- Ammonium nitrate particles formed in upper
- ² troposphere sourced from ground ammonia
 - during Asian monsoons

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The rise of ammonia emissions in Asia is predicted to increase 29 radiative cooling and air pollution by forming ammonium nitrate 30 particles in the lower troposphere. There is, however, a severe 31 lack of knowledge about ammonia and ammoniated aerosol par-32 ticles in the upper troposphere and their possible effects on the 33 formation of clouds. Here we employ satellite observations and 34 high-altitude aircraft measurements, combined with atmospheric 35 trajectory simulations and cloud-chamber experiments, to demon-36 strate the presence of ammonium nitrate particles and also track 37 the source of ammonia forming into the particles. We find that 38 solid ammonium nitrate particles are surprisingly ubiquitous in the 39 upper troposphere from the Eastern Mediterranean to the West-40 ern Pacific during the Asian monsoon period – even as early as 41 1997. We show that this ammonium nitrate aerosol layer is fed 42 by convection that transports large amounts of ammonia from sur-43 face sources into the upper troposphere. Impurities of ammonium 44 sulfate allow for the crystallization of ammonium nitrate even at 45 conditions of high relative humidity prevailing in the upper tropo-46 sphere. Solid ammonium nitrate particles in the upper troposphere 47 play a hitherto neglected role in ice cloud formation and the aerosol 48 indirect radiative forcing. 49

Particulate ammonium nitrate (AN, NH_4NO_3) and ammonium sulfate 50 $(AS, (NH_4)_2SO_4)$ are important species in the tropospheric aerosol system. 51 Ammonia (NH_3) , as the most abundant alkaline gas in the atmosphere, is 52 the major precursor for the formation of these aerosols [1, 2]. Emissions 53 of NH_3 are estimated to keep rising over the coming decades [3, 4, 5, 6]. 54 Model calculations indicate that this will lead to an increase of the reflected 55 solar radiation through AN particles, mainly produced in the lower part 56 of the troposphere [7, 8]. At upper tropospheric levels, the presence of NH_3 57 may strongly enhance new particle formation through stabilization of sulfuric 58 acid clusters especially at low temperatures prevalent at altitudes near the 59 tropopause [9, 10]. Also in the upper troposphere (UT), solid AS particles 60 were identified as potential ice nucleating particles affecting the formation 61 and radiative properties of cirrus clouds [11]. Yet solid AS, as well as AN 62 particles remain unobserved on a global scale in the UT. Furthermore, AN is 63 generally assumed to exist as aqueous solution under UT conditions, so that 64 it cannot promote heterogeneous ice formation [12, 13]. 65

The presence of NH_3 in the UT has recently been revealed by satellite 66 observations during the Asian monsoon [14]. During that season, polluted air 67 masses transported convectively from the boundary layer to high altitudes 68 are confined in the so-called Asian summer Monsoon Anticyclone (AMA), 69 e.g. [15]. In the AMA, pollution accumulates and is dispersed over a large 70 area of the northern hemisphere reaching longitudes of 10–140°E and lati-71 tudes from 10–40°N [16, 17, 18, 19, 20, 21]. Air, which is exported from the 72 AMA, influences the composition of the entire northern hemisphere lower-73 most stratosphere in summer [22, 23]. 74

Connected to the AMA, a layer of enhanced aerosol loading at altitudes 75 of 14–18 km was observed by satellite [24, 25, 26], and balloon-borne instru-76 ments [27]. Although little is known about the composition of the aerosol 77 particles forming this Asian Tropopause Aerosol Layer (ATAL), its particles 78 are currently believed to consist of carbonaceous and sulfate material and 79 mineral dust [28, 29, 30, 31] or nitrate [32]. The ATAL particles are often 80 connected with the presence of pollution markers in the monsoon circula-81 tion and large SO_2 emissions in South East Asia [26, 27]. Observations of 82 the composition of ATAL aerosol particles are, however, sparse. Preliminary 83 data from two single balloon flights indicate a presence of nitrate aerosol 84 particles [27]. 85

Solid AN discovered by satellite and labora tory measurements

⁸⁸ The first evidence of solid AN aerosol particles in the UT is provided by ⁸⁹ satellite observations with the CRISTA (CRyogenic Infrared Spectrometers ⁹⁰ and Telescopes for the Atmosphere) instrument inside the AMA in August ⁹¹ 1997. We identified a distinct spectral signature at 831 cm^{-1} that is only ⁹² present in spectra taken inside the AMA as the $\nu_2(\text{NO}_3^-)$ band of solid AN ⁹³ (Methods, Supplementary Fig. 1) [33]. Neither aqueous AN, nor any other ⁹⁴ aerosol or trace gas could replicate the observed spectral feature.

⁹⁵ We investigated the crystallization behaviour of aqueous AN aerosol par-⁹⁶ ticles at temperature conditions of the UT by AIDA (Aerosol Interaction and

Dynamics in the Atmosphere) aerosol and cloud chamber measurements. In 97 agreement with previous studies that extended down to a temperature of 98 -35°C [12], we did not detect the efflorescence of pure aqueous AN aerosol 99 particles. However, the formation of solid AN was observed for particles 100 composed of aqueous mixtures of AN and AS. At -50°C, already a mole frac-101 tion of 2.9 mol% AS was sufficient to initiate the crystallization of AN. The 102 crystallization rate was dependent on the ambient relative humidity (RH) 103 with respect to supercooled liquid water. On a timescale of several hours, 104 the formation of solid AN particles was even observed at RHs as high as 105 61%, corresponding to an almost ice-saturated environment at -50° C (Meth-106 ods and Supplementary Information). By demonstrating that small amounts 107 of sulfate facilitate the formation of solid AN even at high RH, these exper-108 imental results strongly support the satellite detection of solid AN particles 109 within the AMA. Further, the highly resolved laboratory infrared spectra 110 of the $\nu_2(NO_3^-)$ absorption band at low temperature provide the basis for a 111 quantitative retrieval of AN particle mass density distributions from satellite 112 and airborne observations (Methods). 113

The spatially resolved AN observations with the CRISTA satellite reveal that enhanced concentrations of AN (0.05–0.3 μ g m⁻³) are located only within the AMA (Fig. 1). These observations between 8 and 16 August 1997 indicate that an ATAL was present in the Asian monsoon UT in summer 1997, years earlier than hitherto thought [25].

The seasonal and inter-annual variability of AN concentrations in the AMA between 2002 and 2011 has been derived by analysis of spaceborne MIPAS (Michelson Interferometer for Passive Atmospheric Sounding) ob-

servations (Methods). Enhanced levels of AN appear in the second half of 122 June, increase in areal coverage and concentration until mid-end of August, 123 and reach values comparable to those derived from the CRISTA instrument 124 (Fig. 2a). During these periods, the maximum concentrations are always 125 located within the confines of the AMA, e.g. reaching as far as the Eastern 126 Mediterranean (Supplementary Figs. 5, 6). This seasonal cycle is modulated 127 by a large inter-annual variability with a clear maximum in 2008 with values 128 three to four times higher than during other years. 129

Simultaneously observed concentrations of NH_3 from a new MIPAS dataset 130 (Methods) start to increase by mid-end of June, reach a maximum by mid-131 end of July and vanish until end of August (Fig. 2b and Supplementary 132 Figs. 5, 6). This temporal evolution indicates a delay of 1-2 weeks of the AN 133 maximum with respect to NH_3 . This is consistent with the notion of advec-134 tive upward transport in the monsoon anticyclone at potential temperatures 135 above about 360 K, which is much slower than convection [34]. In spite of 136 this delay, the onset of enhanced values of AN and NH_3 inside the AMA 137 appears simultaneously by end of June. 138

AN particles observed from aircraft in upper troposphere

¹⁴¹ High-altitude aircraft flights from Kathmandu, Nepal in July and August
¹⁴² 2017 within the StratoClim project (http://www.stratoclim.org/) pro¹⁴³ vided the first opportunity to measure the distribution and composition of

aerosols and trace gases inside the Asian monsoon UT with a variety of in situ 144 and remote sounding instruments. Altitude profiles of AN and NH₃ along 145 the flight track with a greatly improved resolution compared to the satel-146 lite products have been obtained from infrared limb-imaging observations 147 with the airborne GLORIA (Gimballed Limb Observer for Radiance Imag-148 ing of the Atmosphere) instrument (Methods) [35, 36]. Figure 3 presents 149 cross-sections of AN and NH₃ along the flight track on 31 July 2017 lead-150 ing from Kathmandu (27.7°N, 85.4°E) southwest-bound to about 21°N, 79°E 151 and back. Due to the limb-sounding instrument's observational direction to 152 the right of the aircraft, the GLORIA measurements sampled air-masses of 153 different origin during the SW-bound and NE-bound flight legs. During the 154 SW-bound flight leg, we observed enhanced concentrations of AN aerosol 155 mass reaching up to $0.7 \,\mu \text{g m}^{-3}$ at around 16–17 km altitude (Fig. 3a,b). The 156 height of this layer coincides with the typical altitude of the ATAL [26] and 157 with our satellite retrievals. During the first part of the flight, a lower region 158 of enhanced AN concentrations of up to $0.5 \,\mu \mathrm{g \, m^{-3}}$ between 12 km and 15 km 159 altitude has been observed. 160

Distinct layers of enhanced values of total aerosol volume density are also 161 observed by in-situ particle size distribution measurements during the ascent 162 and descent of the aircraft on 31 July 2017 (Fig. 4a) [37, 38] (Methods). 163 These correlate very well with maxima in the nitrate mass concentration 164 profiles as measured by a flash vaporization/continuous ionization aerosol 165 mass spectrometer (ERICA-AMS) [39, 40, 41] (Methods) (Fig. 4b). Though 166 there is no exact coincidence between remote sensing and in situ measure-167 ments, the aerosol layer in the upper part of the profiles above about 15 km 168

is present in both kind of observations, as well as a second layer below. The retrieved values of nitrate aerosol mass concentration of around $0.3 \,\mu \text{g m}^{-3}$ from the infrared sounder fit to the independent in situ observations by the mass spectrometer (Supplementary Fig. 7).

Assuming the particle mass density of AN $(1.72 \,\mathrm{g \, cm^{-3}})$ for conversion of the observed aerosol volume densities to NO₃⁻ mass concentrations, the resulting maximum values at 13–14 and 16 km agree with the mass spectrometric data within the error margins. Thus, the hypothesis that a major fraction of the ATAL particle mass consists of AN is consistent with the observed aerosol volume densities at peak NO₃⁻ mass concentrations.

Additionally, the aerosol mass spectrometer detected the simultaneous presence of sulfate and nitrate in the mass spectra of individual single particles (Methods, Supplementary Fig. 9). Such common occurrence of both components was observed in more than 91% of the particles analysed between 10 and 17 km during the flight of 31 July 2017. These findings corroborate the results from the cloud-chamber observations on the crystallization behaviour of aqueous AS/AN mixtures.

¹³⁶ Ground \mathbf{NH}_3 as the source of upper tropo-¹³⁷ spheric AN particles

The region of enhanced AN at 12-15 km altitude during the first part of the flight (Fig. 3a, b) frames a zone of maximum concentrations of NH₃ (Fig. 3c, d). In this region, NH₃ volume mixing ratios reach values of 1 ppbv ¹⁹¹ at 13.5 km altitude. During the previous flight, on 29 July 2017, NH₃ mixing ¹⁹² ratios of even 1.4 ppbv have been encountered at similar altitudes. These ¹⁹³ concentrations are up to 40 times higher than the values reported previously ¹⁹⁴ [14], and are, thus, by far the highest concentrations of NH₃ ever observed ¹⁹⁵ in the UT.

High AN mass concentrations observed closely to the region of enhanced 196 mixing ratios of NH_3 might either be due to similar source regions or by 197 newly formed AN particles through gas to particle conversion within the 198 plume of elevated NH₃ concentrations. Conversion of NH₃ volume mixing 199 ratios to equivalent AN mass concentrations leads to maximum values of 200 up to $1 \,\mu g \, m^{-3}$. Thus, the GLORIA measurements of that particular day 201 show sufficiently high values of the gas phase NH₃ mixing ratios to explain 202 concentrations up to $0.7 \,\mu \mathrm{g \, m^{-3}}$ of AN. 203

We have studied the possible origin of the elevated NH₃ concentrations in 204 the UT by trajectory analyses in combination with satellite data of total col-205 umn amounts of NH_3 [42] (Methods, Supplementary Figs. 10 and 11). During 206 the days prior to our aircraft measurements, enhanced column amounts of 207 NH_3 have been observed at lower atmospheric levels in the region of NW-208 India and NE-Pakistan, a region known as a hot-spot for NH₃ emissions 209 [6, 43, 44]. Trajectory simulations suggest that the high amounts of NH_3 210 observed by the GLORIA instrument have initially been transported con-211 vectively to altitudes of 12–14 km. Subsequently, they were advected by the 212 anticyclonic upper tropospheric monsoon circulation to the location of the 213 airborne observations within a few days. These measurements prove that 214 NH₃ reaches the UT in amounts sufficient to explain the mass density of AN 215

observed. The relevant processes reducing the impact of washout of NH_3 216 during convection might involve low acidity of convective rain droplets [45], 217 or a release of NH_3 during the freezing process of cloud particles [46, 47, 48]. 218 We uncover that a considerable part of the aerosol particles in the AMA 219 consists of solid AN being formed through convective uplift of NH₃ from 220 intense surface emissions. Due to its longer lifetime, AN is subsequently 221 transported vertically and horizontally inside the AMA, thereby already in-222 fluencing a large area of the northern hemisphere. Air masses with enhanced 223 AN are not confined within the AMA but are transported either into the 224 lowermost stratosphere at mid-latitudes or into the tropical belt. Here, due 225 to dilution, the AN concentrations are no longer visible in our satellite data. 226 It is highly probable that in the UT, these solid particles act as effective ice 227 nucleating particles as has previously been demonstrated for solid AS [11]. 228

In the future, rising emissions of NH_3 will probably lead to a change 229 of AN particles also in the UT with potential consequences for the Earth's 230 radiative budget. For quantitative assessments using chemical-dynamical 231 models, a better quantification of the NH₃ surface emissions over the Indian 232 sub-continent as well as a deeper understanding of the interaction processes 233 of NH₃ with liquid water and ice during convection is needed. Furthermore, 234 it is most important to characterize the ice nucleating capacity of solid AN 235 particles. 236

$_{237}$ Methods

²³⁸ Cloud chamber observations

The AIDA aerosol and cloud chamber is a highly instrumented, 84.5 m^3 -239 sized aluminium vessel that can be operated at temperatures between +60240 and -90°C [49]. The ambient relative humidity can be controlled by evap-241 orating a specified amount of water vapour into the chamber. The number 242 concentration and size distribution of aerosol particles added to the chamber 243 are measured with a condensation particle counter (CPC, model 3010, TSI), 244 a scanning mobility particle sizer (SMPS, model 3071A, TSI), and an aero-245 dynamic particle spectrometer (APS, model 3321, TSI). Relative humidity 246 is measured in situ by tunable diode laser (TDL) absorption spectroscopy, 247 detecting a rotational-vibrational water vapour absorption line at $1.37 \,\mu m$ 248 [50]. Infrared extinction spectra of the aerosol particles are recorded in situ 249 with a FTIR spectrometer (model IFS66v, Bruker) coupled to a multiple re-250 flection cell (optical path length $166.8 \,\mathrm{m}$, wavenumber range $6000-800 \,\mathrm{cm}^{-1}$, 251 $0.5 \,\mathrm{cm}^{-1}$ resolution) [51]. With polarization-resolved in situ laser light scat-252 tering measurements, the backscattering linear depolarization ratio of the 253 aerosol particles at a scattering angle of 178° and a wavelength of 488 nm254 are determined [52]. Due to the different infrared spectral signatures of liq-255 uid and solid AN, as well as the different depolarization levels of spherical, 256 aqueous AN solution droplets and aspherical, crystalline AN particles, both 257 the infrared and depolarization measurements allow for studying the phase 258 change behaviour of the added aerosol particles. A schematic of the AIDA 259 facility and further details about its optical instrumentation can be found in 260

261 [53].

For our experiments, the AIDA chamber was held at -50°C and two dif-262 ferent relative humidities with respect to supercooled water, i.e., at 22% and 263 61% RH, in order to simulate dry and humid upper tropospheric air masses. 264 The aerosol particles were generated from aqueous solutions of 100 mol% AN, 265 $90 \mod \% AN + 10 \mod \% AS$, and $97.1 \mod \% AN + 2.9 \mod \% AS$. The bulk 266 solutions were prepared by dissolving the respective chemicals (AN, VWR) 267 Chemicals, 99%; AS, Merck, 99.5%) in high purity water (GenPure Pro UV 268 ultrapure water system, Thermo Scientific), using an overall solute concentra-269 tion of 10 wt%. Aqueous solution droplets were produced with an ultrasonic 270 nebulizer (GA 2400, SinapTec). The aerosol flow first passed a set of sil-271 ica gel diffusion dryers (Topas GmbH) to reduce the RH in the aerosol flow 272 to $\leq 3\%$ and was then injected into the AIDA chamber through a stainless 273 steel tube. The spectrum of liquid AN shown in Supplementary Fig. 1 was 274 recorded after injection of the 100 mol% AN solution into the AIDA chamber 275 at 61% RH, whereas the spectrum of the crystalline phase was recorded for 276 the 97.1 mol% AN + 2.9 mol% AS mixture at 22% RH. Pertinent details of 277 these experiments are further discussed in the Supplementary Information. 278

In order to derive mass-specific absorption coefficients of the recorded $\nu_2(NO_3^-)$ infrared absorption band of crystalline AN (Supplementary Fig. 1), which were then used as input for the aerosol mass density retrievals from the satellite observations, we computed the total mass concentration of the aerosol particles suspended in the AIDA chamber. In the first step, the volume size distribution of the aerosol particles was computed from the combination of the SMPS and APS size spectra, employing a particle density

of $1.72 \,\mathrm{g\,cm^{-3}}$ and a dynamic shape factor of 1.1 to convert the mobility 286 and aerodynamic diameters from the SMPS and APS measurements into the 287 equal-volume sphere diameter (Supplementary Fig. 3). The integrated vol-288 ume concentration was then multiplied with the particle density to yield the 289 total mass concentration of the crystallized particle ensemble. Finally, the 290 smaller mass fraction of AS, which does not contribute to the signature of 291 the $\nu_2(NO_3^-)$ absorption band at 831 cm⁻¹, was subtracted to yield the mass 292 concentration of AN only. The uncertainty for the AN mass density, mainly 293 arising from the uncertainties associated with the employed values for the 294 particle density and the dynamic shape factor, was estimated to $\pm 30\%$. 295

²⁹⁶ Remote sensors

CRISTA: The Space Shuttle experiment CRISTA (CRyogenic Infrared Spec-297 trometers and Telescopes for the Atmosphere) was flown on the Shuttle Pallet 298 Satellite in November 1994 (STS 66) and August 1997 (STS 85) [54, 55, 56]. 299 The instrument was equipped with three optical telescopes and four spec-300 trometers covering spectral ranges from 140 to $2500 \,\mathrm{cm}^{-1}$ using 26 detectors. 301 The spectral resolution was $\lambda/\Delta\lambda \approx 500$, with wavelength λ . In the spectral 302 range around $800 \,\mathrm{cm}^{-1}$, this corresponds to a resolution of $\approx 1.5 \,\mathrm{cm}^{-1}$. Dur-303 ing CRISTA-2 (STS85) the main measurement mode covered tangent heights 304 from 11 to 75 km with a vertical sampling step of 2 km and a vertical field of 305 view width of about 1.5 km. The horizontal sampling pattern of this mode 306 was 200 km along track and 650 km across track. 307

MIPAS: The limb-sounder MIPAS (Michelson Interferometer for Passive

Atmospheric Sounding) was in operation between 2002 and 2012 on board 309 the polar orbiting satellite Envisat [57]. It recorded atmospheric emission 310 spectra in the thermal infrared region between 685 and $2410 \,\mathrm{cm}^{-1}$ with spec-311 tral resolutions of $0.025 \,\mathrm{cm}^{-1}$ in the first measurement period from July 2002 312 to March 2004 and $0.0625 \,\mathrm{cm}^{-1}$ during the second period from January 2005 313 to April 2012. In nominal rearward viewing limb-scan mode, tangent al-314 titudes covered regions between 7 and 72 km with a vertical sampling of 315 3 km up to 42 km in the period 2002-2004 and 1.5 km up to 22 km altitude 316 during 2005–2012, respectively. The along-track sampling distance between 317 limb-scans was 550 km during the first and 420 km during the second mea-318 surement period. Retrievals are performed on basis of level-1B calibrated 319 radiances version 5.02/5.06 provided by ESA. 320

GLORIA: The airborne limb-imaging spectrometer GLORIA (Gimballed 321 Limb Observer for Radiance Imaging of the Atmosphere) covers the mid-322 infrared spectral range from 780 to $1400 \,\mathrm{cm}^{-1}$ [35, 36]. During the StratoClim 323 monsoon campaign, GLORIA was deployed on the Geophysica high-altitude 324 aircraft performing limb observations with an azimuth of 90° to the right 325 hand side. Geophysica is a Russian high-altitude aircraft of the type M55, 326 built and operated by Myasishchev Design Bureau, Zhukovsky, Moscow Re-327 gion. GLORIA takes one limb-image of 128 vertical \times 48 horizontal pixels 328 with a spectral resolution of $0.0625 \,\mathrm{cm}^{-1}$ within 13 s. The vertical elevation 329 coverage is 4.1°. The spectra of horizontal pixel rows are averaged for noise 330 reduction. In combination with the typical velocity of the aircraft of 700 331 to $750 \,\mathrm{km}\,\mathrm{h}^{-1}$ this corresponds to an along track sampling of around 3 km. 332 The generation of calibrated spectra from GLORIA observations has been 333

 $_{334}$ described in [58].

Identification and retrieval of ammonium nitrate (AN) from infrared limb observations

In previous infrared limb observations, the infrared ν_2 out-of-plane deforma-337 tion band of NO_3^- in nitric acid trihydrate (NAT) at 821 cm^{-1} [59, 60, 61] has 338 been used to detect and quantify NAT particles within polar stratospheric 339 clouds [62, 63, 64]. This spectral band is especially suited for detection from 340 limb-viewing sensors since it is located in one of the window regions in the 341 mid-infrared least affected by trace gas signatures. Further, compared to 342 other spectral bands of solids, the nitrate ν_2 band is so sharp that it can 343 easily be identified. 344

The $\nu_2(NO_3^-)$ - band of ammonium nitrate (AN) has frequently been assigned in laboratory spectra to wavenumbers around 831 cm⁻¹: 830 cm⁻¹ [65], 831–833 cm⁻¹ [66], 830–832 cm⁻¹ [60], 831 cm⁻¹[33]. The only field observations where the $\nu_2(NO_3^-)$ band was identified in collected samples of ambient aerosols have been reported in [67] and, tentatively, in [68].

Supplementary Figures 1 and 4 show the detection of the $\nu_2(NO_3^-)$ band of AN in infrared spectra of the spaceborne limb sounding instruments CRISTA and MIPAS, as well as observed by GLORIA during the StratoClim aircraft campaign. From difference spectra of observations at similar tangent altitudes but at different locations, a spectral band with maximum close to 831 cm⁻¹ and width of about 3 cm⁻¹ could clearly be identified - exactly at the position of the $\nu_2(NO_3^-)$ band of AN. An intensive search for an alternative

explanation of this signature, either by other kinds of aerosol particles or by 357 gas phase species was not successful. For example, AS, ammonium bisulfate 358 $(NH_4)HSO_4$ and letovicite $(NH_4)_3H(SO_4)_2$ do not show any band around 359 $830 \,\mathrm{cm^{-1}}$ [69, 70], as well as mineral dust [71] and dicarboxylic acids like 360 oxalic acid $(C_2H_2O_4)$, malonic acid $(C_3H_4O_4)$ [72] or succinic acid $(C_4H_6O_4)$ 361 [73]. Apart from NAT and AN, other nitrates, do show the sharp peak of the 362 $\nu_2(NO_3^-)$ -band in the 820-840 cm⁻¹ region of the infrared spectrum, however, 363 not at the position of AN, e.g. sodium nitrate (NaNO₃) at $836 \,\mathrm{cm}^{-1}$ [74, 75], 364 or potassium nitrate (KNO₃) at $825 \,\mathrm{cm}^{-1}$ [74]. 365

As shown in our AIDA measurements as well as in [33], an aqueous solu-366 tion of AN also has a spectral signature in that spectral region (Supplemen-367 tary Fig. 3). However, it is clearly distinct from the solid AN peak since its 368 maximum is located at $829 \,\mathrm{cm}^{-1}$ and its width is about twice the width of 369 the solid AN signature. Thus, it does not fit to our observed spectra (Supple-370 mentary Figs. 1 and 4). We, therefore, conclude that the spectral signature 371 consistently observed inside the AMA by the three different instruments is 372 caused by aerosol composed of AN in solid state. 373

The absorption coefficients as derived from the aerosol chamber experi-374 ments have been used for the retrieval of vertical profiles of AN mass density 375 from observations of CRISTA, MIPAS and GLORIA. In case of MIPAS and 376 GLORIA the same processing scheme as used for NH_3 (see below) has been 377 applied. For analysis of CRISTA observations, the retrieval employs the pro-378 cessing scheme of the airborne successor instrument already used for deriving 379 PAN in the AMA [76, 19]; the measurement density was improved by synop-380 tically interpolating the measurements of five measurement days to a single 381

point of time in the middle by using trajectory calculations. The spectral 382 window between $828 \,\mathrm{cm}^{-1}$ and $835 \,\mathrm{cm}^{-1}$ has been commonly evaluated. Un-383 certainties for AN mass concentrations are estimated to $\pm 0.03\,\mu\mathrm{g\,m^{-3}}$ (single 384 profile precision) $\pm 10\%$ (systematic instrumental/retrieval) $\pm 30\%$ (spectro-385 scopic AN absorption coefficients from the AIDA observations) for GLORIA, 386 $\pm 15\%$ (single profile precision) $\pm 15\%$ (systematic instrumental/retrieval) 387 $\pm 30\%$ (spectroscopy AN) for CRISTA, and, $\pm 0.015 \,\mu \text{g m}^{-3}$ (single profile pre-388 cision) $\pm 30\% \pm 0.01 \,\mu \text{g m}^{-3}$ (systematic instrumental/retrieval) $\pm 30\%$ (spec-389 troscopy AN) in case of MIPAS. The values of vertical resolutions are 0.8 km 390 for GLORIA, $2.2 \,\mathrm{km}$ for CRISTA, and $4.5 \,\mathrm{km}/3.5 \,\mathrm{km}$ for the retrieval from 391 the MIPAS observational period 1 and 2, respectively. 392

³⁹³ Identification and retrieval of ammonia (NH_3) from in-³⁹⁴ frared limb observations

Passive remote sounding is especially suited for detection of NH₃ since it 395 avoids problems connected to wall-effects [77, 78]. Retrievals of ammonia 396 vertical volume mixing ratio profiles from MIPAS mean radiance spectra 397 which were averaged over three months and 10° latitude $\times 10^{\circ}$ longitude have 398 been published in [14]. In order to obtain better resolution in time and space, 399 here we have performed retrievals of NH_3 from single MIPAS limb-scans. The 400 profile retrievals have been performed as described in [14]. The inversion 401 scheme is a nonlinear least squares fitting in a global-fit approach on 1 km 402 spaced altitude levels [79] regularized by a first-order smoothing operator [80]. 403 The selected spectral windows with NH_3 signatures are 965.1-965.6 cm⁻¹ 404

and 966.6–967.5 cm⁻¹ for the first MIPAS period 965.125–965.625 cm⁻¹, and 966.625–967.5 cm⁻¹ for the second period. Resulting profiles of NH₃ volume mixing ratio are characterized by a vertical resolution of 4–4.5 km/3.5-4 km (1st/2nd period). Uncertainties are estimated according to [14] to ± 5 pptv (single profile precision) and $\pm 15\%$ (accuracy).

As in the case of MIPAS, inversion of vertical profiles of NH₃ volume 410 mixing ratios from GLORIA calibrated limb-radiances has been performed by 411 application of a constrained nonlinear least-squares fitting scheme, whereby 412 measurements from all infrared spectra of one GLORIA image are used [81, 413 82]. The retrievals employ an altitude grid spacing of 0.25 km. For the 414 spectral fit, the same spectral windows and inversion procedure as in the 415 case of the MIPAS second measurement period have been used (see above). 416 Supplementary Fig. 8 shows the spectral evidence for NH_3 from GLORIA 417 observations taken at a tangent altitude of 13.7 km during the flight on 31 418 July 2017 at 4:14 UTC where the retrieval resulted in a value of 920 pptv at 419 the 13.75 km altitude level. Here the spectral emission lines of NH_3 are clearly 420 identified by comparing the simulations including NH₃ to the simulations 421 without consideration of NH₃. This proves unambiguously the presence of 422 NH_3 with concentrations near 1 ppbv in the UT. The vertical resolution of 423 retrieved profiles is about 0.8 km. The vmr profile uncertainties are estimated 424 based on various parameter uncertainties as described in [81, 82]. Thev 425 amount to $\pm 8 \text{ pptv}$ (single profile precision) $\pm 20\%$ (accuracy). 426

427 In situ sensors on board the Geophysica

ERICA (ERc Instrument for the chemical Composition of Aerosols): The 428 mass spectrometer ERICA is a newly developed in situ instrument com-429 bining a laser ablation aerosol mass spectrometer (ERICA-LAMS) similar 430 to [83, 84] with a flash vaporization/electron impact ionization mass spec-431 trometer (ERICA-AMS) based on the Aerodyne Aerosol Mass Spectrometer 432 (AMS) principle [39]. For a quantitative comparison of the aerosol chemical 433 composition, mostly data from the AMS component of ERICA was consid-434 ered in this study. Here small ensembles of the sampled aerosol particles 435 with size diameters between 60 nm and approximately $2 \,\mu m$ are vaporized at 436 600°C followed by electron impact ionization of the vapour [39]. The pos-437 itive ions are detected by a Time-Of-Flight Mass Spectrometer (TOFMS) 438 from which the particle chemical composition can be inferred [40]. While the 439 reproducibility in the laboratory is better than 5% from calibration to cali-440 bration, the uncertainty for the aircraft deployment is estimated to $\pm 30\%$. 441 Single particle aerosol composition was determined by the LAMS compo-442 nent of ERICA covering a size range from $150 \,\mathrm{nm}$ to $2 \,\mu\mathrm{m}$. For each ablated 443 and analysed aerosol particle, the positive and negative ion mass spectra are 444 concurrently recorded by two TOFMSs. This way it is possible to unambigu-445 ously identify individual particles which contain both sulfate and nitrate, as 446 can be seen from the example mass spectra in Supplementary Fig. 9. The 447 ERICA-LAMS and ERICA-AMS instrument combination itself and the cor-448 responding sampling inlet system were custom designed similar to [41] for 449 operation on the high altitude research aircraft Geophysica. 450

UHSAS/COPAS: The ambient aerosol volume density was obtained from 451 particle size distribution measurements by means of an in-house modified UH-452 SAS optical particle counter [37] (mounted underneath the aircraft's wing) 453 and the high-altitude COPAS condensation particle counter [38]. The UH-454 SAS covers a particle size diameter range from $65 \,\mathrm{nm}$ to $1 \,\mu\mathrm{m}$ and has been 455 modified for operation under stratospheric conditions. One more size bin 456 extending from 10 nm to 65 nm was derived as composite from the COPAS 457 and the UHSAS data. 458

⁴⁵⁹ MAS: The Multiwavelength Aerosol Scatterometer obtained in situ mea-⁴⁶⁰ surements of the aerosol backscatter ratio at 532 nm [85].

⁴⁶¹ Trajectory calculations

Two different models for estimation of convective events along backward
trajectories starting at the GLORIA measurement locations have been used,
TRACZILLA and ATLAS.

The TRACZILLA diffusive back-trajectories [86] have been calculated 465 based on ERA5 reanalysis data $(0.25^{\circ} \times 0.25^{\circ})$ horizontal resolution, 137 ver-466 tical levels, hourly temporal resolution) using kinematic vertical winds. Clus-467 ters of 1000 parcels are released along each tangent point with a time resolu-468 tion of 1 hour. The probability of convective influence along the trajectories is 469 estimated from the high-frequency MSG1 and Himawari geostationary satel-470 lites observations of Brightness Temperature (BT). Both satellite images are 471 used, jointed at 90°E longitude, to cover the whole AMA region extent. The 472 trajectory is assumed to encounter a convective event when advected below 473

a deep convective cloud, the top pressure of which is estimated from the BT,
following the same approach as presented in [87].

Trajectories from the ATLAS model [88] are driven by ECMWF ERA 476 Interim reanalysis data $(0.75^{\circ} \times 0.75^{\circ})$ horizontal resolution, 6 h temporal res-477 olution) and use a log-pressure coordinate (kinematic trajectories). The tra-478 jectory model includes a detailed stochastic parameterization of convective 479 transport driven by ERA Interim convective mass fluxes and detrainment 480 rates. At every measurement location of GLORIA, 1000 backward ensem-481 ble trajectories are started, which take different paths due to the stochastic 482 nature of the convective transport scheme. 483

Supplementary Figs. 10 and 11 show backward trajectories starting at the 484 GLORIA measurement locations on 31 Jul 2017 calculated by TRACZILLA 485 and ATLAS, respectively. For the SW-bound flight leg, exemplary 30 day 486 non-convective backward trajectories are shown, starting at locations with an 487 NH_3 concentrations enhancement larger than 0.4 ppbv. For the NE-bound 488 leg, exemplary trajectories starting in the same altitude range (12-14 km)489 are shown for comparison. The trajectories show that the history of the 490 airmasses corresponding to enhanced NH₃ concentrations is different from the 491 one where no NH₃ has been detected. Further, consistently in both models, 492 maxima in the convective event density along the trajectories with enhanced 493 NH_3 , as indicated by white contour lines, are located over the Himalayas 494 and the Tibetan plateau as well as over NW-India and NE-Pakistan. The 495 latter region is clearly correlated with a region of high column amounts as 496 observed by the IASI satellite instruments during the days before the aircraft 497 observations. Thus, we conclude that it is very likely that the high NH_3 498

- ⁴⁹⁹ concentrations observed by GLORIA have been injected into the UT over
- $_{500}$ NW-India and NE-Pakistan by strong convection in a region where, near the
- $_{501}$ ground, high NH₃ concentrations are prevalent.

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4556 Author contributions

M.H. conducted the analysis of MIPAS and GLORIA data, produced Figs. 2– 859 4, and wrote the paper with all authors contributing. J.U. conducted the 860 analysis of CRISTA data, helped with analysis of GLORIA data and pro-861 duced Fig. 1. A.D., S.M., A.B., O.A., A.H., and S.Bo. performed and 862 analysed the aircraft in situ measurements of ERICA. C.M. and R.We. pre-863 pared the analyses for Fig. 4a, O.A. for Fig. 4b. C.M. and R.We. conducted 864 the measurements and data analyses for UHSAS and COPAS, respectively. 865 R.W., H.S., O.M., and T.L. conceived and performed the AIDA experiments 866 and contributed to their interpretation. R.S. discovered the AN emission fea-867 ture in CRISTA data for the first time. M.Ri. conceived the reanalysis of the 868 CRISTA data wrt. signals of the ATAL. G.S. contributed to the analysis of 869 MIPAS data. B.L. and S.Bu. conducted the TRACZILLA trajectory calcu-870 lations. F.C. performed the MAS aircraft observations and conducted their 871 analysis. F.F.V. conducted the GLORIA aircraft observations. S.J. analysed 872 the trajectory datasets in combination with the IASI measurements. S.J. and 873 L.K. helped with analysis of GLORIA data. P.P. contributed to the CRISTA 874 and GLORIA data analysis. T.N. helped with performing the GLORIA ob-875 servations. R.M. contributed to the interpretation of the observations. J.O. 876 contributed to the interpretation of spectroscopic issues of AN and NH₃. F.S. 877 and M.Re. defined the flight region, the general approach, general flight pat-878 terns and instrumentation of the aircraft campaign and organized it. I.W. 879 developed the ATLAS model and provided the trajectory calculations from 880 it, with contributions from M.Re. 881

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Competing interests

⁸⁸⁶ The authors declare no competing interests.

887 Data availability

The datasets generated and analysed during the current study are available from the corresponding author on request. Additionally, the CRISTA dataset of ammonium nitrate is publicly available at: https://datapub. fz-juelich.de/slcs/crista/an/. MIPAS and GLORIA data of ammonia and ammonium nitrate as well as trajectory information and AIDA spectra can be downloaded from the KITopen archive at: https://doi.org/?. IASI data on NH₃ are available at http://iasi.aeris-data.fr/NH3/.



Fig. 1: Ammonium nitrate observed by CRISTA in the UT 1997. Cross sections of synoptically interpolated AN mass concentrations, **a**, at 380 K potential temperature (16–17 km altitude), as well as, **b**, at 30°N derived from CRISTA observations between 8 and 16 August 1997. The cyan line in panel **a** shows the 4.1 PVU potential vorticity contour line, a rough measure for the extent of the AMA core, white hexagons indicate missing data and the black line the position of the cross-section of panel **b**. Red lines in panel **b** show potential temperature (K) and blue dots the location of the lapse rate tropopause.



Fig. 2: Time series of AN and NH₃ in the AMA. Time series of total mass of \mathbf{a} , AN, and \mathbf{b} , NH₃ as derived from MIPAS observations for the region 10°E–110°E, 20°N–40°N and 13–17 km altitude. CRISTA observations of AN are included in \mathbf{a} as a black cross. Error bars are Gaussian combinations of the estimated retrieval errors (see Methods) and the standard deviations of the mean values.



Fig. 3: Airborne limb-imaging observations of AN and NH₃ in the UT above India during the 2017 Asian monsoon season. a,b, altitude-time horizontal projections of AN mass densities, and c,d, NH₃ volume mixing ratios derived from GLORIA measurements during the Geophysica flight on 31 July 2017. Horizontal projection at tangent points (a,c) and vertical projection (b,d). Vertical resolution and estimated uncertainty (precision, accuracy): AN: 0.8 km, $\pm 0.03 \,\mu \text{g m}^{-3} \pm 30\%$; NH₃: 0.8 km, $\pm 8 \text{ pptv} \pm 20\%$; . Red lines: aircraft position (a,c) and altitude (b,d), light blue lines (b,d): lapse rate tropopause.



Fig. 4: Airborne in-situ aerosol observations in the Asian mon-soon UT on 31 July 2017. a, UHSAS and COPAS derived aerosol volume density during ascent (dashed) and descent (dotted) at Kathmandu.
b, ERICA-AMS particulate NO₃⁻ mass concentrations. The values are averages in vertical bins of 0.25 km. For the nitrate content of ERICA-AMS, the uncertainty is estimated to 30%. The estimated uncertainty of the UH-SAS and COPAS data is below 30%. Light blue lines indicate the lapse rate tropopause.

Ammonium nitrate particles formed in upper troposphere sourced from ground ammonia during Asian monsoons

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Supplementary discussion of the AIDA cloud chamber observations

Supplementary Figure 1 shows the infrared spectrum of supercooled liquid AN solution droplets in the regime of the $\nu_2(NO_3^-)$ absorption band (blue line), which was recorded after nebulization of the 100 mol% AN solution, drying of the aerosol flow to RH < 3%, and injection into the AIDA chamber at 61% RH and 223 K. The complete mid-infrared spectrum (shown as trace A in Supplementary Fig. 2, top panel: 6000–800 cm⁻¹, lower panel: expanded view from 1600 to 800 cm⁻¹) proves the liquid phase of the injected aerosol particles by showing the characteristic infrared signature of liquid water centred at about 3500 cm⁻¹ [1, 2]. Furthermore, the recorded depolarization ratio of the aerosol particles at 488 nm and 178° scattering angle (see Methods) was less than 1%, as indicative of spherical, aqueous solution droplets. The strategy of this experiment was similar to the one adopted by [1], namely to confirm the crystallization of AN by exploiting the hysteresis effect in the deliquescence and efflorescence behaviour. According to the Aerosol Inorganics Model (AIM), the deliquescence relative humidity (DRH) of AN at 223 K is 97% [3]. If the AN solution droplets had fully crystallized at \leq 3% RH prior to the injection into the AIDA vessel, the particles should not have taken up any liquid water in the chamber, given that the prevalent RH was well below the DRH. The AN particles were kept for an observation period of almost four hours in the AIDA chamber at 223 K and 61% RH, without showing any increase in the depolarization ratio or change in the infrared spectral signature. Our experiment thus corroborates previous studies that have demonstrated the strong inhibition of efflorescence for pure AN solution droplets [4, 5, 1]. The infrared spectrum of liquid AN (Supplementary Fig. 1), however, clearly fails to reproduce the measurement by the CRISTA limb-sounder.

Whereas pure AN solution droplets have a strong tendency to remain in the supercooled liquid state, the crystallization of AN is facilitated by impurities in the aqueous solutions [5], the presence of insoluble inclusions like mineral dust [6], and/or the admixture of compounds that easily crystallize and thereby serve as heterogeneous nuclei to promote the formation of solid AN [1]. Additionally, contact efflorescence initiated by externally located solid particles has been established as a pathway for the crystallization of supersaturated aqueous AN particles [7]. Internal mixtures of ammonium, nitrate, and sulfate are considered as a common aerosol particle type in the troposphere, and their crystallization behaviour has been thoroughly investigated in aerosol flow tube studies conducted at room temperature [1, 8]. For example, it was shown that the addition of just 10 mol% ammonium sulfate (AS) promoted the crystallization of AN in mixed AN/AS solution droplets [1]. We were able to reproduce this finding in our AIDA experiments. When using a 90 mol% AN + 10 mol% AS bulk solution for aerosol generation, the infrared spectrum of the injected particles (Supplementary Fig. 2, trace C) was devoid of any liquid water signature. Instead, the spectral changes in the band system between 1500 and 1300 cm⁻¹ as well as the appearance of the sharp peak at 831 cm⁻¹ indicated the presence of crystalline AN. In accordance with the prevalence of crystalline particles, the back-scattering linear depolarization ratio adopted a value of 17%. The sulfate admixture was seen by the absorption band at about 1100 cm⁻¹, which can be due to pure crystalline AS or the double salts 2AN·AS and 3AN·AS [9, 1].

We also investigated whether smaller admixtures of AS were still sufficient to trigger the crystallization of mixed AN/AS solution droplets. As shown by spectrum B in Supplementary Fig. 2, we were able to produce crystalline AN particles even for an AS mole fraction as low as 2.9 mol%. For this mixture, however, the AIDA chamber had to be kept at a lower RH (22%) in order to observe the instantaneous formation of crystalline particles upon aerosol injection. When the AIDA chamber was kept at 61% RH, the injected aerosol particles generated from the 97.1 mol% AN + 2.9 mol% AS bulk solution initially showed the infrared signature representative of supercooled solution droplets with a depolarization ratio < 1%. In contrast to the experiment with pure AN solution droplets, the mixed AN/AS particles gradually crystallized while suspended in the AIDA chamber at 223 K and 61% RH for four hours, as evidenced by an increase of the depolarization ratio from < 1 to 5% and a concomitant change of the infrared spectral signature. Spectrum D in Supplementary Fig. 2 illustrates the difference in the spectral signature after the observation time of four hours, demonstrating the formation of the crystalline AN phase. From the infrared spectra, we estimated that about 25% of the aerosol population had crystallized within four hours of observation. Supplementary Figure 1 demonstrates that the $\nu_2(NO_3^-)$ band of crystalline AN recorded in the AIDA chamber (orange line, enlarged view of spectrum B from Supplementary Fig. 2) shows a very good match with the CRISTA observation. To the best of our knowledge, the AIDA spectrum represents the first infrared measurement of suspended, crystalline AN particles at temperature conditions of the UT. The crystallization experiments underline that small amounts of sulfate impurities are sufficient to enable the formation of the

solid phase of AN.

Trace E in Supplementary Fig. 2 (bottom) shows the previously recorded infrared spectrum of crystalline AN/AS particles with 10 mol% AS at room temperature [1]. In comparison with the corresponding AIDA spectrum recorded at 223 K (trace C), there are small changes in the fine structure of the band system between 1500 and 1300 cm⁻¹ with contributions from the $\nu_4(NH_4^+)$ and $\nu_3(NO_3^-)$ modes. These spectral changes could be indicative of a thermal phase transition, given that five different phases of crystalline AN have been reported at ambient pressure [10, 11]. In particular, the phase transition from phase IV to V, which is supposed to occur at 255 K, could account for the temperature-dependent changes in the spectral habitus at 1500 –1300 cm⁻¹. In contrast to this wavenumber regime, previous infrared studies with thin films of AN have shown that the peak position of the $\nu_2(NO_3^-)$ mode is rather insensitive to the transition between phase IV and V [12, 13, 14]. This is corroborated by our findings shown in the left part of Supplementary Fig. 3, where we show the spectral regime of the ν_2 band of AN particles in the liquid and solid phase recorded during the AIDA experiments at 223 K and the flow-tube studies performed at 293 K by [1]. Obviously, the spectral habitus of the $\nu_2(NO_3^-)$ mode does not change with temperature, both for the liquid and the solid phase.

As outlined in the Methods section, the infrared optical depths recorded could be quantitatively scaled by the mass concentration of the suspended aerosol particles, thereby yielding mass specific absorption coefficients of the $\nu_2(NO_3^-)$ mode of AN. A representative volume size distribution of the aerosol particles from the crystallization experiment with 97.1 mol% AN + 2.9 mol% AS at 223 K and 22% RH, which was used to derive the aerosol mass concentration, is shown in the right part of Supplementary Fig. 3.

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Supplementary Figures 1–11



Supplementary Figure 1: **Infrared spectra of AN.** The $\nu_2(NO_3^-)$ infrared band of AN as observed in laboratory (AIDA) and inside the AMA by the CRISTA satellite instrument [15]. The CRISTA signal with 1.5 cm^{-1} spectral resolution has been derived from all spectra measured during the flight in August 1997 within a tangent altitude range of 16-18 km, $20^\circ-50^\circ\text{N}$ and $30^\circ-180^\circ\text{E}$ (see Methods). The laboratory spectra of optical depth (liquid AN: blue, solid AN: orange) and the satellite radiances (green) are scaled and offset to facilitate their comparison.



Supplementary Figure 2: **AIDA laboratory spectra of liquid and solid AN particles.** Mid-infrared spectra of aerosol particles composed of AN and mixtures of AN and AS. Top panel: Spectral regime from 6000 to 800 cm⁻¹ (with CO₂ absorption regime between 2450 and 2150 cm⁻¹ excluded). Bottom panel: Expanded view from 1600 to 800 cm⁻¹. Spectra A to D were recorded at 223 K in the AIDA chamber, whereas spectrum E, recorded at 293 K, was taken from the literature [1]. All spectra were scaled and offset to facilitate their comparison. See Supplementary discussion for details.



Supplementary Figure 3: **Comparison of laboratory spectra of liquid and solid AN and volume size distribution.** Left panel: Comparison of the $\nu_2(NO_3^-)$ infrared absorption mode of liquid and solid AN particles, recorded at 223 K during the present AIDA experiments (expanded view of spectra A and B from Supplementary Fig. 2) and at 293 K during previous aerosol flow-tube experiments ("SchlMar05", digitized graphs from the inserts in Figs. 2 and 5 of [1]). All spectra were scaled and offset to facilitate their comparison. Right panel: Volume-size distribution of crystalline aerosol particles generated from the 97.1 mol% AN + 2.9 mol% AS bulk solution after injection into the AIDA chamber at 223 K and 22% RH.



Supplementary Figure 4: **Detection of the** $\nu_2(NO_3^-)$ **infrared band of ammonium nitrate in infrared limb spectra.** Measured spectra with (orange) and without (blue) the spectral band of AN and their difference (green). The spectra have been taken from observations obtained at different locations with high and low concentrations of AN, respectively. **a**,**d**, CRISTA data are averaged spectra selected over the whole Space Shuttle flight in August 1997 within a tangent altitude range of 16–18 km, 20– 50° latitude and 30–180° longitude. **b**,**e**, MIPAS spectra between 15 km and 16 km tangent altitude from 8 August 2008, 7:15:30 UTC (orange) and 3:52:17 UTC (blue). **c**,**f**, mean GLORIA spectra from flight 31 July 2017 at tangent altitudes between 16.5 and 16.75 km at 4:15–4:21 UTC (orange) and 3:55–3:59 UTC (blue).



Supplementary Figure 5: Ammonium nitrate and ammonia observed by MIPAS in the UT during the Asian monsoon 2008. Monthly mean AN mass concentrations (**a**–**d**) and NH₃ volume mixing ratios (**e**–**h**) at 16 km altitude derived from MIPAS observations from June to September 2008. Data are monthly averages of single retrieved profiles within a 10° latitude \times 10° longitude grid.



Supplementary Figure 6: Ammonium nitrate and ammonia observed by MIPAS in the UT during the Asian monsoon 2008. Monthly mean AN mass concentrations (**a**–**d**) and NH₃ volume mixing ratios (**e**–**h**) at 30–40°N derived from MIPAS observations from June to September 2008. Data are monthly averages of single profiles within a 10° latitude \times 10° longitude grid.



Supplementary Figure 7: **Airborne aerosol observations in the Asian monsoon UT on 31 July 2017. a**, UHSAS and COPAS derived aerosol volume density during ascent (dashed) and descent (dotted). **b**, ERICA-AMS particulate NO₃⁻⁻ mass concentrations. **c**, MAS backscatter ratios (Bsr) at 532 nm during ascent (dashed) and descent (dotted). **d**, GLO-RIA NO₃⁻⁻ mass concentration profiles from AN retrievals. **a**–**c**, have been obtained during ascent and descent at Kathmandu. The values are averages in vertical bins of 0.25 km. GLORIA data are mean profiles averaged during the flight between 3:50–3:55 UTC (orange) and between 4:05-4:10 UTC (green). Shaded areas indicate 25 and 75 percentiles for MAS and standard deviations in case of the other datasets. For the nitrate content of ERICA-AMS, the uncertainty is estimated to 30%. The estimated uncertainty of the UHSAS and COPAS data is below 30%. The uncertainty of MAS Bsr is estimated to 0.05. The estimated accuracy of the GLORIA data is 30%.



Supplementary Figure 8: **Detection of NH**₃ **emission lines in GLO-RIA limb infrared spectra.** Example of a GLORIA spectrum at a tangent height of 13.7 km on 31 July 2017, 4:15 UTC (blue) in comparison to the spectral fit including NH₃ spectral lines (orange) and a calculation without consideration of NH₃ (green). The two strongest lines belong to CO_2 .



Supplementary Figure 9: **Particles containing nitrate and sulfate.** Average single particle mass spectra taken by ERICA-LAMS during the whole StratoClim campaign with simultaneously present spectral lines of sulfate and nitrate ions indicating that the individual particles of this category contain both, sulfate and nitrate components.



Supplementary Figure 10: **The origin of high NH**₃ **concentrations in the UT.** Exemplary subsets of backward TRACZILLA trajectories starting at GLORIA observations with enhanced NH₃ concentrations (SW-bound flight leg) and at no NH₃ enhancement (NE-bound leg) on 31 July 2017. The trajectories are colour-coded according to time. White contour lines highlight regions where the density of convective events, for the NH₃enhancement trajectories release, is larger than 0.5%. This quantity is computed as the ratio, on a 1° latitude × 1° longitude grid, of the number of encountered convective events with respect to the total number of released trajectories. Mean IASI column amounts of NH₃ for the time period 26–30 July 2017 are indicated by the colours of the left colour bar. IASI measurements less than 5% of the maximum value are not plotted. The aircraft flight path is shown as a green line.



Supplementary Figure 11: The origin of high NH_3 concentrations in the UT. Same as Supplementary Fig. 10, but with trajectory data from the ATLAS model.