Two-Dimensional Self-Assembly in Diblock Copolymers

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Diblock copolymers confined to a two-dimensional surface may produce uniform features of macro-molecular dimensions (~10–100 nm). We present a mathematical model for nanoscale pattern formation in such polymers that captures the dynamic evolution of a solution of poly(styrene)-b-poly(ethylene oxide), PS-b-PEO, in solvent at an air-water interface. The model has no fitting parameters and incorporates the effects of surface tension gradients, entanglement or vitrification, and diffusion. The resultant morphologies are quantitatively compared with experimental data.

The functional flexibility and stability of polymeric films plays a crucial role in many commercial applications from protective coatings to dielectric layers in solid state electronics. While the stability properties of such films have been studied for many decades [1], a recent resurgence of interest in these systems [2–7] has arisen, driven partially by the miniaturization of technology. In order to better predict and control the self-assembled, two-dimensional structures that arise in polymeric films, we develop a new mathematical model that captures the evolution of a diblock copolymer monolayer at an air-water interface. Unlike the structures observed in three-dimensional and thin film phase segregation, the patterns in this two-dimensional system are dynamically selected and may be trapped in configurations that do not necessarily reflect thermodynamic equilibrium structures.

It is well known that diblock copolymers with immiscible blocks are excellent materials for the self-assembly of three-dimensional patterns due to the microphase separation that can produce features that are naturally nanometers in scale [8,9]. However, confining the material to a monolayer increases the interfacial contributions due to the substrate and the environment, and changes the mechanisms responsible for such features. Here we study the spreading of a diblock copolymer, poly(styrene)-b-poly(ethylene oxide), PS-b-PEO, at an air-water interface and the two-dimensional structures that result [10–13].

The proposed mechanism for this two-dimensional self-assembly differs fundamentally both from the microphase segregation that occurs in three-dimensional pattern formation [14] and from van der Waals driven spinodal decomposition, commonly associated with dewetting and rupture in ultrathin films. In classic phase segregation, an energy cost is associated with the interface between the two types of polymers; hence the blocks associate to form equilibrium structures such as lamallae or micelles that minimize this interfacial cost. In contrast, the final structures in the present PS-b-PEO system are kinetically trapped. When a droplet of polymer is deposited on the water surface via a solubilizing agent, the polymer rapidly adopts a bilamellate configuration with PEO at the air-water interface and the PS above in the air (see Fig. 1). This initial three-dimensional orientation process, driven by the interface, corresponds to the classic mode of phase segregation and occurs almost instantaneously. Subsequently, the droplet spreads at the aqueous interface, simultaneously increasing the local polymer solution concentration as the solvent evaporates and decreasing the local surface density as more area is covered.

Properties of the PS-b-PEO used in the experimental study [10] are listed in Table I. Solutions of a variety of diblock copolymers in chloroform were deposited on an ultrapure water surface in a Langmuir-Blodgett trough. The resultant monolayer films were compressed to a surface pressure corresponding to a PEO density below the phase transition to three dimensions. The film was then transferred to a silicon substrate and imaged using surface force microscopy (SFM). Sample images are shown in Fig. 2.

Depending on the percentage of PEO in the polymer and the initial concentration of the solution, a variety of morphologies were observed including dots, stripes, and larger structures known as continents. The predominant features that result exhibit characteristic length scales on the order of tens of nanometers and can be controlled by a judicious choice of relative block size and spreading solution concentration.

![FIG. 1. (a) When the solution contacts the water, individual polymers rapidly orient with the PEO at the air-water interface (black) and the hydrophobic PS (gray) in the solvent. (b) Collapsed PS chains after the solvent has evaporated.](image-url)
chains, and diffusion of the polymers along the air-water interface. Poly(ethylene oxide), PEO, is a surfactant that remains localized at the surface in a two-dimensional “pancake” provided the surface density does not exceed ~1 PEO monomer/20 Å² (see Fig. 1). Forces resulting from gradients in surface tension (Marangoni), due to variations in PEO concentration, are proportional to the local concentration gradient, $F_{ST} = \sigma(c) \nabla c$, where the coefficient, $\sigma(c)$, is measured experimentally [11]. The function $c(x, y, t)$ describes the local polymer concentration field, namely, the two-dimensional density of polymers at the interface. Surface tension as a function of concentration, $\gamma(c)$, was approximated, as $\gamma_{H_2O} - \gamma(c) = 5(1 - \tanh[1/(cR_{PEO}) - 4])$ mN/m, by fitting data from [10]. Integrating this function around the perimeter of a PEO pancake and Taylor expanding about a uniform initial concentration, $c_0$, we find the Marangoni force acting on a single polymer [16]:

$$F_{ST} = \pi R_{PEO}^2 \frac{\partial \gamma}{\partial c} \left| \frac{\nabla c}{c_0} \right| \equiv \sigma(c) \nabla c. \quad (1)$$

As the polymers spread, solvent continues to evaporate and the PS chains associate and begin to vitrify. This entanglement-vitrification force between two polymers is modeled as a non-Hookean elastic network [18], with the force between two polymers given by $f_E = 3kT(1/r - r/r_0^3)$ [19] where $T$ is the temperature, $k$ is Boltzmann’s constant, $l$ is the length of one monomer, $N$ is the number of PS monomers, and $r_0 = Nl^2$. Two polymers are considered “entangled” if the PS blobs overlap. Note that entanglement is indistinguishable from vitrification in this physical picture and the detailed mechanism for the connection between two polymers is irrelevant.

The radii of the PS blobs are constantly fluctuating due to thermal effects; thus the probability that a PS blob has an instantaneous radius, $r_i$, is given by $P(r_i) = 4\pi r_i^2(3/2\pi Nl^2)^{3/2} e^{-3r_i^2/2Nl^2}$. The expected value of the force between two polymers may then be written as $f(r) = \int \int \hat{f}(r)P(r_1)P(r_2)dr_1dr_2$ where $\hat{f}(r) = f_E(r_1) + f_E(r_2)$ if the polymers overlap and zero otherwise. Here $r$ is the separation between the centers of the two interacting polymers, and $r_1$ and $r_2$ are the instantaneous radii of the PS blobs. This expected force value, $f(r)$ [shown in Fig. 3(a) for an interaction between two 51 k polymers] is integrated over all pairwise interactions to obtain the effective spring force at a point $r_0$. The concentration field is then Taylor expanded about $r_0$ and only the two lowest order nonzero

| TABLE I. Properties of pure polymers used in experiments. $N_{PEO}$ is the number of PEO monomers. |
|---------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Total MW (in kDaltons) | 141.1 | 250.3 | 62.5 | 288 | 38.2 | 89.6 | 80 | 129.6 | 200 | 479 | 375 |
| % PEO (by mass) | 11 | 16 | 18 | 21 | 22 | 34 | 36 | 55 | 60 | 90 | 92 |
| MW PS (in thousands) | 125 | 211.5 | 51 | 227 | 29.8 | 58.6 | 51 | 58.6 | 80 | 46.5 | 30 |
| $N_{PEO}$ | 350 | 910 | 255 | 1375 | 190 | 690 | 655 | 1620 | 2725 | 9800 | 7840 |
contributions are retained, yielding the force due to vitrification or entanglement between two polymers:

$$F_E(r_0) = \pi \left( \alpha_2 \nabla^2 c + \frac{\alpha_4}{8} \nabla^4 c \right).$$  \hspace{1cm} (2)$$

where $\alpha_n$ is defined as the $n$th moment of the pairwise force function, $\alpha_n = \int_0^\infty r^n f(r) dr$. We expect this approximation to be valid provided the concentration field varies over a characteristic length scale that is long compared to the typical length scales associated with $f(r)$.

As the Reynolds number is very small ($< 10^{-5}$), the system is overdamped, and we neglect inertia of both polymer and fluid phases. In such a system, any velocity is a direct consequence of the instantaneous forces. We model an individual polymer as a sphere of radius $R_{PS}$ moving through chloroform:

$$v = \frac{F_{SP}(c) + F_E(c)}{6 \pi \eta R_{PS} \mu},$$

(3)

where $\mu$ is the viscosity of the chloroform and $R_{PS}$ is the radius of gyration of the PS.

Finally, this velocity in inserted into a two-dimensional transport equation for polymer concentration. Evaporation decreases the mobility of the polymers and can be included by multiplying the velocities by a phenomenological mobility envelope that decreases monotonically with time (e.g., $e^{-\beta t}$). Again, as with the entanglement force function, the resulting morphologies and length scales are not sensitive to the precise form of the mobility envelope. The final component of the model is to introduce a cutoff function, $f_c = [1 - \tanh(c - c_{\text{max}})]/2$, that reflects the fact that the PEO is incompressible, i.e., the maximum allowable surface concentration is set by the area of the PEO pancakes. Defining $\tau = (1 - e^{-\beta t})/(6 \pi \beta R_{PS} \mu)$ yields our final evolution equation:

$$\frac{\partial c}{\partial \tau} = \nabla \cdot \left[ f_c \left( \sigma - \pi \alpha_2 \nabla^2 c - \frac{\pi}{8} \alpha_4 c \nabla^4 c \right) + kT \nabla c \right],$$

(4)

where the diffusivity of the polymers along the air-water interface has been approximated as the Einstein diffusivity, $kT/(6 \pi \mu R_{PS})$.

Equation (4) may be reduced to a standard Cahn-Hilliard [21,22] form,

$$\frac{\partial c}{\partial \tau} = \nabla \cdot \left[ M(c) \nabla \left\{ \frac{\partial q}{\partial c} - K \nabla^2 c \right\} \right],$$

(5)

by defining $M(c) = cf_c$, $K = \frac{\pi}{8} \alpha_4$, and $q = (\sigma - \pi \alpha_2) c^2 + kT \ln c$. It is well known that Eq. (5) can lead to spinodal decomposition; i.e., uniformly mixed states may be metastable and evolve into phase segregated domains. However, it is important to note that the Cahn-Hilliard equation is often treated as a phenomenological model used to generically mimic phase separation, whereas the coefficients in Eq. (4) have been completely determined by ensemble averaging over microscopic physical states. Thus all coefficients are fixed by well-defined physical parameters, and the observed characteristic length scales may be predicted from known quantities.

Typical numerical results obtained by evolving the full nonlinear Eq. (4) are shown in Fig. 2. Parameters used in the simulations are summarized in Table I. The numerics capture both the dot and the stripe morphologies and the transitions between the two (i.e., pearling). At the moderate concentrations used in the theoretical treatment, the features evolve quickly to dilute stripes and, if possible, transition into denser dots as observed in the experiment. There are two control parameters that determine whether the stripe-to-dot transition is allowed for a fixed percentage PEO; the cutoff function, defined by the maximum surface density allowed by the presence of the incompressible PEO, and the initial bulk concentration of polymer in solvent. In the absence of a cutoff function, the polymers can always collapse into dots spaced in a regular hexagonal lattice as there is no limit to the packing density; stripes can freely coalesce into energetically favorable dots. We expect the polymers to become locked in the stripe morphologies when the maximum concentration attained within the stripes approaches the maximum allowed concentration. This indicates that stripes will be observed if the initial bulk concentration of polymer in solvent is sufficiently high, consistent with experimental observations. In addition, polymers are mobile only in the presence of solvent. If there is insufficient solvent at the onset of the experiment, the chloroform rapidly evaporates, freezing the polymer in the stripe morphology before the transition to dots is...
complete. In intermediate regimes, in which the stripes begin to break into dots but are arrested by the cutoff, pearling is observed as shown in Fig. 2.

Finally, we consider the linear stability of (4). Characteristic dot diameters, determined by the fastest growing mode, are given by

$$\lambda_{\text{crit}} = \frac{2\pi}{|k|_{\text{crit}}} = \frac{\pi\alpha_4}{\left(\pi\alpha_2 - \sigma'(c_0) - kT/c_0\right)^{1/2}}. \quad (6)$$

This wavelength can be converted into an aggregate number, $AN = \pi(\lambda_{\text{crit}}/2)^2c_0$, the expected number of polymers/dot. Experimental data and linear stability predictions are summarized in Fig. 3(b). Parameters used in the stability analysis are all known (summarized in Table I) with the exception of $c_0$, the initial surface density, which was estimated as $c_0 = 0.4/N_{\text{PEO}}$ molecules/Å$^2$, consistent with experimental observations. Unlike many phenomenological models that produce dot and stripe morphologies, the present study is quantitative in the sense that dot size and aggregation number are completely predicted by known microscopic properties of the polymer such as number and length of the monomers and by macroscopic properties of the system (e.g., temperature and initial concentration). The most influential parameter in determining the aggregate number is the ratio of PS to PEO and by varying the percentage of PEO, one can continuously tune the size of the observed structures [23].

While there are many other physical mechanisms that may lead to phase separation, the most common do not produce our experimentally observed scalings. For example, spinodal decomposition driven by van der Waals effects could produce similar patterns with similar length scales. However, van der Waals forces are several orders of magnitude smaller than entanglement-vitrification effects for all thicknesses sampled by the film during evaporation [see inset in Fig. 3(a)] and the wavelengths of van der Waals driven structures scale as $h^2$ [3], which does not reflect the dependence on the percentage of PEO that is observed experimentally [Fig. 3(b)].

The ability to control patterned structures by self-assembly on the molecular scale by noncovalent forces can serve as a powerful tool in developing nanoscale technologies. We have developed a system in which the behavior of diblock copolymers in two dimensions can be optimized to produce regular, uniform features of macro-molecular dimensions ($\sim 10-100$ nm) and derived a new dynamic model that quantitatively captures the experimentally observed length scales and diversity of features. It is hoped that this new understanding will eventually lead to insights into mechanisms that could provide more detailed control of the observed structures using macroscopic manipulations (such as shear in the underlying fluid layer) to control microscopic properties.

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[16] Although it is not at all obvious that a continuum concept of surface energy can be extended down to molecular scales, experimental evidence in conjunction with simple scaling arguments presented in [17] indicate that this picture remains valid even for an isolated molecule.
[19] The observed patterns and length scales are not sensitive to the exact form of the entanglement-vitrification force function, and variations such as non-Hookean entropic springs [20] have also been tested. All of these models yield quantitatively similar results provided the force function scales linearly with $kT$.
[23] The average spacing between the dots can be controlled independently by the depositional pressure. Higher pressures correspond to lower mean molecular areas and higher packing densities.