A molecular perspective for global modeling of upper atmospheric NH₃ from freezing clouds

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Ammonia plays a key role in the neutralization of atmospheric acids such as sulfate and nitrates. A few in situ observations have supported the theory that gas-phase NH₃ concentrations should decrease sharply with altitude and be extremely low in the upper troposphere and lower stratosphere (UTLS). This theory, however, seems inconsistent with recent satellite measurements and is not supported by the aircraft data showing highly or fully neutralized sulfate aerosol particles by ammonium in the UTLS in many parts of the world. Here we reveal the contributions of deep convective clouds to NH₃ in the UTLS by using integrated cross-scale modeling, which includes molecular dynamic simulations, a global chemistry transport model, and satellite and aircraft measurements. We show that the NH₃ dissolved in liquid cloud droplets is prone to being released into the UTLS upon freezing during deep convection. Because NH₃ emission is not regulated in most countries and its future increase is likely persistent from agricultural growth and the warmer climate, the effect of NH₃ on composition and phase of aerosol particles in the UTLS can be significant, which in turn can affect cirrus cloud formation, radiation, and the budgets of NOₓ and O₃.

Ammonia, the most abundant gas-phase alkaline species in the atmosphere, contributes considerably to the neutralization of atmospheric acids produced by the oxidation of sulfur and nitrogen oxides with major anthropogenic sources (1, 2). NH₃ mainly comes from fertilizers and animals, and other sources include industry, fossil fuels, oceans, crops, soils, and biomass burning (3, 4). Primarily emitted from land surfaces, ammonia has a lifetime of several hours due to its high affinity to water, which facilitates its effective removal via atmospheric scavenging and its incorporation in aequous and acid particles (1, 2). Hence, gas-phase NH₃ concentrations are expected to decrease sharply with altitude and to be extremely low in the upper troposphere and lower stratosphere (UTLS), especially over the ocean (1, 5–7). This expectation is supported by several in situ observations (1, 5–7). However, in situ measurements of NH₃, a sticky, semivolatile compound with ambient concentrations that vary by several orders of magnitude, may have large uncertainties in the UTLS (8). Recent satellite measurements give different insights and show significant amounts of NH₃ (above 15 pptv) in the UTLS over the subtropical regions of the southeastern Asian continent (28–30), and limb-sounding techniques have been proposed to measure the vertically resolved profiles of NH₃ in the UTLS (28–30). Before 2016, no data from in situ measurements and limb-sounding remote-sensing data (9) showed ammonia to be present at altitudes above 8 km. Höpner et al. (9) presented the first evidence of the presence of ammonia in the UTLS.

Tropical deep convection is a critical mechanism for the transfer of surface air masses and insoluble trace gases (such as CO) from the lower to the upper atmosphere with many important implications for atmospheric chemistry (12, 13). Liu and Zipser (14) found that the deepest convection was located over South America and Africa. Penetration of these trace gases to high altitudes (16–18 km) has also been observed in northern Australia (15, 16) and over tropical cyclones (10, 17, 18). Southeast Asia is one of the tropical regions with frequent deep convection associated with the Asian summer monsoon anticyclone (19). Major source regions of NH₃ are located in southeast China and northern India (9, 20, 21). Very few measurements of NH₃ have been made in the middle and upper atmosphere (9). Before 2008, measurements of ammonia were almost exclusively based on in situ technologies, and most of them focused on ground-level concentrations with few observations above the boundary layer (22). By contrast, satellite measurements of infrared spectra have been used in the past decade to derive the amounts of NH₃ in the atmosphere (3, 23–27), and limb-sounding techniques have been proposed to measure the vertically resolved profiles of NH₃ in the UTLS (28–30). Before 2016, no data from in situ measurements and limb-sounding remote-sensing data (9) showed ammonia to be present at altitudes above 8 km. Höpner et al. (9) presented the first evidence of the presence of ammonia in the UTLS.

The retentions of gases upon freezing of water from available studies are conflicting. Using molecular dynamic simulations, we have revealed that the retention efficiency of NH₃ upon freezing cloud is close to 0 rather than 1 during deep convection. Our results further provide theoretical framework to explain the first-ever satellite-observed high concentration of NH₃ in the upper troposphere and lower stratosphere (UTLS). As NH₃ emission continues to increase, this study calls upon the emergent need to assess the role of NH₃ in UTLS chemistry, especially in the deep convective regions. Methodologically, the molecular dynamics simulation is demonstrated as a tool for improving parameterization of interactions between trace gases and cloud (including ice) particles in global atmospheric models.

Significance

The retentions of gases upon freezing of water from available studies are conflicting. Using molecular dynamic simulations, we have revealed that the retention efficiency of NH₃ upon freezing cloud is close to 0 rather than 1 during deep convection. Our results further provide theoretical framework to explain the first-ever satellite-observed high concentration of NH₃ in the upper troposphere and lower stratosphere (UTLS). As NH₃ emission continues to increase, this study calls upon the emergent need to assess the role of NH₃ in UTLS chemistry, especially in the deep convective regions. Methodologically, the molecular dynamics simulation is demonstrated as a tool for improving parameterization of interactions between trace gases and cloud (including ice) particles in global atmospheric models.


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above 10 km over the Asian summer monsoon system and compared against several global modeling results with either low horizontal resolutions (in range of ~5–10') or low vertical resolutions (with total number of vertical layers ranging from 9 to 26). Höpfner et al. (9) concluded that no literature regarding locally resolved model results of NH3 during the monsoon period over Asia is available to compare with their novel observations.

The scavenging of NH3 by monsoon convection is hypothesized to incompletely deplete atmospheric NH3, thereby leading to the direct transport of NH3 from the boundary layer to the UTLS via convection (9). Our study seeks another pathway for the vertical transport of NH3 via the interactions between NH3 and clouds. Ammonia gas has a large proton affinity and can therefore serve as a base to neutralize liquid particles (either aerosols or cloud droplets) that are often weakly acidic. The ammonia gas can dissolve in the liquid cloud. During the upward movements driven by deep convection, the liquid droplets of clouds are first supercooled and then frozen in the UTLS before they precipitate with atmospheric downdrafts. During this process of liquid-to-ice conversion, some gases in the liquid cloud droplets might be released into the atmosphere upon freezing; the relative amount of the gas that is retained in this conversion process is characterized by the retention coefficient of that gas (31). Past studies have derived the retention coefficients of different gas species through several laboratory experiments, aircraft measurements, and theoretical calculations but showed controversial results. SO2 is one of the most studied species, and its retention coefficients vary from 0.02 (13, 32), to 0.03–0.14 (33) to 0.25–0.9 (34, 35). For H2O2, the retention coefficient is reported as being between 0.64 (31) and 1 (34). As a highly soluble trace gas, HNO3 has been reported close to 1 (31, 36).

The retention coefficients of some other species have also been reported: 1 for H2SO4 (36); 0 for O3, NO, NO2, NOx, N2O, and CO2; 0.02 for OH, CH3O2, and CH2OOH; and 0.65 for HO2, HNO2, HNO3, HCHO, HCOOH, and CH2COOH (37). These studies show that the direct measurement of retention efficiency can be affected by other factors in the laboratory and/or ambient environment, including temperature, strength of dissociation, wet/dry ice cloud growth conditions, riming intensity, concentrations, drop sizes, air speeds, ventilation, etc. (13, 31, 32, 37, 38). Indeed, for NH3, the retention coefficients reported in the literature fall within wide ranges, such as <0.01 (39), 0.29–1 (31), and 1 (32).

In the numerical simulations of the interactions between atmospheric chemistry and freezing clouds, retention is the main process that decides the in-cloud scavenging of trace gases by cloud ice particles (40). Salzmann et al. (41) conducted simulations that assumed that H2O2 was either completely released or completely retained during freezing because H2O2 retention coefficients of 0 and 1 were both reported (35, 42). They concluded that the inefficient scavenging of H2O2 by the ice in clouds of tropical storms in combination with an upper air chemical source can contribute to the observed increases of the mixing ratios of H2O2 in deep convective outflows. Michael and Stuart (43) calculated the retentions of six trace chemicals, SO2, H2O2, NH3, HNO3, CH3O2, and HCOOH, and found that the overall ranges of the simulated retention fractions were between 1 × 10−6 and 1 in different scenarios. Further, these authors proposed using dynamic retention calculations to represent the changing regimes of cloud models. Ervens (44) emphasized that the removal rates of ice and precipitation are still the most uncertain processes in cloud models due to the uncertainties associated with droplets freezing and the retention of molecule species (40). Several chemistry transport models either do not consider this retention process for NH3 or treat NH3 as being completely scavenged (2).
Results and Discussion

MD Modeling. Herein, we present MD simulations to gain molecular-level insights into the behavior of ammonia in supercooled water during ice growth (SI Appendix, Fig. S1). Our focus is mainly on the molecular processes occurring between NH$_3$ and air–water–ice interface(s). Such a simple scenario is especially true in the regions with high NH$_3$ sources. To model this scenario using the MD simulation, we placed 16 ammonia molecules in the liquid water slab (containing 409 water molecules) to mimic the very high initial concentrations of ammonia. Our subsequent MD simulation results show that there is no evaporation of ammonia molecules during the individual ice particle growth process (SI Appendix, Fig. S2). Within this period, the ammonia molecules are pushed to the air–liquid interface concurrently with the continued growth of the ice crystals (Fig. 1 A–C and SI Appendix, Fig. S2). According to our simulations, after the ice growth stops and the ice crystallization reaches the air–liquid interface, only 3 out of 16 ammonia molecules are evaporated within a very short period (SI Appendix, Fig. S2). However, no additional ammonia evaporation events are observed as the system reaches an equilibrium state. Other simulations are also conducted for a total of 6, 8, 10, or 12 ammonia molecules in the supercooled water droplets, respectively. All MD simulations demonstrate that the surface air–water layer can accommodate 94–100% ammonia molecules, a feature consistent with several laboratory measurements (31, 32). In other words, in realistic atmospheric situations (such as winter stable cloud condition) where the ammonia concentrations are orders of magnitude lower than those used in the MD simulation, the ammonia molecules trapped in the surface air–water layer of the ice particles cannot be easily evaporated under nonconvective conditions.

Furthermore, the computed mean square displacements (MSDs) of the water and ammonia molecules at the interface suggest that the ice core is wet by a thin liquid-like layer (Fig. 1D). In addition, to quantitatively measure the tendencies of the ammonia evaporation from the ice particle, the free-energy barrier of the ammonia evaporation is estimated (Fig. 1E). To this end, we computed the free-energy profile of an ammonia molecule moving from the ice–water interface, passing through the liquid water layer, toward the air–liquid interface. Before the ice growth, the ammonia molecules located at the air–liquid interface can either diffuse into the liquid water layer or out into the air. Due to the much lower energy barrier (0.04 eV) of the liquid layer, ammonia molecules at the air–liquid interface tend to diffuse into the liquid layer rather than into the air, explaining why no ammonia evaporation is observed before and during the ice growth. Once the entire liquid layer turns into ice, only a single free-energy minimum is observed (red line in Fig. 1), suggesting that the ammonia molecule tends to be located at the air interface rather than being trapped inside the ice phase. This behavior is further confirmed by the snapshots shown in Fig. 1C and SI Appendix, Movie S2. Hence, once the ice growth ceases, the ammonia molecules located at the thin air–liquid interface can only diffuse into the air and cannot move into the ice phase.

![Fig. 2. Vertical profile comparisons between the measured [NH$_4^+$]/[2 SO$_4^{2-}$] and the GEOS-Chem simulated counterparts. Measured data are the averages of measurements from seven global aircraft campaigns in the years of 1991–2006 to represent the climatology. The simulation data are from the year of 2005 and are sampled along the aircraft tracks.](image)

![Fig. 3. Comparing the simulations with the satellite-measured seasonal averaged (June, July, and August) NH$_3$ values at (Top) 12 km, (Middle) 15 km, and (Bottom) 18 km over Asia. The modeled NH$_3$ shows the 6 y average (2003 and 2007–2011) and from the R1 and R0.05 simulations. The MIPAS-measured NH$_3$ values are averaged and shown only for grid boxes with more than 4 y valid data.](image)
Comparing the simulations with MIPAS-measured seasonal (June, July, and August) vertical profile of NH$_3$ over Asia over 6 y (2003 and 2007–2011). The profile of NH$_3$ volume mixing ratio is averaged over the eastern part of the Asian monsoon area (30°–40°N, 70°–110°E). The NH$_3$ simulations are from the R1, R0.05, and R0 modeling results. The SDs are shown as error bars, and also shown are root mean squared error (RMSE).

**High Concentration of NH$_3$ over Asia.** To interpret Höpfner’s (9) satellite observation, we conduct the simulations with and without considering the new pathway for UTLS NH$_3$ through ice cloud formation. Fig. 3 shows the distribution of the NH$_3$ volume mixing ratios at altitudes of 12, 15, and 18 km, which is in the range of upper troposphere and lower stratosphere (UTLS) over the subtropical region that has the tropopause in average of 12–13 km in summer (SI Appendix, Fig. S7). The modeled seasonal (June, July, and August) NH$_3$ is the average for 6 y (2003 and 2007–2011) from the R1 and R0.05 simulations, respectively. The satellite-observed seasonal NH$_3$ values are from Michelson Interferometer for Passive Atmospheric Sounding (MIPAS) (9) and are averaged for only for grid boxes with more than 4 y valid data. MIPAS showed that the enhancement of NH$_3$ over the Asian monsoon region is prominent at both 12 and 15 km altitudes with ∼10 pptv against the global background of <5 pptv. The simulations of both R1 and R0.05 show (Fig. 3) the relative enhancement of NH$_3$ over the Asian monsoon region. Only the R0.05 simulation is comparable with the MIPAS observations, with values larger than 12 pptv at both 12 and 15 km altitudes. The simulation from R1 shows a very low enhancement of NH$_3$, with maximum values of up to 1.6 pptv at an altitude of 12 km and up to 0.64 pptv at an altitude of 15 km, both of which are a factor of 6–10 lower than the observed counterparts. The same findings hold for the results at an altitude of 18 km.

The retention efficiency of NH$_3$ may vary in different altitudes and in different specific cases depending on the efficacy of breaking up the air–water interface of the ice particles in the convective updrafts. To consider this variation, simulations with R value of 0 are conducted. By sampling model results to match the valid MIPAS values for each year and each grid box, the profile of the NH$_3$ volume mixing ratios is averaged within the eastern part of the Asian monsoon area (30°–40°N, 70°–110°E) and compared with the MIPAS observed profile (Fig. 4). The performance of R0 is similar to that of R0.05, with slightly better (worse) performance above (below) 15 km altitude (Fig. 4). In contrast, the performance of simulation with R = 1 overall is poorer than that of R0.05. The profile of the MIPAS NH$_3$ revealed that the maximum concentrations of NH$_3$ in the Asian monsoon area reached ∼15 pptv at an altitude of 11–13 km. The R0.05 simulation reproduces this feature with averaged NH$_3$ of 20 pptv at the same altitude range, whereas the R1 simulation shows less than 2 pptv from the altitude of 10 to 15 km. Statistically,
the results with lower retention efficiency are robust. The averaged NH$_3$ value for 10–20 km altitude is 0.29 pptv (R1), 9.50 pptv (R0.05), and 10.99 pptv (R0) versus the observed value of 8.63 pptv. Further comparison (SI Appendix, Fig. S4) with satellite-measured NH$_3$ at 12, 15, and 18 km over Asian area (30–40°N, 70–110°E) is made for simulations with $R = 0.05$, and the modeled NH$_3$ is sampled to pair with the valid MIPAS NH$_3$ data for each summer and each grid box. Only the grid boxes with valid data for at least 4 y are considered here. The correlation between multisummer mean measurements and simulation ($R = 0.05$) is 0.86. The averaged NH$_3$ is 8.75 pptv from MIPAS and 8.77 pptv from the simulation ($R = 0.05$); both values are above the MIPAS detection limit (3–5 pptv), and their difference is within the error (near and below 1 pptv) of MIPAS retrievals (9). The year to year variations (error bars in SI Appendix, Fig. S4) from both MIPAS and the simulation could be up to 10 pptv that may in part reflects interannual changes of summer convection and meteorology.

During nonsummer seasons, we do not find any enhancement of NH$_3$ over Asia (not shown) from both the simulation and MIPAS measurement values, which indicates that both high NH$_3$ emissions and frequent deep convection events are two integral parts for the presence of NH$_3$ in UTLS. Hence, the differences between observed and simulated NH$_3$ in UTLS can be partly attributed to emission uncertainties and partially to uncertainty in convection in the model. Although few studies have investigated NH$_3$ emission over India, NH$_3$ emission inventory of Streets et al. (46), as used in this study, shows a strong peak in June over East Asia including India. In contrast, Huang et al. (47) suggest a weak summer peak of NH$_3$ emissions in China, whereas the Magnitude and Seasonality of Agricultural Emissions (MASAGE) inventory (20) indicates largest emissions in April and July. The bottom-up NH$_3$ emissions like Street et al. (46) inventory are estimated ~15% higher than the top-down estimates in May and June and ~20% lower in other months. Nevertheless, our model simulations show a large NH$_3$ emission in Indian summer boundary layer (11), which suggests the possible presence of NH$_3$ gas in the cloud droplets. Assuming that the NH$_3$ emission is underestimated by 100%, the simulated UTLS NH$_3$ concentration for R1 would be increased by a factor of 2 only, which is still lower than MIPAS measurements by at least a factor of 10. Regarding the uncertainties from deep convection, as discussed above. Huang et al.'s (48) finding, our simulated NH$_3$ is also lower than MIPAS measurements above 15 km by about 50% on average. This uncertainty caused by convection contributes the discrepancy between R0.05 simulation and the observation above 15 km, although it cannot explain the difference of NH$_3$ between R1 simulation and MIPAS observations in 10–15 km.

The wet scavenging process in the GEOS-Chem model could be another reason to cause uncertainties in the gas or aerosol contents in the UTLS (49), which allows for greater NH$_3$ survival in the uplift and decreasing the amount of NH$_3$ lost due to precipitation. We tested the response of NH$_3$ to wet scavenging efficiency by reducing the wet deposition by 10% (the average range of relative uncertainty in wet deposition in GEOS-Chem) (50) and compared the results from the simulations with wet deposition unchanged. Both simulations had an $R$ value of 1. As expected, more NH$_3$ is lifted to 12 and 15 km in the simulation with 90% wet deposition than in the simulations with 100% wet deposition (Fig. 3 and SI Appendix, Fig. S5); however, the NH$_3$ values still remained much lower than those of the MIPAS observations. For example, at an altitude of 12 km, the high value of the NH$_3$ mixing ratio over the Asian monsoon area is ~4 pptv with 90% wet deposition and is ~1.6 pptv with wet deposition unchanged. Both the MIPAS and R0.05 simulations with wet deposition unchanged have NH$_3$ of 10 pptv and larger. In summary, direct transport of NH$_3$ during deep convection has a minor contribution to high concentration of NH$_3$ in UTLS. If we consider doubling NH$_3$ emissions, it could increase NH$_3$ at 12 km from 1.6 to 3.2 pptv. Decreasing wet scavenging to 90% increases NH$_3$ at 12 km from 1.6 to 4.0 pptv (SI Appendix, Fig. S5). Hence, consideration of combined uncertainties in NH$_3$ emission and wet deposition could result in the maximum value of NH$_3$ at 12 km of 5.6 pptv, which is still less than 10 pptv as shown in the MIPAS measurements and R0.05 simulations. Among the uncertainties addressed above, although there is a possibility for NH$_3$ (gas) to survive in the wet scavenging and to be uplifted to UTLS, the low retention effects of NH$_3$ in ice clouds over the Asian monsoon region are a dominant (albeit not exclusive) contributor to the high NH$_3$ concentrations over the Asian monsoon region. This result is further supported by the observation of the homogeneous freezing of supercooled water droplets at temperature of 235,15 K or lower in the case of strong updrafts (51), due to shorter transit times that allow the liquid drops to reach high altitudes (51, 52). Hence, the conversion from liquid to ice cloud particles can occur in UTLS during deep convection conditions over Asia where air temperature at tropopause (~12 km) is on average ~226 K (SI Appendix, Fig. S7).

Summary and Discussion

The presence of NH$_3$ in the UTLS suggests that stratospheric aerosol particles are not ubiquitously 75% sulfate acid as often believed (53) or assumed in the global models for climate prediction (54), and their composition can be in the form of ammonium sulfate and ammonium nitrate, at least over Asia in summer. Indeed, ammonium sulfate in stratospheric aerosols (up to 28 km) was reported by Bigg (55) over Wyoming, United States, and by Gras (56) over Mildura, Australia. Overall, the summary of early in situ data by Turco et al. (57) estimated that (NH$_4$)$_2$SO$_4$ may weigh 10% in total stratospheric aerosol mass but admitted that the presence of ammonia in UTLS is “an open question.” This study reveals the importance of release of NH$_3$ from freezing clouds (as summarized in Fig. 5) in the deep convective systems for the presence of NH$_3$ in the UTLS by integrating MD modeling, global transport modeling, and limited satellite measurements. Although our analysis focuses on NH$_3$ over the western Asian summer monsoon region, this pathway for UTLS NH$_3$ also has implications for other trace gases and locations. Many regions are affected by gas pollution that encounters deep convection systems, such as North America, tropical Africa, and Australia. For example, the simulation shows (SI Appendix, Fig. S6) the enhancement of NH$_3$ concentrations over tropical Africa during the boreal winter season at 12 km in our R0.05 simulations but not in the R1 simulations. Furthermore, in UTLS where relative humidity (with respect to liquid water) is low, ammonium sulfate is more likely in the solid phase, which may facilitate the formation of cirrus clouds (58) and affect the Earth’s radiative energy budget (45). The presence of NH$_3$ can also lead to formation of nitrate aerosols through its reaction with nitric acid, which can in turn affect NOx-catalyzed mechanism for O$_3$ loss in the stratosphere. Therefore, explicit consideration of role of NH$_3$ in both chemistry transport and climate models is emergently important, especially after considering that NH$_3$ emission continues to grow in part due to the increased use of fertilization in agriculture (3, 4) and in part due to nonexistence of policy to regulate NH$_3$ emissions in many parts of the world. Such advancement in earth system models can be carried out and evaluated with more observation data of trace gases and (aerosol and cloud) particle composition in UTLS and through the adoption of MD simulation as a tool for improving parameterization of interactions between trace gases and particles at air–particle interface.
Materials and Methods

MD Simulation. The MD simulations are carried out using the Groningen Machine for Chemical Simulations 4.5 program. More detailed methods and models are given in SI Appendix.

Global Chemical Transport Model. A global 3D CTM, GEOChem (59), is used to simulate the transport, deposition, oxidation, and other related chemical processes of trace gases and aerosols. A more detailed model is given in SI Appendix.

Aircraft Measurements. Aircraft measurements of the NH\textsubscript{3} and SO\textsubscript{2} from seven field campaigns are analyzed in this study. More detailed information is given in SI Appendix.

Satellite NH\textsubscript{3} Data. The seasonal (3-monthly) NH\textsubscript{3} averages within the bins of 10° latitude by 10° longitude and a 1-km altitude are provided at www.imk-asf.kit.edu/english/308.php by Höfler et al. (9). More detailed information is given in SI Appendix.

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