Model of Surface Instabilities Induced by Stress

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We propose a model based on a Ginzburg-Landau approach to study a strain relief mechanism at a free interface of a nonhydrostatically stressed solid, commonly observed in thin-film growth. The evolving instability, known as the Grinfeld instability, is studied numerically in two and three dimensions. Inherent in the description is the proper treatment of nonlinearities. We find that these nonlinearities can lead to competitive coarsening of interfacial structures, corresponding to different wave numbers, as strain is relieved. We suggest ways to experimentally measure this coarsening.

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Elastic effects can strongly influence the morphology of materials, and hence influence material properties. If nonequilibrium elastic energies build up, there are different ways for solids to release that energy. One is by plastic deformation, involving dislocations, and another is by elastic deformation, which is commonly seen in thin-film growth. A nonhydrostatically strained solid which is in contact with its own melt or vapor can partially release its elastic energy by a morphological instability at the interface. This strain relief mechanism gives rise to what appears to be a buckling of the surface into trenches, or interface. This strain relief mechanism gives rise to what appears to be a buckling of the surface into trenches, or isolands, of a particular spacing. It was first predicted by its elastic energy by a morphological instability at the interface. This strain relief mechanism gives rise to what appears to be a buckling of the surface into trenches, or isolands, of a particular spacing. It was first predicted by Asaro and Tiller [1]. Experimentally, it has been observed and studied by Torii and Balibar [2] who strained He crystals nonhydrostatically as well by Berrêhar et al. [3] in polymer crystals. Furthermore, it is often associated with the dislocation-free Stranski-Krastanov growth mode (also called island-on-layer mode) of epitaxially grown thin films as being observed for Ge/Si [4], InGaAs/GaAs [5], and InGaAs/InP [6]. Since the independent rediscovery of the instability by Grinfeld [7] and Srolovitz [8], it is often referred to as the Grinfeld instability.

Several approaches have been employed to study the instability. They are based either on static energy minimization calculations by a variational principle [7] or on a dynamical interface equation which describes mass transport, mainly surface diffusion, under the influence of the chemical potential which comprises surface free energy and elastic energy [8–14]. Linear stability analysis [8,10,11] predicts conditions for the onset of instability. Spencer and Meiron [13], and Yang and Srolovitz [14] studied the nonlinear evolution numerically, whereby the surface profile evolved to smooth flat peaks with sharp deep grooves. The numerical studies have been limited to dimension $d = 2$. Within the interface formulation, sharp cusps form within the grooves, leading to numerical stability problems [13] in this regime, where the interface formulation is potentially inapplicable. To make connection to the experiments, one requires a model which comprises a full nonlinear description, and which can be used in three dimensions.

In this paper, we present a Ginzburg-Landau phase-field model of the phenomena. An order parameter field $\phi(\vec{r})$ determines whether one is in a hard solid phase, which supports shear, or a soft disordered phase, hereafter called the liquid phase, which does not. The position of the interface coincides with the rapid variation of this field. Such an approach has been applied successfully to other moving-boundary-value problems, such as phase segregation and crystal growth [15]. Indeed, our model is numerically robust, can be implemented in three dimensions, and is readily generalizable. We show below that we recover the Grinfeld instability in linear and highly nonlinear regimes. We furthermore probe the transient dynamics during the morphological instability, finding that competitive coarsening of interface structures takes place. We suggest ways to measure this experimentally.

The physical mechanism for the stress-driven morphological instability can be understood easily. A stressed solid can partially relieve its stress by differentially moving material from valleys to hills, buckling at a particular wave number. In the less constrained peaks, lateral relaxation occurs, unlike in the more constrained valleys. The resulting stress gradient drives the instability by creating deeper valleys, thereby increasing the stress gradient, and sustaining the growth of the perturbation. At sufficiently small length scales, capillarity prevents the formation of sharp cusps.

The model we propose is based on a Ginzburg-Landau approach in which the elastic strain is a subsidiary tensor variable coupled to a nonconserved scalar order parameter. This approach is related to that of Onuki [16], Onuki and Nishimori [17], and Sagui, Somozza, and Desai [18], which was used to analyze elastic effects in phase-separating alloys [19]. The coarse-grained Ginzburg-Landau free energy is

$$F(\phi, u_{ij}) = \int d\vec{r} \left[ f(\phi, u_{ij}) + \frac{l^2}{2} |\nabla \phi|^2 \right],$$

where $f(\phi, u_{ij})$ is a function of the order parameter and the strain, and $l$ is a characteristic length scale of the instability.
where integration over \( \vec{r} \) is indicated by the subscript on the integral, \( u_{ij} = \frac{1}{2} (\partial u_i / \partial x_j + \partial u_j / \partial x_i) \) is the strain, and \( u_i \) is the displacement field. The bulk free energy density \( f(\phi, u_{ij}) \) is given by

\[
    f(\phi, u_{ij}) = \frac{1}{a} \phi^2 (\phi^2 - 1)^2 + \frac{e^2}{2\kappa} \nabla \cdot \vec{u} + f_e(\phi, u_{ij}), \tag{2}
\]

where the first term describes a three-well potential with \( \phi = 0 \) being the liquid and \( \phi = \pm 1 \) being the solid phase, ensuring that the liquid-solid phase transition is first order. The potential depths are determined by the model parameter \( a \) which, together with the parameter \( l \) being proportional to the surface tension, determines the interfacial thickness. The second term shifts the energy so that, for constant elastic coefficients, solid and liquid are at coexistence. The convenient choice \( g(\phi) = \frac{1}{2} \phi^2 - \frac{1}{4} \phi^4 \) guarantees [15] that both bulk phases keep their equilibrium values \( \phi = 0 \) (liquid) and \( \phi = \pm 1 \) (solid). The coupling constant \( e \) is related to the externally applied stress. The trace of the strain tensor is \( \nabla \cdot \vec{u} \), and \( f_e(\phi, u_{ij}) \) is the isotropic elastic free energy [20]:

\[
    f_e(\phi, u_{ij}) = \frac{\kappa}{2} (\nabla \cdot \vec{u})^2 + g(\phi) \mu \times \sum_{ij} \left( u_{ij} - \frac{\delta_{ij}}{d} \nabla \cdot \vec{u} \right)^2, \tag{3}
\]

where \( \kappa \) is the compressibility and \( \mu \) is the shear modulus in the solid phase alone. By construction, the shear modulus in the soft liquid phase is zero, whereas it stays nonzero and constant in the hard solid phase. Since the solid phase supports shear, whereas the liquid phase does not, our phase-field order parameter has a transparent meaning in the context of the liquid-solid transition.

It is reasonable to suppose that the elastic field relaxes much faster than \( \phi \). The elastic field can then be solved in terms of the order parameter using the condition of local mechanical equilibrium: \( \delta F / \delta u_i = \nabla_j \sigma_{ij} = 0 \), where a summation convention over repeated indices is implicit. The stress tensor, \( \sigma_{ij} = \delta F / \delta u_{ij} \), is then given by

\[
    \sigma_{ij} = \frac{\kappa}{\kappa} (\nabla \cdot \vec{u}) \delta_{ij} + 2 \mu g(\phi) \times \left( u_{ij} - \frac{\delta_{ij}}{d} \nabla \cdot \vec{u} \right). \tag{4}
\]

The solution of this to first order in the shear modulus is

\[
    \nabla \cdot \vec{u} = \text{Tr} \mathbf{A} - \frac{\epsilon}{\kappa} g(\vec{r}) + 2 \mu \frac{\epsilon}{\kappa^2} \int_{\vec{r}} \int_{\vec{r}'} G(\vec{r}, \vec{r}') \nabla_i \nabla_j \left[ g(\vec{r}') M_{ij}(\vec{r}', \vec{r}'') g(\vec{r}'') \right], \tag{5}
\]

where \( g(\vec{r}) = g(\vec{\phi}(\vec{r})) \),

\[
    \nabla_i u_j = A_{ij} - (\epsilon / \kappa) \nabla_i \nabla_j \int_{\vec{r}} G(\vec{r}, \vec{r}') g(\vec{r}'), \tag{6}
\]

\( \nabla^2 G(\vec{r}, \vec{r}') = \delta(\vec{r} - \vec{r}') \), and \( M_{ij}(\vec{r}, \vec{r}') = \nabla_i \nabla_j G(\vec{r}, \vec{r}') - (\delta_{ij}/d) \delta(\vec{r} - \vec{r}') \). In the absence of external strain, that is \( A_{ij} = 0 \), the solid will be stressed, whereas the liquid is stress-free. For a flat surface \( \phi = \phi(y) \), the solution of Eq. (6) in two dimensions is \( u_{xx} = u_{yy} = 0 \) and \( u_{yy}(y) = -(\epsilon / \kappa) g(y) \). Therefore, the solid will be uniaxially strained with \( \epsilon \) determining the strength of that strain.

The elastic field can now be expressed in terms of the order parameter. Substituting the solution for the strain \( \vec{g} = \phi \vec{y} \), the integral,

\[
    u_i = \int_{\vec{r}} \nabla_i G(\vec{r}, \vec{r}') g(\vec{r}') \, d\vec{r}', \tag{7}
\]

where \( G(\vec{r}, \vec{r}') \) is the Green function, results in an integral that is determined by the strength of that strain.

Numerical simulations on a discrete lattice were performed in two and three dimensions. Euler’s method was used for the integration in time. The Green function was solved in Fourier space. For all simulations presented here, the mesh size \( \Delta x = 0.01 \) or 0.005, the time step \( \Delta t = 0.01 \) or 0.05, \( \beta = 1.0, \) and \( \epsilon = 0.01 \). This choice of \( \Delta t \) and \( \epsilon \) guarantees that the surface is resolved by at least eight points. The parameter set \( (L_x, L_y, L_z, Y_0, c) \) will be specified below, where \( Y_0 \) gives the initial amplitude of the surface. Length scales will be measured in units of \( \Delta x \). Periodic boundary conditions were employed in all directions. Thus, the solid was in contact with its liquid phase at the bottom and at the top. It was ensured that the solid was sufficiently thick so that the interfaces at the top and bottom acted independently.

A numerical linear stability analysis was performed in two dimensions. The system was prepared with a small amplitude sinusoidal surface profile \( Y(x, t = 0) = Y_0 \sin(qx) \), where \( q \) is the wave number, and its subsequent
evolution was monitored. We found that the growth of the amplitude of the Fourier modes was initially independent and exponential, obeying \( \exp[\omega(q) t] \), followed by slower constant velocity growth. The fitted dispersion \( \omega(q) \) is consistent with \( \omega = Aq - Bq^2 \), where \( A \approx 28.2 \) and \( B = 1 \) (see Fig. 1). Perturbations with a wave number larger than a critical wave number are stabilized by surface tension, whereas wave numbers smaller than the critical wave number are unstable, therefore being a long wavelength instability. The flat interface, however, is stable. This agrees with the linear stability analysis carried out by Srolovitz [8] for the case where evaporation-condensation is the material transport mechanism, which is appropriate for our model. We have done a multiple-scale analysis to obtain the equations corresponding to the sharp-interface limit of our continuous field model [21]. The coefficients we obtain numerically for linear stability are the same, within our numerical accuracy, as those we obtain analytically.

Linear stability analysis predicts only the condition of the onset of instability. To study the later-stage morphology, a complete nonlinear description has to be employed. One advantage of the phase-field description is that nonlinearities are taken into account implicitly. A typical set of configurations is shown in Fig. 1. The nonlinear effect gives rise to a clear asymmetry between peaks and valleys, wherein deep grooves appear in the valleys. This behavior has been observed experimentally, as well as in previous theoretical studies [9,12–14]. Unlike previous studies, no numerical instabilities limit the study of the formation of the grooves here. However, we have not made a detailed investigation of regimes where grooves are pre-dominant. We expect that other phenomena not included in our model, such as the nucleation of dislocations, could become important there. It is interesting to note that in the early stages of the instability we can fit the interfacial profile with a simple function \( K = \sum_i a_i(t) Y^i \), where the curvature \( K = (Y''(x))/(1 + Y'(x)^2)^{3/2} \) is a low-order polynomial function of the height \( Y(x) \) of the interface.

Experimentally, random fluctuations in the interface will give rise to the competitive growth of different structures corresponding to different wave numbers. To study this, we prepared the system with an interfacial profile consisting of a superposition of \( p \) linearly unstable modes, \( Y(x) = Y_0 \sum_{i=1}^{p} \cos(q_i x + \phi_i) \) with \( q_i < q_c \) and \( \phi \) being a uniformly distributed random variable in the interval \([0, 2\pi]\). We did 100 runs over 500 time steps of a two-dimensional system with 100 unstable modes, where \((L_x, L_y, L_z, Y_0, c) = (1024, 512, 0, 0.24, 12.3)\). Figure 2 shows the Fourier transform of the equal-time height-height correlation function, which we shall call the structure factor \( S(q, t) \). For early times, there is a strong similarity between this behavior and early-stage spinodal decomposition in long-range force systems [22]; we show the results of a linear Cahn-Hilliard-type theory of the modes in the figure as well. Note that the structure factor vanishes for \( q \to 0 \) due to elasticity, not a conservation law. For later times, when the linear theory no longer describes the data, coarsening is evident: The location of the peak of the structure factor \( q_{\text{max}}(t) \) moves to smaller wave numbers, as the peak height increases and sharpenes. The peak height follows \( S(q_{\text{max}}, t) \sim t^{\alpha+1} \), where \( \alpha = 2 \), while the peak width sharpens with time as \( w \sim t^{-\gamma} \), where \( \gamma = 0.5 \). The former dependence is due to the total interface length increasing linearly with time for any unstable wave number. The latter dependence is due to competitive ordering between different wave numbers.
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