Influence of acidity on liquid–liquid phase transitions of mixed SOA proxy–inorganic aerosol droplets

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13 Abstract. Phase state and morphology of aerosol particles play a critical role in determining their effect on climate. While aerosol acidity has been identified as a key factor affecting the multiphase chemistry and phase transitions, the impact of 14 15 acidity on phase transition of multicomponent aerosol particles has not been extensively studied in situ. In this work, we 16 employed an aerosol optical tweezer (AOT) to probe the impact of acidity on the phase transition behavior of levitated aerosol 17 particles. Our results revealed that higher acidity decreases the separation relative humidity (SRH) of aerosol droplets mixed 18 with ammonium sulfate (AS) and secondary organic aerosol (SOA) proxy, such as 3-methylglutaric acid (3-MGA), 1,2,6-19 hexanetriol (HEXT) and 2,5-hexanediol (HEXD) across aerosol pH in atmospheric condition. Phase separation of organic 20 acids was more sensitive to acidity compared to organic alcohols. We found the mixing relative humidity (MRH) was 21 consistently higher than the SRH in several systems. Phase-separating systems, including 3-MGA/AS, HEXT/AS, and 22 HEXD/AS, exhibited oxygen-to-carbon ratios (O:C) of 0.67, 0.50, and 0.33, respectively. In contrast, liquid-liquid phase 23 separation (LLPS) did not occur in the high O:C system of glycerol/AS, which had an O:C of 1.00. Additionally, the 24 morphology of 42 out of the 46 aerosol particles that underwent LLPS was observed to be a core-shell. Our findings provide 25 a comprehensive understanding of the pH-dependent LLPS in individual suspended aerosol droplets and pave the way for 26 future research on phase separation of atmospheric aerosol particles.

27 1 Introduction

28 Atmospheric aerosol particles can directly and indirectly impact climate by absorbing and scattering light and acting as cloud condensation nuclei (Rosenfeld et al., 2014). Particle morphology is a critical factor influencing the physiochemical properties 29 30 of aerosols such as their optical properties, chemistry, and nucleation processes (Freedman et al., 2009; Corral Arroyo et al., 31 2022; Cosman et al., 2008; Lam et al., 2021; Petters and Kreidenweis, 2007; Mikhailov et al., 2021). Morphology can be 32 broadly categorized into single-phase homogeneous morphology and phase separation morphology (Bertram et al. 2011; 33 Ciobanu et al. 2009), based on the phase state of the particle. For droplets with a phase separation morphology, the two main 34 equilibrium morphologies are a fully engulfed (core-shell) structure and a partially engulfed structure (Reid et al. 2011). 35 Droplets can undergo phase transition processes and thus the morphology would be changed. The composition and mass of 36 inorganic and organic components impact the phase transition characteristics of a particle. With a decrease of particle water 37 content, a transition occurs from single homogenous liquid phase to two separated liquid phases, which is known as liquidliquid phase separation (LLPS; Freedman et al., 2017). The relative humidity (RH) when the LLPS occurs is defined as 38 39 separation relative humidity (SRH). The reverse process, in which two liquid phases mix into a single homogenous liquid

40 phase, is referred to as liquid-liquid phase mixing and the corresponding RH is the mixing RH (MRH; You et al., 2014;

- 41 Gorkowski et al., 2017).
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43 The phenomenon of LLPS has garnered considerable attention from the atmospheric research community due to its potential 44 role in affecting the physiochemical properties of atmospheric aerosols (Kucinski et al., 2019; Ott et al., 2020; Freedman, 45 2020). Song et al. (2012) using optical microscopy studied the relationship between LLPS and the oxygen-to-carbon ratio 46 (O:C) and discovered that LLPS was consistently observed when O:C < 0.56, while it was never observed when O:C > 0.80. 47 For O:C between 0.56 and 0.80, the occurrence of LLPS was influenced by the types of organic functional groups. Gorkowski 48 et al. (2020) utilized experimental results of previous studies on LLPS and morphology, observing a general trend in 49 morphology from partially engulfed to core shell and finally homogeneous as oxidation increased. More recently, it is found 50 that submicrometer-sized aerosol particles had a lower SRH compared to micrometer-sized droplets (Kucinski et al., 2021; 51 Ohno et al., 2021). Meanwhile, Stewart et al. (2015) employed aerosol optical tweezer (AOT) to investigate the morphologies 52 of aqueous droplets. They found in the polyethylene glycol (PEG)/ammonium sulfate (AS) system, droplets formed 53 predominately core-shell particles when the AS content was high and partially engulfed when the PEG content was high.

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55 One factor that could influence the phase transitions of aerosol particles is the aerosol pH. The pH values for misty cloud and 56 fog droplets generally range between 2 and 7, whereas continental and marine aerosol particles exhibit a wider range of pH 57 values, from -1 to 5 and 0 to 8, respectively (Pye et al., 2020; Angle et al., 2021; Weber et al., 2016; Tilgner et al., 2021; Zheng 58 et al., 2020). Meanwhile, aerosol pH is size-dependent, with the fine mode showing lower 1-4 pH units than the coarse mode 59 (Fang et al., 2017; Young et al., 2013; Guo et al., 2017). Losey et al. (2018) studied six organic components and discovered 60 that phase separation may be hindered by the addition of sulfuric acid, while the SRH of 3-methylglutaric acid/ammonium 61 sulfate system was found to decrease with the addition of sodium hydroxide (Losey et al. 2016), as the deprotonation of organic 62 component or difference in salting out ability of inorganic may change the SRH. More recently, Tong et al. (2022) investigated 63 the effect of acidity on phase separation in single suspended microdroplets using AOT. Their results showed that the pH can 64 affect the miscibility of the mixture and high acidity results in a reduced SRH of 1,2,6-hexanetriol.

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66 Our aim with this work is to gain a comprehensive understanding of the influence of pH on phase transitions in suspended 67 droplets. To that end, we investigated pH-dependent SRH and MRH, as well as morphologies of aqueous droplets using AOT, 68 meanwhile discussed the effect of O:C on phase separation behavior. Compared to substrate-based measurement techniques, 69 AOT can suspend droplets without any substrate contact, providing a more realistic simulation of the behavior of aerosols in 70 the atmosphere (Wang et al., 2021; Cui et al., 2021; Redding et al., 2015; Gong et al., 2018; Rafferty et al., 2023). We measured 71 droplets containing AS and a range of organic compounds with varying O:C. Our findings provide insight into the mechanisms 72 behind pH-dependent phase transitions in levitated droplets, and have implications for fields such as climate science. Overall, 73 our study highlights the importance of considering pH as a key factor in the phase transition behavior of micron-sized droplets 74 and underscores the need for further research to fully understand the complex interactions between pH and phase transitions 75 in these atmospherically relevant systems.

76 2 Methods

77 2.1 Aerosol generation

Four organics components: glycerol (GL), 3-methylglutaric acid (3-MGA), 1,2,6-hexanetriol (HEXT), and 2,5-hexanediol (HEXD), were chosen as they are commonly-used secondary organic aerosol (SOA) proxies (Lam et al., 2021; Gorkowski et al., 2020). O:C of the selected chemicals varied from 1 to 0.33 (**Table 1**), which is similar to the real atmospheric SOA

81	(Canagaratna et al., 2015; Mahrt et al., 2021). AS was chosen as the inorganic salt component due to its widespread occurrence
82	in the atmospheric environment. All concentrations of organics and AS in the mother solutions were 50 g/L. The pure organic
83	and inorganic components were dissolved in ultrapure water (Millipore, resistivity of 18.2 M Ω) to create solutions with
84	organic-to-inorganic mass ratio (OIR) of 1:1. The pH of studied systems were adjusted within the range of 0.48 to 6.53 by
85	using either concentrated sulfuric acid (SA) or sodium hydroxide (NaOH) solution (5.29 mol/L). Sodium hydroxide, a strong
86	base, allowed for pH adjustment with minimal usage (Losey et al., 2016). However, it is necessary to acknowledge that the
87	addition of NaOH changed the composition of the inorganic part of the solution, potentially affecting the SRH values measured.
88	The pH of each solution was measured using a pH meter (Mettler Toledo Instruments Co., Ltd., Shanghai, China). The purity
89	and supplier of the compounds used in this study are summarized in Table S1 .

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- 91

Table 1. Information of the solutions used to generate aerosol droplets.

Solution ID	Organic component	O:C ratio	pН
GL	glycerol	1.00	5.24±0.01
3-MGA-I			0.48 ± 0.01
3-MGA-II			1.19 ± 0.01
3-MGA-III	3-methylglutaric acid	0.67	2.70 ± 0.01
3-MGA-IV		0.07	3.70 ± 0.01
3-MGA-V			5.21 ± 0.02
3-MGA-VI			6.53 ± 0.02
HEXT-I			$0.92{\pm}0.01$
HEXT-II	1,2,6-hexanetriol	0.50	2.02 ± 0.01
HEXT-III	1,2,0-ilexallet101	0.50	3.14 ± 0.01
HEXT-IV			5.11 ± 0.02
HEXD-I			1.39 ± 0.01
HEXD-II			$2.03{\pm}0.01$
HEXD-III	2,5-hexanediol	0.33	2.71±0.01
HEXD-IV			3.13±0.01
HEXD-V			5.01 ± 0.01

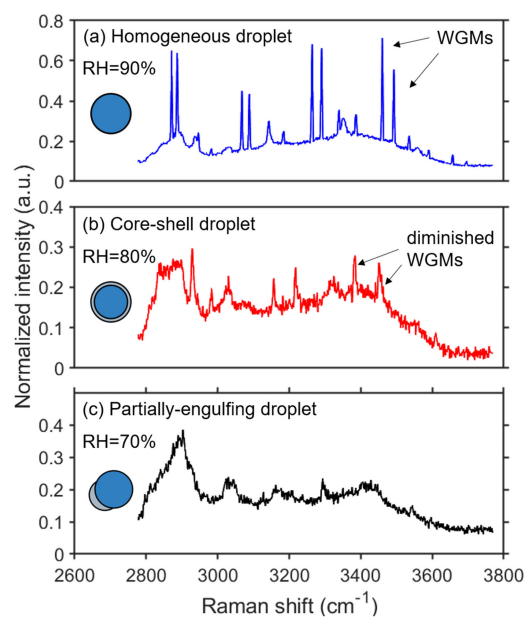
93 2.2 Experimental setup

94 A schematic illustration of the experimental setup is presented in Fig. S1. The aerosol optical tweezer system consists of a 95 custom-made levitation chamber that integrates the optical trapping system, the illumination and imaging system, and the aerosol generation system. A 532 nm (Opus 532-2W) laser was used to create an optical trap with a 100x oil immersion 96 97 objective (Olympus, UPLFLN100XO, NA 1.30) pressed against a glass coverslip (Nest, thickness 160-190 µm). The illumination and imaging system includes a 450 nm LED (Daheng Optics, GCI060404) and a camera (Thorlabs, CS165CU/M) 98 99 to illuminate and image the particle. Two low pass filters (Andover, 500FL07-25) were used in front of the camera lens to 100 remove the influence of back scattered light of the 532 nm laser to photograph clear image of the particle. The Raman scattered 101 light passed through two 50:50 beam splitters (CVI Laser Optics, BTF-VIS-50-2501M-C) and a notch filter (Semrock, NFD01-102 532-25x36) and was focused into the Raman spectrograph. A spectrograph (ZOLIX, Omni- λ 5004i) was used to measure the 103 Stokes shifted Raman spectrum. A 20 µm entrance slit width and 1200 groove/mm diffraction grating with a blaze wavelength 104 of 500 nm were used to achieve a spectral resolution of 0.021 nm. The wavelength position of spectrograph was calibrated 105 with a Hg-laser. The Raman scattered light was recorded every 4 seconds with range of 624.24-665.40 nm.

107 As droplets are introduced continuously into the chamber from a medical nebulizer (LANDWIND, PN100), smaller droplets 108 undergo a process of collision and coalescence, leading to the formation of larger droplets that can be readily trapped near the 109 focal point of the laser. In most cases, droplets can be successfully captured within 30 s after the introduction of an aerosol plume into the cell. Air with relative humidity (RH) of 100% and 0% were mixed to produce wet air with a desired RH. The 110 111 flow rates of the humidified and dry air streams were regulated by mass flow controllers (MFCs, Tianjin Gastool Instruments Co., Ltd., Tianjin, China, GT130D), with a combined flow rate of 0.3 L/min in total. Two humidity sensors (Sensirion, SHT85) 112 113 were utilized, with a precision of $\pm 1.5\%$. Since the sensor located behind the chamber was positioned in close proximity (~80 114 mm) to the droplet, its observed values were used as a surrogate for measuring the RH inside the chamber. The RH values 115 were reduced in increments of 5% every 30 minutes (Tong et al., 2022; Stewart et al., 2015) until droplet phase separation 116 occurred. The measured values of RH given by the sensors were used as the phase separation RH. Subsequently, the RH level 117 was set to 100%, to investigate the phase mixing of the droplets. The entire experiment was repeated 2-4 times for each system.

118 **2.3 Determination of phase transitions**

119 When a transparent or weakly absorbing spherical particle is trapped, it can behave as a high-quality factor optical cavity that 120 supports sharp optical resonances, resulting in cavity-enhanced Raman scattering. These resonances can be observed as peaks 121 in the Raman spectrum of a particle and are often referred to as whispering gallery modes (WGMs). In principle, particle morphology can be deduced from the WGMs, as inhomogeneities in the refractive index can disrupt the circulation of the 122 123 WGMs (Lin et al., 1992; Mitchem et al., 2006). Raman spectra measurements of single droplets in various morphological 124 states are presented in Figure 1. When the droplet was in a homogeneous phase morphology, the droplet acted as a high-125 quality microcavity and sharp WGM peaks overlapped with the spontaneous Raman spectrum (Fig. 1a). When the droplet was 126 in a state of a core-shell structure, observed WGMs were clearly diminished in measured spectra (Fig. 1b). The origin of the 127 damping of the WGMs is the radial homogeneity that is present when the particle is separated into a hydrophilic core and a 128 hydrophobic shell. As a result, when fitting the Raman spectra with the Mie scattering model for homogeneous droplets, the 129 error in the best-fits greatly increase. Examination of the retrieved radius and refractive index reveals a clear break with fits 130 for that of a homogeneous sphere. Therefore, the point at which a significant break in particle size and refractive index occurred 131 can be used as the point at which core-shell phase separation occurs. As illustrated in Fig. 1c, when the droplet was partially-132 engulfed and non-spherical, WGM peaks in the spectrum were absent (Reid et al., 2011). The origin of the spontaneous Raman 133 peaks at 3300 cm⁻¹ and ~3050 cm⁻¹ are identified as the spurious or weakened WGM peaks and the vibration of N-H bond, 134 respectively. Overall, the results of this analysis demonstrate the dynamic changes in the Raman spectra of single droplets as 135 they undergo morphological transitions (Sullivan et al., 2020; Stewart et al., 2015; Tong et al., 2022).



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Figure 1. Raman spectra of 3-MGA-II microdroplets: (a) a homogenous droplet (RH = 90%); (b) a core-shell droplet (RH = 80%); (c) a partially-engulfed droplet (RH=70%). The WGMs are marked by black arrows. The normalization of the peak is achieved by dividing it by the maximum value of the spectrum's intensity, respectively.

141 The peak finding method used in this study is based on the ipeak code developed by O'Haver (2022). In short, the code first 142 smooths the first derivative of the signal and identified downward-going zero-crossings that met a certain predetermined minimum slope and amplitude threshold. By adjusting the corresponding parameters, it is possible to accurately detect the 143 144 desired peaks. The algorithm used to fit WGM peaks in spectra from homogenous spheres in this study was proposed by 145 Preston and Reid (2013) and Preston and Reid (2015). The algorithm compares observed peak positions to expected positions 146 calculated using a resonance condition from Mie theory. Error is minimized by varying particle size and refractive index (i.e. 147 the parameters of best-fit). The method has been demonstrated to provide a rapid determination of the fitted radius and 148 refractive index with an accuracy of ± 2 nm and ± 0.0005 , respectively. During the experiment with reduced RH, we had to 149 adjust the laser power to ensure the stable capture of droplets, which will affect the peak intensity. To eliminate this effect, as 150 demonstrated by Tong et al. (2022), we normalized all Raman spectra used in this study by the area below the spontaneous 151 Raman signals.

152 **3 Results and discussion**

153 3.1 Phase behaviors of droplets mixed SOA proxy with AS

154 Fig. 2 presents the results of time-resolved Raman spectra of aerosol droplets produced from a 3-MGA-II solution under 155 continuously varying RH, as well as the corresponding particle size and refractive index values. To enable temperature and 156 RH to stabilize, the chamber was conditioned with airflow for 50 minutes after trapping a particle. During the dehumidification 157 process, the particle diameter decreased from 11.85 µm to 9.03 µm and the refractive index increased from 1.379 to 1.475 158 when RH decreased from 93.0% to 70.0%. The particle size and water content decreased with RH due to the equilibrium 159 partitioning of water molecules between vapor and droplets. Meanwhile, the refractive index of the droplets gradually increased 160 as the water content decreases. When LLPS occurred, the droplets changed from a symmetrical homogeneous phase to either 161 an asymmetrical partially engulfed structure which led to the disappearance of the WGMs, or the formation of a core-shell 162 structure. As RH in the reaction chamber was reduced, the LLPS was initiated, marked by the variations of the WGM signal 163 (See Fig. 1b). This was achieved by reducing setting RH (setting values) by 5% at 30-minute intervals until the organic phase 164 separated from the water-rich phase and then continuing decreasing RH by 10%-15%. Fig. 2a illustrates how the fitting of the 165 droplet diameter and the refractive index deteriorated as the shell develops, indicating phase separation. The refractive index's 166 shift results from a significant change in the radial profile due to the formation of a core-shell structure. Additionally, the 167 persistence of strong WGMs indicates that the morphology of the droplet remains spherical following LLPS and is core-shell. 168 During the RH increased from 70% to 95%, the reappearance of the continuously shifting WGM signal was observed, 169 suggesting that the inorganic phase has mixed with the organic phase, and droplet returned to a homogeneous phase. During 170 the humidification process, there is an opposite trend observed in the particle size and refractive index of the droplet compared 171 to the dehumidification process. In conclusion, the variations of the WGM signal can serve as a reliable indicator of the 172 occurrence of liquid-liquid phase separation or mixing, and the RH at these points can be considered as the SRH or MRH, 173 respectively. The observed phase transitions of droplets produced from HEXT-IV and HEXD-V solutions are shown in Fig. 3 174 and Fig. 4 respectively.

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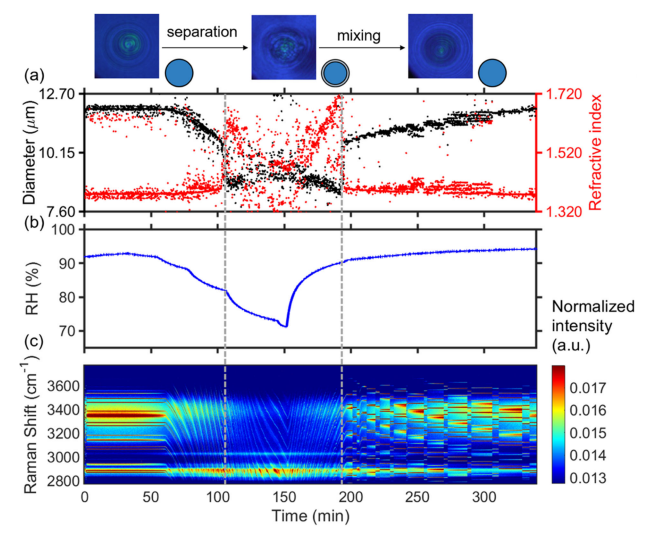
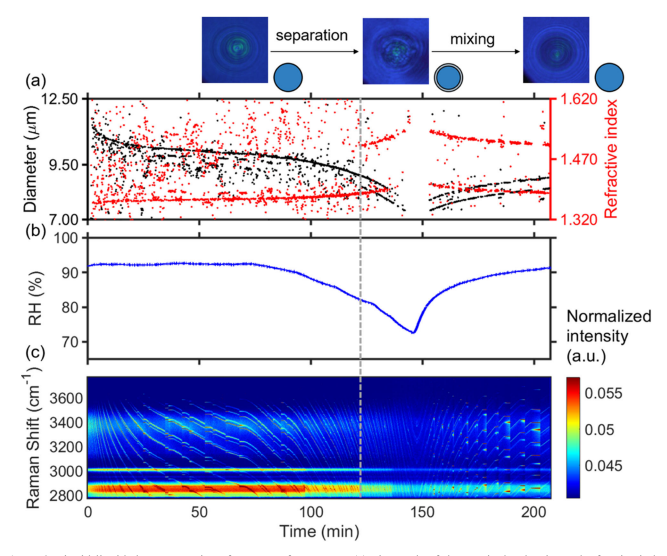


Figure 2. Liquid-liquid phase separation and mixing of aqueous 3-MGA-II. Schematic diagram of phase states is on the top of the figure. (a) Timescale of changes in droplet size and refractive index, determined from fitting the Raman shift positions of the WGMs. (b) RH variation after the trapping chamber during the humidity changing process. (c) Time-resolved Raman spectra. The cessation of the random motion of inclusions within the droplet and the resultant formation of a core-shell structure are indicated by the grey dashed line on the left. The grey dashed line on the right serves as an indication of the point at which the droplet morphology transited from a state of separated phases to a homogeneous phase. The Raman spectra at 53 min, 113 min, 130 min are shown in Fig. 1(a), (b), (c), respectively. Fitting errors of the WGMs was presented in Fig. S3.



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Figure 3. Liquid-liquid phase separation of aqueous of HEXT-IV. (a) Timescale of changes in droplet size and refractive index, determined from fitting the Raman shift positions of the WGMs. (b) RH variation after the trapping chamber during the humidity changing process with time. (c) Time-resolved Raman spectra. The cessation of the random motion of inclusions within the droplet and the resultant formation of a core-shell structure are indicated by the grey dashed line on the left. The grey dashed line on the right serves as an indication of the point at which the droplet morphology transitions from a state of phase separation to a homogeneous phase morphology. This transformation is characterized by the occurrence of phase mixing.

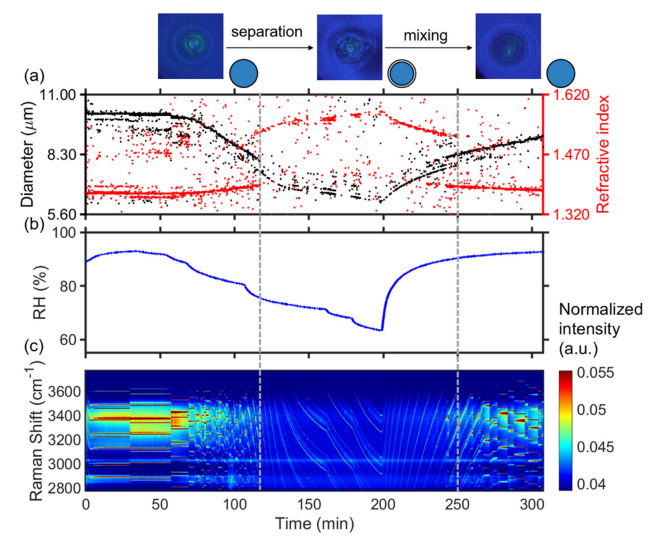


Figure 4. Liquid-liquid phase separation of aqueous of HEXD-V. (a) Timescale of changes in droplet size and refractive index, determined from fitting the Raman shift positions of the WGMs. (b) RH variation after the trapping chamber during the humidity changing process. (c) Time-resolved Raman spectra. The cessation of the random motion of inclusions within the droplet and the resultant formation of a core-shell structure are indicated by the grey dashed line on the left. The grey dashed line on the right serves as an indication of the point at which the droplet morphology transitions from a state of phase separation to a homogeneous phase morphology. This transformation is characterized by the occurrence of phase mixing.

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201 Fig. S2 presents the results of time-resolved Raman spectra of aerosol droplets produced from GL/AS solution under 202 continuously varying RH, as well as the corresponding particle diameter and refractive index values. At the start of the 203 experiment, the chamber RH was held at 93% for approximately 75 minutes. The spectrum during this period revealed a clear 204 bright trend, indicative of the presence of many WGMs in the newly captured droplets. As the chamber RH dropped to a 205 minimum value of 71.5% at around 200 minutes, the position of the WGMs in each spectral snapshot shifted continuously, 206 following the same trend as the chamber RH. This observation suggests that the droplet was homogeneous and that no phase 207 separation occurred in the experimental RH range. The phenomenon regarding the GL/AS system is consistent with the 208 conclusion by Song et al. (2013) and Gorkowski et al. (2020).

209 3.2 Effect of pH on SRH and MRH of different systems

210 The SRH and MRH of aerosol droplets produced from 3-MGA-I~VI solution are shown in **Fig. 5a** and **Table S2**. The pH of

the 3-MGA/AS solution without the addition of an acid or base was 2.70. For solutions with a lower pH (1.19 and 0.48), SA

was added, while NaOH was added to solutions with a higher pH (3.70, 5.21, and 6.53) to adjust their pH levels. The SRH 212 213 values were 92.7%, 89.5%, 80.6%, 79.7%, 76.2% and 69.7% at pH of 6.53, 5.21, 3.70, 2.70, 1.19 and 0.48, respectively. It is 214 worth mentioning that when the pH of the 3-MGA system is 0.48, only two sets of valid parallel experimental data are available, 215 even though we had repeated the experiment several times. Because in other parallel experiments, the SRH of the droplet is 216 lower than the capture range of AOT, the AOT would not be able to continue the capture when the particle size decreases to 217 $\sim 6 \mu m$. Therefore, the actual SRH may be a bit lower at this pH, but this does not affect the results we discuss later. With a 218 decrease in pH, ammonium sulfate transforms into ammonium bisulfate. Our results are consistent with the hypothesis that 219 ammonium bisulfate exhibits a weaker salting out effect compared to ammonium sulfate and thus hinders the ability of organic 220 matter to precipitate out of the solution (Losey et al., 2018). The MRH values at pH 6.53, 5.21, 2.70, 1.19 and 0.48 were 87.6%, 221 89.5%, 87.3%, 83.9% and 83.5%, respectively, and are generally higher than corresponding SRH, especially in the low pH 222 range (<5.00). The SRH was higher than the MRH at pH 6.53, which was abnormal because a lower SRH is commonly 223 expected due to the activation barrier. We do not have a specific explanation for this phenomenon, while it should be noted 224 that the observed values were relatively close to each other, indicating that the higher SRH at pH 6.53 might potentially be 225 attributed to experimental error. The hysteresis between SRH and MRH existed because the SRH process has an activation 226 barrier while the MRH process does not, and lower RH is needed for the aerosol droplet to overcome the activation barrier to 227 form two phases (Freedman, 2020). Similar results were also observed in HEXT/AS and HEXD/AS systems. Additionally, 228 the pH-dependent SRHs obtained in this study were compared to those reported by Losey et al. (2018), as depicted in Fig. 5a. 229 It is worth mentioned that the solute concentration used in our study (50g/L) is comparable to Losey et al. (2018) (5.0 wt%), 230 allowing for meaningful comparison of results. Overall, the SRHs of 3-MGA obtained in this study was higher than the results 231 of Losey et al. (2018). When the pH was lower than 3.70, in 3-MGA system, the present study followed a similar trend as the 232 results of Losey et al. (2018), with the SRH decreasing as the pH decreased. However, when the pH was greater than 3.70, our 233 study showed an opposite trend compared to the results of Losey et al. (2018). The observed discrepancy may be attributed to 234 the distinct ambient conditions of the droplets. The laser levitation, resulting in a spherical morphology, while the optical 235 microscopy involves substrate deposition, leading to a morphology resembling a spherical crown (Tong et al., 2022; Zhou et 236 al., 2014). The underlying reasons for these differences are currently unclear, and further investigations are needed.

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238 In addition to 3-MGA, we also studied two organic/AS systems to investigate how acidity affects SRH and MRH of aerosols 239 of differing composition. These results are shown in Fig. 5 and tabulated in Table 2. The separation diameter (SD) of 3-240 MGA/AS ranges from 7.23µm to 9.74µm, with a corresponding separation refractive index (SRI) ranging from 1.362 to 1.515. 241 For HEXT/AS, the SD ranges from 9.01µm to 9.90µm, while the SRI ranges from 1.396 to 1.421. Lastly, the SD of HEXD/AS 242 ranges from 7.45µm to 8.97µm, with the SRI ranging from 1.382 to 1.406. The data suggests that acidity did not have a 243 noticeable effect on the MRH of the various systems. The pH of the HEXT/AS solution without the addition of any acid was 244 5.11, and SA was utilized to adjust the pH to lower levels (3.14, 2.02 and 0.92). The SRH values of HEXT/AS system 245 (O:C=0.50) decreased as the pH decreased, with values of 78.3%, 76.6%, 76.4% and 75.7% at pH values of 5.11, 3.14, 2.02 and 0.92, respectively. The trend is similar to the 3-MGA (O:C=0.67) system, and the reason why SRH decreased may be due 246 to the acid enhancing the miscibility of organic alcohols and inorganic substances, resulting in a greater difficulty in separating 247 248 the hydrophobic phase from the water-rich phase (Tong et al., 2022). Still, we observed SRH was not strongly dependent on 249 pH for HEXT/AS, compared to 3-MGA/AS system. This is likely due to the fact that organic alcohols have a large pK_a (e.g. 250 the p K_a of HEXT is 14.3) and therefore exhibit minimal ionization in the pH range studied here (Wade and Simek, 2020). 251 Additionally, the relative molecular interactions between alcohols and water are weaker than those of acids, leading to a weaker 252 dependence of salting out ability of AS in the HEXT/AS system. The results of Losey et al. (2018) and Tong et al. (2022) were 253 also depicted in Fig. 5b. Our results differ from those of Losey et al. (2018), who observed a significant decline in SRH as the 254 pH decreased. The specific reason for the discrepancy remains unclear, but we speculate it may due to different condition of 255 droplet. In contrast to the findings of Tong et al. (2022), our study observed a less pronounced trend in the values of SRH, and 256 a narrower range in the distribution of SRH compared to literature values. The difference in OIR between this study (1:1) and 257 Tong et al. (2022) (2:1) may account for the discrepancy in SRH. Previous studies (Ma et al., 2021; Stewart et al., 2015; Song 258 et al., 2012) indicated that OIR differences could affect SRH, but SRH was not significantly dependent on OIR. The 259 discrepancy in SRH may also be due to the variations in experimental conditions, such as laser power, experimental duration, 260 etc. For HEXD/AS (O:C=0.33) system, the pH of the HEXD/AS solution without the addition of any acid was 5.01, and SA was used to adjust the pH to lower levels (3.13, 2.71, 2.03 and 1.39). SRH decreased significantly when the pH was less than 261 262 2.00, while acidity had no significant effect on SRH when pH is greater than 2.00, with values of 79.4%, 81.0%, 77.7%, 82.8% 263 and 70.9% at pH values of 5.01, 3.13, 2.71, 2.03 and 1.39, respectively. This phenomenon was attributed to a mechanism 264 similar to that observed in HEXT/AS. To our knowledge, this is the first investigation on the pH-dependent phase transition 265 of HEXD/AS at the single particle level in a contact-free environment.

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267 The pH values of misty cloud and fog droplets typically fall within the range of 2 to 7, whereas continental and marine aerosol 268 particles display a broader spectrum of pH values (Pye et al., 2020; Tilgner et al., 2021). Our research suggests that in real 269 atmospheric conditions, phase separation behavior of droplets may be influenced significantly by their acidity. It is challenging 270 to measure the droplet pH of the investigated system using AOT. However, previous studies (Coddens et al., 2019; Li et al., 271 2023) have shown that at high RH (90%-100%), the difference in the pH values between droplets and bulk solution is relatively 272 small. Therefore, we used bulk solution pH as an indicator of pH at droplet phase transition. This study focused on volatile 273 organics and was conducted over a relatively long period, which may have affected our results. Nevertheless, the organic 274 compounds used in this study have low volatility. For instance, the vapor pressure of 3-MGA is 7.41×10^{-7} to 2.92×10^{-4} mmHg 275 (DTXSID50871000, United States Environmental Protection Agency), compare to normal volatile organic components of 276 atmospheric aerosol, such as 2-Methyl-1-propanol with vapor pressure of 10.5 to 16.4 mmHg (DTXSID0021759, United States 277 Environmental Protection Agency). Volatility information of other organics are provided in the Table S5. Also, the influence 278 of droplet size change in our system can be neglected. For example, as shown in Fig. 2, the droplet size was basically same at 279 the beginning and the end of the experiment at the same RH 93.0% (11.85 µm at the beginning and 11.79 µm at the end).

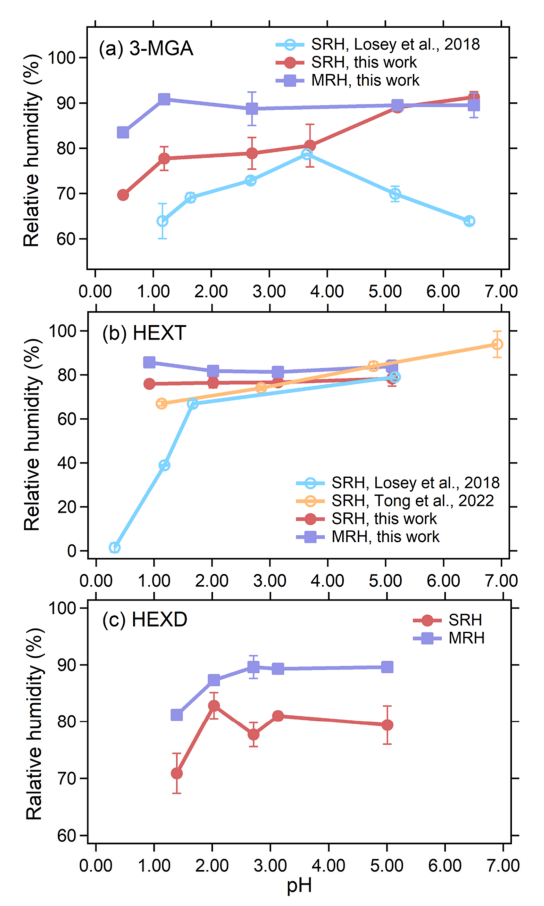


Figure 5. SRHs and MRHs as a function of pH for (a) 3-MGA/AS system, (b) HEXT/AS system, (c) HEXD/AS system.

Hollow circles represent data from Losey et al. (2018) and Tong et al. (2022). The error bars of SRHs and MRHs are derived
 from multiple measurements.

285 Table 2. SRH information for each pH studied as well as initial diameter, separation diameter (SD), separation refractive index

3-MGA/AS system (O:C=0.67)									
Initial	Initial	SRH (%)	SD (µm)	SRI	MRH (%)	MD (µm)	MRI		
pН	Diameter(µm)	()	()	()	N /	(λ=650 nm)		(i)	(λ=650 nm)
0.48	10.97 ± 1.57	69.7±0.2	7.23 ± 1.72	1.515 ± 0.086	83.5	6.82	1.540		
1.19	11.23 ± 1.20	77.7±2.6	8.68 ± 2.38	1.454 ± 0.100	90.8 ± 0.2	9.08 ± 1.64	1.394 ± 0.009		
2.70	12.02 ± 2.94	78.9 ± 3.5	7.88 ± 1.21	$1.493 {\pm} 0.082$	88.7±3.7	6.81±2.76	1.506 ± 0.094		
3.70	10.87 ± 1.87	80.6±4.7	$7.24{\pm}1.00$	$1.491 {\pm} 0.088$					
5.21	11.06 ± 1.63	89.0±0.9	8.93±0.16	1.362 ± 0.014	89.5	7.89	1.381		
6.53	13.73 ± 0.41	91.3±1.2	9.74 ± 0.36	$1.444{\pm}0.187$	89.5±2.7	7.89 ± 0.06	1.383 ± 0.01		
HEXT/AS system (O:C=0.50)									
0.92	13.52±1.6	75.9±0.2	9.90±0.76	1.421 ± 0.017	85.7	10.83	1.420		
2.02	12.88 ± 1.0	76.4±2.3	9.09 ± 0.46	1.409 ± 0.007	81.8	9.34	1.410		
3.14	12.31 ± 0.8	76.6±1.5	9.01 ± 0.47	$1.408 {\pm} 0.002$	81.3	9.04	1.409		
5.11	13.53 ± 0.4	78.3±3.4	9.15±0.35	1.396 ± 0.014	83.9±2.8	9.04 ± 0.73	1.412		
HEXD/AS system (O:C=0.33)									
1.39	11.48 ± 0.78	70.9±3.5	7.45 ± 0.77	1.406 ± 0.008	81.2	7.93	1.406		
2.03	10.54 ± 0.57	82.8±2.3	$7.90{\pm}0.99$	$1.382{\pm}0.007$	87.3	8.83	1.392		
2.71	14.55 ± 1.36	77.7±2.1	8.30 ± 0.28	$1.391{\pm}0.009$	89.6±2.0	8.53 ± 0.32	$1.388 {\pm} 0.010$		
3.13	11.02 ± 0.62	$81.0{\pm}0.7$	8.97 ± 0.22	1.384 ± 0.016	89.3	9.14	1.384		
5.01	12.22±2.73	79.4±3.4	8.33±0.40	1.384 ± 0.019	89.6±0.1	8.38±0.54	1.390 ± 0.004		

286 (SRI), MRH, mixing diameter (MD), and mixing refractive index (MRI) data.

288 **3.3 Effect of O:C on phase separation behavior in different systems**

289 Our findings provide evidence that phase separation of droplets persists even when the organic-inorganic system is adjusted 290 to a specific level of acidity. An important determinant of whether droplets undergo phase separation is the O:C. To illustrate 291 this, we have included a plot in Fig. S4, which show cases the experimental system used in our study alongside relevant 292 literature values. One point that needs to be declared is Fig. S4 only plotted for systems with no additional H₂SO₄ or NaOH. 293 As shown in Fig. S4, our findings, as well as those from previous studies (You et al., 2013; O'Brien et al., 2015), indicated 294 that there is no correlation between the occurrence of LLPS and the hydrogen-to-carbon (H:C) ratios of the organics, which is 295 consistent with results in previous findings (Bertram et al., 2011; Song et al., 2012). However, a clear trend was observed 296 between LLPS occurrence and the O:C of the organic components. We observed that droplets of 3-MGA/AS, HEXT/AS and 297 HEXD/AS systems with O:C between 0.33 and 0.67 undergo LLPS. With the decrease of water content in the droplets, two 298 distinct phases were formed: an organic-rich phase and a salt-rich aqueous phase, under both acidic and neutral conditions. By 299 contrast, no LLPS occurred in the GL/AS system, as shown in Fig. S2. In general, particles with low O:C are more prone to 300 undergo LLPS. This observation is consistent with the findings of Song et al. (2012) who reported that LLPS was never 301 observed when O:C > 0.80 and always observed when O:C < 0.56.

302

As shown in **Fig. 2** and **Table S2**, for most spectra, WGMs remained after LLPS occurred for droplets of 3-MGA/AS. This phenomenon indicates that the droplets undergo LLPS with a core-shell morphology in most conditions, which is consistent with the prediction of Gorkowski et al. (2020). Meanwhile, morphology of phase-separated droplets containing either HEXT or HEXD were also core-shell shape mostly, as depicted in **Fig. 3/4** and **Table S3/S4**. It is attributed to the lower interfacial tension observed at higher O:C, leading to higher possibility condition for forming core-shell shaped droplets (Gorkowski et al., 2020). These findings support the idea that the O:C plays a crucial role in determining the morphology of phase-separated

309 particles in organic/inorganic mixed aerosols.

310 4 Conclusion

311 The aim of this study is to investigate the effect of pH and O:C on phase transition behavior of levitated particles using the 312 AOT. Our results show that across aerosol pH in atmospheric condition, the presence of sulfuric acid inhibited the LLPS of 313 aerosol droplets that contained organics (3-MGA, HEXT, HEXD) and AS. Additionally, the MRHs were found to be higher 314 than the SRHs. The O:C of phase-separating systems is 0.67, 0.50, 0.33, and by contrast, LLPS of the high O:C system (GL, 315 O:C=1.00) did not occur. Meanwhile, the morphology of levitated aerosol particles was studied and we found that 42 out of 316 46 droplets that underwent LLPS for a core-shell structure. The SRH of all experimental systems ranged from $\sim 70\%$ to 90%. 317 In certain cases, as the RH decreased, the droplet morphology changed from core-shell to partially engulfed, similar to the 318 findings reported by Kucinski et al. (2020). However, as the RH further decreased, the droplet particle size became smaller 319 than 6 µm, making it impossible to capture them using AOT. Consequently, in most instances, we were unable to observe the 320 droplet morphology at RH levels below 70%. The results presented here provide new insights into the behavior of different 321 types of aerosol droplets, and the findings have important implications for our understanding of physical and chemical 322 processes that occur in the atmosphere. It is anticipated that future studies will be carried out to examine the OIR-dependent 323 phase separation in real acidified ambient aerosols. Such research will provide insights into the morphological characteristics 324 of real aerosols and the ways in which these characteristics influence important properties such as hygroscopicity and 325 homogenous chemistry. Such information will be helpful in furthering our understanding of the impacts of ambient aerosols 326 on the environment and human health.

327

Additionally, in-situ pH measurement or pH estimation methods, such as the real-time AOT analysis in microdroplets reported by Boyer et al. (2020) could be combined with SRH measurements for a more accurate and comprehensive analysis. Furthermore, our study used a surrogate for SOA instead of in situ measurements of real SOA, which can be addressed in future work using SOA generated from a smog chamber or real SOA precursors and oxidized species.

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333 Data availability. The data used in this paper can be obtained from the corresponding author upon request.

334 *Author contributions.* YC built the instrument, performed the experiments, analyzed the data, plotted the figures, and wrote

the original draft. XP conceptualized the study, contributed to instrumentation, data analysis, discussion, and reviewed the

336 manuscript. HL and CX contributed to the instrumentation and discussion. YM contributed to the experiments and discussion.

337 ZX, FZ contributed to the discussion and manuscript review. TCP contributed to data analysis and manuscript review. ZW

administrated the project, conceptualized the study, reviewed the manuscript, and contributed to funding acquisition.

339 *Competing interests.* The contact author has declared that none of the authors has any competing interests.

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