Anomalous roughening of Hele-Shaw flows with quenched disorder

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The kinetic roughening of a stable oil–air interface, moving in a Hele–Shaw cell which contains a quenched columnar disorder (tracks) has been studied. A capillary effect is responsible for the dynamic evolution of the resulting rough interface, which exhibits anomalous scaling. The three independent exponents needed to characterize the anomalous scaling are determined experimentally. The anomalous scaling is explained in terms of the initial acceleration and subsequent deceleration of the interface tips in the tracks coupled by mass conservation. A phenomenological model that reproduces the measured global and local exponents has been introduced.

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The kinetic roughening of surfaces and interfaces in nonequilibrium conditions has received a great deal of attention in last years. This is a subject of technological importance in surface growth, and of fundamental relevance as an example of scaling far from equilibrium.

Roughening originates from the competition between stabilizing and destabilizing mechanisms at different length scales. As a result of the competition, the correlation length of the interfacial fluctuations along an initially flat front increases monotonously with time, until it reaches the system size and the roughness saturates. The statistical properties of the roughening front are usually given in terms of the (rms) width \( w \) of the interfacial fluctuations. In the absence of characteristic scales, many systems follow the dynamic scaling hypothesis of Family–Vicsek (FV) [2], in which short and long length scales have the same scaling behavior: \( w(l) \sim t^\beta \) for \( t < t_1 \), \( w(l) \sim l^\alpha \) for \( t > t_x \) and \( t_x \sim t^z \). Here \( l \) is the lateral size of the window used to measure \( w \), \( t_x \) is the saturation time, \( \beta \) the growth exponent, \( \alpha \) the roughness exponent, and \( z \) the dynamic exponent. The scaling relation \( \alpha = z\beta \) reduces to two the number of independent exponents that characterize the universality class of the problem under study.

In recent years, however, experiments on molecular beam epitaxy [3], sputtering [4], fracture mechanics [5], and electrodeposition [6] have revealed a rich variety of situations in which the scaling of the global and the local interface fluctuations are substantially different. Long length scales are then characterized by the same FV scaling exponents, while short length scales follow the so-called anomalous scaling ansatz [7]: \( w(l, t) \sim t^\beta \) for \( t < t_1 \), \( w(l, t) \sim l^{(1-\beta)z} \) for \( t_1 < t < t_x \), and \( w(l, t) \sim l^{(1-\alpha)z} \) for \( t_x < t \). The local time \( t_1 \sim t^z \), and the saturation time \( t_x \sim L^z \), where \( L \) is the system size in the direction along the front. Here \( \beta^* \) and \( \alpha_{loc} \) are new local growth and roughness exponents, respectively, which characterize the scaling at short length scales \( (l \ll L) \). There is a new scaling relation \( \alpha - \alpha_{loc} = z\beta^* \). Since now there are five exponents and two scaling relations, the anomalous scaling is characterized by three independent exponents. Usually the roughness exponents are determined through the power spectrum, which is less sensitive than \( w \) to finite size effects. The spectrum scales as \( S(k, t) = k^{-(2\alpha+1)}s_A(kt^{1/2}) \), where \( s_A \) obeys \( s_A(u) \sim u^{2\beta + 1} \) when \( u \ll 1 \) and \( s_A(u) \sim u^{2\beta} \) when \( u \gg 1 \). Here \( \theta = \alpha - \alpha_{loc} \). The anomalous scaling leads to the usual FV scaling when \( \theta = 0 \) (short and long length scales are characterized by the same roughening exponents) and \( \beta^* = 0 \) (the temporal correlations of the short length scales disappear). Recently it has been shown that the anomalous scaling can be related to the presence of high slopes in the front that have nontrivial dynamics [8].

![Fig. 1. Sequence of typical oil-air interfaces. The silicone oil moves upwards in the picture, and the disorder pattern is represented in grey. The experimental parameters are \( b = 0.36 \) mm and \( v = 0.04 \) mm/s.](image_url)
In this Letter we report the first observations of anomalous roughening in experiments of Hele–Shaw flows with quenched disorder. We consider an initially flat front between oil and air, driven at a small constant average velocity $v$ in a horizontal Hele–Shaw cell with a columnar quenched disorder. The disorder consists of continuous copper tracks on a fiberglass substrate in the advancing direction of growth and randomly distributed in the perpendicular direction (Fig. 1). In this situation, the correlation of the disorder in the advancing direction is infinite. The local motion relative to the average position of the front is driven by capillary forces, caused by the different curvatures of the advancing front in the third dimension, depending on whether the oil is on a copper track or on the fiberglass substrate. An analysis of the $rms$ width of the rough interface at short and long length scales plus a collapse of the power spectra of the interfacial fluctuations enables an independent determination of the three exponents characteristic of anomalous scaling.

We associate the anomalous roughening with the initial acceleration and subsequent deceleration of the liquid on the copper tracks, caused by capillarity, and the coupling of the motion over tracks and over fiberglass due to mass conservation. The anomaly decreases as $v$ increases, and disappears when $v$ is large enough to override the local acceleration and deceleration $[9]$. To explain the experimental results, we propose a phenomenological model which gives global and local exponents in good agreement with the experimental exponents.

The experiments have been performed in a horizontal Hele–Shaw cell, $190 \times 550 \ (L \times H) \ mm^2$, made of two glass plates $20 \ mm$ thick. A fiberglass substrate with copper tracks, manufactured using printed circuit technology, is attached to the bottom plate. The unit copper track is $d = 0.06 \pm 0.01 \ mm$ high and $1.5 \pm 0.04 \ mm$ wide. The tracks are placed randomly in a grid of $1.5 \ mm$ lattice spacing, along the $x$ direction, up to a $35\%$ occupation. There are no lateral correlations of the disorder at distances larger than the width of the unit track. The distance between the top plate and the substrate defines the gap spacing $b$, which has been varied in the range $0.16 \leq b \leq 0.75 \ mm$. We have used 4 different disorder configurations and carried out 2 identical runs per disorder configuration. A silicone oil (kinematic viscosity $\nu = 50 \ mm^2/s$, density $\rho = 998 \ kg/m^3$, and surface tension oil–air $\sigma = 20.7 \ mN/m$ at room temperature) is injected into one side of the cell at constant volumetric injection rate. The oil wets similarly and almost completely the surfaces which it is in contact with (glass, substrate, and copper). The evolution of the front is driven by capillary forces, caused by the different curvatures of the advancing front in the third dimension, depending on whether the oil is on a copper track or on the fiberglass substrate. An analysis of the $rms$ width of the rough interface at short and long length scales plus a collapse of the power spectra of the interfacial fluctuations enables an independent determination of the three exponents characteristic of anomalous scaling.

The temporal evolution of the power spectrum is presented in Fig. 2(a). At very short times we observe a regime with $S(k) \sim k^{-3.4}$. It is a super–rough transient regime created when the fluid on the tracks reaches its maximum velocity after coming into contact with the disorder. The main regime appears further on, with $S(k) \sim k^{-2.1}$, and a vertical shift that progressively decreases with time and disappears at saturation ($t \gtrsim 450 \ s$). The vertical shift of the power spectra is a sign of intrinsic anomalous scaling, in which the spectral exponent $[12]$ can be identified with the local roughness exponent, so that $\alpha_{loc} = 0.55 \pm 0.10$.

Using the above procedure, the three independent exponents $\beta$, $\beta^*$, $\alpha_{loc}$ have been obtained directly from the experiments. The other two exponents $\alpha$, $z$ can be obtained from the scaling relations. In order to verify the whole scaling, however, we prefer to determine the critical exponents by performing the best collapse of the spectra compatible with the experimental results. The

\[ W(l,t) = \frac{1}{2} \left( w^2(l,t) - w^2(l,0) \right)^{1/2} \]

FIG. 2. Experimental determination of $\beta$ and $\beta^*$. The window size $l$ decreases from $L$ to $L/256$. The experimental parameters are $b = 0.36 \ mm$ and $v = 0.08 \ mm/s$. 

In Fig. 3 we present the results for the interfacial width as a function of time at different length scales $l$, from $L$ to $L/256$. We plot $W(l,t) = \left[ w^2(l,t) - w^2(l,0) \right]^{1/2}$ to minimize the influence of the initial condition $[10]$. We obtain clear power law dependencies, providing $\beta = 0.52 \pm 0.02$ at the longest length scales and $\beta^* = 0.26 \pm 0.03$ at the two shortest length scales. It is worth mentioning that a different method, based on the growth of the local slopes $[5]$, gives the same value of $\beta^*$. The fact that this local exponent is significantly different from 0 is a sign of anomalous scaling of the measured fronts.

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collapse, presented in Fig. 3(b), leads to the following set of scaling exponents:

\[
\beta = 0.50 \pm 0.04, \beta' = 0.25 \pm 0.03,
\alpha = 1.0 \pm 0.1, \alpha_{loc} = 0.5 \pm 0.1, z = 2.0 \pm 0.2.
\] (1)

We have also looked into the possibility of multiscaling through the scaling of the generalized correlations of order \(q\), of the form \(\{h(x + l, t) - h(x, t)\}^{1/q} \sim l^\alpha\). Different exponents for \(\alpha_q\): \(\alpha_2 \approx 0.6, \alpha_4 \approx 0.3\), and \(\alpha_6 \approx 0.2\) were obtained, which confirm the existence of multiscaling in the experiment.

To understand the origin of anomalous scaling, we should look at the interfacial dynamics in greater detail. An initially flat oil–air interface, moving at a nominal average velocity \(v\), experiences local accelerations at those points that touch a copper track for the first time. The local velocity of these points jumps to a maximum in a characteristic time of 1 s, and then relaxes as \(t^{-1/2}\) to \(v\). The maximum velocity reached in a copper track increases with track width and decreases with gap spacing. As shown in Fig. 4, the average interface velocity over copper tracks or over fiberglass channels follows \(v_\pm = v \pm (v_M - v)t^{-1/2}\) respectively, where \(v_M\) is the maximum of the average velocity over tracks. This functionality varies slightly with gap spacing due to the increasing correlation between neighboring tracks. It does not apply for \(b \gtrsim 0.6\) mm. This behavior of different velocities \(v_+\) and \(v_-\) is directly correlated with the shift of the power spectra, indicative of anomalous scaling: the shift is maximum at very short times and disappears close to saturation, similarly to the difference between the average velocities shown in Fig. 4. When we inhibit the relaxation of the velocity over copper tracks by either injecting at \(v > v_M\) or using large \(b\), the anomalous scaling disappears. This scenario has been studied experimentally by exploring different velocities and gap spacings. Fig. 5 shows a phase diagram representing the regions where the anomalous scaling is present (grey region, \(\theta \neq 0\)) or not (white regions, \(\theta = 0\)). The solid line represents the function \(v_M(b)\), and the vertical line the limit between strong and weak capillary forces.

FIG. 3. (a) Temporal evolution of the power spectrum for the same experiments as Fig. 3. The vertical line gives the value of \(k\) associated with the lateral size of the unit copper track. (b) Collapse of the experimental power spectra for \(t > 10\) s. (c) Collapse of the power spectra obtained from numerical simulations of the interfacial equation.

FIG. 4. Average interface velocity over copper tracks and over fiberglass, for \(b = 0.36\) mm and \(v = 0.08\) mm/s.

The critical exponents of the anomalous scaling depend on gap spacing and velocity. We have seen that the anomalous exponent \(\theta\) tends to \(\theta = 0\) as we increase either the velocity or the gap spacing. This exponent
is particularly sensitive to variations of the gap spacing. For $b \gtrsim 0.6 \text{ mm}$, $\theta = 0$ at any velocity. The experimental parameters $b = 0.36 \text{ mm}$ and $v = 0.08 \text{ mm/s}$, deep in the grey region of Fig. 3 give the appropriate conditions to fully characterize the anomalous scaling experimentally. Smaller gap spacings mean that the correlation length in the $x$ direction cannot grow to scales larger than the average track width, and larger gap spacings give too weak an anomalous scaling because the $t^{-1/2}$ law is only obeyed on the widest tracks.

Finally, in order to reproduce the experimental behavior in a phenomenological way, we have developed an interfacial equation of the form:

$$\partial_t h = \partial_x D(x) \partial_x h + v + (v_M - v) t^{-1/2} \eta(x), \quad (2)$$

where $h = h(x,t)$, and $\eta(x)$ is a dichotomous noise with values $+1$ and $-1$ (columnar disorder). The diffusive coupling accounts for the dynamics with $z = 2$ found in experiments, and the other two terms account for the behavior of $v_{\pm}$ observed experimentally. Although the term $t^{-1/2}$ introduces some nonlocality in the model, this is essentially different from non-local models [14] introduced recently to account for mass conservation. The simplest version (constant $D$) reproduces the experimental behavior at long length scales ($\alpha = 1$, $\beta = 0.5$, $z = 2$), but not at short length scales, since numerical simulations of the model give $\beta^{*} = 0$ and $\alpha_{loc} = \alpha = 1$. The reason is that the simulated interfaces do not display the large fluctuations at the copper track edges observed in the experimental fronts (Fig. 3). This can be imposed analytically in Eq. (2) through an inhomogeneous diffusion coefficient $D(x)$. Numerically, we have introduced this effect by spatially averaging the interface inside each track once each $n$ time steps. We thus recover interfaces that are morphologically analogous to experimental interfaces. Now the exponents derived from the scaling of the power spectra, shown in Fig. 3(c), and the multiscaling exponents, reproduce the values determined experimentally. In summary, Eq. (2) enables the relevant effects in the experiment to be both identified and calibrated.

In conclusion, we have presented the first experimental evidence of anomalous scaling in the dynamic roughening of a fluid interface in a Hele-Shaw cell with quenched disorder, and we have studied the experimental conditions for the appearance of the observed anomaly. The exponents $\beta$, $\beta^{*}$, and $\alpha_{loc}$ have been measured independently and their value has been refined by imposing the scaling relations through a collapse of the power spectra. Finally we have introduced an interfacial equation (2) that models the capillary phenomena observed in the experiments, at a phenomenological level. The model reproduces both the morphology of the interfaces and the values of the anomalous scaling exponents.

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FIG. 5. Phase diagram $v(b)$ indicating the regions where the anomalous scaling is observable. The symbols represent the different regions explored experimentally, and the arrow indicates the parameters used in the paper.

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