Role of Contamination in Optimal Droplet Production by Collective Bubble Bursting

B. Néel 1, M. A. Erinin 1, and L. Deike 1,2

1 Department of Mechanical & Aerospace Engineering, Princeton University, Princeton, NJ, USA, 2 High Meadows Environmental Institute, Princeton University, Princeton, NJ, USA

Abstract Gas bubbles bursting at the sea surface produce drops, which contribute to marine aerosols. The contamination or enrichment of water by surface-active agents, of biological or anthropogenic origin, has long been recognized as affecting the bubble bursting processes and the spray composition. However, despite an improved understanding of the physics of a single bursting event, a quantitative understanding of the role of the physico-chemical conditions on assemblies of bursting bubbles remains elusive. We present experiments on the drop production by millimetric, collective bursting bubbles, under varying surfactant concentration and bubble density. We demonstrate that the production of supermicron droplets (with radius larger than 35 µm) is non-monotonic as the surfactant concentration increases. The bursting efficiency is optimal for short-lived, sparsely distributed and non-coalescing bubbles. We identify the combined role of contamination on the surface bubble arrangement and the modification of the jet drop production process in the bursting efficiency.

Plain Language Summary Understanding the production of primary ocean spray aerosol and its dependence on meteorological and environmental variables is critical for modeling potentially health-adverse effects from dispersants used to break down oil spills, as well as radiative processes and cloud microphysical properties. We demonstrate experimentally that the coupling between the chemical properties of contaminated water and the physical mechanism of droplet production when a bubble bursts controls the efficiency of aerosols production during collective bubble bursting, and reveals an optimal production regime at intermediate contamination. This fundamental study paves the way to improved predictions of aerosol production in contaminated water.

1. Introduction

A gas bubble sitting at the surface of water ends its life in a burst. When it pops, it may eject drops, according to two mechanisms: the cap film puncturing, retraction and destabilization into a mist of film drops (Blanchard, 1963; Lhuissier & Villermaux, 2012), or the later collapse of the cavity into a vertical upwards jet that destabilizes into jet drops (Brasz et al., 2018; Duchemin et al., 2002; Ghabache et al., 2014; Ghabache & Séon, 2016; Spiel, 1994, 1997; Woodcock et al., 1953). These two production mechanisms have been extensively studied and documented for single bubbles, leading to various scalings laws to describe the mean size, distribution and number of ejected film (Lhuissier & Villermaux, 2012) and jet drops (Berny et al., 2021; Deike et al., 2018; Gañán-Calvo, 2017; Gañán-Calvo & López-Herrera, 2021; Gordillo & Rodríguez-Rodríguez, 2019; Lai et al., 2018) as a function of the controlling non-dimensional length scales $R_b/l_p = L$ (Laplace number) and $R_b/l_1 = \sqrt{Bo}$ (Bo is the Bond number), with $K_b$ the bubble size, $l_p = \mu^2/\gamma \rho$ the visco-capillary length, $l_1 = \sqrt{\gamma \rho g}$ the gravity-capillary length, $\mu$ and $\rho$ the liquid viscosity and density, $\gamma$ the surface tension and $g$ gravity.

These scaling laws derived from a single bursting event can in principle be extrapolated to the production of droplets in a marine environment, using estimations of the number of bubbles at the ocean surface through the whitecap coverage method or the breaking wave distribution (Deike, 2022; Lewis & Schwartz, 2004; Veron, 2015). However, large uncertainties remain due to the wide range of scales involved as well as an incomplete understanding of the collective behavior of bubbles at the water surface (Modini et al., 2013; Néel & Deike, 2021; Sellergren et al., 2006) and the complex role of water contamination on individual bursting processes (Constance-Amores et al., 2021; Poulain et al., 2018). Therefore, a physical description of the collective bursting of bubbles in contaminated water is necessary, since it controls the formation, stability, and decay of foams on the ocean surface (Garrett, 1967; Peltzer & Griffin, 1987). Implications are far reaching, from remote sensing of the ocean surface (Callaghan et al., 2017), which is central to the study of marine aerosols, sources of cloud condensation nuclei...
and light scattering particles, production from bursting bubbles (Lewis & Schwartz, 2004; Wang et al., 2017); as well as to better understand the generation and composition of aerosols associated with chemical agents used in oil-spill mitigation (Afshar-Mohajer et al., 2018; Feng et al., 2014; Sampath et al., 2019).

The seemingly basic question: “do bubbles burst collectively the same way as they burst individually?” often assumed in existing models, remains unanswered. Although the physico-chemical properties of water have long been identified in modulating the droplet production and final sea spray aerosol composition (Cochran et al., 2017; Collins et al., 2014; Sellegri et al., 2006; Wang et al., 2017), its precise effects on the surface bubbles assembly, or the drops production mechanisms remain poorly understood in the context of large bubbles rafts (Néel & Deike, 2021). Various experimental studies have attempted to quantify the role of water contamination by surfactant, biological activity or seawater temperature, and have yielded results sometimes contradicting each other (Frossard et al., 2019; Modini et al., 2013; Prather et al., 2013). The trends of production efficiency with contamination for assemblies of bubbles drifting at the surface, directly observed in laboratory experiments mimicking field conditions (Prather et al., 2013), produced from bubbling device in seawater (Frossard et al., 2019; Sellegri et al., 2006), or simulated in the laboratory from breaking waves experiments (Modini et al., 2013), remain unclear.

Here we present the results of a laboratory experiment which studies the spray production by collective bubble bursting at the surface of water, contaminated by surfactant. The experiment is designed to (a) produce narrow, normal distributions of millimetric bubbles in the bulk, (b) observe the arrangement and collective dynamics and statistics of the bubbles at the water surface, and (c) measure the drop production directly, immediately above the water surface. We control the water surface contamination by minute additions of surfactants Triton X-100 and sodium dodecyl sulfate (SDS). This follows our work quantifying how the surface bubble size distribution and raft dynamics depends on the surfactant concentration and the rate of bubbles arriving at the water surface (Néel & Deike, 2021).

The existence of an optimal drop production with respect to contamination, a key finding of this study, is illustrated in §2. §3 investigates the statistics of the drop production, in direct correlation with the surface bubbles assembly. §4 generalizes by comparing the bursting efficiency of two surfactants, and §5 discusses the mechanisms of drop production.

### 2. Contamination Controlled Optimal Production

Bubbles are produced at the bottom of transparent acrylic tank by way of 16–48 needles (inner diameter 203 or 305 μm) connected to a pressurized air chamber and arranged on a ring, and producing each a regular stream of quasi-monodisperse, millimetric bubbles. They are visualized and detected in the bulk of the tank, from the side, and characterized by their volume-equivalent radius $a$ (Figure 1a). The tank, with dimensions $L60 \times W60 \times H25$ cm$^3$, is open at the top, leaving the bubbles unconstrained once they reach the surface (Figure 1a). A top-down imaging allows for a direct measurement of their location and size, as illustrated in Figures 1b–1e and detailed in a recent study focusing on the surface bubble dynamics (Néel & Deike, 2021). Drops, produced by the bursting surface bubbles, are imaged and detected from the side by means of telecentric shadow imaging, which preserves the true drop size across the depth of field. The observation window, close to the surface, ensures that we measure the totality of the drop production. The drops are eventually characterized in a statistical way, for those with radius $d > 35$ μm (see Figures 1f–1i and technical details in Supporting Information S1).

Surface contamination is investigated by varying the surfactant concentration $c$ of two surfactants (SDS and Triton X-100, from Sigma-Aldrich), in levels below, around and above the coalescence transition $c_*$ (Oolman & Blanch, 1986; Yang & Maa, 1984), above which the merging of bubbles in contact is prevented by a shielding action from the surfactants ($c_* = 12 \mu$M for SDS, $c_* = 4 \mu$M for Triton X-100, as measured in Néel and Deike 2021).

Figures 1b–1i illustrates how the surface contamination (top row) affects the droplet production (bottom row) as SDS concentration increases. It demonstrates visually that the knowledge of the bulk bubbles alone (size and production rate being unchanged) is not sufficient in order to predict the drops size and production rate. Figures 1b–1e (top row) shows representative instantaneous states of the bubbles at the surface. The concentrations of SDS we highlight (0, 16, 128 and 2,000 μM) span across the coalescence transition ($c_* = 12 \mu$M), and the surface
bubbles exhibit a large variety of arrangements, despite coming from identical gaussian bulk bubble populations. At low contamination (Figure 1b), surface bubbles merge and burst quickly, resulting in a small number of large, short-lived surface bubbles, typically $A_{10^1}$ bubbles per $A_{cm^2}$ (see also Figure 3a below). At high contamination (Figure 1e), coalescence is prevented and bubbles form dense rafts drifting at the water surface, with surface densities reaching up to $10$ bubbles per $A_{cm^2}$. In between, an intermediate regime (Figures 1c and 1d) is characterized by similar sized, short-lived bubbles that do not merge, assembling at most in small clusters before bursting, with between 0.1 and 1 bubbles per $cm^2$. This intermediate regime is due to the decoupled evolutions of the bubble merging and bursting rates with respect to the surfactant concentration, the coalescence transition occurring at a lower concentration than the onset of the stabilizing effect (Néel & Deike, 2021). The measured bursting and merging rates for these conditions are given in the Supporting Information S1 and Néel and Deike (2021).

Figures 1f–1i (bottom row) shows time-integrated images of the spray production corresponding to the four surface states shown in the top row of images. Very clearly, the drop production (in the detected range of radii $r_d > 35\ \mu m$) is optimal in the intermediate regime where surface bubbles do not coalesce anymore, and form sparse rafts of few elements with relatively short life-times (Figures 1g and 1h). Many drops are ejected, with a wide size distribution. For a contamination below the coalescence transition $c < c_2$, fewer and larger bubbles are present at the surface, and less drops are produced in the $r_d > 35\ \mu m$ range (Figure 1f). Far above the concentration threshold $c \gg c_2$, (Figure 1i), the drops production decreases significantly, due to the increase in surface bubble lifetime, which leads to the formation of three-dimensional rafts.

3. Surface Transfer Function and Droplet Distribution

The number of spray drops and their size distribution is determined by the rearrangements of the bubbles on the surface. This surface arrangement controls the bubble sizes, their bursting mechanism (and their efficiency at producing drops), effectively acting as a surface transfer function. Figure 2 shows bulk bubble $R_b$, surface bubble $R_s$ and drop $r_d$ radius distributions, under identical bubbling conditions and increasing concentrations of SDS. It demonstrates the importance of considering the statistical details of the surface rearrangements: an identical normal distribution of bubbles in the bulk leads, under a varying surface contamination, to various size distributions of the surface bubbles and of the ejected drops.
Figure 2. From monodisperse bulk bubble size distribution to drops production: a surface transfer function, under an increasing concentration of surfactant (SDS from 1 μM to 1 mM). (a) Bulk bubble distribution $N_b(R_b)$ is almost unchanged when increasing the contamination (colored lines), with mean diameter $\langle R_b \rangle = 1.5 \pm 0.1$ mm. (b) Surface bubble distribution $N_s(R_s)$, shifting from a broad distribution at low contamination when bubbles are able to coalesce to a narrower distribution close to $N_s(R_s)$ at high contamination. (c) Corresponding ejected drop distribution $N_d(r_d)$. The number and size of drops ejected evolve with the increased contamination and surface bubble distribution.

Figure 2a shows the bubble distribution in the bulk for the same bubbling conditions and increasing concentrations of surfactant SDS. The concentration is shown in colors and spans from $c = 0$ and 1 μM to $c = 1$ mM a fraction of the critical micellar concentration (CMC $c_m = 8.2$ mM), crossing the coalescence transition $c_c = 12$ μM. The different bulk bubble size distributions $N_b(R_b)$ (for bubbles larger than $R_b > 400$ μm) all exhibit the same gaussian statistics, with a mean size $\langle R_b \rangle = 1.5$ mm which remain unchanged under a variation of the SDS concentration over three orders of magnitude: the addition of surfactant has no noticeable influence on the bubble production in the bulk.

Figure 2b shows the corresponding surface bubble size distributions $N_s(R_s)$. As the surfactant concentration $c$ is increased, the surface distributions are modified in two ways. First, they narrow down around the bulk injection size $\langle R_b \rangle$, a direct consequence of the suppression of bubble coalescence events for concentrations $c > c_c$. Second, the magnitude of the distribution increases, with more bubbles populating the surface, a consequence of the increased stability, or lifetime, of the surface bubbles.

Figure 2c shows the drop size distribution $N_d(r_d)$, for drops in the range $r_d \geq 35$ μm. Deionized and barely contaminated water (red and yellow lines) produce two categories of drops. Above $r_d \geq 200$ μm, a narrow peak centered around $r_d = 250$ μm is immediately identified. For drops below $r_d \leq 200$ μm, the smaller the drops, the more of them are produced (down to the measurement cutoff $r_d = 35$ μm). The total number of drops produced at low contamination is lower than for higher surfactant concentrations, a consequence of the corresponding low surface bubbles density. When the surfactant concentration is increased to 4 μM of SDS (green line), the drop size distribution starts to transition, following the changes in the surface bubble population (Figure 2b). At this transition point, the total number of drops below $r_d < 200$ μm begins to increase with the concentration, with the peak initially around 250 μm gradually shifting to drop sizes close to 150 μm at the highest $c$. For SDS concentrations in the range $c = 10500$ μM, the drop size distributions are broader, and do not exhibit two distinct modes. For these concentrations, drops are produced in larger numbers, up to three orders of magnitude more when compared to the clean water case. When the surfactant concentration is increased further, above $c \geq 1$ mM of SDS, the number of drops decreases at all sizes $r_d > 35$ μm (dark blue line). This corresponds to a surface state where bubbles accumulate at the surface too quickly, when compared to their lifetime. They form large drifting rafts or even three-dimensional foams, which ultimately inhibit the drop ejection. Similar results are observed when using a different surfactant (Triton X-100) and for various bubble injection rates (see Supporting Information S1).

4. Bursting Efficiency and Mean Drop Size

We now relate the number of drops produced to the number of bubbles at the surface, as a function of the surface contamination. Figure 3a plots the surface bubble density $\rho_s = \int_{R_b} N_b(R_b) dR_b$ (per unit area, black circles, left axis) and the drop density $\rho_d = \int_{r_d>35 \mu m} N_d(r_d) dr_d$ (per unit volume of air, green crosses, right axis) as a function of SDS concentration $c$, under the same bubbling conditions. As $c$ is increased from 0 to $c \approx 0.5$ mM, a fraction
of the CMC, both \( \rho_b \) and \( \rho_d \) increase. However, they do so at different rates: the surface bubble density \( \rho_b \) first undergoes an intermediate plateau around the coalescence transition \( c_c = 12 \ \mu\text{M} \). Then \( \rho_b \) rises sharply to the maximal value \( 1/2 \sqrt{3}(R_0)^2 \), the density of a close packing of disks with radius \( R_0 \) where bubbles saturate the surface entirely. If the stability (lifetime) of the surface bubbles is increased even further for the same bubbling rate and total surface area available, they can start to pile up and form three-dimensional foams. For a given surfactant concentration the onset of foaming can be delayed in the experiment by providing a larger surface area over which bubbles can collect on, or changing the bubbling rate, as illustrated in the Supporting Information S1. In parallel to the evolution of \( \rho_b \), the drop density \( \rho_d \) increases sharply between \( c = 4 \) and \( 10 \ \mu\text{M} \) followed by a more gradual increase until \( c \approx 800 \ \mu\text{M} \). Next, when \( \rho_b \) saturates for \( c \geq 800 \ \mu\text{M} \), the drop density \( \rho_d \) decreases abruptly, where the close packing of the bubbles, and the formation of a three-dimensional foam, inhibit the drops ejection.

We combine surface bubble density \( \rho_b \) and drop density \( \rho_d \) into a non-dimensional bursting efficiency \( e = H \rho_b / \rho_s \), where \( H \) is a vertical height above the water surface where drops are measured (see Supporting Information S1). The bursting efficiency \( e \) captures how different regimes of contamination affect drop ejection efficiency, under the same bulk bubble production, within the range of drop size measured (\( r_d > 35 \ \mu\text{m} \)). The bursting efficiency is plotted as a function of the surfactant concentration in Figure 3b, for two surfactants SDS and Triton X-100, with \( c \) normalized by \( c_c \), the coalescence transition concentration (\( c_c = 12 \ \mu\text{M} \) for SDS, \( c_c = 4 \ \mu\text{M} \) for Triton X-100, Néel and Deike (2021)). The evolution of \( e \) with respect to \( c/c_c \) exhibits similar features for both surfactants at low bubbling rate (light blue symbols): the efficiency first decreases as \( c \) increases before reaching a local maximum around \( c \approx c_c \) (Triton X) to \( c \approx 8c_c \) (for SDS). The efficiency decreases at high contamination corresponding to the transition to a three-dimensional foam. For a higher bubbling rate (dark blue symbols), the behavior for Triton X-100 is identical while for SDS, no local maxima is observed due to the occurrence of packing at the surface at a lower concentration. This result indicates that while the overall behavior is similar for both surfactants, the range of optimal drop production is dependent on the type of surfactant, as well as the transition to packing at the surface. These dynamical features at the surface are themselves controlled by the interplay between the bubbling rate and the bubbles bursting and merging rates.

Figure 3c shows the mean drop radius \( \langle r_d \rangle \) as a function of \( c/c_c \) for both SDS and Triton X-100. For SDS under two different bubble production rates, \( \langle r_d \rangle \) continuously decreases with an increasing surfactant concentration, from \( \langle r_d \rangle \approx 200 \ \mu\text{m} \) for gently contaminated cases, to \( \langle r_d \rangle \approx 70 \ \mu\text{m} \) at high concentrations. In the case of Triton X-100, the mean drop size follows a similar trend as SDS until the coalescence concentration \( c_c \), then reaches a very narrow local maximum around \( c/c_c \approx 2 \) when the bursting efficiency is the highest. Drop sizes thus seem to be only affected by the surfactant type and its concentration, as experiments performed under different bubbling rates exhibit the same trend.

5. Discussion and Production Models

We show in the previous sections and Néel and Deike (2021) that the ability of the surface bubbles to merge, form clusters and large rafts, pile up in three-dimensional foams, or burst isolated, depends on the interplay between surface contamination and bubble flux to the surface. The surface contamination, in particular, induces different collective effects on the bubbles (merging, or clustering), which in turn modifies the drop production, both in size and number. The evolution of the bursting efficiency as a function of the surfactant concentration highlights a complex behavior, with the existence of an optimal regime of production. We find that this optimal efficiency

\[ \langle r_d \rangle = \frac{\langle r_d \rangle}{c_c} \]

where \( \langle r_d \rangle \) is the mean drop radius, and \( c_c \) is the coalescence transition concentration.
occurs when bubbles burst in small rafts of few elements (see Figures 1b and 1c) sparsely distributed at the surface, suggesting that collective effects in large rafts tend to inhibit droplet production by bubble bursting. In turn, the effect of surfactants on the mean drop radius and total number distributions suggests a modification of the spray production mechanisms as the surface becomes more and more contaminated.

The production of drops from an assembly of surface bubbles can be analyzed within the framework proposed by Lhuissier and Villermaux (2012) for film drops and extended to jet drops by Berny et al. (2021), which integrates scalings for the number $n(R_s)$ and size of drops (mean size $\langle r_d \rangle$ and distribution $p(r_d/\langle r_d \rangle, R_s)$) produced by a single bubble bursting over the distribution of bursting bubbles $N_d(R_s)$:

$$N_d(r_d) = \int_{R_s} \frac{N_e(R_e)n(R_e)}{\langle r_d \rangle(R_e)} p\left(\frac{r_d}{\langle r_d \rangle}, R_s\right) dR_e.$$  

(1)

Figure 4 shows examples of the surface bubble distribution $N_e(R_e)$ in clean and contaminated regimes (Figure 4a) and the corresponding drop distribution $N_d(r_d)$ (Figure 4b), together with corresponding models associated with film drops (Lhuissier & Villermaux, 2012):

$$n(R_e) = \chi_f \left(\frac{R_e}{\ell_f}\right)^2 \frac{R_e}{h_s} \frac{\eta_f R_s^{5/8} h_s^{5/8}}{\langle r_d \rangle}.$$  

(2)

and jet drops (Berny et al., 2021):

$$n(R_e) = \chi_j \left(\frac{R_e}{\ell_j}\right)^{1/3} \frac{\eta_j h_s}{\langle r_d \rangle}.$$  

(3)

$h_s$ is the surface bubble cap film thickness, related to the bubble radius $h_s = R_s^2 / \mathcal{L}$, with $\mathcal{L} \approx 20 \text{ m}$ (Lhuissier & Villermaux, 2012; Poulain et al., 2018). $\chi_f, \eta_f, \chi_j, \eta_j$ are non-dimensional prefactors for the film ($F$) and jet ($J$) drop number and mean radius, respectively, and distributions $p(r_d/\langle r_d \rangle)$ are described by Gamma (1) distributions.

For clean water (red lines), surface bubbles are sorted into two categories: surface bubbles with sizes close to $\langle R_s \rangle = 1.5 \text{ mm}$ are known to produce between 1 and 3 jet drops when bursting in clean water (Berny et al., 2020; Ghabache & Séon, 2016; Spiel, 1994). This result is confirmed in our experiments by complementary dynamical observations shown in Figures 4c and 4e. The vertical upwards motion of the drops is consistent with the jet drop production mechanism (since the expected motion of film drops is mostly sideways). These jet drops explain the narrow peak around $\langle r_d \rangle = 250 \mu \text{m}$ (Figure 4b solid red line), and are well described by the jet drop model (dotted red line) derived from single bubble bursting experiments (Berny et al., 2021; Gañán-Calvo, 2017; Lai et al., 2018). In the clean water case, larger bubbles in the exponentially-tailed surface distribution (Figure 4a dotted red line) are the result of at least one merging event. These bubbles are larger than the capillary length $R_c > \ell_c$ so that no jet drops are being produced when they burst. The resulting drops are assumed to be film drops, and the red dashed line shows the corresponding model derived from single bubble bursting experiments (Lhuissier & Villermaux, 2012). In both the jet (red dotted line) and film (red dashed line) models, we use the measured bubble surface distribution for $R_c$ in the appropriate range of production. The agreement between the measured (Figure 4b solid red line) and predicted (dashed red line) distributions is good using the coefficients derived from single bursting experiments (Berny et al., 2021; Lhuissier & Villermaux, 2012).

For moderately contaminated water (blue line), surface bubbles do not coalesce and burst either isolated or in small rafts (Figures 1b, 1c and 4a). Their surface distribution is centered around $\langle R_s \rangle = 1.5 \text{ mm}$, and they produce mostly jet drops (in the detected range, Figure 4b), as suggested by the upwards vertical motion of the ejected drops (Figures 4d and 4f). Unlike the clean water case, each bubble produces more and smaller drops, which are scattered in sizes, as illustrated in the representative sequence (Figure 4d). The drop size distribution can still be modeled by integration over the narrow surface bubbles distribution (Figure 4a dotted blue line) but now requires to modify the coefficients for the mean drop size and number of ejected drop, following the experimental results from Figures 3b and 3c, yielding the broad drop size distribution (Figure 4b dotted blue line).

The role of contamination and surface bubble rearrangement can be summarized as follow: A broad distribution of surface bubbles generates a narrow production of jet drops (in clean water), whereas a narrow distribution of
surface bubbles leads to a jet drop distribution with a significantly broader dispersion (in contaminated water). Supported by the dynamical observations of Figures 4c–4f and the satisfactory comparison to a mechanistic model, this suggests that surfactants directly modify the bubble bursting mechanism: For contaminated water, more jet drops are being produced and of smaller sizes for the same original bubble. Furthermore, we note that the distribution amplitudes are orders of magnitude apart, linking back to the bursting efficiency previously defined: For bubble radii around $R_b \approx 1.5 \text{ mm}$, the most efficient drop production mechanism seems to be jet drops.

We show that even with a precisely controlled bubble production, the drop production differs greatly between different concentrations and types of surfactants. This results from collective re-arrangements of the bubbles at the surface (characterized by the merging and bursting rates) as well as a modification of the jet drop production.
process, impacting both the size and number of ejected drops. Determining whether the influence of surfactants on the bursting processes results from collective effects or a modification of the individual jetting mechanism remains to be investigated. The resulting drop size distributions can be, and are, rationalized using the framework developed by Lhuissier and Villermaux (2012) and extended by Berny et al. (2021), but the knowledge of the surface transfer function and the changes in drop size and number being ejected for given contaminated conditions remain to be theoretically explained. The existence of an optimal production regime controlled by the interplay between the bubble production, bursting and merging rates could explain some of the apparent contradictions in the literature where these characteristics were unknown. Finally, while the present study focused on millimetric bubbles and drops above 30 μm, similar studies on smaller bubbles and drops, involving how the drop production might be affected by contamination and collective effects, may extend the present findings to sizes even more relevant to atmospheric applications, such as the origin for cloud condensation nuclei.

Data Availability Statement

Open research all data used in preparing this work are publicly available at https://doi.org/10.34770/pjmm-cp20.

References


References From the Supporting Information

