Dynamic van der Waals theory

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We present a dynamic van der Waals theory starting with entropy and energy functional with gradient contributions. The resultant hydrodynamic equations contain the stress arising from the density gradient. It provides a general scheme of two-phase hydrodynamics involving the gas-liquid transition in nonuniform temperature. Some complex hydrodynamic processes with evaporation and condensation are examined numerically. They are (i) adiabatically induced spinodal decomposition, (ii) piston effect with a bubble in liquid, (iii) temperature and velocity profiles around a droplet in heat flow, (iv) efficient heat transport at small liquid densities (the mechanism of heat pipes), (v) boiling in gravity with continuous bubble formation and rising, and (vi) spreading and evaporation of liquid on a heated boundary wall.

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I. INTRODUCTION

The van der Waals theory provides a simple description of gas-liquid phase transitions in one-component fluids [1]. It is an equilibrium mean-field theory for hard sphere particles with long-range attractive interaction [2]. Moreover, in his pioneering paper in 1893, van der Waals introduced a gradient term in the Helmholtz free energy density to describe a gas-liquid interface. Such a gradient term began to be widely used in statistical mechanics of nonuniform states, since seminal papers by Ginzburg and Landau for type-I superconductors [3] and by Cahn and Hilliard for binary alloys [4].

In most phase transition theories, including those of dynamics, the temperature $T$ is a given parameter independent of space [5,6]. The Ginzburg-Landau theory is based on a free energy functional with homogeneous $T$. It goes without saying that there are a variety of situations, where phase transitions occur in inhomogeneous $T$ or in heat flow. In fluid systems, wetting dynamics [7–11], boiling processes [12–14], and droplet motion [15,16] are strongly influenced by applied heat flux. Rayleigh-Bénard convection is very complex in liquid crystals with first-order phase transition [17] and in binary fluid mixtures with phase separation [18–20]. In He4, the superfluid transition can be non-linearly influenced by heat flux, where a sharp interface is produced separating superfluid and normal fluid [21,22]. Understanding such a problem is very difficult. We need to construct a dynamical model first to study nonequilibrium effects, where phase transitions and hydrodynamics are inseparably coupled.

In 1901, Korteweg proposed hydrodynamic equations for binary fluid mixtures including the stress induced by composition gradients [23]. Diffuse interfaces between two-phases then arise in solutions of the hydrodynamic equations in the presence of the gradient stress [24–26]. In this line, a number of two-dimensional simulations of diffuse-interface models have been presented to describe two-phase hydrodynamics in one-component fluids [6,14] and in binary mixtures [19,20,26–28]. One-dimensional numerical solutions are themselves highly nontrivial in heat flow [29,30]. Note that diffuse-interface models or phase field models have been used in numerical analysis of dendrite instability in crystal growth [31–33]. As another line of research, critical dynamics in classical fluids was studied with inclusion of the gradient contributions of the mass density and the composition, which are relevant even in one-phase states with growing of the correlation length near the critical point [5,6,34–38]. The so-called model H for near-critical binary mixtures was originally devised to describe dynamics of the thermal fluctuations [5], but has also been used to describe phase separation processes [39] and steady states under shear flow [6,40]. We mention Kawasaki’s hydrodynamic equations for van der Waals fluids with long-range interaction [36], where the stress is of a nonlocal convolution form but reduces to the well-known form in critical dynamics in the gradient approximation [41]. For He4, on the other hand, a set of nonlinear hydrodynamic equations with the gradient contributions was established in an early period [42], whose simplified version near the superfluid transition was later used in critical dynamics [5,6] and in studying nonlinear effects of heat flow [21].

Recently, we presented a dynamic van der Waals theory on the basis of entropy and energy functionals containing the gradient contributions [15]. We then constructed hydrodynamic equations with the gradient stress and numerically solved them to examine droplet motion in heat flow. One of our findings is that the temperature becomes homogeneous within a droplet under applied heat flux without gravity. Latent heat transport is so efficient such that the interface region is nearly on the coexistence curve $T=T_{cx}(p)$ even in nonequilibrium, where the pressure $p$ is uniform outside the droplet. This picture was confirmed in a subsequent hydrodynamic theory [16]. As a result, the Marangoni effect arising from inhomogeneous surface tension is not operative in one-component (pure) fluids, while it gives rise to much faster fluid and interface motions in mixtures even at very small impurity concentrations [43].

In this paper, we will present a general scheme of the dynamic van der Waals theory in one-component fluids in a general form and give numerical results of some fundamental but complicated processes with evaporation and condensation. Section II will be a theoretical part, while Sec. III will present some numerical illustrations of such processes.
II. THEORY

A. van der Waals theory

We summarize the main results of the van der Waals theory \[1,6\], which will be needed in constructing our theory. As a function of the number density \( n \) and the temperature \( T \), the Helmholtz free density \( f(n, T) \) for monoatomic molecules is written as

\[
f = k_BTn[\ln(\lambda_0^d n) - 1 - \ln(1 - v_0 n)] - \epsilon v_0 n^2,
\]

where \( v_0 = a/d \) is the molecular volume and \( \epsilon \) is the magnitude of the attractive potential with \( d \) being the space dimensionality. The molecular radius is given by \( a = v_0^{1/d} \). The \( \lambda_0^d = h/(2\pi/mT)^{1/2} \) is the thermal de Broglie length with \( m \) being the molecular mass. As functions of \( n \) and \( T \), the internal energy density \( e \), the entropy \( s \) per particle, and the pressure \( p \) are given by \[44\]

\[
e = nk_BT/2 - \epsilon v_0 n^2, \tag{2.2}
\]

\[
s = -k_B \ln(\lambda_0^d n/(1 - v_0 n)) + k_B(d + 2)/2, \tag{2.3}
\]

\[
p = nk_BT/(1 - v_0 n) - \epsilon v_0 n^2. \tag{2.4}
\]

In this paper, we will make use of the expressions for the sound velocity \( c = [(\partial p/\partial n)/m]^{1/2} \) and the specific heat ratio \( \gamma = C_p/C_v \),

\[
c = (k_BT/m)^{1/2}(1 + 2ld - T/\gamma T), \tag{2.5}
\]

\[
\gamma = 1 + 2ld(1 - T/\gamma T). \tag{2.6}
\]

where \( C_V = nk_BT/2 \) is the constant-volume specific heat and \( C_p \) is the isobaric specific heat per volume. We introduce the spinodal temperature,

\[
T_s = 2\epsilon v_0 n/(1 - v_0 n)^2/k_B. \tag{2.7}
\]

Maximization of \( T_s(n) \) as a function of \( n \) yields the critical temperature \( T_c = 8\epsilon/77k_B \) and the critical density \( n_c = 1/3\epsilon v_0 \).

As mentioned in Introduction, van der Waals introduced the gradient free energy density,

\[
f_{\text{gra}} = \frac{1}{2}M|\nabla n|^2, \tag{2.8}
\]

We consider the equilibrium interface density profile \( n(x) \) varying along the \( x \) axis. The chemical potential per particle \( \mu(n, T) = (\partial f/\partial n)_T \) changes as

\[
\mu - \mu_{\text{cx}} = \frac{M d^2n}{2 dx^2} + \frac{d M dn}{dx 2 dx}, \tag{2.9}
\]

where \( \mu_{\text{cx}}(T) \) is the chemical potential on the coexistence curve. From \( dp = nd\mu \) at constant \( T \), the van der Waals pressure \( p(n, T) \) in Eq. \[2.4\] is expressed as \[6\]

\[
p - p_{\text{cx}} = \frac{M d^2n}{2 dx^2} + \frac{d M dn}{dx 2 dx} - M \left( \frac{dn}{dx} \right)^2, \tag{2.10}
\]

where \( p_{\text{cx}}(T) \) is the pressure on the coexistence curve. In Sec. II D, a stress tensor \( \Pi_{ij} \) including gradient contributions will be introduced, in terms of which Eq. \[2.10\] is equivalent to \( \Pi_{xx} = \rho_{xx} \) [see discussions below Eq. \[2.50\]]. The surface tension \( \gamma \) is expressed as \[6,45,46\]

\[
\gamma = \int_{-\infty}^{\infty} dx M(dn/dx)^2. \tag{2.11}
\]

We note that Eqs. \[2.9\]–\[2.11\] hold even if \( M \) depends on \( n \) as \( M = M(n) \) (while \( M \) was a constant in the original theory \[1\]).

B. Gradient entropy and energy

We generalize the van der Waals theory by including gradient contributions to the entropy and the internal energy as

\[
S_b = \int dr \left[ n_s(n, e) - \frac{1}{2}C|\nabla n|^2 \right], \tag{2.12}
\]

\[
\mathcal{E}_b = \int dr \left[ e + \frac{1}{2}K|\nabla n|^2 \right], \tag{2.13}
\]

where the space integrals are within the container of the fluid. The entropy and energy contributions at the boundary walls will be introduced in Eq. \[2.28\] below. The entropy density and the internal energy density including the gradient contributions are thus written as

\[
\dot{S} = n_s - \frac{1}{2}C|\nabla n|^2, \tag{2.14}
\]

\[
\dot{e} = e + \frac{1}{2}K|\nabla n|^2. \tag{2.15}
\]

The gradient terms represent a decrease of the entropy and an increase of the energy due to inhomogeneity of \( n \). They are particularly important in the interface region. For simplicity, we neglect the gradient terms proportional to \( (\nabla n) \cdot \nabla e \) and \( (\nabla e)^2 \). The gravitational energy is also neglected, but its inclusion is trivial [see discussions around Eq. \[2.51\]]. In this work, in constructing a general theory, we allow that \( C \) and \( K \) depend on \( n \)

\[
C = C(n), \quad K = K(n). \tag{2.16}
\]

However, in our simulations in the next section, we will set \( K = 0 \) and assume that \( C \) is independent of \( n \).

We then define the local temperature \( T = T(n, e) \) by

\[
\frac{1}{T} = \left( \frac{\partial S_b}{\partial e} \right)_n = n \left( \frac{\partial e}{\partial n} \right)_n, \tag{2.17}
\]

where \( n \) is fixed in the derivatives. This definition of \( T \) is analogous to that in a microcanonical ensemble. We will use it even for inhomogeneous \( n \) and \( e \) in nonequilibrium. We also define a generalized chemical potential \( \tilde{\mu} \) per particle including the gradient contributions by

\[
\dot{\tilde{\mu}} = -T \left( \frac{\partial S_b}{\partial n} \right)_e = \mu - T \nabla \cdot \frac{M}{T} \nabla n + \frac{M'}{2} |\nabla n|^2, \tag{2.18}
\]

where the internal energy density \( \dot{\tilde{e}} \) in Eq. \[2.15\] is fixed so we set \( \Delta e = -\partial K|\nabla n|^2/2 \) in the functional derivative. The
\( \mu = (e + p)/n - Ts \) is the usual chemical potential per particle and
\[
M = CT + K. \tag{2.19}
\]
Obviously, \( M \) is the coefficient of the gradient term in the Helmholtz free energy defined in equilibrium (see the next section). Under Eq. (2.16), \( M = M(n, T) \) depends on \( n \) and \( T \), so \( M' = \partial M/\partial n \) in Eq. (2.18) is its derivative with respect to \( n \). Now, regarding \( S_b \) as a functional of \( n \) and \( \hat{e} \), we consider small changes \( \hat{e} \) of \( n \) and \( \hat{e} \) which yield an incremental change of \( S_b \). Using Eq. (2.18) and \( \hat{e} = \hat{e} - 8(K|\nabla n|^2)/2 \), we obtain
\[
\delta S_b = \int dr \left( \frac{1}{T} \hat{e} - \frac{\hat{S}}{T} \right) - \int da M_T (\mathbf{v} \cdot \nabla n) \delta n. \tag{2.20}
\]
The second term is the surface integral, where \( da \) is the surface element and \( \mathbf{v} \) is the outward normal unit vector at the surface.

We remark on the following two papers. Fixman proposed the energy and entropy functionals to include the velocity field in the solidification problem [33]. Anderson et al. started with energy and entropy functionals to include the velocity field in the solidification problem [33].

C. Equilibrium conditions

In equilibrium, we maximize \( S_b \) at a fixed particle number \( N = \int d\hat{e} n \) and a fixed energy \( \mathcal{E}_b = \int d\hat{e} e \) for a fluid confined in a cell. First, we are interested in the bulk equilibrium. To this end, we introduce
\[
W = S_b/k_b + \nu N - \beta \mathcal{E}_b, \tag{2.21}
\]
where \( \nu \) and \( \beta \) are the Lagrange multipliers. The maximization condition \( (\partial W/\partial \delta n)_e = 0 \) with respect to \( e \) at fixed \( n \) yields the homogeneity of the temperature,
\[
T = 1/k_b \beta \tag{2.22}
\]
and then
\[
W = (-\mathcal{F} + k_b T \nu N)/k_b T \tag{2.23}
\]
becomes a functional of \( n \). We introduce the Helmholtz free energy functional,
\[
\mathcal{F} = \mathcal{E}_b - TS_b = \int dr \left( f(n, T) + \frac{M}{2} |\nabla n|^2 \right), \tag{2.24}
\]
where the integrand is \( \hat{e} - TS_b \), so \( f = e - Ts + M \) and \( M \) is defined by Eq. (2.19). In the Ginzburg-Landau theory, we furthermore minimize \( W \) with respect to \( n \) to obtain
\[
\hat{\mu} = k_b T \nu = \text{const}, \tag{2.25}
\]
where \( \hat{\mu} \) is defined by Eq. (2.18). In the thermodynamic limit we find \( W = p V/k_b T \) where \( p \) is the pressure and \( V \) is the total volume of the fluid.

In a two-phase state with a planar interface at \( x = 0 \), the surface tension \( \gamma \) is expressed as Eq. (2.11). Here we introduce the grand potential density,
\[
g(n, T) = \int \left( f(n, T) - \mu_c n + p_c \right), \tag{2.26}
\]
where \( \mu_c \) and \( p_c \) are the values of \( \mu \) and \( p \) on the coexistence curve. It is nonvanishing in the interface region, where \( n = n(x) \) changes from a gas density \( n_g \) to a liquid density \( n_l \) as a function of \( x \). From Eq. (2.25) it holds the relation
\[
\gamma(T) = \int_{n_g}^{n_l} dn \left( \frac{2M g(n, T)}{\gamma} \right)^{1/2}. \tag{2.27}
\]
This expression holds even when \( M \) depends on \( n \) [6].

In real fluids we need to consider the boundary condition on the surface of the container. For simplicity, we consider the surface entropy and energy of the forms
\[
S_s = \int da \sigma_s(n), \quad \mathcal{E}_s = \int da \mathcal{E}_s(n), \tag{2.28}
\]
where \( da \) is the surface element and the areal densities \( \sigma_s \) and \( \mathcal{E}_s \) are assumed to depend only on the number density \( n \) on the surface. We need to maximize the total entropy \( S_{tot} = S_b + S_s \) under fixed \( N \) and \( \mathcal{E}_{tot} = \mathcal{E}_b + \mathcal{E}_s \). To this end, we introduce
\[
W_{tot} = W + \int da (\sigma_s - \beta \mathcal{E}_s), \tag{2.29}
\]
where \( \beta \) is common to that in Eq. (2.22). Maximization of the surface part of \( W_{tot} \) additionally yields
\[
M \mathbf{v} \cdot \nabla n + (\partial f/\partial \delta n)_T = 0, \tag{2.30}
\]
as the surface boundary condition. Here we use Eq. (2.20) and define the Helmholtz surface free energy density,
\[
f_s(n, T) = e_s - T \sigma_s. \tag{2.31}
\]
In equilibrium, the temperature is homogeneous and we should minimize the total Helmholtz free energy,
\[
\mathcal{F}_{tot} = \mathcal{F} + \int da f_s(n, T), \tag{2.32}
\]
in order to calculate the surface density profile [7,48]. The surface free energy is usually expanded to second order in \( n - n_c \) as
\[
f_s = -a_s (n - n_c) + b_s (n - n_c)^2/2, \tag{2.33}
\]
where \( a_s \) and \( b_s \) are constants. Then the boundary condition (2.30) is written as
\[
M \mathbf{v} \cdot \nabla n - a_s + b_s (n - n_c) = 0. \tag{2.34}
\]

D. Generalized hydrodynamic equations

We propose hydrodynamic equations taking account of the gradient entropy and energy. They are of the same forms as those of compressible fluids in the literature [47] except that the stress tensor contains gradient contributions. The guiding principle of their derivation is the non-negative definiteness of the entropy production rate in the bulk region. In the previous theories [24,26,28] the gradient entropy was not
assumed and $M=K$ in Eq. (2.19). We suppose that a fluid is in a solid container with controllable boundary temperatures and the velocity field $v$ vanishes on the boundary. We hereafter include a gravity $g$ applied in the downward direction along the $z$ axis.

First, the number density $n$ obeys

$$\frac{\partial n}{\partial t} = -\nabla \cdot (nv). \tag{2.35}$$

The mass density is defined by $\rho = mn$, where $m$ is the particle mass. Second, the momentum density $\rho v$ obeys

$$\frac{\partial}{\partial t} (\rho v) = -\nabla \cdot (\rho vv) - \nabla \cdot (\tilde{\Pi} - \tilde{\sigma}) - \rho \dot{e}_v, \tag{2.36}$$

where the last term arises from the gravity with $e_v$ being the (upward) unit vector along the $z$ axis. The $\tilde{\Pi} = \{\Pi_{ij}\}$ is the reversible stress tensor containing the gradient contributions, which is invariant with respect to time reversal $v \rightarrow -v$. The $\tilde{\sigma} = \{\sigma_{ij}\}$ is the dissipative stress tensor of the form,

$$\sigma_{ij} = \eta (\nabla v_j + \nabla v_i) + (\zeta - 2\eta d) \delta_{ij} \nabla \cdot v \tag{2.37}$$

with $\nabla = \partial / \partial x$. The $\eta$ and $\zeta$ are the shear and bulk viscosities, respectively. Third, including the kinetic part, we define the (total) energy density,

$$e_T = \dot{e} + \rho v^2 / 2. \tag{2.38}$$

We assume that the energy current is of the usual form in terms of the stress tensor $\tilde{\Pi} - \tilde{\sigma}$ and the thermal conductivity $\lambda$. Then $e_T$ is governed by [47]

$$\frac{\partial}{\partial t} e_T = -\nabla \cdot [e_T v + (\tilde{\Pi} - \tilde{\sigma}) \cdot v] + \nabla \cdot (\lambda \nabla T) - g \rho v_z. \tag{2.39}$$

Equivalently, we may rewrite the above energy equation in terms of the internal energy density $\dot{e}$ using Eqs. (2.35) and (2.36) as

$$\frac{\partial}{\partial t} \dot{e} = -\nabla \cdot (\dot{e} v) - \tilde{\Pi} \nabla v + \dot{e}_v + \nabla \cdot (\lambda \nabla T), \tag{2.40}$$

where there is no term proportional to $g$. On the right-hand side of Eq. (2.40), the second term represents $-\sum_j \Pi_{ij} \nabla v_j$ and $\dot{e}_v = \sum_i \sigma_{ij} \nabla v_j$ is the viscous heat production rate. Use of Eq. (2.37) gives

$$\dot{e}_v = \frac{1}{2} \sum_j \left[ \nabla v_j + \nabla v_j - 2 \xi (\delta_{ij} \nabla \cdot v \epsilon T) \right]^2 + \xi (\nabla \cdot v)^2 \geq 0. \tag{2.41}$$

From Eq. (2.20) the time derivative of $S_b$ becomes composed of bulk and surface contributions as

$$\frac{dS_b}{dt} = \int dr \left[ \frac{1}{T} \frac{\partial T}{\partial t} - \frac{\dot{\mu}}{T} \frac{\partial T}{\partial t} \right] - \int da (v \cdot \nabla n) \frac{M \partial n}{T \partial t}. \tag{2.42}$$

From our hydrodynamic equations the first bulk term becomes

$$\left( \frac{dS_b}{dt} \right)_{\text{bulk}} = \int dr \left[ v \cdot \left( \frac{\tilde{\Pi}}{T} \frac{\partial T}{\partial t} + \dot{\epsilon}_v \frac{1}{T} - n \frac{\dot{\mu}}{T} \right) + \frac{1}{T} (\nabla \cdot \lambda \nabla T + \dot{e}_v) \right]. \tag{2.43}$$

If there is no heat flow from outside, the above quantity should be non-negative definite. We thus require

$$\sum_j \nabla \cdot \left( \frac{1}{T} \Pi_{ij} \right) = - \dot{e}_v + \frac{n}{T} \frac{\partial T}{\partial t} + \frac{\partial \mu}{\partial T}, \tag{2.44}$$

Under Eq. (2.44) we obtain \( dS_b / dt \)_{bulk} = $\int dr (\dot{\epsilon}_v + \dot{\epsilon}_\theta) / T \geq 0$ without heat input from the boundary [47], where the thermal heat production rate is

$$\dot{\epsilon}_\theta = \lambda (\nabla T)^2 / T \geq 0. \tag{2.45}$$

Now we seek the symmetric tensor $\Pi_{ij}$ which satisfies Eq. (2.44). Notice the thermodynamic identity,

$$d(p/T) = -e d(1/T) + n d(\mu/T), \tag{2.46}$$

following from the Gibbs-Duhem relation $d\mu = -sdT + dp/n$ [6], where $(\cdot \cdot \cdot)$ denotes an infinitesimal change. If the gradient stress is neglected, we have $\Pi_{ij} = n \delta_{ij}$ as should be the case, where $p = p(n,T)$ is the van der Waals pressure in Eq. (2.4). If it is included, some calculations yield

$$\Pi_{ij} = (p + p_1) \delta_{ij} + M (\nabla v) (\nabla v), \tag{2.47}$$

where $p_1$ is a diagonal gradient part,

$$p_1 = (nM' - \frac{|\nabla n|^2}{2} - Mn^2 - Tn (\nabla n) \cdot \nabla M / T \neq 0. \tag{2.48}$$

In the first line of Eq. (2.48), the last term does not exist in the previous theories [26]. The second line indicates that Eq. (2.48) is a generalization of the thermodynamic identity $p = n\mu - e - T n^2$. In Appendix A, we will present another derivation of the above expression for $\tilde{\Pi}$ using Eqs. (2.35) and (2.40). With the above form of $\tilde{\Pi}$, the total entropy density $\hat{S}$ in Eq. (2.14) obeys

$$\frac{\partial}{\partial t} \hat{S} = -\nabla \cdot \left[ \hat{S} v - \frac{M}{T} n (\nabla \cdot v) \nabla n - \frac{\lambda}{T} \nabla T \right] + (\dot{\epsilon}_v + \dot{\epsilon}_\theta) / T. \tag{2.49}$$

The additional entropy flux $-T^{-1} M (\nabla \cdot v) \nabla n$ is reversible and is nonexistent in the usual hydrodynamics [47]. The last term on the right-hand side is the usual entropy production rate per unit volume.

We make some further comments. (i) Gouin [24] and Anderson et al. [26] also started with the entropy production rate with $M=K=\text{const}$ [see Eq. (10) of Ref. [24] and Eq. (15) of Ref. [26]], but without assuming Eqs. (2.39) and (2.44) they included a reversible contribution (in our notation),

$$J_{GA} = M (\nabla \cdot v) n \nabla n, \tag{2.50}$$

in the energy flux. We do not assume such a new energy flux in Eq. (2.39); instead, we have the last term \( \approx \nabla (M/T) \) in
$p_1$ in Eq. (2.48) and the additional entropy flux written as $T^{-1}J_{GA}$ in Eq. (2.49) [49]. (ii) When $M$ and $T$ are constants independent of space, our expression for $\Pi$ reduces to that used near the critical point [6,37]. The off-diagonal components $M\nabla_i p_i (i \neq j)$ give rise to the critical anomaly of the shear viscosity [5,6,34] and the surface tension contribution to the shear viscosity in sheared two-phase states [40]. (iii) We consider the one-dimensional equilibrium case, where all the quantities vary along the $z$ axis. We then obtain $\mu=0$, $T=$const, and the equilibrium relation,

$$
\frac{d}{dz}\Pi_{zz} = -g\rho,
$$

(2.51)

which leads to Eq. (2.10) in two phase coexistence (by changing $z$ to $x$ and setting $g=0$). Using Eq. (2.44) we also obtain the equilibrium chemical potential,

$$
\hat{\mu} + mgz = \text{const},
$$

(2.52)

which is a generalization of Eqs. (2.9) and (2.25) in gravity. In addition, we find $\Pi_{zz}-\Pi_{xx}=\Pi_{zz}-\Pi_{yy}=M(dn/dz)^2$. This is consistent with the surface stress tensor $\gamma(n_i n_j - \delta_{ij})\delta_x$ in the thin interface limit [50], where $\nu$ is the normal unit vector and $\delta_x$ is the $\delta$ function on the surface.

As the simplest boundary condition on the container surface, we may use the equilibrium condition (2.30) even in nonequilibrium and even when the temperature is not homogeneous on the boundary walls. This is justified when the equilibration at the boundary is much faster than that in the bulk. Then, under Eqs. (2.30) and (2.44) the total entropy $S_{tot}=S_B+S_s$ changes in time as

\[\frac{d}{dt}S_{tot} \leq 0.\]
the vertical direction, with 0 boundary condition for the density, we assume Eq. 2.20

\[ v = 202 \]

for the form

\[ x = \frac{L}{2} \]

\[ t / t_0 = 200 \]

Fig. 4. \((T - T_b) / (T_b - T_c)\), \(p / p_c\), and \(v_0 n\) vs \(y / L\) at \(t / t_0 = 200\) and at \(x = L / 2\), with \(T_b = 0.895 T_c\) and \(T_c = 0.875 T_c\), in the same run as in Fig. 3. The thermal diffusion layers are reaching at the bubble due to the small system size.

\[ \frac{dS_{tot}}{dt} = \int dr \frac{\dot{e}_\theta + \dot{e}_v}{T} + \int da \frac{\boldsymbol{v} \cdot \nabla T + \dot{e}_s}{T}, \quad (2.53) \]

where \(\dot{e}_s = \partial e_s / \partial t = (\partial e_s / \partial n) \partial n / \partial t\). On the right-hand side, the first term is the entropy production rate in the bulk, while the second term is due to the heat flux from outside and surface energy change. Note that the relation (2.53) holds even when the temperature is inhomogeneous on the boundary walls.

III. NUMERICAL RESULTS

A. Method

We integrated the hydrodynamic equations (2.35), (2.36), and (2.39) in two dimensions on a 202x202 lattice. We assume that \(C\) in Eq. (2.12) is a constant independent of \(n\) and \(K\) in Eq. (2.13) vanishes. The mesh size of our simulation cell is chosen to be

\[ \ell = (C/2k_B v_0)^{1/2}. \]

The interface thickness in two phase coexistence is of order \(\ell\) far from the critical point and is given by \([8T_c/27(T_c - T)]^{1/2}\) close to the critical point. The system length is \(L = 202 \ell\), so our system is very small. In the following figures, the x axis is in the horizontal direction, while the y axis is in the vertical direction, with \(0 \leq x \leq L\) and \(0 \leq y \leq L\). The velocity field \(\boldsymbol{v}\) is assumed to vanish on the walls; then, the average number density \(\langle n \rangle = \int d\boldsymbol{r} n / V\) is a conserved quantity. Hereafter \(\langle \cdots \rangle\) denotes taking the space average. As the boundary condition for the density, we assume Eq. (2.34) in the form

\[ 2\sqrt{2} \ell \nu \cdot \nabla n = n - 5n_c / 2, \]

on all the walls. This yields \(a_s = \ell k_B T / 2\sqrt{2}\) and \(b_s = 2v_0 \mu_0\) in Eq. (2.33). Then the boundary walls are wetted by liquid in equilibrium. The boundary condition for the temperature is as follows. In Figs. 1 and 2, we assume a common boundary temperature on all the walls. In Figs. 3–15, the bottom boundary temperature \(T_b\) at \(y = 0\) and the top boundary temperature \(T_c\) at \(y = L\) will be taken to be different, while the side walls are thermally insulating or \(\boldsymbol{v} \cdot \nabla T = 0\) at \(x = 0\) and \(L\). We use the words “top” and “bottom” even in the absence of gravity.

The transport coefficients strongly depend on the density \([30, 51]\). That is, the shear viscosity and the thermal conductivity of liquid are usually much larger than those in gas except in the vicinity of the critical point. For simplicity, we assume them to be proportional to \(n\) as

\[ \eta = \xi = \nu_0 mn, \]

\[ \lambda = k_B \nu_0 n. \]

Here the kinematic viscosity \(\nu_0 = \eta / \rho\) is assumed to be independent of \(n\). For example, the liquid and gas densities are \(n_l = 1.70 n_c\) and \(n_g = 0.27 n_c\) in two phase coexistence at \(T\).
The right-hand side gives the deformation rate of a droplet due to the surface tension if $Q=q^2 \sim 2\pi \ell / R$ with $R$ being the droplet radius.
B. Phase separation by adiabatic expansion at $g = 0$

First, we investigate phase separation induced by the piston effect [52] taking place upon lowering of the boundary temperature. In such situations, due to contraction of thermal boundary layers, the interior can be expanded into metastable or unstable states adiabatically [6]. The piston effect is intensified near the critical point owing to the divergence of isobaric thermal expansion. With this method in near-critical one-component fluids, nucleation experiments were performed by Moldover et al. [53], while spinodal decomposition experiments were realized in space by Beysens et al. [54]. Recently Miura et al. [55] measured adiabatic density changes of order $10^{-7}n_c$ in the interior caused by sound pulses in supercritical states. In this paper, however, we present an example of adiabatic spinodal decomposition far below the critical point caused by the growth of a wetting layer.

We prepared a supercritical state at $T = 1.1T_c$ and $\langle n \rangle = n_c$ in equilibrium, where $\langle n \rangle = \int dr n/V$ is the average density. We then lowered the temperature of all the boundary walls from $1.1T_c$ to $0.9T_c$ at $t=0$. Subsequently, a liquid wetting layer thickened and acted as a piston adiabatically expanding the interior. Here the interior was in a gaseous state with $n < n_c$ (see Fig. 2). In Fig. 1, we show snapshots of the density $n$, where instability and coarsening proceeded from the boundary into the interior on a slow time scale. It is remarkable that the interior can be adiabatically quenched on a rapid time scale. In Fig. 2, we display $T$, $n$, $s$, and the inverse isothermal compressibility,

$$\frac{1}{K_T} = n \left( \frac{\partial p}{\partial n} \right)_T = \frac{k_B n}{(1 - v_0 n)^2} (T - T_s),$$

at the cell center $x = y = L/2$ in the early stage when the interior was still homogeneous. It demonstrates that the adiabatic changes were caused by sounds emitted from the boundary walls. In the present case, the isothermal compressibility be-
came negative or brought below the spinodal \( T_s/T_c(n) \) in the early stage. The acoustic time scale \( L/c \) is of order \( \alpha^{1/2}t_0L/\ell \sim 10t_0 \) from Eqs. (3.8) and (3.9), which agrees with the oscillation period in Fig. 2.

We make some comments. (i) Though the interior was transiently close to the critical point in the adiabatic quenching in Fig. 2, we neglected the critical fluctuations and the critical anomaly of the transport coefficients. (ii) Though not shown here, temperature variations are significant among domains in the presence of latent heat, while previous simulations of spinodal decomposition have been performed in the isothermal condition [6,27,39]. (iii) In our case, a wetting layer acts as a piston because of its thickening in deep quenching. In near-critical fluids a thermal diffusion layer can well play the role of a piston in supercritical states. If the bulk region is in a liquid state below \( T_c \), a cooled thermal diffusion layer can even cause phase separation in the interior [54]. By cooling the boundary wall at fixed volume, Moldover et al. [53] could realize homogeneous nucleation in the bulk at liquid densities but could not do so at gas densities, where the latter result was ascribed to the preexistence of a liquid wetting layer. The growth process of a liquid wetting layer upon cooling itself has not yet been examined in the literature.

C. Piston effect with a bubble in liquid at \( g=0 \)

Let us consider the piston effect in the presence of a bubble in liquid. When \( T_b \) is changed by \( \delta T_b \), sound waves emitted from the boundary cause a pressure change \( \delta p \). After many traversals of the sounds in the cell, \( \delta p \) soon reaches of the following order [52]:

\[
\delta p \sim A_{st}^{-1}\delta T_b, \tag{3.13}
\]

where \( A_{st} \) is the liquid value of the adiabatic coefficient \( A_s = (\partial T/\partial p)_s \). From Eqs. (2.3) and (2.4) the van der Waals theory gives

\[
A_s = \left( \frac{\partial T}{\partial p} \right)_s = \frac{1/\nu_0}{k_B[(1 + d^2) - dT/2T]}, \tag{3.14}
\]

If \( \delta T_b \sim 0.1T_c \) and \( T/T_c \sim 0.9 \), we find \( \delta p \sim 0.1p_c \), which is 10 times larger than the Laplace pressure difference \( \gamma/R \) for \( R \sim 50\ell [46] \). Here, due to the different values of \( A_s \) in the two phases, there arises a considerable temperature difference in the two regions,

\[
(\Delta T)_{ge} = (A_{ge} - A_{ae})\delta p, \tag{3.15}
\]

in the early stage. For example, \( A_{ge}k_B/\nu_0 \) is equal to 7.1 for gas and 0.65 for liquid at \( T=0.875T_c \) from Eq. (3.14) in two dimensions. On the other hand, near the critical point, \( A_{ge} - A_{ae} \) tends to zero as \( 2\gamma_s^{1/2}(\partial T/\partial p)_s \), where \( \gamma_s \) is the specific heat ratio [6]. Thus, the bubble temperature becomes higher (lower) than in the surrounding liquid for positive (negative) \( \delta T_b \). For the case of a liquid droplet, as will be shown in Fig. 6, the droplet temperature becomes lower (higher) than in the surrounding gas for positive (negative) \( \delta T_b \).

In Figs. 3 and 4, we prepared a bubble with radius \( R = 50\ell = L/4 \) in equilibrium at \( T=0.875T_c \) under zero gravity. We then changed \( T_b \) to 0.895\( T_c \) for \( t>0 \) with \( T_t \) held unchanged. In the upper plate of Fig. 3, the bubble center is more heated than in the liquid at \( y=L/5 \) and \( 4L/5 \) in the early stage in accord with Eq. (3.15). We notice that the oscillation persists within the bubble for a longer time than in the liquid region. The lower panel of Fig. 3 shows the velocity field induced by the sounds at \( t=55t_0 \), which is larger within the bubble than outside it. The typical velocity within the bubble is a few times larger than \( \sqrt{\langle v^2 \rangle} = 0.011\ell/t_0 \) at this time. It indicates the presence of appreciable mass flux through the interface and suggests existence of collective modes with evaporation and condensation localized near a bubble. In Fig. 4, we show \( (T-T_c)/(T_b-T_t) \), \( p/p_c \), and \( v_0t/v_c \) vs \( y/L \) at \( t=200t_0 \) and \( x=L/2 \). Here, the oscillation is mostly damped, while the temperature inside the bubble is not yet flat. We also notice that the thermal diffusion layer is thickened up to the bubble, since the distance between the bottom and the bubble is narrow. It would take an extremely long time for macroscopic separation. While the temperature is continuous, the pressure is slightly higher inside the bubble than outside it by \( \gamma/R \approx 0.01p_c \) and exhibits a singularity in the interface region. The latter behavior can be seen in the equilibrium relation (2.10), from which the integral of \( p \) in the interface region in the normal direction is equal to \(-3\gamma/2 \) for constant \( M \).

D. Two-phase states in heat flow at \( g=0 \)

In our previous work [15,16], we examined the temperature and velocity field around a droplet in heat flow. We summarize the main results here. Under zero gravity, we showed numerically and analytically that a gas (liquid) droplet slowly moves to the warmer (cooler) boundary and that the temperature becomes nearly homogeneous within the droplet. Let a gas droplet (bubble) with radius \( R \) be placed in a heat flux \( Q=-\lambda_2T_s/T_c \), where \( \lambda_2 \) is the liquid thermal conductivity and \( T_s \) is the temperature gradient far from the droplet. A convective velocity of order

\[
v_c = Qn_2T_s\Delta s = -\lambda_2T_s/n_2T_s\Delta s, \tag{3.16}
\]

is induced within the droplet, where \( \Delta s=s_g-s_c \) is the entropy difference per particle between the two phases. The velocity \( v_c \) characterizes latent heat flow and is independent of the bubble radius. The Reynolds number \( Re=v_0p_cR/\eta_k \) is less than unity for \( Re<\eta_kT_s\Delta s/mQ \), where \( p_c=mn_2 \) and \( \eta_k \) are the mass density and the shear viscosity, respectively, in the gas phase.
In our simulation $Re$ is smaller than unity. In Figs. 5–8, we examine heat-conducting two-phase states at zero gravity, where the top temperature at $y=L$ is fixed at $T_\text{r}=0.875T_c$ and the bottom temperature $T_b$ at $y=0$ is changed to a higher value. In Fig. 5, we initially prepared a bubble with radius $R=50\ell$ in equilibrium at $T_c=0.875T_c$ at the cell center. We then raised $T_b$ to a high value, $9.457T_c$, for $t>0$ to induce droplet motion toward the bottom. The time scale of migration was about $10^3t_0$. Eventually, the droplet partially wetted the bottom plate with an apparent contact angle. However, Fig. 5 shows the presence of a thin gas layer sandwiched between them. The typical velocity in Fig. 5 is given by $\sqrt{\langle \mathbf{v}^2 \rangle}=0.029\ell/t_0$, which is of order $v_\infty$ in Eq. (3.16). We can see that $T$ is flat inside the bubble and continuous across the interface. The generalized chemical potential $\hat{\mu}$ in Eq. (2.18) exhibits the same behavior (not shown here). The pressure is nearly constant outside the bubble at $p=0.76p_c$, while it is slightly higher inside by $\sqrt{\gamma/R-0.01p_c}$. With further increasing $T_b$, the apparent contact angle decreases and the bottom wall is eventually wetted completely by gas. See such an example of complete wetting at $T_b=0.97T_c$ in Ref. [15]. As a related experiment, Garbaros et al. observed that gas spreads on a heated wall initially wetted by liquid and exhibits an apparent contact angle [10].

Next, we initially prepared a liquid droplet in gas with $R=50\ell$ at the cell center at $T_c=0.875T_c$ and then raised $T_b$ to $0.91T_c$ for $t>0$. In Fig. 6, we show $T$, $\mathbf{v}$, and $n$ around a liquid droplet sustained in gas at $t=4565t_0$. Here the velocity field is nearly steady and, as in the lower panel of Fig. 3, it is much smaller in the liquid than in the gas due to the viscosity difference from Eq. (3.3). The average velocity amplitude is given by $\sqrt{\langle \mathbf{v}^2 \rangle}=0.015\ell/t_0$ at this time. The temperature inside the droplet is nearly homogeneous and is lowest in the cell, while the pressure inside it is again only slightly higher than outside it by $\sqrt{\gamma/R-0.01p_c}$. Hence, condensation is taking place at the interface and the temperature inside it is slowly increasing in time as $0.849T_c$, $0.851T_c$, and $0.855T_c$ at $t/t_0=4565$, 7770, and 15820, respectively, on the time scale of thermal diffusion ($\sim R^2/D\sim \ell^2/c^2$). Note that the initial temperature lowering inside the liquid droplet can be explained by the adiabatic relation (3.15) (see the arguments around it). In Fig. 7, we show three snapshots of the density at later times in the same run as in Fig. 6. The droplet migration to the cooler wall is very slow, but after its encounter with it the liquid spreads over it rather rapidly under the wetting condition (3.2). In this process the warmer boundary remains dried.

### E. Heat conduction in steady two-phase states

Even when the liquid volume fraction is small, efficient heat transport is realized in the presence of evaporation on the warmer boundary and/or condensation on the cooler boundary. This is the mechanism of heat pipes widely used in refrigerators and air conditioners. In such processes, the warmer boundary dries out if evaporation proceeds without supply of liquid. Here we demonstrate that steady heat transport with evaporation and condensation can be achieved in a closed cell in the presence of a liquid wetting layer and/or a gravity.

In our situation, we define the effective thermal conductivity through the cell by

$$\lambda_{\text{eff}} = \int_0^L dx Q_b(x)/(T_b - T_t).$$  \hspace{1cm} (3.17)

As the heat flux, we choose its value at the bottom $y=0$,

$$Q_b(x) = - (\lambda \partial T_b/\partial y)_{y=0},$$  \hspace{1cm} (3.18)

where $\mathbf{v}$ vanishes. It is the heat flux from the bottom boundary wall to the fluid. In convection, a normalized effective thermal conductivity, called the Nusselt number, has been used to represent the efficiency of heat transport. We define it by

$$Nu = \lambda_{\text{eff}}/\lambda_c.$$  \hspace{1cm} (3.19)

We take $\lambda_c$ as the thermal conductivity at $n=1.7n_c$, which is the liquid density in two-phase coexistence at $T_c=0.875T_c$. [see the sentences below Eq. (3.4)]. In one-phase states without convection, we have $Nu=n/1.7n_c$ under Eq. (3.4). Therefore, $Nu$ is small for gas without convection.

In Fig. 8, we show $T$, $\mathbf{v}$, and $n$ in a steady heat conducting state with $T_b=0.89T_c$ and $T_t=0.875T_c$ at zero gravity. A characteristic feature in this case is that a liquid layer covers all the boundary walls. The average density $(n) = 0.22v_0^{-1}$ is relatively small. The typical velocity is given by $\sqrt{\langle \mathbf{v}^2 \rangle} = 0.008\ell/t_0$. Remarkably, the temperature profile is nearly flat in the majority gas region, due to latent heat convection through the cell. As a result, $Nu=5.0$ is realized, despite the fact that the liquid region is only near the boundary walls.

Under gravity, the warmer boundary does not dry out even for large heat flux. In Fig. 9, we realized a steady state with $(n)=0.206v_0^{-1}$ and $T_b=0.91T_c$ under gravity at $g^* = 10^{-4}$, where $\sqrt{\langle \mathbf{v}^2 \rangle} = 0.0119\ell/t_0$ and $Nu=2.0$. Again a liquid layer covers all the boundary walls, while the bottom part is thicker due to the gravity. These features are very different from those in the right panel of Fig. 7 without gravity. In Fig. 9, the temperature gradient is large everywhere on the side walls and widely in the region $y/\ell \leq 0.25$ near the bottom, while the liquid-gas interface is at $y/\ell \approx 0.1$ apart from the side walls. Due to these features we obtain the smaller value of $Nu$ than for the case in Fig. 8. In Fig. 10, we raised $T_b/T_c$ from 0.91 to 1.1 starting with the steady state in Fig. 9. We recognize that this is the case of strong heating, where bubbles emerge from the bottom to break the liquid into smaller pieces. As the gas flow from the bottom is warmer, condensation occurs on the side and top boundary walls, leading to growth of the liquid wetting layer there.

### F. Boiling in gravity

Boiling phenomena are very complex and still remain virtually unexplored in physics [6,12–14]. Usually, the typical bubble size is macroscopic and the associated Reynolds number is large. In the present case, we realized boiling under gravity at $g^* = 10^{-4}$ even for our small system size, where bubbles from the heated bottom wall were small but could rise upward with small Reynolds numbers.

In Figs. 11 and 12, we set $(n)=0.34v_0^{-1}$ and held $T_b$ at 0.775$T_c$. Starting with an equilibrium two-phase state at $t$
FIG. 11. Changeovers of boiling with increasing \( T_b \) at \( T_c =0.775T_c \), where \( \phi \approx 10^{-4} \). Upper plates: nonboiling state at \( T_b =0.915T_c \) (left), and weakly boiling state with a single bubble at \( T_b =0.945T_c \) (middle and right). Middle plates: transition from weak to strong boiling after a change of \( T_b \) from 0.945\( T_c \) to 1.05\( T_c \) at \( t =34245t_0 \). Lower plates: dynamical steady state with continuous bubble formation and rising.

=0, we reached a steady nonboiling state at \( T_b =0.915T_c \), where \( \sqrt{(\nu)} =0.023\ell/t_0 \) and \( Nu =3.2 \) owing to latent heat convection in the gas region. See the left snapshot at \( t =10^4t_0 \) in the upper panels of Fig. 11. We then raised \( T_b \) to 0.945\( T_c \) to produce a weakly boiling state, where a single bubble periodically appeared with a period about \( 10^3t_0 \). At \( t=27700t_0 \) a bubble is at the center of the liquid region, while at \( t=28010t_0 \) it is being absorbed into the gas region. Here \( \sqrt{(\nu)} =0.033\ell/t_0 \) and \( Nu =4.5 \) at \( t=27700t_0 \). Finally, at \( t=t_0 =34245t_0 \), we again raised \( T_b \) to 1.05\( T_c \). The middle panels of Fig. 11 display transition from weak to strong boiling. The bottom plates of Fig. 11 are those in fully developed boiling, which was reached for \( t-t_b \approx 5000t_0 \). In Fig. 12, we show \( T, n, \text{ and } \nu \) in fully developed boiling at \( t=63047t_0 \), where \( \sqrt{(\nu)} =0.055\ell/t_0 \) and \( Nu =9.2 \). We can see steep temperature gradients near the boundary walls and large velocity deviations around the rising bubbles. In Fig. 13, we show that the heat flux \( Q_b(x) \) on the bottom in Eq. (3.18) is large where rising bubbles are being produced. The upper panel at \( t=34\,600t_0 \) is at the inception of strong boiling, exhibiting a sharp maximum where a bubble is being detached. The lower panel of Fig. 13 at \( t=63\,047t_0 \) is taken in the fully developed boiling state, exhibiting multiple peaks.

In our boiling simulation, the bubble sizes are of order \( R \approx 20\ell \) and the Reynolds number \( Re = \sqrt{(\nu)}R/\nu_0 \) on the droplet scale is of order 0.1 in the lower panels in Fig. 11 and in Fig. 12. Thanks to the high efficiency of latent heat transport, the temperature gradient is mostly localized near the boundary walls. Generally in gravity, there should remain a small temperature gradient in the bulk region in boiling and in stirring. It is given by the so-called adiabatic temperature gradient [6,14],

\[
\left( \frac{dT}{dx} \right)_{\text{bulk}} = -\rho g \left( \frac{dT}{dp} \right)_{\text{sat}},
\]

which is the minimum for the onset of convection for compressible fluids [47] and is equal to \(-0.27 \text{ mK/cm} \) for CO2 near the critical point. In our case, the right-hand side of Eq. (3.20) is of order \(-g' T_c/\ell \) and is consistent with the average gradient seen in the upper panel of Fig. 12. In usual Rayleigh-Bénard convection in one-phase fluids, the average temperature gradient in the bulk region can be suppressed only in turbulence by rapid convective motion of thermal plumes, where the Reynolds number exceeds unity even on the scale of the plumes [56].

G. Wetting dynamics

Wetting dynamics has been studied mostly for involatile droplets [7], but it has not yet been well understood for volatile droplets or in the presence of evaporation and condensation [8–10]. The previous approaches in the latter case were by solving the hydrodynamic equations with phenomenological surface boundary conditions [8] and by performing molecular dynamics simulations of nanometer-size liquid droplets [9]. Here we present its preliminary numerical study in our scheme.

At \( t=0 \), we placed a semi-circular liquid droplet with \( R =40\ell \) on the bottom, where the interface was on the curve \((x−L/2)^2+y^2=R^2 \). The initial temperature was 0.875\( T_c \) throughout the cell. In the upper panels of Fig. 14, where both \( T_b \) and \( T_c \) were held unchanged, spreading of the liquid region occurred over the walls on a time scale of \( 10^3t_0 \). In the middle panels of Fig. 14, we set \( T_b =0.915T_c \) for \( t>0 \) to induce slow evaporation of the liquid region. We can see only a small amount of liquid left on the bottom at \( t =4000t_0 \) (right). In the bottom panels of Fig. 14, where \( T_b =0.975T_c \) for \( t>0 \), a gas droplet appeared within the liquid region and, as it grew, the liquid region itself was finally detached from the boundary (right). In these processes, a considerable amount of liquid was evaporating even for the first case of \( T_b =T_c =0.875T_c \) and the latent heat needed for evaporation was supplied from the equitemperature boundary wall. In all these examples, the liquid region was itself cooled even below the top temperature \( T_c =0.875T_c \) by an amount typically of order 0.05\( T_c \). In Fig. 15, we show the velocity \( \nu \) (upper panel) and the heat flux \( Q_b(x) \) (lower panel) at \( t=500t_0 \) obtained from the run giving the series (a) in Fig. 14. We can see a circulating flow in the gas and large heat absorption at the contact points. The peak height of \( Q_b(x) \) gradually decreased with spreading. These stem from evaporation at the contact points and condensation on the top of the droplet. The flow pattern here is very different from that for involatile droplets [7].
We mention two theoretical papers in accord with our results. Koplik et al. found a large temperature decrease of a nanometer-scale liquid droplet on a heated wall using molecular dynamics simulation. Nikolayev et al. solved the heat conduction equation around a contact point of a bubble in two dimensions to find a divergence of \( Q_b(x) \) there on an equitemperature heated wall.

**IV. SUMMARY**

We have developed a general scheme of the dynamic van der Waals theory. Our hydrodynamic equations including the gradient stress can readily be integrated numerically. We may then gain intuitive ideas of complicated but fundamental hydrodynamic processes in two-phase states. In particular, emphasis has been put on adiabatic changes and latent heat transport in two-phase states, which have rarely been discussed in the literature. Though in two dimensions and for small system size, we have presented a number of simulation results to demonstrate the usefulness of our scheme. Examples treated are (i) adiabatically induced spinodal decomposition, (ii) piston effect with a bubble in liquid, (iii) temperature and velocity profiles around a gas and liquid droplet in heat flow, (iv) efficient latent heat transport at small liquid densities, (v) boiling in gravity with continuous bubble formation and rising, and (vi) spreading and evaporation of a liquid region on a heated boundary wall.

We make some comments below.

(i) Our numerical analysis in this work is very fragmentary. We should investigate each problem in more detail over wide ranges of the parameters. In particular, we have used the special boundary condition for which liquid is strongly attracted to the boundary walls. We should perform more simulations for various \( a_i \) and \( b_j \) in Eq. (2.33).

(ii) The mesh size of our simulation cell is on the order of the interface thickness away from the critical point. If \( \ell \) is a microscopic length in our numerical analysis, we have treated hydrodynamics on small scales (\( \approx 10 \) nanometers), as in the molecular dynamics simulation. We have not examined crossovers taking place with increasing the spatial scales, but general features of adiabatic changes and latent heat transport revealed in this work should be independent of the spatial scales.

(iii) In integrating our hydrodynamic equations, we do not assume phenomenological boundary conditions at the interface, while they are needed in the hydrodynamic calculation in the thin interface limit. To clarify the relationship between these two approaches, we should examine the...
thin interface limit of our model [26–28]. Moreover, in the phase field model of crystal growth [31–33], the interface motion can be insensitive to the interface thickness \( \ell \) under some conditions among the parameters in the model. Also in our fluid problems, we should clarify such conditions.

(iv) In our numerical analysis, we evidently see smooth variations of the temperature \( T \) and the generalized chemical potential \( \mu \) in Eq. (2.18) across the interface (not shown for \( \mu \)). In our hydrodynamic theory [16], we assumed the continuity of these quantities together with the stress balance and the mass and energy conservation at the interface. In the literature [8,29], however, discontinuities in the temperature and the chemical potential have been supposed to arise across the interface between liquid and dilute gas (which originate from the long mean free path in the gas phase).

(v) Since the Reynolds number is small in this work, we should examine the high-Reynolds-number regime by enlarging the droplet size \( R \) in a much larger system [see the discussion below Eq. (3.16)].

(vi) Figure 3 indicates complex nonspherical dynamic response of bubbles to sound waves in the presence of evaporation and condensation. This aspect seems to have not yet been investigated, while large sound attenuation of liquid containing bubbles has long been discussed [6].

(vii) We should study hydrodynamics near the critical point, where the singularities of the thermodynamic and dynamical properties largely influence the dynamics [6,10,12,14,54]. It is well-known that the piston effect gives rise to critical speeding-up of thermal equilibration [52], while the thermal diffusion is slowed down. In near-critical one-phase convection, significant are overall pressure and temperature fluctuations propagating as sounds on birth and arrival of thermal plumes at the boundary walls [56]. In near-critical two-phase states, such adiabatic fluctuations could be stronger in boiling and phase separation.

(viii) We are also interested in droplet motion and two-phase hydrodynamics in binary fluid mixtures. Marek and Straub examined the effect of a small amount of noncondensable gas doped in the gas phase [43], which drastically alters thermocapillary convection. Pooley et al. [20] took account of latent heat transport in binary mixtures to calculate the Nusselt number in two-phase convection.

(ix) In the Landau and Lifshitz book [47], the linearized hydrodynamic equations are treated as Langevin equations, where newly added are random stress and random heat flux related to the transport coefficients via the fluctuation-dissipation relations [6]. They describe the thermal fluctuations of the hydrodynamic variables. The random source terms in the dynamic equations are particularly important near the critical point [5,6]. We should clarify how our hydrodynamic equations with the gradient stress can be treated as Langevin equations.

(x) We mention the lattice-Boltzmann method devised to treat two-phase flow and phase separation [20,57,58]. In particular, Palmer and Rector included thermal effects to calculate a nonequilibrium temperature profile around a gas-liquid interface [58].

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APPENDIX A: DERIVATION OF THE REVERSIBLE STRESS

We derive the reversible stress tensor \( \mathbf{\Pi} \) following the method in Appendix 6A of Ref. [6]. Adopting the Lagrange picture of fluid motion, we suppose a small fluid element
with volume $V$ at position $r$ and at time $t$. Due to the velocity field the element is displaced to a new position, $r'=r+u$ with $u=\mathbf{v}\cdot\partial t$, after a small time interval $\partial t$. Hereafter, we neglect the dissipation ($\eta=\nu=\lambda=0$). The density $n$ and the volume of the fluid element $V$ are changed by

$$\delta n = -n \nabla \cdot u, \quad \delta V = V \nabla \cdot u. \quad (A1)$$

The change of the internal energy in the volume element is written as $\delta E = \int dr' \epsilon' - \int dr \epsilon = V \delta \epsilon$, where $dr'=dr(1+\nabla \cdot u)$. From Eq. (2.39) we have

$$\delta \epsilon = -(\epsilon\mathbf{I} + \mathbf{P}) \nabla \cdot u, \quad (A2)$$

where $\mathbf{I}=\{\delta_{ij}\}$ is the unit tensor. We shift the coordinates as $r'=r+u$; then the space derivatives are changed as

$$\frac{\partial}{\partial x_i} = \frac{\partial}{\partial x_i} - \sum_j \frac{\partial u_j}{\partial x_i} \frac{\partial}{\partial x_j}. \quad (A3)$$

As a result, $|\nabla n|^2/2$ is changed by $\Sigma_i (\nabla \epsilon) (\nabla \delta n) - \Sigma_i (\nabla n) \times (\nabla \epsilon) (\nabla n) \nabla n: \nabla n$. We then calculate the entropy change in the volume element $\delta S = \int dr' \delta s - \int dr \delta s$ as

$$\frac{T}{V} \delta S = [(T \mathbf{S} - \delta + n \delta \mathbf{P}) - M \nabla n] \nabla n. \quad (A4)$$

Without dissipation we should require $\delta S=0$ to obtain the reversible stress $\mathbf{I}$ in Eqs. (2.47) and (2.48).

**APPENDIX B: SCALED HYDRODYNAMIC EQUATIONS**

We rewrite our hydrodynamic equations into dimensionless forms in two dimensions using Eqs. (3.3) and (3.4). The scaled space and time, $\ell^{-1}r$ and $t_0^{-1}t$, will be simply written as $r$ and $t$, respectively. Here we introduce the scaled density, temperature, and velocity,

$$\phi = \nu_0 n, \quad \Theta = \frac{k_B}{\epsilon} T, \quad V = \frac{t_0}{\ell} \mathbf{v}. \quad (B1)$$

The continuity equation (2.35) is written as $\partial \phi / \partial t = -\nabla \cdot (\phi \mathbf{V})$. The momentum equation (2.36) reads

$$\sigma \left[ \frac{\partial}{\partial t} \phi \mathbf{V} + \nabla \cdot (\phi \mathbf{V}) \right] = -\nabla \cdot \mathbf{M} - \sigma \phi \mathbf{V}. \quad (B2)$$

where $\sigma$ and $\sigma^*$ are the constants defined by Eqs. (3.6) and (3.7). Including the viscous stress we define

$$M_{ij} = \left[ \frac{\Theta \phi}{1 - \phi} - \phi - 2 \Theta \nabla^2 \phi - \Theta (\nabla \phi)^2 \right] \delta_{ij} + 2 \Theta (\nabla \phi)(\nabla \phi) - \sigma \phi \nabla \phi \cdot \nabla \phi. \quad (B3)$$

We define a scaled total internal energy density by $E_T = e_T \nu_0 / \epsilon$ to obtain

$$E_T = \phi \Theta - \phi - \frac{\sigma}{2} \phi \nabla^2 \phi, \quad (B4)$$

which is governed by

$$\frac{\partial}{\partial t} E_T + \nabla \cdot (E_T \mathbf{v} + M \cdot \mathbf{V}) = \nabla \cdot \phi \nabla T - \sigma \phi \nabla V. \quad (B5)$$

The boundary condition (3.2) is rewritten as

$$\nu \cdot \nabla \phi = (\phi - 5/6) / 2 \sqrt{\gamma}. \quad (B6)$$

Equations (2.35), (B2), and (B5) were integrated using the Euler scheme with the time mesh size $\Delta t=0.01 [59]$. 


[15] A. Onuki, Phys. Rev. Lett. 94, 054501 (2005). In this paper the gradient stress is written with the opposite sign. Also there is an error in the computer program such rotational motion of flow is incorrectly superposed.


For diatomic molecules such as O$_2$ the kinetic energy contribution to $\varepsilon$ should be $5nk_BT/2$ due to the two rotational degrees of freedom. For polyatomic molecules such as CO$_2$ it should be $3nk_BT$.


[46] H. Kitamura and A. Onuki, J. Chem. Phys. 123, 124513 (2005). The approximation $\gamma=A\varepsilon^{1-\delta(1-T/T_c)^{3/2}}/2A$ holds within a few percent in the range $0.6\leq T/T_c < 1$ for the van der Waals model. Here $M$ is assumed to be independent of $n$. For example, $\gamma = 0.02\varepsilon/v_0$ at $T/T_c = 0.85$ in terms of $\ell$ in Eq. (3.1).


[49] From the argument of the entropy production rate only, our heat flux $J_q$ in Eq. (2.39) and our diagonal pressure $p_1$ in Eq. (2.48) may be generalized to $J_q = J_q + \alpha J_{GA}$ and $p_1 = p_1 + \alpha n \nabla \cdot (M/T)$, where $\alpha$ is arbitrary. The expressions in Refs. [24,26] follow for $\alpha = 1$. However, the arguments in Appendix A lead to $\alpha = 0$.


[59] As a technique increasing the numerical stability, we defined $q = \lambda \nabla T$ on the midpoints $[i+1/2,j+1/2]$, where $[i,j]$ represent the lattice points. For example, $q_i$ at the middle point $[i+1/2,j+1/2]$ was defined by $(\lambda[i+1,j+1]+\lambda[i,j+1]+\lambda[i+1,j]+\lambda[i,j])/(T[i+1,j+1]+T[i+1,j]+T[i,j+1]+T[i,j])/8$. Then $\nabla \cdot \lambda \nabla T$ at the point $[i,j]$ was set equal to $(q_i[i+1,j+1/2]-q_i[i-1,j+1/2]+q_i[i+1/2,j-1/2]-q_i[i-1/2,j-1/2])/2$. The same technique was also used for $\nabla \cdot \eta \nabla v_k$.