Dynamic capillary wetting studied with dissipative particle dynamics

Claudio Cupelli\textsuperscript{1,4}, Björn Henrich\textsuperscript{2,3}, Thomas Glatzel\textsuperscript{1}, Roland Zengerle\textsuperscript{1}, Michael Moseler\textsuperscript{2,3} and Mark Santer\textsuperscript{1,3}

\textsuperscript{1} Laboratory for MEMS applications, Department of Microsystems Engineering (IMTEK), University of Freiburg, Georges-Koehler-Allee 106, 79110 Freiburg, Germany
\textsuperscript{2} Freiburg Materials Research Center (FMF), Stefan-Meier-Straße 21, 79104 Freiburg, Germany
\textsuperscript{3} Fraunhofer Institute for Mechanics of Materials IWM, Wöhlerstraße 11, 79108 Freiburg, Germany
E-mail: cupelli@imtek.de

\textit{New Journal of Physics} 10 (2008) 043009 (16pp)
Received 22 August 2007
Published 10 April 2008
Online at http://www.njp.org/
doi:10.1088/1367-2630/10/4/043009

\textbf{Abstract.} We present a study on dynamic capillary wetting in the framework of dissipative particle dynamics (DPD) based on a novel wall model for wetting on solid boundaries. We consider capillary impregnation of a slit pore in two situations: (i) forced (piston-driven) steady state flow and (ii) capillarity driven imbibition out of a finite reservoir. The dynamic contact angle behavior under condition (i) is consistent with the hydrodynamic theories of Cox under partial wetting conditions and Eggers for complete wetting. The flow field near the contact line shows a region of apparent slip flow which provides a natural way of avoiding a stress singularity at the triple line. The dynamics of the capillary imbibition, i.e. condition (ii), is consistently described by the Lucas–Washburn equation augmented by expressions that account for inertia and the influence of the dynamic contact angle.

\textsuperscript{4} Author to whom any correspondence should be addressed.
1. Introduction

Dynamic wetting phenomena such as spreading of droplets and thin films, or the impregnation of capillary tubes and channels is of great importance in many areas of applied and engineering science [1], microfluidics [2] or thin film lubrication [3]. Studying the temporal evolution of dynamic capillary menisci involving moving liquid fronts by analytical or numerical methods, however, is much more difficult than the static counterpart, where, e.g. the shape of a droplet at rest is completely determined by the equilibrium (static) contact angle $\theta_0$. A moving liquid front, however, will result in an (apparent) dynamic contact angle $\theta_d (\theta_d \neq \theta_0)$, which is phenomenologically a function of the local marching velocity $v$ or, in dimensionless terms, the capillary number $Ca = \eta v/\sigma$ (measuring the opposing influences of driving capillary and damping viscous forces) [4]. The behavior of $\theta_d(Ca)$ governs the global evolution of free surface flows, and is thus also of great practical relevance. In industrial processes such as curtain coatings it determines under which process parameters air entrainment will set in [5, 6].

Stating a general and complete relation of $\theta_d(Ca)$ is by no means trivial, since details of the microscopic three-phase coexistence region where the liquid meets a moving surface, the ‘contact line’ (CL), become crucial. This manifests itself in an unphysical stress singularity in the solution of the Navier–Stokes-equations (NSE) by applying standard boundary conditions (i.e. ‘stick’ at the solid–liquid (sl) interface, and zero stress at the free liquid surface) to this problem [7, 8]. Various physical mechanisms have been suggested supplementing the continuum description such as slip laws [4, 9], diffuse interface approaches [10], van der Waals forces [1], chemical rate processes and diffusion kinetics [11, 12] or a combination of molecular kinetics and hydrodynamic effects [5, 13, 14].

Experimentally, $\theta_d$ is sensed as the slope of the liquid front some small finite distance away from the CL. Hence, a theoretical description must comprise a global solution for the flow field, turning dynamic wetting into a complicated multi-scale problem [15, 16]. Hydrodynamic theory can provide at most perturbation expansions in $Ca$, and a global solution the NSE is sought by rather involved asymptotic matching of the solutions on various length scales [4, 9, 17–19]. At best, then, one can arrive at a useful relation $\theta_d = f(Ca, L_1, L_2, \ldots, \chi_1, \chi_2, \ldots)$ containing as arguments the relevant length scales $L$ and microscopic model parameters $\chi$ [5]. But the problem seems more involved than that. As was pointed out only recently [9], the length scales might themselves be functions of $Ca$. In addition, recent experimental [5] and numerical works [6] on the curtain coating problem suggest that the flow...
field itself might influence the wetting dynamics rendering the dynamic contact angle problem non-local.

Simulation methods such as molecular dynamics (MD) are in principle capable of representing dynamic wetting problems as a whole, and have extensively been used in the past to shed more light on the molecular processes close to the CL region [20]–[22]. Their application to realistic problems is numerically, however, very costly and therefore limits the size and timescale of the problems to be studied. On the other hand pure continuum methods such as computational fluid dynamics (CFD) methods based on finite elements can conveniently be used to solve the global flow problems in general. However, in the specific problem of free surface flows there is no unique way of coupling the relevant molecular microscopic processes to continuum fluid dynamics [6], [23]–[25].

In recent years novel simulation schemes such as the grid-based Lattice–Boltzmann method (LBM) (see [26] and references therein), kinetic Monte Carlo approaches such as direct simulation Monte Carlo (DSMC) [27, 28] or dissipative particle dynamics (DPD) [29, 30] have emerged that offer alternatives to address the above mentioned problems. The DPD method was originally invented to model the dynamics of solvents on mesoscopic time and length scales, hydrodynamics is represented by the collision dynamics of an ensemble of pseudo-atoms. Fluid properties but also flow boundary conditions arise as a consequence of the collision dynamics or particle interactions. One may think of these methods as coarse-grained versions of an MD, that still retain some molecular characteristics. Thus, when applied to dynamic wetting problems, they are likely to provide some reasonable model for CL motion, such as Navier-slip [31]. But equally important, they are much more efficient than, e.g. MD in solving the global flow problem. In the present work, we apply a multi-body-DPD (MDPD) approach to a nontrivial transient problem of capillary dynamics, namely, the impregnation of a slit channel out of a finite reservoir (section 5). For representing the sl-system, we employ a recently developed model for solid boundaries, which is reviewed in section 3, and some important results on capillary flow induced by a piston at constant capillary number, referred to as forced wetting are explained in section 4 in greater detail. In the following section, we outline the MDPD method used in this work originally proposed by Warren [32].

2. A brief account of the scheme

DPD is a mesh-free particle based simulation method that was first devised by Hoogerbrugge and Koelman [29] in 1992 as a generic method for modeling fluid dynamics. In the DPD-scheme point particles interact via short range forces [29, 33] such that momentum and mass are conserved, which is responsible for the hydrodynamic behavior of a liquid at large scales [34]. A common interpretation is that each particle is considered as a lump of molecules [34]. In this picture the DPD method can be regarded as a coarse-graining of MD and as such is capable of modeling larger length and timescales.

The dynamics of the DPD particles is governed by Newton’s second law: \( \mathbf{F}_i = m_i \ddot{\mathbf{r}}_i \), where \( m_i \) is the mass of the DPD particle. Particles interact via pairwise central forces consisting of a random, dissipative and conservative part: \( \mathbf{F}_{ij} = \mathbf{F}_{ij}^R + \mathbf{F}_{ij}^D + \mathbf{F}_{ij}^C \). If \( \mathbf{r}_i \) denotes the particle position, the conservative force in standard DPD reads: \( \mathbf{F}_{ij}^C = A w_C(r_{ij}) e_{ij}, \) with the repulsion parameter \( A > 0, r_{ij} = \mathbf{r}_i - \mathbf{r}_j, r_{ij} = |r_{ij}| \) and \( e_{ij} = r_{ij}/r_{ij} \). The weight function \( w_C(r) = (1 - r/r_c) \) vanishes for an inter-particle distance \( r_{ij} \) larger than a cut-off radius \( r_c \). The random force reads: \( \mathbf{F}_{ij}^R = q w^R(r_{ij}) \xi_{ij} \mathbf{r}_{ij}, \) while the dissipative force damps the relative
velocity normal to the connecting line between the particles, \( v_{ij} \cdot \mathbf{e}_{ij} = (v_i - v_j) \cdot \mathbf{e}_{ij} \), which leads to: \( F^D_{ij} = -\gamma w^D(r_{ij})(v_{ij} \cdot \mathbf{e}_{ij})e_{ij} \). The random and dissipative force act as a thermostat if the amplitudes \( q \) of the random white noise \( \xi_{ij} \) and the dissipation constant \( \gamma \) satisfy a fluctuation–dissipation theorem: \( q^2 = 2\gamma k_B T \) and \( w^R(r)^2 = w^D(r) \). The usual choice for the weight functions is \( w^R = w^C \). In this context \( k_B \) is the Boltzmann constant and \( T \) is the temperature determined by the DPD thermostat. The resulting force \( F_i \) on the particle \( i \) is the sum of all interactions with the neighbors within the cut-off range \( r_c \): \( F_i = \sum_{j \neq i} (F^R_{ij} + F^D_{ij} + F^C_{ij}) \).

Over the past years several authors have investigated and refined this numerical scheme. Español showed that the Navier–Stokes equations are correctly represented by the DPD-method [35] and also showed how energy conservation can be incorporated if needed [36]. Marsh [37] used kinetic theory in order to relate macroscopic properties of the DPD-fluid to the model parameters. As a recent development several groups [30, 32, 38] devised an extension of the simple conservative force law in order to generate more complex equations of state (EOS) showing a van der Waals loop. The latter is commonly realized by augmenting the conservative force \( F^C_{ij} \) by a density-dependent contribution. This class of schemes is termed MDPD. The density-dependent part can, e.g. be derived from a mean-field approximation to the excess free energy [38]. In an analogous manner though technically different (eventually density gradient terms must be evaluated), also LBM or DSMC can be extended to represent two-phase fluids [27, 39].

In the approach of Warren [32] adapted in this work the density-dependent contribution is rather introduced empirically, and has a different cut-off range \( r_d \):

\[
F^C_{ij} = A_{ij} w^C(r_{ij}) \mathbf{e}_{ij} + B(\bar{\rho_i} + \bar{\rho_j}) w^d(r_{ij}) \mathbf{e}_{ij}
\]

(1)

with an additional weight function \( w^d(r) = 15/(2\pi r^3_d)(1 - r/r_d)^2 \). In contrast to standard DPD, the density-independent force (with an interaction range \( r_c \)) is an attractive force, i.e. \( A_{ij} < 0 \), while \( B > 0 \) applies for the density-dependent part, rendering it repulsive. The local density \( \bar{\rho}_i \) at the location of particle \( i \) is estimated by the instantaneously weighted average \( \bar{\rho}_i = \sum_{j \neq i} w^d(r_{ij}) \) [32]. Warren showed that this approach produces stable free capillary surfaces. An appropriate choice of parameters results in sharply defined liquid–vapor (lv) interfaces that can easily be kept track of in simulations. There are virtually no particles in the vapor phase, which is thus effectively neglected. In the simulation, the numerical cost is only on the dense phase. In this respect, the MDPD approach is easier to handle than, e.g. LBM as there extra effort is required to deal with two-phase flows when the phases have significantly different densities [26]; for a solution to represent an air/water-like interface with a density ratio of 1 : 1000, see [40]. In addition to the original scheme, we introduce a species-dependent force constant \( A_{sl} \), defining the interaction between liquid (l) particles and those of the solid (s) wall in order to establish adjustable adhesive properties of the sl-interaction. All relevant quantities of the MDPD liquid (similar to those in [32]) are summarized in table 1. As in many MD schemes, in DPD dimensionless units are used, i.e. the thermal energy \( \epsilon_T = k_B T \), the length scale \( r_c \) and the mass of the DPD particles \( m_i \) are all set to unity. Reference to dimensional units can be made by defining the length scale \( r_c \), the mass of a DPD particle and the mean thermal energy \( \epsilon_T \) of a DPD particle. Different procedures for matching length and timescale have been discussed [43]. One way of matching the length scale is to match the dimensionless isothermal compressibility \( \kappa = \kappa_T / \kappa^{(0)} \), which assures that relative density fluctuations within the DPD-fluid are matched to the real liquid. Here, \( \kappa_T = -1/V (\partial V / \partial p)_{T} \) is the isothermal compressibility of the real fluid computed from the change of the volume \( V \) under variation.

New Journal of Physics 10 (2008) 043009 (http://www.njp.org/)
Table 1. Parameters used in the simulations. The surface tension has been obtained in a planar geometry, by integrating the lateral part of the pressure tensor in normal direction [41]. The viscosity $\eta$ was obtained by a separate simulation following [42]. We use a velocity Verlet algorithm for the numerical integration of the MDPD equations using a time step $\Delta t = 0.01$.

| Parameter                        | Symbol | $\mu$   |
|----------------------------------|--------|---------|
| Fluid particle density           | $\rho$ | 6.00    |
| Interaction range (attr.)        | $r_c$  | 1.0     |
| Interaction range (rep.)         | $r_d$  | 0.75    |
| Amplitude of random noise        | $q$    | 6.00    |
| Surface tension                  | $\sigma$ | 7.51 ± 0.04 |
| Viscosity                        | $\eta$ | 7.47 ± 0.05 |
| Attraction parameter             | $A_{ss} = A_{hi} = A$ | $-40.0$ |
| Repulsion parameter              | $B$    | 25.0    |

of the pressure $p$ using isothermal conditions and $\kappa^{(0)}$ is the isothermal compressibility of an ideal gas. An alternative way of matching the DPD-simulation to a real system is to calculate the system specific dimensionless numbers, e.g. the Reynolds number $Re = \rho v l / \eta$ and capillary number $Ca = \eta v / \sigma$ and to generate the corresponding situation in the DPD-simulation. In this work all results are given in model specific, i.e. DPD units, and are used for calculating the dimensionless numbers wherever needed.

3. A model for solid boundaries with adjustable adhesive properties

Modeling sl-interfaces in mesoscopic simulation approaches is particularly difficult: even on a molecular scale, the transition from a liquid to the solid is very sharp and abrupt. Naturally, the true behavior of a sl-interface cannot be represented, but it is possible to capture selected aspects phenomenologically. For instance, in LBM stick boundary conditions can be imposed by the so-called ‘back reflection’ prescription [31]. The solid boundary is represented as a mathematical plane, and particles crossing it are reflected by reversing their momentum. This principle has also been adopted in DPD approaches [44], but care must be exercised in order not to introduce numerical artefacts [45, 46]. Complications also arise if, in order to represent wetting phenomena [41], also the boundaries are constructed from particle assemblies, analogous to MD [47]. A dense impenetrable particle configuration may lead to spurious density oscillations [48] or artificial slip [49].

An elegant and simple solution for flat interfaces has been presented by Merabia and Pagonabarraga only recently [50]. They replace the solid by yet another liquid phase with similar properties. For both phases, bounce-back conditions apply at the boundary plane so that they remain separate. Otherwise, the inter-particle interaction is left unchanged even across the boundary. This way, distortions in the fluid phase directly at the sl-interface are avoided. With this system they investigated the spreading of droplets on solid boundaries, whereas they did not investigate the behavior under forced wetting as considered in the next section of this work.
Figure 1. Density distribution for a mutual attraction strength $A_{\text{sl}} = -40.0$ within a slit of full width $b = 20 \sigma$ (only the right half of the slit is depicted). A rigid wall with an fcc arrangement of particles (dotted dashed line) shows spurious density oscillations. The ‘thermalized’ wall (solid line) clearly reduces density oscillations and maintains a constant temperature at a preset value of $T = 1$.

Similar in spirit, we have recently suggested a wall model that provides a smooth transition from the liquid to the solid with respect to the forces specified in equation (1) [51]. The main difference in our approach is that the solid is not defined by a mathematical description (e.g. a predefined plane), but instead consists of an ensemble of particles, which are pinned to the initial position via a harmonic spring which largely enhances the possibility to create complicated channel geometries or textured surfaces. For this to be achieved, we allow for a thin diffuse interface between both phases to develop, and we shall now briefly summarize the most important aspects. The wall regions consist of an amorphous configuration of particles at the same density as in the liquid. The particles are attached to fixed sites by harmonic forces, while the inter-particle interaction is a ‘normal’ liquid–liquid interaction given by equation (1) ($A_{\text{ss}} = A_{\text{ll}} = -40.00$). Thus, in our approach solid particles oscillate around the initial position over time leading to thermal roughness. In order to prevent particles of the liquid phase from diffusing into the wall indefinitely, an additional soft repulsive force normal to the wall surface is applied. Since the solid phase is made of DPD particles of the same density as the liquid phase a smooth transition from the liquid to the solid, see figure 1, is established with constant density of the dissipative particles in the liquid region. In contrast to a rigid wall, spurious density oscillations as well as temperature oscillations near the wall can essentially be reduced. Figure 1 illustrates the difference between a rigid wall (with a (100)-fcc arrangement of the wall particles) and the ‘thermalized’ wall presented here. In summary, the wall model used here provides a numerically simple scheme avoiding local distortions of liquid properties at the sl-interface.

Note that the only property imposed on the wall model is the affinity of the solid surface, governed by the parameter $A_{\text{sl}}$, which is larger than $A_{\text{ll}}$ and $A_{\text{ss}}$ ($A_{\text{sl}} \geq A_{\text{ll}}, A_{\text{sl}} \geq A_{\text{ss}}$). Qualitatively this is similar to the methodology in LBM, where an attractive external potential is used to tune the interfacial free energy in a van der Waals/mean field framework [52, 53].
In a pure liquid, a value $A_{sl} > -40.0$ would lead to a separation into immiscible phases (see also the work of Pagonabarraga and Frenkel on mixtures of non-ideal fluids [30] or Diaz-Herrera et al [54], who also achieve a broad miscibility gap in a binary mixture of Lennard–Jones fluids by varying the attractive part of the interaction). In the boundary model described above the choice of $A_{sl}$ thus assists in shaping the sl-interface accordingly. Interestingly, even at large $A_{sl}$, when the repulsive (i.e. hydrophobic) interaction dominates, still no particle layering develops. This seems similar to the behavior of water close to hydrophobic surfaces, as studied by Grigera [55], where also a very smooth transition from the bulk density to zero directly at the interface is found. In that situation, it is believed that the depletion of liquid originates on account of entropic reasons, due to a reorganization of hydrogen bonds [56]. It might be of some interest to consider generic interface models as developed here in order to explore under which other conditions layering/depletion may arise (see also the discussion in [32] on layering at a free capillary surface).

There is a rather simple dependence of the static contact angle on the parameter $A_{sl}$. The set-up for extracting the static contact angle is illustrated in figure 2. The meniscus was segmented into 20 slices, and the center of mass was determined for each slab separately in order to extract the shape of the meniscus and thus the static contact angle $\theta_0$. After achieving stationary conditions the center of mass was averaged 10 000 time steps for each slice separately in order to determine the static contact angle. Figure 3 shows the static contact angle as a function of the attraction parameter $A_{sl}$. The wetting behavior of the wall can be tuned arbitrarily between hydrophilic and hydrophobic by changing the mutual attraction $A_{sl}$. Furthermore, the resulting static contact angle is consistent with the expected value given by Young’s law: $\theta_0 = \arccos((\sigma_{sv} - \sigma_{sl})/\sigma_{lv})$, with $\sigma_{sv}$, $\sigma_{sl}$, $\sigma_{lv}$ being the interfacial tensions of the solid–vapor (sv), sl- and lv-interface. The surface tensions were measured independently.

Figure 2. Scheme of the set-up used for determining the static contact angle. The center of mass in each slab is determined separately and averaged over 1000 time steps in order to determine the shape of the meniscus.
in a planar geometry, by integrating the lateral part of the pressure tensor $p$ in normal, i.e. $z$-direction, to the interface, $\sigma_{ij} = \int \left( p_{zz} - \frac{1}{2} (p_{xx} + p_{yy}) - \phi \right) \, dz$, ($\sigma_{ij} \in \{ \sigma_{sl}, \sigma_{sv}, \sigma_{lv} \}$) as described in detail in [33]. $\phi$ as defined by Nijmeijer et al. [47] accounts for the contribution of an external potential. In our case this is the soft repulsive force preventing particles of the liquid phase from diffusing into the wall over time. The contribution to the surface tension is less than 5% for all cases of $A_{sl}$. Within the numerical accuracy the two data sets agree implying that the ‘measured’ contact angle $\theta_0$ is in agreement with the thermodynamic prediction.

4. Behaviour of dynamic contact angles under forced flow in a thin slit

In the following, we demonstrate that the DPD scheme provides a global solution for the moving CL-problem, which can be interpreted in terms of effective continuum theories [4]. For investigating the dynamic contact angle behavior, we consider a situation of driven capillary flow in a thin slit, similar to the experiments of Hoffman [57]. This is termed here as forced wetting, as the liquid front is made to move with constant velocity across the solid without changing shape, the velocity is determined by the piston. This facilitates experimental but also the theoretical or numerical analysis, since one is essentially dealing with a stationary situation, in contrast to many natural wetting phenomena such as droplet spreading, where the time varying shape of the capillary meniscus must be taken into account.

For the channel system, we generate a stationary situation as illustrated in figure 4(a). The liquid is placed between two plates that are moved with a relative velocity $v_{\text{wall}}$, while the plug is kept fixed by a piston. In the reference frame of the liquid the walls move with velocity $-v_{\text{wall}}$ relative to a piston that keeps the liquid in place. $v_{\text{wall}}$ determines the capillary number $Ca = \eta v_{\text{wall}}/\sigma$. We may thus extract the shape of the meniscus by the procedure described above for the static case, see figure 2, and obtain a relation $\theta_d(Ca)$. Note that this procedure will in general yield an apparent dynamic contact angle $\theta_d$, as we do not measure the local inclination of the meniscus with the boundary and the data points at the interface are not considered for the

Figure 3. Comparison of Young’s law (circles) and the result of the MDPD-simulation (squares). Vertical axis in degrees.
Figure 4. (a) Schematic picture of the set-up used for studying forced wetting in a capillary slit. (b) Shows the flow field resulting from the MDPD-simulation for $\theta_0 = 0^\circ$ and $Ca = 0.05$. This value corresponds to a dynamic contact angle of $\theta_d = 50^\circ$. (c) Shows the stream function solution according to Cox [4]. The continuum model predicts pronounced convection (rolling) next to the CL in agreement with experimental observations [8] and the numerical result obtained from the MDPD-simulation.

The slip occurs in combination with pronounced caterpillar motion of the fluid, which has been demonstrated in an experiment by Dussan [8]. Figure 4(c) shows the streamlines as obtained from solving the Navier–Stokes equations for an infinite fluid wedge, with constant velocity at the free surface and a no-slip condition at the sl-interface:

$$\psi(\phi, r) = r \left((C_A \phi + D_A) \cos(\phi) + (E_A \phi + F_A) \sin(\phi)\right), \quad (2)$$

with the origin of the reference frame being the position of the CL. The coefficients $C_A$, $D_A$, $E_A$ and $F_A$ are given in [4]. Equation (2) leads to a logarithmic singularity of the viscous stress as $r \to 0$. In contrast figure 4(b) shows that the DPD-method provides slip as a mechanism for removing the stress singularity.

According to Cox [4] the singularity can be avoided by identifying at least two distinct regions: (i) a region close to the CL, where slip predominates the flow and (ii) a no-slip region
Figure 5. Comparison of the constitutive law for dynamic contact angles according to Cox [4] (equation (4)) and the corresponding characteristics obtained by the MDPD-simulations. Vertical axis in rad$^3$.

(and (iii) sometimes an intermediate region is also needed). The asymptotic matching of the different regions provides a solution for the constitutive law of the dynamic contact angle to the leading order of $Ca$ [4]:

$$g(\theta_d) - g(\theta_0) = Ca \ln(\epsilon^{-1})$$

with the ratio $\epsilon = L_{\text{micro}}/L_{\text{macro}}$ independent of $Ca$ and $g(\theta) = \int_0^\theta \frac{x \sin(x) \cos(x)}{2 \sin(x)} \, dx$. $L_{\text{micro}}$ is a microscopic length scale over which a slip boundary condition is assumed to dominate the flow field. $L_{\text{macro}}$ signifies a macroscopic length. By studying forced wetting of silicone oils in thin glass tubes, Hoffman [57] found empirically a relationship that almost quantitatively agrees with equation (3) and that complies with the Taylor expansion of equation (3) for small angles ($\theta_d < 100^\circ$) [9]:

$$\theta_d^3 - \theta_0^3 = 9 \, Ca \ln(\epsilon^{-1}).$$

In the case of partial wetting, see figure 5, the data sets can well be fitted to equation (4). If we take $9 \cdot \ln(\epsilon^{-1})$ as a compound fit parameter we find $9 \cdot \ln(\epsilon^{-1}) = 20.8 \pm 0.3$.

In contrast, the case of complete wetting has a more complex characteristics, see figure 6. Although for $Ca > 0.04$ the slope attained is comparable to the case of partial wetting, a significant deviation from the linear behavior occurs for small $Ca$. Eggers [9] pointed out that in the case of $\theta_0 = 0^\circ$ the microscopic length $L_{\text{micro}}$ actually depends on $Ca$. Eggers showed that the solution of the lubrication equation yields an additional dependency such that $L_{\text{micro}} = 0.54 \lambda Ca^{-1/3}$, if slip is assumed as the mechanism responsible for removing the stress singularity. In this context $\lambda$ is the Navier slip length such that, $v - v_{\text{wall}} = \lambda \cdot (\partial v/\partial x)$, if $v$ is the fluid velocity at the wall and $v_{\text{wall}}$ is the velocity of the moving boundary. This leads to the constitutive law for $\theta_0 = 0^\circ$:

$$\theta_d^3 = 9 \, Ca \ln(\alpha Ca^{1/3}),$$

New Journal of Physics 10 (2008) 043009 (http://www.njp.org/)
Figure 6. Comparison of the constitutive law for dynamic contact angles according to Eggers [9] (equation (5)) and the corresponding characteristics obtained by the MDPD-simulations. For reference the model of Cox [4] is depicted. Vertical axis in rad$^3$.

where $\alpha = L_{\text{macro}}/(0.54 \lambda)$. The additional weak logarithmic dependency only becomes relevant for small Capillary numbers. We compare the constitutive law, i.e. equation (5), with the result obtained by the MDPD-method. Figure 6 shows that within the numerical accuracy the MDPD-method reproduces the behavior given by equation (5) for $\theta_0 = 0^\circ$. The numerical value obtained by the fit is $\alpha = (16.7 \pm 0.3)$.

We should stress that all that really is required for the exponent of 1/3 to apply is the absence of an extended precursor foot, present only for strongly wetting systems with a large excess interfacial free energy. A numerical value of 1/3 is not strictly related to the appearance of slip, but instead may originate from some other mechanism, that removes the stress singularity. In this context the diffuse nature of the sl-interface (see also for use of diffuse Cahn–Hillard interfaces [58]) has been pointed out as one possible mechanism to avoid a stress singularity. Furthermore, it must be expected that along the slip region, especially at the foremost liquid front, the interface needs some time to form, which yet would be a refinement of a diffuse-interface mechanism. As a matter of fact, a possible interface relaxation plays a major role in an elaborate continuum theory put forward by Shikhmurzaev [10, 59]. Exploring the present boundary model along these lines further could be of considerable interest.

5. Dynamic capillary impregnation

Finally, we address the wetting behavior of a transient problem, the capillary impregnation of a slit pore. The asymptotic limit ($t \rightarrow \infty$) of this problem can effectively be described by a one-dimensional equation, the so-called Washburn equation [60], which describes the penetration depth $z$ over time $t$, i.e. $z(t) \propto \sqrt{t}$. This behavior has been confirmed experimentally over a wide range of time and length scales [60]–[62] down to the nano-scale. Also MD simulations have shown that the characteristics can be described in the framework of a one-dimensional
continuum description [63, 64] even for short time and length scales. We pursue this path by studying a system as shown in figure 7. We consider the case where $\theta_0 = 0^\circ$, i.e. $A_{sl} = -40.0$. The asymptotic limit for $t \to \infty$ cannot directly be applied since inertial effects and the presence of the dynamic contact angle, i.e. $\theta_d \neq \theta_0$, alter the dynamics of the wetting process.

Before investigating the dynamics of the impregnation, the analytical solution of this problem is derived by extending the Washburn law [60] in order to account for the dynamic contact angle contribution and inertia. The dynamics of the capillary impregnation is governed by balancing capillary forces $F_{cap}$, viscous forces $F_\eta$ and inertial forces due to the momentum change $d\rho(t)/dt$ generated by the mass $M(t)$ which is moved in the $z$-direction during the process of impregnation:

$$\frac{dp(t)}{dt} = F_{cap} - F_\eta. \quad (6)$$

Assuming that the energy is dissipated in the capillary equation (6) gives:

$$\frac{1}{bh} \frac{dp(t)}{dt} = \frac{2\sigma_{sl}}{b} \cos(\theta_d) - \frac{12\eta}{b^2} \frac{dz(t)}{dt}, \quad (7)$$

where $b$ denotes the slit width, see figure 7, $h$ denotes the width in the $y$-direction ($12r_c$) and $z$ is the penetration depth into the pore. Equation (7) corresponds to the Washburn law [60] (adapted to a slit) if inertia is set to zero, i.e. $dp(t)/dt = 0$, and the dynamic contact angle is replaced by the static contact angle, i.e. $\theta_d = \theta_0$. This corresponds to the special case in the asymptotic limit $t \to \infty$. The momentum change in $z$-direction can directly be expressed in terms of the momentum of each particle, $d\rho(t)/dt = d(M_{sl}(z(t))/dt = d(\rho bh z(t)/dt(t))/dt$, while the momentum change in the reservoir is more difficult to determine (due to the complex flow field) and therefore was measured during the course of the simulation and used as input to equation (7). Also the dynamic contact angle $\theta_d$ was monitored during the simulation. Figure 8 shows the observed dynamic contact angle over time during the MDPD-simulation. During the timescale studied here the resulting dynamic contact angle clearly exceeds the

Figure 7. Set-up for capillary impregnation. The meniscus penetrates in the $z$-direction due to the driving capillary pressure. Periodic boundary conditions are applied in the $y$-direction and in the $x$-direction. The shape of the meniscus and the penetration depth is monitored by segmenting the pore into 20 slabs and evaluating the center of mass in each slab separately as in figure 2.
Figure 8. Dynamic contact angle behavior $\theta_d$ obtained by the MDPD-simulation versus time during the capillary filling process. Vertical axis in degrees.

Figure 9. Dynamics of the penetration into the pore over time $z(t)$ obtained by the MDPD-simulation compared to equation (7). The dynamic contact angle is measured online during the simulation and inserted into equation (7) for the numerical integration.

static contact angle ($\theta_0 = 0^\circ$). Over time the dynamic contact angle slowly decreases as the penetration velocity into the slit decreases, see figure 9. Interestingly, the inertial timescale [64] $\tau = \sqrt{\rho b^3/\sigma} \approx 80$ roughly corresponds to the timescale needed for the meniscus to relax from its initial configuration (starting conditions corresponds to the static contact angle $\theta_0 = 0^\circ$). Since inertia has a considerable influence on the dynamics of the impregnation, the dynamic contact angle is directly extracted from the simulation and inserted into equation (7) in order to check if the MDPD-simulation can be consistently described by equation (7).
Integrating equation (7) numerically with the data set of figure 8 results in excellent quantitative agreement with the MDPD-simulation, see figure 9. As a reference in figure 9 the Washburn law \([60]\) is shown, which clearly overestimates the penetration dynamics as inertia and the reduction of the capillary pressure due to the dynamic contact angle are missing.

6. Summary and conclusions

We presented a study on capillary wetting under static as well as dynamic conditions in the framework of DPD. The measured static contact angle is in agreement with Young’s law. The characteristics of the dynamic contact angle was investigated under stationary conditions and compared to the hydrodynamic models of Cox [4] and Eggers [9], respectively. The dynamic contact angle is in agreement with the theoretical prediction of Eggers [9] for a static contact angle \(\theta_0 = 0^\circ\) with an additional logarithmic dependency on \(Ca\) compared to the model of Cox [4]. We find that for a static contact angle \(\theta_0 > 0^\circ\) the microscopic length \(L_{\text{micro}}\) is independent of the capillary number in agreement with theory [1, 4, 9]. As a last step a transient problem, the wetting into a slit was studied. The dynamics can be described by a simple approach balancing capillary and viscous forces under consideration of inertia. This analytical solution was compared to the result of the MDPD-simulation resulting in good quantitative agreement. In summary, we have shown that the MDPD-method can successfully be applied for studying dynamic stationary as well as transient capillary wetting phenomena. The MDPD-approach therefore has the potential to become a valuable explorative tool as an alternative or supplement to CFD or analytical approaches.

Acknowledgment

We gratefully acknowledge the financial support of the Deutsche Forschungsgemeinschaft (DFG) within the priority program SPP 1164.

References

[1] de Gennes P G 1985 Wetting: statics and dynamics Rev. Mod. Phys. 57 827–63
[2] Herminghaus S et al 2000 J. Phys.: Condens. Matter 12 A57
[3] Oron A, Davis S H and Bankoff S G 1997 Long-scale evolution of thin liquid films Rev. Mod. Phys. 69 931–80
[4] Cox R G 1986 The dynamics of the spreading of liquids on a solid-surface 1. Viscous-flow J. Fluid Mech. 168 169–94
[5] Blake T D, Bracke M and Shikhmurzaev Y D 1999 Experimental evidence of nonlocal hydrodynamic influence on the dynamic contact angle Phys. Fluids 11 1995–2007
[6] Wilson M C T, Summers J L, Shikhmurzaev Y D, Clarke A and Blake T D 2006 Nonlocal hydrodynamic influence on the dynamic contact angle: slip models versus experiment Phys. Rev. E 73 041606
[7] Huh C and Scriven L E 1971 Hydrodynamic model of steady movement of a solid/liquid/fluid contact line J. Colloid Interface Sci. 35 85
[8] Dussan E B 1979 Spreading of liquids on solid-surfaces—static and dynamic contact lines Annu. Rev. Fluid Mech. 11 371–400
[9] Eggers J and Stone H A 2004 Characteristic lengths at moving contact lines for a perfectly wetting fluid: the influence of speed on the dynamic contact angle J. Fluid Mech. 505 309–21
[10] Shikhmurzaev Y D 1997 Moving contact lines in liquid/liquid/solid systems J. Fluid Mech. 334 211–49

New Journal of Physics 10 (2008) 043009 (http://www.njp.org/)
[11] Blake T D and Haynes J M 1969 Kinetics of liquid/liquid displacement J. Colloid Interface Sci. 30 421
[12] Oshanin G, de Coninck J, Cazabat A M and Moreau M 1998 Dewetting, partial wetting, and spreading of a two-dimensional monolayer on solid surface Phys. Rev. E 58 R20–23
[13] de Ruijter M J, Blake T D and De Coninck J 1999 Dynamic wetting studied by molecular modeling simulations of droplet spreading Langmuir 15 7836–47
[14] de Ruijter M J, Charlot M, Voue M and De Coninck J 2000 Experimental evidence of several time scales in drop spreading Langmuir 16 2363–8
[15] Tanner L H 1979 Spreading of silicone oil drops on horizontal surfaces J. Phys. D: Appl. Phys. 12 1473
[16] Hocking L M 1992 Rival contact-angle models and the spreading of drops J. Fluid Mech. 239 671–81
[17] Hocking L M and Rivers A D 1982 The spreading of a drop by capillary action J. Fluid Mech. 121 425–42
[18] Cox R G 1998 Inertial and viscous effects on dynamic contact angles J. Fluid Mech. 357 249–78
[19] Eggers J 2004 Toward a description of contact line motion at higher capillary number Phys. Fluids 16 3491–94
[20] Thompson P A and Robbins M O 1989 Simulations of contact-line motion-slip and the dynamic contact-angle Phys. Rev. Lett. 63 766–9
[21] Thompson P A and Robbins M O 1990 Shear flow near solids: epitaxial order and flow boundary conditions Phys. Rev. A 41 6830
[22] Gentner F, Ogonowski G and De Coninck J 2003 Forced wetting dynamics: a molecular dynamics study Langmuir 19 3996–4003
[23] Hadjiconstantinou N G 1999 Combining atomistic and continuum simulations of contact-line motion Phys. Rev. E 59 2475–8
[24] Qian T, Wang X-P and Sheng P 2003 Molecular scale contact line hydrodynamics of immiscible flow Phys. Rev. E 68 016306
[25] De Fabritiis G, Delgado-Buscalioni R and Coveney P V 2006 Multiscale modeling of liquids with molecular specificity Phys. Rev. Lett. 97 134501
[26] Sbragaglia M, Benzi R, Biferale L, Succi S, Sugiyama K and Toschi F 2007 Generalized lattice Boltzmann method with multirange pseudopotential Phys. Rev. E 75 026702
[27] Bastea S and Lebowitz J L 1997 Spinodal decomposition in binary gases Phys. Rev. Lett. 78 3499–502
[28] Hadjiconstantinou N G, Garcia A L and Alder B J 2000 The surface properties of a van der Waals fluid Physica A 281 337–47
[29] Hoogerbrugge P J and Koelman J M V A 1992 Simulating microscopic hydrodynamic phenomena with dissipative particle dynamics Europhys. Lett. 19 155–60
[30] Pagonabarraga I and Frenkel D 2001 Dissipative particle dynamics for interacting systems J. Chem. Phys. 115 5015–26
[31] Latva-Kokko M and Rothman D H 2007 Scaling of dynamic contact angles in a lattice-Boltzmann model Phys. Rev. Lett. 98 254503
[32] Warren P B 2003 Vapor-liquid coexistence in many-body dissipative particle dynamics Phys. Rev. E 68 066702
[33] Groot R D and Warren P B 1997 Dissipative particle dynamics: bridging the gap between atomistic and mesoscopic simulation J. Chem. Phys. 107 4423–35
[34] Espanol P, Serrano M and Zuniga I 1997 Coarse-graining of a fluid and its relation with dissipative particle dynamics and smoothed particle dynamics Int. J. Mod. Phys. C 8 899–908
[35] Espanol P 1995 Hydrodynamics from dissipative particle dynamics Phys. Rev. E 52 1734–42
[36] Espanol P 1997 Dissipative particle dynamics with energy conservation Europhys. Lett. 40 631–6
[37] Marsh C A, Backx G and Ernst M H 1997 Fokker–Planck–Boltzmann equation for dissipative particle dynamics Europhys. Lett. 38 411–5
[38] Trofimov S Y, Nies E L F and Michels M A J 2002 Thermodynamic consistency in dissipative particle dynamics simulations of strongly nonideal liquids and liquid mixtures J. Chem. Phys. 117 9383–94
Swift M R, Osborn W R and Yeomans J M 1995 Lattice Boltzmann simulation of nonideal fluids Phys. Rev. Lett. 75 830–3

Inamuro T, Ogata T, Tagajima S and Konishi N 2004 A lattice Boltzmann method for incompressible two-phase flows with large density differences J. Comp. Phys. 198 628–44

Jones J L, Lal M, Ruddock J N and Spenley N A 1999 Dynamics of a drop at a liquid/solid interface in simple shear fields: a mesoscopic simulation study Faraday Discuss. 129 129–42

Backer J A, Lowe C P, Hoefsloot H C J and Iedema P D 2005 Poiseuille flow to measure the viscosity of particle model fluids J. Chem. Phys. 122 154503

Lowe C P 2004 Simulating the dynamics of mesoscopic systems Novel Methods in Soft Matter Simulations (Lecture Notes in Physics vol 640) (Berlin: Springer) pp 39–68

Revenga M, Zuniga I, Espanol P and Pagonabarraga I 1998 Boundary models in DPD Int. J. Mod. Phys. C 9 1319–28

Revenga M, Zuniga I and Espanol P 1999 Boundary conditions in dissipative particle dynamics Comp. Phys. Commun. 121–122 309–11

Visser D C, Hoefsloot H C J and Iedema P D 2005 Comprehensive boundary method for solid walls in dissipative particle dynamics J. Comp. Phys. 205 626–39

Nijmeijer M J P, Bruin C, Bakker A F and van Leeuwen J M J 1990 Wetting and drying of an inert wall by a fluid in a molecular-dynamics simulation Phys. Rev. A 42 6052–9

Pivkin I and Karniadakis G E 2006 Phys. Rev. Lett. 96 206001

Pivkin I V and Karniadakis G E 2005 A new method to impose no-slip boundary conditions in dissipative particle dynamics J. Comp. Phys. 207 114–28

Merabia S and Pagonabarraga I 2006 A mesoscopic model for (de)wetting Eur. Phys. J. E 20 209–14

Henrich B, Cupelli C, Moseler M and Santer M 2007 An adhesive DPD wall model for dynamic wetting Europhys. Lett. 80 60004

Zhang J and Kwok D Y 2004 Apparent slip over a solid-liquid interface with a no-slip boundary condition Phys. Rev. E 70 056701

Davies A R, Summers J L and Wilson M C T 2006 On a dynamic wetting model for the finite-density multiphase lattice Boltzmann method Int. J. Comput. Fluid Dyn. 20 415–25

Díaz-Herrera E, Moreno-Razo J A and Ramirez-Santiago G 2004 Wetting phenomenon in the liquid–vapor phase coexistence of a partially miscible Lennard-Jones binary mixture Phys. Rev. E 70 051601

Grigera J R, Kalko S G and Fischbarg J 1996 Wall–water interface. A molecular dynamics study Langmuir 12 154–8

Chandler D 2005 Interfaces and the driving force of hydrophobic assembly Nature 437 640–7

Hoffman R L 1975 A study of the advancing interface. I. Interface shape in liquid gas systems J. Colloid Interface Sci. 50 801–17

Seppacher P 1996 Moving contact lines in the Cahn–Hilliard theory Int. J. Eng. Sci. 34 977

Shikhmurzaev Y D 1993 Int. J. Multiphase Flow 19 589–610

Washburn E W 1921 The dynamics of capillary flow Phys. Rev. 17 273–83

Tas N R 2004 Capillary filling speed of water in nanochannels Appl. Phys. Lett. 85 3274

Quere D 1997 Inertial capillarity Europhys. Lett. 39 533–538

Martic G, Gentner F, Seveno D, Coulon D, De Coninck J and Blake T D 2002 A molecular dynamics simulation of capillary imbibition Langmuir 18 7496–502

Dimitrov D I, Milchev A and Binder K 2007 Capillary rise in nanopores: molecular dynamics evidence for the Lucas–Washburn equation Phys. Rev. Lett. 99 054501