Prewetting transition on a weakly disordered substrate: evidence for a creeping film dynamics

X. MÜLLER, J. DUPONT-ROC

Laboratoire Kastler Brossel (∗), Ecole Normale Supérieure, 24 rue Lhomond, F75231 Paris cedex 05, France

PACS. 68.10.Gw – Interface activity, spreading.
PACS. 68.15.+e – Liquid thin films.
PACS. 68.35.Rh – Phase transitions and critical phenomena.

Abstract. – We present the first microscopic images of the prewetting transition of a liquid film on a solid surface. Pictures of the local coverage map of a helium film on a cesium metal surface are taken while the temperature is raised through the transition. The film edge is found to advance at constant temperature by successive avalanches in a creep motion with a macroscopic correlation length. The creep velocity varies strongly in a narrow temperature range. The retreat motion is obtained only at much lower temperature, conforming to the strong hysteresis observed for prewetting transition on a disordered surface. Prewetting transition on such disordered surfaces appears to give rise to dynamical phenomena similar to what is observed for domain wall motions in 2D magnets.

Introduction. – Over the past twenty years, prewetting transition has attracted a lasting interest since its first consideration by Ebner, Saam and Cahn [1–3]. Considering an equilibrium between two fluid phases, the prewetting transition on a surface is simply the continuation off coexistence of the wetting transition. The surface coverage corresponding to the wetted state is simply finite rather than infinite. Apart from very special matching conditions [4,5] for which critical wetting transition has indeed been observed [6], first order transitions are expected, resulting from long range Van der Waals forces between the surface and the various fluid components [2–4]. In the simple case of a liquid/vapor phase equilibrium in presence of a surface, the surface coverage changes from a thin coverage to a thick liquid film. Experimental observations of these transitions on various systems have been made over the last decade [7–22]. A particular goal of experimentalists has been to check the first order nature of the transition, to determine the prewetting line in the $T - \Delta \mu$ plane and to locate the prewetting critical point. As a matter of fact, experiments have raised further questions about hysteresis phenomena, about the consequences of surface disorder, about the nucleation of the new phase at the transition and its subsequent growth. Presently there is no unique interpretation of the various experimental observations of prewetting transitions, especially at low temperatures. To our best knowledge, nobody has ever provided a clear

(∗) Unité mixte de recherche du CNRS, de l’Ecole Normale Supérieure et de l’Université Pierre et Marie Curie

© EDP Sciences
experimental evidence for the coexistence at the prewetting transition of areas with thick and thin coverages, separated by an interface which can be moved in a reversible way, as in a bona fide first order transition such as the liquid/vapor one. The transition reversibility has not always been tested. When it has been looked for, hysteresis has been found quite systematically \[7, 13, 17\]. This hysteresis has received diverging interpretations. For some, it reveals a nucleation barrier which can be overcome only by over-passing the equilibrium line and approaching the spinodal line \[23, 24\]. Such a barrier is indeed expected for instance for the dewetting branch of the transition when the initial surface is completely covered by a thick film and that the thin film phase is to be nucleated. For others, hysteresis is related to surface disorder, such as local variations of the surface binding properties. They are equivalent to a random field coupled to the surface coverage, which is the order parameter. This random field is expected to destroy the first order transition. Dynamical transitions may occur for the invasion or drying of the surface \[25, 27\]. Then experimental data are to be related to numerous studies about growth mechanisms of 2D phases \[28, 30\]. Both interpretations may be true, depending of the substrate. While in demixing liquids, the liquid-vapor interface play the role of a perfectly smooth substrate for which the first interpretation is relevant, solid surfaces are thought to exhibit frozen disorder. Thus detailed experimental data are still desired to justify one or the other interpretation. In many experiments only the overall coverage of the surface, or another surface averaged, quantity is measured. Hence a detailed view of the transition is not accessible. Conversely some experiments \[14, 16\] have provided detailed images or movies of spreading films, but clearly out of equilibrium. In this letter, we report the first observation of the coverage map during a prewetting transition. The system used is liquid helium on a cesium metal surface, which includes some frozen disorder. The invasion of the surface by the thick film at the transition is found to take place through a creeping motion of the thick film edge.

**Experimental procedure.** – Cesium metal surfaces are prepared by cold deposition from SAES Getters dispensers on a gold evaporated mirror in conditions similar to other experiments \[1, 2, 3, 21\]. Some care is taken to avoid impurities on the substrate: dispensers, as well as the cell and the pumping tubes, are outgazed at room temperature. To improve its uniformity, the cesium film is deposited from two dispensers at \(\pm 45^\circ\) with respect to the surface normal. It extends over a \(2.5 \times 4 \text{ mm}^2\) rectangle determined by a mask located 0.4 mm above the center of the gold mirror. For the experiments reported here, the cesium layer was 40±5 nm thick. Last layers were deposited at low rate (0.3 nm/min). Differently prepared cesium substrates have also been used in wetting experiments with liquid helium \[33, 34\]. For these surfaces, contact angles and hysteresis properties for the wetting transition have been reported to be markedly different from those obtained with the type of surface used here. Prewetting transitions on such substrates have not however been investigated.

Cesium thickness and helium coverage are measured through an improved Nomarski microscope adapted to low temperatures \[35, 36\]. It provides a map of the local gradient along a given direction of the local surface height. The corresponding image looks like a side illuminated picture of the true surface landscape with a nanometer height scale, and with the microscope resolution (about 4 \(\mu\)m in our case). If a transparent coverage is added on the surface, it produces an optical retardation which appears as an additional negative height. Film edges appear as bright or dark lines. With a 6 s integration time, the microscope is sensitive to optical thickness differences as small as 0.05 nm over a distance of 4 \(\mu\)m. That is sufficient to detect 10 nm thick helium films which correspond to a 0.26 nm optical retardation \((n_{\text{He}} - 1 = 0.026)\).

Below 1.8 K, liquid helium does not wet the cesium surface. When filling the cell with
helium liquid at low temperatures (typically 1.4 K), the vertical cesium mirror surface does not get wetted. When the liquid level is raised onto the surface, it makes a non-zero advancing contact angle. When afterwards the level is lowered by pumping part of the helium out of the cell, the contact angle becomes zero and a metastable thick film remains pinned on the surface. The film thickness is determined by its altitude above the liquid level which can be varied from 1 mm to 20 mm. The corresponding measured thicknesses are 70 nm and 20 nm. These values are consistent with those computed from the cesium/helium non-retarded Van der Waals force [38]. The mirror surface around the small rectangle covered with cesium is of course also covered by a wetting helium film. When temperature is raised, the prewetting transition temperature is eventually met and the helium film should cover the originally dry area. Afterwards, lowering the temperature should cause dewetting. Because the initial state contains both thin and thick phases in contact, nucleation metastability is avoided. Interface pinning by the disorder is the only hysteresis mechanism at work in this situation.

Prewetting transition through thick film invasion. – In a first run, temperature is increased slowly, typically at a rate less than 1 mK/min, until a change is observed in the coverage. Then temperature is stabilized during the time of the experiment within a few mK. In subsequent runs, temperature is set at different values around this initial one. It is observed that the straight line of the initial film border becomes irregular because it starts to move at a few places. Then the film advances over the dry area with a slow creeping motion by successive avalanches. Intermediate states of the surface invasion at 1.817 K are shown in Figure 1. Because the observed area is near the right edge of the cesium rectangle, part of invasion comes also from the right. This explains the overall L-shaped film front. The contrast of the film edge does not change significantly, indicating a constant thickness.

The morphology of the film border is characterized by a macroscopic correlation length, clearly visible in Figure 1, much larger than typical lengths characterizing the cesium layer such as its thickness 40 nm, or its morphological length scale which is probably smaller. This is a clear indication of a cooperative (i.e. weak) pinning situation for the film edge. To quantify the correlation length, one considers the r.m.s deviation of the film edge position $w(l)$ taken over a length $l$ along the mean front direction. It is plotted versus $l$ in Figure 2. At short distance, the plot starts from a non-zero value due to existing overhangs and dry holes inside the film. $w(l)$ increases over long distances. Cross-over between the two regimes occurs for the correlation length $l_c \approx 10 \, \mu m$ and a front depth $w(l_c) = 50 \, \mu m$. For zero temperature
Fig. 2 – $w(l)$, r.m.s front position fluctuations on length $l$ along a line approximating the front. The average is taken over several successive positions of the film border and on several experimental runs. Non vanishing values at short distances are due to overhangs.

Fig. 3 – Creep velocity as a function of temperature for 50 ± 8 nm thick films

models, Robbins et al. point out that this cross-over prevents from getting the roughness exponent 0.5 expected for a self-affine growth [25]. We are not aware of predictions for $w(l)$ taking into account both finite temperatures and important contributions of overhangs.

No nucleation of wet patches in front of the film seems to occur. It is unlikely that wet parts smaller than the microscope resolution exist. In particular, some defects (presumably submicronic dust grains) give rise to ‘helium hills’ inside the film by a local thickening, as can be seen in Figure 1. These ‘hills’ appears only when the film edge reaches them. This suggests that the surface ahead the film is indeed free of thick liquid film.

After the cesium surface is completely invaded, the cell is cooled down as much as possible. Partial dewetting was observed around 1.23 K. The film split into two parts separated by a dry horizontal line, then one of this part completely disappeared. Dewetting, as wetting, was anisotropic. As anticipated from previous studies, the film edge dynamics is strongly hysteretic.

The film advances at constant temperature by successive avalanches with sizes on the order of $l_c$. The advance velocity is defined as the wet area increase between two images divided by the time interval and the average front length. For comparison, velocities are always measured on the same area of the surface. Two facts hints at a thermally activated process. First, the dry holes inside the film fill up after a while. This means that no part of the border is definitely pinned. They have a chance to move if one waits for a sufficiently long time. A probabilistic mechanism is at work to get over the higher barriers, probably thermal fluctuations since intentionally produced vibrations have no effect. Secondly the film creep velocity increases strongly with temperature as shown in Figure 3. Higher temperatures indeed increase the spreading forces compared to the pinning ones. This lowers the barriers and makes avalanches more probable. Film dynamics exhibits also some peculiarities. The advance is much faster from the right to the left than upwards, and the speed of the front
motion somewhat decreases while it sweeps across the picture area. This may be due to a slight non uniformity and anisotropy of the cesium layer related to the deposition geometry, but this remains to be checked.

To summarize, in the case presented here, prewetting transition takes place through the displacement of the thick film border by a random, probably thermally activated process, with a macroscopic correlation length indicating a cooperative pinning. The creep velocity changes from an vanishing low value to observable ones on a very small temperature interval, around what appears as a unbinding threshold. The inverse process, the drying transition, occurs through a similar process significantly shifted to lower temperatures.

Discussion. – These properties strongly suggests that the normal first order prewetting transition is hindered by the pinning of the thick film border by the substrate disorder. The thick film advance or retreat takes place only when the spreading force is sufficiently strong to overcome the corresponding barriers. The observed transition is thus related to the film border dynamics. This is consistent with the observation of different advancing and receding angles on similar surfaces below the wetting temperature \[39–41\]. The fact that no nucleation ahead of the front is observed means that the pinning is not strong enough to bring the observed dynamical transition up to the spinodal line of the first order wetting transition \[42\].

The growth itself is nearly compact. These features are in contrast with the observations in a previous experiment presented in \[43\]. With thicker cesium layers obtained at higher deposition rates, no motion of the film edge was observable. Only a continuous increase of the coverage ahead of the front was detected when temperature was increased above a given threshold. This was interpreted as the growth of a film non compact at a scale below the microscope resolution. The substrate disorder was probably more important, and thus the film correlation length much shorter. It is also interesting to remark that the wetting threshold was somewhat higher in temperature (1.9 K instead of 1.82 K here) and that dewetting was not observable, suggesting an overall larger hysteresis in term of spreading force. A similar case was recently studied for cesium surfaces with a controlled large scale disorder \[44\]. The correlation length of the film border is then closer to the disorder length scale indicating a strong pinning case.

Concerning the dynamics, the motion of a line in a disordered 2D medium has been described by a thermal activation law for the velocity, with a stretched exponential form \[32, 45\] : $v \sim \exp[-Cf^{-\mu}/T]$ where $C$ is a constant, $f$ is the driving force per unit length of line and $\mu$ is an exponent depending the system dimensionality and disorder. Such a law has been verified recently for the triple line motion \[32\], and for domain wall motion in 2D magnets \[30\] with $\mu = 1/4$. In the present case, the system is in the same universality class as the 2D random field Ising model so that $\mu$ is expected to take the value 1. The range of velocity observed is however not wide enough to check this prediction.

The spreading of a helium prewetting film was already observed by Reinelt et al. \[14\]. A droplet falling on an inclined surface was accompanied by a spreading helium film, 30 nm thick. Its spreading velocity was about 0.5 cm/s at the temperature 1.85K. This large velocity may indicate a larger temperature difference from the threshold. It is remarkable, but probably a coincidence that linear extrapolation of the data of Figure \[3\] to 1.85 K yields the same order of magnitude. Note that with such a thermally activated law for the velocity the temperature at which the prewetting transition is actually observed depends somewhat on the observation time. Changing this time from 1 s to 4 hour shifts the observed prewetting temperature only from 1.85 K to 1.82 K. Hence it is tempting to suggest that the situations appearing as metastable in the temperature range between the observed prewetting temperature and the dewetting one actually require exponentially large and unrealistic observation times to
reach equilibrium. Then the threshold at which the invasion begins does not correspond to a real dynamical transition as in zero temperature models\textsuperscript{[26,27]}. It is rather a cross-over between a low temperature regime with large and rare avalanches to a high temperature regime, not reached in this experiment, with an overall motion of the border line limited by some dissipation mechanism.

**Conclusion.** – The observations presented show clearly how the frozen disorder of the surface interfere with the prewetting transition in the weak disorder situation. Collective pinning of the border line between the thick and the thin film phases prevents the transition from taking place through its displacement in favor of one or the other of the two phases. Experimental observations of the transition correspond then to the unbinding of the line at a finite spreading power. This relates prewetting transitions on disordered substrates to other phenomena exhibiting domain wall motions in two dimensions, such as 2D magnetism.

∗ ∗ ∗

We are indebted to E. Rolley, A. Prevost and C. Guthmann for many helpful discussions.

REFERENCES

[1] Ebner C. and Saam W., *Phys. Rev. Lett.*, **38** (1977) 1486.
[2] Cahn J. W., *J. Chem. Phys.*, **66** (1977) 3667.
[3] Ebner C. and Saam W., *Phys. Rev. B.*, **35** (1987) 1822.
[4] See for instance Dietrich S., *Phase transitions and critical phenomena*, edited by C. Domb and J. L. Lebowitz, Vol. **12** (Academic Press, London) 1988, p. .
[5] Indekeu J. O., Ragil K., Bonn D., Broseta D. and Meunier J., *J. Stat. Phys.*, **95** (1999) 1009.
[6] Ragil K., Meunier J., Broseta D., Indekeu J. O. and Bonn D., *Phys. Rev. Lett.*, **77** (1996) 1532.
[7] Ketola K. S., Wang S., and Hallock R. B., *Phys. Rev. Lett.*, **68** (1992) 201
[8] Rutledge J. E. and Taborek P., *Phys. Rev. Lett.*, **69** (1992) 937
[9] Kellay H., Bonn D., and Meunier J., *Phys. Rev. Lett.*, **71** (1993) 2607
[10] Demolder B. and J. Dupont-Roc, *Physica B*, **194-196** (1994) 975
[11] Demolder B., Bigelow N., Nacher P.J. and Dupont-Roc J., *J. Low Temp. Phys.*, **98** (1995) 91
[12] Cheng E., Mistura G., Lee H.C., Chan M.H.W., Cole M.W., Carraro C., Saam W.F. and Toigo F., *Phys. Rev. Lett.*, **70** (1993) 1854
[13] Phillips J.A., Ross D., Taborek P. and Rutledge J. E., *Phys. Rev. B.*, **58** (1998) 3361
[14] ReineLT D., Gau I., Herminghaus S. and Leiderer P., *Czech. J. Phys.*, **46** (1996) 432
[15] Herminghaus S., Vorberg J., Gau H., Conradt R., ReineLT D., Ulmer H., Leiderer P. and PrzyrembEl M., *Ann. Physik*, **6** (1997) 425
[16] ReineLT D., Klier J. and Leiderer P., *J. Low Temp. Phys.*, **113** (1998) 805
[17] ReineLT D., Engel A. and Klier J., *Physica B*, **284-288** (2000) 149
[18] Yao M. and Hensel F., *J. Phys. Condens. Matter*, **8** (1996) 9547
[19] Hensel F. and Yao M., *Euro. J. of Solid and Inorg. Chem.*, **34** (1997) 861
[20] Kozhevnikov V. F., Arnold D. I., Naurzakov S. P. and Fischer M. E., *Phys. Rev. Lett.*, **78** (1997) 1735
[21] Wynblatt P. and Chatain D., *Ber. Buns. Ges.: Phys. Chem. Chem. Phys.*, **102** (1998) 1142
[22] Lucht R. and Bahr C., *Phys. Rev. Lett.*, **80** (1998) 3783
[23] Bonn D., Kellay H. and Wegdam G. H., *Phys. Rev. Lett.*, **69** (1992) 1975
[24] Herminghaus S., Jacobs K., Mecke K., Bischof J., Fehy A., Ibn-Elhaj M. and Schlagowski S., *Science*, **282** (1998) 916
[25] Nolle C.S., Koiller B., Martys N., and Robbins M.O., Physica A, 205 (1994) 342 and ref. in.
[26] Blossey R., Kinoshita T. and Dupont-Roc J., Physica A, 248 (1998) 247
[27] Blossey R., Müller X., Kinoshita T. and Dupont-Roc J., J. Low Temp. Phys., 110 (1998) 665
[28] Léger L. and Joanny J. F., Rep. Prog. Phys., 55 (1992) 431
[29] Voue M., De Coninck J., Villette S., Valignat M. P. and Cazabat A. M., Thin Solid Films, 313-314 (1998) 819 and ref. in.
[30] Lemerle S., Ferré J., Chappert C., Mathet V., Giamarchi T. and Le Doussal P., Phys. Rev. Lett, 80 (1998) 849
[31] Phillips J. A., Taborek P. and Rutledge J. E., J. Low Temp. Phys., 113 (1998) 829
[32] Rolley E., Prevost A. and Guthmann C., J. Low Temp. Phys., 113 (1998) 787
[33] Klier J., Stefanyi P. and Wyatt A. F. G., Phys. Rev. Lett., 75 (1995) 3709
[34] Klier J. and Wyatt A. F. G., J. Low Temp. Phys., 110 (1998) 919
[35] Gleyzes P., Boccara A. C. and Saint-Jalmes H., Optics Letters, 22 (1997) 1529.
[36] Müller X., Kinoshita T. and Dupont-Roc J., J. Low Temp. Phys., 113 (1998) 823
[37] Müller X., Kinoshita T. and Dupont-Roc J., Rev. Sci. Instr., (April 2001)
[38] Vidal G., Ihm G., Kim H. and Cole M. W., Surf. Sci. Rep., 12 (1991) 133
[39] Rolley E. and Guthmann C., J. Low Temp. Phys., 108 (1997) 1
[40] Ross D., Taborek P. and Rutledge J. E., Science, 278 (1997) 664
[41] Phillips J.A., Ross D., Taborek P. and Rutledge J. E., J. Low Temp. Phys, 113 (1998) 811
[42] Bonn D., Kelly H. and Meunier J., Phys. Rev. Lett, 73 (1994) 3560
[43] Müller X. and Dupont-Roc J., Physica B, 284-288 (2000) 143
[44] Prevost A., Poujade M., Rolley E. and Guthmann C., Physica B, 280 (2000) 80
[45] Chauve P., Giamarchi T. and Le Doussal P., Phys. Rev. B, 62 (2000) 6241