Reaction of SO₃ with H₂SO₄ and Its Implication for Aerosol

Particle Formation in the Gas Phase and at the Air-Water

Interface

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Abstract. The reactions between SO_3 and atmospheric acids are indispensable in improving the formation of aerosol particle. However, relative to those of SO_3 with organic acids, the reaction of SO_3 with inorganic acids has not received much attention. Here, we explore the atmospheric reaction between SO_3 and H_2SO_4 , a typical inorganic acid, in the gas phase and at the air-water interface by

- using quantum chemical (QC) calculations and Born-Oppenheimer molecular dynamics simulations.
 We also report the effect of H₂S₂O₇, the product of the reaction between SO₃ and H₂SO₄, on new particle formation (NPF) in various environments by using the Atmospheric Cluster Dynamics Code kinetic model and the QC calculation. The present findings show that the gas phase reactions of SO₃ + H₂SO₄ without and with water molecule are both low energy barrier processes. With the involvement of interfacial water molecules, H₂O-induced the formation of S₂O₇²⁻…H₃O⁺ ion pair,
- HSO₄⁻ mediated the formation of HSO₄⁻···H₃O⁺ ion pair and the deprotonation of H₂S₂O₇ were observed and proceeded on the picosecond time-scale. The present findings suggest the potential contribution of SO₃-H₂SO₄ reaction to NPF and aerosol particle growth as the facts that *i*) although H₂S₂O₇ is easily hydrolyzed with water to form H₂SO₄, it can directly participate in H₂SO₄-NH₃-
- 25 based cluster formation and can present a more obvious enhancement effect on SA-A-based cluster formation; and *ii*) the formed interfacial $S_2O_7^{2-}$ can attract candidate species from the gas phase to the water surface, and thus, accelerate particle growth.



Graphic abstract

30 **1 Introduction**

Sulfur trioxide (SO₃) is a major air pollutant (Zhuang and Pavlish, 2012; Chen and Bhattacharya, 2013; Cao et al., 2010; Kikuchi, 2001; Mitsui et al., 2011) and can be considered as the most important oxidation product of SO₂ (Starik et al., 2004). As an active atmospheric species, SO₃ can lead to the formations of acid rain and atmospheric aerosol (Sipilä et al., 2010; Mackenzie

- et al., 2015; England et al., 2000; Li et al., 2016; Renard et al., 2004) and thus plays a well-documented role in regional climate and human health (Zhang et al., 2012; Pöschl, 2005; Zhang et al., 2015; Pöschl and Shiraiwa, 2015; Haywood and Boucher, 2000; Lohmann and Feichter, 2005). In the atmosphere, the hydrolysis of SO₃ to product H₂SO₄ (SA) is the most major loss route of SO₃ (Morokuma and Muguruma, 1994; Akhmatskaya et al., 1997; Larson et al., 2000; Hazra and Sinha,
- 40 2011; Long et al., 2013a; Torrent-Sucarrat et al., 2012; Ma et al., 2020). As a complement to the loss of SO₃, ammonolysis reaction of SO₃ in polluted areas of NH₃ can form H₂NSO₃H, which not only can be competitive with the formation of SA from the hydrolysis reaction of SO₃, but also can enhance the formation rates of sulfuric acid (SA)-dimethylamine (NH(CH₃)₂, DMA) clusters by about 2 times. Similarity, the reactions of SO₃ with CH₃OH and organic acids (such as HCOOH)
- 45 were reported (Liu et al., 2019; Hazra and Sinha, 2011; Long et al., 2012; Mackenzie et al., 2015; Huff et al., 2017; Smith et al., 2017; Li et al., 2018a), and both processes can provide a mechanism for incorporating organic matter into aerosol particles. However, the reaction mechanism between SO₃ and inorganic species are still unclear.
- As a major inorganic acidic air pollutant (Tilgner et al., 2021), SA can act as an important role
 in the new particle formation (Weber et al., 1995; Weber et al., 1996; Weber et al., 2001; Sihto et al., 2006; Riipinen et al., 2007; Sipilä et al., 2010; Zhang et al., 2012) and acid rain (Calvert et al., 1985; Finlayson-Pitts and Pitts Jr, 1986; Wayne, 2000). The source of gas-phase SA is mainly produced by the gas-phase hydrolysis reaction of SO₃. For the direct reaction between SO₃ and H₂O, it takes place hardly in the atmosphere due to high energy barrier (Chen and Plummer, 1985; Hofmann and Schleyer, 1994; Morokuma and Muguruma, 1994; Steudel, 1995). However the
- addition of a second water molecule (Morokuma and Muguruma, 1994; Larson et al., 2000; Loerting and Liedl, 2000), the hydroperoxyl radical (Gonzalez et al., 2010), formic acid (Hazra and Sinha, 2011; Long et al., 2012), sulfuric acid (Torrent-Sucarrat et al., 2012), nitric acid (Long et al., 2013a),

oxalic acid (Lv et al., 2019) and ammonia (Bandyopadhyay et al., 2017) have been reported to

- 60 catalyze the formation of SA from the hydrolysis reaction of SO₃ as they can promote atmospheric proton transfer reactions. Similarity, as SA can give out protons more readily than H₂O, which in turn is more conducive to the proton transfer, thus we predict that the addition reaction involving the proton transfer between SO₃ and SA is much easier under atmospheric conditions than that between SO₃ and H₂O. However, this gas-phase reaction has not been investigated as far as we know. Previous studies have shown that the concentration of water vapor decreases significantly with increasing altitude (Anglada et al., 2013), leading to longer atmospheric lifetimes of SO₃. The gas phase reaction of SO₃ with H₂SO₄ may contribute significantly to the loss of SO₃ in dry areas where [H₂SO₄] is relatively high (especially at lower temperatures) and at higher altitude. So, it is
- 70 hydrolysis of SO₃. Meanwhile, in many gas phase reactions, single water molecule can play a catalyst role by increasing the stability of pre-reactive complexes and reducing the activation energy of transition states (Kanno et al., 2006; Stone and Rowley, 2005; Chen et al., 2014; Viegas and Varandas, 2012, 2016). For example, single water molecule in the H₂O····HO₂ + SO₃ reaction can catalyze the formation of HSO₅ (Gonzalez et al., 2010). Thus, it is equally important to study the

important to study the reaction mechanism of SO3 with H2SO4 and its competition with H2O-assisted

- SO₃ + SA reaction without and with H₂O. In addition to the gas phase reactions, many new atmospheric processes and new reaction pathways have been observed at the air-water interface (Zhong et al., 2017; Kumar et al., 2017; Kumar et al., 2018; Zhu et al., 2016; Li et al., 2016; Zhu et al., 2017). Such as, the organic acids reacting with SO₃ can form the ion pair of carboxylic sulfuric anhydride and hydronium at the air-water interface (Zhong et al., 2019). This mechanism is different from the gas phase reaction in which the organic acid either serves as a catalyst for the hydrolysis of SO₃ or acts as a reactant reacting with SO₃ directly. So, water droplets may play important roles in atmospheric behaviors between SO₃ and SA. Thus, it is also important to study the interfacial
 - mechanism between SO₃ and SA, and to compare its difference with the corresponding gas-phase reaction.
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Previous experimental studies (Otto and Steudel, 2001; Abedi and Farrokhpour, 2013) found that disulfuric acid ($H_2S_2O_7$, DSA) is the product of the reaction between SO₃ and SA. From the perspective of structure, DSA possesses two HO functional groups. Both HO groups can act as hydrogen donors and acceptors to interact with atmospheric particle precursors. It has been shown that the reaction between SO₃ and some important atmospheric species (Li et al., 2018a; Yang et al.,

- 2021; Liu et al., 2019; Rong et al., 2020) not only can cause appreciable consumption of SO₃ and thus reduce the abundance of SA from the hydrolysis of SO₃ in the atmosphere, but also can promote NPF process by their products. For example, the products of NH₂SO₃H, HOOCOOSO₃H, CH₃OSO₃H and HOCCOOSO₃H from the reactions of SO₃ with NH₃ (Li et al., 2018a), H₂C₂O₄ (Yang et al., 2021), CH₃OH (Liu et al., 2019) and HOOCCHO (Rong et al., 2020) all have a catalytic effect on the formation of new particles in aerosols. However, whether DSA produced by the reaction between SO₃ and SA contributes to aerosol formation or not is still unclear. Thus, another main question that we intend to address here is the role of DSA in atmospheric SA-NH₃ (A)
- nucleation, which have been recognized as dominant precursors in highly polluted areas, especially
 in some megacities in Asia.
 100 In this work, using quantum chemical calculations and Master Equation, we first studied the
- gas-phase reaction between SO₃ and SA to product DSA with H₂O acting as a catalyst. Then, we use the Born-Oppenheimer Molecular Dynamic (BOMD) simulations to evaluate the reaction mechanism of SO₃ with SA at the air-water interface. Finally, we used Atmospheric Clusters Dynamic Code (ACDC) and quantum chemical calculations to investigate atmospheric implications of SO₃-SA reaction to the atmospheric particle formation. Particular attention of this work is focused on the study of *i*) the mechanism difference of the SO₃ + SA reaction in the gas phase and at the air-water interface; *ii*) the fate of DSA in atmospheric NPF and its influence at

2 Computational details

various environmental conditions.

2.1 Quantum chemical calculation. The M06-2X functional has been proved to be one of the best functionals to describe the noncovalent interactions and estimate the thermochemistry and equilibrium structures for atmospheric reactions (Elm et al., 2012; Mardirossian and Head-Gordon, 2016). So, for the SO₃ + SA reaction without and with water molecule in the gas phase, the optimized geometries and vibrational frequencies of reactants, pre-reactive complexes, transition states (TSs), post-reactive complexes and products were calculated using M06-2X method (Zhao and Truhlar, 2008; Elm et al., 2012) with 6-311++G(2df,2pd) basis set by Gaussian 09 packages (Frisch, 2009). It is noted that the calculated

bond distances and bond angles at the M06-2X/6-311++G(3df,2pd) level (Figure S1) agree well with the available experimental values (Kuczkowski et al., 1981). At the same level, the connectivity

- 120 between the TSs and the suitable pre- and post-reactant complexes was performed by intrinsic reaction coordinate (IRC) calculations. Then, single point energy calculations were calculated at the CCSD(T)-F12/cc-pVDZ-F12 level (Adler et al., 2007; Knizia et al., 2009) by using ORCA (Neese, 2012).
- A multistep global minimum sampling technique was used to search for the global minima of 125 the $(DSA)_x(SA)_y(A)_z$ ($z \le x + y \le 3$) molecular clusters. Specifically, a multistep global minimum sampling technique was used to search for the global minima of the (SA)x(A)y(DSA)z (0< $y \le x + y = x + x + y \le x + y \le x + y = x + x + x +$ $z \leq 3$) clusters. Specifically, the initial n*1000 (1 < n < 5) configurations for each cluster were systematically generated by the ABCluster program (Zhang and Dolg, 2015), and were optimized at the semi-empirical PM6 (Stewart, 2013) methods using MOPAC 2016 (Stewart, 2013; Stewart, 2007). Then, up to n*100 structures with relatively lowest energy among the n*1000 (1 < n < 5) 130 structures were selected and reoptimized at the M06-2X/6-31+G(d,p) level. Finally, n*10 lowestlying structures were optimized by the M06-2X/6-311++G(2df, 2pd) level to determine the global minimum. To obtain the reliable energies, single-point energy calculations were refined at the DLPNO-CCSD(T)/aug-cc-pVTZ level based on the optimized geometries at the M06-2X/6-135 311++G(2df,2pd) level. The optimized structures and the formation Gibbs free energy of the stable clusters were summarized in Figure S9 and Table S8 of the SI Appendix, respectively.

2.2 Rate constant calculations. Using the Rice-Ramsperger-Kassel-Marcus based Master Equation (ME/RRKM) model (Miller and Klippenstein, 2006), the kinetics for the SO₃ + SA reaction without and with water molecule were calculated by adopting a Master Equation Solver for Multi Energy-well Reactions (MESMER) code (Glowacki et al., 2012). In the MESMER calculation, the rate coefficients for the bimolecular barrierless association step (from reactants to pre-reactive complexes) were evaluated by the Inverse Laplace Transform (ILT) method (Horváth et al., 2020), meanwhile the unimolecular step was performed by the RRKM theory combined with the asymmetric Eckart model. The ILT method and RRKM theory can be represented in Eq (1) and Eq (2), respectively.

$$k^{\infty}(\beta) = \frac{1}{Q(\beta)} \int_0^{\infty} k(E) \rho(E) \exp(-\beta E) dE$$
(1)

$$k(E) = \frac{W(E - E_0)}{h\rho(E)}$$
(2)

Where *h* is denoted as Planck's constant; $\rho(E)$ is denoted as the active density of state of the reactant at energy level *E*; *E*₀ is denoted as the reaction threshold energy and *W*(*E*-*E*₀) is denoted as the sum

- 150 of the rovibrational states of the transition state (TS) geometry (excluding the degree of freedom related to passing the transition state). The input parameters for electronic geometries, vibrational frequencies, and rotational constants were calculated at the M06-2X/6-311++G(2*df*,2*pd*) level and single-point energy calculations were refined at the CCSD(T)-F12/cc-pVDZ-F12 level for the modeling.
- 2.3 Born-Oppenheimer Molecular Dynamic (BOMD) simulation. The CP2K code (Hutter et al., 2014) was used in the BOMD simulations. The Becke-Lee-Yang-Parr (BLYP) functional (Becke, 1988; Lee et al., 1988) was chosen to treat with the exchange and correlation interactions, and the Grimme's dispersion was carried out to account for the weak dispersion interaction (Grimme et al., 2010). The Goedecker-Teter-Hutter (GTH) conservation pseudopotential (Goedecker et al., 1996; Hartwigsen et al., 1998) with the Gaussian DZVP basis set (VandeVondele and Hutter, 2007) and the auxiliary plane wave basis set was applied to correct the system valence electrons and the core electrons, respectively. For the plane wave basis set and Gaussian basis set, the energy cut off (Zhong et al., 2017; Zhong et al., 2018; Zhong et al., 2019) were set to 280 and 40 Ry, respectively. For each simulation in the gas phase, a 15 × 15 Å³ supercell with periodic boundary condition was adopted with a time step of 0.5 fs. As the droplet system with 191 water molecules are sufficient to describe the interfacial
- mechanism (Zhong et al., 2017), the air-water interfacial system here included 191 water molecules, SO_3 and SA in the BOMD simulation. It is pointed out that the droplet system with 191 water molecules has been equilibrated before SO_3 and H_2SO_4 was added at the water surface.
- 170 The details of the equilibrium process for the droplet system with 191 water molecules are shown in the *SI Appendix* Part 4. To avoid periodic interactions between adjacent water droplets, the size of the simulation box (Kumar et al., 2017; Kumar et al., 2018; Ma et al., 2020) was set as $35 \times 35 \times 35$ Å³ with a time step of 1.0 fs. Notably, the timestep of 1.0 fs has been proved to achieve sufficient energy conservation for the water system (Zhong et al., 2015; Li et al., 2016;
- 175 Zhu et al., 2016; Kumar et al., 2017). For all the simulations in the gas phase and at the air-water

interface, the Nose-Hoover thermostat (Zhong et al., 2017; Zhong et al., 2018; Zhong et al., 2019; Kumar et al., 2017; Kumar et al., 2018; Ma et al., 2020) was selected the NVT ensemble to control the temperature around 300 K. To eliminate the influence of the initial configuration on the simulation results of interfacial reaction, 40 BOMD simulations for the air-water interface reactions were carried out.

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2.4 Atmospheric Clusters Dynamic Code (ACDC) model

The Atmospheric Cluster Dynamics Code (ACDC) (McGrath et al., 2012) was used to simulate the cluster formation rates and mechanisms of $(DSA)_x(SA)_y(A)_z$ ($z \le x + y \le 3$) clusters at different temperatures and monomer concentrations. The thermodynamic data of quantum 185 chemical calculation at the DLPNO-CCSD(T)/aug-cc-pVTZ//M06-2X/6-311++G(2df,2pd) level of theory can be used as the input of ACDC. The birth-death equation (Eq. 3) for clusters solves the time development of cluster concentrations by numerical integration using the ode15s solver in MATLAB program (Shampine and Reichelt, 1997).

$$\frac{dc_i}{dt} = \frac{1}{2} \sum_{j < i} \beta_{j,(i-j)} C_j C_{(i-j)} + \sum_j \gamma_{(i+j) \to i} C_{i+j} - \sum_j \beta_{i,j} C_i C_j - \frac{1}{2} \sum_{j < i} \gamma_{i \to j} C_i + Q_i - S_i \quad (3)$$

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Where c_i is the concentration of cluster *i*; $\beta_{i,j}$ is the collision coefficient between clusters *i* and j; $\gamma_{(i+j) \rightarrow i}$ is the evaporation coefficient of cluster i+j evaporating into clusters i and j, and Q_i is all other source term of cluster i. (See more details of β and γ in SI Appendix Part 4). Besides, a constant coagulation sink coefficient 2×10^{-2} s⁻¹ (corresponding to the median observed in contaminated areas) was used for taking into account external losses (Yao et al., 2018; Zhang 195 et al., 2022; Liu et al., 2021b). The boundary conditions in the ACDC require that the smallest clusters outside of the simulated system should be very stable so that not to evaporate back immediately (McGrath et al., 2012). Based on cluster volatilization rate (shown in Table S10) and the formation Gibbs free energy of the clusters (shown in Table S8), the cluster boundary conditions simulated in this study were set as $(SA)_4 \cdot (A)_3$, $(SA)_4 \cdot (A)_4$, $SA \cdot (A)_3 \cdot (DSA)_3$, $(SA)_3 \cdot (A)_4 \cdot (DSA)_1$ and 200 $(SA)_2 \cdot (A)_3 \cdot (DSA)_2$. According to field observations, the concentration of SA and A was respectively set in a range of 10⁶-10⁸ molecules cm⁻³ and 10⁷-10¹¹ molecules cm⁻³ (Almeida et al., 2013; Kuang

et al., 2008; Bouo et al., 2011; Zhang et al., 2018). As the prediction in Table S7, the concentration of DSA is set to 10⁴-10⁸ molecules cm⁻³. However, DSA is easily hydrolyzed with abundant water in the troposphere to form H₂SO₄, the concentration of DSA listed in Figure S9 was overestimated.

So, the maximum concentration of DSA (10^8 molecules·cm⁻³) was not included in the effect of H₂S₂O₇ on new particle formation (NPF) in various environments. Besides, the temperature was set to be 218.15-298.15 K, which span most regions of the troposphere and the polluted atmospheric boundary layer.

3 Results and discussion

210 **3.1 Reactions in the gas phase**

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The addition reaction involving the proton transfer between SO₃ and SA (Channel DSA) proceeded through the formation of SO₃····H₂SO₄ complex followed by unimolecular transformation through transition state TS_{DSA} to form H₂S₂O₇ (Figure 1(a)). The reactant complex SO₃····H₂SO₄ was a double six-membered ring complex with a relative Gibbs free energy of -1.6 kcal·mol⁻¹. After the formation of SO₃····H₂SO₄ complex, Channel DSA overcame a Gibbs free energy barrier of 2.3 kcal·mol⁻¹, which was lower by 4.2 kcal·mol⁻¹ than that of H₂O-catalyzed hydrolysis of SO₃ (Figure S1). Rate constant for the SO₃ + SA reaction was calculated at various temperatures (Table 1). Within the temperature range of 280-320 K, the rate constants for the SO₃ + SA reaction were calculated to be 2.57×10^{-12} - 5.52×10^{-12} cm³·molecule⁻¹·s⁻¹, which were larger by 3.43-4.03 times than the corresponding values of H₂O-catalyzed hydrolysis of SO₃. Therefore, it can be said that the direct reaction between SO₃ and SA occurs easily under atmospheric conditions.

The SO₃ + H₂SO₄ reaction with H₂O produced two distinct products, labeled (*i*) H₂S₂O₇ (DSA, Channel DSA_WM) and (*ii*) H₂SO₄ (SA, Channel SA_SA). A single water molecule in (*i*) acted as a catalyst, while it played as a reactant in (*ii*). The schematic potential energy surface for the SO₃ + H₂SO₄ reaction with H₂O was shown in Figure 1. As the probability of simultaneous collision (Pérez-Ríos et al., 2014; Elm et al., 2013) of three molecules of SO₃, SA and H₂O was quite low under realistic conditions, both Channel DSA_WM and Channel SA_SA can be considered as a sequential bimolecular process. In other words, both Channel DSA_WM and Channel SA_SA occurred via the collision between SO₃ (or H₂SO₄) and H₂O to form dimer (SO₃…H₂O and H₂SO₄…H₂O) first, and then the dimer encountered with the third reactant H₂SO₄…H₂O were respectively 0.8 kcal·mol⁻¹ and -1.9 kcal·mol⁻¹, which were respectively consistent with the previous values (the

range from -0.2 to 0.62 kcal·mol⁻¹ for SO₃…H₂O complex (Bandyopadhyay et al., 2017; Long et

al., 2012) and the range from -1.82 to -2.63 kcal·mol⁻¹ for $H_2SO_4\cdots H_2O$ complex (Long et al., 2013b;

- Tan et al., 2018)). The Gibbs free energy of H₂SO₄···H₂O was lower by 2.7 kcal·mol⁻¹ than that of 235 SO₃···H₂O, thus leading to that the equilibrium constant of the former complex was larger by 1-2 orders of magnitude than that of the latter one in Table S2. Additionally, the larger equilibrium constant of H_2SO_4 ... H_2O complex leaded to its higher concentration in the atmosphere. For example, when the concentrations of SO₃ (Yao et al., 2020), H₂SO₄ (Liu et al., 2015) and H₂O (Anglada et al., 2013) were 10^6 , 10^8 and 10^{17} molecules cm⁻³, respectively, the concentrations of SO₃···H₂O and 240 H₂SO₄···H₂O were 2.41 × 10³-2.01 × 10⁴ and 5.01 × 10⁵-3.01 × 10⁸ molecules cm⁻³ within the temperature range of 280-320 K (see Table S3), respectively. So, we predict that Channel DSA_WM and Channel SA SA mainly take place via the collision of H₂SO₄…H₂O with SO₃. In order to check this prediction, the effective rate constants for two bimolecular reactions of H_2SO_4 ... $H_2O + SO_3$ and 245 SO₃···H₂O + H₂SO₄ were calculated, and the details were shown in *SI Appendix*, Part 3 and Table 1. As seen in Table 1, the $SO_3 \cdots H_2O + H_2SO_4$ reaction in both Channel DSA_WM and Channel SA SA can be neglected as their effective rate constants were smaller by 16.7-48.5 and 1.02-3.05 times than the corresponding values in the $H_2SO_4\cdots H_2O + SO_3$ reaction within the temperature range of 280-320 K, respectively. Therefore, we only considered the $H_2SO_4\cdots H_2O + SO_3$ bimolecular
- reaction in both Channel DSA_WM and Channel SA_SA.
- As for Channel DSA_WM, the H₂SO₄···H₂O + SO₃ reaction occurred in a stepwise process as displayed in Figure 1(b), which was similar to the favorable routes in the hydrolysis of COS, HCHO and CH₃CHO catalyzed by sulfuric acid (Long et al., 2013b; Li et al., 2018b; Tan et al., 2018). When the H₂SO₄···H₂O complex and SO₃ served as reactants, the reaction was initiated by complex IM_{DSA_WM}' where a van der Waals interaction (S2···O4, 2.75 Å) was found between the O4 atom of SA moiety in H₂SO₄···H₂O and the S atom of SO₃. After complex IM_{DSA_WM}', the ring enlargement from IM_{DSA_WM}' to SO₃···H₂SO₄····H₂O complex occurred through transition state TS_{DSA_WM}' with a Gibbs free energy barrier of 1.2 kcal·mol⁻¹. Complex IM_{DSA_WM} was 6.1 kcal·mol⁻¹ lower in energy than IM_{DSA_WM}'. In IM_{DSA_WM}, SO₃ acted as double donors of hydrogen bond to form a cage-like hydrogen bonding network with H₂SO₄····H₂O. Then, starting with IM_{DSA_WM} with a Gibbs free barrier energy of 0.5 kcal·mol⁻¹ to form a quasi-planar network complex, H₂S₂O₇····H₂O. TS_{DSA_WM} was in the middle of a double proton transfer, where H₂O played as a bridge for proton transfer, along with

the simultaneous formation of the O4...S2 bond. In order to estimate the catalytic ability of H₂O in

265 the SO₃ + SA reaction, the effective rate constant $(k'_{DSA_WM_s})$ of the H₂SO₄···H₂O + SO₃ reaction were compared with the rate constant (k_{DSA}) of the SO₃ + H₂SO₄ reaction. As seen in Table 1, under the experimental concentration (Anglada et al., 2013) ([H₂O] = 5.20 × 10¹⁶-2.30 × 10¹⁸ molecules · cm⁻³) within the temperature range of 280-320 K, the calculated $k'_{DSA_WM_s}$ was 1.03 × 10^{-11} -4.60 × 10^{-12} cm³·molecule⁻¹·s⁻¹, which was larger by 1.79-1.86 times than that of k_{DSA} . This result shows that H₂O exerts catalytic role in promoting the rate of the SO₃ + H₂SO₄ reaction.

Regarding Channel SA_SA, the stepwise reaction occurred firstly via the ring enlargement from six-membered ring complex IM_{SA_SA} ' to a cage-like hydrogen bonding network IM_{SA_SA} , and then took place by going through a transition state, TS_{SA_SA} , to from the product complex $(H_2SO_4)_2$. TS_{DSA_WM} was in the middle of a double hydrogen transfer, where H_2SO_4 acted as a bridge of

- 275 hydrogen atom from the H₂O to SO₃ along with O1 atom of H₂O addition to the S atom of SO₃. It was worth noting that the energy barriers of two elementary reactions involved in the stepwise route of Channel SA_SA were only 1.8 and 0.6 kcal·mol⁻¹, respectively, showing that the stepwise route of Channel SA_SA is feasible to take place from energetic point of view. To check whether Channel DSA_WM is more favorable than Channel SA_SA or not, their rate ratio listed in Eq. 4 has been aclaulated in Table 1. The calculated rate ratio N
- calculated in Table 1. The calculated rate ratio v_{DSA_WM}/v_{SA_SA} shows that Channel DSA_WM is more important than Channel SA_SA because the rate ratio v_{DSA_WM}/v_{SA_SA} is 1.53-3.04 within the temperature range of 280-320 K. So, we predicted that the SO₃ + H₂SO₄ reaction with H₂O producing H₂S₂O₇ is more favorable than that forming H₂SO₄.

$$\frac{v_{\text{DSA_WM}}}{v_{\text{SA_SA}}} = \frac{v_{\text{DSA_WM_s}} + v_{\text{DSA_WM_o}}}{v_{\text{SA_SA_s}} + v_{\text{SA_SA_o}}} = \frac{k_{\text{DSA_WM_s}} \times K_{\text{eq(H_2SO_4 ••• H_2O)}} + k_{\text{DSA_WM_o}} \times K_{\text{eq(SO_3 ••• H_2O)}}}{k_{\text{SA_SA_s}} \times K_{\text{eq(H_2SO_4 ••• H_2O)}} + k_{\text{SA_SA_o}} \times K_{\text{eq(SO_3 ••• H_2O)}}}$$
(4)

3.2 Reactions at the air-water interface

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The mechanism for the $SO_3 + SA$ reaction at the air-water interface was lacking. Notably, SO_3 , SA and DSA molecules can stay at the interface for 35.8%, 30.1% and 39.2% of the time in the 150 ns simulation (Figure S2), respectively, revealing that the existence of SO₃, SA and DSA at the air-water interface cannot be negligible. So, the BOMD simulations were used to evaluate the reaction mechanism of SO₃ with SA at the aqueous interfaces. Similar with the interfacial reaction of SO₃ with organic acids (Cheng et al., 2023; Zhong et al., 2019), the reaction between SO₃

and SA at the aqueous interface may occur in three ways: (*i*) SO₃ colliding with adsorbed SA at the air-water interface; (*ii*) SA colliding with adsorbed SO₃ at the aqueous interface; or (*iii*) the SO₃-SA complex reacting at the aqueous interface. However, due to the high reactivity both of SO₃ and SA at the air-water interface, the lifetimes of SO₃ (Zhong et al., 2019) and SA (Figure S3) (on the order of a few picoseconds) on the water droplet were extremely short and can be formed SA⁻ ion quickly. Besides, as the calculated result above, SO₃····H₂SO₄ complex can be generate DSA easily before it approaches the air-water interface. So, two possible models were mainly considered for the SO₃ + SA reaction on the water surface: (*i*) gaseous SO₃ colliding with SA⁻ at the air-water interface and (*ii*) the DSA (the gas-phase product of SO₃ and SA) dissociating on water droplet.

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Gaseous SO₃ colliding with SA⁻ at the air-water interface. At the water droplet's surface, the interaction between SO₃ and SA⁻ included two main channels: (*i*) H₂O-induced formation of S₂O₇²⁻ ····H₃O⁺ ion pair (Figure 2, Figure S4 and Movie S1) and (*ii*) SA⁻-mediated formation of SA⁻····H₃O⁺ ion pair (Figure 3, Figs. S5-S6 and Movies S2-S3). The BOMD simulations for H₂O-induced formation of S₂O₇²⁻····H₃O⁺ ion pair was illustrated in Figure 2, the H1 atom of SA⁻ ion can combine with a nearby interfacial water molecule at 8.18 ps by hydrogen bond (d_(O3-H1) = 1.17 Å) interaction, thus forming hydrated hydrogen sulfate ion (SA⁻····H₂O). Then, the H1 atom of SA⁻ ion was moved to the O3 atom of the interfacial water molecule at 8.28 ps, revealing the formation of SO₄²⁻····H₃O⁺ ion pair. Additionally, SO₄²⁻ gradually approached to SO₃ molecule with the shortening of S1-O1
bond. At 9.26 ps, the S1-O1 bond length was 1.84 Å, which was close to the length of S-O1 (1.65

Å) bond in $S_2O_7^{2-}$ ion (Figure S8), revealing the formation of $S_2O_7^{2-}$... H_3O^+ ion pair.

Both direct (Figure 3(a), Figure S5 and Movie S2) and indirect (Figure 3(b), Figure S6 and Movie S3) forming mechanisms were observed in SA⁻-mediated formation of SA^{-…}H₃O⁺ ion pair. The direct SA⁻-mediated formation of SA^{-…}H₃O⁺ ion pair was a loop structure mechanism, which was consistent with gas phase hydrolysis of SO₃ assisted by acidic catalysts of HCOOH, HNO₃,

- was consistent with gas phase hydrolysis of SO₃ assisted by acidic catalysts of HCOOH, HNO₃, H₂C₂O₄ and SA in the previous works (Long et al., 2012; Long et al., 2013a; Torrent-Sucarrat et al., 2012; Lv et al., 2019), and the interfacial reactions of HNO₃-mediated Criegee hydration (Kumar et al., 2018) and the hydration of SO₃ via the loop-structure formation (Lv and Sun, 2020). As for the direct formation mechanism of SA⁻…H₃O⁺ ion pair seen in Figure 3(a) and movie S2, an eight-
- 320 membered loop complex, $SO_3 \cdots H_2O(1) \cdots SA^-$, was found at 1.46 ps with the formations of two hydrogen bonds ($d_{(O3 \cdots H2)} = 2.13$ Å; $d_{(O4 \cdots H3)} = 2.18$ Å) and a van der Waals interaction ($d_{(S1 \cdots O1)} =$

2.14 Å). Subsequently, SO₃ and interfacial $H_2O(1)$ were close to each other. At 1.59 ps, a transition state-like loop structure was observed and proton transfer from interfacial $H_2O(1)$ to another suspended $H_2O(2)$ was found, where the bond lengths of S1-O1, O1-H1 and H1-O2 were 1.94 Å,

325 1.19 Å and 1.32 Å, respectively. At 1.70 ps, the bond lengths of S-O1 and H1-O2 were reduced to 1.73 Å and 1.01 Å, while the bond length of H1-O2 was extended to 1.61 Å, showing the formation of SA⁻···H₃O⁺ ion pair. During the direct formation route of SA⁻···H₃O⁺ ion pair, SA⁻ played as a spectator, while interfacial water molecules acted as both a reactant and a proton acceptor. As compared with the hydration reaction mechanism of SO₃ at the air-water interface reported by Lv
330 et al. (Lv and Sun, 2020), the loop-structure formation with proton transferred in the loop was not observed in the direct mechanism of SA⁻····H₃O⁺ ion pair. This was

probably because SA⁻ ion was more difficult to give the proton.

As seen in Figure 3(b) and Movie S3, the indirect forming process of SA^{····}H₃O⁺ ion pair contained two steps: (*i*) SO₃ hydration along with SA formation and (*ii*) SA deprotonation. Specifically, as for step (*i*), at 0.70 ps, a transition state like structure of SO₃ hydration was observed with SO₃, SA⁻ and an interfacial water molecule involved. Note that at this time the H1 atom in interfacial H₂O molecule migrated to the O2 atom of SA⁻ ion instead of the surrounding water molecule. At 0.96 ps, the O1-H1 bond of H₂O was broken with the length of 1.56 Å, while the S1-O1 bond was formed with the length of 1.75 Å, demonstrating the completion of hydrolysis reaction of SO₃ and the formation of SA molecule. Then, at 8.08 ps, the H2 proton transferred from SA to the O4 atom of SA⁻ ion and to the O5 atom of the nearby water molecule was occurred, where the O3-H2 and O1-H3 bonds extended to 1.13 Å and 1.22 Å, and the length of O4-H2 and O5-H3 bonds shortened to 1.45 Å and 1.20 Å. Finally, SA deprotonation was completed at 8.23 ps with the formation of SA⁻···H₃O⁺ ion pair. During the whole indirect forming process of SA^{-···}H₃O⁺ ion pair,

SA⁻ played as protons donor and acceptor, and water molecules acted as hydration reactants and proton acceptors. Compared with the direct mechanism of SA⁻···H₃O⁺ ion pair, the indirect forming process of HSO_4^{-} ···H₃O⁺ ion pair required more time. This was consistent with the interfacial reactions of CH₂OO + HNO₃ (Kumar et al., 2018) and the hydration of SO₃ (Lv and Sun, 2020) where the direct forming mechanism needed less time than indirect forming mechanism.

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The $H_2S_2O_7$ *dissociating on water droplet.* In addition to the gaseous SO₃ colliding with SA⁻ at the air-water interface, DSA, the product of the barrierless reaction between SO₃ and SA, can

further quickly react with interfacial water molecule at the air-water interface. As seen in Figure 4, Figure S7 and Movie S4, DSA was highly reactive at the air-water interface and can undergo two deprotonations to form S₂O₇²⁻ ion. Specifically, the DSA can firstly form a H-bond with interfacial water and produced HS₂O₇⁻ and H₃O⁺ ions. The formed HS₂O₇⁻ ion can survive for ~3 ps on water droplet. At 4.14 ps, the H2 atom of HS₂O₇⁻ ion moved to O4 atom of nearby interfacial water molecule and produced the formation of S₂O₇²⁻…H₃O⁺ ion pair, which was stable at the air-water interface over a simulated time scale of 10 ps. Note that the second deprotonation of DSA indeed needs more time than its first deprotonation as the pK_a1 (pK_a1 = -16.05) of DSA is much smaller than its pK_a2 (pK_a2 = -4.81) (Abedi and Farrokhpour, 2013). In brief, at the air-water interface, both these two routes of the formation of S₂O₇²⁻…H₃O⁺ ion pair occurred on the picosecond time scale.

3.3 Atmospheric implications

The application of the $SO_3 + SA$ *reaction in atmospheric chemistry.* In the gas-phase, the 365 main sink route of SO₃ was H₂O-assisted hydrolysis of SO₃ (Morokuma and Muguruma, 1994; Akhmatskaya et al., 1997; Larson et al., 2000; Hazra and Sinha, 2011; Long et al., 2013a; Torrent-Sucarrat et al., 2012; Ma et al., 2020). To study the atmospheric importance of the SO₃ + SA reaction without and with H₂O, the rate ratio (v_{DSA}/v_{SA}) between the SO₃ + SA reaction and H₂O-assisted hydrolysis of SO₃ was compared, which was expressed in Eq. (5).

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$$\frac{v_{\text{DSA}}}{v_{\text{SA}}} = \frac{k_{\text{DSA}} \times [\text{SO}_3] \times [\text{H}_2 \text{SO}_4] + k_{\text{DSA}_\text{WM}_s} \times \text{K}_{\text{eq1}} \times [\text{SO}_3] \times [\text{H}_2 \text{SO}_4] \times [\text{H}_2 \text{O}]}{k_{\text{SA}_\text{WM}} \times \text{K}_{\text{eq2}} \times [\text{SO}_3] \times [\text{H}_2 \text{O}] \times [\text{H}_2 \text{O}]}$$
(5)

In Eq. (5), K_{eq1} and K_{eq2} were the equilibrium constant for the formation of H₂SO₄···H₂O and SO₃····H₂O complexes shown in Table S2, respectively; *k*_{DSA}, *k*_{DSA_WM_s} and *k*_{SA_WM} were respectively denoted the bimolecular rate coefficient for the H₂SO₄ + SO₃, H₂SO₄····H₂O + SO₃ and SO₃····H₂O + H₂O reactions; [H₂O] and [H₂SO₄] were respectively represented the concentration of H₂O and SA taken from references (Anglada et al., 2013; Liu et al., 2015); The value of *v*_{DSA}/*v*_{SA} was listed in Table S7 (0 km altitude) and Table S8 (5-30 km altitude). As seen in Table S7, the hydrolysis reaction of SO₃ with (H₂O)₂ dominates over the SO₃ + H₂SO₄ reaction at 0 km altitude as the [H₂O] (10¹⁶-10¹⁸ molecules[•]cm³) was much larger than that of [H₂SO₄] (10⁴-10⁸ molecules[•]cm³). Although the concentration of water molecules decreased with the increasing of altitude in Table S8, the concentration of [H₂O] was still much greater than that of [H₂SO₄], resulting

in the SO₃ + H₂SO₄ reaction cannot compete with H₂O-assisted hydrolysis of SO₃ within the altitude range of 5-30 km. Even considering of high H₂SO₄ concentration at the end and outside the aircraft engine and flight at 10 km (Curtius et al., 2002), the SO₃ + H₂SO₄ reaction was not the major sink route of SO₃. Notably, as the concentration of sulfuric acid was even greater than that of water vapor in the atmosphere of Venus, the SO₃ + SA reaction was probably favorable than the H₂O-assisted hydrolysis of SO₃ in the Venus' atmosphere. To check whether the SO₃ + H₂SO₄ reaction was more favorable than H₂O-assisted hydrolysis of SO₃ or not in the Venus' atmosphere, the rate ratio of v_{DSA}/v_{SA} listed in Eq. 4 has been calculated in Table 2. It can be seen from Table 2 that the rate ratio of v_{DSA}/v_{SA} was 3.24×10^8 -5.23 $\times 10^{10}$ within the altitude range of 40-70 km in the Venus' atmosphere, which indicates that the SO₃ + H₂SO₄ reaction is significantly more favorable than the hydrolysis reaction of SO₃ + (H₂O)₂ within the altitudes range of 40-70 km in the Venus' atmosphere.

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Enhancement effect of DSA on NPF. From the multistep global minimum sampling technique, for $(DSA)_x(SA)_y(A)_z$ ($z \le x + y \le 3$) molecular clusters, 27 most stable structures in the present system have been found (Figure S11). To evaluate the thermodynamic stability of these clusters, Gibbs formation free energies (ΔG) at 278.15 K and evaporation rate coefficient (y, s⁻¹) for 395 $(DSA)_x(SA)_y(A)_z$ ($z \le x + y \le 3$) molecular clusters were calculated in Figure 5 and Table S11-12, respectively. As for dimers formed by SA, A and DSA, the ΔG of $(A)_1 \cdot (DSA)_1$ was -16.1 kcal·mol⁻ ¹, which was lowest in all dimers followed by $(SA)_2$ (-8.5 kcal·mol⁻¹) and then $(SA)_1$ ·(A)₁ (-6.3 kcal·mol⁻¹), meanwhile, the γ of (A)₁·(DSA)₁ (1.17 × 10⁻³ s⁻¹) was lower than those of (SA)₂ (3.81 400 $\times 10^2$ s⁻¹) and (SA)₁·(A)₁ (4.19 $\times 10^4$ s⁻¹). Regarding for the SA-A-DSA-based clusters, the values of ΔG and γ of SA-A-DSA-based clusters containing more DSA molecules were relatively lower than the corresponding values of other SA-A-DSA-based clusters with the same number of acid and base molecules. In the free-energy diagram for cluster formation steps of the SA-A-DSA system (Figure 5), thermodynamic barriers were weakened mainly by the subsequential addition of A or 405 DSA monomer. Also, the SA-A-DSA-based growth pathway was thermodynamically favorable with decreasing ΔG . These results indicate that DSA not only can promote the stability of SA-A-DSA-based clusters but also may synergistically participate in the nucleation process.

The potential enhancement influence of DSA to the SA-A-based particle formation was shown in Figure 6. The formation rate $(J, \text{cm}^{-3} \cdot \text{s}^{-1})$ of SA-A-DSA-based system illustrated in Figure 6 was negatively dependent on temperature, demonstrating that the low temperature is a key factor to accelerate cluster formation. It is noted that, at low temperatures of 218.15 K (Figure S12) and 238.15 K (Figure S13), the actual ΔG of clusters has been calculated to ensure meaningful cluster dynamics of the 3 × 3 systems, where the actual ΔG surface represented that the simulated set of clusters always included the critical cluster. In addition to temperature, the *J* of SA-A-DSA-based system shown in Figure 6 rise with the increase of [DSA]. More notably, the participation of DSA can promote *J* to a higher level, indicating its enhancement on SA-A nucleation. Besides, there was significantly positive dependence of the *J* of SA-A-DSA-based system on both [SA] and [A] in Figure 7 (238.15 K) and Figure S15-Figure S18 (218.15, 258.15, 278.15 and 298.15 K). This was because the higher concentration of nucleation precursors could lead to higher *J*. Besides, Figure S19 showed the nucleation rate when the sum ([SA] + [DSA]) was kept constant. *J*_{DSA/SA} at substituted condition was higher than that at unsubstituted condition. These results indicated that DSA may can greatly enhance the SA-A particle nucleation in heavy sulfur oxide polluted atmospheric boundary layer, especially at an average flight altitude of 10 km with high [DSA].

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- Two main cluster formation pathways, the pure SA-A-based cluster (i) and DSA-containing 425 cluster (ii), at different [DSA] and different temperatures (218.15 K, 238.15 K and 258.15 K) were shown in Figure 8(a). As seen, the DSA molecule exhibited an ability to directly participate in cluster formation under high [SA] and [DSA], and median [A]. Interestingly, at different temperature and different [DSA], the DSA molecule showed different effect mechanism and contribution in SA-A system. As seen in Figure 8(b) and Figure S20(b), the cluster growth pathways 430 were dominated by DSA-containing cluster formation under the conditions of 238.15 K ([DSA] is 10⁶-10⁷ molecules cm⁻³), 258.15 K ([DSA] is 10⁵-10⁷ molecules cm⁻³), 278.15 K ([DSA] is 10⁴-10⁷ molecules \cdot cm⁻³) and 298.15 K ([DSA] is 10⁴-10⁷ molecules \cdot cm⁻³). By the way, the cluster growth pathways were completely dominated by the DSA-containing cluster at 298.15 K where [DSA]= 10^{5} - 10^{7} molecules cm⁻³, and its contribution for growth flux out of the system reached to 100% 435 (Figure S22). In short, on one hand, the contribution of the DSA participation pathway has been increased with increasing temperature. On the other hand, the contribution of the pathway with participation of DSA increased with increasing [DSA], while the number of DSA molecules contained in clusters $[(SA)_2 \cdot (A)_3 \cdot DSA, SA \cdot (A)_2 \cdot DSA, SA \cdot (A)_3 \cdot (DSA)_2, and (A)_3 \cdot (DSA)_3]$ that can contribute to cluster growth had a positive correlation with [DSA]. These results suggested that DSA
- 440 has the ability to act as a potential contributor to SA-A-based NPF in the atmosphere at low *T*, low

[SA], high [A] and high [DSA], and the DSA participation pathway can be dominant in heavy sulfur oxide polluted atmospheric boundary layer and in season of late autumn and early winter.

At the air-water interface, important implication of the BOMD simulations was that the reaction between SO₃ and SA at the air-water interface can be accomplished within a few picoseconds, among which the interfacial water molecules played a significant role in promoting 445 the formation of $S_2O_7^{2-}$... H_3O^+ and SA^- ... H_3O^+ ion pairs. Furthermore, the adsorption capacity of the $S_2O_7^{2-}$, H_3O^+ and SA^- to gasous precursors in the atmosphere was further investigated by the calculated interaction free energies. Herein, the species of SA, NH₃, HNO₃ and (COOH)₂ have been regarded as the candidate species. (Kulmala et al., 2004; Kirkby et al., 2011). Our calculated Gibbs free energies in Table 3 showed that the interactions of S2O72-...H2SO4, S2O72-...HNO3, S2O72-450 ···(COOH)₂, H₃O⁺···NH₃, H₃O⁺···H₂SO₄, SA⁻···H₂SO₄, SA⁻···(COOH)₂, and SA⁻···HNO₃ were stronger than those of H₂SO₄...NH₃ (major precursor of atmospheric aerosols) with their binding free energies enhanced by 18.6-42.8 kcal·mol⁻¹. These results reveal that interfacial $S_2O_7^{2-}$, SA⁻ and H₃O⁺ can attract candidate species from the gas phase to the water surface. Moreover, we evaluated whether $S_2O_7^{2-}$ could lead to increased particle growth on SA-A cluster by considering geometrical 455 structure and the formation free energies of the $(SA)_1(A)_1(S_2O_7^{2-})_1$ clusters. As compared with $(SA)_1(A)_1(X)_1$ (X = HOOCCH₂COOH, HOCCOOSO₃H, CH₃OSO₃H, HOOCCH₂CH(NH₂)COOH and HOCH₂COOH) clusters (Zhong et al., 2019; Zhang et al., 2018; Rong et al., 2020; Gao et al., 2023; Liu et al., 2021a; Zhang et al., 2017), the number of hydrogen bonds in $(SA)_1(A)_1(S_2O_7^{2-})_1$ 460 cluster presented in Figure S8 increased and the ring of the complex was enlarged. It was demonstrated that $S_2O_7^{2-}$ has the highest potential to stabilize SA-A clusters and promote SA-A nucleation in these clusters due to its acidity and structural factors such as more intermolecular hydrogen bond binding sites. Subsequently, comparing to $(SA)_1(A)_1(X)_1$ clusters (Table 2), the Gibbs formation free energy ΔG of $(SA)_1(A)_1(S_2O_7^{2-})_1$ cluster was lower, showing $S_2O_7^{2-}$ ion at the air-water interface has stronger nucleation ability than X in the gas phase. Therefore, we predict that 465 $S_2O_7^{2-}$ at the air-water interface would lead to increased particle growth.

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4 Summary and conclusions

In this work, we employed QC calculations, BOMD simulations and ACDC kinetic model to characterize the SO_3 -H₂SO₄ interaction in the gas phase and at the air-water interface and to study

- 470 the effect of $H_2S_2O_7$ on H_2SO_4 -NH₃-based clusters. Results revealed that the energy barrier of the gas phase $SO_3 + H_2SO_4$ reaction without and with H_2O was less than 2.3 kcal·mol⁻¹. Rate constants indicated that though the $SO_3 + H_2SO_4$ reaction cannot compete with H_2O -assisted hydrolysis of SO_3 within the temperature range of 280-320 K, its rate constant was close to the upper limits for bimolecular reactions and H_2O exerted obvious catalytic role in promoting the reaction rate.
- 475 Moreover, ACDC kinetic simulations showed that DSA has unexpected facilitate effects on the NPF process and can present a more obvious enhancement effect on SA-A-based cluster formation in polluted atmospheric boundary layer. Of particular note, DSA can directly participate in the SA-A-based cluster formation pathway and the contribution of the pathway with participation of DSA increases with increasing [DSA] in regions with atmospheric pollution boundary layer of high concentrations of SO₃, especially in late autumn and early winter.

At the air-water interface, H₂O-induced the formation of $S_2O_7^{2-\cdots}H_3O^+$ ion pair, SA⁻ mediated the formation of SA⁻…H₃O⁺ ion pair and the deprotonation of H₂S₂O₇ were observed, both of which can occur within a few picoseconds. The formed interfacial $S_2O_7^{2-}$, SA⁻ and H_3O^+ can attract candidate species (such as H₂SO₄, NH₃, and HNO₃) for particle formation from the gas phase to the water surface, and thus accelerated the growth of particle. Moreover, potential of X ($X = S_2O_7^{2-}$, HOOCCH₂COOH, HOCCOOSO₃H, CH₃OSO₃H, HOOCCH₂CH(NH₂)COOH and HOCH₂COOH) in ternary SA-A-*X* cluster formation indicated that $S_2O_7^{2-}$ has the highest potential to stabilize SA-A clusters and promote SA-A nucleation among *X*.

The present work will expand our understanding of new pathway for the loss of SO₃ in acidic polluted areas. Moreover, this work will also help to reveal some missing sources of metropolis industrial regions NPF and to understand the atmospheric organic-sulfur cycle more comprehensively.

Data availability

All data presented in this study are available upon request from the corresponding author.

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Declaration of competing interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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Table 1. The rate constant (cm³·molecule⁻¹·s⁻¹) for the SO₃ + H₂SO₄ reaction and the effective rate constant (cm³·molecule⁻¹·s⁻¹) for the SO₃ + H₂SO₄ reaction with H₂O (100%RH) within the temperature range of 280-320 K

<i>T/</i> (K)	280 K	290 K	298 K	300 K	310 K	320 K
$k_{\rm DSA}$	5.52×10^{-12}	4.60×10^{-12}	3.95×10^{-12}	3.80×10^{-12}	3.13 × 10 ⁻¹²	2.57×10^{-12}
$k'_{ m DSA_WM_o}$	2.12×10^{-13}	2.68×10^{-13}	2.88×10^{-13}	2.89×10^{-13}	2.89×10^{-13}	2.75×10^{-13}
<i>k'</i> dsa_wm_s	1.03 × 10 ⁻¹¹	8.55 × 10 ⁻¹²	7.42×10^{-12}	7.11 × 10 ⁻¹²	5.79 × 10 ⁻¹²	4.60 × 10 ⁻¹²

 k_{DSA} is the rate constant for the SO₃ + H₂SO₄ reaction; $k'_{\text{DSA}_WM_o}$ and $k'_{\text{DSA}_WM_s}$ are respectively the effective rate constants for H₂O-assisted SO₃ + H₂SO₄ reaction occurring through one-step and stepwise routes.

unitides in the autosphere of venus								
H (km)	<i>T</i> (K)	P (Torr)	[H ₂ O]	$[H_2SO_4]$	$k_{\rm DSA}$	$k_{DSA_WM_s}$	$k_{\rm SA_WM}$	$v_{\rm DSA}/v_{\rm SA}$
40	410	2025	1.08×10^{15}	6.15×10^{13}	5.22×10^{-12}	1.43×10^{-12}	2.31×10^{-13}	3.24×10^{8}
50	340	750	5.17×10^{14}	1.23×10^{14}	$1.12\times10^{\text{-}12}$	$3.87\times10^{\text{-}12}$	5.43×10^{-13}	3.81×10^{10}
60	320	104	1.72×10^{14}	1.85×10^{14}	1.23×10^{-12}	7.80×10^{-12}	1.37×10^{-12}	5.12×10^{10}
70	270	19	8.61×10^{13}	8.61×10^{13}	1.07×10^{-12}	8.61×10^{-12}	1.82×10^{-12}	5.23×10^{10}

Table 2. The rate ratio between the $SO_3 + H_2SO_4$ reaction and the hydrolysis of SO_3 at different altitudes in the atmosphere of Venus

 k_{DSA} , $k_{\text{DSA}_WM_s}$ and k_{SA_SA} are respectively the rate constants for SO₃ + H₂SO₄ reaction, H₂O-assisted SO₃ + H₂SO₄ reaction occurring through stepwise route and the hydrolysis reaction of SO₃ + (H₂O)₂.

Table 3. Gibbs free energy (ΔG , kcal·mol⁻¹) for the formation of S₂O₇²⁻····H₂SO₄, S₂O₇²⁻···HNO₃,S₂O₇²⁻···(COOH)₂, H₃O⁺····NH₃, H₃O⁺····H₂SO₄, HSO₄⁻····(COOH)₂, HSO₄⁻···(COOH)₂, HSO₄⁻···(COOH)₂, HSO₄⁻····(COOH)₃,HNO₃, H₂SO₄····NH₃, SO₇²⁻····H₂SO₄····NH₃, HOOCCH₂COOH····H₂SO₄····NH₃,HOCCOOSO₃H····H₂SO₄····NH₃, CH₃OSO₃H····H₂SO₄····NH₃ andHOOCCH₂CH(NH₂)COOH····H₂SO₄····NH₃ at 298 K

	$S_2O_7^2$ -····H ₂ SO ₄	$S_2O_7^2$ -···HNO ₃	S ₂ O ₇ ²⁻ ···(COOH) ₂	$H_3O^+\cdots NH_3$	H_2SO_4 ···N H_3
ΔG	-46.3	-30.6	-39.9	-51.7 (-49.2) ^a	-8.9 (-8.9)ª
	$H_3O^+\cdots H_2SO_4$	HSO4H2SO4	HSO ₄ (COOH) ₂	HSO4 ⁻ ··· HNO3	$S_2O_7^{2-}$ ····H_2SO_4····NH_3
ΔG	-27.5 (-27.0) ^a	-41.6	-33.6	-27.8	-40.1
	HOOCCH ₂ COOH …H ₂ SO ₄ …NH ₃	HOCCOOSO ₃ H …H ₂ SO ₄ …NH ₃	CH ₃ OSO ₃ H …H ₂ SO ₄ …NH ₃	HOOCCH ₂ CH(NH ₂)COOH …H ₂ SO ₄ …NH ₃	HOCH ₂ COOH …H ₂ SO ₄ …NH ₃
ΔG	-13.1 (-13.6) ^b	-20.4 (-22.5)°	-18.8 (-20.7) ^d	-13.2 (-14.0) ^e	-12.8 (-13.5) ^f

Energies are given in kcal·mol⁻¹, and calculated at the M06-2X/6-311++G(2*df*,2*pd*) theoretical level. References are as follows: [a] Zhong et al., 2019.; [b] Zhang et al., 2018.; [c] Rong et al., 2020.; [d] Gao et al., 2023.; [e] Liu et al., 2021a; [f] Zhang et al., 2017.



Figure 1. Schematic potential energy surface for the $SO_3 + H_2SO_4 \rightarrow H_2S_2O_7$ reaction; Distances is in angstrom at the M06-2X/6-311++G(2*df*,2*pd*) level, while the energy values correspond to the calculations at the CCSD(T)-F12/cc-pVDZ-F12//M06-2X/6-311++G(2*df*,2*pd*) level. The pre-reactive complex and TS for the route of DSA formation from the SO₃ + H₂SO₄ reaction with H₂O was denoted by "IM_{DSA_WM}" and "TS_{DSA_WM}", respectively, while the corresponding pre-reactive complex and TS for the process of SA formation from the hydrolysis of SO₃ with H₂SO₄ was respectively labeled as "IM_{SA_SA}" and "TS_{SA_SA}".



Figure 2. Top panel: Snapshot structures taken from the BOMD simulations, which illustrate H₂Oinduced the formation of $S_2O_7^{2-}$ ···H₃O⁺ ion pair from the reaction of SO₃ with HSO₄⁻ at the air-water interface. Lower panel: time evolution of key bond distances (S-O1, O2-H1, and O3-H1) involved in the induced mechanism.



Figure 3. Top panel: Snapshot structures taken from the BOMD simulations, which illustrate the hydration reaction mechanism of SO_3 mediated by HSO_4^- at the air water interface. Lower panel: time evolution of key bond distances (S-O1, O1-H2, O5-H2, O2-H1, O3-H4 and O4-H3) involved in the hydration mechanism.



Figure 4. Top panel: Snapshot structures taken from the BOMD simulations, which illustrate the deprotonation of $H_2S_2O_7$ at the air water interface. Lower panel: time evolution of key bond distances (O1-HI, O1-H2, O3-H2 and H2-O4) involved in the hydration mechanism.



Figure 5. The Gibbs free energy (kcal·mol⁻¹) diagram of $(DSA)_x(SA)_y(A)_z$ ($z \le x + y \le 3$) clusters at 278.15K and 1 atm. "A" refers to sulfuric acid, "D" refers to disulfuric acid and "N" refers to ammonia.



Figure 6. Cluster formation rates J (cm⁻³ s⁻¹) against the of DSA monomer concentration (unit: molecules·cm⁻³) under different temperatures (218.15, 238.15, 258.15, 278.15 and 298.15 K) where [SA] = 10⁷ molecules·cm⁻³ and [A] = 10⁹ molecules·cm⁻³.



Figure 7. Simulated cluster formation rates J (cm⁻³ s⁻¹) as a function of (a) [SA], (b) [A], with different concentrations of disulfuric acid [DSA] of 10^4 (red), 10^5 (blue), 10^6 (green) , 10^7 (purple) and 0 molecules cm⁻³ (black, pure-SA-A), at T = 238.15 K.



Figure 8. (a) The main pathways of clusters growing out of the research system under the conditions where 218.15 K,238.15K and 258.15 K, $[SA] = 10^8$ molecules \cdot cm⁻³, $[A] = 10^9$ molecules \cdot cm⁻³, and $[DSA] = 10^6$ molecules \cdot cm⁻³; (b) The contribution of different concentrations of DSA to the main cluster formation pathway at 218.15 K, 238.15 K and 258.15 K is shown in the pie charts.