Universality in fluid domain coarsening: The case of vapor-liquid transition

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Abstract – Domain growth during the kinetics of phase separation is studied following vapor-liquid transition in a single component Lennard-Jones fluid. Results are analyzed after appropriately mapping the continuum snapshots obtained from extensive molecular-dynamics simulations to a simple cubic lattice. For near-critical quench interconnected domain morphology is observed. A brief period of slow diffusive growth is followed by a linear viscous hydrodynamic growth that lasts for an extended period of time. This result is in contradiction with earlier reports of late-time growth exponent 1/2 that questions the uniqueness of the non-equilibrium universality for liquid-liquid and vapor-liquid transitions.

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Introduction. – Understanding non-equilibrium evolution during the phase separation in a system is of fundamental importance [1–4] and of much research interest, both theoretically and experimentally [5–22]. Upon quenching from a high-temperature homogeneous state to a temperature below the critical point, the system becomes unstable to fluctuations and starts phase separating with the formation and usually non-linear growth of particle-rich and particle-depleted domains. Such coarsening is a scaling phenomenon [23], e.g., the shape functions characterizing the morphology obey the scaling relations [1–3,20,23]

\[ C(r,t) \equiv \tilde{C}(r/\ell(t)), \]

\[ S(k,t) \equiv \ell(t)^d \tilde{S}(k\ell(t)), \]

\[ P(\ell_d,t) \equiv \ell(t)^{-1} \tilde{P}(\ell_d/\ell(t)), \]

where \( C(r,t), S(k,t) \) and \( P(\ell_d,t) \) are, respectively, the two-point equal-time correlation function, structure factor and domain size distribution function. In eqs. (1)–(3) \( \tilde{C}(x), \tilde{S}(y) \) and \( \tilde{P}(z) \) are the master functions independent of time \( (t) \)-dependent average domain size \( \ell(t) \) that typically grows in a power law manner,

\[ \ell(t) \sim t^\alpha. \]  

The growth exponent \( \alpha \), in eq. (4), depends upon the transport mechanism driving the phase separation. For diffusive dynamics, associating the domain growth with the chemical potential gradient as

\[ \frac{d\ell(t)}{dt} \sim |\nabla \mu| \sim \frac{\gamma}{\ell(t)^2}, \]

\( \gamma \) being the interfacial tension, one obtains \( \alpha = 1/3 \). This is referred to as the Lifshitz-Slyozov [24] (LS) law. While the LS law is the only expected behavior for phase-separating solid mixtures, one expects much faster growth, at large length scales, for fluids and polymers, where hydrodynamics is important. Compared to the kinetics in vapor-liquid phase separation, domain coarsening in binary fluids received much more attention [5–7,11,13,14,19,25–28], where following consensus for the behavior of late-stage growth [29–32] has been reached in d = 3. Considering a balance between the surface energy density \( (\gamma/\ell(t)) \) and the viscous stress \( (6\pi \eta v \ell/\ell(t)), v \ell \) being the interface velocity and \( \eta \) the shear viscosity) for an interconnected domain structure, one can write

\[ \frac{d\ell(t)}{dt} = v_\ell = \frac{\gamma}{\eta}. \]  

Solution of eq. (6) predicts a linear growth \( (\alpha = 1) \), a picture that holds for low Reynolds number. However, for \( \ell(t) \gg \ell_m = (\eta^2/(\rho \gamma)) \), \( \rho \) being the density, the inertial
length, the surface energy density is balanced by kinetic energy density \( \rho v^2 \ell \), so that
\[
\frac{d\ell(t)}{dt} \sim \frac{1}{\ell(t)^{1/2}},
\]
giving \( \alpha = 2/3 \). Note that \( \alpha = 1 \) is referred to as the viscous hydrodynamic growth and \( \alpha = 2/3 \) as inertial hydrodynamic growth. While crossover from diffusive to viscous hydrodynamic regime was observed in molecular-dynamics (MD) simulations [19,28] and experiments [33–35], both viscous and inertial growths were observed in lattice Boltzmann simulations [3].

While non-equilibrium universality for vapor-liquid phase separation is expected to be the same as the liquid-liquid one, rare MD simulations that exist [15,36–39] for the vapor-liquid transition report an exponent \( \alpha = 1/2 \) for late-time domain coarsening. Thus, even though our recent focus has turned to systems with realistic interactions and physical boundary conditions [5–11], our basic knowledge and understanding of segregation kinetics in bulk fluids still remains a challenge. In this letter, we present results from large-scale MD simulations for kinetics of vapor-liquid phase separation in a single-component Lennard-Jones (LJ) fluid. This work reports the first observation of viscous hydrodynamic growth via MD simulation, thus computationally confirming that the non-equilibrium universality class of vapor-liquid and liquid-liquid phase separation is indeed the same as in the case of equilibrium critical phenomena for both static and dynamic properties.

**Model and method.** – We consider a model where particles interact via
\[
V(r_{ij}) = U(r_{ij}) - U(r_c) - (r_{ij} - r_c) \left( \frac{dU(r_{ij})}{dr_{ij}} \right)_{r_{ij}=r_c},
\]
with
\[
U(r_{ij}) = 4\epsilon \left[ \left( \frac{\sigma}{r_{ij}} \right)^{12} - \left( \frac{\sigma}{r_{ij}} \right)^6 \right],
\]
being the standard LJ potential. In eqs. (8) and (9) \( r_{ij} \) is the scalar distance between the \( i \)-th and \( j \)-th particles, \( \sigma \) is the particle diameter and \( \epsilon \) is the interaction strength. The cut-off and shifting of the potential at \( r_c \) were used to facilitate faster computation. Further, introduction of the third term [40] on the right-hand side ensures that both the effective potential and force are continuous at \( r_c \) which was set to 2.5\( \sigma \). All particles were assigned equal mass \( m \). For the sake of convenience, below we set \( m, \sigma, \epsilon \) and \( k_B \) to unity.

The phase diagram of the model without the force correction term is known [41], the critical temperature \( T_c \) and critical density \( \rho_c \) being 1.085 ± 0.005 and 0.317 ± 0.006, respectively. It is understood, of course, that the introduction of force correction will somewhat change the value of \( T_c \), which, from our experience [42–44] with a similar model that was used to study phase separation in binary mixture, could be approximately 10% lower. In this work we have chosen the temperature low enough so that even without the complete knowledge of the critical parameters, we are safely below the co-existence curve, as is also clear from the nice phase-separating domain pattern shown in fig. 1. The interconnected nature of the morphology also ensures that our choice of overall density \( \rho \) is close to the critical value. In a future correspondence,
of course, we will undertake the understanding of the phase behavior of this model in conjunction with the presentation of detailed results on the phase separation kinetics at various temperatures and densities.

A total of 265421 particles were confined in a periodic 3-dimensional box of linear dimension $L = 96$. Thus, we worked with a density $\rho = 0.3$. The MD runs were performed using the Verlet velocity algorithm [40,45] with an integration time step of $\Delta t = 0.005\tau$, $\tau = (m\sigma^2/\epsilon)^{1/2} = 1$. The temperature $T$ was controlled by using a Nosé-Hoover thermostat (NHT) [45], which is known to preserve hydrodynamics well. At $t = 0$ homogeneous initial configurations, prepared by equilibrating the system at $T = 5$, were quenched to $T < T_c$. All the results were obtained by averaging over 5 independent runs at a quench temperature $T = 0.7$.

The left frames of fig. 1(a) are typical 3-dimensional evolution pictures from two different times where the dots represent a particle. They show a nice interconnected structure of segregating domains growing with time. The average density of liquid and vapor domains are, respectively, 0.72 and 0.02. A direct method to characterize the structure is to calculate the radial distribution function, $g(r)$, from these continuum configurations. However, we are interested in looking at $C(r, t)$, its Fourier transform $S(k, t)$ and $P(\ell_d, t)$ which have been more standard tools in the study of phase ordering dynamics. To facilitate the calculation of latter quantities, we map the continuum snapshots to a lattice one in such a way that a pure domain structure is obtained where all lattice sites inside a liquid domain have occupancy one while sites inside the vapor domain have occupancy zero. An appropriate renormalization procedure [12,16,19] is applied for this noise removal exercise. We further map this system into spin-1/2 Ising system where an occupied lattice site has spin value $S_i = 1$ and for a vacant site $S_i = -1$. The right frames in fig. 1(a) show such mapped configurations. In fig. 1(b) we demonstrate a similar exercise by showing cross-sectional views at two other times. Which may be difficult to appreciate from the 3d snapshots, these 2d pictures clearly show that no significant structural change has occurred during this transformation that can hamper the analysis. Also during the entire duration of study, we find that the composition of up and down spins remains conserved within a tolerable limit of 5% fluctuations.

From the mapped configurations, the correlation function $C(r)$ is calculated as

$$C(r, t) = \langle S_i S_j \rangle - \langle S_i \rangle^2,$$

where the angular brackets stand for statistical averaging. The average domain size $\ell(t)$ from the decay of $C(r, t)$ was estimated as

$$C(r = \ell(t), t) = h,$$

where $h$ was set to the first zero of $C(r, t)$. Other methods employed for the calculation of $\ell(t)$ are

$$\ell(t) = \frac{1}{\int dk kS(k)}$$

(12)

and

$$\ell(t) = \int d\ell_d \ell_d P(\ell_d, t).$$

(13)

In eqs. (12) and (13), it is assumed that $S(k, t)$ and $P(\ell_d, t)$ are appropriately normalized. In eq. (13), $\ell_d$ is obtained from the separation between two successive domain interfaces in x-, y- or z-directions.

**Results.** In fig. 2(a) we show the scaling behavior of $S(k, t)$ on a log-log plot in accordance with eq. (2), for which four different times were chosen, as indicated on the figure. While collapse of data from the earliest time with the other ones is not good, excellent data collapse for three later times is indicative that a scaling regime is reached.
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Fig. 3: Plot of $\ell(t)$ vs. $t$ on a log-scale. Note that we have presented $\ell(t)$ calculated both via eq. (12) and eq. (13). All data were obtained after removing the noise as described in the text. Here different lines correspond to growth laws $\ell(t) \sim t^{1/3}, t^{1/2}, t^{2/3}$ and $t$, as indicated.

The parallel nature of the tail region to the continuous line confirms consistency with the Porod law [46,47]. Figure 2(b) shows the scaling plot of $P(\ell_d, t)$ on a semi-log plot where the linear behavior of the tail region is consistent with an exponential decay [16,20]. Note that in both the cases, $\ell(t)$ used was obtained from eq. (13). The nice data collapse obtained in both the quantities by using $\ell(t)$ from a single measure speaks about the consistency of different methods.

Coming to the central objective of the paper, viz., quantifying the dynamics, in fig. 3 we present plot of $\ell(t)$, obtained via eqs. (12) and (13), vs. $t$ on a log scale. There different lines correspond to various growth laws in domain coarsening problem. While the early time data is more consistent with a diffusive growth ($\alpha = 1/3$), there appears a gradual crossover to a faster growth. The intermediate time data, of course, look consistent with $t^{1/2}$ as reported in previous works [15,36–39] and at a later time there seems to be a further crossover. However, due to large offset at the crossover region(s), which will be clearer from further analysis and discussions, we warn the reader not to take this conclusion seriously. For similar reasons, an apparently $t^{1/3}$ looking behavior during early time should also not be taken for granted. In fact, due to this and other practical difficulties, it is advisable not to reach a firm conclusion from looking at the behavior on a log-scale (which, of course, is a standard practice followed in the literature), unless one has results over several decades in time and length. Note that this plot again confirms equivalence of different methods for calculating $\ell(t)$. Thus, from now on, we will use data obtained only from eq. (13).

Fig. 4: Plot of $\ell(t)$, calculated using eq. (13), vs. $t$ on a linear scale. All data were obtained after removing the noise as described in the text. The solid line corresponds to the expected linear growth in the viscous hydrodynamic regime. Upper inset: plot of the growth exponent $\alpha$, as a function of $1/t_0$, obtained by fitting the data to the form (14) in the range [$t_0, 350$]. Lower inset: plot of system size $L$ vs. saturation time $t_s$. For MD results, the time is in $10^3$ LJ unit while for the MC simulation it is in unit of $10^6$ Monte Carlo steps.

Figure 4 shows a plot of $\ell(t)$ vs. $t$ on a linear scale where the data starting from $t \approx 100$ look quite linear all the way till the end. The deviation from this linear behavior at early time ($t < 100$) could be attributed to slower diffusive growth, as discussed above. A fitting to the form,

$$\ell(t) = \ell_0 + Bt^{\alpha}, \quad (14)$$

however, in the range $t \in [0,100]$ gives $\alpha = 0.66$. Since this diffusive regime is short lived and also accompanied by gradual crossover to a faster growth regime, it is in fact not practical to search for growth exponent $\alpha = 1/3$ in this region. On the other hand, a similar fitting to the range $t \in [100,350]$ gives $\alpha \simeq 0.9$, quite consistent with the predictions for viscous hydrodynamic growth. In the upper inset of fig. 4 we present a plot of the exponent $\alpha$ as a function of $1/t_0$ obtained by fitting the data to the form (14) in the range [$t_0, 350$]. This exercise conclusively says that the exponent at late time certainly is larger than $1/2$ which was reported earlier.

Note that for $t > 350$ we expect to see the finite-size effects for this system size. In the lower inset of fig. 4 we plot the system size $L$ as a function of time $t_s$ needed for the domain size to reach the saturation limit. The linear behavior of the data is again consistent with the viscous hydrodynamic growth. This should be compared with the corresponding result obtained from Kawasaki exchange Monte Carlo simulation of the Ising model [48] where, because of diffusive dynamics, $L$ should scale as

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t^{1/3}\) (dashed line) in accordance with the LS law. While we already have enough evidence for a linear behavior of the time dependence of \(\ell(t)\), for the sake of completeness and to further strengthen our claim we take the following route which, in addition to being a nice exercise, could be useful for other complex situations.

In fig. 5, we present a plot of the instantaneous exponent \(\alpha_i\) calculated as \([49]\)

\[
\alpha_i = \frac{1}{3} \left[ \frac{d \ln \ell(t)}{d \ln t} \right], \quad (15)
\]

vs. \(1/\ell(t)\). Due to the off-set \((\ell_0)\) at \(t=0\), as discussed in the context of fig. 3, \(\alpha_i\) is less than unity for the whole range. However, it seems to be approaching unity in a non-linear fashion. The dashed line there with an arrow at the end serves as a guide to the eyes. The lower part in this figure represents the corresponding result obtained from MD simulation using Andersen thermostat (AT) \([45]\). Due to the stochastic nature of this thermalization algorithm, it is expected that a diffusive growth will be seen for all time and lengthscales and indeed is true as will be clear from the following discussion.

Using (14) and (15), one obtains \([12]\)

\[
\alpha_i = \frac{1}{3} \left[ 1 - \frac{\ell_0}{\ell(t)} \right] ; \quad \alpha = \frac{1}{3}, \quad (16)
\]

where \(\ell_0\) corresponds to the characteristic length scale at the beginning of a scaling regime from which the time measurement starts and not necessarily the length immediately after quench. Thus, the observation of \(\alpha_i < 1/3\) for finite domain lengths is a mathematical artifact coming due to non-zero value of \(\ell_0\) that provides a non-zero slope \((-\ell_0/3)\) and does not necessarily mean that there is correction to scaling at early time, as previously understood \([49]\). In situations where one does not expect multiple scaling regimes (as is the case with AT) \(\ell_0\) can be nicely estimated from a data-collapse experiment in finite-size scaling analysis \([12,48]\). Our experience with Ising model, however, tells us that the system enters the scaling regime rather early for a critical quench \([12]\) and so, here we take \(\ell_0\) to be the length at the time of quench. The consistency of the AT data with eq. (16) (dashed line) confirms this to be a good approximation.

Coming to the point of hydrodynamic preserving capability of NHT, of course, much better thermostats are available these days. However, from the difference seen in fig. 5 for the results coming from NHT and AT, it is clear that NHT is preserving it rather well. Thus, the validity of the methodology adopted in this work is justified. But the situation is more complex with NHT, as far as playing with \(\ell_0\) is concerned, for the crossover to hydrodynamic regime. However, considering that the crossover, at time \(t_c\), under discussion is to a linear regime, confirmation about it as well as a linear behavior for the growth of \(\ell(t)\) could be obtained by using the following simpler method.

To start with, we assume that the growth obeys a power law behavior as a function of the shifted time \(t' = t - t_c\) as

\[
\ell'(t') = \ell(t) - \ell(t_c) = At'^{\alpha} \quad (17)
\]

and calculate the instantaneous exponent

\[
\alpha_i = \frac{d \ln \ell'(t')}{d \ln t'} \quad (18)
\]

Equation (17) is invariant under an arbitrary choice of \(t_c\) if we are in a linear growth regime and one should obtain a constant value \(\alpha = 1\) for all choices of \(t_c\) in the post crossover regime. In the inset of fig. 5, we present \(\alpha_i\) for two different choices of \(t_c\). For \(t_c > 110\), \(\alpha_i\) oscillates around the mean value 1 which is consistent with the viscous hydrodynamic regime. This oscillation (which could as well be appreciated from main part of fig. 5) is also observed in other studies \([12,25]\). In a finite system, at later time only a few domains of comparable size exist which are separated from each other by large distances. Thus, as time increases, it takes longer time for them to merge which causes the exponent to come down. Finally merging of two huge domains after a long interval suddenly enhances the value of the exponent.

**Conclusion.** – We have presented results for the kinetics of vapor-liquid phase separation from the molecular-dynamics (MD) simulation of a simple Lennard-Jones fluid using both Andersen (AT) and Nosé-Hoover (NHT) thermostats. It is observed that the NHT is reasonably useful for studying hydrodynamic effects in the fluid phase separation. A brief period of diffusive coarsening was followed by a linear viscous hydrodynamic growth. Our results are in contradiction with few previous MD studies which reported an exponent much less than unity; however, they
are consistent with the results of binary fluid phase separation. One requires a much larger system size to observe the inertial hydrodynamic growth. This we leave out for a future exercise, in addition to the study of coarsening of liquid droplets for off-critical quench. For the latter problem one expects a single asymptotic exponent 1/3 from the droplet diffusion-coagulation mechanism [23,50]. Also, as discussed, MD with AT and a Monte Carlo simulation of the same system will provide a single diffusive 1/3 growth for all compositions. For off-critical composition the latter exercise should be able to provide information on the curvature-dependent interfacial tension which is expected to bring in early-time correction in the LS law. This will certainly be interesting to compare with the corresponding results obtained from equilibrium studies [51].

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