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Breakup of particle-laden droplets in airflow

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The atomisation of a suspension containing liquid and dispersed particles is prevalent in many applications. Previous studies of droplet breakup mainly focused on homogeneous fluids, and the heterogeneous effect of particles on the breakup progress is unclear. In this study, the breakup of particle-laden droplets in airflow is investigated experimentally. Combining synchronised high-speed images from the side view and the $45°$ view, we compare the morphology of particle-laden droplets with that of homogeneous fluids in different breakup modes. The results show that the higher effective viscosity of particle-laden droplets affects the initial deformation, and the heterogeneous effect of particles appears in the later breakup stage. To evaluate the heterogeneous effect of particles quantitatively, we eliminate the effect of the higher effective viscosity of particle-laden droplets by comparing cases corresponding to the same inviscid Weber number. The quantitative comparison reveals that the heterogeneous effect of particles accelerates the fragmentation of the liquid film and promotes localised rapid piercing. A correlation length that depends on the particle diameter and the volume fraction is proposed to characterise the length scale of the concentration fluctuation under the combined effect of the initial flattening and later stretching during the droplet breakup process. Based on this correlation length, the fragment size distributions are analysed, and the scaling results agree well with the experimental data.

Key words: [brea](#page-28-0)kup/coalescence, [drops,](#page-27-0) suspens[ions](#page-29-0)

1. Introduction

The atomisation of a suspension is a fundamental issue for many natural phenomena and practical applications, such as slurry-fuel atomisation (Gvozdyakov & Zenkov 2021; Noh *et al.* 2022), spray coating (Suhag *et al.* 2020; Koivuluoto 2022; Sun *et al.* 2022), [respiratory drug](https://doi.org/10.1017/jfm.2023.847) delivery (Dietzel 2016) and ocean spray (Veron 2015). In these processes, the fluids contain dispersed objects such as metal or coal particles, polymer or catalyst and pollutant plastics, which complicate their atomisation processes. Fluid atomisation

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usually involves several stages, of which droplet breakup in the airflow is an important process. The droplet breakup produces n[umero](#page-28-1)us small frag[ments](#page-29-1), and the fragm[ent siz](#page-29-2)e is affected by the dispersed particles, which is crucial to the subsequent heat/mass transfer and reaction. For example, the breakup of slurry fuel droplets directly affects the ignition and combustion of the fuel (Padwal, Natan & Mishra 2021). In addition, interaction between the fluid and the airflow during the droplet breakup process also ex[ists in](#page-29-1) primary break[up pro](#page-29-2)cesses. Therefore, the study of the breakup of particle-laden droplets is useful for understanding the atomisation process of suspensions.

Droplet breakup of homogeneous fluids has been studied by many researchers (Guildenbecher, López-Rivera & Sojka [2009;](#page-28-2) Theofanous 2011; [Sharm](#page-29-3)a *et al.* 2022). The breakup process is mainly controlled by the aerodynamic force of the airflow, and the viscosity and surface tension of the liquid. The aerodynamic force of the airflow drives the deformation and breakup of the droplet. As the aerodynamic force increases, droplets in the airflow exhibit different breakup morphologies (Theofanous 2011; Sharma *et al.* 2022), including bag breakup, multimode breakup, shear-[strippi](#page-28-1)ng breakup, etc. In contrast, the viscosity of the droplet inhibits the droplet deformation, resulting in a stronger aerodynam[ic forc](#page-29-1)e being re[quired](#page-29-4) for high-viscosity fluids to achieve the same breakup morphology (Radhakrishna *et al.* 2021; Xu, Wang & [Che](#page-28-3) [2023\).](#page-28-4) The magnitude [of the](#page-28-5) aerodynamic force of the airflow and the viscous force of the droplet relative to the droplet surface tension can be characterised by the Weber number (W_{e_g}) and the Ohnesorge number (*Oh*), respectively. In addition to the breakup morphologies, the droplet breakup processes can be quantified with their time characteristics (such as the initial deformation time and the total breakup time (Guildenbecher *et al.* 2009)), kinematic and dynamic characteristics (such as the velocity, acceleration and drag coefficient of the droplet (Theofanous 2011; Yang *et al.* 2016)) and shape character[istics \(](#page-28-1)such as the d[roplet](#page-29-1) width and thi[cknes](#page-29-5)[s, and](#page-29-2) the fragment size (Jackiw & Ashgriz 2021, 2022; Joshi & Anand 2022)).

Due to the multiphase interaction and the complex mo[rpholo](#page-29-6)gy exhibited during the droplet breakup, the underlyin[g mech](#page-29-7)anisms of dr[oplet b](#page-29-8)reakup are often interpreted from different perspectives in different studies. From the perspective of surface instability, different breakup morphologies em[erge u](#page-29-6)nder the comp[etition](#page-29-5) [of the](#page-29-2) middle piercing controlled by the Rayleigh–Taylor (RT) instability and the peripheral stripping controlled [by th](#page-28-6)e Kelvin–Helmholt[z \(KH](#page-28-7)) instability (Guild[enbech](#page-28-3)er *et al.* 2009; Theofanous 2011; Sharma *et al.* 2021, 2022). For the RT instability, in addition to the simplified RT instability [being](#page-28-8) used, the RT instability considering the finite thickness of the liquid was developed for the analysis of the droplet piercing (Theofanous *et al.* 2012) and the breakup of the bag film (Villermaux & Bossa 2009; Villermaux 2020). For the KH ins[tabilit](#page-29-9)y, different velo[city pr](#page-29-10)ofiles at the li[quid–a](#page-28-9)ir interface were d[iscuss](#page-28-10)ed and used in the shear-stripping breakup process (Theofanous *et al.* 2012; Sharma *et al.* 2021, 2022). Moreover, from the perspective of droplet internal flow, the two-point approach (O'Rourke & A[msden](#page-28-11) 1987; Kulkarni [& Soj](#page-27-1)ka 2014; Ja[ckiw &](#page-29-11) Ashgriz 2021) (mathematically consistent with the Taylor analogy breakup model) and the energy approach (Ibrahim, Yang & Przekwas 1993) (i.e. the droplet deformation and breakup model) are proposed to describe the droplet deformation and breakup. Different variations and improvements have been developed based on this perspective, but mostly for low Weber numbers (Wang *et al.* 2015; Stefanitsis *et al.* 2019; Rimbert *et al.* 2020; Obenauf & Sojka 2021). In addition, from the perspective [of](https://doi.org/10.1017/jfm.2023.847) [droplet](https://doi.org/10.1017/jfm.2023.847) [ex](https://doi.org/10.1017/jfm.2023.847)ternal airflow, the formation and shedding of vortices outside the droplet are found to significantly affect the droplet breakup at high Weber numbers (Liu *et al.* 2018; Dorschner *et al.* 2020; Wang *et al.* 2020). Finally, besides these perspectives, other models

of droplet deformation and breakup have been proposed based on global principles, including the virtual work principle (Sichani & Emami 2015) and the maximum e[ntropy](#page-28-12) formali[sm \(K](#page-28-13)uo & Trujillo 2022).

Different from homogeneous fl[uids,](#page-28-14) [s](#page-28-14)uspensions exhibit s[ome](#page-28-15) [u](#page-28-15)nique properties that depend primarily on the length scales of the process and t[he vol](#page-27-2)[ume f](#page-27-3)raction of the particles (Tropea, Yarin & Foss 2007). At a length scale much larger than the particle size, the suspension can be treated as a homogeneous fluid. In addition, its rheological properties vary with the volume fraction of the particles, exhibiting Newtonian behaviours at low volume fractions, shear-thickening or shear-thinning behaviours at high volume fractions and jamming near the maximum packing fraction (Guazzelli & Pouliquen 2018; Morris 2020; More & Ardekani 2021; Ness, S[eto &](#page-27-4) Mari 2022). In contrast, at the [length](#page-27-5) scale close to the pa[rticle](#page-29-12) size, the heterogeneous [effect](#page-29-13)s of the particles in the suspensio[n need](#page-29-14) to be taken into account. Furbank & Morris (2004, 2007), first reported the pinch-off dynamics of particle-laden liquids and proposed a two-stage description of [the d](#page-28-16)roplet formation from [suspe](#page-27-6)nsions, including the early stage, where the suspensions behave as an effective Newtonian fluid, and the later stage, where the detachment is accelerated due to the heterogeneous [effects](#page-28-17) of the particles. The detailed process, stage transition and underlying mechanisms of the pinch-off dynamics of particle-laden liquids were further developed by Bonnoit *et al.* (2012), Château, Guazzelli & Lhuissier (2018), Rubio-Rubio *et al.* [\(20](#page-28-18)18), Thieve[naz &](#page-27-7) Sauret (2022) and Thi[evena](#page-28-19)z, Rajesh & Sauret (2021). Similarly, for the breakup of suspension jets, the heterogeneity of the particles affects the shape and the breakup length of the jets (Roche, Kellay $\&$ Stone 2011; Château & Lhuissier 2019). Moreover, for the spreading and fragmentation of the suspension films, the presence of particles changes the capillary flows of films and leads to more complex situations. Raux *et al.* (2020) experimentally studied the fragmentation of particle-laden liquid sheets generated by a droplet impacting a cylindrical target, and found the presence of particles modifies the thickness and reduces the stability of the liquid sheet. Jeong *et al.* (2022[\), Gans](#page-29-15) *et al.* (2019) and Pa[lma &](#page-29-16) Lhuissier (2019) reported the dip coating of a substrate withdrawn from a suspension bath and classified different coating regimes based on the entrainment degree of the particles on the liquid film, which is governed by the film thickness relative to the particle diameter. As for the breakup of the particle-laden droplet considered in this study, the droplet deformation and breakup process under the action of airflow exhibits complex morphologies, including films, ligaments, nodes, etc. For the breakup of these complex morphologies, a variation in length scale will occur as the droplet deforms and breaks up and the effect of particles will be complex. Zhao *et al.* (2011*b*) and Wang *et al.* (2021) studied the effect of the yield stress and the discontinuous shear thickening exhibited by suspensions on droplet breakup and found some new breakup modes, such as hole breakup, tensile breakup and hardened deformation. These studies treated suspensions as homogeneous fluids, but the heterogeneous effect of particles on droplet breakup has not been studied.

In this study, the breakup of particle-laden droplets in airflow is investigated. We first discuss the changes in the deformation morphology and breakup mode of particle-laden droplets compared with homogeneous fluids. To quantitatively analyse the effect of particles on the breakup process, we distinguish the effects of the particles via the effective viscosity and via the heterogeneity by comparing at the same breakup mode, i.e. the same [inviscid Weber](https://doi.org/10.1017/jfm.2023.847) number. Based on this, we further investigate the heterogeneous effects of particles on the typical morphological features and fragment sizes in different breakup modes.

Figure 1. Schematic diagram of the experimental set-up.

2. Experimental set-up and materials

The experimental set-up is shown in figure 1. A continuous airflow was used to study the droplet breakup, which has been adopted in many studies (Jackiw & Ashgriz 2021; Radhakrishna *et al.* 2021; Wang *et al.* 2021). The airflow originated from compressed-air storage and was regulated by a mass flow controller (Alicat MCRQ, maximum flow rate 3000 standard litres per minute, estimated uncertainty $\pm 0.8\%$ of reading and $\pm 0.2\%$ of full scale). The airflow then passed through a honeycomb (a hexagon with a side length of 1 mm) and two-layer meshes (100 mesh) to reduce turbulence, and was finally ejected from a rectangular nozzle (the outlet cross-section is 20 mm in width and 30 mm in height). The range of the airflow velocity was $u_g = 8.9$ –70 m s⁻¹. Droplets entered the airflow from above by gravity. To minimise the effect of the jet shear layer, the [fall velo](#page-3-0)city of the droplet was adjusted to ensure that deformation and breakup of the droplet occurred in the uniform velocity region of the jet for all airflow velocities considered in this study. The experiments were performed at room temperature (22 ◦C) and atmospheric pressure with air density $\rho_g = 1.2 \text{ kg m}^{-3}$.

High-speed cameras with different arrangements were used to film the droplet deformation and breakup for different purposes. To capture the complete morphology of the droplet breakup, two synchronised cameras (Photron Fastcam SA1.1) were used to take images from the side view and $45°$ view, as illustrated in figure 1. From the side view, a camera with a spatial resolution of 90 μ m pixel⁻¹ acquired the images of the side of the droplet. From the 45◦ view, a camera with a spatial resolution of $66 \mu m$ pixel⁻¹ obtained the information on the windward side of the droplet. Both cameras were synchronised at a frame rate of 10 000 frames per second (f.p.s.). In addition, to obtain the fragment size after the droplet breakup, two synchronised cameras (Phantom [v1612](https://doi.org/10.1017/jfm.2023.847) [for](https://doi.org/10.1017/jfm.2023.847) [th](https://doi.org/10.1017/jfm.2023.847)e upstream, and Photron Fastcam SA1.1 for the downstream) were arranged together on the side. By combining the shooting areas of the two cameras, we were able to capture all fragments with a sufficient spatial resolution of 60 μ m pixel⁻¹. Both cameras were set at the frame rate of 5000 f.p.s. For all camera arrangements, macro lenses (Nikon

Figure 2. (*a*) Particle size distribution of $d_p = 90 \,\mu$ m. The inset is an image of particles under the microscope. (*b*) Relative effective shear viscosity of the suspensions (η/η_f) for different particle volume fractions (ϕ). The error bars of t[he](https://doi.org/10.1017/jfm.2023.847) effective viscosity were calculated from the data in the [shear](https://doi.org/10.1017/jfm.2023.847) [rate](https://doi.org/10.1017/jfm.2023.847) [range](https://doi.org/10.1017/jfm.2023.847) [of](https://doi.org/10.1017/jfm.2023.847) $100-1000 s^{-1}$ $100-1000 s^{-1}$ $100-1000 s^{-1}$.

AF 100 mm f/2.8D) with a small aperture (F16 or F22) were used to obtain a large depth of field with low optical distortion, and two high-power (280 W) light-emitting diode lights diffused by ground glasses were used as the backgroun[d](#page-4-0) [light](#page-4-0) [so](#page-4-0)urces to ensure sufficient brightness for the high-speed imaging. The high-speed images were analysed via a customised image-processing program in Matlab, and the details are given in the supplementary material and movies available at https://doi.org/10.1017/jfm.2023. 847. The fluids used for droplet breakup were suspensions of solid particles in interstitial liquids. The solid particles were spherical polystyrene particles with a density of $\rho_p =$ 1050 kg m⁻³. The particle sizes were measured from the images taken under an inverted microscope (Nikon Ti-U with a spatial resolution of 2 μ m pixel⁻¹), and figure 2(*a*) shows a typical particle size distrib[ution](#page-27-5) with $d_p = 90 \mu$ [m.](#page-28-18) [I](#page-28-18)n our experiments, three different particle sizes ($d_p = 40$, 90 and 180 μ m) were used and their relative standard deviation was approximately 12 %. The interstitial liquid was silicone oil (phenyl methyl silicone oil from Macklin), which had a viscosity $\eta_f = 35$ mPa s, a density $\rho_f = 1050$ kg m⁻³ and a surface tension $\sigma = 27$ mN m⁻¹. The suspensions were repeatedly stirred and rested to remove air bubb[les. The pa](#page-26-0)[rticles are](#page-4-0) totally wetted by the silicone oil and did not modify the liquid–gas interface, so the change of surface tension caused by the particle is negligible (Château *et al.* 2018; Jeong *et al.* 2022). Due to the matching in density between t[he par](#page-28-12)ticles and the interstitial liquid, the effect of particle sedimentation was limited during the experiment.

The effective shear viscosity (η) of each suspension was measured by a rotating rheometer (Anton Paar MCR 501). The particle volume fraction (ϕ) of the suspensions was varied from 0 % to 44 % in the rheological measurements. The details of the measurement can be found in the Appendix. Figure $2(b)$ presents the relative effective shear viscosity (η/η_f) of the suspensions for different particle volume fractions, indicating that the relative effective shear viscosity can be described well by the Maron–Pierce model (Guazzelli & [Pouliquen](https://doi.org/10.1017/jfm.2023.847) 2018)

$$
\frac{\eta}{\eta_f} = \left(1 - \frac{\phi}{\phi_m}\right)^{-2},\tag{2.1}
$$

Figure 3. Image sequences of the bag breakup mode at $W_{eg} = 20.3$, $Oh = 0.32$: (*a*) suspension with $d_p =$ 90 μm and $\phi = 0.25$; (*b*) silicone oil with $\eta = 100$ mPa s. The corresponding video can be found in the supplementary material and movie 1.

where ϕ_m is the maximum packing fraction and $\phi_m = 0.64$ in our experiment. To separate the influence of the effective viscosity from the heterogeneous effects of the particles, we chose silicone oils of different viscosities (phenylmethyl silicone oil of $\eta = 35$, 100, 200 mPa s from Macklin) as a control group, which had a surface tension $\sigma = 27 \pm 1$ mN m⁻¹. The particle volume fraction (ϕ) in the droplet breakup experiments was varied from 0% to 35% because the droplets tend to be non-spherical at high particle volume fractions. The droplet diameter used in this study was $d_0 = 3.35 \pm 0.05$ mm. Correspondingly, the range of the gas Weber number ($W_e = \rho_g u_g^2 d_0 / \sigma$) in this study was 11.8–718, and the range of the Ohnesorge number $(Oh = \frac{\eta}{\sqrt{\rho_f d_0 \sigma}})$ $(Oh = \frac{\eta}{\sqrt{\rho_f d_0 \sigma}})$ [was](#page-5-0) [0.1](#page-5-0)1–0.65.

3. Results and dis[cussion](#page-5-0)

3.1. *Droplet morphology during the deformation and breakup*

3.1.1. *Bag and bag-stamen breakup*

The bag breakup mode is one of the main modes of droplet breakup. Figure 3(*a*) shows the bag breakup process of a particle-laden droplet. For comparison, the bag breakup process of a silicone oil droplet is shown in figure 3(*b*), which has the same effective viscosity as the particle-laden droplet. The breakup process can be divided into an initial deformation [stage](https://doi.org/10.1017/jfm.2023.847) $(0-13 \text{ ms in figure 3})$, and a bag development and breakup stage (after 13 ms in figure 3). During the initial deformation stage, the particle-laden droplet deforms almost the same as the silicone oil droplet. This indicates that during the initial deformation stage, the particle-laden droplet can be regarded as a homogeneous fluid; and compared

Figure 4. Synchronised images of the ba[g-stamen b](#page-5-0)reakup at $We_g = 28.7$, $Oh = 0.32$: (*a*) suspension with $d_p = 90 \mu$ m and $\phi = 0.25$, *(b)* silicone oil with $\eta = 100$ mPa s. The corresponding video can be found in the su[pplementar](#page-5-0)y material and movie 2.

with the interstitial liquid, the [particle](#page-28-17)s lead to a higher effective viscosity of the droplet. However, during the bag development and breakup stage, the heterogeneous effect of particles appears. The particles agglomerate on the liquid film and accelerate the breakup of the liquid film (23.2–30 [ms i](#page-12-0)n figure 3*a*). In contrast, the liquid film of the silicone oil droplet is significantly stretched and deformed, and the breakup starts later (35 ms in figure 3*b*). The development and breakup of the suspension bag film are similar to the fragmentation of particle-laden liquid sheets generated by a droplet impacting a cylindrical target described by Raux *et al.* (20[20\). How](#page-6-0)ever, compared with the particle-laden liquid sheets described by Raux *et al.* (2020), the particle distribution on the suspension bag film is more un[even due](#page-6-0) [to](#page-6-0) [the](#page-6-0) [ac](#page-6-0)celerated stretching of the bag film caused by the continuous aerodynamic force. Further quantitative comparisons for the stretch and breakup of the [liquid fil](#page-6-0)m will be given in § 3.3.

As *Weg* increases, the bag-stamen breakup mode occurs. Different fro[m](#page-6-0) [the](#page-6-0) [bag](#page-6-0) breakup, the droplet morphology in bag-stamen breakup mode consists of the peripheral bag and the middle stamen. The comparison of the breakup process of the particle-laden droplet and the silicone oil droplet is shown in figure 4. The initial deformation of the particle-laden droplet $(0-10 \text{ ms in figure } 4a)$ is similar to that of the silicone oil droplet under the same conditions (figure 4*b*). But after the initial deformation, compared with that of the silicone oil droplet, the pe[riphera](#page-29-3)l bag of the particle-laden droplet breaks up quickly (20 ms in figure 4*a*), and the elongation of the peri[pheral rin](#page-7-0)g is weaker (25 ms in figure 4*a*).

3.1.2. *[Multim](#page-7-0)ode b[reakup](#page-7-0)*

The multimode breakup mode is another main mode of droplet breakup. Compared with bag breakup, the multimode breakup occurs at a higher Weber number and has new features. Xu *et al.* (2023) classified the multimode breakup into low-order and high-order modes according to the piercing times in the middle of the droplet. A typical process of the low-order multimode breakup is shown in figure 5, while the morphology of the high-order [multimode](https://doi.org/10.1017/jfm.2023.847) [brea](https://doi.org/10.1017/jfm.2023.847)kup is shown in figure 6.

For comparison, figure 5(*a*) shows the breakup process of a particle-laden droplet and figure 5(*b*) shows the breakup process of a silicone oil droplet under the same conditions. The process of the low-order multimode breakup can be divided into the

Figure 5. Synchronised images of the low-order multimode breakup at $W_{e} = 71.7$, $Oh = 0.32$: (*a*) suspension with $d_p = 90 \mu$ m and $\phi = 0.25$; (*b*) silicone oil with $\eta = 100$ mPa s. The corresponding video can be found in the supplementary material and movie 3.

initial deformation stage $(0-4 \text{ ms in figure 5})$, the peripheral retraction stage $(4-6.5 \text{ ms})$ in figure 5(*a*) and 4–8 ms in figure 5*b*) and the subsequent piercing and breakup stage. During the initial deformation stage, the deformation is similar for both fluids, and the particle-laden droplet can be regarded as a homogeneous fluid. During the [peripher](#page-7-0)al retraction stage, the silicone oil dropl[et tends to](#page-7-0) form a peripheral sheet, and the peripheral sheet deflects to the leeward of the droplet before the sheet destabilisation develops, i.e. the sheet retraction (8 ms in figure 5*b*). In contrast, for the particle-laden droplet under the same conditions, the azimuthal destabilisation of the droplet periphery develops and fo[rms fing](#page-8-0)erings at the periphery (6.5 ms in figure 5*a*). Hence, the formation of fingers in[dicates th](#page-8-0)e emergence of particle heterogeneity as the peripheral sheet be[comes thi](#page-8-0)nner. In addition, for the piercing in the middle of the droplet, the particle-laden droplet tends to form lo[cal piercin](#page-8-0)g with a faster speed and less lateral expansion (9.8 ms in figure 5*a*) than the silicone oil droplet (11 ms in figure 5*b*).

Different from the low-order multimode break[up](#page-9-0) [where](#page-9-0) the droplet middle is pierced once, the droplet m[iddle](#page-14-0) is pierced multiple times under the high-order multimode breakup. The comparison of the particle-laden droplet and the silicone oil droplet is shown in figure 6. The time interval between two piercings of the particle-laden droplet (7–9.6 ms in figure $6a$) is shorter than that of the silicone oil droplet $(7-11 \text{ ms in figure } 6b)$. In addition to the major piercings, the particle-laden droplet forms some minor piercings (9.6 ms in figure 6*a*). Similar minor piercings [also](#page-29-1) occur in the cas[e of th](#page-29-6)[e low-ord](#page-10-0)er multimode breakup with large particles (9.2 ms in figure 7). These piercings i[n the midd](#page-10-0)le have important effects on the breakup morphology and fragment size, which will be discussed further in § 3.4.

3.1.3. *Shear-stripping breakup*

The shear-stripping breakup occurs at a much higher Weber number and is the terminal [regime](https://doi.org/10.1017/jfm.2023.847) [cons](https://doi.org/10.1017/jfm.2023.847)idered by some studies (Theofanous 2011; Theofanous *et al.* 2012). Figure 8(*a*) shows the shear-stripping breakup process of a particle-laden droplet, and figure 8(*b*) shows the breakup process of a silicone oil droplet under the same conditions for comparison. Being different from the bag and multimode breakup modes, here the

Figure 6. Synchronised images of the high-order multimode breakup at $W_{e} = 103.7$, $Oh = 0.32$: (*a*) suspension with $d_p = 90 \mu$ m and $\phi = 0.25$, (*b*) silicone oil with $\eta = 100$ mPa s. The corresponding video can be found in the supplementary material and movie 4.

heterogeneous effect of particles appears at the beginning of the breakup process. In the early stage of the breakup of the particle-laden [droplet \(0](#page-10-0)–3 ms in figure 8*a*), the interstitial liquid is rapidly pushed downst[ream by](#page-10-0) the airflow, thus particles protrude from the windward side in the middle of the droplet forming a rough surface, while some particles are stripped directly from the periphery of the droplet. In co[ntrast, for](#page-10-0) the silicone oil droplet, the windward side in the middle is relatively smoother due to the tension generated by the outward stripping (2 ms in figure 8*b*), while some ripples on the edge of the windward surface of the silicone oil droplet are formed. After the flattening of the droplet middle, massive piercings occur in the middle of the droplet. The particle-laden [droplet](https://doi.org/10.1017/jfm.2023.847) [forms](https://doi.org/10.1017/jfm.2023.847) [s](https://doi.org/10.1017/jfm.2023.847)ome large particle clusters (4.5 ms in figure 8*a*), while the silicone oil droplet forms some liquid clumps (4.5 ms in figure 8*b*). Finally, the particle clusters or the liquid clumps may be further pierced by the airflow. The particles and the interstitial liquid in the particle-laden droplet are almost completely separated (6 ms in figure 8*a*), while the

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Figure 7. Synchronised images of the low-order multimode breakup at $W_{eg} = 69.9$, $Oh = 0.32$. The fluid used is a suspension with $d_p = 180 \,\mu\text{m}$ and $\phi = 0.25$. The corresponding video can be found in the supplementary material and movie 5.

silicone oil droplet forms mists of tiny droplets (6 ms in figure 8*b*). Overall, compared with the continuous breakup of the silicone oil droplet, the breakup of the particle-lad[en drople](#page-5-0)t is more discrete, like t[he shatter](#page-6-0)ing of a solid.

3.2. *Regime map of droplet breakup*

The morphology of droplet brea[kup](#page-6-0) [is](#page-6-0) [a](#page-6-0) [c](#page-6-0)omplex process with different breakup patterns in different conditions. [Here, acc](#page-7-0)ording to the criteria for viscous droplets (Xu *et al.* 2023) [\(i.e. base](#page-7-0)d on the piercing time in the droplet middle and the retraction in the droplet periphery), we classify the droplet breakup modes into no breakup, bag breakup (figure 3), bag-stamen breakup (figure 4), low-order multimode breakup (figure 5), high-order multimode breakup (figure 6) and shear-s[tripping b](#page-7-0)reakup (figure 8). The transition from bag breakup to bag-stame[n breakup](#page-8-0) corresponds to the formation of a stamen at the middle of the droplet (24 ms in figure 4*b*). The transition from bag-stamen breakup to [low-orde](#page-10-0)r multimode breakup corresponds to the formation of a sheet at the periphery of the droplet (8 ms in figure 5*b*) and the piercing at the middle of the d[roplet \(](#page-10-1)11 ms in figure 5*b*). The transition from [low-ord](#page-11-0)er multimode breakup to high-order multimode breakup corresponds to the change of the piercing times in the droplet middle. After the sheet formation at the periphery, the middle part of the droplet is pierced once in [the low](#page-27-8)-order multimode breakup mode (figure 5) and multiple times in the high-order multimode breakup mode (figure 6). Finally, the transition from high-order multimode breakup to sh[ear-strip](#page-29-17)ping breakup corresponds to the o[ccurre](#page-29-3)nce of peripheral stripping (figure 8), i.e. the appearance of KH instability waves at the periphery of the droplet. The main morphology features of different breakup modes are summarised in table 1.

The regime map is shown in figure 9. The transition of different modes is determined by the comparison between the droplet width after the initial flattening (d_w) and the [most-amplifi](https://doi.org/10.1017/jfm.2023.847)ed wavelength of the RT instability (λ_{RT}) ; λ_{RT} was obtained by Aliseda *et al.* (2008), and d_w can be obtained by balancing the kinetic energy of the airflow and the surface energy of the droplet. By linking these processes, a theoretical model was proposed by Zhao *et al.* (2011*a*) and further developed by Xu *et al.* (2023) for the transitions between

Figure 8. Synchronised images of the shear-stripping breakup at $We_g = 459.5$, $Oh = 0.32$: (*a*) suspension with $d_p = 180 \mu$ m and $\phi = 0.25$; (*b*) silicone oil with $\eta = 100$ mPa s. The insets show enlarged images of the windward surface of the droplets. The corresponding video can be found in the supplementary material and movie 6.

No breakup Deformation and oscillation Bag breakup Bag with a toroidal rim Bag stamen Stamen Peripheral bag

Low-order multimode Single piercing Peripheral bag High-order multimode Multiple piercings Peripheral sheet [She](https://doi.org/10.1017/jfm.2023.847)ar stripping Massive piercings Stripping

Breakup mode Middle of the droplet Periphery of the droplet

Peripheral bag or sheet

Table 1. Main morphological features of different breakup modes.

Figure 9. Regime map of droplet breakup. The hollow symbols correspond to the breakup of silicone oil droplets and the solid symbols correspond to the breakup of particle-laden droplets. For particle-laden droplets, *Oh* is calculated based on the effective viscosity (2.1) at the corresponding particle volume fraction. The solid lines are based on the theoretical model (3.1) for the transitions of different regimes. The stars along the dashed lines correspond to the cases for the comparison under the same $We_{c,0}$.

different breakup modes

$$
\left(\frac{W e_{c,0}}{W e_c}\right)^{1/2} + C W e_{c,0}^{2/3} \left(\frac{Oh^2}{W e_c}\right)^{1/3} = 1, \tag{3.1}
$$

where W_{e} is t[he critica](#page-11-0)l Weber number for the transition between different regimes. Here, *C* is equal to 0.24 and is universal f[or](#page-11-1) *Oh* < 2.15, which can be considered as a weight coefficient of the viscous effect relative to the surface tension in th[e drople](#page-11-0)t breakup process (Xu *et al.* 2023). When *Oh* appr[oaches ze](#page-11-0)ro, *Wec* is equal to the critical Weber number in the inviscid case ($We_{c,0}$). Under the same $We_{c,0}$, the degree of initial deformation of the droplet is similar, and the inhibition of the initial deformation by the higher viscosity of the droplets is counteracted by a higher Weber number (Cohen 1994). Here, we use this model to obtain the transitions of different breakup modes, as shown by the solid lines in figure 9. The critical Weber number (*Wec*) of both the suspension droplets and the silicone droplets is obtained through (3.1) .

Comparing the breakup modes of silicone oil droplets (the hollow symbols in figure 9) [and](https://doi.org/10.1017/jfm.2023.847) [particle](https://doi.org/10.1017/jfm.2023.847)-laden droplets (the solid symbols in figure 9), we can see that the effect of particles on the transition of breakup mode is manifested by a higher viscosity compared with the interstitial fluid. This is because the transition of the breakup mode is controlled by the interfacial instability underlying the initial deformation. For the initial flattening

deformation, the suspension can be regarded as a homogeneous fluid, as described in § 3.1. For the interfacial instability, the instability wavelength $(\lambda_{RT} \sim d_0)$ is much larger than the particle size, so the heterogeneous effect of particles on interfacial instability is minor. Therefore, the breakup mode can [be](#page-11-1) classified ba[sed](#page-11-1) on the effective viscosity of the particle-laden droplet, and the heterogeneous effect of particles appears only in the later breakup stage.

Since the addition of particles into the liquid can increase the [effe](#page-11-1)ctive viscosity of the fluid, the contribution of the higher viscosity sh[ould](#page-11-1) be eliminated when we analyse the heterogeneous effect of the particles. Otherwise, the effects of the higher viscosity and the effects of the particle heterogeneity are mixed, hindering us from understanding the underlying mechanisms. In this regard, suspensions with different concentrations should be compared under the same $We_{c,0}$ in (3.1). Equation (3.1) is t[o obtain](#page-11-0) the transitions between different breakup modes, but the transitions between different breakup modes are essentially the change of breakup morphology. Therefore, (3.1) with different $We_{c,0}$ can reflect the change of breakup morphology, that is, the droplet has a similar breakup morphology under the condition obtained by (3.1) with a certain $We_{c,0}$. In this way, we can compare the conditions with different particle volume fractions and isolate the effects of particle heterogeneity. In the subsequent sections, we choose two typical breakup morpholo[gies for c](#page-13-0)omparison, i.e. the bag breakup ($W_{c,0} = 12.2$) and the low-order multimode breakup ($We_{c,0} = 36$), as shown by the dashed lines in figure 9.

3.3. *Stretching and fragmentation of liquid film*

[The](#page-13-0) [stretc](#page-13-0)hing and fragmentation of the liquid film is the main morphological feature of droplet breakup, especially for the bag and bag-stamen breakup modes. Here, we compare the fragmentation process o[f the b](#page-28-17)ag film under the same $We_{c,0} = 12.2$, as shown in figure 10. It can be seen that, as the particle volume fraction increases, the bag fragmentation occurs earlier in the stretching process. For example, the morphology corresponding to the fragmentation moment (t_0) in figure $10(c)$ is similar to the morphology corresponding to 3.5 ms before the fragmentation moment $(t_0 - 3.5)$ in figure 10(*a*). [This](#page-29-8) is because particles occupy a part of the droplet, so the liquid content in the droplet is reduced compared with the case without particles. In addition, the liquid enclosed in particle clusters is not available for the stretching of the liquid film, which is similar to the observation of Raux *et al.* (2020) in the fragmentation of [partic](#page-29-8)le-laden liquid sheets. Therefore, the liquid used for the bag stretching is less, which is not conducive to the stretching of the bag film. Moreover, when the thickness of the bag film is smaller than the diameter of the particle clusters, capillary stress will form around the particle clusters. The local stress induced by the partic[les ma](#page-29-8)kes the liquid film susceptible to fragmentation (Villermaux 2020). In addition, the fragmentation processes of the bag film are different. For the silicone oil droplet, the fragmentation of the liquid film starts by forming holes at one or several [locations](#page-14-1) where the film is the thinnest, and then the holes expand. In this process, the kine[matic thinn](#page-14-1)ing leads to the film rupture (Villermaux 2020). However, for the particle-laden droplet, many holes are initiated on the liquid film due to the presence of the particles, forming a web-like fragmentation. This fragmentation process of the liquid film is the spontaneous hole formation caused by internal defects, i.e. the local stress induced by the particles (Villermaux 2020).

[Through](https://doi.org/10.1017/jfm.2023.847) [dig](https://doi.org/10.1017/jfm.2023.847)ital image processing, we can obtain the droplet thickness (l_t) along the direction of airflow, i.e. the width of the vertical bounding box of the droplet (as shown in the inset of figure 11*a*). The evolution of the droplet thickness in the bag development stage is shown in figure $11(a)$. The stretching speed of the liquid film of a particle-laden

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Figure 10. Image sequences of bag fragmentation processes. The cases correspond to the stars along the dashed line with $W_{e_0} = 12.2$ in figure 9: (*a*) silicone oil with $\eta = 35$ mPa s, $W_{e_8} = 15.7$, $Oh = 0.11$; (*b*) suspension with $\phi = 0.10$, $W_{eg} = 16.6$, $Oh = 0.17$; (*c*) suspension with $\phi = 0.25$, $W_{eg} = 20.3$, $Oh = 0.32$; (*d*) suspension with $\phi = 0.35$, $\overline{W}e_g = 24.3$, $Oh = 0.52$. The size of the particles within the suspensions is $d_p = 90 \mu$ m. Here, t_0 correspon[ds to the mom](#page-14-1)ent when the bag fragmentation begins. The images with similar morphology are conne[cted by dash](#page-14-1)ed lines. For comparison with other panels of the figures, the time sequence in (*a*) is from the bottom le[ft to the top](#page-14-1) left and then to the top right.

droplet (so[lid](#page-14-1) [symbo](#page-14-1)ls in figure 11*a*) is closer to that of the interstitia[l](#page-14-1) [liquid](#page-14-1) [\(h](#page-14-1)ollow square symbols in figure $11a$) than that of liquid with the same effective viscosity (hollow triangular [symbols](#page-13-0) in figure 11*a*). This is because the interaction between the particles is insigni[ficant](#page-27-6) during the stretching process of the liquid film of the bag, while the viscosity of the interstitial fluid is dominant. Moreover, we calculate the total breakup time, which is the time when all fragmentation has ceased (Pilch $\&$ Erdman 1987), as shown in figure $11(b)$. The total breakup time is smaller for particle-laden droplets (solid [symbols in](https://doi.org/10.1017/jfm.2023.847) figure 11*b*) than for silicone oil droplets (hollow symbols in figure 11*b*) and also decreases slightly with increasing particle volume fraction. This reduction in the total breakup time is mainly because the particles accelerate the fragmentation of the bag (as shown in figure $10a$) and the ring (similar to the jet breakup described by Château & Lhuissier 2019).

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Figure 11. (*a*) Evolution of the droplet thickness along the direction of airflow in the bag development stage. The time here is defined from the end of initial deformation (when the droplet is flattened to a disk shape with the largest aspect ratio) to the beginning of bag fragmentation (when holes appear in the bag). (*b*) Total breakup time under different particle volume fractions. The cases correspond to the stars along the dashed line with $W_{\ell,c,0} = 12.2$ in figure 9, and the particle size in the suspensions is $d_p = 90 \,\mu\text{m}$.

Figur[e 12](#page-9-1). (*a*) Evolution of local piercing in the bag-stamen breakup. The whole breakup process is shown in figure 4(*a*). (*b*) First piercing in the high-order multimode breakup. The whole breakup process is shown in figure $6(a)$.

3.4. *Piercing by RT i[nstability](#page-14-2)*

The piercing by the RT instability wave is another main morphological feature of the droplet. The piercing times in the middle of the droplet that determines the breakup mode are mainly affected by the effective viscosity of the particle-laden droplet, as described in § 3.2. However, in the later stages of piercing development, the heterogeneity of the particle-laden droplets becomes important and affects the droplet morphology.

[To](https://doi.org/10.1017/jfm.2023.847) [clearly](https://doi.org/10.1017/jfm.2023.847) [dis](https://doi.org/10.1017/jfm.2023.847)play the piercing development process, we select the local piercing process of the bag-stamen breakup mode, as shown in figure $12(a)$. After the formation of circumferential depression on the windward of the droplet, the local piercing with a faster speed occurs in the interstitial liquid as the droplet becomes thinner. The bag produced

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by the local piercing contains fewer particles, whereas particles aggregate at the edge of the bag. Inspired by the comparison with the study of Raux *et al.* (2020), we suggest that the particle aggregation at the junction of the bag film and the ring is caused by the accelerated stretching of the bag film. Let us take t[he case of](#page-14-2) figure $12(a)$ as an example. At the beginning of bag stretching (i.e. the end of droplet flattening), the particle-based Weber number is $W e_p = \rho_p d_p u_f^2 / \sigma \approx 0.77$, where u_f i[s the velo](#page-14-2)city of the bag film and can be estimated as $u_f \sim u_g \sqrt{\rho_g/\rho_f} \approx 0.47 \text{ m s}^{-1}$ at the beginning of bag stretching. Meanwhile, the Stokes number of the particle is $St = \rho_p \gamma d_p^2 / \eta_f \approx 0.136$, where $\dot{\gamma}$ is the shear rate of the droplet and can be estimated $\dot{\gamma} \sim u_f / \dot{d}_0$. So at the early stage of the bag development, the particle motion is in the viscous regime ($St \ll 1$), and the particles can follow the stretching of bag film (as shown at t_0 in figure 12*a*). However, as the bag film stretches and becomes thinner, the velocity and the shear rate of the bag film increase rapidly ($u_f \approx 4.3$ m s⁻¹ and $\dot{\gamma} \sim u_f/d_p$ at $t_0 + 1$ ms in figure 12*a*) due to the continuous aerodynamic force. Here, *Wep* and *St* increase rapidly and can be expected to exceed 1. The motion of the particle s[hifts](#page-14-2) [to](#page-14-2) [an](#page-14-2) inertia-dominated regime gradually, and the ability of the particles to follow the stretching of the bag film weakens. Therefore, the particles are retained at the ju[nction of](#page-14-2) the bag film and the ring to form particle aggregation. Moreover, as the particles are retained at the junction, a viscosity difference develops between the tip and the root of the bag, which further increases the velocity difference. This form[s a mutua](#page-5-0)l[ly](#page-7-0) reinforcing process of particle aggregation at the root of the bag film and acceleration at the tip of the bag film. Finally, a local piercing with a faster speed forms (as shown $t_0 + 2$ ms in figure 12*a*). The piercing is faster due to the lower viscosity of the interstitial liquid compared with [the effect](#page-7-0)ive vis[cosi](#page-14-2)ty of the suspension. Meanwhile, the piercing is localised due to particle aggregation at the bag edge (highlighted by the arrows at $t_0 + 2$ ms in figure 12*a*) inhibiting the lateral expansion of the bag. In addition, another piece of evidence that can reflect the effect of the stretching acceleration of t[he bag](#page-29-1) film is that w[ith the](#page-29-2) increase of the stretching acceleration of the bag film (i.e. increasing We_g , from figures 3*a*–5*a*), the number of particles on the bag film decreases. This indicates that the ability of the particles to follow the stretching of the bag film weakens with the increase of the stretching acceleration of the bag film. As *Weg* increases, local piercings become more abundant, as shown in figures 5(*a*) and 12(*b*).

The development of local piercing for particl[e-laden](#page-8-0) [drop](#page-9-0)lets will prod[uce](#page-8-0) [som](#page-8-0)e new features. For the homogeneous liquid, the RT inst[ability w](#page-8-0)ave with the largest growth rate (i.e. the most-amplified w[ave\) dom](#page-8-0)inates the droplet breakup (Theofanous 2011; Sharma *et al.* 2022), and the lateral expansion of the most-amplified wave can suppress the development of instability waves with other wavelengths. However, for the particle-laden droplet, the most-amplified wave tends to further produce local piercing, while its lateral extension weakens. Therefore, the instability waves with other wavelengths may develop and form some minor piercings, such as 13.7 ms in figure 7 and 9.6 ms in figure $6(a)$. In addition, in the high-order multimode breakup (figure 6), the time interval between two piercings of the particle-laden droplet (7–9.6 ms in figure 6*a*) is shorter than that of the silicone oil droplet (7–11 ms [in](#page-29-8) figure 6*b*), which also indicates that the most-amplified wave of the particle-laden droplet has a weaker suppression effect on other instability waves. Overall, the piercing phenomenon of the particle-laden droplet is more abundant and random.

3.5. *Analysis of fragment size*

Understanding and controlling the fragment size of droplet breakup is important for numerous fields (Villermaux 2020). In this section, we will discuss the effect of particles

on the fragment size after droplet breakup. The fr[agmen](#page-28-4)t size is measured through post-processing of the images obtained by two synchronised cameras, and the details of the image-processing procedure are given in the supplementary material and [movies](#page-29-13).

The fragment originates from the rupture of elementary ligaments, which is affected by m[any fa](#page-27-9)ctors. For the [breaku](#page-29-18)p of silicone o[il dro](#page-27-5)plets, the rupture of [eleme](#page-29-13)ntary ligaments is controlled by the RT or Rayleigh–Plateau instabilities, so the instability wavelength determines the breakup length of the ligament and further the fragment size, which has been studied in detail by Jackiw & Ashgriz (2022) . However, for particle-laden droplets, there is a correlation length (ξ) that controls the stability of the suspension through a concentration fluctuation. The correlation length can be seen as a characteristic length scale over which a fluctuation will be damped (Thievenaz & Sauret 2022). The scaling relation of the correlation length is different for different flow conditions (Bonnoit *et al.* 2010; Rognon *et al.* 2014; Château *et al.* 2018; Thievenaz & Sauret 2022[\). For](#page-28-20) the breaku[p of p](#page-28-15)article-laden droplets, before the final rupture of ligaments, the droplet undergoes initial flattening and then stretching. The correlation length is determined by the combined effects of the flattening and stretching processes. In the initial flattening process, the droplet is sheared strongly. The confined space and rapid shearing increase the particle–particle interaction, producing a condition similar to the crowded condition even at a relatively lower particle volume fraction. In the crowded condition, there exists a typical correlation length (ξ_1) of th[e veloc](#page-27-10)ity field (or the non-affine velocity field in the case of a sheared system), over which particles move cooperatively (Olsson & Teitel 2007; Ness *et al.* 2022). The range of the correlated motion depends on the collective degrees of freedom in the system, which can be characteri[sed by](#page-28-15) $\phi_m - \phi$. The typical correlation length sc[ale can](#page-28-20) be found as

$$
\xi_1 \sim d_p(\phi_m - \phi)^{-n},\tag{3.2}
$$

where ϕ_m is the maximum packing fraction and $\phi_m = 0.64$, *n* is a constant which depends on the properties of the particles and their interactions (DeGiuli *et al.* 2015). To determine *n*, we need to know the flow regime of the particles, which depends on the Stokes number (*St*). The range of *St* in the initial flattening process is approximately 0.005–0.22, so a visco[us reg](#page-29-13)ime ($St \ll 1$) is expected (Ness *et al.* 2022). In the viscous regime, Olsson & Teitel (2007) reported $n = 0.6$ based on their numerical results. DeGiuli *et al.* (2015) predicted $n = 0.43$ based on a scaling analysis and argued that the effect of inter-particle friction is weak in the viscous regime. The exact value of *n* is obscure, and here, we choose an intermediate value $n = 0.5$.

After the initial flattening, the droplet is stretched to produce fragments. The basic thinning dynamics of the stretching process is similar to that of the pinch-off process of a suspension droplet, that is, the liquid between two portions of suspension is pulled apart. For this basic thinning [dynam](#page-29-13)ics process, Thievenaz & Sauret (2022) proposed a dislocation mechanism for the thinning dynamics to reveal the heterogeneity effect of particles. By balancing the power associated with the capillary forces and the viscous dissipation, they obtained a scaling law for the thinning dynamics. The applicable minimum volume fraction of this scaling law ranges from less than 2% for $20 \mu m$ particles to 20 % for 500 μ m particles, which is generally satisfied in our experiments. Here, considering that the cooperative motion of particles in the flattening process further affects the thinning dynamics in the stretching process, the particle diameter in the original [model](https://doi.org/10.1017/jfm.2023.847) [of](https://doi.org/10.1017/jfm.2023.847) [Thiev](https://doi.org/10.1017/jfm.2023.847)enaz & Sauret (2022) is replaced with ξ_1 , that is

$$
\frac{\xi}{\xi_1} \sim \left(\frac{l_c}{\xi_1}\right)^{2/3} \left(1 - \frac{\phi}{\phi_m}\right)^{-2/3},\tag{3.3}
$$

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where ξ_1 is the size of the particle cluster moving cooperatively in the initial flattening and can be obtained from (3.2). In this manner, we can obtain a correlation length for the breakup of particle-laden droplets by coupling the effects of the initial flattening and later stretching. The correlation length represents the scale of concentration fluctuations caused by the particle heterogeneity, and controls the stability of the suspension after the flattening and stretching process. The capillary length *lc* is

$$
l_c = \sqrt{\frac{\sigma}{\rho_f a}},\tag{3.4}
$$

where *a* is the st[retchin](#page-29-19)g acceleration of the droplet caused by the airflow and can be estimated as

$$
a \sim \frac{\frac{1}{2} C_d \rho_g u_g^2 \left(\frac{\pi}{4} d_w^2\right)}{\frac{\pi}{6} d_0^3 \rho_d} = \frac{3C_d}{4} \frac{\rho_g u_g^2 d_w^2}{\rho_d d_0^3},\tag{3.5}
$$

where C_d is the drag coefficient and can be considered as a constant $C_d = 1.2$ for simplicit[y \(W](#page-17-0)hite 2003). Here, d_w is the width of the droplet after the initial flattening and can be estimated as $d_w \sim d_0 We_{c,0}^{1/2}$ (Xu *et al.* 2023). By substituting (3.2), (3.4) and (3.5) into (3.3), we can obtain a dimensionless correlation length for the breakup of particle-laden droplets

$$
\frac{\xi}{d_0} \sim (4\phi_m^2/3C_dWe_{c,0}^2)^{1/3}(\phi_m - \phi)^{-5/6} \left(\frac{d_p}{d_0}\right)^{1/3}.
$$
 (3.6)

From (3.6), we can know that the correlation length is mainly controlled by the volume fraction ϕ and the particle size d_p . Based on this correlation length, we will further analyse the fragment size distribution in two typical breakup modes, including the bag breakup and the low-order multimode breakup. As for the shear-stripping breakup, the fragments consist mainly of i[ndividual](#page-18-0) particles and mists of the interstitial liquid. Since the particle size is known, and the mists are too small to measure, the fragment size in the shear-stripping breakup mode will not be analysed here.

3.5.1. *Fragments in bag breakup mode*

To quantify the fragment size, we use the probability density of fragment size weighted by volume, which represents the ratio of the volume of fragments with a certain diameter to the total volume. As shown in figure 13, these distributions of volume probability density in the bag breakup mode contain three distinct peaks. Combined with the morphological evolution of bag breakup, we can know that the three peaks, from small to large, correspond to the fragmentation of different parts of the droplet, including the bag, ring and nodes. Compared with silicone oil droplets, the fragment size distributions of the particle-laden droplets show distinct differences. The positions of th[e two p](#page-29-8)eaks for small droplets shift to the right, while the position of the peak for large droplets remains almost unchanged. In addition, the heights of different peaks also change, corresponding to the variation of the volume fractions of different parts.

To further analyse the effect of particles on the fragment size distribution, we need [to](https://doi.org/10.1017/jfm.2023.847) [distingui](https://doi.org/10.1017/jfm.2023.847)sh the effects of particles on different parts of the droplet (i.e. the bag, ring and nodes). For each part, the fragments essentially come from the rupture of threads or ligaments, so the fragment size distribution corresponding to each part of the droplet can be fitted with a gamma distribution (Villermaux 2020; Xu, Wang &

Figur[e](#page-29-20) [13.](#page-29-20) [V](#page-29-20)olume probability density distribution of the fragments in the bag breakup. The corresponding cases are the stars along the dashed line with $W_{e,0} = 12.2$ in figur[e 9: \(](#page-29-21)*a*) silicone oil with $\eta = 35$ mPa s, $W_{eg} = 15.7, Oh = 0.11$; *(b)* suspensi[on](#page-29-8) [with](#page-29-8) $\phi = 0.10$, $W_{eg} = 16.6$, $Oh = 0.17$; *(c)* suspension with $\phi = 0.25$, $W_{eg} = 20.3$, $Oh = 0.32$; (*d*) silicone oil with $\eta = 100$ mPa s, $W_{eg} = 20.3$, $Oh = 0.32$. $d_0 = 3.35$ mm. The dashed lines are the compound gamma distributions with the best weight fitting. Each plot shows the statistical results of approximately 60 repeated experiments under the same conditions.

Che 2022). The parameters of t[he](#page-28-4) [ga](#page-28-4)mma distribution corres[pondin](#page-27-11)g to different parts depend on the average fragment size and the corrugation of the ligaments from which the fragments originate (Villermaux, Marmottant & Duplat 2004; Villermaux 2020). Hence, the parameters of the gamma distribution are different for different parts. The different gamma distributions for different parts can be combined to form a compound gamma distribution. The overall size distribution is fitted by this compound gamma distribution, and different peaks of the overall size distribution correspond to the different sub-gamma distributions (Jackiw & Ashgriz 2022; Ade, Chandrala & Sahu 2023). In this regard, we use a compound gamma distribution to fit the overall size distribution

$$
g(x) = \sum_{i=1}^{3} w_i g_{i,v}(x),
$$
 (3.7)

where $x = d/d_0$ is the dimensionless fragment size and d_0 is the initial diameter of the droplet, w_i is the weight of each part and $w_1 + w_2 + w_3 = 1$, $g_{i,v}(x) =$

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 $\beta_i^{\alpha_i} x^{\alpha_i-1} e^{-\beta_i x}/\Gamma(\alpha_i)$ is the volume probability density function for each part, Γ is the gamma function, α_i and β_i are the shape [parameter](#page-18-0) and the rate parameter of $g_{i,v}$, respectively. Since the compound gamma distribution contains many parameters, a direct fitting using the gamma distribution may yield different results and thus make quantitative comparisons difficult. Her[e, for](#page-28-4) the fragment size distribution of the bag breakup, the interaction between the peaks is small due to the large difference in the size of the fragments produced by different parts. So we can determine α_i and β_i of each sub-gamma distribution based on the shape and position of each peak, and then obtain the weight *wi* of each peak by fitting. In this way, the compound gamma distributions are in good agreement with the measured results, as shown by the dashed lines in figure 13. Based on the parameters of each volume probability density function, we can obtain the shape parameter and the rate parameters of each number probability density function as α_i − 3 and β_i (Jackiw & Ashgriz 2022), and further estimate the average fragment size of the corresponding part as

$$
d_i = \frac{\alpha_i - 3}{\beta_i}, \quad i = 1, 2, 3,
$$
\n(3.8)

where d_1 , d_2 and d_3 are the average fragment sizes corresponding to the bag (d_b) , ring (d_r) an[d](#page-20-0) [node](#page-20-0) (d_n) (d_n) , respectively. Moreover, the weight w_i of each sub-gamma distribution is the volume fraction of that part of the fragments.

The average fragment size corresponding to different parts of the droplet obtained by (3.8) is shown in figure 14. The fragment size of particle-laden droplets is generally larger than that of silicone oil droplets, and the scaling relations between the correlation length and the fragment size are different for different parts. For the fragmentation of the bag, the liquid film retracts towards particle clusters to form the fragments, as illustrated in the inset of figure 14(*a*). The sizes of the particle clusters can be characterised by the correlation length. This is because the particle clusters essentially correspond to the concentr[ation](#page-19-0) fluctuation (i.e. the particle agglomeration or separ[ation\),](#page-20-0) [and](#page-20-0) the coherence length reflects the range of concentration fluctuation caused by the heterogeneous effect of particles. Hence, the fragment size of the bag (d_b) [can be esti](#page-20-0)mated as

$$
\frac{d_b}{d_0} \sim \frac{\xi}{d_0}.\tag{3.9}
$$

The relation is in good agreement with the data in figure $14(a)$, where the scaling in (3.9) is shown as the solid line with a prefactor of $C_1 = 1.12$.

For the fragmentation of the ring, each fragment corresponds to a segment of the liquid ring, as illustrated in the inset of figure 14(*b*). The initial diameter of the liquid ring, which is approximately equal to the thickness of the droplet (*h*) after the [initia](#page-28-3)l flattening, is larger than the correlation length (ξ) . But as the liquid ring is stretched and thinned, the diameter of the liquid ring (h_1) is smaller than the correlation length. At this condition, the concentration fluctuation caused by the particle heterogeneity can lead to the destabilisation of the ring and then produce fragments. Therefore, the correlation length reflecting the range of concentration fluctuation can characterise the wavelength of the destabilisation, which is the length of the segment of the liquid ring where a fragment [is](https://doi.org/10.1017/jfm.2023.847) [produced](https://doi.org/10.1017/jfm.2023.847). In addition, the diameter of the liquid ring (h_1) is mainly controlled by the rim expansion ratio, which is affected by the Weber number (Jackiw & Ashgriz 2021). Hence, h_1 can be considered as a constant due to the same inviscid Weber number in our experiments. Therefore, the fragment size of the ring can be estimated based on the

Figure 14. Average fragment sizes of (*a*) the bag, (*b*) the ring and (*c*) the node under different particle volume fractions and particle sizes. The scatters are the fragment size of the particle-laden droplets obtained from (3.8). The dashed and dotted lines correspond to the fragment sizes of silicone oil droplets with $\eta = 35$ and 100 mPa s, respectively. Here, ξ/d_0 is calculated according to (3.6), $d_0 = 3.35$ mm. The corresponding cases are the stars along the dashed line with $W_{e,c,0} = 12.2$ in figure 9. Each data point shows the statistical results of approximately 60 repeated experiments under the s[ame conditi](#page-20-0)on. The insets illustrate the fr[agmen](#page-20-1)t mechanisms for the corresponding parts.

mass conservation as

$$
\frac{d_r}{d_0} \sim \left(\frac{\xi}{d_0}\right)^{1/3}.\tag{3.10}
$$

The relation is in good agreement with the data in figure $14(b)$, where the scaling in (3.10) is shown as the solid line with a prefactor of $C_2 = 0.78$.

Finally, for the fragmentation of the nodes, the fragment size is almost constant unless the correlation length ξ is large ($\xi/d_0 > 0.1$), as shown in figure 14(*c*). This is because the effect of particle heterogeneity occurs when the characteristic scale is smaller than the correlation length. For the nodes with a large scale, the size of the node fragments will be affected only when the correlation length is large, that is, a high volume fraction or a large [particle size. F](https://doi.org/10.1017/jfm.2023.847)or particle-laden droplets with $\xi/d_0 < 0.1$, we find that the fragment size of the node obeys

$$
\frac{d_n}{d_0} = C_3 \frac{h}{d_0},\tag{3.11}
$$

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Figure 15. Variation of the volume fraction of each part with (*a*) the particle volume fraction and (*b*) the particle size. The corresponding cases are the stars along [the dashed](#page-20-0) line with $W_{c,0} = 12.2$ in figure 9. Each data point shows the statistical results of approximately 60 repeated experiments under the same conditions. (*c*) Illustration of the droplet morphology after the initial flattening. (*d*) Illustration of the liquid flow in the bag development process and the effect of particles on flow inhibition.

where the thickness o[f the d](#page-29-22)roplet after the initial flattening *h* is [a](#page-28-21) [con](#page-28-21)stant for the cases in this study, and C_3 is a constant equal to 1.55. In figure $14(c)$, the dashed line for silicone oil droplets with $\eta = 35$ mPa s corresponds to $C_3 = 1.53$ ($d_n/d_0 = 0.481$), and the dotted line for silicone oil droplets with $\eta = 100$ mPa s corresponds to $C_3 = 1.47$ ($d_n/d_0 = 0.460$). This law is consistent with the end-pinching mechanism proposed by Schu[lkes \(1996](#page-21-0)) for free liquid jets (with $C_3 = 1.55$), that is, the droplets are shed via end pinching caused by the retraction of ligament tips, as illustrated in the inset of figure $14(c)$. A similar law was also found in the breakup of Worthington jets (Gordillo $\&$ Gekle 2010) and unsteady sheets (Wang & Bourouiba 2018), which have $C_3 = 1.5{\text -}1.6$. Here, our results show that the fragmentation of the node also follows this scaling law [for the en](#page-21-0)d-pinching mechanism.

In addition to the average fragment size, the volume fraction of each part is also important for understanding the fragment size distribution. The volume fraction of each part obtained by the weight w_i of each sub-gamma distribution is shown in figure 15(*a*,*b*). To analyse the reason of the volume fraction of each part, we need to consider the mass [distribution](https://doi.org/10.1017/jfm.2023.847) in the droplet after the initial flattening. This is because the evolution of the droplet shape and the subsequent breakup process determine the volume fraction of each part. After the initial flattening, the droplet is deformed into a disk with a thickness of $h \approx 1.05$ mm and a width of $d_f \approx 5.6$ mm, as shown in figure 15(*c*). Correspondingly, the

volu[me fr](#page-20-2)actions at the rim and middle of the droplet are roughly 56% and 44% . respectively (based on volu[me calcula](#page-21-0)tion from figure 15*c*). Then the mid[dle of t](#page-28-4)he droplet develops into the bag, and the rim develops into the ring and nodes. During this process, liquid flows from the bag to the ring and from the ring to the nodes (Jackiw & Ashgriz 2022), as illustrated in figure 15(*d*). These processes ultimately determine the volume fraction of each part. For the silicone oil droplet ($\phi = 0$ in figure 15*a*), the flow from the bag to the ring reduces the volume fraction of the middle bag from the initial 44 % to 25 % after the breakup. The volume fraction of the nodes calculated from the fragment size (3.11) and number (usually approximately 3) is 28 %, which is in agreement with the experimental data (27 %) in figure 15(*a*). In addition, Jackiw & Ashgriz (2022) predicted the fragment size distribution of water droplets, and found the volume fraction of the bag, ring and nodes were 12 %, 45 % and 38 %, respectively. Compared with their prediction, the volume fraction of the bag in our results is larger, while the volume fraction of the node is smaller. Besides the effect of a slight difference in *Wec*,0, this difference may be due to the higher viscosity of the droplet in our experiments, which inhibits the liquid flow between the pa[rts. These](#page-21-0) comparisons also indicate that the volume fractions obtained from the compound gamma distribution are reasonable.

Compared with the silicone oil droplet, there are several differences in the volume fractions of different parts of the particle-laden droplet. First, the volume fraction of the bag of the particle-laden droplet is much larger, i.e. approximately 40 %. This is because, during the [bag develo](#page-21-0)pment process, the particles tend to be retained at the junction of the bag film and the ring. This results in a higher local particle concentration and correspondingly a [higher vi](#page-21-0)scosity, which inhibits the liquid flow from the bag to the ring, as illustrated in figure 15(*d*). In additi[on, the pa](#page-20-0)rticle clusters on the liquid film can trap some liquid. Therefore, the bag volume fraction of the particle-laden droplet is maintained at about 40 %, i.e. the initial volume fraction in the middle of the droplet. Second, as the p[article volu](#page-21-0)me fraction increases, the volume fraction of the nodes decreases first and then increases, while the volume fraction of the ring increases first and then decreases, as shown in figure 15(*a*). This is because the higher particle volume fraction inhibits the liquid flow from the ring to the nodes. However, when the particle volume fraction is very large (ϕ = 0.35 in figure 15*a*), the presence of the particles makes the fragment size of the node increase significantly (as shown in figure 14*c*), and the corresponding volume fraction of the node also increases. Third, as the [partic](#page-28-22)le size increases, the volume fraction of the nodes increases gradually, and the volume fractions of the bag and ring decrease, as shown in figure 15(*b*). This is because the larger particle tends to reside on the larger-scale nodes rather than the smaller-scale bag and ring.

In addition to considering the effects of particles on different parts, the fragment size distribution can also be described by some representative diameter to quantify the overall atomisation effect. A representative diameter is the Sauter mean diameter (SMD), which is the diameter of a droplet whose ratio of volume to surface area is the same as that of the entire fragments (Lefebvre & McDonell 2017). The SMD can be calculated as

$$
SMD = \frac{\sum n_i d_i^3}{\sum n_i d_i^2} = \frac{1}{\sum w_i / d_i},
$$
\n(3.12)

where n_i is the number of fragments corresponding to a size of d_i and equal to $w_i d_0^3 / d_i^3$. Combining the fragment sizes (obtained by (3.9) , (3.10) and (3.11)) and the corresponding

Figure 16. Sauter mean diameter of fragments at different particle volume fractions and sizes in the bag breakup mode. The solid line is (3.13) with a prefactor of 1.4. The corresponding cases are the stars along the dashed line with $W_{e_0} = 12.2$ in figure 9. Here, ξ/d_0 is calculated according to (3.6), $d_0 = 3.35$ mm. Each data point shows the statistical results of approximately 60 repeated experiments under the same conditi[ons.](#page-19-0)

[volum](#page-20-1)e frac[tions](#page-20-2) of different parts, we can estimate [the S](#page-23-0)MD as

$$
\frac{SMD}{d_0} \sim \frac{1/d_0}{w_1/d_b + w_2/d_r + w_3/d_n} = \left[\frac{w_1}{C_1} \left(\frac{\xi}{d_0} \right)^{-1} + \frac{w_2}{C_2} \left(\frac{\xi}{d_0} \right)^{-1/3} + \frac{w_3}{C_3} \left(\frac{h}{d_0} \right)^{-1} \right]^{-1}.
$$
\n(3.13)

By substituting the prefactors $C_1 = 1.12$, $C_2 = 0.78$, $C_3 = 1.55$ (obtained from (3.9), (3.10) and (3.11)) [and the w](#page-24-0)eights $w_1 = 0.405$, $w_2 = 0.345$, $w_3 = 0.25$ (obtained by averaging all the cases considered in this study) into (3.13) , we can see that the relationship is [in good a](#page-24-0)greement with the experimental data, as shown in figure 16, where (3.13) is shown as the solid line with a prefactor of 1.4.

3.5.2. *Fragments in multimode breakup*

The volume probability density distribution [of the frag](#page-24-0)ments in the low-order multimode breakup is shown in figure 17. The fragments are also formed by the fragmentation of three parts, including the middle node, the peripheral nodes and the peripheral ring, as illustrated in figure $17(c)$. The fragmentation of the middle node forms a [distinct p](#page-25-0)eak with the largest fragment size. But the fragment sizes formed by the fragmentation of the peripheral nodes and ring are close, and the corresponding peaks are indistinguishable. Therefore, we still use the compound of three gamma distributions to fit the data, but do not distinguish the effects of particles on different parts. We only use the fitting curves to identify the changes in the overall size distributions. As shown in figures $17(a,b)$, the fragment size distribution [shifts](https://doi.org/10.1017/jfm.2023.847) [to](https://doi.org/10.1017/jfm.2023.847) [the](https://doi.org/10.1017/jfm.2023.847) right as the particle volume fraction and particle size increase, that is, the overall fragment size of the particle-laden droplets becomes larger.

The SMDs of the fragment size distributions are shown in figure 18. For the multimode breakup, the droplet is pierced by RT instability waves to form fragments with a typical

Breakup of particle-laden droplets in airflow

Figure 17. Variation of the volume probability density distribution in the low-order multimode breakup with (*a*) the particle volume fraction and (*b*) the particle size. The corresponding cases are the stars along the dashed line with $W_{e_0} = 36$ $W_{e_0} = 36$ $W_{e_0} = 36$ in figure 9, i.e. fluids used are silicone oil with $\eta = 35$ mPa s, $W_{e_g} = 51.6$, *Oh* = 0.11, and suspensions with $\phi = 0.10$, $W_{e} = 56.3$, $Oh = 0.17$ and $\phi = 0.25$, $W_{e} = 71.7$, $Oh = 0.32$. Here, $d_0 = 3.35$ mm. The solid lines are the compound gamma distributions with the best weight fitting. Each plot shows the statistical results of approximately 90 repeated experiments under the same conditions. (*c*) Droplet morphology at 14.5 ms in low-order multimode breakup mode, correspo[nding](#page-28-23) [t](#page-28-23)o the whole breakup [process](#page-28-24) shown in figur[e 5.](#page-29-23)

size (d_s) . The breakup is extremely unstable, and the heterogeneity of the particle-laden fluid will intensify the breakup randomness, as described in $\S 3.4$. For this condition, we can refer to the full-wave approach to predict the SMD. The full-wave approach assumes that there is a spectrum of wavelengths which could be excited to appreciable amplitudes and then produce fragments of corresponding size (Mayer 1961; Qin, Yi & Yang 2018; Yang *et al.* 2020). Following this idea, in the condition of the multimode breakup, the fragments originate from the piercing caused by the RT instability wave, so the sizes of the fragments (d_s) can be estimated as

$$
d_s = C_s \lambda_{RT},\tag{3.14}
$$

where C_s is a constant and λ_{RT} is the RT instability wavelength. In addition, the number probability of fragments with size d_s is required to calculate SMD. However, due to the close size of the fragments originating from different parts, the exact size distribution of [fragments](https://doi.org/10.1017/jfm.2023.847) [is](https://doi.org/10.1017/jfm.2023.847) [di](https://doi.org/10.1017/jfm.2023.847)fficult to be quantified by the compound gamma distributions. Since we mainly focus on the effect of particles, we, for simplicity, neglect the variation of the volume probability density with the fragment size d_s , that is, the volume of the initial droplet is divided uniformly into *N* parts and each of which is broken into fragments with

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Figure 18. Sauter mean diameter of fragments at different particle volume fractions and particle sizes in the low-order multimode breakup. The corresponding cases are the stars along the dashed line with $W_{e,c,0} = 36$ in figure 9. Here, ξ/d_0 is calculated according to (3.6), $d_0 = 3.35$ mm. The solid line is (3.17) with $C_s = 0.764$. Each point shows the statistical results of approximately 90 repeated experiments un[der th](#page-24-1)e same [condit](#page-25-1)ion.

size d_s . Therefore, the number probability (*n*) of fragments with size d_s is estimated as

$$
n \sim \frac{d_0^3}{Nd_s^3}.\tag{3.15}
$$

By integrating over the spectrum of wavelengths and combining (3.14) and (3.15), the SMD can be written as

$$
SMD = \frac{\int_{\lambda_{min}}^{\lambda_{max}} nd_s^3 d\lambda}{\int_{\lambda_{min}}^{\lambda_{max}} nd_s^2 d\lambda} = C_s \frac{\lambda_{max} - \lambda_{min}}{\ln \lambda_{max} - \ln \lambda_{min}},
$$
(3.16)

where λ_{max} and λ_{min} are the upper and lower limits of the spectrum of wavelengths. For droplet breakup, the wavelength must be smaller than the droplet diamet[er, i.e](#page-25-3). λ_{max} = *d*0. The correlation length of the particle-laden droplet is the typical length over which a fluctuation will be damped, so short-wavelength instabilities will be s[uppressed](#page-25-0) by the [partic](#page-25-3)les, i.e. $\lambda_{min} = \xi$. Therefore, the SMD in (3.16) can be written in dimensionless form as

$$
\frac{SMD}{d_0} = C_s \frac{1 - \xi/d_0}{-\ln(\xi/d_0)}.
$$
\n(3.17)

[Through](https://doi.org/10.1017/jfm.2023.847) [th](https://doi.org/10.1017/jfm.2023.847)e full-wave approach with a lower limit ξ , the relation in (3.17) reflects the effect of the particle in intensifying the breakup randomness and suppressing the short-wavelength instabilities, and is in good agreement with the data in figure 18, where (3.17) is the solid line with $C_s = 0.764$.

4. Conclusions

In this study, we [inv](#page-11-1)estigate the breakup of particle-laden droplets in airflow experimentally. By combining high-speed images from the side view and the $45°$ view, the morphologies of particle-laden droplets and homogeneous fluid droplets are compared in detail. In the bag breakup mode, the particles agglomerate on the liquid film and accelerate the breakup of the liquid film. In the multimode breakup, the particles induce localised rapid piercing. In the shear-stripping breakup, the breakup process is more discrete, and the particles and interstitial liquid are almost completely separated. A breakup regime map is produced, and transitions between different breakup modes are obtained based on a unified model (3.1). The effect of particles on the transition of breakup modes is manifested by a higher effective viscosity compared with the interstitial fluid.

Furthermore, to quantitatively evaluate the effects of particles, we eliminate the effect of the higher effective viscosity for particle-laden droplets by comparing the cases with the same $W_{\mathcal{C}_c,0}$. For the stretching and fragmentation of the liquid film, the particles accelerate the breakup of the film due to the less liquid content in the film and the local stress caused by the particles. For the piercing in the droplet middle, the particles induce local piercing due to the change in effective viscosity when the droplet thickness is reduced close to the particle size, which further leads to more abundant and random piercing phenomena of the particle-laden droplet.

Finally, considering the coupling effects of the initial flattening and later stretching in the droplet breakup process, we propose a correlation length (ξ) that controls the stability of the suspension through a concentration fluctuation. Based on the correlation length, the fragment size distributions in the bag breakup and low-order multimode breakup are analysed theoretically. For the bag breakup, the fragment sizes corresponding to different parts of the droplet obey different scaling relations with the correl[ation length,](https://doi.org/10.1017/jfm.2023.847) [and the volume f](https://doi.org/10.1017/jfm.2023.847)ractions of different parts also change with the particle diameter and the volume fraction. By combining the fragment sizes and the corresponding volume fractions of different parts, the SMD of fragments of the particle-laden droplet in the bag breakup mode is obtained. For the multimode breakup, the overall fragment size of the particle-laden droplet is larger, and its SMD can be predicted by the full-wave approach with the correlation length as the lower limit.

[Su](https://orcid.org/0000-0002-8625-3329)pplement[ary](https://orcid.org/0000-0002-8625-3329)[material](https://orcid.org/0000-0002-8625-3329)[and](https://orcid.org/0000-0002-8625-3329)[movies.](https://orcid.org/0000-0002-8625-3329) Supplementary material and movies are available at https://doi.org/ [10](https://orcid.org/0000-0002-0682-0603).1017/jfm.2023.847.

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Appendix. Rheological measurements of the suspensions

The rheology of the suspensions was measured using a rotational rheometer (Anton Paar MCR 501) with a rough parallel plate. The plates had a diameter of 25 mm and the gap [between](https://doi.org/10.1017/jfm.2023.847) [the](https://doi.org/10.1017/jfm.2023.847) [pla](https://doi.org/10.1017/jfm.2023.847)tes was 1 mm. Before the measurements, the suspension was pre-sheared at 0.01 r.p.s. for three minutes. During the measurements, each data point was obtained by maintaining the corresponding shear rate for 10 s. The test temperature was controlled at 22 \degree C. The results are presented in figure 19. As the particle volume fraction increases, the

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Figure 19. Viscosity of the suspensions with different particle volume fractions (ϕ) .

suspension gradually exhibits non-Newtonian properties at low shear rates. But at a high shear rate, the suspensions with different particle volume fractions maintain an almost constant viscosity. For droplet breakup in the airflow, the droplet experiences a high shear rate. It can be estimated by the characteristic shear rate $\dot{\gamma} = u_g \sqrt{\rho_g/\rho_f}/d_0$ (Li *et al.* 2022), which is in the range of 90–700 s⁻¹ in our experiments. Therefore, we used the averaged viscosity at shear rates of 100–1000 s⁻¹ as the effective viscosity of the suspensions.

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