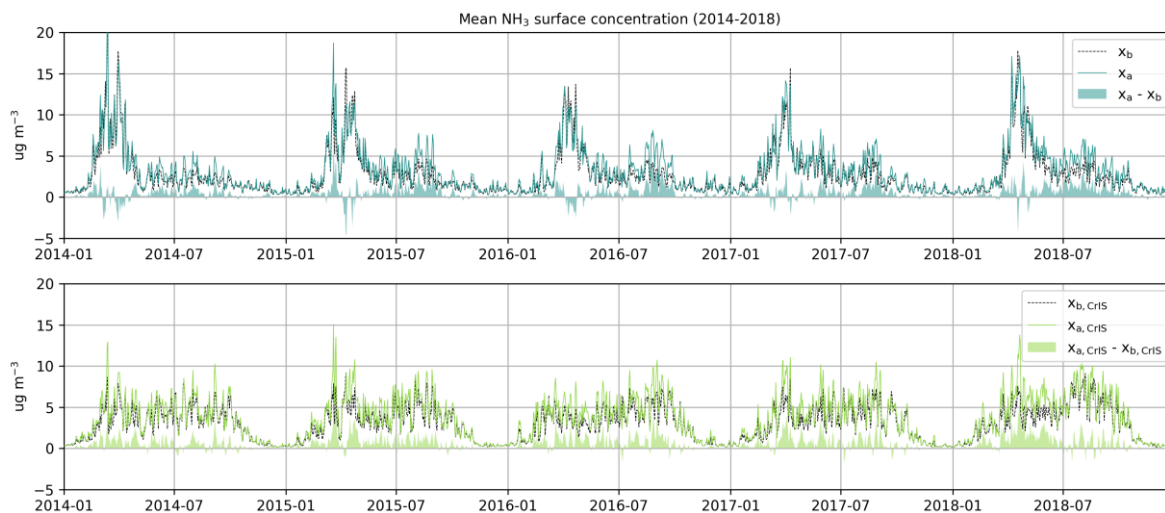


Figure 9: Timeseries of the daily grid-averaged NH_3 surface concentrations in the background and analysis runs, and their absolute difference. The top figure (blue) represents the default background (x_b) and analysis run (x_a). The bottom figure (green) the background ($x_{b,\text{CrIS}}$) and analysis run ($x_{a,\text{CrIS}}$) with the CrIS-based NH_3 time factors.

415 3.3.2. Effect on NH_x deposition in LOTOS-EUROS

The modelled total NH_x budgets from 2014 to 2018 from the two background runs (x_b and $x_{b,\text{CrIS}}$) and analysis runs (x_a and $x_{a,\text{CrIS}}$) are shown in Fig. 10. Overall, the modelled NH_x budget shows the same spatial pattern as the NH_3 emissions. Like the NH_3 emissions, the relatively largest spatial differences between the background and analysis runs took place in the south of the Netherlands, the west of Belgium and northeast Germany. Compared to the default runs, the relative changes in total NH_x

420 budget were slightly larger in the runs with the CrIS-based NH_3 time factors ($x_{b,\text{CrIS}}$ and $x_{a,\text{CrIS}}$).

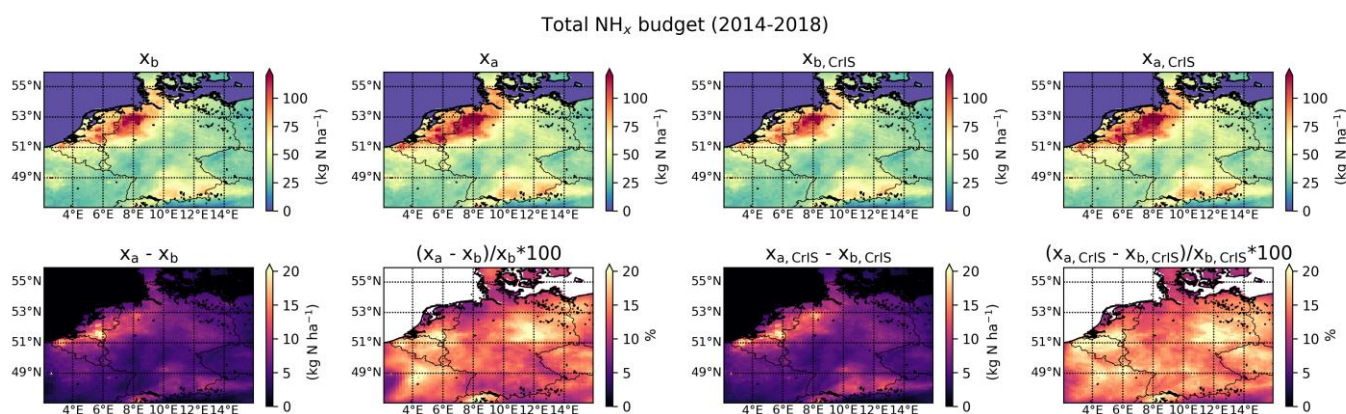
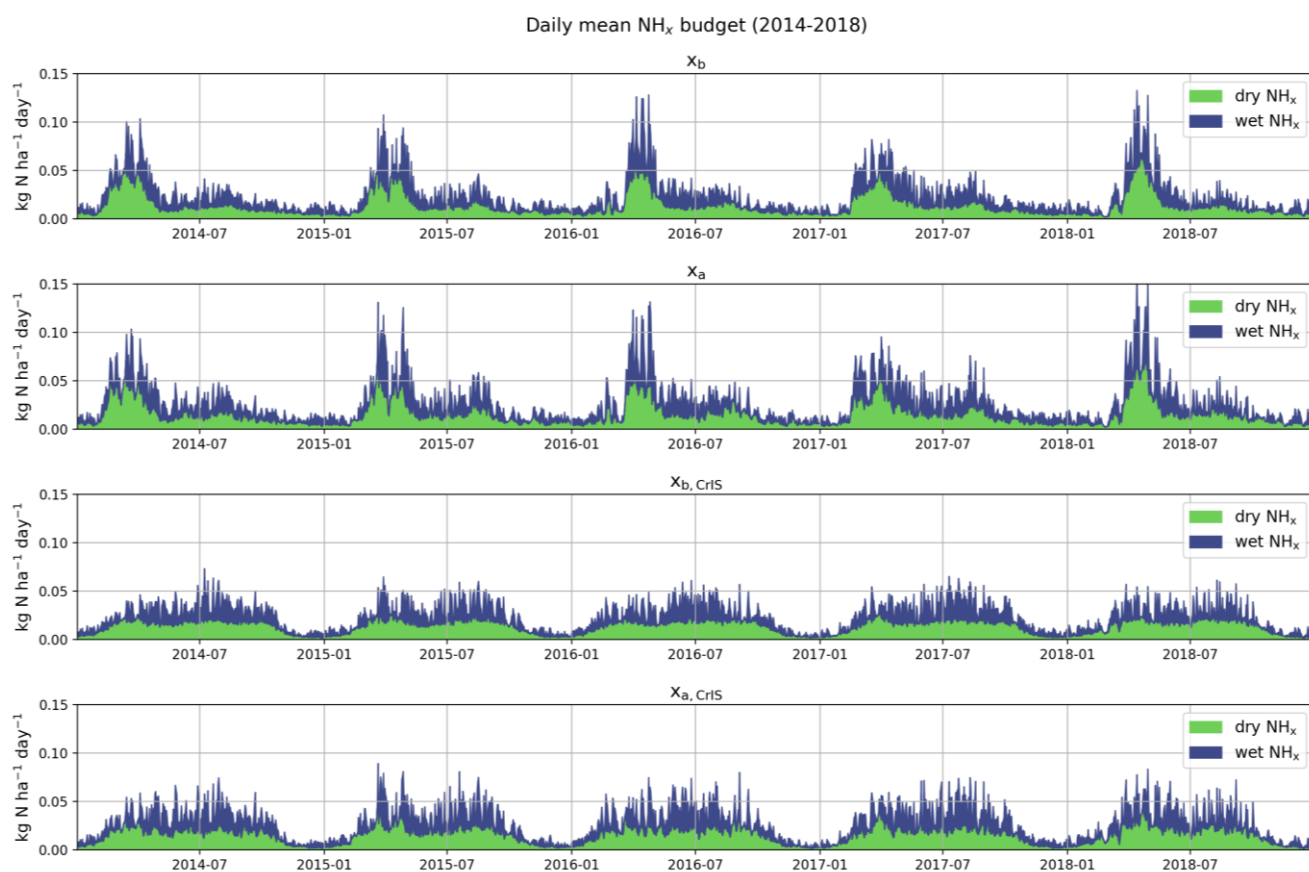


Figure 10: The total NH_x budget from 2014-2018 in the background (x_b and $x_{b,\text{CrIS}}$) and analysis (x_a and $x_{a,\text{CrIS}}$) model runs in LOTOS-EUROS, and their absolute and relative difference.

The modelled NH_x deposition follows the temporal distribution of the NH_3 emissions, too. Timeseries of the daily wet and dry deposition amounts in the domain are shown in Fig. 11. The wet and dry deposition in the default runs (x_b and $x_{b,\text{CrIS}}$) versus the analysis runs (x_a and $x_{a,\text{CrIS}}$) per month is shown in Fig. S14 in the supplement. In the default background run (x_b), the total NH_x deposition peaks in March and April. In the analysis run (x_a), the dry and wet deposition both increased and decreased during spring (March to May). Later in the year, the wet and dry NH_x deposition mostly increased in the analysis run, particularly in August and September. In the background runs with the CrIS-based NH_3 time factors ($x_{b,\text{CrIS}}$ and $x_{a,\text{CrIS}}$), the modelled dry and wet deposition fields are much less variable. Following the NH_3 emission updates, both the dry and wet deposition mostly increased in the analysis run, especially in March and April. Moreover, the use of the CrIS-based NH_3 time factors resulted in a redistribution of the ratio of wet and dry deposition over the year. As a result of the relatively lower spring NH_3 surface concentrations, there is a reduction of the dry deposition during spring. The relatively higher summer NH_3 total column concentrations led to a shift in wet deposition, too, from spring to summer.



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Figure 11: Timeseries of the average amounts of dry (green) and wet (blue) NH_x deposition in the different model runs. The top two figures represent the default background (x_b) and analysis (x_a) run and the bottom two figures the background ($x_{b,\text{CrIS}}$) and analysis ($x_{a,\text{CrIS}}$) run with the CrIS-based NH_3 time factors.

3.4. Comparison to in-situ observations

440 The modelled NH_3 surface concentration and NH_4^+ wet deposition fields are compared with in-situ observations. First, the spatial distribution is evaluated by comparing the modelled NH_3 surface concentration and NH_4^+ wet deposition fields to the observed annual averages per measurement site. Second, the temporal distribution is evaluated by comparing the modelled NH_3 surface concentration and NH_4^+ wet deposition fields to the same set of observations, but on a monthly basis. The comparisons are done per type of observation, e.g., all available wet-only measurements simultaneously. To differentiate
445 between different NH_3 emission regimes, the results are plotted separately for either all hourly observations or for the passive samplers. The results are shown in Fig. 12 and 13. The Dutch site with the highest NH_3 surface concentrations, Vredepeel, is excluded from this comparison because of the large model-observation discrepancies here (see Fig. S18). This site is located near agricultural emission sources and therefore less representative of a larger region. In these figures, the first column shows the comparison for the default background run (x_b), the second column shows the background run with CrIS-based NH_3 time factors ($x_{b,\text{CrIS}}$), the third column shows the analysis run with the default NH_3 time factors (x_a) and, finally, the fourth column shows the analysis run with CrIS-based NH_3 time factors ($x_{a,\text{CrIS}}$).

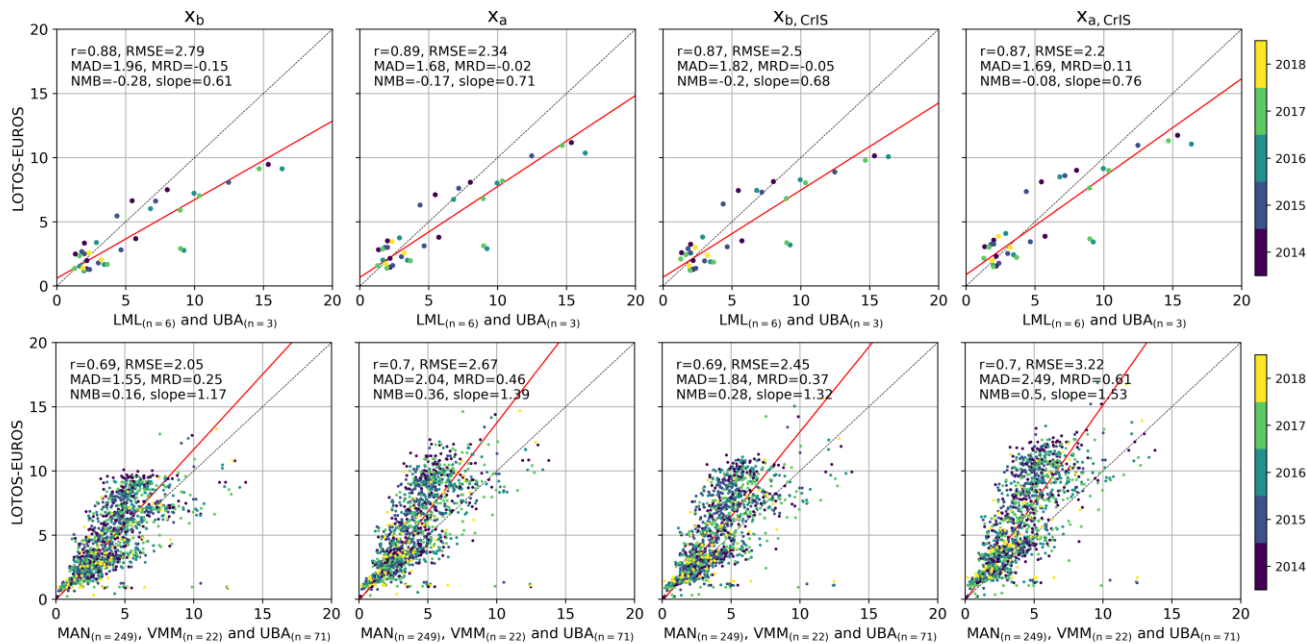
3.4.1. Spatial distribution

Fig. 12 shows the scatterplots of the annual averages per site per year. The annual average NH_3 surface concentrations (top row) in the default run x_b show a strong correlation ($r = 0.88$) with the observed concentrations at the hourly observation sites
455 (LML and UBA). Here, the NH_3 surface concentrations are generally underestimated (slope = 0.61). The annual average NH_3 surface concentrations (middle row) at the passive sampler sites (MAN, VVM and UBA) are generally overestimated (slope = 1.17), with a lower, but still relatively strong correlation is observed ($r = 0.69$). The modelled annual average NH_4^+ wet deposition budgets (bottom row) are moderately correlated with the observations from wet-only samplers ($r = 0.45$), and are generally lower than the observed wet deposition (slope = 0.81). When using the CrIS-based NH_3 time factors, the annual
460 average NH_3 surface concentrations and NH_4^+ wet deposition budgets are slightly increased. This led to a slight, overall increase in slope between all observations and the background run with the CrIS-based NH_3 time factors ($x_{b,\text{CrIS}}$). As the annual totals, and herewith the spatial distribution of the NH_3 emissions, remained the same in this run, the other measures (r , RMSE, MAD, MRD, NMB) didn't change drastically on a yearly basis.

465 The comparison with annual average NH_3 surface concentrations from the passive sampler networks from both analysis runs (x_a and $x_{a,\text{CrIS}}$) slightly worsened compared to the background runs (x_b and $x_{b,\text{CrIS}}$). The comparison at the hourly observation and wet-only sampler sites, on the other hand, showed clear improvements. Here, virtually all statistical measures improved, illustrating an overall improvement in modelled NH_3 surface concentration and NH_4^+ wet deposition field spatially. Of all runs, the analysis run with the CrIS-based NH_3 time factors ($x_{a,\text{CrIS}}$) compared the best with the hourly observation and wet-only
470 sampler network. The differences between the modelled and observed NH_3 surface concentrations at the hourly observation

were distinctly smaller, compared to the default background run (x_b : {RMSE = 2.79, MAD = 1.96, MRD = -0.15, NMB = -0.28} versus $x_{a,CrIS}$: {RMSE = 2.2, MAD = 1.69, MRD = -0.11, NMB = -0.08}). Here, also the slope largely improved (x_b : slope = 0.61 versus $x_{a,CrIS}$: slope = 0.76). The same is observed for the modelled NH_4 wet deposition fields, where the slope improved particularly (x_b : {RMSE = 0.95, MAD = 0.63, MRD = -0.13, NMB = -0.22, slope = 0.81} versus $x_{a,CrIS}$: {RMSE = 0.92, MAD = 0.61, MRD = -0.02, NMB = -0.11, slope = 0.95}).

NH_3 surface concentration ($\mu g m^{-3}$)



NH_4 wet deposition ($mg m^{-2}$)

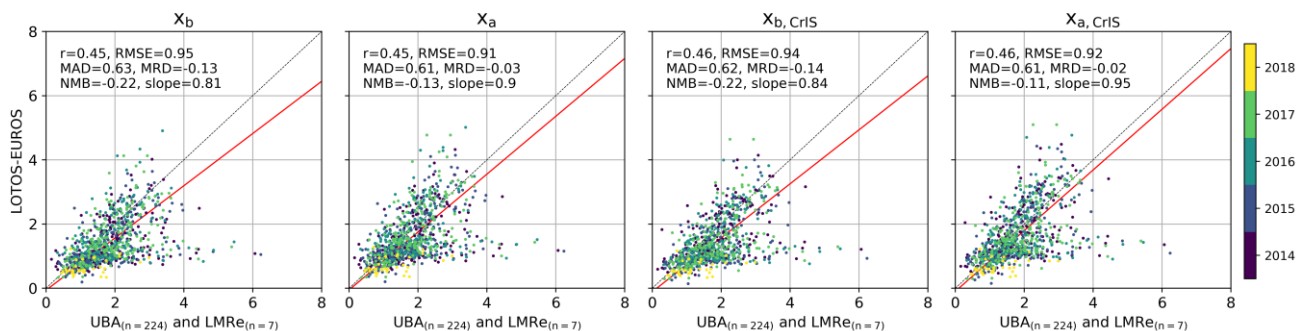


Figure 12: Comparison of the modelled annual average NH_3 surface concentrations and NH_4^+ wet deposition fields to in-situ observations.

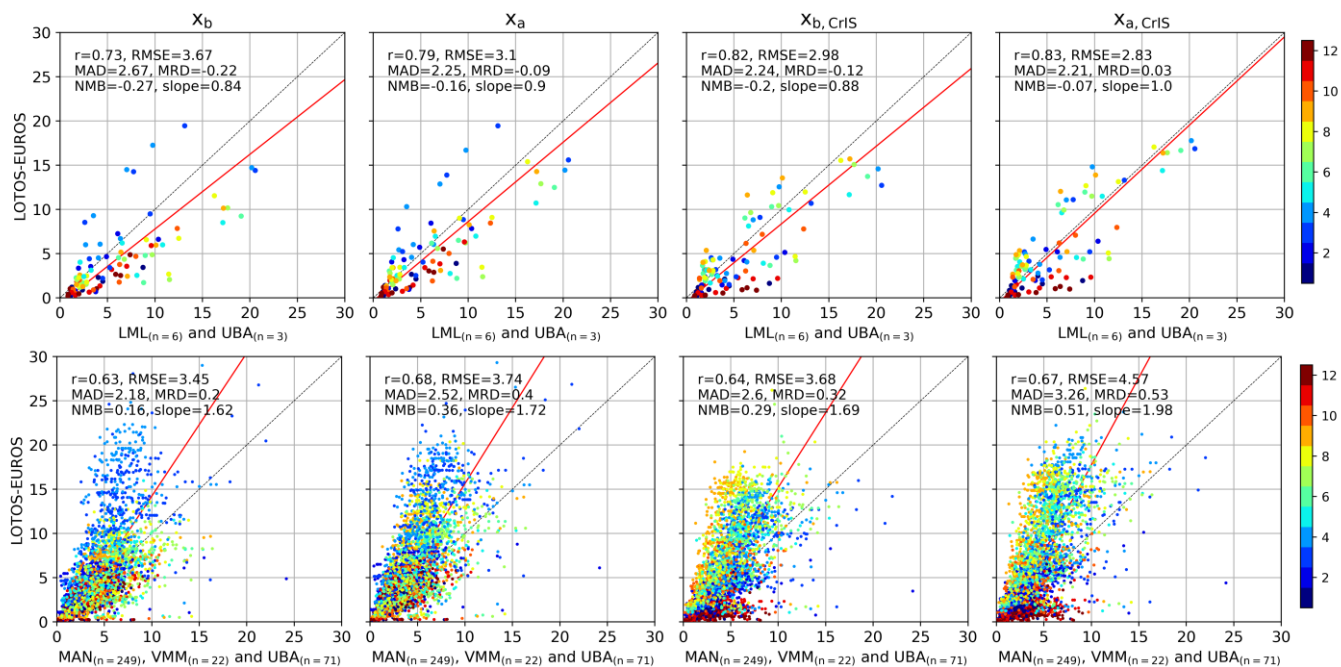
3.4.2. Temporal distribution

480 Fig. 13 shows the scatterplots of the monthly means per site. The modelled monthly NH_3 surface concentrations from the default background run (x_b) are strongly correlated with the hourly observation network ($r = 0.73$), and with the passive sampler network ($r = 0.63$). Both comparisons show a distinct overestimation of the NH_3 surface concentration in March and April. The observed NH_3 surface concentrations at the hourly observation sites are higher than the modelled ones during the rest of the year. At the passive sampler sites, the observed versus modelled monthly NH_3 surface concentrations during the rest of the year lie more around the one-on-one line. Here, too, the modelled NH_3 surface concentrations are slightly underestimated at 485 the beginning of summer (June and July). The NH_4^+ wet deposition is moderately correlated with monthly observations from wet-only samplers ($r = 0.44$). At these sites, a similar pattern is observed. The modelled NH_4^+ wet deposition is overestimated in spring (especially March and April), and underestimated during the rest of the year. In general, this comparison indicates an overestimation of the NH_3 spring peak emissions in the default model runs, particularly in March and April, and an 490 underestimation of the NH_3 emission during the rest of the year, mainly during summer (June, July, August).

The use of the CrIS-based NH_3 time factors ($x_{b,\text{CrIS}}$) led to an overall improvement at the hourly observation and wet-only sampler sites. Compared to the default background run (x_b), higher correlations and lower differences (RMSE, MAD, MRD, NMB) are observed. At the hourly observation sites, the comparison improved the most (x_b : $\{r = 0.73, \text{RMSE} = 3.67, \text{MAD} =$ 495 $2.67, \text{MRD} = -0.22, \text{NMB} = -0.27, \text{slope} = 0.84\}$ versus $x_{b,\text{CrIS}}$: $\{r = 0.82, \text{RMSE} = 2.98, \text{MAD} = 2.24, \text{MRD} = -0.12, \text{NMB} = -0.20, \text{slope} = 0.88\}$). Compared to observations from the passive sampler and wet-only sampler networks, the modelled monthly NH_3 surface concentration and NH_4^+ wet deposition fields now generally lie around the one-on-one line during spring (March, April, May). There is, on the other hand, an overestimation in July and August now. Moreover, as a result of the decrease in CrIS-based NH_3 time factors to zero during winter, the NH_3 surface concentration and NH_4^+ wet deposition in 500 December is underestimated in the $x_{b,\text{CrIS}}$ run.

Compared to the background runs (x_b and $x_{b,\text{CrIS}}$), the two analysis runs (x_a and $x_{a,\text{CrIS}}$) show higher correlations with all types of measurements. The differences (RMSE, MAD, MRD, NMB) between the observed and modelled monthly NH_3 surface concentrations at the passive sampler sites are now, on the other hand, larger in the two analysis runs (x_a and $x_{a,\text{CrIS}}$), illustrating 505 an overall overestimation of the NH_3 concentrations in background regions. Although a large shift in the temporal distribution of the monthly NH_4^+ wet deposition is observed, the differences between the observed and modelled values remained similar. At the hourly observation sites, the comparison improved the most in the analysis run with the CrIS-based NH_3 time factors ($x_{a,\text{CrIS}}$). Here, compared to the default background run (x_b), higher correlations and smaller differences were found (x_b : $\{r = 0.73, \text{RMSE} = 3.67, \text{MAD} = 2.67, \text{MRD} = -0.22, \text{NMB} = -0.27, \text{slope} = 0.84\}$ versus $x_{a,\text{CrIS}}$: $\{r = 0.83, \text{RMSE} = 2.83, \text{MAD} =$ 510 $2.21, \text{MRD} = 0.03, \text{NMB} = -0.07, \text{slope} = 1.0\}$).

NH₃ surface concentration (ug m⁻³)



NH₄ wet deposition (mg m⁻²)

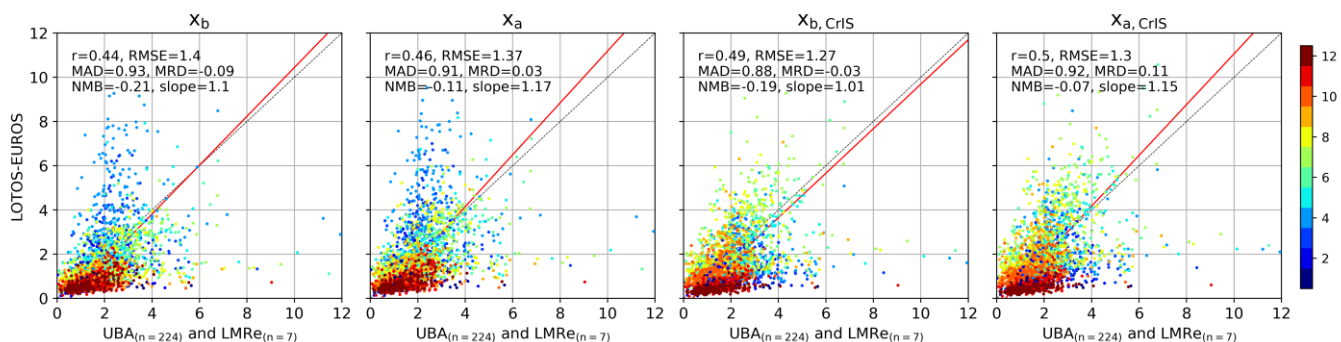


Figure 13: Comparison of the modelled monthly mean NH₃ surface concentrations and NH₄⁺ wet deposition fields to in-situ observations. The colors indicate the month.

515 3.4.3. Regional patterns

The modelled NH₃ surface concentrations were compared to observations from each passive sampler network separately. Fig. S15, S16 and S17 show comparison with the MAN network in the Netherlands, the UBA network in Germany and the VMM network in Belgium, respectively. In the default background run (x_b), the Dutch sites with relatively higher NH₃ surface concentrations are overestimated, most of which are located along the eastern border of the Netherlands. The highest correlation coefficients and lowest differences (RMSE, MAD) are found at the VMM network in Belgium. Here, the lower

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NH₃ surface concentration sites are overestimated and the higher NH₃ concentrations sites are underestimated in the default background run (x_b). At the German UBA stations, the comparison lies more around the one-on-one line. The mean NH₃ surface concentrations at the sites close to the western border of Germany are generally overestimated in the default background run (x_b).

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The use of the CrIS-based NH₃ time factors ($x_{b,CrIS}$) led to an overall increase in modelled mean NH₃ surface concentrations compared to the default background run (x_b). This led to a slight, overall increase in differences (RMSE and MAD) at all networks. Furthermore, steeper slopes were found at all three networks, i.e., the modelled NH₃ surface concentrations increased relatively more at sites with already higher concentrations. The same is observed in the two analysis runs (x_a and $x_{a,CrIS}$), but to a greater extent. Compared to background runs (x_b and $x_{b,CrIS}$), the differences (RMSE, MAD) between the modelled and observed concentrations were relatively higher at all networks. At the Dutch MAN network, a slightly higher correlation coefficient is observed.

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Fig. S18 of the supplementary materials shows another comparison of the modelled and observed NH₃ surface concentrations at the hourly observation stations at daily resolution. Here, the correlation coefficient, root-mean-squared error RMSE, the mean difference MD and the slope are shown per site. The stations are located in different NH₃ emission regimes and are sorted by increasing NH₃ surface concentrations. The modelled NH₃ surface concentrations in the default background run (x_b) are generally overestimated at stations with low NH₃ emission regimes and underestimated at stations with medium to high NH₃ emission regimes. The use of the CrIS-based NH₃ time factors ($x_{b,CrIS}$) led to an improved comparison (higher correlation coefficient and lower RMSE) at the Dutch stations, but a worse comparison at the German stations. On a monthly basis, the comparison to the German UBA sites slightly worsened after the use of the CrIS-based NH₃ time factors ($x_{b,CrIS}$) (Fig. S19). The modelled NH₃ surface concentrations in the background run with the CrIS-based NH₃ time factors ($x_{b,CrIS}$) were, on the other hand, closer to the observations of the Dutch LML network in most months, with a lower differences (RMSE, MD) and slopes closer to 1. Here, the largest increase in correlation coefficients were found in March and April. In both analysis runs (x_a and $x_{a,CrIS}$), the correlation coefficient improved and lower model-observation differences were found at all sites. Here, no clear distinction between sites located in different NH₃ emission regimes can be seen.

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Compared to the default background run (x_b), the modelled NH₃ surface concentrations in the background run with the CrIS-based NH₃ time factors ($x_{b,CrIS}$) thus improved the most at Dutch stations located in medium to high NH₃ emission regimes. Most of the Dutch stations are located in the proximity of agricultural hotspots. The German stations, on the other hand, are located in background areas in central Germany, further away from major agricultural hotspots for NH₃. Fig. S8 of the supplementary materials shows the fitted CrIS-based NH₃ time factors at each site. The fitted NH₃ time factors at the majority of the Dutch stations show clear, identifiable peaks, in particular the spring peak. Moreover, most Dutch sites show clear year-to-year variations. For the German stations, on the other hand, the fitted NH₃ time factors are much flatter and show much less

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555 interannual variation. This indicates that the observed CrIS-NH₃ surface concentrations at these locations remained around the same level, and thus that no clear (inter)annual patterns were present in the CrIS data.

In the Netherlands, the CrIS-based NH₃ time factors led to an improvement in the representation of the NH₃ spring peak. A time-series of the observed daily NH₃ surface concentrations at LML sites Valthermond and Zegveld are plotted in Fig. S20
560 of the supplementary materials. The modelled NH₃ surface concentrations in the default background run (x_b) start to rise too early in the year, particularly in 2014. In the background run with the CrIS-based NH₃ time factors ($x_{b,CrIS}$), both the start and the duration of the spring peak in NH₃ concentration improve. Here, the onset of the spring peak is delayed, better matching the observed NH₃ timeseries.

4.1. Summary

565 In this study, the CrIS-NH₃ product is integrated into the LOTOS-EUROS chemical transport model using two different methods. In the first method, the CrIS-NH₃ surface concentrations were used to fit spatially varying NH₃ time factors to redistribute the NH₃ emission inputs in LOTOS-EUROS over the year. In the second method, the CrIS-NH₃ columns were assimilated to adjust NH₃ emissions through local Ensemble Transform Kalman filtering in a top-down approach.

570 The fitted NH₃ time factors based on the CrIS-NH₃ surface concentrations led to a major temporal redistribution of the NH₃ emissions. In most regions, the updated NH₃ time profiles became flatter, with an overall decrease in spring (March to May) NH₃ emissions and an increase in NH₃ emissions in June to October. As a result, the mean modelled NH₃ fields between 2014 and 2018 spatially changed by up to +25% in NH₃ surface concentrations, -5 to +5% in NH₃ total column concentrations and -5 to +5% in NH_x budget. The CrIS-based NH₃ time factors added an extra interannual variation of up to ±25% in the annual
575 mean NH₃ concentrations and deposition fields. Data assimilation of the CrIS-NH₃ columns with the LETKF led to a unanimous increase in total NH₃ emissions. The modelled NH₃ fields between 2014 and 2018 changed with up to +30% in NH₃ surface concentrations, up to +20% in NH₃ total column concentrations and +10 to +25% in NH_x budget. The largest increases in NH₃ emissions (+30%) were found over the south of the Netherlands (Brabant), the west of Belgium (West-Vlaanderen) and a large region in northeastern Germany. The temporal distribution of the NH₃ emissions wasn't largely
580 adjusted by the LETKF. The largest positive NH₃ emission updates took place in late summer and the beginning of autumn (July to September) and both increases and decreases in NH₃ emissions were observed in spring (March to May).

The modelled NH₃ surface concentration and NH₄⁺ deposition fields were compared to in-situ observations. **The statistics are summarized in Table 2.** Our results illustrate that the strength of the first method, i.e., CrIS-based NH₃ time factors, primarily
585 lies in improving the temporal distribution of the NH₃ emissions. Compared to in-situ networks, an overall increase in correlation coefficient and clear decrease in differences (RMSE, MAD, MRD, NMB) at the hourly observation and the wet-only sampler sites was observed. Moreover, time-series of observed daily NH₃ surface concentrations illustrate that using the

CrIS-based NH₃ time factors resulted in a better representation of both the onset and duration of the spring NH₃ peak in the Netherlands. The second method, data assimilation of the CrIS-NH₃ columns with the LETKF, improved the comparability to in-situ observation both spatially and temporally. Here, higher correlations with both annual and monthly observed mean NH₃ surface concentrations and NH₄⁺ wet deposition were observed. This method also led to a decrease in differences (RMSE, MAD, MRD, NMB) at the hourly observation and the wet-only sampler sites. The mean NH₃ surface concentrations at the passive sampler sites, on the other hand, were more strongly overestimated in both methods. The comparison to in-situ observations improved the most, both spatially and temporally, in the run where both methods are combined (x_{a,CrIS}). This illustrates that an initial, observation-based, rescaling of the NH₃ emissions enhances the adaptability of the LETKF, herewith thus improving the modelled NH₃ surface concentration and NH₄⁺ wet deposition fields.

		hourly observations				passive samplers				wet-only samplers			
		Xb	Xa	Xb,cris	Xa,cris	Xb	Xa	Xb,cris	Xa,cris	Xb	Xa	Xb,cris	Xa,cris
annual mean	r	0.88	0.89	0.87	0.87	0.69	0.70	0.69	0.70	0.45	0.45	0.46	0.46
	RMSE	2.79	2.34	2.50	2.20	2.05	2.67	2.45	3.22	0.95	0.91	0.94	0.92
	MAD	1.96	1.68	1.82	1.69	1.55	2.05	1.84	2.49	0.63	0.61	0.62	0.61
	MRD	-0.15	-0.02	-0.05	0.11	0.25	0.46	0.37	0.61	-0.13	-0.03	-0.14	-0.02
	NMB	-0.28	-0.17	-0.20	-0.08	0.16	0.36	0.28	0.50	-0.22	-0.13	-0.22	-0.11
	slope	0.61	0.71	0.68	0.76	1.17	1.39	1.32	1.53	0.81	0.90	0.84	0.95
monthly mean	r	0.73	0.79	0.82	0.83	0.63	0.68	0.64	0.67	0.44	0.46	0.49	0.50
	RMSE	3.67	3.10	2.98	2.83	3.45	3.74	3.68	4.57	1.40	1.37	1.27	1.30
	MAD	2.67	2.25	2.24	2.21	2.18	2.52	2.60	3.26	0.93	0.91	0.88	0.92
	MRD	-0.22	-0.09	-0.12	0.03	0.20	0.40	0.32	0.53	-0.09	0.03	-0.03	0.11
	NMB	-0.27	-0.16	-0.20	-0.07	0.16	0.36	0.29	0.51	-0.21	-0.11	-0.19	-0.07
	slope	0.84	0.90	0.88	1.00	1.62	1.72	1.69	1.98	1.10	1.17	1.01	1.15

Table 2: Summary of the computed statistics (correlation coefficient (r), root mean square error (RMSE), mean absolute difference (MAD), mean relative difference (MRD), normalized mean bias (NMB) and slope) for each type of instruments from Figure 12 and 13.

4.2. Discussion

4.2.1. CrIS-based NH₃ time factors

The temporal redistribution of the NH₃ emissions after using the fitted CrIS-based NH₃ time factors led to a significantly better representation of the temporal variation in NH₃ emissions, especially the spring peak. Compared to in-situ observations, however, the NH₃ surface concentrations were overestimated in late summer and autumn (August to October). Further fine-tuning of the fitting algorithm could help to reduce the current overestimation and potentially improve the fitted NH₃ time

factors. For example, data filtering and selection criteria could be adapted. Narrowing the selection radius used for selecting the CrIS-NH₃ observations could for instance lead to a better representation of the NH₃ concentrations locally. This, however, may not always be possible, as a minimum number of observations is needed for a converging fit. Furthermore, the fitting algorithm currently doesn't allow for NH₃ area emissions during winter, because of the limited number of available CrIS observations at this time. As a result, the fitted NH₃ time factors show a relatively steep increase at the beginning of spring and a decrease at the beginning of winter. This could lead to step-like functions in areas where clear NH₃ peaks in the CrIS-NH₃ data are absent. However, as this mainly occurs in areas with little to no NH₃ emissions, this shouldn't severely impact the modelled NH₃ concentrations in this study.

615 **4.2.2. Local Ensemble Transform Kalman Filter**

The NH₃ emission updates computed by the Local Ensemble Transform Kalman Filter (LETKF) always remain tied to the initial model fields by a certain uncertainty range. As such, data assimilation of the CrIS-NH₃ columns with the LETKF is mainly suitable for fine-tuning NH₃ emissions in regions where the NH₃ emissions are already relatively well known. The chosen LETKF configuration is for instance not able to correct for missing NH₃ emissions in areas where little or no initial NH₃ emissions are present. Furthermore, the LETKF is unable to resolve temporal patterns well without sensible input, as was illustrated in an experiment with homogeneous NH₃ emission fields (supplement section S1).

The LETKF filter settings used in this modelling setup ($\rho = 15$ km, $\sigma = 0.5$, $\tau = 3$ days) led to a maximum emission increase of roughly ~30% on the initial NH₃ emissions for long-term simulations. The choice of these filter settings affects the adaptability of the LETKF, i.e., the achievable emission adjustments by correction factors. In this study, a temporal length scale τ of 3 days was chosen as a compromise between short time scales needed for irregular emissions (e.g., fertilizer application) and longer time scales needed for regular emissions (e.g., stables and other point sources). Moreover, it matches the average satellite revisiting time per grid cell given the number of CrIS-NH₃ observations left after data selection (Fig. S21). A spatial correlation of $\rho = 15$ km was chosen because it matches the footprint size of the satellite. Furthermore, as shown in section S1 in the supplement, increasing standard deviation σ leads to larger, positive β factors. To prevent further overestimations in background regions, a σ of 0.5 was used for this region.

The current LETKF settings could for instance be improved by using spatially varying τ values. The choice of τ could be optimized for each emission category in the model. Locations with fertilizer application as dominant NH₃ emission source could for instance be set to lower τ values than locations with predominantly regular NH₃ sources. Another way to optimize the filter settings would be to study timeseries of the model-satellite discrepancies in more detail and base the choice of τ on this. A more apparent memory effect (i.e., higher τ) would be useful in areas with consistent model-satellite discrepancies, whereas in areas with incidental differences a lower τ would be more appropriate. Similarly, statistical analysis of the already computed emission perturbation factors β could be performed. In this study, the model uncertainty follows a normal

640 distribution in the current model setup. The distribution of the NH₃ concentrations, however, is, in reality, better approximated by a log-normal distribution. It would therefore be more realistic to use a log-normal distribution for the model uncertainty as well. This would incidentally allow for larger correction factors when high NH₃ peaks are observed, for instance after fertilizer application.

645 In the current LETKF model setup, only the NH₃ emissions are perturbed. Thus, the discrepancies between the observed and modelled NH₃ concentrations are currently thus fully assigned to errors in the underlying model NH₃ emissions. However, other model uncertainties could also cause these discrepancies, for instance uncertainties in other model inputs (e.g., other trace gas emissions) or parameterizations (e.g., deposition routines). In a follow-up study, it would be interesting to further investigate to the effect of an inverted LETKF setup, where model sink terms are perturbed instead of the source terms.
650 However, the current setup is the most obvious as the NH₃ emissions are likely the largest source of model uncertainty in this area. It would also be interesting to assimilate in-situ observations and/or other satellite products (for instance IASI-NH₃) simultaneously in a follow-up study.

4.2.3. Data products

Direct comparison of the observed and simulated NH₃ columns showed distinctly lower NH₃ total column concentrations in
655 LOTOS-EUROS. This discrepancy could be the result of a systematic underestimation of the input NH₃ emission in LOTOS-EUROS, or other model uncertainties. It could, on the other hand, also be partially related to the CrIS observations themselves. Here, only CrIS observations with the highest quality flag (QF=5) were used, which for instance could have resulted in a bias towards observations with higher NH₃ concentrations or during good weather (e.g., no clouds), as these observations usually have a lower uncertainty. Moreover, an offset of approximately $\sim 0.5 \times 10^{16}$ molecules/cm² is observed. This seems to be the
660 effect of the detection limit of the CrIS instrument, which is unable to detect very low NH₃ concentrations. Furthermore, this, too, could be enhanced by the relatively strict data selection criteria used in this study, which favors higher NH₃ concentrations that usually have a lower uncertainty. In the next version of the CrIS-NH₃ product, which was not yet available for this study, these non-detects are addressed. This might lead to lower NH₃ concentrations in background regions and partially solve this discrepancy. Moreover, this could also result in a better comparison with observations of the passive sampler networks.

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The differences between the modelled and observed NH₃ concentrations and NH₄⁺ wet deposition fields are partially related to limitations in the spatial representativeness of the in-situ observations. The comparison of the different model runs to in-situ observations showed an overall overestimation at the passive sampler sites. These sites are mainly located in nature areas and therefore assumed to be representative of background regions with little to no NH₃ emissions. However, especially in the
670 Netherlands, the landscape layout is very heterogenous and the nature areas are relatively small. As a result, at the current model grid size, each model pixel is likely to include other landscape elements than nature alone. The larger model scale averages out the small-scale effects, thus leading to an overestimation. The opposite is true for the hourly observation sites

located in medium to high NH₃ emission regimes. Especially at sites close to NH₃ emission sources, an underestimation is expected.

675 **4.2.4. Conclusions**

To conclude, satellite observed CrIS-NH₃ timeseries are helpful in improving NH₃ emissions, both spatially and temporally. Our results illustrated that CrIS-NH₃ surface concentrations can be successfully used to fit spatially variable NH₃ time factors, which allows us to improve temporal NH₃ emission distributions relatively easy in a forward modelling setup. Comparison with in-situ NH₃ surface concentrations and NH₄⁺ wet deposition observations showed that the fitted CrIS-based NH₃ time factors were particularly useful for adjusting the timing and duration of the NH₃ spring peak in medium to high NH₃ regimes. Moreover, the comparison showed that the CrIS-based NH₃ time factors improve the temporal distribution of the modelled NH₃ surface concentrations and NH₄⁺ wet deposition fields. Our results show that data assimilation of the CrIS-NH₃ columns data with the Local Ensemble Transform Kalman Filter (LETKF) improves the comparability with in-situ observations both spatially and, to a lesser extent, temporally, too. As the adaptability of the LETKF is limited by the uncertainty in the modelled fields, the strength of this method primarily lies in fine-tuning pre-existing NH₃ emission patterns. As a result, this method proved particularly useful for improving spatial NH₃ distributions in long-term simulations. Moreover, our results illustrated that combining both methods enhanced the adaptability of the LETKF, and led to the largest improvements in modelled NH₃ surface concentration and NH₄⁺ wet deposition fields compared to in-situ observations.

690 *Author contributions.* SvdG worked on the observation-based NH₃ time factors. The CrIS-NH₃ dataset was provided by ED and MWS. SvdG, ED, AS and RK worked on the modelling and data assimilation. JWE, ED, AS, MWS, RK and MS helped with the interpretation of the results. SvdG wrote the paper with contributions from all co-authors.

Competing interests. The authors declare that they have no conflict of interest.

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