



Characterizing efflorescence regimes in organic–inorganic aerosols using thermodynamically modeled viscosity

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Abstract. Atmospheric aerosols, especially internally mixed organic–inorganic aerosols, exhibit complex phase behaviors that affect their size evolution, optical properties, and chemical reactivity, ultimately impacting climate and human health. Although parameterizations for secondary organic aerosol phase state exist, quantitative descriptions of efflorescence behavior in organic–inorganic aerosols remain underdeveloped. Herein, we evaluated chemical parameters, including O : C ratio, organic mass fractions, glass transition temperature (T_g), and viscosity (η), and identified aerosol viscosity as the primary variable associated with efflorescence relative humidity (ERH) in organic–inorganic aerosols. We developed a viscosity–ERH framework, which defines boundary conditions that separate viscosity–humidity space into three empirical regions. Above the fitted line, aerosols remain in an aqueous state, whereas below the line, efflorescence occurs. Additionally, efflorescence is unlikely to occur when η exceeds a threshold value ($> 4.76 \times 10^2$ Pa s). The predicted boundary line was validated using an independent dataset, and a multivariate regression model incorporating η and T_g improved the statistical performance of the boundary line but was limited by T_g parameterization for complex organic–inorganic mixtures. Our findings highlight the association between aerosol viscosity and efflorescence and emphasize the need to develop improved aerosol viscosity measurement techniques to better constrain aerosol phase behavior in atmospheric models.

1 Introduction

Atmospheric aerosols significantly influence climate and human health through both direct and indirect effects. They directly influence the energy balance by absorbing and scattering solar radiation and indirectly affect the climate by acting as cloud condensation nuclei (CCN), thereby altering cloud microphysical properties such as droplet size, lifetime, and optical characteristics (Boreddy et al., 2014; Fan et al., 2010; Knopf and Alpert, 2023; Novo et al., 2021; Pöschl, 2005; Shrivastava et al., 2017; Smith et al., 2021). Among aerosol physicochemical properties, phase state (liq-

uid, solid, or semi-solid) is particularly important, as it governs hygroscopicity, optical properties, and heterogeneous reactivity (Fard et al., 2018; Freedman et al., 2024; Novo et al., 2021). Moreover, aerosol phase transition significantly affects viral activity during transmission (Oswin et al., 2022). Consequently, aerosol phase states substantially impact climate, air quality, and human health (Reid et al., 2018).

Atmospheric aerosols, particularly internally mixed secondary aerosols composed of organic and inorganic components, exhibit diverse phase states, including liquid, liquid–liquid phase-separated, semi-solid, amorphous solid, and crystalline solid, which govern their physicochemical prop-

erties (Novo et al., 2021). For instance, aerosols undergoing efflorescence (i.e., crystallization) and deliquescence exhibit hysteresis in particle size and water content in response to relative humidity (RH) variations (Choi and Chan, 2002; Guo et al., 2020). In contrast, amorphous organic aerosols generally do not exhibit the pronounced discontinuous hysteresis associated with efflorescence and deliquescence of crystalline particles. However, semi-solid or glassy amorphous particles may still show gradual, kinetically driven hysteresis in water uptake and release because of slow diffusion and equilibration (Koop et al., 2011). Thus, effloresced (or crystallized) aerosols display fundamentally different hygroscopic behaviors from amorphous ones, introducing uncertainties in predicting mixing aerosol water uptake and particle size under varying RH conditions because phase transitions and kinetic limitations can lead to different water contents and growth factors during hygroscopic cycles (Marcolli and Krieger, 2006; Mikhailov et al., 2009).

Aerosol particles consisting of more than a single phase may have crystalline as well as amorphous phases. It is important to note that, in internally mixed organic–inorganic aerosols, the onset of phase transitions such as deliquescence and efflorescence typically occur only in specific components rather than the entire particle. For example, crystalline inorganic salts or hydrates may undergo deliquescence or crystallization, while the remaining particle matrix can persist in an amorphous, semi-solid, or aqueous state (Hodas et al., 2016).

However, internally-mixed organic–inorganic aerosols are chemically complex, often containing thousands of distinct organic species, of which only about 10 % have been identified (Hallquist et al., 2009). This limited chemical characterization constrains our ability to accurately predict aerosol phase states. Existing parameterization models based on organic aerosol properties, such as molecular weight and oxygen-to-carbon ratio (O : C ratio) (Shiraiwa et al., 2017), elemental composition (DeRieux et al., 2018), and volatility (Li et al., 2020; Zhang et al., 2019) to predict the glass transition temperature (T_g), have been established to predict the phase state of organic aerosols but these models typically neglect efflorescence. Consequently, current parameterizations fail to distinguish between crystalline and amorphous solid states of internally mixed organic–inorganic aerosols and overlook the hysteresis in particle size and water uptake during efflorescence–deliquescence cycles. This may lead to biases in describing atmospheric aerosol phase behavior, hygroscopicity, and optical properties, affecting our understanding of particle aging via both physical and chemical processes in the atmosphere.

While laboratory studies have provided substantial data on efflorescence in mixed organic–inorganic aerosols, a comprehensive understanding of the factors influencing efflorescence remains limited (Cai et al., 2017; Ma et al., 2021b; Wang et al., 2017). Efflorescence involves the formation of crystalline nuclei from a supersaturated solution and pro-

ceeds through kinetically controlled nucleation and crystal growth that requires overcoming an energy barrier. This process is governed by a combination of thermodynamic driving forces and kinetic limitations.

Component diffusion regulates the mobility of ions and molecules and therefore affects the rate at which crystalline nuclei can form and grow; higher viscosity generally suppresses diffusion and delays efflorescence (Ji et al., 2017; Mikhailov et al., 2009). Chemical composition, including solute concentration, solubility, and vapor pressure, influences the degree of supersaturation and thus the thermodynamic driving force for crystallization (Cohen et al., 1987; Gupta et al., 2015). In addition, molecular interactions within the particle phase can either facilitate or hinder the structural rearrangement required for nucleation, while interfacial properties such as surface tension affect the nucleation energy barrier by altering the energetic cost of forming a new phase interface (Davis et al., 2015; Ma et al., 2021a; Mikhailov et al., 2009).

Thermodynamic constraints further require that efflorescence and crystallization can only occur when the aqueous phase becomes supersaturated with respect to the relevant solid phase; otherwise, efflorescence is thermodynamically inhibited. Once supersaturation is achieved, nucleation and crystal growth proceed through kinetically controlled processes (Bouzidi et al., 2022; Marcolli and Krieger, 2006). The corresponding transition point, described by the efflorescence relative humidity (ERH), is therefore strongly influenced by these factors, which are modulated by the properties of organic components, and the hydration state of the inorganic salts.

The O : C ratio is widely used as an indicator of organic compound polarity (Yu et al., 2021). Previous studies have shown that increasing the mass fraction of organic compounds (ω_{org}), particularly with high-viscosity organics, can significantly alter mass transport processes, and thereby affect phase transition (Parsons et al., 2004; Shiraiwa et al., 2013; Virtanen et al., 2010; Wang et al., 2017; Zobrist et al., 2011). Under such conditions, efflorescence may be kinetically suppressed because of limited molecular diffusion, causing aerosol particles to remain in an amorphous semisolid or glassy state; in this case, their phase-state evolution is more appropriately characterized with reference to the glass transition temperature (T_g) (Berkemeier et al., 2016; Koop et al., 2011; Song et al., 2016b; Tong et al., 2011). However, direct and reliable measurements of the viscosity of atmospheric aerosols under atmospherically relevant conditions remain challenging (Fitzgerald et al., 2016; Hosny et al., 2016), particularly because water uptake can continuously modify particle composition, phase state, and molecular mobility (Sheldon et al., 2023; Zobrist et al., 2008). This has resulted in limited observational data and motivated the use of thermodynamic models, such as AIOMFAC, to predict the viscosity of organic–inorganic mixed aerosols (Lilek and Zuend, 2022).

In this study, we develop a framework to predict efflorescence in organic–inorganic aerosols by integrating literature-reported ERH data and evaluating key physicochemical parameters, with particular emphasis on aerosol viscosity (η) and the combined effects of T_g , O : C ratio, and ω_{org} . This framework constrains the efflorescence phase transition in viscosity–humidity space and provides a quantitative description of boundaries separating aqueous, efflorescence and non-efflorescence states. The result improves the representation of aerosol phase behavior by using ambient RH and thermodynamically modeled viscosity, and offers constraints for atmospheric processes such as gas–particle partitioning, heterogeneous chemistry, and cloud formation.

2 Methods

2.1 The dataset for the parameterization

We compiled aerosol ERH data from our previous studies and peer-reviewed literature identified through Web of Science using keywords: aerosol, hygroscopic, and efflorescence. All studies were laboratory-based, with ERH measured using IR spectroscopy (Braban and Abbatt, 2004; Cai et al., 2017; Ghorai et al., 2014; Ji et al., 2017; Ma et al., 2021b; Shi et al., 2017; Wang et al., 2017; Wu, 2017; Xu et al., 2022), Raman spectroscopy (Chang, 2020; Hu et al., 2025; Ushijima et al., 2021; Wang, 2018; Yu et al., 2012), optical microscopy (Bertram et al., 2011; Huang et al., 2024; Parsons et al., 2004) or Hygroscopicity Tandem Differential Mobility Analyzer (HTDMA) (Laskina et al., 2015). In multicomponent organic–inorganic particles, multiple ERH may exist due to different salts crystallizing at distinct RH levels, and efflorescence typically occurs over a RH range. In this study, “aerosol ERH” is defined as the mean value of this transition range, serving as a simplified but effective macroscopic parameter for the overall phase transition of the particle.

A total of 102 ERH data points for organic–inorganic mixture aerosols were collected: 66 ERH points for the training set (Table S1 in the Supplement) and the remaining 36 points for the validation set (Table S2). The training set includes aerosol compositions containing 14 organic compounds (composed of C, H, and O) and 4 inorganic salts (ammonium sulfate, ammonium nitrate, sodium chloride, and potassium chloride) mixed in varying ratios. All ERH data in the training set were from our previous studies, and all corresponding viscosity data were derived using the AIOMFAC model. The particles investigated in our previous studies typically range from 1–10 μm in diameter. The validation set contains 36 ERH data from systems involving 12 organic compounds and 4 inorganic salts (Table S2), selected from studies that either reported both aerosol ERH and viscosity or observed no efflorescence. For dataset included in this study, experiments were conducted over several hours with slow RH change rate ($< 0.1\% \text{ s}^{-1}$) to ensure

near-equilibrium conditions. Although these timescales are shorter than typical atmospheric aerosol lifetimes (days to weeks), previous studies have shown that nucleation induction times in high-viscosity systems can be extremely long due to diffusion limitations. (Raes, 2000; Wang et al., 2017) Therefore, the suppression of efflorescence at high viscosity reported in the cited studies is likely a fundamental kinetic constraint rather than an artifact of limited experimental duration. Moreover, the empirical data utilized to derive the regression model in this study were all obtained within a temperature range of $298 \pm 10 \text{ K}$.

Key physicochemical parameters for describing aerosol ERH include the O : C ratio of the organic compound in the mixture:

$$\text{O : C} = \frac{N_{\text{O,org}}}{N_{\text{C,org}}} \quad (1)$$

where $N_{\text{O,org}}$ and $N_{\text{C,org}}$ are the number of oxygen and carbon atoms in the organic compound, respectively. Other parameters include ω_{org} , fractional T_g at ERH ($T_g(\omega_{\text{org}})$), and viscosity.

2.2 AIOMFAC model

Aerosol viscosity at the corresponding ERH and 298 K was estimated using the Aerosol Inorganic–Organic Mixtures Functional Groups Activity Coefficients (AIOMFAC) model. (Zuend et al., 2008, 2011) AIOMFAC is a thermodynamics-based group-contribution framework developed for aqueous organic–inorganic mixtures. To predict the viscosity of complex organic–inorganic mixtures, the AIOMFAC-VISC framework conceptually divides the mixture into two distinct sub-systems. For the aqueous inorganic (electrolyte) sub-system, it employs a semi-empirical model based on Eyring’s absolute rate theory, which relates viscosity to the molar Gibbs energy of activation for viscous flow (Δg^*) taking into account temperature, ion activities, and ionic strength. Conversely, for the aqueous organic sub-system, the model relies on a combinatorial-activity-weighted approach based on functional group contributions. (Gervasi et al., 2020) Adopting the UNIFAC concept, AIOMFAC segments organic molecules into functional subgroups, treating the liquid mixture as a “solution of groups” to manage chemical complexity. (Fredenslund et al., 1975; Hansen et al., 1991) In our study, the viscosity of mixed organic–inorganic phases was estimated using the default “aquelec” (electrolyte-aware water) approach in AIOMFAC-web. This method represents the effects of ions by first computing the viscosity of the aqueous electrolyte mixture (excluding organics), and then using that sub-system’s viscosity to represent a modified viscosity of water in the calculation of the electrolyte-free organic sub-system. Validation against experimental data (e.g., Fig. 11 in Lilek and Zuend, 2022) has demonstrated that the aquelec approach accurately characterizes the viscosity of internally mixed aerosols such as sucrose–nitrate systems, establishing

it as a robust tool for our predictions (Lilek and Zuend, 2022; Song et al., 2021; Zuend et al., 2011).

Accordingly, viscosity calculations were performed by specifying the functional groups of the organic components, inorganic salt composition, organic-to-inorganic ratio (OIR, molar ratio of organic compounds to inorganic salts, excluding water), and a series of organic molar fractions in the solute mixture, x_{org} , where the solute consists of organic compounds and inorganic salts, ranging from 0.01–0.99 in increments of 0.01. For each system, the viscosity corresponding to the water activity closest to the literature reported ERH (within $\pm 0.5\%$ RH) was selected. For example, in a 1,2,6-hexanetriol and ammonium sulfate (1 : 1) mixture, we defined 1,2,6-hexanetriol as 2 CH₂(OH), 1 CH(OH), 1 C and 3 OH. The viscosity of the mixture system was evaluated at a water activity of 0.47, which closely matches the ERH value in the literature (47.4 %). Functional group definitions for all organic components are listed in Table S3. To assess the uncertainty associated with the thermodynamic state predictions, the sensitivity of viscosity ($\pm \log_{10}(\eta/[\text{Pa s}])$) was calculated for each mixture. This metric quantifies the variation in predicted viscosity resulting from a $\pm 2\%$ perturbation in aerosol water mass fraction and serves as a proxy for uncertainty related to compositional variability. The corresponding values are listed in Table S1.

2.3 Calculation of the glass transition temperature ($T_g(\omega_{\text{org}})$)

Multiple parameterizations exist for estimating the glass transition temperature (T_g) of organic compounds, utilizing predictors such as chemical structure, volatility, or melting point (T_m) (Armeli et al., 2023; Galeazzo and Shiraiwa, 2022; Koop et al., 2011; Li et al., 2020). Recent machine-learning models based on chemical structure do not strictly require T_m information, though including it as an optional parameter can further improve predictive accuracy (Armeli et al., 2023). The framework proposed by Armeli et al. comprises four distinct modes, contingent upon the specific input parameters available for a given compound: the Functional Group Mode (FGM) and the SMILES Mode (SM). Each mode features a variant that includes melting temperature as an additional feature (FGM with T_m ; SM with T_m) and a version that operates without it (FGM no T_m ; SM no T_m). The T_g values and parameters calculated according to the methodology of Armeli et al. (2023) are summarized in Tables S4–S7.

The volatility-based parameterization developed by Li et al. (Eq. 2) predicts T_g from the saturation mass concentration (C^0).

$$T_g = 288.7 - 15.33 \times \log_{10}(C^0) - 0.33 \times \left[\log_{10}(C^0) \right]^2 \quad (2)$$

C^0 was estimated as a function of C, O, N, S numbers of the organic compound ($\log_{10}C^0 = f(n_C, n_O, n_N, n_S)$, Eq. 3) (Donahue et al., 2011; Li et al., 2016):

$$\log_{10}C^0 = \left(n_C^0 - n_C \right) b_C - n_O b_O - 2 \frac{n_C n_O}{n_C + n_O} b_{CO} - n_N b_N - n_S b_S \quad (3)$$

Here, n_C^0 is the reference carbon number; n_C , n_O , n_N , and n_S denote the number of carbon, oxygen, nitrogen, and sulfur atoms, respectively; b coefficients (b_C , b_O , b_N , and b_S) denote atomic contributions fitted via multi-linear least squares analysis from 30 000 compounds across multiple classes (CH, CHO, CHN, CHON, CHOS, and CHONS). (Li et al., 2016) $b_{(CO)}$ represents the carbon–oxygen nonideality.

A comparison between the experimentally measured T_g values of the organic compounds and the predictions derived from the methods of both Li et al. (2020) and Armeli et al. (2023) is presented in Table S7.

The presence of inorganic salts (e.g., sodium nitrate) has been demonstrated to modulate the T_g of organic–inorganic mixtures; specifically, depending on the effective T_g of inorganic salt, the T_g of the mixture can be either elevated or depressed. (Dette and Koop, 2015) Typically, the T_g of organic–inorganic mixtures is calculated using the Gordon–Taylor equation (Gordon and Taylor, 1952):

$$T_g = \frac{\omega_1 T_{g1} + \frac{1}{k} \omega_2 T_{g2}}{\omega_1 + \frac{1}{k} \omega_2} \quad (4)$$

where T_g represents the glass transition temperature of the dry binary mixture, T_{g1} and T_{g2} are the T_g values of the respective pure compounds, ω_1 and ω_2 denote the mass fractions of the two components, and k is the Gordon–Taylor constant. However, the experimental data of glass transition temperature of inorganic compounds is limited, we thus employ the glass transition temperature of the organic–water system ($T_{g,\text{org-water}}$) as a proxy for the overall mixture T_g when analyzing the correlation between efflorescence relative humidity (ERH) and T_g . In Eq. (4), the parameters ω_2 and T_{g2} are substituted with the mass fraction ω_{org} and the glass transition temperature $T_{g(\omega_{\text{org}})}$ of the organic component, respectively, where ω_{org} is the organic mass fraction calculated via the AIOMFAC model at aerosol ERH. $T_{g,w}$ is the glass transition temperature of pure water (136 K) (Kohl et al., 2005), while k_{GT} (the Gordon–Taylor constant) is assumed to be 2.5 for organic–water mixtures (Koop et al., 2011; Zobrist et al., 2008). These two values are substituted for T_{g1} and k in the equation, with ω_1 calculated as $(1 - \omega_{\text{org}})$. As reported by Koop et al. (Dette and Koop, 2015), the T_g of NaNO₃ is specified as 290 K. Using this reported T_g value for NaNO₃ together with $k_{GT} = 0.5$, we calculated the T_g for dry glycerol/NaNO₃ and dry sucrose/NaNO₃ mixtures across a range of mixing ratios (Fig. S1 in the Supplement). It should be noted that assuming a constant Gordon–Taylor parameter ($k_{GT} = 2.5$) for all organic–water mixtures represents a

generalized simplification. The k_{GT} value essentially parameterizes the interaction strength between water and the specific organic solute. Consequently, for organic mixtures with highly complex or varying oxygenated functional groups, the actual k_{GT} value may deviate from this assigned average (typically estimated within a range of 2.5 ± 1.0). This variability introduces a potential, albeit minor, source of uncertainty into the T_g -based model predictions presented in this study.

In addition to the approach used here for estimating $T_g(\omega_{org})$, the glass transition temperature of the entire mixture, including contributions from inorganic ions, could also be derived from the predicted mixture viscosity. This can be achieved by combining AIOMFAC-VISC viscosity predictions with an assumed fragility parameter within the Vogel–Tammann–Fulcher framework. (DeRieux et al., 2018) However, the pure-component viscosity parameterization for organics implemented in AIOMFAC-web may be less accurate than more recent approaches (Armeli et al., 2023). Moreover, for the purposes of the present regression model, estimating T_g from mixture viscosity is likely redundant, as it does not provide additional predictive power beyond viscosity itself.

3 Results and discussion

3.1 Correlation between ERH and physicochemical parameters

Herein, we focus on the efflorescence of the inorganic component in internally mixed organic–inorganic aerosols. Therefore, we examined the correlation between mean ERH and key physicochemical parameters describing the organic components in organic–inorganic mixture aerosols: O : C ratio, ω_{org} , and $T_g(\omega_{org})$ (Fig. 1).

Linear regression analysis using the training set (66 data points) showed no significant correlations between aerosol ERH and any of O : C, ω_{org} , or $T_g(\omega_{org})$ individually ($R^2 < 0.15$; Table S8). Multivariate regression considering the combined influence of three parameters provided only a limited improvement ($R^2 = 0.18$, Table S9). According to the nested cross-validated model evaluation reported by Armeli et al. (Armeli et al., 2023) for the full dataset, the SMILES mode with T_m showed the highest predictive accuracy among the four modes. Therefore, the T_g values calculated using the SMILES mode with T_m were selected for comparison with the parameterization developed by Li et al. (2020). The results, presented in Fig. 1c, demonstrate that neither methodology yields a statistically significant correlation. Furthermore, analyses employing alternative FGM and SMILES models for T_g estimation, and T_g values for organic–inorganic mixtures were also investigated (Fig. S2). Similarly, no discernible correlation with ERH was observed.

When we color-coded the training dataset in Fig. 1b–d based on aerosol viscosity (the logarithm of viscosity estimated using the AIOMFAC, $\log_{10}\eta$), a clear trend emerged:

aerosols with lower viscosity generally exhibited higher ERH, while those with higher viscosity tended to have lower ERH. This trend is further supported by the scatter plot of aerosol ERH versus viscosity (Fig. 1a), which shows an apparent negative correlation. These results indicate that viscosity may play a key regulatory role in aerosol efflorescence and could be a critical predictor for ERH. We also note that the RH dependence of aerosol viscosity can contribute in the observed viscosity–ERH correlation, and we will discuss this in detail in the following section.

3.2 Characterizing Efflorescence regimes in viscosity–humidity space

Building on the correlation between viscosity and mean ERH values presented in Fig. 1, we develop a viscosity–ERH framework (Fig. 2) that establishes quantitative boundary lines in viscosity–humidity space. The boundary lines distinguish efflorescence and non-efflorescence regions, providing a simplified yet physically constrained representation of aerosol phase-transition behavior. The fitted equation for the boundary line is

$$\text{ERH}(\%) = (-10.23 \times \log_{10}(\eta/(\text{Pa s})) + 27.40) \times 100\%. \quad (5)$$

The R^2 of the linear fit is 0.65, which suggests a significant correlation between aerosol viscosity and aerosol ERH. This indicates that aerosol viscosity serves as a useful descriptor of efflorescence behavior across diverse organic–inorganic systems. Interestingly, we found that although aerosol efflorescence occurs due to the crystallization of aqueous inorganic salt, the correlation between aerosol viscosity and the normalized aerosol ERH defined as $\text{ERH}/\text{ERH}_{\text{Ing}}$ (where ERH_{Ing} is the efflorescence relative humidity of the corresponding inorganic salt system) was even weaker ($R^2 = 0.59$, Fig. S3). Moreover, the linear fit for organic–ammonium sulfate systems exhibits a similar correlation between ERH and viscosity, with comparable R^2 , slope, and intercept to those obtained for the full dataset (Fig. S4). Since the AIOMFAC calculation accounts for inorganic salts, these results suggest that the observed viscosity–ERH relationship is broadly consistent across the inorganic salts represented in this study.

The resulting viscosity–ERH framework provides a quantitative boundary for characterizing efflorescence regimes in viscosity–humidity space and for assessing the likelihood of efflorescence under a given combination of viscosity and ambient RH. Notably, the fitted relationship intersects $\text{ERH} = 0$ at a viscosity reaches $4.76 \times 10^2 \text{ Pa s}$, with a corresponding 95 % confidence interval is $[1.05 \times 10^2, 3.24 \times 10^3] \text{ Pa s}$. Within the dataset examined here, aerosols with viscosities above this threshold did not exhibit measurable efflorescence. Accordingly, this threshold can be interpreted as a practical boundary separating efflorescence and non-efflorescence regions, suggesting that organic–inorganic

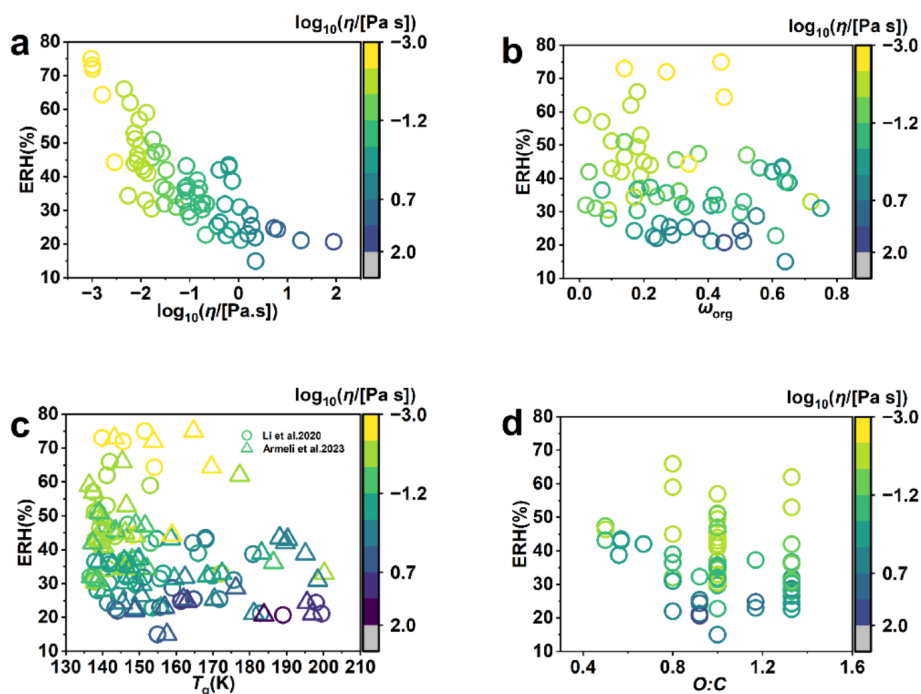


Figure 1. Relationships between the ERH of organic–inorganic mixed aerosols. (a) the logarithm of viscosity ($\log_{10}(\eta/[\text{Pa s}])$), (b) the organic mass fraction (w_{org}), (c) the glass transition temperature ($T_g(w_{\text{org}})$, K) calculated using the methods of Li et al. (2020) and Armeli et al. (2023), respectively, and (d) O : C ratio. The color bar denotes the aerosol viscosity in ($\log_{10}(\eta/[\text{Pa s}])$). All empirical ERH values were measured within a temperature range of 298 ± 10 K, and all viscosity values were calculated at 298 K.

aerosols with viscosities exceeding 4.76×10^2 Pa s are unlikely to undergo efflorescence. The complete viscosity–ERH model is expressed as:

$$\text{ERH}(\eta) = \begin{cases} (-10.23 \times \log_{10}\eta + 27.4) \times 100\%, & \eta < 4.76 \times 10^2 \text{ Pa s} \\ 0, & \eta \geq 4.76 \times 10^2 \text{ Pa s}. \end{cases} \quad (6)$$

It should be emphasized that the empirical data utilized to derive this regression model were obtained at a constant temperature of 298 K. Given the well-established temperature dependence of both ERH and viscosity, the applicability of the current framework is limited to standard room temperature conditions (298 K). Extrapolating this model to broader temperature regimes would necessitate further temperature-specific calibrations.

The uncertainty of the framework reflects the scatter of the experimental ERH data, the stochastic nature of nucleation, and the uncertainty in AIOMFAC-predicted viscosity. The current analysis is mainly applicable to internally mixed organic–inorganic aerosols containing ammonium sulfate, ammonium nitrate, sodium chloride, or potassium chloride, with organic components primarily consisting of water-soluble C, H, and O, containing compounds, such as dicarboxylic acids and sugars.

To further examine the structure underlying the observed relationship, we analyzed (i) viscosity variations for fixed chemical systems across RH (20 %–60 %) and (ii) the relationship between viscosity evaluated at fixed RH (from 20 %–90 %) and ERH across different compositions. The slopes of the viscosity–RH relationships for representative systems range from -10 to -60 (Table S12 and Fig. S7); therefore, the fitted relation between ERH and viscosity can be interpreted as an apparent representation of the combined effects of compositional variability across different aerosol compositions.

The correlation between viscosity and ERH is greatly reduced when RH is held constant (Fig. S8). However, we found that the reduction is not uniform across RH. The highest correlations occur at intermediate RH values that overlap with the ERH range of many atmospheric aerosol systems ($R^2 = 0.29$ at RH = 60 % and $R^2 = 0.24$ at RH = 50 %), whereas substantially weaker correlations are observed at RH values well below or above the typical ERH range (e.g., $R^2 = 0.15$ at RH = 20 % and $R^2 = 0.08$ at RH = 90 %). Although these correlations remain weak overall and are insufficient to establish an independent predictive relationship, they suggest that viscosity may still contribute to variations in observed ERH under conditions relevant to efflorescence. We therefore interpret these results as indicating that the RH dependence of viscosity explain the majority of the observed viscosity–ERH relationship, while the potential contribution

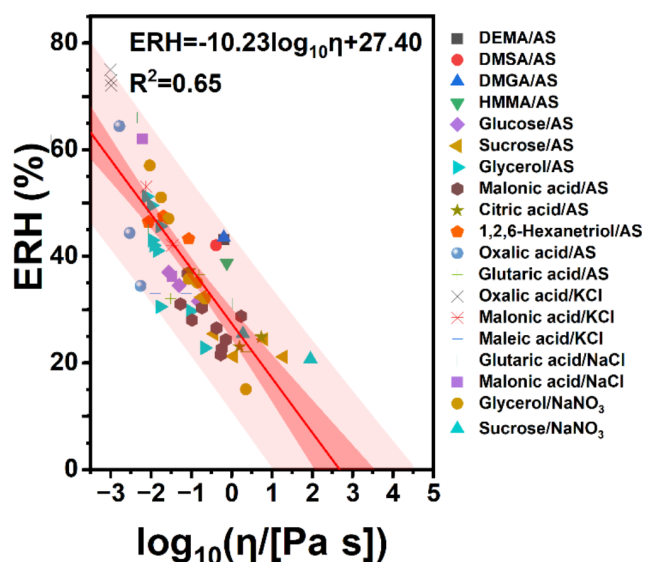


Figure 2. Linear fitting of aerosol ERH for different organic–inorganic aerosols by using viscosity ($\log_{10}\eta$) as the predictor. Data points are color- and shape-coded to distinguish between different organic and inorganic aerosol species. The red solid line represents the linear regression fit between viscosity and ERH. The dark pink band indicates the 95 % confidence interval, while the light pink band shows the prediction interval. AS refers to ammonium sulfate, DEMA represents Diethyl malonic acid, DMSA represents 2,2-Dimethyl succinic acid, DMGA represents 3,3-Dimethyl glutaric acid, HMMA represents DL-4-Hydroxy-3-methoxymandelic acid and DHBA represents 2,5-Dihydroxybenzoic acid. All empirical ERH values were measured within a temperature range of 298 ± 10 K, and all viscosity values were calculated at 298 K.

from viscosity itself cannot be excluded and warrants further investigation.

Accordingly, this consolidates the interpretation of the viscosity–ERH framework as an empirical boundary representation in viscosity–humidity space that captures variability arising from organic composition and organic–inorganic interactions. In this framework, aerosol viscosity is not regarded as a standalone predictor of ERH, rather, ambient RH and aerosol viscosity are treated as independent inputs, and their position relative to the fitted boundary is used to characterize the expected aerosol state (aqueous, efflorescing, or highly viscous/non-efflorescing).

We further performed multivariate regression incorporating viscosity ($\log_{10}\eta$), O : C ratio, ω_{org} , and $T_g(\omega_{\text{org}})$ as independent variables. The four-variable model demonstrated an improved R^2 of 0.69, which is significantly different from the ERH–viscosity model (Sig. F change < 0.05, Table S10) due to the major contribution of viscosity and $T_g(\omega_{\text{org}})$ (both showed $p < 0.05$, Table S10). We therefore established a bivariate viscosity– T_g –ERH model, and the analysis of variance (ANOVA) showed a Sig. F change (0.949) much greater than 0.05 with the same R^2 of 0.69. This indicated that

there was no significant difference between the viscosity– T_g –ERH model and the four-variable model (Table S11), and O : C and ω_{org} did not significantly improve predictive performance. The viscosity– T_g –ERH model is expressed as:

$$\text{ERH}(\eta, T_g) = (0.25 \times (T_g(\omega_{\text{org}})/K) - 12.58) \times \log_{10}(\eta/(\text{Pa s})) - 13.28 \times 100 \% \quad (7)$$

Although adding parameters typically increases the coefficient of determination (R^2) by capturing additional variance, the inclusion of $T_g(\omega_{\text{org}})$ improved R^2 by only 0.04. In contrast, the viscosity–ERH framework achieves comparable explanatory capability while requiring fewer assumptions. As a result, it provides a simpler and more broadly applicable approach for characterizing efflorescence regimes and phase-transition boundaries across diverse internally mixed organic–inorganic aerosols.

Moreover, the viscosity– T_g –ERH model exhibits practical limitations. The $T_g(\omega_{\text{org}})$ does not account for the effects of inorganic salts due to the limited availability of apparent T_g values for inorganic components. As shown in Fig. S1, we estimated the T_g of organic/NaNO₃ mixtures by incorporating the T_g of pure NaNO₃ into the Gordon–Taylor framework, using a k_{GT} value of 0.5 extrapolated from Dette and Koop. (Dette and Koop, 2015) The result indicates that, regardless of whether the T_g of pure inorganic salt or water is used, the calculated $T_g(\omega_{\text{org}})$ exhibits no apparent correlation with ERH.

We also note that other parameters, such as aerosol pH, can affect the ERH of organic/inorganic aerosol (Sun et al., 2023); however, to the best of our knowledge, no consistent trend has been established regarding the direction or magnitude of this effect, since the influence of pH is highly dependent on the specific organic composition. Given that the modeled viscosity accounts for organic functional groups, the influence of pH may be implicitly reflected within the overall model uncertainty. Future studies should incorporate more inorganic T_g data and systematically investigate pH effects to provide more accurate $T_g(\omega_{\text{org}})$ and pH values and better evaluate the role of both parameters in the viscosity–ERH relationship.

We then validated the established viscosity–ERH model using an independent dataset (32 data points) compiled from the literature. As depicted in Eq. (6), a threshold viscosity of 4.76×10^2 Pa s distinguishes between crystallized and amorphous solid phases in atmospheric aerosols. Figure 3a compares the experimentally measured ERH in the validation set with model predictions. For viscosities < 4.76×10^2 Pa s (data points to the left of the red dashed line), all validation data fall within the predicted confidence intervals, supporting the consistency of the viscosity–ERH relationship across independent datasets.

The viscosity–ERH relationship defines a boundary line and a threshold value in viscosity–humidity space, which can be used in combination with ambient RH value and aerosol

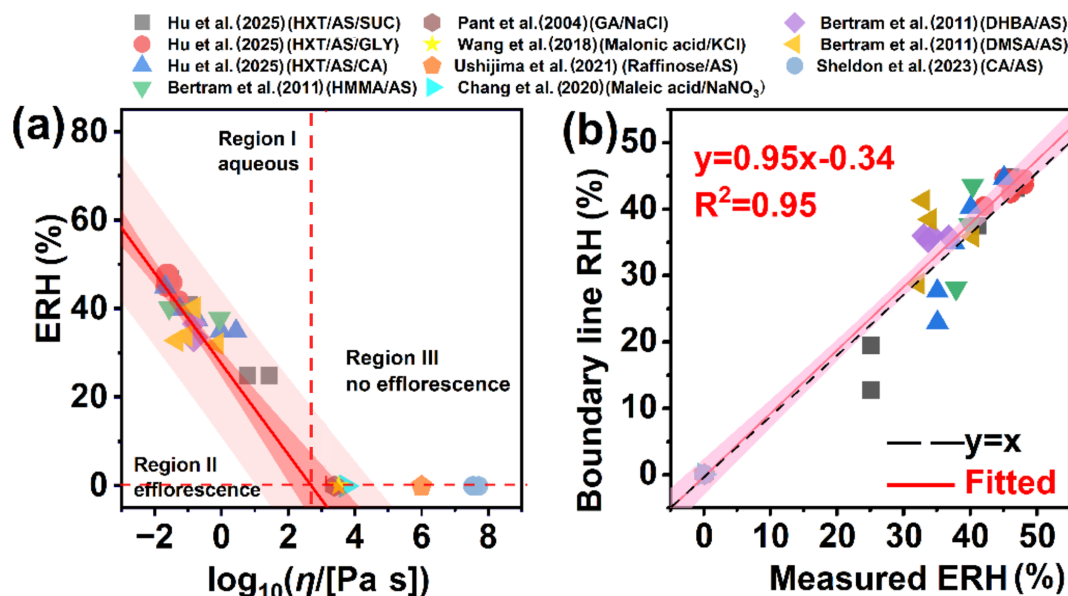


Figure 3. (a) Comparison between the validation dataset and viscosity–ERH boundary framework (Eq. 5), showing three distinct regions in viscosity–humidity space: Region I (aqueous state), Region II (efflorescence-favorable conditions), and Region III (non-efflorescence or highly viscous state). (b) Comparison between experimentally measured ERH values and the boundary line RH values derived from the viscosity–humidity relationship. The black dashed line denotes the $y = x$ reference line, and the red line represents the linear regression fit to the data, along with the corresponding coefficient of determination, $R^2 = 0.95$. The pale pink shading represents the 95 % confidence interval. DEMA represents Diethyl malonic acid, DMSA represents 2,2-Dimethyl succinic acid, DMGA represents 3,3-Dimethyl glutaric acid, HMMA represents DL-4-Hydroxy-3-methoxymandelic acid and DHBA represents 2,5-Dihydroxybenzoic acid, CA represents citric acid, SUC represents Sucrose, GA represents glutaric acid, HXT represents 1,2,6-hexanetriol. All empirical ERH values were measured within a temperature range of 298 ± 10 K, and all viscosity values were calculated at 298 K.

viscosity to interpret aerosol phase behavior within three distinct regions (Fig. 3a). In Region I (above the red regression line), aerosols remain in an aqueous state. In Region II (below the regression boundary but with $\eta < 4.76 \times 10^2$ Pa s), aerosols are kinetically favorable for nucleation and crystal growth, corresponding to efflorescence. In Region III ($\eta > 4.76 \times 10^2$ Pa s), aerosols remain in a highly viscous state and do not exhibit measurable efflorescence, instead forming amorphous solids.

To validate the viscosity–ERH boundary framework derived from linear regression (Eq. 6), we compared the experimentally measured ERH values with the boundary line RH estimated from the fitted relationship (Fig. 3b). The data points cluster around the $y = x$ reference line (black dashed line), with the fitted line (red solid line) showing close alignment and a slope near unity ($R^2 = 0.95$), indicating strong consistency between the empirical ERH values and the viscosity–ERH boundary representation. When excluding the (0,0) data points (Fig. S5), the slope remains close to unity, while the coefficient of determination decreases but remains significant ($R^2 = 0.68$), reflecting increased scatter at low ERH conditions. In addition, the viscosity– T_g –ERH framework was evaluated using the same dataset, excluding systems for which T_g could not be estimated. This extended

framework shows comparable consistency with the experimentally reported ERH values (Fig. S6).

3.3 Physical interpretation of the viscosity–ERH framework

Aerosol efflorescence is a kinetically controlled process involving: (1) formation of critical crystal nuclei in supersaturated droplets; (2) subsequent crystal growth, and (3) water evaporation (Zhang et al., 2012). In highly viscous aerosols, mass transfer limitations significantly alter nucleation dynamics, making nucleus formation the rate-limiting step (Ji et al., 2017; Wang et al., 2017). Turnbull and others showed that the nucleation rate can be expressed as the product of a pre-exponential factor and two thermally activated rate terms (Hollomon and Turnbull, 1953; Kelton et al., 1983; Turnbull, 1969):

$$\tau_x^{-1} = v \exp \left[\frac{-W(T)}{kT} \right] \exp \left[\frac{-\Delta G(T)}{kT} \right] \quad (8)$$

where $W(T)$ represents the kinetic barrier associated with atomic rearrangement, and $\Delta G(T)$ is the thermodynamic free energy barrier for nucleus formation. The term $\exp(W(T)/kT)$ represents the thermally activated atomic rearrangement rate (i.e., atomic mobility) in liquids and is in-

versely related to viscosity (η^{-1}). Here, we introduce an approximate proportional relationship between the kinetic barrier term and the logarithm of viscosity:

$$\frac{-W(T)}{kT} = -A \log_{10} \eta \quad (9)$$

where A is a constant linking $-W(T)/kT$ with $-\log_{10} \eta$. Substituting this relation into Eq. (8) and taking $\log_{10}$ gives:

$$\log_{10}(\tau_x^{-1}) = -A \log_{10} \eta + \log_{10} v - \frac{\Delta G(T)}{2.303kT}. \quad (10)$$

The viscosity–ERH framework established in this study is qualitatively consistent with Eq. (10), in which increasing viscosity is associated with reduced molecular mobility and lower effective nucleation rates. In this context, $\log_{10} \eta$ can be viewed as a practical indicator for the kinetic energy barrier $W(T)$ on nucleation and crystal growth. The intercept achieved in Eq. (5) can therefore be associated with $\log_{10} v - \Delta G(T)/2.303kT$. However, as discussed in Sect. 3.2, the observed viscosity–ERH relationship is strongly influenced by the RH dependence of viscosity. Therefore, the present results should not be simply interpreted as establishing a unique mechanistic relationship between viscosity and ERH. Rather, the viscosity–ERH framework provides an empirical description of the conditions under which efflorescence may occur for a broad range of internally mixed organic–inorganic aerosol systems. At the critical viscosity threshold, the aerosol transitions from a liquid to a semi-solid phase, creating an immense kinetic barrier to molecular diffusion. Consequently, the nucleation induction time is prolonged so significantly that efflorescence is effectively inhibited, even over the extended residence times typical of aerosols in the atmosphere (days to weeks).

Thermodynamics-based approaches, such as that described by Hodas et al. (2016), predict salt crystallization and ERH based on the degree of supersaturation relative to the ion activity product (IAP) at the point of saturation of the species involved in the crystalline phase (i.e., IAP_{sat}). These frameworks capture the fundamental requirement that crystallization can only occur when the aqueous phase becomes thermodynamically supersaturated with respect to the relevant solid phase.

In this study, we consider aerosol conditions near ERH, where particles are expected to exhibit high ionic strength due to reduced water content. Under these conditions, efflorescence can be viewed as a coupled thermodynamic–kinetic process. Thermodynamic supersaturation provides the driving force necessary for crystallization, whereas nucleation and crystal growth remain subject to kinetic limitations. Thus, thermodynamic and kinetic perspectives are complementary: thermodynamic conditions determine whether crystallization is possible, while kinetic factors influence when crystallization occurs. In this context, aerosol viscosity is considered as a practical descriptor of molecular mobility

and diffusion limitations, rather than a standalone determinant of efflorescence.

While the viscosity–ERH framework characterizes distinct efflorescence regimes across diverse aerosol systems, its application to predicting aerosol phase behavior in viscosity–humidity space has inherent limitations. As highlighted by previous studies (Ciobanu et al., 2010; Kuwata and Martin, 2012), in complex organic–inorganic systems, liquid–liquid phase separation (LLPS) frequently occurs, forming core–shell morphologies. In such core–shell aerosols, the inner aqueous inorganic-rich phase may maintain a low, liquid-like viscosity comparable to that of pure inorganic droplets, yet still exhibit a suppressed ERH. This suppression can arise because the organic coating alters the interfacial energy or acts as a physical barrier, independent of the inner core’s viscosity (Ciobanu et al., 2010). Consequently, the occurrence of efflorescence is influenced by a combination of thermodynamic driving force, particle morphology, interfacial properties, particle size, and molecular mobility. The viscosity–ERH framework presented here should therefore be viewed as an empirical boundary description of observed aerosol behavior rather than a complete mechanistic model of crystallization.

4 Conclusions and implications

Our results demonstrate that aerosol viscosity provides a useful framework for characterizing efflorescence behavior in internally mixed organic–inorganic aerosols. Although inclusion of both viscosity and T_g slightly improves model performance, the viscosity–ERH framework offers broader applicability and greater simplicity for diverse aerosol systems. The framework remains valuable for describing aerosol phase transitions because it organizes observed behavior into distinct regimes in viscosity–humidity space and defines boundary lines separating different phase states. Within the dataset examined here, three behavioral regions are identified: (i) an aqueous regime above the fitted boundary line, (ii) an efflorescence regime below the boundary line when viscosity remains below the threshold value, and (iii) a non-efflorescence regime at viscosities exceeding approximately 4.76×10^2 Pa s. This provides a physically constrained representation of aerosol phase-transition behavior.

Regional viscosity predictions by Zhang et al. suggest that in southern and northeastern China (Zhang et al., 2024), SOA are predominantly liquid or have low viscosities ($< 10^4$ Pa s); in central and northeastern China, SOA typically exhibit semi-solid viscosities ($10^5 - 10^8$ Pa s); while highly viscous or glassy SOA ($\eta > 10^8$ Pa s) are mainly present in the northwest. Although SOA viscosity may significantly increase by several orders of magnitude as water content (and consequently aerosol water activity) decreases (Hosny et al., 2016; Renbaum-Wolff et al., 2013; Saukko et al., 2012; Song et al., 2016a; Yli-Juuti et al., 2017), SOA tend to become more oxi-

dized and internally mixed with secondary inorganic aerosols during atmospheric aging. This aging process significantly lowers the overall viscosity of SOA due to the greater hygroscopicity of more oxygenated organics and inorganic salts (Dette and Koop, 2015). Consequently, in more developed and urbanized regions like southern and northeastern China, atmospheric aerosols are more likely to undergo efflorescence and crystallization, leading to hysteresis in aerosol size and water contents during humidification and dehumidification processes. From a broader atmospheric perspective, the implications of aerosol phase state should be considered on regional to global scales. Shiraiwa et al. (2017) showed that SOA phase state varies strongly with relative humidity and temperature, with SOA in the planetary boundary layer being predominantly liquid in humid tropical and polar regions, semi-solid in mid-latitudes, and solid over dry lands, while SOA in the middle and upper troposphere are expected to be mostly glassy. The hysteresis induced by aerosol efflorescence has important implications for modeling aerosol size evolution, optical properties, and chemical reactivity.

It is important to note that the aerosol viscosities in this study were derived from thermodynamic model predictions, as in-situ measurement techniques for atmospheric aerosol viscosity remain largely unavailable. Although substrate-based methods – such as poke-flow cytometry and fluorescence lifetime imaging microscopy (FLIM) – have been successfully employed to determine SOA viscosity under laboratory conditions, substrate effects must be carefully considered (Grayson et al., 2015; Hosny et al., 2013). This highlights an urgent need to develop techniques that can directly measure aerosol viscosity at a given RH in ambient environments and to validate viscosity-dependent phase transition predictions. Optical tweezers is a promising tool for quantitative viscosity measurements of levitated particles; however, the technique requires further advancement to measure sub-micron organic aerosols in real atmospheric conditions (Reid et al., 2014). Analyses in this study suggest a possible contribution of viscosity, particularly at RH values relevant to efflorescence, but the available evidence is not sufficiently conclusive to establish such a relationship. Therefore, whether viscosity can ultimately serve as a more direct mechanistic predictor of ERH remains an important topic for future investigation. Furthermore, quantitative evaluation is needed to assess the extent to which viscosity-based predictive tools for aerosol phase state influence simulations of climate and air quality.

Code and data availability. The data supporting this article have been included in the main text and as part of the Supplement: Tables S1–S12, Fig. S1–S8.

Supplement. The supplement related to this article is available online at <https://doi.org/10.5194/acp-26-9967-2026-supplement>.

Author contributions. SC and QH analyzed and visualized the data and wrote the original draft. SC conducted the experiments and model calculations. QH created the original research framework. QH and YL reviewed and edited the paper. QH, PL, and YHZ supervised the study. All authors discussed the results and contributed to the article editing.

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

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