1	Unchanged PM2.5 levels over Europe during COVID-19 were
2	buffered by ammonia
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16 Abstract

The coronavirus outbreak in 2020 had devastating impact on human life, albeit a positive effect 17 for the environment reducing emissions of primary aerosols and trace gases and improving air quality. 18 In this paper, we present inverse modelling estimates of ammonia emissions during the European 19 20 lockdowns of 2020 based on satellite observations. Ammonia has a strong seasonal cycle and mainly originates from agriculture. We further show how changes in ammonia levels over Europe, in 21 conjunction with decreases in traffic-related atmospheric constituents modulated PM2.5. The key 22 result of this study is a -9.8% decrease in ammonia emissions in the first half of 2020 compared to 23 the same period in 2016–2019 attributed to restrictions related to the global pandemic. We further 24 calculate the delay in the evolution of the ammonia emissions in 2020 before, during and after 25 lockdowns, by a sophisticated comparison of the evolution of ammonia emissions during the same 26 time periods for the reference years (2016-2019). Our analysis demonstrates a clear delay in the 27 evolution of ammonia emissions of -77 kt, that was mainly observed in the countries that suffered the 28 strictest travel, social and working measures. Despite the general drop in emissions during the first 29 half of 2020 and the delay in the evolution of the emissions during the lockdown period, satellite and 30 ground-based observations showed that the European levels of ammonia increased. On one hand, this 31 was due to the reduction of SO_2 and NO_x (precursors of the atmospheric acids with which ammonia 32 reacts) that caused less binding and thus less chemical removal of ammonia (smaller loss - higher 33 lifetime); on the other, the majority of the emissions persisted, because ammonia mainly originates 34 35 from agriculture, a primary production sector that was influenced slightly by the lockdown restrictions. Despite the projected drop in various atmospheric aerosols and trace gases, PM2.5 levels 36 37 stayed unchanged or even increased in Europe, due to a number of reasons attributed to the complicated NH_3 - H_2SO_4 - HNO_3 system. Higher water vapour during the European lockdowns 38 favoured more sulfate production from SO_2 and OH (gas phase) or O_3 (aqueous phase). Ammonia 39 first reacted with sulfuric acid also producing sulfate. Then, the continuously accumulating free 40 ammonia reacted with nitric acid shifting the equilibrium reaction towards particulate nitrate. In high 41 free ammonia atmospheric conditions such as those in Europe during the 2020 lockdowns, a small 42 reduction of NO_x levels drives faster oxidation toward nitrate and slower deposition of total inorganic 43 nitrate causing high secondary PM2.5 levels. 44

46 1 Introduction

Ammonia (NH3), the most abundant gas, has played a vital role in the evolution of human 47 population through the Haber-Bosch process (Chen et al., 2019). However, today it is recognized to 48 have significant negative influence, not only for the environment (Stevens et al., 2010), but also for 49 human population (Cohen et al., 2017; Pope and Dockery, 2006) and the climate (De Vries et al., 2011). 50 As an alkaline molecule, ammonia regulates the pH of clouds, while its excessive atmospheric 51 deposition and terrestrial runoff affect natural reservoirs creating algae blooms and degrading water 52 quality (Camargo and Alonso, 2006; Krupa, 2003). When emitted to the atmosphere, it reacts with 53 54 the abundant sulfuric and nitric acids (Malm, 2004) forming sulfate, nitrate, and ammonium and contributing up to 50% to the total aerosol mass (Anderson et al., 2003). The latter has implications 55 56 for human health (Gu et al., 2014) as aerosols penetrate the human respiratory system and accumulate in the lungs (Pope III et al., 2002) causing premature mortality (Lelieveld et al., 2015). Furthermore, 57 through secondary aerosol formation (Pozzer et al., 2017), ammonia has a significant impact (i) on 58 regional climate (Bellouin et al., 2011) causing visibility problems and contributing to haze effect, 59 and (ii) on global climate directly by scattering incoming radiation (Henze et al., 2012) and indirectly 60 as cloud condensation nuclei (Abbatt et al., 2006) altering the Earth's radiative balance. 61

The largest portion of atmospheric ammonia originates from the synthesis of nitrogen 62 fertilizers, which are in high demand for agriculture (Erisman et al., 2007). The expansion of intensive 63 agriculture during the 20th century has increased atmospheric ammonia above natural levels (Erisman 64 et al., 2008), while the projected growth of the global population will likely create larger nutritional 65 needs that are expected to further increase ammonia emissions during the 21st century (Pai et al., 66 67 2021). Other sources of ammonia include emissions from livestock (Sutton et al., 2000a), industry, ammonia-rich watersheds (Sørensen et al., 2003), traffic (Kean et al., 2009), sewage (Reche et al., 68 69 2012), humans (Sutton et al., 2000b), biomass and domestic combustion (Sutton et al., 2008; Fowler 70 et al., 2004) and volcanic eruptions (Sutton et al., 2008).

In the past years, atmospheric ammonia observations were mostly limited to ground-based 71 measurements with relatively sparse monitoring networks. This resulted in large emission 72 uncertainties in regions poorly covered by measurements (Heald et al., 2012). Today, satellite 73 products are capable to record daily ammonia column concentrations providing useful information 74 75 on its atmospheric abundance. Recently, Van Damme et al. (2021) analyzed Infrared Atmospheric Sounding Interferometer (IASI) retrievals and showed increased ammonia levels over most of Europe 76 after 2015. Then, suddenly the COVID-19 outbreak came in 2020 creating a unique situation 77 (Baekgaard et al., 2020), which affected all segments of life in a detrimental way (Chakraborty and 78 79 Maity, 2020; Sohrabi et al., 2020). As a measure to inhibit further spread of the virus, authorities took

strict social, travel and working restrictions for months, which resulted in lower traffic-related 80 emissions and improved air quality (Bauwens et al., 2020; Dutheil et al., 2020; Sicard et al., 2020). 81 Illustrating the impact on emissions, Guevara et al. (2021) reported average emission reductions in 82 Europe to be 33% for NO_r , 8% for non-methane volatile organic compounds (NMVOCs), and 7% 83 84 for SO_x during the strictest lockdowns in 2020, while more than 85% of the total reduction is attributed to road transport. CO₂ emissions were also decreased by 11% over Europe during the first 85 lockdowns (Diffenbaugh et al., 2020), so as aerosols did; notably Black Carbon (BC) emissions 86 dropped by 11% (Evangeliou et al., 2020) and Aerosol Optical Depth (AOD) decreased up to 20% 87 over Central and Northern Europe (Acharya et al., 2021). 88

While the COVID-19 lockdown impact on emissions for primary aerosols and trace gases has 89 been studied extensively, how ammonia emissions were affected in Europe is unknown. The latter is 90 91 very important and may have largely moderated the atmospheric levels of particulate matter (Giani et al., 2020; Guevara et al., 2021; Matthias et al., 2021), because of ammonia's contribution to 92 93 secondary PM2.5 (particulate matter) formation (Anderson et al., 2003). Here, we make use of satellite measurements of ammonia and a novel inversion algorithm to track how ammonia emissions 94 95 changed before, during and after the European lockdowns in 2020. We examine the reasons behind the estimated changes and validate the results against ground-based observations from the EMEP 96 97 measurement network (<u>https://emep.int/mscw/</u>, Figure S 1). Finally, we calculate the resulting impact of ammonia changes during the European lockdowns on the formation of PM2.5 using a chemistry 98 transport model (CTM) and try to interpret the mechanisms governing these changes. 99

100 2 Methods

101 2.1 Cross-Track Infrared Sounder (CrIS) ammonia measurements

The CrIS sensor onboard the NASA Suomi National Polar-orbiting Partnership provides 102 atmospheric soundings at high spectral resolution (0.625 cm-1) (Shephard et al., 2015) resulting in 103 improved vertical sensitivity for ammonia at the surface (Zavyalov et al., 2013). The CrIS fast 104 physical algorithm (Shephard and Cady-Pereira, 2015) retrieves ammonia at 14 vertical levels using 105 a physics-based optimal estimation retrieval, which also provides the vertical sensitivity (averaging 106 kernels) and an estimate of the retrieval errors (error covariance matrices) for each measurement. 107 Shephard et al. (2020) reports a total column random measurement error of 10-15%, with total 108 random errors of $\sim 30\%$. The individual profile random errors are 10–30%, while total profile random 109 errors increase above 60% due to the limited vertical resolution (Shephard et al., 2020). Vertical 110 111 sensitivity and error calculations are also important when using CrIS observations in satellite inverse modelling applications (Li et al., 2019; Cao et al., 2020) as a satellite observational operator can be 112

generated in a robust manner (see next sections). The detection limit of CrIS measurements has been calculated down to 0.3–0.5 ppbv (Shephard et al., 2020) and the product has been validated extensively against ground-based observations (Dammers et al., 2017; Kharol et al., 2018) showing small differences and high correlations.

Daily CrIS ammonia satellite measurements (version 1.6.2) were gridded on $0.5^{\circ} \times 0.5^{\circ}$ covering 117 all Europe (10°W–50°E, 25°N–75°N) from 1st January to 30th June 2020. Data were screened prior 118 to its use with Quality Flag \geq 4, as recommended in the CrIS documentation, and Cloud Flag \neq 1. The 119 latter excludes retrievals that are performed under thin cloud conditions and are not as reliable as 120 retrievals performed under cloud-free conditions (White et al., 2023). Gridding was chosen to limit 121 the large number of observations (around 10,000 per day per vertical level for 2550 retrievals January 122 to June 2020), hence the need for a large number of source-receptor matrices (SRMs), which is 123 computationally inefficient. Specifically, day and night-time observations from CrIS were averaged 124 in each 0.5° resolution grid-cell daily from 1st January to 30th June 2020. This gridding method, 125 although simple, it gives more robust results than classic interpolation methods and presents small 126 standard deviations of the gridded values (see Tichý et al., 2023). Sitwell and Shephard (2021) 127 showed that the averaging kernels of CrIS ammonia are significant only for the lowest six levels (the 128 upper eight have no influence into the satellite observations) and therefore we have considered these 129 six vertical levels (~1018-619 hPa). 130

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2.2 Source-receptor matrix (SRM) calculations

SRMs were calculated for each 0.5°×0.5° grid-cell over Europe (10°W–50°E, 25°N–75°N) 132 using the Lagrangian particle dispersion model FLEXPART version 10.4 (Pisso et al., 2019) adapted 133 to model ammonia. The model releases computational particles that are tracked backward in time 134 using hourly ERA5 (Hersbach et al., 2020) assimilated meteorological analyses from the European 135 Centre for Medium-Range Weather Forecasts (ECMWF) with 137 vertical layers and a horizontal 136 resolution of 0.5°×0.5°. FLEXPART simulates turbulence (Cassiani et al., 2015), unresolved 137 mesoscale motions (Stohl et al., 2005) and convection (Forster et al., 2007). SRMs were calculated 138 for 7 days backward in time, at temporal intervals that matched satellite measurements and at spatial 139 resolution of 0.5°×0.5°. This 7-day backward tracking is sufficiently long to include almost all 140 ammonia sources that contribute to surface concentrations at the receptors given a typical atmospheric 141 lifetime of about a day (Evangeliou et al., 2021; Van Damme et al., 2018). 142

The complicated heterogeneous chemistry of ammonia was modelled with the Eulerian model LMDz-OR-INCA, which couples the LMDz (Laboratoire de Météorologie Dynamique) General Circulation Model (GCM) (Hourdin et al., 2006) with the INCA (INteraction with Chemistry and Aerosols) model (Folberth et al., 2006; Hauglustaine et al., 2004) and with the land surface dynamical

vegetation model ORCHIDEE (ORganizing Carbon and Hydrology In Dynamic Ecosystems) 147 (Krinner et al., 2005). The model has a horizontal resolution of 2.5°×1.3°, and 39 hybrid vertical 148 levels extending to the stratosphere. It accounts for large-scale advection of tracers (Hourdin and 149 Armengaud, 1999), deep convection (Emanuel, 1991), while turbulent mixing in the planetary 150 151 boundary layer (PBL) is based on a local second-order closure formalism. The model simulates atmospheric transport of natural and anthropogenic aerosols and accounts for emissions, transport 152 (resolved and sub-grid scale), and dry and wet (in-cloud/below-cloud scavenging) deposition of 153 chemical species and aerosols interactively. LMDz-OR-INCA includes a full chemical scheme for 154 cycle and nitrate particle formation, as well as a state-of-the-art 155 the ammonia CH₄/NO_x/CO/NMHC/O₃ tropospheric photochemistry (Hauglustaine et al., 2014). The global 156 transport of ammonia was simulated for 2020 with a month of spin-up by nudging the winds of the 157 3-hourly ERA5 (Hersbach et al., 2020) with a relaxation time of 10 days (Hourdin et al., 2006). 158

For the calculation of ammonia's lifetime, LMDz-OR-INCA ran with traditional emissions for 159 anthropogenic, biomass burning and oceanic emission sources from ECLIPSEv5 (Evaluating the 160 CLimate and Air Quality ImPacts of Short-livEd Pollutants), GFED4 (Global Fire Emission Dataset) 161 and GEIA (Global Emissions InitiAtive) (hereafter called "EGG") (Bouwman et al., 1997; Giglio et 162 163 al., 2013; Klimont et al., 2017). FLEXPART uses the exponential mass removal for radioactive 164 species based on the e-folding lifetime (Pisso et al., 2019), which gives the time needed to reduce the species mass to 1/e contribution. We calculated the e-folding lifetime (Kristiansen et al., 2016; Croft 165 166 et al., 2014) of ammonia from LMDz-OR-INCA, assuming that the loss occurs as a result of all processes affecting ammonia (chemical reactions, deposition) with a minimum time-step of 1800 s. 167 Then we calculated the exponential loss of ammonia and the respective loss-rate constant κ (s⁻¹). We 168 point to Tichý et al. (2023) for more details on the methodology to avoid repetition. 169

Ammonia has a complicated atmospheric chemistry and may react with sulfuric and nitric acid 170 producing sulfate and nitrate. However, under certain atmospheric conditions, the equilibrium 171 reaction with nitric acid can be shifted to the left producing free ammonia (Seinfeld and Pandis, 2000). 172 Tichý et al. (2023) showed that production of free ammonia happened very rarely in continental 173 Europe in 2013–2020 period. Nevertheless, we have previously published a full validation of the 174 obtained CTM concentrations against all the available ground-based measurements of ammonia 175 globally (Tichý et al., 2023), from the EMEP network (<u>https://emep.int/mscw/</u>) in Europe, EANET 176 (East Asia acid deposition NETwork) in Southeastern Asia (https://www.eanet.asia/) and AMoN 177 (Ammonia Monitoring Network in the US, AMoN-US; National Air Pollution Surveillance Program 178 (NAPS) sites in Canada) in North America (<u>http://nadp.slh.wisc.edu/data/AMoN/</u>). 179

180 2.3 Inverse modelling of ammonia emissions

The proposed inversion method is based on a comparison of the CrIS satellite observations with the model profile retrievals to estimate the spatiotemporal ammonia emissions. The comparison of remote-sense observations such as CrIS with model (or in-situ) profiles is not straightforward as in the cases of ground-based observations. Here, we used the more rigorous approach of the "instrument operator" (see equation below), after interpolation of the model profile to the first six levels of the satellite product (Rodgers, 2000):

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$$\ln(v^{ret}) = \ln(v^a) + A(\ln(v^{true}) - \ln(v^a))$$
 Eq. 1

where v^{ret} is the retrieved profile concentration vector, v^a is a priori profile concentration vector, 188 v^{true} is the true profile concentration vector, and A is the averaging kernel matrix in logarithmic 189 space (for each 0.5°×0.5° resolution grid-cell). In our inversion setup, we directly compared the 190 retrieved v^{ret} and the observed satellite column concentration v^{sat} that is given by CrIS. In our case, 191 v^{true} is equal to the modelled concentration v^{mod} calculated from the SRMs and a prior emission 192 inventory. The argument for this approach is that v^{ret} is what the satellite would observe if v^{mod} was 193 the true profile. This is a useful technique for evaluating if the retrieval algorithm is performing as 194 195 designed, i.e., is it unbiased and the calculated root mean square error (RMSE) is within the expected variability. Further details about the algorithm and the setup can be found in Tichý et al. (2023). 196

197 The goal of the inversion is to iteratively update prior emissions by minimizing the distance 198 between v^{sat} and v^{ret} by correcting the emission flux x in the term $v^{mod} = srm^{Flex}x^a$ (srm^{Flex} 199 denotes the FLEXPART SRMs), at each grid-cell and each of the six vertical levels that are important 200 for CrIS (Sitwell et al., 2022):

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The inverse problem is constructed for each spatial element of the computational domain. Inspired by the construction of covariance matrix in Cao et al. (2020), we consider 4° surroundings (445 km), expressed by the index set S, of which the column concentrations are considered due to computational effectivity. Note that we observed low sensitivity of resulting emission estimates to this choice. Then, we can formulate the inverse problem for each spatial element as:

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$$\left[v_{s_i}^{sat}; s_i \in \mathbb{S}\right] = \left[v_{s_i}^{ret}; s_i \in \mathbb{S}\right] q^{\mathbb{S}} \qquad Eq. 3$$

where the left side of the equation is formed by the vector with aggregated CrIS observations, vectors $v_{s_i}^{ret}$ form a block-diagonal matrix, and $q^{\$}$ is an unknown vector with correction coefficients for each temporal element of the emission. The inverse problem in Eq. 3 was solved using the least squares with adaptive prior covariance (LS-APC) algorithm (Tichý et al., 2016). The algorithm is based on a Bayesian model which assumes that all coefficients are positive and that the abrupt changes in their neighbouring values are less probable. It is shown that the method is less sensible to manual tunning of regularization parameters (see sensitivity tests in Tichý et al. (2020)) than classical optimization procedures, which is crucial for such a large dataset where each spatial element represents a separate inverse problem.

A detailed description of the algorithm is given in Tichý et al. (2016). Here, we do not describe 217 the algorithm again but explain a few modifications that were necessary for this study. By estimating 218 the correction coefficients $q^{\mathbb{S}}$ for each grid-cell of the spatial domain (10°W–50°E, 25°N–75°N), we 219 can propagate the coefficients through Eq. 2 to update a priori emissions x^a in the model 220 concentration term v^{mod} . We follow Li et al. (2019) and Cao et al. (2020) to bound the ratio between 221 the prior and the posterior emissions. The lower and upper bound of this ratio is set to 0.01 and 100, 222 respectively, to omit the unrealistically low or high emissions. We consider these bounds large 223 enough to allow for new emission sources to be exposed, not presented in the prior emissions. 224

We evaluate the performance of the inversion by using three a priori emission datasets, (i) one 225 based on Van Damme et al. (2018) calculations (Evangeliou et al., 2021) (hereafter denoted as "VD"), 226 (ii) the ECLIPSEv6 inventory (Klimont, 2022; Klimont et al., 2017) (combined with biomass burning 227 228 emissions from GFEDv4 (Giglio et al., 2013)) as the most recent one (denoted as "EC6G4"), and (iii) the average of four emission inventories for ammonia, except for these two mentioned before, "EGG" 229 (see previous section), and "NE" calculated from IASI (Infrared Atmospheric Sounding 230 Interferometer) observation (Evangeliou et al., 2021) (denoted as "avgEENV"). To account for the 231 spatiotemporal impact of the lockdown on the European emissions, we corrected prior emission 232 inventories of ammonia (EGG, EC6G4 and avgEENV) for 2020 using adjustment factors (AFs) 233 adopted from Doumbia et al. (2021). The same was done for SO_2 and NO_x (precursors of sulfuric and 234 nitric acid in the atmosphere) in EGG that was used to calculate ammonia's loss rates using LMDz-235 OR-INCA model (see section 2.2). This dataset provides, for the January-August 2020 period, 236 gridded AFs at a 0.1°×0.1° resolution on a daily resolution for transportation (road, air and ship 237 traffic), power generation, industry and residential sectors. The quantification of AFs is based on 238 activity data collected from different databases and previously published studies. These emission AFs 239 have been applied to the CAMS global inventory, and the changes in emissions of the main pollutants 240 241 have been assessed for different regions of the world in the first 6 months of 2020 (Doumbia et al., 2021). 242

Figure *1* shows the comparison of prior and posterior concentrations against independent observations (observations that were not used in the inversion algorithm) from the EMEP network (<u>https://emep.int/mscw/</u>, Figure S *1*) for January–July 2020. Note that prior concentrations of

ammonia result by coupling the FLEXPART SRMs with prior emissions (from VD, ECLIPSEv6 and 246 avgEENV), while posterior concentrations by coupling the SRMs with the calculated posterior 247 emissions. In Figure 1 it is evident that the most accurate reconstruction of surface concentrations 248 with respect to the EMEP observations was obtained using avgEENV as the a priori information, and 249 250 therefore the results presented hereafter are based on this setup. We performed inversions for the first half of 2020 to assess the effect of lockdown measures on ammonia emissions, as well as the situation 251 after lockdown measures were taken away (rebound period). To have a more generic view, we also 252 performed inverse modelling calculations for the first half of each year between 2016–2019 (reference 253 period). Then, we assess in impact of ammonia changes on aerosol formation (PM2.5), by feeding 254 the posterior emissions to the LMDz-OR-INCA model and calculating the production of PM2.5. 255

256 2.4 Statistical tests

To evaluate the comparisons between modelled and observed concentrations of ammonia, we used the root mean squared logarithmic error (RMSLE) defined as follows:

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$$RMSE = \sqrt{\sum_{i=1}^{N} \frac{(C_m - C_o)^2}{N}}$$
 and $RMSLE = \sqrt{\frac{1}{N} \sum_{i=1}^{N} (\log C_m - \log C_o)^2}$ Eq. 4

where C_m and C_o are the modelled and measured ammonia concentrations and N is the total number of observations. The commonly used squared Pearson correlation coefficient (r) was also used as a measure of linear correlation between two sets of data defined as:

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$$r = \frac{\sum (C_m - \overline{C_m})(C_o - \overline{C_o})}{\sqrt{(C_m - \overline{C_m})^2 (C_o - \overline{C_o})^2}} \qquad Eq. 5$$

where the distance of modelled and measured ammonia concentrations from the mean ($\overline{C_m}$ and $\overline{C_o}$) is computed. Finally, the standard deviation was adopted as a measure of the dispersion of modelled ammonia from the observations, which is the true value:

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$$\sigma = \sqrt{\frac{(c_m - c_o)^2}{N}} \qquad Eq. \ 6$$

The mean fractional bias (*MFB*) was selected as a symmetric performance indicator that gives equal weights to under- or over-estimated concentrations (minimum to maximum values range from -200% to 200%). It was used in the independent validation (validation against measurements that were excluded from the inversion, see section **3.3**) of the posterior concentrations of ammonia during the European lockdowns of 2020 and is defined as:

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$$MFB = \frac{1}{N} \frac{\sum_{i=1}^{N} (C_m - C_o)}{\sum_{i=1}^{N} (\frac{C_m + C_o}{2})} \qquad Eq. 7$$

For the same reason, the mean absolute error was computed normalized (nMAE) over the average of all the actual values (observations here), which is a widely used simple measure of error:

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$$MAE = \frac{\sum_{i=1}^{N} |c_m - c_o|}{\sum_{i=1}^{N} c_o} \qquad Eq. \ 8$$

277 **3 Results**

278 **3.1** Emission changes of ammonia due to COVID-19 restrictions over Europe

The reason behind the selected three priors used in the inversion (EGG, EC6G4 and avgEENV) 279 of ammonia is trifold; (i) they are based on the most recent estimates, (ii) they present different spatial 280 distribution, and (iii) they were derived using different methodologies. More specifically, EC6G4 is 281 282 based on the emission model GAINS (Klimont et al., 2017), while VD uses satellite observations combined with a box model (Evangeliou et al., 2021). As mentioned in the previous section, it is seen 283 284 that the most accurate representation of surface model concentrations was achieved using the avgEENV a priori, which forces posterior concentrations closer to 1×1 line, whereas the obtained 285 statistics are significantly better than using other priors (Figure 1). Therefore, the results presented 286 below have all been obtained using avgEENV as the prior emission dataset keeping results using the 287 other two priors in the Supplements. 288

The total prior emissions of ammonia over Europe for the inversion period (January – June), 289 the posterior emissions for years 2016–2019 and the posterior emissions during the lockdown year 290 2020 (January – June) are plotted in Figure 2 (the results from inversions using EC6G4 and VD prior 291 emissions are illustrated in Figure S 2 and S 3). The total prior ammonia emitted between January 292 and June in Europe were equal to 1061 kt (Figure 2a). To check whether calculated changes in 2020 293 were due to meteorology and avoid misinterpretation of our findings, inverse calculations of ammonia 294 295 were performed for the reference years 2016–2019 (January–June) using respective observations from CrIS and exactly the same set-up as the one described in section 2 (Methods). The total posterior 296 emissions of ammonia over Europe for the reference period (2016 - 2019) were estimated to be 297 1665±330 kt (4-y mean±SD) or 57% higher than the prior (Figure 2b). Finally, for January–June 298 2020 the derived emission estimates were equal to 1568 ± 172 kt (posterior \pm uncertainty) (Figure 2c). 299 This is 48% higher than the prior and 6% lower than the posterior emissions of January–June 2016– 300 2019. 301

The weekly-average evolution of prior and posterior emissions of ammonia over Europe (January to June) for 2016–2019 show a similar pattern with small year-to-year variability (Figure 2d,e), and similar temperatures (Figure S 4) thus insignificant impact from the prevailing

meteorology that would justify change in emissions due to volatilisation. The weekly posterior 305 ammonia emissions over Europe changed during the lockdown period (2020) as compared to the 306 307 reference years (Figure 2f). Satellites and national monitoring measurements of ammonia show that emissions peak in spring (March) and late-summer in Europe (Van Damme et al., 2022) 308 309 corresponding to the two main fertilization periods (Paulot et al., 2014). Ammonia abundances are however high throughout the entire spring-summer period due to agriculture associated with rising 310 temperature (Sutton et al., 2013). Ammonia posterior emissions in 2020 declined by -9.8% as 311 compared to the same period over the previous four years (2016-2019, Figure 2f). Although the 312 obtained posterior emissions for the reference period (dashed grey line and shade) are very similar to 313 those of 2020, (solid blue line and shade in Figure 2f), emissions during lockdown period in 2020 314 dropped substantially, outside of the deviation of the emissions in the reference period (Figure 2f). 315

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3.2 Uncertainty of the posterior emissions

As described in section 2.3 in more detail, we considered 4° surroundings of each spatial 317 318 element of our inversion domain from which the CrIS observations were used in the inverse problem. This means that 45 spatial elements in CrIS space were used, with six vertical levels each, for each 319 320 of the 26 temporal emission elements. To calculate the associated uncertainty of the posterior estimates, we tested two sources of uncertainty: (i) how different surroundings for each spatial 321 322 element affect posterior emissions of ammonia and (ii) how the use of different prior emissions affects posterior ammonia. We organized a series of sensitivity tests using surroundings covering 2°, 3° and 323 324 4° from each grid-cell. This selection is realistic as it was shown previously in Cao et al. (2020) for 325 the construction of prior emission error covariance matrix. For the second source of uncertainty, we performed the same inversion using not only EC6G4 and VD priors, but also adding results using two 326 more datasets for ammonia (in total four), which have 10 times higher emissions, namely EGG and 327 NE (see section 2.3). 328

The calculated absolute uncertainties are depicted in Figure 3a-c together with the relative 329 uncertainty (Figure 3d) with respect to the posterior emissions of ammonia (posterior ammonia is 330 shown in Figure 2c). The first source of uncertainty (different surroundings) slightly affects the 331 resulting posterior emissions of ammonia (Figure 3a) causing an average relative uncertainty below 332 4% in the European emissions. The second source of uncertainty (use of different priors) causes much 333 larger bias as shown in Figure 3b (average relative uncertainty 35%). The reason for this is obviously 334 the large variation of the EGG (Bouwman et al., 1997; Giglio et al., 2013; Klimont et al., 2017) and 335 NE (Evangeliou et al., 2021) prior datasets that have total emissions in the first half of 2020 of 63.5 336 and 53.3 Tg, respectively, in contrast to only 6.2 and 5.7 Tg for EC6G4 and VD. Hence, the results 337 presented here are sensitive to the use of prior emission dataset. The modelled concentrations (that 338 replaces the hypothetical true column concentration in Eq. 1) is calculated by the SRMs and the prior 339

emission and, therefore, play a key role in the comparison of the CrIS value (v^{sat}) and retrieved value 340 (v^{ret}) (see Eq. 2). Also, the modelled concentrations stand as the argument of the natural logarithm 341 weighted by the averaging kernel in logarithmic space. The linearization of this operator as suggested 342 by Sitwell and Shephard (2021) may reduce the dependency on the prior emission term, however, 343 344 this is beyond the scope of this study. Overall, the propagated (absolute and relative) uncertainties of the posterior emissions are shown in Figure 3c and d and are equal to 11% over Europe on average 345 (Figure 3). The latter shows that our calculations are robust on one hand, but dependent on the use 346 of a priori information on the other. 347

348 3.3 Validation of posterior ammonia against independent measurements

349 The optimized emissions of ammonia must be validated against independent observations, because the inversion algorithm has been designed to reduce the model-observation mismatches. 350 Here, the reduction of the posterior concentration differences from the observations from CrIS is 351 determined by the weighting that is given to the observations and, hence, such comparison depends 352 on this weighting (dependent observations). Therefore, the ideal comparison of any posterior 353 emission resulting from top-down methods would be against measurements that were not included in 354 the inversion algorithm (independent observations). Here, we used ground-based observations of 355 ammonia from all EMEP sites (https://emep.int/mscw/) for the period of our study as an independent 356 dataset for validation. All stations are illustrated in Figure S 1. 357

As we mentioned in section 2.3, we evaluated the efficiency of the inversion and the most 358 effective a priori dataset for our purpose by assessing the match between the calculated posterior 359 concentrations against all the available observations from EMEP (N=3957) for the study period 360 (Figure 1). More specifically, after it became evident that the most accurate results were obtained 361 with avgEENV as the prior (relationship closer to unity against measured ammonia), we saw an 362 immediate improvement in the statistical tests used (nRMSE, nMAE and RMSLE) when using the 363 posterior emissions to model ammonia in FLEXPART during the first half of 2020 (Figure 1 – right 364 panel). nMAE decreased from 0.80 using the prior emissions to 0.76 using the posterior ones, 365 accordingly nRMSE of the posterior concentrations dropped to 0.073 as compared to -0.069 using 366 the prior emissions, while the RMSLE decreased from 0.60 using prior emissions to 0.55 using the 367 optimized a posteriori emissions. To get a better insight on how modelled concentrations improved 368 towards ammonia observations, eight random EMEP stations were selected to show timeseries of 369 prior and posterior concentrations in the first half of 2020 (Figure S 5). Although large peaks were 370 not reproduced, all statistics were improved using the posterior emissions of ammonia. 371

372 3.4 Country-level changes due to COVID-19 restrictions

To document the emission changes of ammonia over the different European countries before, during and after the 2020 lockdowns, we report the weekly evolution of the emissions for 16 countries individually (**Figure 4**). Specifically, weekly emissions were averaged for each country based on respective country definitions that are shown in **Figure S** *6* using the avgEENV prior.

Most countries show that ammonia emissions declined or at least stayed less affected by the 377 2020 lockdowns, as compared to the same period during the reference years (2016–2019). Countries 378 with substantial decreases in the 2020 lockdown emissions were The Netherlands (-16%) and 379 Belgium (-23%), both countries with important agricultural activity, as well as Denmark (-20%), 380 Ireland (-18%) and Ukraine (-18%). Smaller changes were recorded in Spain (-2.1%), Czechia (-381 4.0%) and Italy (-6.0%) despite the intensive lockdown measures. This practically shows that 382 383 agricultural activity is insignificantly affected, even in periods of extraordinary austerity, as the last remaining primary production sector, necessary for human life. 384

We note that the largest emissions of ammonia in European countries were seen around 385 March-April (weeks 8-16) and in summer. These coincide with the fertilization periods mentioned 386 previously (Paulot et al., 2014) that control the seasonality of ammonia's emissions. In most European 387 countries, the time of the year when fertilizers can be applied is tightly regulated (Ge et al., 2020). In 388 the Netherlands and Belgium, for instance, the largest ammonia contributing region in Europe, 389 390 application of nitrogen fertilizer is only allowed from February to mid-September. This produces two peak periods, in March and late May (Figure 4). Manure application also follows stringent 391 regulations and is only allowed in the same periods depending on the type of manure (slurry or solid) 392 and the type of land (grassland or arable land) (Van Damme et al., 2022). 393

To understand and position where ammonia emissions changed during the European 394 lockdowns of 2020, we plot the difference of the posterior emissions of ammonia during the lockdown 395 period (15 March – 30 April) for the same period in Figure 5a. We calculate higher emissions of 396 ammonia during the lockdown of +115 kt as compared to the prior emissions. Note that inversion 397 algorithms aim at reducing the mismatches between modelled concentrations and observations (in 398 our case, from CrIS satellite measurements) by correcting emissions. This means that different 399 400 posterior emissions are most likely, due to errors in the prior emissions and do not indicate any impact 401 from the restriction measures.

Therefore, we demonstrate the impact of the COVID-19 lockdowns over Europe in 2020, by calculating the emission anomaly for the lockdown period from 2016–2020 (same period as the 2020 lockdowns, namely 15 March – 30 April) in **Figure 5**b. Emissions during the 2020 lockdowns

dropped by -29 kt with respect to the same period in 2016 - 2020 showing the impact of the COVID-405 19 restrictions. Maximum decreases were seen in The Netherlands and Belgium, both countries 406 comprising high emissions (Figure 5b) that also suffered heavily from the COVID-19 outbreak 407 (Bendz and Aaberge, 2020) and took strict lockdown measures. Other areas where significant changes 408 409 were calculated were Northern Italy, Switzerland and Austria, while Scandinavian countries were not affected. This agrees well with the state of the epidemic in these countries in spring 2020. While 410 North Italy was the first country outside China to suffer high mortality rates and, thus, dramatic social 411 restrictions in spring 2020, Norway, Sweden, Denmark and Finland showed total infected cases far 412 below 1% per capita, mostly suffering higher rates later in 2020 (Gordon et al., 2021). 413

As mentioned previously, ammonia emissions increase in spring (March) and late-summer in 414 Europe (Van Damme et al., 2022; Paulot et al., 2014). Therefore, calculating the difference in the 415 416 calculated emissions during the lockdown from the period before or after is practically meaningless and cannot show the lockdown impact since agricultural activity was slightly affected in 2020. For 417 this reason we quantify the delay in the evolution of the 2020 emissions by calculating emission 418 differences in the lockdowns from the period before (Lock – Prelock) for the lockdown year 2020 419 and emission differences (Lock – Prelock) for the reference years (2016 – 2019). Then, we plot their 420 spatial differences in Figure 5c. Accordingly, we do the same calculation for differences in the 421 422 rebound period (the period after the restrictions were relaxed) from the lockdown period (Rebound -Lock) in 2020 and compare them with Rebound – Lock for the reference years 2016 – 2019 (Figure 423 424 5d). We observe a clear delay in the evolution of ammonia emissions in 2020 of -77 kt (Figure 5c), while only Scandinavian countries show positive changes. Hot-spots of negative evolution were seen 425 in central Europe, mainly in the triptych of Northern Italy, Switzerland and Austria, for the reasons 426 discussed in the previous paragraph. In Poland, social measures affected the daily lives of citizens 427 significantly (Szczepańska and Pietrzyka, 2021) and might be the reason for the decreased evolution 428 of ammonia emissions (Figure 5c). After the measures were relaxed, the evolution of the emissions 429 rebounded slightly with respect to the reference period (2016 - 2019) as shown in Figure 5d. The 430 changes in ammonia during the rebound period were concentrated in countries that were affected 431 most severely from the lockdown restrictions, namely Northern Italy, Switzerland, Austria and 432 Poland. The same has been reported elsewhere for several other pollutant emissions (Davis et al., 433 2022; Jackson et al., 2022). 434

435 **4 Discussion**

436 4.1 Rising ammonia concentrations during the European lockdowns

One issue that has been overlooked is the concentrations of ammonia before, during and after 437 the 2020 lockdowns in Europe. Despite the delay in the emissions during the lockdown period in 438 2020 (section 3.4), satellite ammonia from CrIS showed an increase during the lockdowns and 439 declined after the restrictions were relaxed in almost all European countries (Figure 4). The latter 440 was reported in several studies analysing ground-based measurements. For example, Lovarelli et al. 441 (2021) concluded that contrary to other air pollutants, ammonia was not reduced, when the COVID-442 19 restrictions were introduced in North Italy. They further reported that urban and rural ammonia 443 was the highest compared to previous years during the same months for which the strictest lockdowns 444 occurred (i.e., spring 2020). Rennie et al. (2020) reported a slight decrease of ammonia in the UK, 445 while Xu et al. (2022) observed increased of ambient ammonia during the lockdowns in China. 446 Accordingly, Viatte et al. (2021) found enhanced ammonia during lockdown in Paris. Finally, in a 447 recent study, Kuttippurath et al. (2023) reported increase in ammonia during lockdowns almost 448 everywhere, with maxima in Western Europe, Eastern China, the Indian subcontinent and the Eastern 449 USA. Since atmospheric ammonia has been increasing globally due to various anthropogenic 450 activities, the European lockdowns in 2020 offer a unique opportunity to expose ammonia's sources 451 452 and address the importance of secondary PM2.5 formation.

Figure 6a depicts the modelled atmospheric lifetime of ammonia and its dependence from the 453 calculated loss-rates over Europe for the first half of 2020. Ammonia is a particularly interesting 454 455 substance due to its affinity to react with atmospheric sulfuric and nitric acids producing secondary aerosols. However, the reaction with sulfuric acid is more prevalent due to several factors. For 456 instance, sulfuric acid is a stronger acid than nitric acid, leading to more efficient reactions with 457 ammonia (higher reaction rate constant for ammonia with sulfuric than with nitric acid, thus faster 458 formation of ammonium sulfate) (Behera and Sharma, 2012). Furthermore, ammonium sulfate (final 459 product of ammonia reaction with sulfuric acid) is less volatile and more thermodynamically stable 460 than ammonium nitrate (product of the reaction with nitric acid) favoring the formation and 461 persistence of ammonium sulfate particles in the atmosphere (Walters et al., 2019). Finally, sulfuric 462 acid forms more stable clusters with ammonia, even in the presence of nitric acid (Liu et al., 2018). 463 Results from laboratory and field studies have confirmed that ammonia actually promotes the 464 nucleation of sulfuric acid in the atmosphere (Weber et al., 1999; Schobesberger et al., 2015). The 465 CLOUD (Cosmics Leaving Outdoor Droplets) experiment has also highlighted that ammonia 466 preferentially reacts with sulfuric acid in the atmosphere due to its strong acidity, ability to drive 467 stable aerosol formation, and significant nucleation enhancement effects (Kirkby et al., 2016; Wang 468

et al., 2022). Nitric acid plays a secondary role, primarily forming ammonium nitrate once sulfuricacid has reacted, but its contribution is limited by its volatility.

471 During the lockdown period over Europe, transport and industrial activities mostly stopped, and consequently the related emissions also decreased. This had an immediate effect on SO_2 and NO_x 472 (Guevara et al., 2021; Doumbia et al., 2021). Reductions of SO_2 and NO_x caused less production of 473 atmospheric sulfuric and nitric acids. The latter had a rapid twofold effect on the lifetime of ammonia: 474 (i) Less available atmospheric acids needed less ammonia for reaction towards sulfate (mainly) and 475 476 nitrate aerosols (secondarily) and therefore the loss-rates declined (Figure 6a) leading to accumulation of ammonia in its free form; (ii) ammonia originates mainly from agriculture and 477 livestock, and these activities were slightly affected during the European lockdowns increasing the 478 associated emissions (see Figure 2, though with a lower trend than previous years as discussed in 479 480 section 3.4). The rising levels of ammonia during the COVID-19 lockdowns in Europe have been confirmed by the CrIS observations (Figure 2 and 3) and have been also reported elsewhere 481 482 (Kuttippurath et al., 2023; Viatte et al., 2021; Xu et al., 2022; Lovarelli et al., 2021).

483 **4.2 Disturbance in the secondary formation of PM2.5**

The response of the restriction measures on PM2.5 mass concentrations suggests a 484 relationship that is more complex than expected and beyond road traffic intensity, at least for Europe. 485 It has been reported that there was no systematic decrease in PM2.5 concentrations during COVID-486 19 lockdowns in the USA (Archer et al., 2020; Bekbulat et al., 2021) or even in Chinese cities (Mo 487 488 et al., 2021), where primary sources are abundant and stringent lockdown measures decreased PM levels (Zhang et al., 2023). In a recent study focusing on PM2.5 measurements over 30 urban and 489 regional background European sites, Putaud et al. (2023) showed that the implementation of the 490 lockdown measures resulted in minor increases in PM2.5 mass concentration in Europe of +5±33%. 491 The latter aligns well with several regional studies focusing on the impact of lockdowns to regional 492 pollution (Querol et al., 2021; Shi et al., 2021; Viatte et al., 2021; Thunis et al., 2021; Putaud et al., 493 2021). 494

Figure 6b demonstrates observed PM2.5 from the EMEP stations (78 sites) in comparison 495 with modelled PM2.5 concentrations, both averaged for all sites. In modelled PM2.5 mass 496 497 concentrations, we have separated primary and secondary PM2.5, as secondary PM2.5 is modulated by the chemical state of the atmosphere as defined by the abundance in acids and free ammonia. We 498 499 see that observed and modelled PM2.5 concentrations are in good agreement in the first half of 2020. The good agreement between modelled and observed concentrations can be also confirmed for most 500 501 of the EMEP stations over Europe with high Pearson's coefficients, low RMSE's and low standard deviations in the Taylor plot that is demonstrated in Figure S 7. Furthermore, while secondary PM2.5 502 503 constitute around 20-30% of the total PM2.5 (Dat et al., 2024; Bressi et al., 2013; Li et al., 2023), this

proportion increased during the European lockdowns despite that reactions of ammonia to form
PM2.5 were decelerated (as seen by the declined loss in Figure 6a).

506 Leung et al. (2020) reported that the abatement of nitrate in China is buffered not only by increased oxidant build-up, but also by an increase in free ammonia concentrations through sulfate 507 508 concentration reduction, which favours ammonium nitrate formation. During COVID-19 restrictions in Europe, a significant decrease of NO_x (and SO_2) emissions occurred (Guevara et al., 2021) also 509 confirmed by Doumbia et al. (2021). Thunis et al. (2021) showed that the latter might have increased 510 the oxidative capacity of the atmosphere and, in turn, PM2.5 formation. This is the main reason why 511 PM2.5 concentrations were not decreased during the COVID-19 lockdowns in many European cities 512 (Varotsos et al., 2021; Shi et al., 2021), while the same has been reported elsewhere (Huang et al., 513 2021; Le et al., 2020; Zhang et al., 2022). 514

PM2.5 increased at areas less affected by primary emissions during the 2020 lockdown or at 515 areas where the oxidative atmosphere favours secondary aerosol formation. For instance, reductions 516 in PM2.5 were observed to be less pronounced than those in nitrogen dioxide in several regions (Patel 517 et al., 2020; Shi and Brasseur, 2020), while PM2.5 even increased in others (Wang et al., 2020; Li et 518 al., 2020). Li et al. (2020) indicated that while primary emissions dropped by 15-61% in China, daily 519 average PM2.5 concentrations were still very high (15-79 µg m-3) showing that background and 520 521 residual pollutants were important. In a similar manner, an extreme PM2.5 pollution event during the Chinese lockdown in Nanning that cause public concern was due to secondary aerosol formation (Mo 522 et al., 2021). 523

Here we aim at interpreting the mechanism below this disturbance in PM2.5 formation. As 524 explained in Seinfeld and Pandis (2000) and represented in the LMDZ-INCA model (Hauglustaine 525 et al., 2014), the neutralisation of atmospheric acids by ammonia in the atmosphere occurs through 526 ammonium sulfate formation. Sulfate $(SO_{4(s)}^{2-})$ is also produced from sulfur dioxide $(SO_{2(q)})$ gas 527 phase oxidation by the hydroxyl radical (OH). Note that the hydroxyl radical is mostly formed in the 528 529 atmosphere when ultraviolet radiation (UV) photolyses ozone in the presence of water vapour, hence 530 it is linked to humidity (Figure S 8). Sulfate production can also occur in the aqueous phase (Hoyle et al., 2016) through sulfur dioxide $(SO_{2(aq)})$ oxidation with ozone $(O_{3(aq)})$ or hydrogen peroxide 531 $(H_2O_{2(aq)})$. In both phases, a higher humidity favors sulfate formation (Figure S 8). Ammonia also 532 reacts with nitric acid $(HNO_{3(g)})$ to form ammonium nitrate $(NO_{3(s)})$ in an equilibrium reaction. In 533 that case, as SO_2 is strongly decreased due to the restrictions (Doumbia et al., 2021) and more free 534 535 ammonia accumulates (see previous section), these higher gaseous ammonia levels increase the particulate nitrate formation. This mechanism has been highlighted in China as an unintended 536 consequence of the of NO_x and SO_2 regulation on the PM2.5 levels (Lachatre et al., 2019). 537

Conducting specific experiment in the frame of the CLOUD collaboration, Wang et al. (2022) 538 reported that the NH_3 - H_2SO_4 - HNO_3 system forms particles synergistically, at rates orders of 539 magnitude faster than those the individual reactions of ammonia with sulfuric or nitric acid can give. 540 In addition to this mechanism, as the fraction of the total inorganic nitrate, as particulate $NO_{3(s)}^{-}$ 541 (instead of gaseous $HNO_{3(g)}$), increases, and as NO_x and SO_2 decrease, while NH_3 emissions remain 542 543 high, a small increase in the particulate fraction greatly slows down deposition of total inorganic $NO_{3(s)}^{-}$ and hence drives particulate $NO_{3(s)}^{-}$ to increase (Zhai et al., 2021). Thus, although NO_{x} 544 emissions decreased during COVID-19 lockdowns in Europe, secondary PM2.5 stayed unchanged, 545 because NO_x emissions reduction drives faster oxidation of NO_x and slower deposition of total 546 inorganic $NO_{3(s)}^{-}$. 547

548 **5** Conclusion

We have examined the impact of lockdown measures in Europe due to COVID-19 on the atmospheric levels and emissions of ammonia using high-resolution satellite observations combined with a dispersion model and an inverse modelling algorithm. We find that ammonia emissions in 2020 declined by -9.8% as compared to the same period in previous years (2016–2019). However, this decrease is insensitive to the meteorological conditions, as the 2020 ammonia emissions during the European lockdowns dropped outside of the deviation of the emissions in the reference period (2016–2019), while temperature, humidity and precipitation showed limited variability.

556 While ammonia emissions generally increase in spring and late summer in Europe due to 557 fertilisation, during the 2020 lockdowns, a clear delay in the evolution of the emissions of -77 kt was 558 calculated, mostly in the central European countries, which suffered by the stringent restrictions. The 559 evolution of ammonia emissions slightly rebounded after the restrictions were relaxed.

During the COVID-19 lockdowns of 2020 over Europe the atmospheric levels of ammonia were drastically increased, as confirmed by ground-based and satellite observations. The reason for this is twofold; first, the European lockdown measures reduced atmospheric emissions and levels of SO_2 and NO_x and their acidic products (H_2SO_4 and HNO_3) slowing down binding and chemical removal of ammonia (lifetimes increased), and thus accumulating free ammonia; second, the prevail of agricultural activity during the lockdowns increased ammonia emissions (though at a lower rate).

566 Surprisingly, despite all the travel, working and social restrictions that the European 567 governments took to combat the outbreak of COVID-19, ambient pollution levels did not change as 568 expected. PM2.5 levels were modulated by the chemical state of the atmosphere through secondary 569 aerosol formation. Secondary PM2.5 rather increased during the European lockdowns despite that the

precursors of H_2SO_4 and HNO_3 declined. More sulfate was produced from SO_2 and OH (gas phase) 570 or O_3 (aqueous phase), while both atmospheric reactions were favoured by higher water vapour 571 (humidity) during the lockdown period. The accumulated ammonia reacted with H_2SO_4 first 572 producing sulfate. Then, as SO_2 decreased during the European lockdowns and more free ammonia 573 accumulated, the high excess gaseous ammonia reacted with HNO₃ shifting the equilibrium reaction 574 towards conversion to particulate nitrate causing unintended increase in the PM2.5 levels. While NO_x 575 emissions declined during the European lockdowns by -33%, this reduction drives faster oxidation of 576 NO_x and slower deposition of total inorganic nitrate causing high secondary PM2.5 levels. 577

The present study gives a comprehensive analysis of the atmospheric $NH_3 - H_2SO_4 - HNO_3$ system. It also proves the complicated relationship of secondary PM2.5 formation with the abundant atmospheric gases. The general drop of emissions during the first consistent lockdowns of 2020 in Europe offers a unique opportunity to study atmospheric chemistry under extreme conditions of fast pollutant emission drop equivalent to "The Clean Air Action" of the Chinese government.

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Data availability. All data from study available from 584 this are for download 585 https://datadryad.org/stash/share/Wgbc9UiXwtMH44366myWh2bt7MQc92JKhJBz7UwQlgY (reserved doi: 10.5061/dryad.12jm63z1q). The EMEP measurements of ammonia can be downloaded 586 587 from https://ebas.nilu.no. The remote sensing data for ammonia can be retrieved from https://hpfx.collab.science.gc.ca/~mas001/satellite ext/cris/snpp/nh3/v1 6 4/ or upon request to Dr. 588 M. W. Shephard. FLEXPART version 10.4 model be downloaded 589 can from https://www.flexpart.eu/downloads. 590

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592 Supplement. The supplement related to this article in available online at.

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594 **Author contributions.** NE led the overall study, analysed the results and wrote the paper. OT 595 developed the inverse modelling algorithm and performed the inversions. MSO processed CrIS 596 ammonia on a grid. SE developed FLEXPART version 10.4 model to account for the loss of ammonia 597 from the chemistry transport model LMDz-OR-INCA. YB and DH set up and ran the chemistry 598 transport model LMDz-OR-INCA. All authors contributed to the final version of the manuscript.

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600 **Competing interests.** The authors declare no competing interests.

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Financial support. The work was supported by the COMBAT (Quantification of Global Ammonia
 Sources constrained by a Bayesian Inversion Technique) project funded by ROMFORSK – Program

- 604 for romforskning of the Research Council of Norway (Project ID: 275407, website:
- 605 <u>https://prosjektbanken.forskningsradet.no/project/FORISS/275407?Kilde=FORISS&distribution=A</u>
- 606 <u>r&chart=bar&calcType=funding&Sprak=no&sortBy=date&sortOrder=desc&resultCount=30&offse</u>
- 607 <u>t=0&ProgAkt.3=ROMFORSK-Program+for+romforskning</u>). Dr. Ondřej Tichý was supported by the
- 608 Czech Science Foundation, grant no. GA24-10400S.
- 609

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1078 FIGURE LEGENDS

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1081 Figure 1. Scatter plots of prior and posterior concentrations against independent observations

- 1082 (observations that were not included in the inversion algorithm) from the EMEP network
- 1083 (<u>https://emep.int/mscw/</u>, **Figure S 1**) from January to July 2020. Three statistical measures
- 1084 (nRMSE, nMAE and RMSLE) were used to assess the performance of each inversion using three
- 1085 different prior emission inventories for ammonia (EC6G4, VD and avgEENV).



Figure 2. (a) Total a priori emissions of ammonia over Europe for the inversion period (January – 1088 1089 June). The emissions correspond to avgEENV prior, and the total emitted amount is equal to 1061 kt. (b) Total a posteriori emissions of ammonia over Europe for the inversion period (January – June) 1090 for the reference period 2016 - 2019 (using avgEENV prior) that amount 1665 kt. (c) Total posterior 1091 emissions of ammonia over Europe for January - June 2020 (1568 kt) using the avgEENV as the 1092 prior. (d) Timeseries of weekly-average prior emissions of ammonia over Europe (January to June 1093 2020) from avgEENV prior. (e) Timeseries of weekly-average posterior emissions of ammonia over 1094 Europe for the reference years 2016–2019 (January to June) (yellow, green, cvan, magenta colors). 1095 (f) Timeseries of weekly-average posterior emissions of ammonia with the associated uncertainties 1096 1097 over Europe in 2020 resulting from inversions using the avgEENV prior are plotted together with the 1098 CrIS observations averaged over Europe (red line) and the mean ammonia emissions with the calculated standard deviations for the reference period (2016-2019). The single top number -9.8% 1099 shows percentage change in ammonia emissions during the 2020 lockdown as compared to the same 1100 1101 period in reference years, whereas two bottom ones show the corresponding changes in ammonia 1102 emissions (i) during the 2020 lockdown as compared to the period before lockdown (+13.3%), and (ii) the period after lockdown finished as compared to the lockdown period $\pm 18.3\%$, known as 1103 rebound period. 1104

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Absolute uncertainty of NH_3 emissions (mg m⁻²)

Figure 3. (a) Absolute uncertainty from use of different surrounding grid area for each spatial element 1108 of our inversion domain in the sensitivity tests; 2° to 4° grid-cells were considered resulting in a mean 1109 relative uncertainty of 4%. (b) Absolute uncertainty from use of four different prior emission 1110 estimates, namely EC6G4, VD, EGG and NE (see section 2.3). Here, a much larger uncertainty was 1111 calculated, due to the use of tenfold different prior emission datasets. (c) Propagated absolute 1112 1113 uncertainty from the different sensitivity tests, and (d) relative uncertainty with respect to the posterior emissions (Figure 2c). The average uncertainty in the inversion domain for the first half of 2020 was 1114 estimated to be 11%. 1115



1118 Figure 4. Timeseries of weekly-average posterior emissions of ammonia with the calculated uncertainties in different European countries in 2020 resulting from inversions using prior 1119 information from avgEENV plotted together with the CrIS observations averaged over Europe (red 1120 1121 line) and mean emissions with the calculated standard deviations for the reference period (2016-2019). The single top numbers show the change in ammonia emissions during the 2020 lockdowns 1122 (15 March – 30 April) as compared to the same period the years before (2016-2019), whereas the two 1123 bottom ones show the respective changes in ammonia emissions during the 2020 lockdown as 1124 1125 compared to the period before the lockdown, and after lockdown finished compared to the lockdown period (rebound period). 1126

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Figure 5. (a) Difference of posterior from prior emissions of ammonia during the European

- 1131 lockdowns of 2020 (15 March 30 April) using the avgEENV emissions as the prior. (b) Emission
- anomaly relative to the 2020 lockdowns from the 2016-2020 period (15 March 30 April).
- 1133 Difference in posterior ammonia (c) during the 2020 lockdowns (15 March 30 April, Lock) from
- the period before (1 January 14 March) and (d) after the 2020 lockdowns (1 May 31 June, Reb)
- from the period during the 2020 lockdowns (15 March 30 April, Lock) compared with the
- 1136 reference years (2016–2019).
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- 1138



Figure 6. (a) Modelled lifetime (blue) and loss-rates (red) of atmospheric ammonia averaged over 1141 Europe for January – June 2020. The lockdown period (15 March – 30 April) is shaded in yellow. 1142 Right after COVID-19 restrictions were applied, loss-rates of ammonia (shown in red) were 1143 disturbed due to reported decreases on SO_2 and NO_x (Guevara et al., 2021; Doumbia et al., 2021), 1144 precursors of sulfuric and nitric acids (with which ammonia reacts to form PM2.5) and the constant 1145 accumulation of atmospheric ammonia. This had an effect on the lifetime of ammonia (plotted in 1146 blue), which started increasing in Europe leading to further accumulation of ammonia. (b) 1147 1148 Observations of PM2.5 from the EMEP stations (78 stations) plotted against modelled PM2.5 concentrations, both averaged over Europe, from primary sources and secondary formation. It is 1149 evident that right after lockdown (yellow shade), secondary PM2.5 formation maintained high 1150 concentrations across Europe. 1151