Temporally and spatially resolved investigation on the efflorescence process of a mixed droplet of ammonium sulfate and ammonium nitrate

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Temporally and spatially resolved investigations of the efflorescence of equimolar mixtures of ammonium sulfate (AS) and ammonium nitrate (AN) aerosol droplets were performed by confocal Raman spectroscopy combined with high-speed photography. Information was obtained on the dynamic process of morphological changes and the heterogeneous distribution of efflorescence products. The supersaturated AS/AN mixed droplet was found to first precipitate as the double salt 3AN·AS when the relative humidity was 50%. The excess AS was irregularly distributed on the surface of the double salt as an inclusion structure. No formation of 2AN·AS was observed throughout the efflorescence process.

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Ammonium sulfate (AS) and ammonium nitrate (AN) are critical inorganic components in atmospheric aerosols [1]. Mixtures of compounds have also been shown to form double salt structures with various compositions in the aerosol during the efflorescence process. Martin and Schlenker et al. [2−4] investigated the efflorescence of $\text{SO}_4^{2−}/\text{NO}_3^{−}/\text{H}^{+}/\text{NH}_4^{+}$ by Fourier transform infrared spectroscopy/aerosol flow tube (FTIR-AFT) analysis. They found two double salts, 3AN·AS and 2AN·AS, formed along with pure AN and AS. The relative contents of these salts were dependant on the concentration ratio of AN/AS. Chan et al. [5] used an electrodynamic balance (EDB) in combination with Raman spectroscopy to study the efflorescence of equimolar AN/AS mixed in a single droplet, and found that the initial product of efflorescence was mainly metastable 3AN·AS. However, FTIR-AFT and EDB-Raman have some limitations and these studies have not provided in-depth descriptions on the spatial distribution of efflorescence products and its dynamic aspects. For FTIR-AFT, the statistical information is from a great number of aerosols droplets and it does not look at the differences in individual droplets, which makes it difficult to identify the mixing conditions of components such as the internal/external mixing states [6]. EDB-Raman has been widely applied for measurements of single particles. However, only the fundamental information of the droplet can be obtained and the growth status and spatial distribution of AN/AS in the mixed droplet throughout efflorescence process cannot be studied in detail. The confocal technique has high spatial resolution and has been applied to studies on aerosol particles with micro-scale spatial resolution [7,8]. A high-speed camera and optical microscope have been used in combination for a temporally resolved study on the morphological changes of aerosol droplets during efflorescence [9]. In the present study, confocal Raman spectroscopy was combined with high-speed photography for a temporally and spatially resolved study on the efflorescence of a mixed AS/AN droplet. This allowed monitoring of the dynamic process of morphological...
changes and the heterogeneous distribution of efflorescence products.

Equimolar mixtures of AN/AS solutions with a concentration of 0.5 mol/L for each component was prepared and deposited on the quartz substrate using a micro syringe. The substrate was then placed in a sample cell. The relative humidity (RH) in the sample cell could be adjusted by controlling the flow rates of water vapor saturated with nitrogen and dry nitrogen gas into the cell. In situ monitoring of the morphological changes during the efflorescence process was achieved by high-speed photography (MS55K, MegaSpeed Corp, Canada). Confocal Raman spectroscopy (inVia, Renishaw, Britain) was used for spatial resolved analysis of the heterogeneous distribution of efflorescence products. The instrument settings and process have been described elsewhere [8,9].

The Raman spectra of pure AN, AS, 3AN·AS and 2AN·AS crystals are shown in Figure 1. The spectra of pure AN and AS were acquired from the corresponding analytical grade chemicals, while the 3AN·AS and 2AN·AS for Raman analysis were obtained from the precipitants of different ratio mixtures of AN/AS solutions. These precipitants were screened to identify crystals that gave spectra consistent with those for 3AN·AS and 2AN·AS described in an earlier study [5]. In this earlier study, the Raman peaks of nitrate were located at 716 and 1045 cm\(^{-1}\), while those of sulfate were located at 451, 614, and 975 cm\(^{-1}\). No distinct Raman peaks for ammonium were observed between 400 and 1100 cm\(^{-1}\). The nitrate peak at 1051 cm\(^{-1}\) could be clearly distinguished in the Raman spectra, and the peak position and shape were used in component identification and analysis for the AN/AS mixed droplet.

The time-dependent morphological changes of the AN/AS mixed droplet with a diameter of 100 \(\mu\)m are presented in Figure 2. At 50% RH, no obvious changes were observed in the droplet in the earlier stages. Metastable crystallization began to occur on the right edge of the droplet after 5 min. For convenience, the moment that crystallization first took place was regarded as starting point. Following the appearance of the crystal seed, a number of primary branches began to grow with time. Among these branches, two distinct branches had larger growth velocities and continuously grew until the whole droplet was covered by the crystal at 10.48 s. The crystal growth was accompanied by obvious evaporation of water to equilibrate the humidity of the system, and this occurred throughout the crystal growth process. The final crystal morphology of the droplet is shown in the last picture in Figure 2.

To obtain an accurate image of the distribution of chemical compositions in the freshly formed crystal, the ambient RH was adjusted to 30%, which suppressed the transformation of double salts from 3AN·AS to 2AN·AS [5]. Using micro-Raman spectroscopy, the Raman signals of the freshly formed crystal in different regions were acquired. Four typical regions marked as A, B, C and D were selected in reference to the dynamic images discussed above. Figure 3 presents the morphologies and Raman spectra of these different regions in the formed crystal. Regions A and B (Figure 3(a)) were located at the center and the edge of the largest branch, respectively. Region C was located at the edge of the region between the two large branches, and region D was located at the position where the crystallization first occurred. The Raman spectra at these four locations are shown in Figure 3(b). In the spectrum for region D, a strong peak appeared at 1051 cm\(^{-1}\) and a weak peak at 979 cm\(^{-1}\). Compared with the reference spectra (Figure 1), the crystal in region D was pure 3AN·AS. For the same reasons, the crystal in region C could also be considered as 3AN·AS even though the signals were weak. In the spectrum of region B, a strong peak at 1051 cm\(^{-1}\) indicated that 3AN·AS was present, and a peak at 975 cm\(^{-1}\) showed pure AS was present. The relatively weak intensity of the 975 cm\(^{-1}\) peak and its asymmetrical shape indicated that the content of pure AS was small. In the spectrum of region A, the crystal showed the highest purity of AS and only a small amount of 3AN·AS. The peak at 975 cm\(^{-1}\) was strong and almost symmetrical, and the peak at 1051 cm\(^{-1}\) was weak. Because of the low ambient RH, no characteristic peak for 2AN·AS was observed in any of the four regions or elsewhere (data not presented) for all the Raman spectra of the crystal.

In combination, the morphology and spatial-resolved
Raman spectroscopy results can be used to deduce a reasonable mechanism for the crystallization process of an equimolar AN/AS mixed droplet. In this mechanism, the concentration of the droplet continues to increase and reaches supersaturation following the decrease in ambient humidity. As the RH reaches 50%, metastable crystallization of 3AN·AS starts to occur on the edge of droplet. Accompanying water evaporation, results in the crystallization of 3AN·AS until the nitrate content remaining in the mixed droplet is negligible. The excess sulfate in the solution will then crystallize to form pure AS, which will be deposited irregularly on the surface of previously precipitated 3AN·AS to form an inclusion structure. Formation of 2AN·AS was not observed throughout the crystallization process.

Figure 4(a) and (b) shows the Raman spectra of the crystal at different depths in regions D and A, respectively. There was no change in the spectra in region D (Figure 4(a)) when the scanning depth was increased, which indicates that 3AN·AS was present at all depths and there were no other components. For region A (Figure 4(b)), the spectra showed the peak at 1051 cm\(^{-1}\) gradually increased compared with the peak at 975 cm\(^{-1}\), which suggests that the content of 3AN·AS increased gradually as the depth increased. The appearance of a 979 cm\(^{-1}\) shoulder on the peak at 975 cm\(^{-1}\) caused asymmetry, and provides further information about the crystallization process.
evidence for this increase in 3AN·AS.

The application of temporally and spatially resolved analysis of single particles of a multi-component aerosol droplet was useful to further understand the mechanisms of phase separation and transformation. The results can be used to evaluate the common principles of atmospheric aerosol phase transformation.

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1 Seinfeld J H, Pandis S N. Atmospheric Chemistry and Physics. New York: John Wiley and Sons Inc., 2006
2 Martin S T, Schlenker J C, Malinowski A, et al. Geophys Res Lett, 2003, 30: 2102
3 Schlenker J C, Malinowski A, Martin S T, et al. J Phys Chem A, 2004, 108: 9375–9383
4 Schlenker J C, Martin S T. J Phys Chem A, 2005, 109: 9980–9985
5 Ling T Y, Chan C K. Environ Sci Technol, 2007, 41: 8077–8083
6 Rosenoern T, Schlenker J C, Martin S T. J Phys Chem A, 2008, 112: 2378–2385
7 Everall N. Appl Spectrosc, 2008, 62: 591–598
8 Wang F, Shou J J, Zhang Y H. Chinese Sci Bull, 2008, 53: 2414–2416
9 Wang F, Zhao L J, Zhang Y H. Chinese Sci Bull, 2008, 53: 2140–2145