Dewetting of thin polymer films: Influence of interface evolution

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received 7 April 2009; accepted 22 April 2009
published online 20 May 2009

PACS 68.60.-p - Physical properties of thin films, nonelectronic
PACS 68.15.+e - Liquid thin films
PACS 83.10.-y - Rheology: Fundamentals and theoretical

Abstract – The dewetting dynamics of ultrathin polymer films, e.g., in the model system of polystyrene on a polydimethylsiloxane-covered substrate, exhibits interesting behavior like a fast decay of the dewetting velocity and a maximum in the width of the built-up rim in the course of time. These features have been recently ascribed to the relaxation of residual stresses in the film that stem from the nonequilibrium preparation of the samples. Recent experiments by Coppée et al. on PS with low molecular weight, where such stresses could not be evidenced, showed however similar behavior. By scaling arguments and numerical solution of a thin-film viscoelastic model we show that the maximum in the width of the rim can be caused by a temporal evolution of the friction coefficient (or equivalently of the slip length), for which we discuss two possible mechanisms. In addition, the maximum in the width is affected by the sample age. As a consequence, knowing the temporal behavior of friction (or slip length) in principle allows to measure the aging dynamics of a polymer-polymer interface by simple dewetting experiments.

Introduction. – Dewetting experiments [1–3] have been proven to be a simple tool to investigate thin polymer films on various substrates [4,5]. Especially the model system of the incompatible polymers polystyrene (PS) on polydimethylsiloxane (PDMS) grafted on a silicium substrate has attracted numerous experimental investigations. There is consensus that friction and slippage at the interface, as well as the viscoelasticity of the dewetting film are crucial to understand the experiments [6,7]. To account for the occurrence of a maximum in the width of the rim in the course of time, and to interpret the effects of sample aging on this peculiar feature of the thin films’ dewetting dynamics, residual stresses in the PS film have been shown to be important [8–10]. These stresses are supposed to stem from the sample preparation process (usually spin-coating) and represent an additional degree of freedom that can (partially) relax during dewetting. For high enough molecular weights, residual stresses have been clearly evidenced in various systems [8,11,12]. However, their relaxation well below the glass temperature, needed to interpret the effects of aging [8], is currently debated. An additional puzzle is the fact that for films of low molecular weight no such relaxation of stress could be evidenced, while a maximum in the rim width and its dependence on aging have still been found. Very recently, in parallel to dewetting experiments the interface between the two polymer species has been studied by neutron reflectometry [13] and interdiffusion of the PS-PDMS interface has been found, for samples aged well below the PS bulk glass temperature.

In this letter we show by scaling arguments and demonstrate by numerical solution of a thin-film model that a slow increase in the friction coefficient or, equivalently, a decrease of the slip length can give rise to a maximum in the rim width. This evolution of friction/slip might be due to several mechanisms: first, as suggested by the neutron reflectometry measurements [13], a slow roughening of the film-substrate (PS-PDMS) interface with concomitant increase in friction may be responsible. A second possibility would be that (for the lowest PDMS grafting densities) a small number of PS chains may enter the PDMS brush and become attached to the substrate. It is known that minute amounts of such “connectors” [14–16] already decrease the slip length.

A second finding is that the value of the maximum rim width decreases monotonously with the friction coefficient at the beginning of the dewetting process. This could explain the experimentally observed decrease of the

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maximum with the sample age. Thus knowing the evolution of the friction coefficient allows in principle to relate the maximum width to the aging history. As studied in refs. [8,9,17,18], a similar connection between rim width and aging time can also be caused by relaxing internal residual stresses. One would expect that stress relaxation is the dominating mechanism for high-molecular-weight films, while for low-molecular-weight films interface changes are responsible. In general, the two nontrivial processes, stress relaxation in bulk and evolution of the interface, might be present at a time, which leaves interpretation of experimental results an intricate task. The specific dynamics of the PDMS brush-PS melt interface being unknown, drawing conclusions from the dependence of the rim width on the sample aging, as suggested above, is only a rough estimate at present. Dynamic self-consistent field techniques or molecular simulations would be needed to establish this dynamics.

What processes can lead to a maximum in the rim width? A scaling argument. – First we want to give a simple argument, why a slow evolution of the interface makes the occurrence of a maximum in the rim width possible. For dewetting of a viscoelastic film, scaling arguments have been used to qualitatively understand the dynamics [17,19]. The balance of the work done by the driving force and the dissipation by friction reads approximately \( |S|V(t) \sim \zeta W(t)V(t)^2 \) or

\[
\frac{|S|}{\zeta} \approx W(t)V(t). \tag{1}
\]

Here \( V(t) \) is the velocity at the dewetting edge and \( W(t) \) is the width of the rim. The driving force acting on the rim is the negative of the spreading parameter \( S = \gamma_{sv} - \gamma_{sl} - \gamma \) (where the \( \gamma \)'s are the interface energies for the substrate-vapor, substrate-liquid and liquid-vapor interfaces, respectively), and \( \zeta \) is the friction coefficient associated to the polymer-polymer interface. In general, the friction will be nonlinear (see below), but for the simple argument given here this is not of importance. In addition to this balance, one needs the fact that the driving force is monotonously decreasing —there is no mechanism, and also no experimental evidence, for a speed-up of the dewetting process.

What possibilities does eq. (1) allow for \( W(t) \) to have a maximum? Clearly, if \( W \) has passed through a maximum, it has to decrease for some time simultaneously with the monotonously decreasing velocity. This is not possible for eq. (1) if the left-hand side is a constant. A necessary condition for a maximum in the width is thus that the effective driving force \( |S|/\zeta \) decreases in the course of time. There are several possibilities to achieve such a decrease: The first one is motivated by residual stresses stemming from spin-coating. Introduction of a residual stress \( \sigma(t) \), that is allowed to relax in the course of time, leads to a renormalization of the driving force \( |S| \to |S| + h_0\sigma(t) \) [17,18] (where \( h_0 \) is the film thickness). Accordingly, upon stress relaxation the driving force decreases.

The second possibility, suggested by the interface evolution evidenced in ref. [13], is an increase in the friction coefficient \( \zeta \to \zeta(t) \) due to roughening of the liquid-substrate interface. Additional to the increase in friction, interface roughening also leads to a decrease of the driving force \( |S| \), as discussed for autophobic dewetting in [20], and can even lead to a cross-over to wetting in that case. For the incompatible system PS-PDMS, where the PS-air surface energy is dominating, this latter effect is however not probable for the expected microscopically small roughnesses.

A third possibility, also associated to tiny changes in the interface, is the successive attachment of a few PS chains to the substrate where the PDMS-coverage is low or defective. This process leads to a decrease in the driving force \( |S| \to |S| - \nu(t)I_f \), where \( \nu \) is the areal density of such “connector” chains, \( t \) the length they are stretched upon pull-out and \( f \) the pull-out force [14,15]. Additionally the friction increases due to the connectors (in simplest approximation \( \zeta \to \zeta + \nu(t)\kappa \) with \( \kappa \) the friction coefficient of connectors; for a more accurate treatment, see [21]).

In the following we would like to exemplify the second case further. We first give an estimate of how the interface evolution gives rise to an increasing friction coefficient. Later on this is implemented in a thin-film model and its influence on the dewetting is studied.

Interface evolution. – Polymer-polymer interfaces are well studied [22,23], although mostly at equilibrium. The equilibrium width of an immiscible interface is given by \( \delta \propto a/\sqrt{N} \) [24], where \( a \) is the Kuhn length and \( \chi \) is the Flory interaction parameter. A simple scaling argument [25], consistent also with a full statistical treatment [26], states that the slip length \( b \) at the interface is given by the ratio of bulk and interfacial viscosity and scales as \( b \propto N/\delta^2 \). The factor \( N \) is the degree of polymerization and leads to the well-known large slip lengths for polymers [27]. Alternatively, the slippage length can be defined as \( b = \eta/\zeta \), implying that the friction coefficient scales like \( \zeta \propto \delta^2 \). This scaling should hold for the interface between two immiscible melts, for a brush-melt interface the situation is slightly different. The structure of brushes in various matrix polymers has been studied in ref. [28], and the width for the immiscible case has been found to be narrow, but larger than the width between melts. The effect of the grafting density on the interface width is unclear, but one would expect that the grafting density of the PDMS-brush should not be too high to prevent total stretching and allow for mobility of the brush chains. Indeed, as reported in ref. [13], roughening was pronounced only for the lowest grafting densities where the interface width increased from 1 to 4 nm in the course of time. It should also be mentioned that polymer interfaces display thermally activated capillary waves [29], which lead to a higher apparent interface width.

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As far as the dynamics is concerned, the evolution of the interface towards its equilibrium value due to chain interdiffusion and its influence on the dewetting via the slip length or friction is difficult to determine. Strictly speaking it requires to solve a hydrodynamic problem for a system with an evolving boundary, to be determined by a model for an immiscible polymer brush-polymer melt interface and which in turn will be influenced by the flow. This scope clearly demands extensive microscopic modeling and we rather use a simple estimate, motivated by experiments. Interdiffusion dynamics for the symmetric case has been studied for both a melt and a network of PS on a PS-brush. The interface width has been found to change by a factor of 3-4 on the time scale (the time $t^*$) and displayed a logarithmic behavior in time. For the immiscible system PS-PMMA, also a logarithmic growth in time has been found [32], such as observed for rubber-brush friction [33]. The exponent $\alpha$ of eq. (2) is empirical and a microscopic foundation would be desirable. However, as shown below, the effects of an increasing friction on dewetting turn out to be very robust with respect to variations of the detailed form of $\zeta(t)$.

**Modeling.** In order to demonstrate that a time evolution of interfacial properties indeed can give rise to a maximum in the rim width and a concomitant dependence on sample age, we performed a numerical study for the model established recently in refs. [9,17,18]. This model is a lubrication approximation for a Jeffrey-type viscoelastic fluid film on slippery substrate. As motivated by recent experiments on polymer-polymer friction [33], nonlinear friction with the substrate is considered. In the model, one accounts for three fields: the velocity in the film $v(x,t)$, the stress field $\sigma(x,t)$, and the height profile $h(x,t)$ (assumed to be small compared to the slip length). A simple edge geometry is considered, where the edge of the film is initially at $x=0$ and the dewetting occurs in the $+x$-direction. As governing equations one has the momentum equation (3), a Jeffrey-type constitutive law (4), and volume conservation (5):

$$ \frac{\partial^* v}{\partial t} = \partial_x (h \sigma),$$

$$ \sigma + \tau_1 \partial_x \sigma = G\tau_1 [\partial_x v] + \tau_0 \partial_t [\partial_x v],$$

$$ \partial_t h = -\partial_x (vh).$$

**Equation (3)** is the balance of the frictional force at the film-substrate interface and the divergence of the total stress inside the film. The exponent $\alpha\in [0,1]$ characterizes the nonlinear behavior of friction and is chosen to be $\alpha=0.8$, a value obtained from recent experiments on high-molecular-weight PS dewetting on PDMS-covered substrates [9] and consistent with measurements of rubber-brush friction [33]. $\sqrt{v}$ is a characteristic velocity that enters the characteristic scales, see eq. (6) below. In eq. (4), $G$ is the elastic modulus and $\tau_0$ an additional friction coefficient.

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The friction coefficient evolves as given by eq. (2) (parameters $t^* = 0.1$, $\zeta_1 = 0.03$) and in panel (b) by eq. (8) (parameters $\tau_0 = 500$, $\zeta_\infty = 5$). Viscoplastic time scales $\tau_1 = 1$, $\tau_2 = 100$ and friction exponent $\alpha = 0.8$. $\zeta_0 = 1$ (upper curve) and $\zeta = 5$ (lower curve).

Fig. 1: (Color online) The width $W$ of the dewetting rim as a function of time in semi-logarithmic scale for different initial values of the friction coefficient, $\zeta_0$. In panel (a) the friction coefficient evolves as given by eq. (2) with constant friction coefficients $\zeta$ and for different $W$ values of $0.1$, $1$, $10$, $100$, $1000$. The dashed curves in panel (b) have been obtained with constant friction coefficients $\zeta = \zeta_0 = 1$ (upper curve) and $\zeta = \zeta_\infty = 5$ (lower curve).

Numerical results for the temporal evolution of the width of the rim are shown in fig. 1. One can see that the two different evolution laws for the friction coefficient, eqs. (2) and (8), amount only to quantitative differences. The mechanism is robust and the qualitative behavior, i.e. the occurrence of a maximum in the width and ii) its monotonic decrease as a function of increasing $\zeta_0$, is unchanged. For the friction law with saturation, eq. (8), it is instructive to compare with the cases of non-evolving friction coefficients fixed at $\zeta_0$ and $\zeta_\infty$, see the upper and lower dashed curves, respectively, in fig. 1(b). In both cases, there is no maximum in the width. The two curves enframe the ones with evolving friction. The curves with evolving friction will finally join for long times with the curve obtained with $\zeta_\infty$ (however, the present model should not be applied for such long times). Figure 2(a) shows the dewetting velocity as a function of time. A fast decay of the velocity in the elastic regime, for $\tau_0 < t < \tau_1$, is obtained as also observed experimentally. This signature is due to the decrease of the “effective” driving force $|S|/\zeta$, and has also been observed in the model when including a relaxing residual stress [17,18].

Let us now discuss the effect of aging. Even if kept well below the bulk-PS glass transition, the PS-PDMS interface evolves towards its equilibrium width, as evidenced in [13]. This is most probably due to the fact that motion at the interface is easier than in bulk and that, for the aging temperatures used, the PDMS is not glassy. Consequently this means that aging time translates into increasing values of $\zeta_0$, i.e. of the value of the friction coefficient at the moment where the dewetting process is initiated (experimentally by increasing the temperature above $T_g$). As one can see from fig. 1, increasing $\zeta_0$ leads to a monotonic decrease in the maximum of the width, in accordance with the experiments [13]. Figure 2(b) shows the dependence of the maximum in the width as a function of the initial value of the friction coefficient, $\zeta_0$. There is consistent decrease for both proposed evolutions for the friction coefficient and for various parameter values.

The numerically obtained results can also be interpreted by a scaling estimate: the typical scale for the characteristic width, eqs. (6), (7), implies a dependence on viscosity $\eta$ and friction $\zeta$ as

$$W \propto \eta^{1-\alpha} \zeta^{-1-\alpha},$$

where $\alpha$ is the nonlinear friction exponent. At short times, one has to use $\eta_0$ and $\zeta_0$ in this formula, while at times $t > \tau_1$ one has $\eta_1$ and $\zeta(t)$. This provides a simple picture for the occurrence of the maximum: The increase of the viscosity leads to an increase in the width of the rim up to a time of the order of $\tau_1$. Later on, the width will decrease slowly since the viscosity stays at $\eta_1$, while the friction coefficient $\zeta$ still slowly increases. Assuming the maximum $W^*$ to be roughly at $\tau_1$, cf. fig. 1, one can estimate $W^* \propto [\zeta(\tau_1)]^{-1-\alpha}$. This expression was used to fit the dependence of $W^*$ as a function of $\zeta_0$ to obtain the solid lines in fig. 2(b), with a fixed value $\alpha = 0.8$ for the exponent, as used in the numerical solution. Although the window of $\zeta_0$-values is too small to establish a real power law, the agreement of this simple estimate is very good.

The numerical study and the simple scaling thus suggest that —once the temporal increase in friction due to roughening is established by a microscopic theory— it would be possible to measure the roughness evolution below the glass temperature, i.e. the aging of the buried polymer-polymer interface, by tracing the maximum in the rim width in a dewetting experiment.
We should mention that in case of the interface evolution studied here, a maximum in the width of the rim can already be obtained for a linear friction law, i.e. for $\alpha = 0$ in the momentum equation (3). This is in contrast to the case where relaxing residual stresses are responsible for the maximum, as discussed in detail in refs. [17,18], where nonlinear friction was crucial. The underlying reason for this difference is the fact that the friction coefficient already influences $W$ for a linear friction law, see eq. (9) with $\alpha = 0$, while for the case with residual stress the nonlinearity in the friction was needed to introduce a dependence on residual stress in the scaling for $W$. However, looking at the scaling law eq. (9), for $\alpha = 0$ the dependence on the friction coefficient is much weaker, $W \propto \zeta^{-1/2}$, as compared to $W \propto \zeta^{-5/6}$ for $\alpha = 0.8$. Hence a much larger increase in friction is needed if the friction law is linear, which might be improbable for the small interface changes that are to be expected for an immiscible polymer-polymer interface. Thus, although nonlinear friction is not essential it amplifies the effect of the interface evolution with respect to the viscosity evolution due to viscoelasticity, as directly shown by eq. (9). Investigating the (non-)linearity of the friction experimentally, similarly as has been done in ref. [9], might thus be helpful to better understand quantitative features of the maximum in the rim width.

Conclusions and perspective. – To conclude, an increase in friction (or a decrease in the slip length) due to an evolution of the substrate-film interface can give rise to a maximum in the rim width upon dewetting. This implies that the – macroscopically observable – maximum in the width may be caused by and is susceptible to minute changes of the interface. In addition, there is a monotonous relation between the maximum of the width and the friction coefficient at the beginning of dewetting. Concerning the experiments on PS-PDMS systems, this friction coefficient in turn is affected by the aging time of the polymer-polymer interface below the glass temperature. As compared to the relaxation of residual stresses put forward recently [8,9,17], our work offers an alternative interpretation of these aging effects, in the case of low-molecular-weight dewetting films. However, at present stage it is not possible to give a direct relation between the maximum in the rim width and the aging time, because neither the exact time evolution of the interface towards its equilibrium value, nor the exact relation (for an asymmetric melt-brush system) between interface width/roughness and friction/slippage is known. We would like to encourage MD simulations or dynamic self-consistent field studies to establish this dependence. Then it would be possible to investigate buried polymer-polymer interfaces by such relatively simple and cheap experiments as dewetting.

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We thank P. DAMMAN for inspiring this work and L. LÉGER and G. REITER for very stimulating discussions.

FZ acknowledges financial support by the German Science Foundation (DFG).

REFERENCES

[1] REITTER G., *Phys. Rev. Lett.*, 87 (2001) 186101.
[2] SEEMANN R., HERMINGHAUS S. and JACOBS K., *Phys. Rev. Lett.*, 87 (2001) 196101.
[3] DAMMAN P., BAUDELET N. and REITTER G., *Phys. Rev. Lett.*, 91 (2003) 216101.
[4] BUCKNALL D. G., *Prog. Mater. Sci.*, 49 (2004) 713.
[5] BLOSEY R., *Phys. Chem. Chem. Phys.*, 10 (2008) 5177.
[6] HERMINGHAUS S., JACOBS K. and SEEMANN R., *Eur. Phys. J. E*, 12 (2003) 101.
[7] VILMIN T. and RAPHAËL E., *Europhys. Lett.*, 72 (2005) 781.
[8] REITTER G., HAMIEH M., DAMMAN P., SCLAVONS S., GABRIELE S., VILMIN T. and RAPHAËL E., *Nat. Mater.*, 4 (2005) 754.
[9] VILMIN T., RAPHAËL E., DAMMAN P., SCLAVONS S., GABRIELE S., HAMIEH M. and REITTER G., *Europhys. Lett.*, 73 (2006) 906.
[10] DAMMAN P., GABRIELE S., COPPE S., DESPREEZ S., VILMERS D., VILMIN T., RAPHAËL E., HAMIEH M., AL AKHRASS S. and REITTER G., *Phys. Rev. Lett.*, 99 (2007) 036101.
[11] YANG M. H., HOU S. Y., CHANG Y. L. and YANG A. C.-M., *Phys. Rev. Lett.*, 96 (2006) 066105.
[12] BODIGUEL H. and FRETTIGNY C., *Eur. Phys. J. E*, 19 (2006) 185.
[13] COPPÉE S., GABRIELE S., JONAS A., JESTIN J. and DAMMAN P., *Influence of Chain Interdiffusion Between Immiscible Polymers on Dewetting Dynamics*, arXiv:0904.1675 (2009).
[14] RAPHAËL E. and DE GENNES P. G., *J. Phys. Chem.*, 96 (1992) 4002.
[15] REITTER G., SCHULTZ J., AYROU P. and AUVRAY L., *Europhys. Lett.*, 33 (1996) 29.
[16] REITTER G. and KIHANNA R., *Phys. Rev. Lett.*, 85 (2000) 2753.
[17] VILMIN T. and RAPHAËL E., *Eur. Phys. J. E*, 21 (2006) 161.
[18] ZIEBERT F. and RAPHAËL E., *Phys. Rev. E*, 79 (2009) 031605.
[19] BROCHARD-WYART F., DEBREGES G., FONDECRAVE R. and MARTIN P., *Macromolecules*, 30 (1997) 1211.
[20] KERLE T., YERUSHALMI-ROZEN R. and KLEIN J., *Macromolecules*, 31 (1998) 422.
[21] AJDARI A., BROCHARD-WYART F., GAY C., DE GENNES P. G. and VIOVY J. L., *J. Phys. II*, 5 (1995) 491.
[22] JONES R. A. L. and RICHARDS R. W., *Polymers at Surfaces and Interfaces* (Cambridge University Press, Cambridge) 1999.
[23] BUDKOWSKI A., *Adv. Polym. Sci.*, 148 (1999) 1.
[24] HELFAND E. and TAGAMI Y., *J. Chem. Phys.*, 56 (1972) 3592.
[25] BROCHARD F., DE GENNES P. G. and TROIANTIN S., *C.R. Acad. Sci. (Paris)*, 310 (1979) 1169.
[26] GOVEAS J. L. and FREDRICKSON G. H., *Eur. Phys. J. B*, 2 (1998) 79.
[27] de Gennes P. G., C.R. Acad. Sci. (Paris), 288 (1979) 219.

[28] Clarke C. J., Jones R. A. L., Edwards J. L., Shull K. R. and Penfold J., Macromolecules, 28 (1995) 2042.

[29] Sferrazza M., Xiao C., Jones R. A. L., Bucknall D. G., Webster J. and Penfold J., Phys. Rev. Lett., 78 (1997) 3693.

[30] Clarke C. J., Polymer, 37 (1996) 4747.

[31] Geoghegan M., Clarke C. J., Boué F., Menelle A., Russ T. and Bucknall D. G., Macromolecules, 32 (1999) 5106.

[32] Sferrazza M., Xiao C., Jones R. A. L. and Penfold J., Philos. Mag. Lett., 80 (2000) 561.

[33] Bureau L. and Léger L., Langmuir, 20 (2004) 4523.

[34] Redon C., Brzoska J. B. and Brochard-Wyart F., Macromolecules, 27 (1994) 468.