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Introduction

New particle formation (NPF) from gaseous precursors is an important source of particles in the atmosphere.^{1,2} These newly formed particles can grow and ultimately act as cloud condensation nuclei (CCN), contributing to climate forcing.^{3–6} Up to half of global CCN is estimated to originate from NPF.^{5,6} In addition, particles have well-known deleterious impacts on human health^{7,8} and visibility.^{9–11} Despite these impacts, the underlying mechanisms and the potential species driving NPF are not well understood. This limits our ability to quantitatively assess the impacts of particles on visibility, human health and climate change as well as to develop effective control strategies.^{1,12}

Field measurements have shown that sulfuric acid (H_2SO_4) is a key species for NPF,^{13–15} possibly with the involvement of organic compounds.^{16–18} The presence of ammonia (NH₃) can stabilize nucleating clusters, and enhance nucleation by orders of magnitude.^{19–24} However, even taking into account this effect, atmospheric concentrations of H₂SO₄ and NH₃ are often not sufficient to explain nucleation rates in the atmosphere.²³

Amines have recently been recognized as additional bases that can participate in NPF. $^{22,24-28}$ Aminium salts from the

New particle formation and growth from methanesulfonic acid, trimethylamine and water

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New particle formation from gas-to-particle conversion represents a dominant source of atmospheric particles and affects radiative forcing, climate and human health. The species involved in new particle formation and the underlying mechanisms remain uncertain. Although sulfuric acid is commonly recognized as driving new particle formation, increasing evidence suggests the involvement of other species. Here we study particle formation and growth from methanesulfonic acid, trimethylamine and water at reaction times from 2.3 to 32 s where particles are 2–10 nm in diameter using a newly designed and tested flow system. The flow system has multiple inlets to facilitate changing the mixing sequence of gaseous precursors. The relative humidity and precursor concentrations, as well as the mixing sequence, are varied to explore their effects on particle formation and growth in order to provide insight into the important mechanistic steps. We show that water is involved in the formation of initial clusters, greatly enhancing their formation as well as growth into detectable size ranges. A kinetics box model is developed that quantitatively reproduces the experimental data under various conditions. Although the proposed scheme is not definitive, it suggests that incorporating such mechanisms into atmospheric models may be feasible in the near future.

reaction of acids and amines have been found to be widely present in particles.^{29–32} Amines have a variety of sources, including animal husbandry, biomass burning, industrial and agriculture activities as well as biological processes in the ocean.³³ Release associated with their use in CO₂ capture and storage could become more important as this technology becomes more widely adopted.^{33–35} Although atmospheric concentrations of amines are typically parts per trillion (ppt), one or more orders of magnitude less than NH₃, amine concentrations near animal husbandry operations can be up to several parts per billion (ppb).^{36–40} Amongst over 150 amines that have been identified in the atmosphere,³³ trimethylamine (TMA, (CH₃)₃N) is one of the most abundant.^{33,40,41}

Results from quantum chemical calculations show that amines are much more strongly bound to H_2SO_4 than NH_3 ,²⁵ and laboratory experiments have demonstrated higher efficiencies of amines in particle formation compared to NH_3 .^{22,26–28} In addition, amines have been shown to displace NH_3 in sulfate particles and clusters.^{42–45} Based on these results, small particles are more likely to be aminium salts even if ammonium salts are initially formed.

While sulfuric acid is often measured in field campaigns and linked to NPF, other sulfur-containing compounds are potentially involved in NPF. For example, methanesulfonic acid (MSA, CH₃SO₃H) is commonly detected in particles,^{29,30,46–49} but only a few studies have focused on its potential role in NPF.^{50–53}

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Atmospheric MSA is formed along with SO₂, the precursor to H_2SO_4 , in the oxidation of organosulfur compounds generated from biological processes, biomass burning and from agricultural, industrial, and domestic activities.^{54–58} The gas-phase concentration of MSA is typically 10–100% of that of H_2SO_4 in the coastal marine boundary layer.^{59,60} Although MSA was shown to be much less efficient than H_2SO_4 in forming particles with water,^{50–52,61} a recent study from this laboratory found that MSA does form particles with dimethylamine (DMA, (CH₃)₂NH) and TMA, but only in the presence of water vapor.⁵³ However, this study used a large volume flow system with reaction times from 4.2 min to ~1 h, so that early stages of particle formation and growth could not be investigated.

In the present study, a newly designed and tested borosilicate glass flow system⁶² was applied to investigate particle formation and growth from MSA and TMA at shorter times, from 2.3 to 32 seconds. The effect of relative humidity (RH) and precursor concentrations on particle formation and growth is explored. A kinetics scheme for particle formation from MSA, TMA and water that reproduces the experimental data reasonably well is developed based in part on previous quantum chemical calculations of likely early intermediates.⁵³ This approach is sufficiently simple that it could be incorporated into atmospheric models of new particle formation in a relatively straightforward manner.

Experimental

The flow system, described in detail elsewhere,⁶² is shown in Fig. 1. Briefly, it is fabricated from borosilicate glass and is designed to cover reaction times from 2.3 to 32 s. The major section of the reactor has a diameter of 7.6 cm and a length of 1.3 m, and is water-jacketed for temperature control. End-caps are mated to each end of the major section and sealed with O-rings. Two perforated hollow glass rings (Fig. 1, ring A and ring B) serve as fixed inlets at the upstream end. Two movable concentric glass tubes are guided by the upstream cap into the reactor, and terminate in two perforated hollow glass "spokes" which serve as the other two inlets (Fig. 1, spokes C and spokes D). A movable stainless steel sampling tube guided by the downstream cap is used to sample particles and gases. The downstream end-cap also has a 1/2" glass joint to vent the majority of the flow.

The reaction time is controlled by changing the distance between the sampling tube and the spoke inlets. The distance is then converted to the corresponding reaction time using a previously determined conversion. 62

The flow reactor was cleaned with Nanopure water $(>18.0 \text{ M}\Omega \text{ cm}; \text{ Model 7146}; \text{ Thermo Scientific})$, and purged with dry purified air at least overnight. Gas-phase MSA and TMA along with dry/humidified air were introduced into the flow reactor from selected inlets to initiate particle formation and growth. Gas flows were controlled by high precision mass flow controllers (Alicat or MKS), and were regularly calibrated (Gilibrator 2; Sensidyne). Dry compressed air was passed through a purge gas generator (Model 75-62; Parker Balston), carbon/alumina media (Perma Pure, LLC), and a 0.1 µm filter (DIF-N70; Headline Filters) for further purification. Relative humidity was adjusted by diverting part of the dry air flow through a water bubbler filled with Nanopure water. An RH probe (Model HMT338; Vaisala) was placed at the downstream end to monitor RH. Gas-phase MSA was generated by directing dry purified air over liquid MSA (99.0%, Fluka). The concentration of MSA in the flow reactor was controlled by adjusting the flow of purified air over the liquid MSA. It was found necessary to condition the flow reactor with a flow of MSA for two to three days prior to each experiment to passivate the walls with respect to MSA uptake. MSA was measured by passing the gas flow from the MSA trap through a 0.45 µm Durapore filter (Millex-HV) for 5-15 min followed by extraction with 10 mL Nanopure water and analysis using ultra performance liquid chromatography coupled with a tandem mass spectrometer (UPLC-MS/MS, Waters). MSA concentrations in the flow system were then calculated based on the concentration exiting the trap and the total gas flow.

Gas-phase TMA (1 ppm in N₂; Airgas) was used without further purification. Although certified by the manufacturer, the concentration of TMA was independently measured by collection onto a weak cation exchange resin for 30–60 min followed by extraction with 10 mL 0.1 M oxalic acid (Fluka) and analysis using ion chromatography as previously described.⁴⁰ The measured concentration of TMA, which was generally lower than the value provided by the manufacturer, was used to calculate its concentration in the flow system. This analysis also confirmed that NH₃ and other amines were not present at significant levels (<0.1% of TMA).

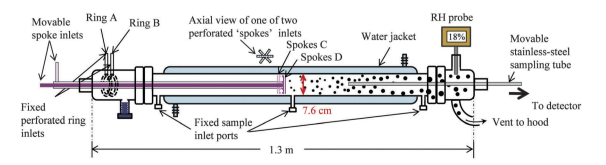


Fig. 1 Schematic of the flow reactor.⁶²

Table 1 The mixing sequence, initial concentrations of gaseous precursors and relative humidity in each experiment

Exp. #	Gas introduced				$[MSA]^a$		$[TMA]^a$		$[H_2O]$	
	Ring A	Ring B	Spokes C	Spokes D	In ppb	In 10^{10} cm^{-3}	In ppb	In 10^{10} cm^{-3}	In % RH	In 10 ¹⁶ cm ⁻³
1	Dry air	Dry air	MSA	TMA	1.8	4.4	2.5	6.2	<2	<2
2	H ₂ O/air	H ₂ O/air	MSA	TMA	1.8	4.4	2.5	6.2	48	37
3^b	H ₂ O/air	H ₂ O/air	MSA	Dry air	1.8	4.4	0.0	0.0	50	38
4	TMA	MSA	Dry air	Dry air	1.7	4.2	5.0	12	$<\!2$	$<\!2$
5	TMA	MSA	H ₂ O/air	H ₂ O/air	1.7	4.2	5.0	12	18	14
6	$TMA + H_2O/air$	MSA	Dry air	Dry air	1.7	4.2	5.0	12	19	15
7	$TMA + H_2O/air$	MSA	Dry air	Dry air	1.7	4.2	5.0	12	8	6
8	H ₂ O/air	H ₂ O/air	MŠA	TMA	1.8	4.4	2.0	4.9	7-56	5-43
9	H_2O/air	H ₂ O/air	MSA	TMA	3.1	7.6	2.0	4.9	9-59	7-45
10	H_2O/air	H ₂ O/air	MSA	TMA	2.7	6.6	1.0	2.5	6-57	5-44
11	H_2O/air	H ₂ O/air	MSA	TMA	2.0	4.9	0.7 - 1.1	1.7 - 2.7	55	42
12	H_2O/air	H ₂ O/air	MSA	TMA	1.4 - 2.0	3.4-4.9	1.1	2.7	55	42
13	Dry air	Dry air	MSA	TMA	2.4	5.9	2.0 - 2.6	4.9-6.4	$<\!2$	$<\!2$
14	H ₂ O/air	H ₂ O/air	MSA	TMA	0.3	0.7	10-26	25-64	55	42
15	H ₂ O/air	H ₂ O/air	MSA	TMA	14.5	35.7	0.5-0.9	1.2 - 2.2	55	42

^{*a*} Sampled from the sources and represent upper limits for the concentrations since some losses may occur between the source and flow tube inlets and in the flow tube itself. ^{*b*} No detectable particles were formed.

All experiments were performed at ambient temperature with a total flow of 17.0 liters per minute (lpm). The mixture of purified air and MSA as well as the mixture of purified air and TMA were separately introduced into the flow system either through the rings or spokes, depending on the experiment. Dry or humidified air was added as a carrier gas and diluent. Particle formation and growth at different reaction times were achieved by changing the position of the sampling tube in the flow system. The relative concentrations of MSA, TMA and water vapor and their order of mixing were adjusted to explore their effects on particle formation and growth. Experiments with varied initial concentrations of gaseous precursors, RH and mixing sequence present in this study are tabulated in Table 1.

Particle size distributions were monitored using a scanning mobility particle sizer (SMPS) consisting of an electrostatic classifier (Model 3080; TSI), a nano-differential mobility analyzer (Model 3085; TSI), and a butanol-based condensation particle counter (CPC, Model 3776; TSI). The manufacturer specified 50% cut-off size of the SMPS based on sucrose particles is ~2.5 nm. While SMPS can detect some particles below 2.5 nm, depending on the composition,⁶³ this portion of the data has high uncertainties and thus was excluded in quantitative analysis. The geometric mean mobility diameters obtained from SMPS are reported as particle diameters.

Kinetics schemes for particle formation from MSA, TMA and H₂O were developed and simulated in a box model, in which ordinary differential equations are integrated with the solver, 'gsl_odeiv2_step_msbdf', provided in the GNU Scientific Library.⁶⁴

Results and discussion

Effect of water on new particle formation and growth

The effect of water on particle formation and growth was first investigated by mixing 1.8 ppb MSA added through the spokes C with 2.5 ppb TMA added through the spokes D (Exp. 1 and 2 in Table 1). Fig. 2 shows particle size distributions as a function of reaction time under dry conditions (Fig. 2a, Exp. 1) and at 48% RH (Fig. 2b, Exp. 2). In the latter case, water vapor was added with air through the upstream inlets of ring A and ring B so that it is present when MSA and TMA react. Further comparisons of particle number concentrations (N_{exp}) and diameters ($D_{p,exp}$) are shown in Fig. 3.

It is noteworthy that we did not observe particle formation from $MSA-H_2O$ in the absence of TMA (Exp. 3), indicating that

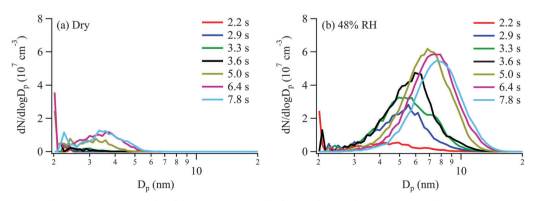


Fig. 2 Size distributions of particles measured by SMPS from the reaction of 1.8 ppb MSA with 2.5 ppb TMA at different reaction times in (a) Exp. 1 under dry conditions, and (b) Exp. 2 at 48% RH.

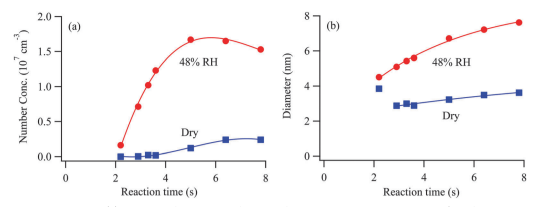


Fig. 3 (a) Number concentrations and (b) diameters of particles as a function of reaction time under dry conditions (blue filled square, Exp. 1) and at 48% RH (red filled circle, Exp. 2) corresponding to Fig. 2. Particles were measured by SMPS from the reaction of 1.8 ppb MSA with 2.5 ppb TMA. The lines between data points are drawn as guides to the eye. Measurements of the diameters have high uncertainties at short reaction times due to the experimental challenges of detecting low particle numbers, and have been disregarded in drawing lines.

there is negligible contamination of ammonia and amines in water. Binary nucleation of MSA–H₂O was reported in previous studies,^{50–53} but the nucleation was not as efficient as for H₂SO₄.^{50,61} It is likely that higher MSA levels and longer reaction times than those used in the current study are required to form detectable particles from MSA + H₂O.

Although MSA mixed with TMA shows particle formation under both dry and humid conditions, particle formation and growth are much more favorable when water is present. Under dry conditions, particles grow relatively slowly, with $D_{p,exp}$ increasing from 2.9 to 3.6 nm in 4.9 s, an average growth rate of 0.14 nm s^{-1} . After a short induction time, the particle number concentration increases. Spikes are seen in the size distributions of particles below the SMPS cut-off of 2.5 nm at the shortest times (Fig. 2). Although signal variability precludes quantification at these small sizes, as indicated earlier, the presence of these peaks suggests that there is a pool of small clusters and their growth into detectable sizes is relatively slow under dry conditions. Preliminary experiments using a recently acquired particle size magnifier (PSM, Model A10; Airmodus) which has a cut-off size of \sim 1.3 nm for ammonium sulfate particles confirms that about 50-70% of particles detected by PSM are smaller than the 2.5 nm cut-off size of SMPS.

The presence of water significantly increases both the number concentration and diameter of particles, with particle growth from 4.5 to 7.6 nm in 5.6 s, an average growth rate of 0.55 nm s⁻¹. Even at the shortest experimentally accessible reaction time when particles are relatively small, particles formed under humid conditions are significantly larger than those under dry conditions. The particle number concentration at 48% RH increases in the initial 5 s, and then slowly decreases. However, particles keep growing throughout the experiment. The slow decrease in N_{exp} and increase in $D_{\text{p,exp}}$ after 5 s is due to a combination of the depletion of the gas-phase precursors, coagulation and wall losses. For example, taking a diameter of 8 nm, a particle concentration of 1.7×10^7 cm⁻³, an assumed density of the condensed phase of 1.3 g cm⁻³ and composition of 1:1:1 MSA:TMA:H₂O, there are 2×10^{10} MSA molecules in the condensed phase per cm³ of air. Even if the

particle has grown primarily through uptake of water to give a particle composition of 1:1:10 MSA:TMA:H₂O, there would be 1×10^{10} MSA molecules per cm³ of air tied up in the condensed phase. Given the difficulty in accurately measuring the concentrations of low volatility species such as MSA, these values are comparable to the initial concentration of MSA (1.8 ppb, 4×10^{10} cm⁻³), suggesting that a large portion of MSA and TMA is in the condensed phase. Because of the high number concentration and increasing diameters of particles, the remaining gaseous MSA and TMA as well as small clusters can be easily scavenged by existing particles before growing into the detectable size range. In addition, coagulation starts playing a role at particle concentrations >10⁷ cm⁻³ on our experimental timescales.⁹

It is possible that water is not involved in the initial formation of clusters but only grows small particles into diameters >2.5 nm that can then be detected. In order to further explore the effect of water, the mixing sequence of MSA, TMA and H₂O in the flow reactor was adjusted (Exp. 4-7). TMA and MSA were separately introduced along with flows of air into the inlets of ring A and ring B at the upstream end of the reactor. The flow from each ring inlet was 7 lpm to make the total flow from the ring inlets 14 lpm. Flows of air through the inlets of spokes C and spokes D were maintained at 2 and 1 lpm, respectively. There is turbulence around the upstream ring inlets that results in significant wall losses of gaseous precursors and particles so that in the configuration of Exp. 4 under dry conditions, the number concentration is smaller than those in Fig. 2a and 3a. However, the effect of water can be probed by comparing the results in the presence of water to those under dry conditions where the addition of MSA and TMA remains constant and just the inlet used to add water vapor and its concentration changes.

Fig. 4 shows the ratios of N_{exp} and $D_{p,exp}$ collected under humid conditions to those collected under dry conditions at the same reaction time for Exp. 4–7. The addition of water through the spokes C and spokes D corresponding to 18% RH after MSA and TMA from the upstream rings had reacted for 11.6 s (Exp. 5) had no significant effect on the number

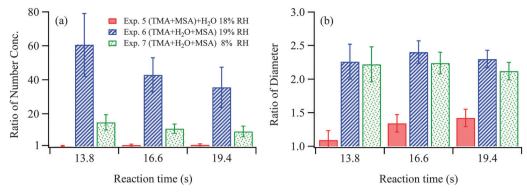


Fig. 4 Ratios of (a) number concentrations and (b) diameters of particles measured by SMPS in Exp. 5–7 to those under dry conditions in Exp. 4 at the same reaction time. Particles are from 1.7 ppb MSA with 5.0 ppb TMA under different mixing scenarios and RH. Exp. 4: MSA and TMA were added separately from ring A and ring B under dry conditions; Exp. 5: with H₂O corresponding to 18% RH added through spokes C and spokes D; Exp. 6: with the H₂O corresponding to 19% RH added through ring A simultaneously with TMA; Exp. 7: with the H₂O corresponding to 8% RH added through ring A simultaneously with TMA; Exp. 7: with the H₂O corresponding to 8% RH added through ring A simultaneously with TMA, Exp. 7: with the H₂O corresponding to 10.1.0) × 10⁴ cm⁻³ at reaction times of 13.8, 16.6 and 19.4 s, respectively. The corresponding diameters of particles were (3.6 ± 0.2) , (3.6 ± 0.1) and (3.9 ± 0.1) nm, respectively. The total reaction time shown on the axes includes the 11.6 s between the ring and spoke inlets. Uncertainties represent two standard deviations from triplicate SMPS measurements.

concentration (Fig. 4a, Exp. 5), but the size of the particles slightly increased (Fig. 4b, Exp. 5). This is consistent with the high hygroscopicity of aminium salts.⁶⁵ In contrast, when water vapor is present simultaneously with MSA and TMA (Exp. 6) the generation of particles in the detectable size range is enhanced. The number concentration and diameter of particles are significantly higher than those formed by adding H_2O after MSA and TMA have reacted. Enhanced particle formation and growth by mixing H_2O simultaneously with MSA and TMA are significant even at 8% RH (Exp. 7). These results show that water is not only involved in particle growth but also plays a central role at the initial stages of particle formation.

The fact that adding water vapor after MSA and TMA have reacted for 11.6 s had no effect on the number of detectable particles (Fig. 4a, Exp. 5) suggests that at this point most of the limiting reagent (MSA in this case) is tied up in clusters and particles with TMA, and little gas-phase MSA is available to form particles with TMA and H_2O .

Particle formation and growth as a function of precursor concentrations

The present study enabled measurements of particle formation and growth up to 56% RH, while MSA and TMA concentrations were held constant. The variation of N_{exp} and $D_{p,exp}$ with reaction time as a function of RH by mixing 1.8 ppb MSA added through the spokes C with 2.0 ppb TMA added through the spokes D is shown in Fig. 5 (Exp. 8). Water continues to promote particle formation and growth up to 56% RH, the highest RH used in this study. Significant uncertainties exist at low RH and short reaction times due to the experimental challenges associated with detecting the low numbers of small particles. It is noteworthy that prior to particle formation, there is an induction period of a few seconds, which grows shorter as the RH increases. We hypothesize that this induction period represents the time required for small clusters to grow into detectable-size particles.

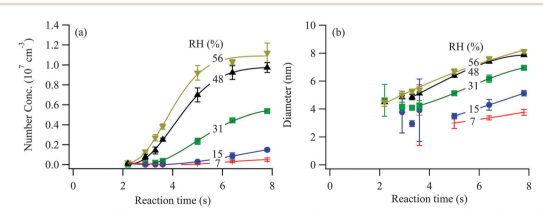


Fig. 5 (a) Number concentrations and (b) diameters of particles measured by SMPS in Exp. 8 from the reaction of 1.8 ppb MSA with 2.0 ppb TMA as a function of reaction time at RH 7% (red plus), 15% (blue filled circle), 31% (green filled square), 48% (black filled triangle) and 56% (brown filled upside-down triangle). The lines between data points are drawn as guides to the eye. Uncertainties represent two standard deviations from triplicate SMPS measurements. Measurements of the diameters have high uncertainties at short reaction times due to the experimental challenges of detecting low particle numbers, and have been disregarded in drawing lines.

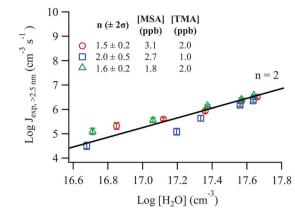


Fig. 6 Particle formation rates with two standard deviations as a function of H_2O observed in the MSA-TMA- H_2O system. Three experiments (Exp. 8–10) were performed under varied concentrations of MSA, TMA and H_2O . The power dependency (*n*) with two standard deviations and the concentrations of MSA and TMA for each experiment are shown.

Formation rates of detectable particles $(J_{exp,>2.5nm})$ can be estimated using the linear regions of the number concentration-time curves in Fig. 5a. The regions that have been used to calculate $J_{exp,>2.5nm}$ are indicated in Fig. S1 in the ESI.[†] The values of $J_{exp, > 2.5nm}$ are plotted as a function of water vapor concentration in Fig. 6. The slopes obtained from three separate experiments (Exp. 8–10) are $1.5 \pm 0.2 (2\sigma)$, $2.0 \pm 0.5 (2\sigma)$ and 1.6 \pm 0.2 (2 σ) with a weighted average of 1.6 \pm 0.2 (2 σ), suggesting that the overall particle formation rate is approximately second order with respect to water. Water vapor has also been reported to play a role in particle formation from H₂SO₄ and amines.^{22,26,27,66,67} The reaction order of 2 with respect to H₂O does not imply an elementary reaction involving two water molecules nor the number of water molecules in a critical cluster, but rather the involvement of water in the rate-determining step(s).⁶⁸ The second order dependence is captured by the individual steps in the kinetics model as discussed below, and is attributed to the involvement of water in forming the reactive MSA hydrates and/or in subsequent rate-determining step(s).

The effects of MSA and TMA on particle formation and growth were further investigated by independently varying the concentration of TMA or MSA at 55% RH as shown in Fig. 7 and 8. Fig. 7 shows N_{exp} and $D_{p,exp}$ as a function of reaction time while the concentration of TMA increased from 0.7 to 1.1 ppb at a constant MSA of 2.0 ppb (Exp. 11). Fig. 8 shows N_{exp} and $D_{p,exp}$ as a function of MSA increased from 1.4 to 2.0 ppb at a constant TMA of 1.1 ppb (Exp. 12). Both N_{exp} and $D_{p,exp}$ increase with increasing MSA and/or TMA levels. An induction period before rapid particle formation was observed in all cases but was shorter at higher precursor concentrations.

Proposed kinetics scheme

For the formation of particles from sulfuric acid and amines, a "birth-death equations" approach has been developed by others in which the formation and loss of molecular clusters through collisions and evaporation are calculated.^{69,70} The addition of a cluster or molecule to a pre-existing cluster is typically taken to occur at the gas-collision rate. The evaporation rate is obtained from the free energy difference between the product cluster and the reacting species/clusters that formed it, calculated using a quantum chemical approach. This allows for the evaporation of not only monomers but also clusters.⁷¹ Much work has gone into quantum chemical calculations of selected hydrated systems, *i.e.* sulfuric acid with amines or ammonia^{66,67,72–74} and organic acids with ammonia,^{75,76} and comparison with a thermodynamic model has been carried out.⁶⁷ The "birth-death equations" approach has yielded prediction of particle formation rates necessary for analysis and comparison with experimental data in limited cases, such as sulfuric acid with ammonia and dimethylamine.²⁸ Only recently, simulations involving hydrated clusters using their Gibbs free energies in the system of sulfuric acid and dimethylamine were carried out to provide predicted particle formation rates for comparison to observed ones.^{74,77,78} Calculation of free energies for all relevant hydrated clusters up to a reasonable size for incorporation into a "birth-death equations" model is not feasible for the MSA-TMA-H₂O system at the

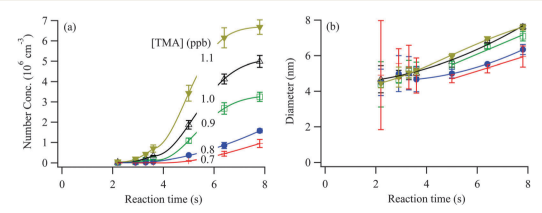


Fig. 7 (a) Number concentrations and (b) diameters of particles measured by SMPS in Exp. 11 from the reaction of 2.0 ppb MSA with variable concentrations of TMA as a function of reaction time at 55% RH. The lines between data points are drawn as guides to the eye. Uncertainties represent two standard deviations from triplicate SMPS measurements. Measurements of the diameters have high uncertainties at short reaction times due to the experimental challenges of detecting low particle numbers, and have been disregarded in drawing lines.

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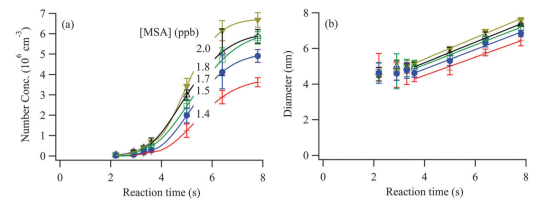


Fig. 8 (a) Number concentrations and (b) diameters of particles measured by SMPS in Exp. 12 from variable MSA concentrations reacting with 1.1 ppb TMA as a function of reaction time at 55% RH. The lines between data points are drawn as guides to the eye. Uncertainties represent two standard deviations from triplicate SMPS measurements. Measurements of the diameters have high uncertainties at short reaction times due to the experimental challenges of detecting low particle numbers, and have been disregarded in drawing lines.

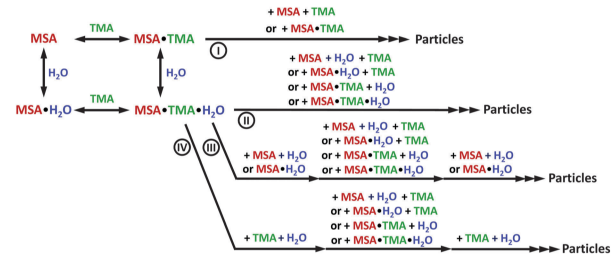


Fig. 9 Proposed four different pathways for particle formation from the MSA–TMA–H₂O system. Particles are mainly formed from pathway (I) under dry conditions, (II) under humid conditions with similar levels of MSA and TMA, (III) under humid conditions with excess MSA and (IV) under humid conditions with excess TMA.

present time. As an alternative, other studies indicate that a kinetics model with adjustable parameters can be used to describe such systems.^{53,79}

A simplified kinetics scheme that was previously developed⁵³ was expanded to include individual steps that lead to particles in the MSA-TMA-H₂O system (Fig. 9), and the expanded kinetics scheme was simulated using a box model. A detailed mechanism with 94 species and 157 reactions that involves stepwise addition of the gaseous reactants or small clusters is tabulated in Table S1 in the ESI.† As in the earlier model, the first few steps (reactions (S1)-(S8) in Table S1, ESI†) were assumed to be diffusion-controlled in the forward direction, and rate constants were calculated using hard-sphere collision theory.⁵³ Rate constants in the reverse direction were calculated using forward rate constants and ΔG of formation obtained from quantum chemical calculations.⁵³ For the subsequent steps, rate constants were adjusted to fit particle number concentrations obtained from experiments under different conditions.

It should be noted that this multistep mechanism and the rate constants are not unique, but simply used to demonstrate that a box model kinetics approach can capture the fundamental behavior of the system. Quantitative treatment of the thermodynamics and kinetics of all of the clusters would be needed to promulgate a definitive mechanism, but is beyond the scope of this work. The basis of the mechanism is the following.

First, MSA is known to form hydrates with water.⁵² Fig. 10 shows the concentrations and fractional distribution of MSA hydrates at various RH for 2 ppb MSA using published equilibrium constants.⁵² The monohydrate and dihydrate (n = 1 or 2) are the two most prominent hydrates at all RH so are included in the model. Both unhydrated and hydrated MSA can react with TMA:⁵³

$$MSA + TMA \rightleftharpoons MSA \cdot TMA \tag{1}$$

$$MSA + H_2O \rightleftharpoons MSA \cdot H_2O \tag{2}$$

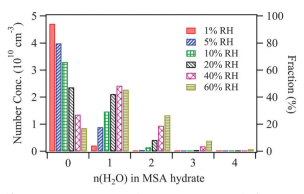


Fig. 10 The concentrations and fractional distribution of MSA hydrates at various RH for 2 ppb MSA using published equilibrium constants.⁵²

N

 $MSA \cdot H_2O + TMA \rightleftharpoons MSA \cdot TMA \cdot H_2O$ (3)

$$MSA \cdot TMA + H_2O \rightleftharpoons MSA \cdot TMA \cdot H_2O$$
(4)

$$MSA \cdot H_2O + H_2O \rightarrow MSA \cdot (H_2O)_2$$
(5)

$$MSA \cdot (H_2O)_2 + TMA \rightarrow MSA \cdot TMA \cdot (H_2O)_2$$
(6)

As stated earlier, reactions (1)-(4) are assumed to be diffusioncontrolled in the forward direction, and reversible.^{52,53} No further evaporation of monomer from large clusters or loss of small clusters from large ones as proposed in the H₂SO₄ system^{69,70} was included. Under dry conditions, there are no significant concentrations of hydrates (the presence of some due to trace amounts of water cannot be excluded) and the reaction with TMA forms stable clusters, MSA·TMA. In a similar vein, H₂SO₄ and DMA have been shown to form an initial H₂SO₄·DMA cluster which grows by stepwise addition of the acid and base to generate clusters of increasing size with 1:1 stoichiometry.^{28,69,70} Addition of DMA to dimethylaminium salt clusters of MSA was observed to be fastest when there was an excess acid available in the cluster to be neutralized.80 This is similar to the case of dry conditions proposed in this study (pathway I in Fig. 9). In our reaction scheme, the clusters further grow by stepwise addition of MSA and TMA or by the addition of MSA-TMA clusters (pathway I in Fig. 9; reactions (S124)-(S151) in Table S1, ESI⁺) into particles that can be measured by SMPS (>2.5 nm). Loss of MSA-TMA from larger clusters may also occur to significantly slow particle formation and growth under dry conditions. The ion pair formed from MSA and TMA is a particularly stable unit in the MSA-TMA system.⁶⁵ The interaction energy between MSA-TMA units is significant, but not as great as that between ions in the ion pair due to the lack of hydrogen bonding. As discussed in Ortega et al.⁷¹ for the sulfuric acid–DMA system, a deep local minimum on the free energy surface can lead to evaporation of clusters that will affect particle formation rates.

As the water vapor concentration increases, hydrated MSA becomes more important so that the initial cluster is now largely MSA·TMA·H₂O. At similar levels of MSA and TMA, the growth of MSA·TMA·H₂O into detectable particles can proceed by adding MSA, hydrated MSA, TMA, MSA·TMA or MSA·TMA·H₂O (pathway II in Fig. 9; reactions (S19)–(S67) in Table S1, ESI⁺).

However, with excess MSA (pathway III in Fig. 9; reactions (S68)-(S102) in Table S1, ESI[†]) or TMA (pathway IV in Fig. 9; reactions (S103)-(S123) in Table S1, ESI⁺) present, the excess species can first add to MSA·TMA·H₂O clusters, followed by the stepwise addition of MSA and TMA. As initial clusters grow to a certain size, further growth can proceed by condensation of the excess species. It is assumed that initial concentrations of the gaseous precursors are those in Table 1, but the gases and clusters are lost through a combination of processes that includes loss to the walls and addition to existing particles (reactions (S152)-(S157) in Table S1, ESI[†]). The rate constants for losses of all clusters except MSA, TMA and MSA·H₂O were assumed to be first order, 0.05 s⁻¹. Loss rates for gaseous MSA, TMA, as well as small cluster MSA·H₂O were assumed to be larger since they will diffuse to the walls faster than larger clusters. In each pathway, the growth processes for initial clusters by addition of each type of molecule/cluster are set to have the same rate constant, reducing the number of independent rate constants to 51. For example, reactions (S20), (S26), (S32), (S38), (S44), (S50), (S56) and (S62) in Table S1 (ESI[†]) showing the growth of clusters by addition of MSA in pathway II are assigned the same rate constant, 4 imes 10^{-10} cm³ molecules⁻¹ s⁻¹. The smallest kinetics model that gives the best fit between laboratory and model predicted number concentrations treats detectable particles as clusters containing \sim 20 molecules for pathways I and IV, and \sim 30 molecules for pathways II and III. Clusters that were treated as particles are shown in bold in Table S1 (ESI[†]).

Fig. 11 compares the experimental and model results for particle formation in the MSA-TMA-H₂O system under a variety of conditions. Overall, the model calculations agree reasonably well with laboratory results throughout the course of the experiments except the case of excess MSA (Fig. 11d, Exp. 15), in which model calculations reproduce the shape of the time profiles of laboratory results but are approximately 35% lower. Although the proposed kinetics scheme is certainly not unique or definitive, it suggests that particle formation from MSA, TMA and H_2O is a very dynamic and complex process that can, however, be simplified through a series of reactions representing initial cluster formation and growth *via* addition of MSA, TMA and H_2O , as well as small clusters.

The model also reproduces well the enhancing effect of water in particle formation. Addition of water into initial clusters proceeds through:

$$MSA \cdot TMA \cdot H_2O + H_2O \rightarrow MSA \cdot TMA \cdot (H_2O)_2$$
(7)

$$MSA \cdot TMA \cdot (H_2O)_2 + MSA \rightarrow (MSA)_2 \cdot TMA \cdot (H_2O)_2 \qquad (8)$$

$$(MSA)_2 \cdot TMA \cdot (H_2O)_2 + H_2O \rightarrow (MSA)_2 \cdot TMA \cdot (H_2O)_3$$
 (9)

$$MSA \cdot TMA \cdot (H_2O)_2 + H_2O \rightarrow MSA \cdot TMA \cdot (H_2O)_3 \quad (10)$$

Reactions (5) and (7) are taken to be rate-limiting steps, resulting in an overall second order dependence of particle formation on water.

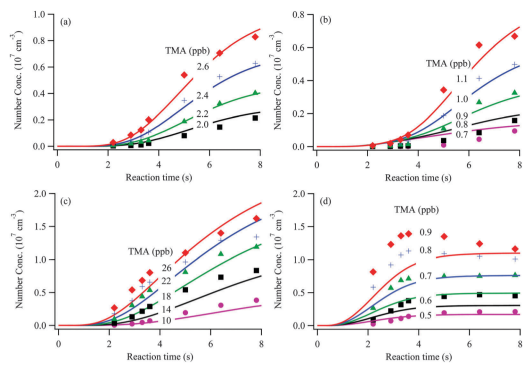


Fig. 11 Comparison of particle formation from laboratory measurements and model calculations in the MSA-TMA- H_2O system under a variety of conditions: (a) under dry conditions with the initial concentration of MSA at 2.4 ppb (Exp. 13); (b) at 55% RH with the initial concentration of MSA at 2.0 ppb (Exp. 11); (c) at 55% RH in the presence of excess TMA with initial concentration of MSA at 0.3 ppb (Exp. 14); and (d) at 55% RH in the presence of excess MSA with initial concentration of MSA at 14.5 ppb (Exp. 15). Symbols represent the experimental data, while solid lines depict the results of the model calculations.

Atmospheric implications

Although H_2SO_4 is commonly recognized as the main species driving NPF, increasing evidence from laboratory studies and field measurements has suggested that other species such as MSA also contribute.^{29,30,46–53} Although the concentration of gas-phase MSA is generally lower than that of H_2SO_4 in the atmosphere, the role of MSA in NPF will potentially become more important with the implementation of stricter environmental regulation on SO_2 emissions which lead to H_2SO_4 formation. In addition, it has been predicted that MSA can enhance cluster formation between sulfuric acid and amines under some conditions.⁸¹

Water clearly plays a critical role in particle formation from the reaction of MSA with TMA at the initial stages. Although the specific case described here only involves MSA and TMA, the enhancing effect of water is expected to be applicable to other bases such as ammonia and other amines. This suggests that new particle formation in this system will depend on RH, all other factors being equal. The same may be the case for sulfuric acid reactions with amines.^{22,26,27,66,67} Although our results suggest that particle formation in this system is quite dynamic and complex, it is still sufficiently straightforward that incorporating such schemes into atmospheric models should be feasible in the near future.

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