Effects of thermal fluctuations in the fragmentation of a nanoligament

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Effects of thermal fluctuations in the fragmentation of a nanoligament

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I. INTRODUCTION

The hydrodynamical description of a nonideal interface at the microscales is generically assessed via the combined effects of viscous dissipation and surface tension forces [1–7]. Pushing such description toward smaller scales presents two main difficulties. From one side, hydrodynamics itself cannot hold true at all scales of motion, and its coarse-grained foundations are inevitably weakened whenever the physics gets closer to the atomistic level. From the other side, if hydrodynamics needs to be corrected by atomistic effects, one faces the issue of determining the appropriate description to adopt. The equations of fluctuating hydrodynamics [8,9] provide a promising route, accounting for molecular collisions via the introduction of stochastic contributions to the nonequilibrium fluxes, as originally proposed by Landau [8]. Such treatment has the major appeal of retaining the generic hydrodynamic approach, since it does not enter into a detailed characterization of the atomistic motion, but rather takes a coarse-grained perspective, where molecular effects are modeled via the fluctuation-dissipation balance [10]. So far, a consistent body of work considered various aspects of interfacial flows in open and confined geometries [2,5,11–14], while there are only very few works considering the effect of thermal fluctuations [15–22]. At small scales, thermal fluctuations promote interface excitations with energy $k_B T$; these are resisted by the surface tension $\gamma$, which opposes a force (per unit length) against the deformation of the interface. This balance determines a new length scale, named the thermal length scale, defined as [23]

$$\ell_T = \sqrt{\frac{k_B T}{\gamma}}.$$

which is typically in the nanometer range (or fractions of it) [24,25]. On general grounds, thermal fluctuations are expected to become increasingly more relevant and produce measurable effects when moving from micrometer scales down to smaller nanometer scales. This is the case, for example, of nanojets, where stochastic hydrodynamic equations have been used to study the interface dynamics [19,20] with extensive comparisons with fully atomistic descriptions [19]. It was convincingly shown that thermal fluctuations impact the breakup properties of nanojets. For the spreading of viscous drops on a solid substrate, thermal fluctuations have been shown to accelerate the spreading in comparison to the deterministic case [26]. This has been studied with the help of fluctuating hydrodynamics in the lubrication limit [15], and later confirmed by mesoscale numerical simulations [21]. Other numerical methods were used to study dewetting of thin liquid films, and it was observed that thermal fluctuations accelerate the rupture of films [17]. Also, experimental studies exist [18,22], concerning the breakup of a satellite droplets [22], but no quantitative characterization of the distribution of droplets size has been provided. In the study [18], the pinch-off process has also been found to be affected by thermal fluctuations, since an initial viscocapillary regime is followed by a fluctuation-dominated regime when the characteristic size of the neck approaches the thermal length.

The focus of this paper is on the quantitative analysis of the effects of thermal fluctuations on the statistics of the breakup times and on the droplet volume distribution, following the

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We study the effects of thermally induced capillary waves in the fragmentation of a liquid ligament into multiple nanodroplets. Our numerical implementation is based on a fluctuating lattice Boltzmann (LB) model for nonideal multicomponent fluids, including nonequilibrium stochastic fluxes mimicking the effects of molecular forces at the nanoscales. We quantitatively analyze the statistical distribution of the breakup times and the droplet volumes after the fragmentation process at changing the two relevant length scales of the problem, i.e., the thermal length scale and the ligament size. The robustness of the observed findings is also corroborated by quantitative comparisons with the predictions of sharp interface hydrodynamics. Beyond the practical importance of our findings for nanofluidic engineering devices, our study also explores a novel application of LB in the realm of nanofluidic phenomena.

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contraction and fragmentation of a liquid nanoligament. Similarly to other studies [19,20], we rely on a hydrodynamical description coupled to the effects of thermal noise. However, we do not solve the continuum equations of interfacial hydrodynamics with a sharp interface directly, but instead we utilize the fluctuating multicomponent lattice Boltzmann (LB) model [27]. The study presented herein contributes to show a realistic application of fluctuating LB for nanofluidic phenomena. Our main findings can be summarized as follows: we confirm that thermal fluctuations are able to accelerate the nanoligament fragmentation process, as it was previously reported [19,20]. Furthermore, we give quantitative information on how much the thermal noise can accelerate the breakup process, and we present the effects of thermal fluctuations on the polydispersity of droplets distribution. Last but not least, we find that the LB simulations with thermal noise are consistent with sharp interface hydrodynamics results. The paper is organized as follows: in Sec. II we briefly review the fluctuating LB used; in Sec. III we present technical details of the numerical simulations; in Sec. III A we report on the destabilization process driven by the Plateau-Rayleigh instability and the qualitative effects of thermal fluctuations; in Sec. III B we discuss results on the statistics of breakup times, while in Sec. III C we report on the statistics of the droplet volumes; in Sec. III D we provide a quantitative comparison between the LB results and the results of sharp interface hydrodynamics; conclusions will follow in Sec. IV.

II. MODEL: FLUCTUATING LATTICE BOLTZMANN FOR MULTICOMPONENT FLUIDS

Beyond the traditional problems of homogeneous hydrodynamics [28–30], LB models have proven particularly suitable for the modeling of complex fluids with multiple phases and/or components [31]. Moreover, stimulated by earlier contributions for homogeneous fluids [32–34], recently there has been a significant work to include the effects of thermal fluctuations in LB for multiphase [35,36] and multicomponent flows [27,37]. Technical details of the fluctuating LB have already been extensively presented in [27], and here we only briefly recall the most important facts for the sake of completeness. We employ the D3Q19 LB model, which discretizes the momentum lattice most important facts for the sake of completeness. We employ probability distributions functions

In Sec. III we present technical details of the numerical simulations; in Sec. III A we report on the destabilization process driven by the Plateau-Rayleigh instability and the qualitative effects of thermal fluctuations; in Sec. III B we discuss results on the statistics of breakup times, while in Sec. III C we report on the statistics of the droplet volumes; in Sec. III D we provide a quantitative comparison between the LB results and the results of sharp interface hydrodynamics; conclusions will follow in Sec. IV.

\[ f_i(x_i,t+1) - f_i(x_i,t) = \mathcal{L} (f_i(x_i,t)) + F_i(x_i,t) + \xi_i(x_i,t), \]

where \( \mathcal{L} \) is a collision kernel, \( F_i \) is a source coming from nonideal forces, and \( \xi_i \) is a stochastic source. For simplicity, we have used a unitary time step. The macroscopic quantities such as density \( \rho \) (one for each component) and global velocity \( v \) are readily evaluated from the distribution functions:

\[ \rho(x_i,t) = \sum_i f_i(x_i,t), \quad v(x_i,t) = \frac{\sum_i f_i(x_i,t)c_i}{\rho(x_i,t)}, \]

where \( \rho = \rho_A + \rho_B \) is the total density. Regarding the collisional operator \( \mathcal{C} \), we use a multirelaxation time (MRT) scheme [34,38,39]. The basic idea behind the MRT scheme is to introduce a vector basis \( e_i \) \( (n = 0, \ldots, 18) \) to decompose the probability distribution functions into “modes,” \( M_n = \Sigma_i e_i f_i \). The lowest-order modes coincide with hydrodynamic modes (density, momentum, stress tensor) while higher-order modes (“ghost” modes) do not contribute to the hydrodynamic behavior of the LB models [34,38]. Each one of the modes is relaxed with its own relaxation frequency toward the corresponding equilibrium mode calculated from the equilibrium distribution,

\[ f_i^{eq}(x_i,t) = \rho_i(x_i,t) c_i, \]

where \( \rho_i \) and \( \omega_i \) are weights associated with the discrete lattice directions. The nonideal forces are chosen in the Shan-Chen formulation [31,40–43]:

\[ F_i(x_i,t) = -G \nabla f_i(x_i,t) = -G \rho_i(x_i,t) \sum_{l \neq i} \omega_l \nabla f_l(x_i,t) c_i, \]

where \( G \) is a coefficient that regulates the strength of the interactions between the two components. The pseudopotential \( \phi_l \) is set equal to the density for the sake of simplicity, i.e., \( \phi_l(x_i,t) = \rho_l(x_i,t) \). When the coupling strength is large enough, the system can show phase segregation with the formation of diffuse interfaces separating bulk regions with the majority of one of the two components. Diffuse interfaces display widths of the order of a few grid sizes and a positive surface tension \( \gamma \), which increases at increasing \( G \). The term \( \xi_{li} \) in Eq. (1) is a noise term that is assumed to be a zero-mean Gaussian random variable, uncorrelated in time and with constant variance (which can, however, be space-dependent). While noise does not introduce stochastic forces on the density modes, it does so in momentum modes with the following correlations:

\[ \langle \xi_{ln}(x_i(t)) \xi_{ln'}(x_i(t')) \rangle = (2\kappa - \kappa^2) k_B T \frac{\rho_l \rho_{l'}}{\rho_l + \rho_{l'}} \delta_{nn'} \delta(x - x') \delta(t - t'), \]

where \( n, n' \) refer to the modes and \( \kappa \) represents the relaxation frequency of the momentum modes in the MRT scheme. The noise correlations on higher modes satisfy

\[ \langle \xi_{ln}(x_i(t)) \xi_{ln'}(x_i(t')) \rangle = (2\kappa - \kappa^2) k_B T \frac{\rho_l \rho_{l'}}{c_s^2} \delta_{nn'} \delta(x - x') \delta(t - t'), \]

where \( n, n' = 4, \ldots, 18 \).
Fig. 1. Ligament breakup with LB simulations. Left column panels: LB deterministic evolution of a liquid ligament destabilized by the Plateau-Rayleigh instability. The system is initialized with a small perturbation on the fastest growing mode with wavelength $\lambda_{\text{fast}}$. The axial system size of the numerical simulations is chosen to accommodate $2\lambda_{\text{fast}}$. After the breakup, we observe that the volumes of the two “mother” droplets are equal to each other, and the same happens for the “satellite” droplets. Middle column panels (“without-TN” protocol): random initial condition followed by LB deterministic evolution (see text for details on preparation). Right column panels (“with-TN” protocol): random initial condition followed by fluctuating hydrodynamics evolution with thermal noise (details are reported in the text).

where $N_n$ are normalization constants fixed by $N_n\delta_{nn'} = \sum_i w_i e_n e_{n'}$. Notice that all other noise correlation vanish.

At hydrodynamical scales, the fluctuating LB allows us to obtain—via the Chapman-Enskog analysis [34,39]—the stochastic hydrodynamic equations for a binary fluid (repeated indexes denote “summed upon”) [9],

\[
\begin{align*}
\frac{\partial}{\partial t} \rho_{\text{tot}} + \frac{\partial}{\partial x} (\rho_{\text{tot}} v_x) &= 0, \\
\frac{\partial}{\partial t} \rho_A + \frac{\partial}{\partial x} (\rho_A v_x) &= \frac{1}{2} \frac{\partial}{\partial x} \left[ D \frac{\partial}{\partial x} \bar{\mu} + \Psi_x \right], \\
\frac{\partial}{\partial t} \rho_{\text{tot}} v_x + \frac{\partial}{\partial x} (\rho_{\text{tot}} v_x v_y) &= -\frac{\partial}{\partial y} P_{xx} + \frac{\partial}{\partial y} \left[ \eta (\frac{\partial_x v_y + \partial_y v_x}{2}) + \Sigma_{\text{fluid}} \right].
\end{align*}
\] (7)

The equilibrium properties are fully encoded in the chemical potential $\mu$ and the pressure tensor $P_{\alpha\beta}$, which depend on the interaction model chosen at the level of LB [40,41] and whose expressions may be found elsewhere [43]. The terms $\Psi_x$ and $\Sigma_{\text{fluid}}$ are stochastic fluxes and tensors, respectively. Specifically, the stochastic vector field $\Psi_x$ is the term due to the thermal noise that must be added to the diffusion flux $D \frac{\partial}{\partial x} \bar{\mu}$ [9], with $D$ the diffusion constant; the stochastic tensor $\Sigma_{\text{fluid}}$ is added to the viscous stress tensor $\eta (\frac{\partial_x v_y + \partial_y v_x}{2})$ [8], with $\eta$ the dynamic viscosity for the bulk. Requiring that the fluctuation-dissipation relation holds for our hydrodynamic problem, and using (5) and (6), one can derive a unique choice for the intensity of the stochastic contributions [9,27]:

\[
\Sigma_{\alpha\beta} = \sqrt{\eta k_B T (W_{\alpha\beta} + W_{\beta\alpha})}, \quad \Psi_x = \sqrt{2Dk_B T} \tilde{W}_x.,
\] (9)

where $k_B$ is the Boltzmann constant, and $W_{\alpha\beta}$ and $\tilde{W}_x$ are random Gaussian tensors and a random Gaussian vector field, respectively, with independent components and variance equal to unity.

III. NUMERICAL SETUP AND RESULTS

Numerical simulations are conducted in a 3D fully periodic domain with sizes $L_x \times L_y \times L_z$. A cylindrical ligament with a majority of phase $A$ and radius $R_0$ is set up with a symmetry axis along the $z$ coordinate (see Fig. 1). By keeping fixed the ratio between the system sizes $L_x, L_y, L_z$ and the ligament initial radius $R_0$, we have performed different numerical simulations at changing the thermal length $\ell_T$ and the domain resolution $L_x \times L_y \times L_z$. For each realization of the thermal length and domain resolution, we performed hundreds of simulations to gather sufficient statistics over the breakup time and the droplet volumes after breakup. For computational reasons, larger resolutions are associated with a smaller number of simulations. All these parameters are summarized in Table I.

We remark that applications of LB in the problem of ligament contraction have already been proposed in the literature. In particular, in [44] a comparison among axisymmetric LB and the predictions of deterministic sharp interface hydrodynamics has been provided. The use of an axisymmetric LB obviously reduces the computational effort; however, in the presence of thermal noise, it does not provide a realistic description of the interfacial fluctuations typical of 3D interfaces [24,25]. For this reason, we have used a fully 3D fluctuating LB without any axial symmetry. In all the simulations that we conducted, the length of the domain size is chosen to be $L_z \approx 18R_0$, which is well suited to accommodate roughly two wavelengths $\lambda_{\text{fast}}$ of the fastest-growing mode of the...
Plateau-Rayleigh instability [45] (see also Fig. 1 for quantitative details). The numerical simulations are then conducted for a set of parameters for which the Ohnesorge number Oh is small (Oh < 1). The Ohnesorge number quantifies the importance of the viscous forces with respect to the inertial and surface tension forces, and it is defined as Oh = η/√ρ°maxγR°max, where ρ°max indicates the maximum density of the ligament and η is the dynamic viscosity. The fastest-growing mode has a wave number k°fastR°0 ≈ 0.697, where λ°fast = k°fast/√ρ°0, independently of Oh [45].

### A. Effects of thermal fluctuations

We are interested to understand the effect of thermal noise on the ligament breakup in combination with the Plateau-Rayleigh instability. To this aim, we have designed three different simulation cases, as shown in Fig. 1. In the first case (left panels), we evolve the ligament without thermal noise, i.e., we use Eqs. (7) and (8) with k°B T = 0. We set an initial small perturbation with wavelength λ°fast and very small amplitude. Due to the Plateau–Rayleigh instability, the unstable mode along the interface grows and eventually determines the breakup of the ligament into droplets. In the second case (middle panels), we evolve the same equations with a different initial condition: beyond the perturbation on the fastest-growing mode, we also add a random Gaussian perturbation on the less unstable Fourier modes; the evolution is kept deterministic, i.e., again we use Eqs. (7) and (8) with k°B T = 0. In the third case (right panels), we show results coming from numerical simulations with the same initial condition used for the middle panels followed by the fluctuating hydrodynamics evolution, i.e., Eqs. (7) and (8) with k°B T > 0. Typically, after a few capillary times t°c = √ρ°maxR°03/γ, the ligament breaks into two “mother” droplets and two “satellite” droplets. However, in the case of thermal fluctuations (right panel in Fig. 1), the enhanced volume polydispersity may cause one of the satellite droplets to be so small that it cannot be resolved with the resolution used. The presence of two mother droplets is clearly due to the fact that we choose an axial length that corresponds to twice the wavelength of the fastest-growing mode. The presence of small satellite droplets is generated by the combined effect of viscosity and surface tension at the late stage of pinch-off, as already described in the literature [13,46,47]. This qualitative feature is robust and independent of the thermal noise and of the initialization protocol used.

For the pure deterministic case (left panels), the fragmentation process evolves in a symmetric way, and it leads to two identical mother droplets and two identical satellite droplets. However, when we add noise either in the initial configuration or during the entire evolution, things become more complicated: the breakup time is a random variable, and a volume polydispersity in both mother and satellite droplets is observed. In particular, the presence of thermal noise in the evolution (right panels) manifestly accelerates the breakup, which is consistent with previous studies [19,20]. This is because the effects of thermal fluctuations dominate at the late stage of the pinch-off regime. It is apparent from Fig. 1 that during the ligament fragmentation process, both the random initial conditions and thermal fluctuations have a role; more importantly, numerical simulations offer the possibility to quantitatively disentangle the two contributions by using two complementary simulations protocols: we can change the random initial condition and integrate a deterministic dynamics without thermal noise (k°B T = 0) in Eqs. (7) and (8) (“without-TN” protocol, as in the middle panels of Fig. 1) or we can include the effects of thermal noise in the dynamics by using k°B T ≠ 0 in Eqs. (7) and (8) (“with-TN” protocol, as in right panels of Fig. 1). Based on these two simulation protocols, in the following we aim at characterizing the statistics of the droplet breakup times and the droplet volume distribution. The quantitative characterization of the droplet volume statistics naturally poses the question of the specificity of the results, due to the fact that LB is a diffuse interface hydrodynamics solver. To address this point, we will perform a quantitative comparison between the droplet volume statistics from the simulations and the predictions of sharp interface hydrodynamics.

### B. Statistics of breakup times

Based on previous experimental and numerical works [18,20,22], we know that the thermal noise results in accelerating the late stage of the pinch-off process. Beyond this accelerated dynamics, here we focus on characterizing the statistics of the breakup times. We have conducted numerical simulations for a fixed ligament radius R°0 = 7 lbu and different thermal lengths t°f, ranging from 7 × 10⁻⁵ to 3 × 10⁻³ lbu, in the two simulation protocols “with-TN” and “without-TN.” To measure the breakup time, we follow the ligament fragmentation evolution and record the time when a discontinuity in the density profile is observed along the ligament axis. In Fig. 2,

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**TABLE I. Summary of the different numerical simulations conducted.** All the numerical simulations that we describe in this paper are performed with a coupling coefficient C = 1.5 LB units (lbu hereafter) in Eq. (4). Inside the ligament, the density for the two components is set to ρA = 2.21 lbu and ρB = 0.09 lbu, with a corresponding total density ρ°tot = 2.3 lbu. The surface tension of the system is γ = 0.1515 lbu. The surface tension is kept fixed in all the numerical simulations, while the noise intensity is varied in Eqs. (7) and (8) to achieve different thermal lengths. The viscosity ratio between the dispersed and continuous phases is set equal to unity.

<table>
<thead>
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<th>R°0 (lbu)</th>
<th>L°x (lbu)</th>
<th>L°y (lbu)</th>
<th>L°z (lbu)</th>
<th>ε°f² (lbu²)</th>
<th>No. of simulations</th>
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<td>48</td>
<td>128</td>
<td>(7×10⁻³)–(3×10⁻³)</td>
<td>1000</td>
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<tr>
<td>10.0</td>
<td>72</td>
<td>72</td>
<td>180</td>
<td>(1×10⁻³)–(3×10⁻³)</td>
<td>500</td>
</tr>
<tr>
<td>14.0</td>
<td>96</td>
<td>96</td>
<td>256</td>
<td>1×10⁻⁴</td>
<td>200</td>
</tr>
<tr>
<td>18.0</td>
<td>122</td>
<td>122</td>
<td>324</td>
<td>1×10⁻⁴</td>
<td>200</td>
</tr>
<tr>
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<td>192</td>
<td>512</td>
<td>1×10⁻⁴</td>
<td>100</td>
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</table>
we study the probability density function (PDF) of the breakup time $t^*$. Overall, we observe that the shapes of the PDFs are similar to the breakup time distribution reported in a previous fluctuating thin-film study [17]. Comparing the “without-TN” protocol (first row) and the “with-TN” protocol (second row) for each thermal length, we see that the thermal fluctuations increase the probability for the ligament to break up sooner, which causes the peaks of the distribution of $t^*$ to move closer to the origin. Similarly, for both simulation protocols, we find a systematic speedup of the breakup time by increasing the amplitude of the thermal fluctuations, as shown by comparing PDFs on the same row at increasing thermal length (from left to right).

In Fig. 3(a), we show the PDFs of $t^*$ for both protocols normalized by their mean $\langle t^* \rangle$ and the standard deviation $\sigma_{t^*}$. The breakup time data for both protocols follow similar trends, which are well reproduced by the log-normal fit

$$f_{\log}(x) = \frac{1}{(x - x_0)\sigma_{\log}\sqrt{2\pi}} e^{-[\log(x - x_0) - \mu]_+^2/2\sigma_{\log}^2}$$

with $\sigma_{\log} = 0.32$, $\mu = 2.82$, and $x_0 = -2.97$. Also, Fig. 3(b) presents results for the average breakup time $\langle t^* \rangle$ as a function of the thermal length squared $\ell_T^2$ for both protocols. We observe a power-law-like behavior

$$\langle t^*/t_{\text{cap}} \rangle \sim \left(\ell_T^2\right)^{-0.07}$$

in both cases “with-TN” and “without-TN,” with the latter case always systematically above the former.

C. Droplet volumes

In this section, we study the droplet volumes after breakup. For this analysis, the droplet volumes are measured using the marching tetrahedra method [48] with PARAVIEW software and a Python interface. This ensures a very accurate measurement of the droplet volume, which is a key ingredient to differentiate the small changes induced by thermal fluctuations. In Fig. 4 we show the PDFs for the droplets volumes for the “with-TN” protocol at changing the thermal length $\ell_T$ for a fixed ligament radius $R_0$ [top row, panels (a)–(c)] or at changing the ligament radius $R_0$ for a fixed thermal length [bottom row, panels (d)–(f)]. We show only the volume distribution of the “mother” droplets (see Fig. 1): this is done to limit the range of $V$ and allow for a more insightful comparison. We have separately analyzed the PDFs of the satellite droplets, and the conclusions drawn for Fig. 4 are valid for them as well. From Fig. 4, we can see that when the thermal length increases, the PDFs develop larger standard deviation [panels (a)–(c)], and the same thing happens when we increase the ligament radius $R_0$ at fixed thermal length [panels (d)–(f)]. The shape of the PDFs and the dependence of the standard deviation $\sigma_V$ on both $\ell_T^2$ and $R_0$ are quantitatively summarized in Figs. 5(a) and 6(a). In particular, in Fig. 5(a) we report the standardized PDFs at changing the thermal lengths for fixed ligament radius for both protocols. We observe that the rescaled PDFs collapse well on the same master curve, independently of the thermal length and the simulation protocol. This indicates that the presence of randomness in the initial condition plays the major role in determining the shape of the distributions. A tendency toward a slight sub-Gaussian behavior is detectable for small (normalized) volume fluctuations, whereas larger fluctuations
FIG. 3. (a) PDF of normalized breakup time $\tilde{t}_f$ at changing thermal length $\ell_T$ for fixed ligament radius $R_0$ for both simulation protocols fitted with log-normal distribution. (b) Dimensionless mean breakup time $\langle \tilde{t}^* \rangle / t_{\text{cap}}$ as a function of thermal length squared for the two different simulation protocols used (see Fig. 1). Red and blue dotted lines are power fits for the dimensionless mean breakup time. Error bars are estimated from the standard deviation (see Fig. 2).

are associated with tails higher than Gaussian. The analysis of the standard deviation $\sigma_V$ [Fig. 5(a)] shows that in the “with-TN” protocol, the droplet polydispersity is enhanced by a factor around 40% with respect to what we obtain in the “without-TN” protocol. In both cases, however, signatures of a scaling law with exponent $\sim 0.14$ are obtained by fitting simulations data. Even though we are not able to provide an analytical explanation for the scaling law, we can confirm the observed scaling law by comparing with the sharp interface hydrodynamics in Sec. III D. In Fig. 6 we show the results of a similar analysis but at changing the ligament radius for a fixed thermal length for the physically relevant “with-TN” protocol. Again, we observe that the rescaled PDFs collapse well on the same master curve. Data are more scattered with respect to Fig. 5 due to the smaller number of simulations used to compute the PDFs. Overall, we notice that the PDFs reported in Figs. 5(a) and 6(a) display fatter tails with respect to a Gaussian distribution. If from one side one could say that the statistics

FIG. 4. PDFs for the droplet volumes after breakup based on the “with-TN” protocol (see Fig. 1) at changing the thermal length $\ell_T$ for fixed ligament radius $R_0 = 7.0$ [top row, panels (a)–(c)] or at changing the ligament radius $R_0$ for fixed thermal length squared $\ell_T^2 = 1 \times 10^{-4}$ [bottom row, panels (d)–(f)]. To make figures comparable at changing $R_0$, we have subtracted the average volume and rescaled by $R_0^3$.

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accumulated on the tails may be not enough to precisely quantify them, from the other side we will also present in Sec. III D data on sharp interface hydrodynamics supporting the view that those tails are definitively non-Gaussian. Regarding the standard deviation [Fig. 6(b)], we observe a scaling law close to 3, $\sigma_V \sim R_0^3$, that is what one would expect based on pure geometrical considerations. Regarding the connection between the distribution of volumes and breakup times, one could say that if the former is Gaussian and the latter log-normal, then an Arrhenius-like scenario could be invoked to connect the two, i.e., $t^* \sim e^{\Delta E/k_B T}$, with $\Delta E$ some energy contribution proportional to the volume variation. However, it must be noted that the volume distributions in Figs. 5(a) and 6(a) are slightly non-Gaussian, hence the above relation would imply also some departure from log-normality for the time distribution; additional data would be needed to clarify this issue.

**D. Comparison with sharp-interface hydrodynamics**

In the previous sections, we observed that rescaled PDFs of droplet volumes display a sub-Gaussian shape for small volume fluctuations, and higher tails for larger fluctuations [see Figs. 5(a) and 6(a)]; moreover, the standard deviation displays signatures of scaling laws in $\ell_T^2$ [see Fig. 5(b)]. To better reveal the origin of the shape of the PDFs and the scaling law for the standard deviation, we conducted additional numerical simulations with deterministic sharp interface hydrodynamics [45,49]. This comparison with sharp interface hydrodynamics can also elucidate the role of the diffuse interfaces that are inherent to the LB approach. We numerically considered an axisymmetric formulation of the lubrication equation of sharp interface hydrodynamics [45,49,50] using a finite-difference scheme with a total variation-diminishing method [50]. In this approach, the periodic axisymmetric ligament is placed at fixed ligament radius $R_0$ for both simulation protocols (“with-TN” and “without-TN”). (b) Standard deviation, $\sigma_V$, as a function of the thermal length squared $\ell_T^2$ at fixed ligament radius $R_0 = 7$ for both simulation protocols. Error bars are estimated from the standard deviation of different groups of the configurations.
along the $z$ axis and the whole evolution is described by its height, $h(z,t)$, and its axial velocity $v(z,t)$. The dimensionless lubrication equation becomes

$$\partial_t h^2 + \partial_z (h^2 v) = 0,$$

and

$$\partial_t v + v \partial_z v = -\partial_z P_{\text{lap}} + 3 \text{Oh} h^{-2} \partial_z (h^2 \partial_z v),$$

where $P_{\text{lap}}$ is the Laplace pressure that can be written as

$$P_{\text{lap}} = \left[ \frac{1}{h[1 + (\partial_z h)^2]} - \frac{\partial_z h}{[1 + (\partial_z h)^2]} \right].$$

To study the droplet size distributions in the sharp interface hydrodynamics approach, we imposed an initial perturbation on the radius in the form $R_0 + \epsilon \mathcal{Z}(z)$, with $\mathcal{Z}(z)$ a random Gaussian variable with unitary variance and zero mean, and $\epsilon$ a small number that is the analog of $\ell_T$ that we have used in the LB simulations. By varying the realization of the variable $\mathcal{Z}(z)$ in the initial conditions, we can compute the PDFs for droplet volumes. In other words, the hydrodynamic solver is the sharp interface counterpart of the “without-TN” protocol. The ensemble that we consider now is made of 4000 simulations, which is larger than what we used for the LB simulations (see Table I). This will allow us to see how much of the observed behavior for the LB simulations can depend on the statistics of thermal fluctuations during the dynamical process of the breakup. From one side, the thermal fluctuations allow the breakup process to speed up [20,22] and to obtain larger polydispersity; from the other side, the shape of the resulting PDF for droplet volume appears to be largely generated by a dynamical process that does not involve fluctuating hydrodynamics. The leading mechanism is that of a fastest-growing mode that is destabilized by the Plateau-Rayleigh instability, and other unstable modes (growing at a smaller rate) that provide—if initialized with random phases and amplitudes—an effective noise broadening the final distributions of droplet volumes. As a future perspective, there are various interesting issues to be investigated. For example, it could be an interesting challenging problem to predict the observed shape for the PDFs directly from sharp interface hydrodynamics, as well as the scaling laws for the standard deviations or the breakup time.

IV. CONCLUSIONS

We have used numerical simulations based on fluctuating multicomponent lattice Boltzmann (LB) models [27] to study the effects of thermal fluctuations on the breakup time of a liquid ligament and the associated polydispersity in droplets volumes after breakup. To quantitatively understand the role of thermal fluctuations during the dynamical process of the breakup, we have designed two different simulation protocols that allowed to evolve with or without thermal fluctuations a random initial condition realized over the ligament interface. From one side, the thermal fluctuations allow the breakup process to speed up [20,22] and to obtain larger polydispersity; from the other side, the shape of the resulting PDF for droplet volume appears to be largely generated by a dynamical process that does not involve fluctuating hydrodynamics. The leading mechanism is that of a fastest-growing mode that is destabilized by the Plateau-Rayleigh instability, and other unstable modes (growing at a smaller rate) that provide—if initialized with random phases and amplitudes—an effective noise broadening the final distributions of droplet volumes. As a future perspective, there are various interesting issues to be investigated. For example, it could be an interesting challenging problem to predict the observed shape for the PDFs directly from sharp interface hydrodynamics, as well as the scaling laws for the standard deviations or the breakup time. We also remark that the thermal lengths that we explored are quite small in comparison to the ligament radius. Hence, it

FIG. 7. (a) Comparison among the standardized PDFs of the droplet volumes following the two LB evolution protocols or the lubrication equations with initial random disturbance (see text for details). (b) Rescaled standard deviation ($\sigma_V / R_0^3$) at changing the rescaled thermal length squared ($\ell_T/R_0^2$) and rescaled Gaussian noise amplitude ($\epsilon/R_0^2$) for lubrication theory. Error bars are estimated from the standard deviation of different groups of the configurations.
could be a challenging computational task to extend our study in a range of parameters with larger thermal lengths [18,22].

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