The Influence of Intermolecular Forces at Critical Point Wedge Filling

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We use microscopic density functional theory to study filling transitions in systems with long-ranged wall-fluid and short-ranged fluid-fluid forces occurring in a right-angle wedge. By changing the strength of the wall-fluid interaction we can induce both wetting and filling transitions over a wide range of temperatures and study the order of these transitions. At low temperatures we find that both wetting and filling transitions are first-order in keeping with predictions of simple local effective Hamiltonian models. However close to the bulk critical point the filling transition is observed to be continuous even though the wetting transition remains first-order and the wetting binding potential still exhibits a small activation barrier. The critical singularities for adsorption for the continuous filling transitions depend on whether retarded or non-retarded wall-fluid forces are present and are in excellent agreement with predictions of effective Hamiltonian theory even though the change in the order of the transition was not anticipated.

The properties of confined and inhomogeneous fluids have received enormous theoretical and experimental attention over the last few decades [18]. These have revealed convincing evidence that the interplay between surface tension, intermolecular forces and the substrate geometry can induce new phase transitions and fluctuation regimes beyond that occurring for bulk fluids. A simple but striking example of a transition induced by a structured surface is the filling transition pertinent to fluid adsorption in a linear wedge formed by the junction of two planar walls with opening angle 2ψ. Simple macroscopic arguments dictate that, at bulk two phase coexistence, a wedge-gas interface is completely filled by liquid for temperatures T > T_{fill}, where the filling temperature T_{fill} is given implicitly by [9–12]

\[
\theta(T_{\text{fill}}) = \frac{\pi}{2} - \psi
\]  

(1)

where \( \theta(T) \) denotes the equilibrium contact angle. Since (1) is exact, all fluids that form drops with a finite contact angle will exhibit a filling transition in a wedge tuned to the appropriate opening angle. The properties of the filling transition were first studied using mesoscopic effective Hamiltonian models which concluded that the transition may be first-order or continuous determined by the qualitative properties of the interfacial binding potential (see below) [13–15]. The critical singularities characteristic of continuous filling depend sensitively on the dimensionality and the range of the intermolecular forces. In 3D, even in the presence of long-ranged forces, they show dramatically enhanced interfacial fluctuation effects compared to wetting [15,17] while in 2D many properties of wetting and filling are precisely related by a symmetry called wedge covariance [18]. Interestingly in 3D the conditions for continuous wedge filling appear to be less restrictive than those for continuous wetting hinting that they should be more easily observed in the laboratory [19]. Several of the early predictions of the effective Hamiltonian theory of filling for systems with short-ranged forces have been confirmed in more microscopic approaches including simulations [16,17], exactly solvable models [20,22] and field theory [23]. However recent studies using microscopic Fundamental Measure Density Functional Theory (FM-DFT) suggest that with long-ranged forces there are new aspects of 3D wedge filling not anticipated by interfacial models [24,25]. In particular there is evidence that in acute wedges or close to the bulk critical temperature \( T_c \) continuous filling transitions are even more prevalent than first thought. This phenomenon of critical point wedge filling is not easily explicable using interfacial models, which are typically limited to open wedges and to low temperatures and neglect completely short-ranged correlations associated with the liquid structure near the walls. In this paper we show that critical point wedge filling is present for a broader class of wall-fluid intermolecular forces and determine numerically the critical singularity characterising the divergence of the adsorption which is found to be in excellent agreement with the predicted value for both retarded and non-retarded van der Waals forces.

To begin we recall the simple effective Hamiltonian theory of wedge filling valid for shallow wedges with small tilt angle \( \alpha = \pi/2 - \psi \) [13,15]. Assuming translational invariance along the wedge, the position of the wall may be approximated \( z_w = \alpha |t| \) where \( t \) denotes the horizontal coordinate (see Fig.1) and where we used a standard shallow-wedge approximation \( \tan \alpha \approx \alpha \). If \( \ell(t) \) denotes the local height of the liquid-gas interface above the horizontal then the free-energy cost of a given interfacial configuration is described by the approximate functional

\[
F[\ell] = \int dt \left\{ \frac{\gamma}{2} \ell'^2 + W(\ell - z_w) \right\},
\]

(2)

Here \( \gamma \) is the liquid-gas surface tension, \( \ell \equiv \frac{dt}{d\tau} \) and \( W(\ell) \) is the interfacial binding potential modelling the local wetting properties of a planar wall-gas interface.
density, diverges as energy cost of filling the wedge with liquid to thickness the wedge apex measured above the apex. Inset shows the qualitative shape of the binding potential $W(\ell)$ for wetting at a planar wall for the cases of first-order wetting (solid line) and continuous (dashed line) wetting transitions.

At mean-field level, simple minimization of $F[\ell]$ determines the equilibrium profile and that the height $\ell_w$ of the interface above the wedge apex satisfies

$$\frac{\gamma}{2}(\alpha^2 - \theta^2) = W(\ell_w).$$

Now suppose we are at bulk two-phase coexistence (chemical potential $\mu = \mu_{\text{sat}}(T)$) on the vapor side and below the wetting temperature $T_w$ so that $\theta > 0$ implying $W(\ell)$ has a global minimum (at $\ell_x$, say) and decays to zero as $\ell \to \infty$. From (3) it follows that if $W(\ell)$ has an activation barrier (see inset Fig. 1) then the filling transition occurring as $\theta \to \alpha^+$ is first-order so that $\ell_w$ jumps discontinuously from a finite to macroscopic value at $T_{\text{fill}}$. If there is no activation barrier then the filling transition is second-order and $\ell_w$ diverges continuously. This condition translates as follows: Second-order wetting implies second-order (continuous) filling while first-order wetting implies first-order filling unless $T_{\text{fill}} < T_s$ where $T_s$ is the spinodal temperature, defined as the lowest temperature at which the barrier in $W(\ell)$ appears.

In this case the filling transition will also be continuous. The present theory also predicts the critical exponents for continuous filling. For long-ranged forces, the binding potential decays as $W(\ell) \sim A\ell^{-p}$ with $A$ negative for continuous filling and $p = 2, 3$ for non-retarded and retarded dispersion forces respectively. This implies that the excess adsorption of liquid near the apex $\Gamma = \int dz \rho(x, z) \approx \ell_w^2$ where $\rho_g$ is the bulk gas density, diverges as

$$\Gamma \sim (\theta - \alpha)^{-\frac{2}{p}}.$$  

The same result for the divergence of $\Gamma$ follows from a different argument which determines directly the free-energy cost of filling the wedge with liquid to thickness $\ell_w$. Young’s equation implies that the surface tensions contribute a term $\propto \sin(\theta - \alpha)\ell_w$ which is balanced by a term $\propto \ell_w^{-p}$ coming directly from the intermolecular forces [15]. If the latter contribution is net repulsive, as it must be for continuous filling, it follows that the film thickness diverges continuously as $\ell_w \sim (\theta - \alpha)^{-\frac{1}{2}}$. These mean-field arguments are believed to be valid for $p < 4$ [15]. For $p \geq 4$ mean-field theory breaks down since interfacial fluctuations are dominant and the adsorption diverges with a universal power-law $\Gamma_w \sim (\theta - \alpha)^{-\frac{1}{2}}$ characteristic of filling in systems with short-ranged forces. This latter prediction is consistent with extensive simulations of filling in the 3D Ising model [16].

An interesting theoretical aspect of wedge filling is that in attempting to understand the transition for more acute wedges we inevitably push effective Hamiltonian theories to the limits of their applicability. It is of course straightforward to improve on the shallow wedge model [2] by replacing the square gradient term with a full “drumhead” expression $\int dt \sqrt{1 + \ell''^2}$ [13]. However to reliably model filling in acute wedges also requires that we abandon the use of the planar wetting potential $W(\ell)$ and adopt a fully non-local description of the interaction of the interface with the non-planar wall [26–28]. This would also account for the self-interaction between the wetting films across the wedge [39]. While this is feasible at least numerically at low temperatures, by for example assuming a sharp-kink approximation for the density profile [3, 28], this is much more difficult if the filling transition occurs close to the bulk critical temperature. To understanding filling in this regime, even at mean-field level, it is necessary and far simpler to move directly to more microscopic DFT methods similar to studies of fluid adsorption in other geometries [29–33].

Within the framework of classical DFT the equilibrium density profile is found by minimizing the grand potential functional [1]

$$\Omega[\rho] = F[\rho] + \int d\mathbf{r} \rho(\mathbf{r}) [V(\mathbf{r}) - \mu]$$

where $F[\rho]$ is the intrinsic free energy functional of the fluid one-body density, $\rho(\mathbf{r})$. Following a perturbative scheme modern DFT usually separates this as

$$F[\rho] = F_{\text{ideal}}[\rho] + F_{\text{hs}}[\rho] + \frac{1}{2} \int d\mathbf{r}_1 d\mathbf{r}_2 \rho(\mathbf{r}_1) \rho(\mathbf{r}_2) u_a(r_{12})$$

where the first two terms on the right hand side are the ideal and hard-sphere contributions respectively. For the latter we use Rosenfeld’s approximate FM theory which accurately models short-ranged repulsive correlations between the fluid atoms [40, 41]. The final term in (6) is a mean-field treatment of the attractive part, $u_a(r)$, of the intermolecular fluid-fluid potential. Following our earlier study we take this to be a Lennard-Jones (LJ) potential $u_a(r) = -4\epsilon(\sigma/r)^6 H(r - \sigma)$ which is truncated at $r_c = 2.5\sigma$, where $\sigma$ is the hard-sphere diameter and $H(x)$ is the Heaviside function.
The external potential $V(\mathbf{r})$ arises from summing over all two-body wall-fluid interactions. That is $V(\mathbf{r}) = \rho_w \int d\mathbf{r}' \phi_{w,f}(|\mathbf{r} - \mathbf{r}'|)$ where the integration is over the whole domain $V$ of the wall which is assumed to be a uniform distribution of atoms, with density $\rho_w$. Here $\phi_{w,f}(r)$ is the wall-fluid two-body interaction which for $r > \sigma$ we suppose is given by the Generalised Long-Range potential $\phi_{w,f}(r) = -4\varepsilon_w \left( \frac{r}{\sigma} \right)^n$. The exponent $n = 4 + p$ for purposes of comparison with the above effective Hamiltonian theory. A hard-wall repulsion is also imposed for $r < \sigma$. For a planar wall occupying the half-space $z < 0$, say, this integration produces the external potential

$$V_\pi(z; n) = \frac{8\pi\varepsilon_w \rho_w \sigma^n}{(n-2)(3-n)} z^{3-n} ; z > \sigma$$

while for a right angle wedge ($\psi = \pi/4$) the potential $V(\mathbf{r}) = V(x, z)$ is a more complicated function of Cartesian $x, z > 0$ although it is translationally invariant along the wedge. Calculation shows that this can be written $V(x, z; n) = V_2(x; n) + V_2(x, z; n)$ where the additional contribution

$$V_2(x; n) = C_n \left[ \frac{J_{n-1}(\frac{z}{\sigma}) + J_{n-1}(\infty)}{x^{n-3}} + \frac{J_{n-1}(\frac{z}{\sigma})}{x^{n-3}} \right]$$

with $C_n = -\frac{4\pi\varepsilon_w \rho_w \sigma^n}{\Gamma(\frac{3}{n})}$ and $I_n(x) \equiv \int dx (1 + x^2)^{-n/2}$. For integer $n$ the latter integral can be easily determined using the recurrence formula

$$nJ_{n+2}(y) = \sinh y \cosh^{-n} y + (n - 1)J_n(y)$$

with $J_2 = 2 \tan^{-1}(e^y), J_3 = \tanh y$ and where we have

$$V_2(x, z; 7) = \frac{1}{15} \varepsilon_w \rho_w \sigma^4 \frac{6 \arctan (\frac{z}{\sigma})}{x^6 + x^4z^2 - x^2z^4 - z^6} - 3\pi x^2z^4 - 3\pi z^6 - 6x^3z^3 - 6xz^5$$

change in sign of the Hamaker constant $A$ is only possible if the wall-fluid and fluid-fluid forces have the same range. Thus there is no spinodal temperature $T_s$ at which the activation barrier disappears. However for wetting (and filling) transitions occurring near $T_s$ the size of the activation barrier may be very small due to the small difference in bulk liquid and gas densities. We will return to this shortly.

Turning attention to the right-angle wedge we determine equilibrium free-energies and density profiles at bulk coexistence from minimization of the functional $\Omega[\rho]$. This is done on an $L \times L$ grid (with $L = 50\sigma$) with discretization size $0.05\sigma$ using the same numerical scheme described in Ref. [42]. According to the thermodynamic prediction [1], the location of the filling transitions can be determined from the intersection of the contact angle curves with $\psi = \pi/4$ and gives $T_{\text{fill}} = 0.72 T_c$ and $T_{\text{fill}} = 0.97 T_c$ as $\varepsilon_w$ decreases in strength. These predictions are in near perfect agreement with our numerical results obtained from minimization of $\Omega[\rho]$ which also determines the order of the filling transition. Starting from

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For the filling transition occurring at $T_{\text{fill}} = 0.97T_c$ on the other hand a unique phase is found for all temperatures indicating that the transition is continuous. Corresponding density profiles are shown in Fig. 3 and show the wedge gradually filling with liquid to a maximum value determined by $L$. A plot of the adsorption $\Gamma$ versus $T$ is shown in Fig. 4 and shows a dramatic but continuous increase in the adsorption near the anticipated $T_{\text{fill}}$. Thus, as with the earlier study with dispersion forces $(n = 6)$, it appears that within this microscopic theory, the filling transition occurring near $T_c$ is continuous or at least effectively continuous. Assuming that the transition is continuous the critical singularity for the adsorption is precisely in accord with the expectation \( (\theta(T)^{\frac{1}{\epsilon}}) \).

The observation of the expected $p$-dependence of the adsorption critical exponent for critical point wedge filling with retarded van der Waals forces is the central result of our paper. This continuous phase transition occurs despite the fact that our model DFT has short-ranged fluid-fluid interactions and that therefore the binding potential $W(\ell)$ for the planar wetting transition still exhibits an activation barrier. Therefore the condition for continuous filling, according to the shallow wedge interfacial model is not met. However numerical determination of the binding potential by constrained partial minimization of the Grand potential functional $\Omega[\rho]$ shows that the activation barrier is extremely small near the filling temperature $T_{\text{fill}} = 0.97T_c$. This can be seen in our final figure (Fig. 5) where the activation barrier is hardly visible. Thus while the condition for continuous filling, according to the shallow wedge interfacial model, is not strictly met, the transition would be predicted to be very weakly first-order. It is possible that in much larger systems the filling transition in our DFT model would also appear first-order and that eventually the adsorption would show a jump to a macroscopic value.

FIG. 2: Temperature dependence of the contact angle for two different wall strengths $\varepsilon_w = 2\varepsilon$ and $\varepsilon_w = 2.5\varepsilon$. The bulk critical temperature is located at $k_BT_c/\varepsilon = 1.41$. The intersection of $\theta(T)$ with the dashed line at $\theta = 45^\circ$ is the thermodynamic prediction for $T_{\text{fill}}$. In the inset is shown a log-log plot illustrating the vanishing of $\theta$ in the vicinity of each $T_{\text{fill}}$; both fitted straight lines have slope equal to $1/2$ consistent with the expected first-order singularity $\theta(T) \sim (T_{\text{fill}} - T)^{\frac{1}{2}}$.

FIG. 3: Coexisting density profiles near a first-order filling transition for wall strength $\varepsilon_w = 2.5\varepsilon$ (corresponding to $k_BT_{\text{fill}}/\varepsilon = 1.05$). The upper panel shows the microscopic configuration in which the interface is tightly bound to the apex. The lower panel shows the macroscopic configuration in which the meniscus is far from the wall and meets each wall at the contact angle $\theta = \pi/4$ in line with the macroscopic condition (1).
FIG. 4: Sequence of density profiles for the wedge with weaker strength ($\varepsilon_w = 2\varepsilon$) showing a continuous increase in the adsorption of liquid as $T$ is increased to the filling transition occurring at $k_bT_{\text{fill}}/\varepsilon = 1.3693$. From top left to bottom right $k_bT/\varepsilon = 1.367, 1.368, 1.369$ and 1.3692.

FIG. 5: Plot of the adsorption (in reduced units) as a function of temperature for the wedge with weaker wall strength $\varepsilon_w = 2\varepsilon$. Inset is a log-log plot and comparison with the prediction for continuous filling with retarded van der Waals interactions $\Gamma \sim (T_{\text{fill}} - T)^{-2}$. However this is not what was observed in our previous studies for non-retarded forces and there is no indication of this in our numerical calculations. It appears to us to be more likely that the shallow wedge interfacial model, which uses a purely local effective interfacial interaction is inadequate close to $T_c$, see Ref. [?]. Whatever scenario, the observed behaviour in our numerical DFT study has all the hallmarks of a second-order phase transition showing the expected $p$-dependence of the critical exponents. This means that the conditions for continuous or effective continuous filling are even more relaxed than initially anticipated. This is something that we hope can be tested in the laboratory similar to experiments on complete wedge filling [19]. If continuous filling is experimentally accessible it would be a means of observing the dramatic and universal enhancement of the interfacial roughness $\xi_{\perp} \propto (\theta - \alpha)^{-\frac{1}{4}}$ which is predicted even in the mean-field fluctuation regime of wedge filling pertinent to systems with retarded or non-retarded forces [19].

FIG. 6: Plot of the numerically determined binding potential for wetting at a planar wall with $\varepsilon_w = 2\varepsilon$ at $k_bT/\varepsilon = 1.393$ close to the filling temperature $T_{\text{fill}}$. Notice that $W(\ell)$ shows a very small activation barrier near $\ell \approx 16\sigma$.

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