Complex shape evolution of electromigration-driven single-layer islands

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The shape evolution of two-dimensional islands through periphery diffusion biased by an electromigration force is studied numerically using a continuum approach. We show that the introduction of crystal anisotropy in the mobility of edge atoms induces a rich variety of migration modes, which include oscillatory and irregular behavior. A phase diagram in the plane of anisotropy strength and island size is constructed. The oscillatory motion can be understood in terms of stable facets which develop on one side of the island and which the island then slides past. The facet orientations are determined analytically.

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The manipulation of nanostructures by macroscopic forces is likely to become a key ingredient in many nanotechnology applications. Understanding the influence of external fields on the shape evolution of nanoscale surface features is therefore of considerable importance. As a first step in this direction we analyze here the effects of an electric current on single-layer islands on a crystalline surface. The islands evolve under surface electromigration, the directed motion of adsorbed atoms due to the slight force transmitted by collisions with the conduction electrons in the sample.

Electromigration along interfaces and grain boundaries is the most persistent and menacing reliability problem in integrated circuit technology. Correspondingly, much work has been devoted to electromigration-induced void formation and breakdown in metallic conductor lines. The capacity for quantitative numerical modeling has been demonstrated at least for simple void geometries. A major obstacle to achieving predictive power in such studies, however, is the insufficient control over the complex internal structure of the polycrystalline samples. Hence an important motivation for investigating electromigration-induced effects on simple, well-controlled nanoscale morphologies, such as step patterns on vicinal surfaces and single layer islands, is to bridge the gap between the microscopic mechanisms of electromigration and their consequences on technologically relevant length and time scales.

Electromigration of islands has been modeled previously using Monte Carlo simulations and continuum theory. The continuum approach to island shape evolution, which treats the island edge as a smooth curve, has been successfully applied to a range of problems including the diffusion and sintering of islands, and the pinch-off of vacancy clusters. Here we focus on the regime of periphery diffusion (PD), where the dominant kinetic process is the migration of atoms along the island boundary. The shape then follows a local evolution law, without coupling to the adatom concentration on the surrounding terrace.

We extend the model of by including crystal anisotropy in the adatom mobility. It was observed recently in the context of step flow growth that crystalline anisotropy can change the behavior of step patterns in a qualitative way. In the present case, it leads to the unfolding of a remarkable richness of dynamic phenomena: In addition to the scenarios of steady motion and island breakup observed in previous work, we find spontaneous symmetry breaking, oscillatory shape evolution, and complex migration trajectories where different modes of motion alternate in a periodic or irregular fashion. This highlights the importance of properly including anisotropy in the modeling of boundary evolutions. Oscillatory shape dynamics has been seen previously in a numerical study of void electromigration, and transitions to quasiperiodicity and chaos are known to occur in directional solidification. To the best of our knowledge, however, our work provides the first example of complex shape evolution for a closed contour subject to purely local dynamics.

In the PD regime, the normal velocity $v_n$ of the island boundary satisfies the continuity equation

$$v_n + \frac{\partial}{\partial s} \Omega \sigma \left[ - \frac{\partial}{\partial s} (\Omega \gamma \kappa) + q^* E_t \right] = 0.$$  (1)

Here $s$ denotes the arclength along the island edge, and $\Omega$ the atomic area. The square bracket multiplied by the edge atom mobility $\sigma$ is the total mass current along the boundary, which is driven by the tangential derivative of the chemical potential $\Delta \mu = \Omega \gamma \kappa$ and the electromigration force $q^* E_t$. $\gamma$ is the edge stiffness, $\kappa$ the local curvature, $q^*$ the effective charge of an edge atom, and $E_t$ the tangential component of the local electric field. The crystal anisotropy of the surface enters through the dependence of $\gamma$ and $\sigma$ on the orientation angle $\theta$ of the island edge.

For atomic layer height islands on the surface of a thick sample, the island boundary has a negligible effect on the
electric field; this is in contrast to the modeling of voids in conductor lines, where the coupling of the void shape to the electric field leads to a manifestly nonlocal problem. Here we can take the field to be of constant strength $E_0$ and aligned along the $x$-axis. Letting $\theta$ denote the angle between the normal of the island edge and the $y$-axis (counted positive in the clockwise direction), this implies $E_i = E_0 \cos(\theta)$.

Together with the specification of $\gamma$ and $\sigma$, to be addressed below, this completes the definition of the local boundary evolution. Comparing the two terms inside the square brackets, we extract the characteristic length scale $l_E = \sqrt{\gamma/S_0^2}$, which gauges the relative importance of capillary and electromigration forces; electromigration dominates on scales large compared to $l_E$. Below all lengths are reported in units of $l_E$.

The isotropic version of (1), with $\gamma, \sigma = \text{const.}$, has been studied previously by Suo and collaborators. A circular island moving at constant velocity is stable for (dimensionless) radii $R < R_c = 3.26$. Beyond the instability a bifurcation to two branches of non-circular stationary solutions occurs. Numerical integration of the time-dependent problem shows that only one of the branches, corresponding to islands elongated in the field direction, is realized. At large radii island breakup occurs, mediated by the outgrowth of a finger of the kind found in [11].

We now turn to the effects of crystal anisotropy. Throughout this paper only the anisotropy of the adatom mobility $\sigma$ will be taken into account, while the edge stiffness $\gamma$ is kept isotropic. This is motivated partly by the fact that the anisotropy in $\sigma$ is found experimentally (to the extent that it has been investigated) to much exceed that of $\gamma$, and partly by our desire to disentangle kinetic (\(\gamma\)) and thermodynamic (\(\delta\)) effects. For the kinetic anisotropy we employ the functional form

$$\sigma(\theta) = \sigma_{\text{max}}(1 + S)^{-1}\{1 + S \cos^2[n(\theta + \alpha)/2]\}. \quad (2)$$

Since the prefactor $\sigma_{\text{max}}$ only sets the time scale, the relevant parameters in (2) are the anisotropy strength $S$, the number of symmetry axes $n$, and the angle $\alpha$ between the symmetry axes and the electric field direction. The natural time unit is $t_E = l_E^2/(\sigma_{\text{max}}\Omega^2)$. The simplest solutions in the anisotropic case are stationary islands moving at constant speed, which satisfy the equation $v_n = V \sin(\theta + \phi)$; here the angle $\phi$ accounts for the fact that the island does not necessarily move in the field direction. A complete analysis of stationary island shapes has been achieved in the limiting case of zero stiffness. For an even number $n$ of symmetry axes smooth stationary shapes are found for small $S$, while for larger anisotropy the shapes develop self-intersections; no stationary shapes exist when $n$ is odd. The migration direction generally lies between the field direction and the symmetry axis of the anisotropy.

Despite their mathematical interest, these results are of limited applicability to real islands, because all stationary shapes are wildly unstable when $\gamma = 0$. In the remainder of the article we therefore focus on the numerical solution of the full, time-dependent problem with $\gamma > 0$ and $\sigma(\theta)$ given by (2). Two complementary numerical algorithms have been employed. For relatively small islands a finite difference scheme described in [2] was found to be most efficient, while for large islands we rely on the better stability properties of a semi-implicit adaptive finite element algorithm. The full mutual consistency of the two approaches has been checked.

Most results have been obtained for $n = 6$ and $\alpha = 0$. This leaves the anisotropy strength $S$ and the initial condition for the deterministic shape evolution to be specified. Extensive calculations show that the dependence on the precise initial shape is minor, and hence the initial condition can be characterized by the radius $R_0$ of a circular island of the same area; in practice, we usually start the calculation from a slightly distorted circle.

The phase diagram in Fig. 1 gives an overview of the observed migration modes in the $S - R_0$-plane. For small islands ($R_0 \leq 2$) the evolution converges to a sta-
FIG. 3: Snapshots of oo motion for $R_0 = 4$ and $S = 1$ (upper panel) and zz motion for $R_0 = 3.5$, $S = 0.5$ (lower panel), taken at time intervals $\Delta t = 20$. In both cases the perimeter displays simple oscillations, as in the bottom panel of Fig.4

stationary shape which moves in the direction of the field. For large $S$ the shapes develop facets \cite{22}, similar to what has been observed for void electromigration \cite{3}. Increasing the island radius the direction of migration starts to deviate from the field direction, and we enter the regime of oblique stationary (os) motion (Fig.2). Since the field is aligned with the symmetry axis of the anisotropy, the appearance of obliquely moving solutions implies that the symmetry of the problem is spontaneously broken. In the os regime, pairs of symmetry-related stationary solutions coexist; which solution is chosen in a given run depends on the initial condition.

Upon further increase of $R_0$ the obliquely moving shapes start to oscillate (Fig.3). Near the onset of oblique oscillatory (oo) motion at radius $R_0^c$ the oscillation period diverges as $T \sim |R_0 - R_0^c|^{-\nu}$ with $\nu \approx 2.5$. For larger radii higher harmonics of the fundamental oscillation period appear and the motion becomes increasingly irregular (Fig.4). This characterizes the complex oscillatory (co) regime, which is exemplified in Fig.5. The direction of island motion displays random shifts, which seem to be triggered by small fluctuations. This behavior is typical for large islands, and it is distinct from the periodic direction changes seen in the zig-zag (zz) regime for moderate sizes and small anisotropies (Fig.3).

The true long time behavior for large islands ($R_0 > 7$) and large anisotropy ($S > 5$) could not be pinned down unambiguously with our current numerical methods. Generally speaking, large islands with small anisotropy break up, while for large $R_0$ and $S$ faceted shapes undergoing irregular motion dominate.

The example shown in Fig.5 provides an important clue to the origin of the complex shape evolution. After an initial transient lasting until $t \approx 300$, the island settles down into a shape consisting of a straight upper edge and a lower edge which has broken up into a faceted hill-and-valley structure. The direction of island motion coincides with the orientation of the upper, straight edge, as shown for a smaller island in the upper panel of Fig.6. The key observation is that the hill-and-valley structure on the faceted edge does not move in the substrate frame. The moving island slides past the static facets, causing the shape to oscillate. Around $t \approx 900$ the roles of the upper and lower edges are seen to reverse, and the direction of motion changes.

Quite generally, large islands in the co regime can be constructed from four selected facet orientations $\theta_1, \theta_2, \theta_3, \theta_4$. Here $\theta_1$ and $\theta_2$ are the possible stable orientations of the upper island edge, which must satisfy

FIG. 4: Time series of the island perimeter showing regular and irregular oscillations. From bottom to top the parameters are $S = 2$, $R_0 = 5$; $S = 5$, $R_0 = 5$; and $S = 3$, $R_0 = 8$. The top panel corresponds to the run shown in Fig.5

FIG. 5: Complex oscillatory motion with $S = 3$, $R_0 = 8$. Light lines show the facet orientations predicted from \cite{9}, dark dashed lines illustrate that the hill-valley structure is static in the substrate frame. Consecutive snapshots are shifted upwards in time.
For the lower edge are obtained by reflection at the x-axis, \( \theta_3 = -\pi - \theta_1 \) and \( \theta_4 = \pi - \theta_2 \). To form a closed shape, at least three orientations must be combined, two of which are those two symmetry-related orientations that are closest to the horizontal direction (\( \theta = 0 \) or \( \pi \)). In Fig.6 we see a transition from a shape with orientations \( \{\theta_1, \theta_3, \theta_4\} \) to a shape with orientations \( \{\theta_1, \theta_2, \theta_3\} \).

The stable facet orientations can be computed along the lines of [27]. The condition \( v_n = 0 \) for a static shape implies that the current in (1) is set to a constant \( j^* \). Using the relation \( \kappa = d\theta/ds \) this can be brought into the form

\[
\Omega \gamma \frac{d^2}{ds^2} \theta = -|j^*/\sigma(\theta) - q^* E_0 \cos(\theta)| \equiv -V'(\theta),
\]

which describes the position \( \theta(s) \) of a particle moving in time \( s \) subject to a potential \( V(\theta) \) determined by the mobility and the electric field strength. As explained in [27], the coexistence of two stable facet orientations corresponds to a particle trajectory moving between two degenerate potential maxima. To determine the selected orientations, \( j^* \) is tuned until two degenerate maxima satisfying the above constraints appear. We have checked that this procedure correctly accounts for the facet orientations observed in the time-dependent calculations throughout the relevant region of the phase diagram (see Fig.5). In general, stable facets can be constructed from (3) only when the anisotropy is sufficiently large [27]. For \( n = 6, \alpha = 0 \) the requirement is \( S > S_c \approx 1.77 \). No stable facets are found when the number of symmetry axes is too small (\( n \leq 3 \)); this may explain why we do not see oscillatory shape evolution for a threefold anisotropy.

We close with two remarks concerning future research. First, we note that the observed island shapes are quite smooth, which implies that the number of circular harmonics involved in the shape evolution is small. It thus seems promising to attempt a description in terms of a low-dimensional dynamical system, in the spirit of [10], to gain a deeper understanding of the various migration modes and the bifurcations connecting them.

Second, we address the experimental conditions under which the predictions of this paper could be realized. As an example, we consider islands on Cu(100), for which most material parameters entering the theory are available. Following [8], we estimate that the electromigration force on an edge atom at a current density of \( 10^6 \text{Acm}^{-2} \) is about 400 eV/cm. Together with the experimentally determined stiffness [17] and mobility [18] for kinked steps at 300 K, this yields a characteristic length of \( L_k \approx 25 \text{ nm} \), and a time scale \( t_k \) on the order of seconds. Thus we expect complex shape dynamics to be observable for island radii around 100 nm and on time scales of a few hundred seconds. As a first step towards a more detailed description of specific surfaces, it would be important to identify oscillatory shape evolution in kinetic Monte Carlo simulations of island electromigration.

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