Pattern formation by phase-field relaxation of bending energy with fixed surface area and volume

Timothy Banham*

West Virginia Wesleyan College, 59 College Ave, Buckhannon, West Virginia 26201, USA

Bo Li†

Department of Mathematics and Center for Theoretical Biological Physics, University of California, San Diego, 9500 Gilman Drive, MC 0112, La Jolla, California 92093-0112, USA

Yanxiang Zhao‡

Department of Mathematics, George Washington University, Washington, DC 20052, USA

(Received 6 December 2013; revised manuscript received 28 July 2014; published 17 September 2014)

We explore a wide variety of patterns of closed surfaces that minimize the elastic bending energy with fixed surface area and volume. To avoid complicated discretization and numerical instabilities for sharp surfaces, we reformulate the underlying constrained minimization problem by constructing phase-field functionals of bending energy with penalty terms for the constraints and develop stable numerical methods to relax these functionals. We report our extensive computational results with different initial surfaces. These results are discussed in terms of the reduced volume and are compared with the known results obtained using the sharp-interface approach. Finally, we discuss the implications of our numerical findings.

DOI: 10.1103/PhysRevE.90.033308
PACS number(s): 02.60.–x, 87.10.Pq, 87.16.dm, 87.17.Aa

I. INTRODUCTION

Bending energy contributes crucially to physical and biological properties of closed surfaces. Examples of such properties in biology include the biconcave shape of a red blood cell and the different equilibrium states of cell membranes [1–6]. Macroscopically the bending energy of a closed surface is often modeled by the surface integral of the square of mean curvature (i.e., the average of two principal curvatures). This integral is the principal term in the widely used Canham-Helfrich functional, an integral over the surface of a quadratic polynomial of mean curvature [1,7]. One of the interesting problems related to the interfacial phenomenon is the minimization of bending energy with fixed surface area and enclosed volume [6,8,9]. In this work we study numerically such a problem to explore a variety of different patterns.

The numerical implementation for minimizing the bending energy of closed surfaces, with or without constraints, is in general very challenging as it amounts to solving a problem of geometrical flow, the Willmore flow [10]. This is a nonlinear fourth-order partial differential equation. With a usual sharp-interface formulation and a fixed finite-difference spatial grid, the numerical discretization of such an equation can be very complicated, and the stability of numerical solution is hard to achieve. An alternative approach is to use a phase-field representation of the surface [11–13]. This means that a phase field, a continuous function defined on the entire computational domain, takes values close to one constant (say, 0) outside the closed surface and another constant (say, 1) inside, but smoothly varies its values from one of the constants to another in a thin transition region that represents the surface. Such an approach has been widely used in studying surface and interface problems arising in many scientific areas, such as materials physics, complex fluids, and biomolecular systems; cf. Refs. [11–27] and the references therein.

In our current work, we develop a phase-field model to minimize the bending energy of a closed surface with fixed surface area and enclosed volume. We use the phase-field description of the bending energy that has been mathematically analyzed thoroughly in Refs. [28–31]. We enforce the surface-area and volume constraints by penalty terms. This is similar in part to the method used in Ref. [30] but is different from some other methods, such as the Lagrange multipliers method used in Refs. [20,22,31]. In Ref. [31] the volume constraint results from a Model-B-like formulation of the underlying relaxation dynamics, involving high-order spatial derivatives. One of the reasons that we use penalty terms is for easier numerical implementation. We minimize our phase-field functional by solving the gradient-flow partial differential equations, using a finite-difference spectral method. We report our extensive numerical results of a wide variety of equilibrium patterns resulting from minimizing the bending energy with fixed surface area and enclosed volume in three-dimensional space (or fixed perimeter and enclosed area in two-dimensional space). In three-dimensional space, which is of most practical interest, these patterns are analyzed using the reduced volume (i.e., the ratio of volume to that of the unit ball). In particular, we compare our results with the known, sharp-interface results analyzed thoroughly in Refs. [28–31]. We enforce the surface-area and volume constraints by penalty terms. This is similar in part to the method used in Ref. [30] but is different from some other methods, such as the Lagrange multipliers method used in Refs. [20,22,31]. In Ref. [31] the volume constraint results from a Model-B-like formulation of the underlying relaxation dynamics, involving high-order spatial derivatives. One of the reasons that we use penalty terms is for easier numerical implementation. We minimize our phase-field functional by solving the gradient-flow partial differential equations, using a finite-difference spectral method. We report our extensive numerical results of a wide variety of equilibrium patterns resulting from minimizing the bending energy with fixed surface area and enclosed volume in three-dimensional space (or fixed perimeter and enclosed area in two-dimensional space). In three-dimensional space, which is of most practical interest, these patterns are analyzed using the reduced volume (i.e., the ratio of volume to that of the unit ball). In particular, we compare our results with the known, sharp-interface results for the three-dimensional axisymmetric case [8].

The rest of this paper is organized as follows: In Sec. II we describe our phase-field energy functionals and the related gradient flows. In Sec. III we present briefly our numerical methods. In Sec. IV we report and analyze our computational results. Finally, in Sec. V we draw conclusions.

II. PHASE-FIELD ENERGY FUNCTIONAL AND RELATED GRADIENT FLOW

We consider the minimization of bending energy of closed surfaces, possibly with multiple connected components, that
have fixed surface area $A$ and fixed volume $V$ enclosed by the surface, where $A$ and $V$ are two positive constants. Let $\varepsilon$ be a positive number such that $\varepsilon \ll 1$. Let $\Omega$ denote our computational domain in $\mathbb{R}^2$ or $\mathbb{R}^3$. We define the phase-field functional of all smooth functions $u = u(x)$ $(x \in \Omega)$:

$$E_\varepsilon[u] = \int_{\Omega} \frac{\kappa}{2\varepsilon} \left[ -\varepsilon \Delta u + \frac{1}{\varepsilon} W(u) \right]^2 \, dx$$

$$+ \frac{1}{2} M_A(\varepsilon) [A_\varepsilon(u) - A]^2 + \frac{1}{2} M_V(\varepsilon) [V_\varepsilon(u) - V]^2,$$

(2.1)

where

$$A_\varepsilon(u) = \int_{\Omega} \left[ \frac{\varepsilon}{2} |\nabla u|^2 + \frac{1}{\varepsilon} W(u) \right] \, dx,$$

(2.2)

$$V_\varepsilon(u) = \int_{\Omega} u(x) \, dx.$$  

(2.3)

The first term in (2.1) approximates the bending energy. In this term the parameter $\kappa > 0$ is the bending modulus, and $W(u) = 18au^2(1-u)^2$ is a double-well potential with the “two wells” 0 and 1. The second term in (2.1) is a penalty term. It enforces the surface area to be $A$. The quantity $M_A(\varepsilon)$ is a function of $\varepsilon$ such that $M_A(\varepsilon) > 0$ and $M_A(\varepsilon) \rightarrow \infty$ as $\varepsilon \rightarrow 0$. The term $A_\varepsilon(u)$ defined in (2.2) is a phase-field description of the area of interface separating the regions where a phase field $u$ takes the values 0 and 1, respectively. With the prefactor chosen precisely to be 18 in (2.4), it is known that $A_\varepsilon[u]$ converges (in certain sense of functional convergence called $\Gamma$-convergence) to the surface area in the sharp-interface limit (i.e., in the limit $\varepsilon \to 0$) [32–34]. The third term in (2.1) is also a penalty term. It enforces the volume to be $V$, since $V_\varepsilon(u)$ defined in (2.3) approximates the volume of the region defined by $u \approx 1$. The quantity $M_V(\varepsilon)$ is positive and depends also on $\varepsilon$. Moreover, $M_V(\varepsilon) \rightarrow \infty$ as $\varepsilon \to 0$.

Heuristically, if the first term (the integral term) in the functional $E_\varepsilon[u]$ is small, then the integrand will be close to 0. That means $u$ solves approximately the Euler–Lagrange equation corresponding to the variational problem of minimizing $A_\varepsilon(u)$ that is defined in (2.2). This minimization leads to $W(u) \approx 0$ for $\varepsilon \ll 1$. Therefore, $u \approx 1$ or 0 in $\Omega$. Mathematical studies indicate that $A_\varepsilon(u)$ is now close to the interfacial area, and $V_\varepsilon(u)$ is close to the volume of the region where $u \approx 1$ [32–34].

To minimize the functional (2.1), we consider the relaxation dynamics, i.e., the gradient flow

$$\frac{\partial u}{\partial t} = -M(\varepsilon) \frac{\delta E_\varepsilon}{\delta u},$$

(2.5)

where $t$ is the time, $M(\varepsilon) > 0$ is a mobility constant that can depend on $\varepsilon$, and $\delta E_\varepsilon/\delta u$ is the variational derivative of $E_\varepsilon$. Direct calculations lead to

$$\frac{\delta E_\varepsilon}{\delta u} = \frac{\kappa}{\varepsilon} \left[ \varepsilon \Delta u - \frac{1}{\varepsilon} W''(u) \right] \varepsilon \Delta u - \frac{1}{\varepsilon} W'(u)$$

$$+ M_A(\varepsilon) [A_\varepsilon(u) - A] \varepsilon \Delta u + \frac{1}{\varepsilon} W'(u)$$

$$+ M_V(\varepsilon) [V_\varepsilon(u) - V].$$

(2.6)

Combining (2.5) and (2.6), we therefore obtain the time-dependent equation that we need to solve

$$\frac{\partial u}{\partial t} = -\frac{M(\varepsilon)\kappa}{\varepsilon} \left[ \varepsilon \Delta u - \frac{1}{\varepsilon} W''(u) \right] \left[ \varepsilon \Delta u - \frac{1}{\varepsilon} W'(u) \right]$$

$$- M(\varepsilon)M_A(\varepsilon)[A_\varepsilon(u) - A] \varepsilon \Delta u + \frac{1}{\varepsilon} W'(u)$$

$$- M(\varepsilon)M_V(\varepsilon)[V_\varepsilon(u) - V].$$

(2.7)

We choose our computational domain $\Omega$ to be a box in $\mathbb{R}^2$ or $\mathbb{R}^3$ and use the periodic boundary condition. Initial solutions $u^{(0)} = u^{(0)}(x)$ $(x \in \Omega)$ will be specified in the numerical computation. As the functional $E_\varepsilon$ is highly nonconvex, different initial solutions can lead to different final, stable, steady-state solutions that are local minimizers of the functional $E_\varepsilon$.

### III. NUMERICAL METHODS

In this section we describe briefly our numerical methods for solving the gradient-flow equation (2.7). We present our methods for the three-dimensional case and omit those, similar and simpler, for the two-dimensional case. We choose our computational box to be a cube $(-L/2, L/2) \times (-L/2, L/2) \times (-L/2, L/2)$ for some $L > 0$ and cover it by a uniform finite-difference grid with grid size $h$ in each direction. We choose a time step $\Delta t > 0$ and denote $t_m = m\Delta t$ $(m = 0, 1, \ldots)$. For a given function $u(x, t)$, we denote by $u^{(m)} = u^{(m)}(x)$ an approximation of $u(x, t_m)$.

We rewrite Eq. (2.7) into

$$\frac{\partial u}{\partial t} = -\varepsilon M(\varepsilon)\kappa \Delta^2 u + B[u],$$

where

$$B[u] = \frac{M(\varepsilon)\kappa}{\varepsilon} \left[ \Delta(W'(u)) + W''(u)\Delta u - \frac{1}{\varepsilon^2} W'(u)W''(u) \right]$$

$$- M(\varepsilon)M_A(\varepsilon)[A_\varepsilon(u) - A] \varepsilon \Delta u + \frac{1}{\varepsilon} W'(u)$$

$$- M(\varepsilon)M_V(\varepsilon)[V_\varepsilon(u) - V].$$

We design accordingly our semi-implicit splitting scheme to discretize the time variable for Eq. (2.7):

$$\frac{u^{(m+1)} - u^{(m)}}{\Delta t} = -\varepsilon M(\varepsilon)\kappa \Delta^2 u^{(m+1)} + B[u^{(m)}],$$

$$m = 0, 1, \ldots,$$

where $u^{(0)}$ is a given initial solution. Here all the nonlinear terms, collected in the $B$ term, are treated explicitly. Rearranging terms, we obtain

$$u^{(m+1)} + \Delta t \varepsilon M(\varepsilon)\kappa \Delta^2 u^{(m+1)} = u^{(m)} + \Delta t B[u^{(m)}],$$

$$m = 0, 1, \ldots$$

(3.1)

For a fixed $m$, we solve Eq. (3.1) using a finite-difference Fourier spectral method. To do so, we discretize spatially both sides of Eq. (3.1), and then use the discrete Fourier transform to obtain the corresponding system of linear equation in the Fourier space. This is a diagonal system so that it can be solved...
directly. Finally, we transform back to the real space to obtain the numerical solution to (3.1).

We choose our constant mobility $M(\varepsilon) = 1/\varepsilon$. To stabilize our numerical computations, we vary the two penalty parameters $M_A(\varepsilon)$ and $M_V(\varepsilon)$ in the time iteration (3.1). For instance, we set

$$M_A(\varepsilon, m) = M_V(\varepsilon, m) = 0.1m \quad (3.2)$$

and replace $M_A(\varepsilon)$ and $M_V(\varepsilon)$ in (2.7) by $M_A(\varepsilon, m)$ and $M_V(\varepsilon, m)$, respectively. After certain number of steps, say, $m = 1000$, we freeze $M_A(\varepsilon)$ and $M_V(\varepsilon)$ in (3.1).

Algorithm

1. Select a tolerance $\delta > 0$. Select an initial solution $u^{(0)}$ and compute the energy $E[u^{(0)}]$. Set $m = 0$.
2. Solve Eq. (3.1) for $u^{(m+1)}$ using the Fourier spectral method.
3. Calculate the energy $E[u^{(m+1)}]$.
4. If $|E[u^{(m)}] - E[u^{(m+1)}]| < \delta$ then stop. Otherwise, change $m$ to $m + 1$ and go back to step (2).

IV. COMPUTATIONAL RESULTS

A. Two-dimensional (2D) results

We first present our 2D computational results, where the volume and area reduce to the 2D perimeter and area, respectively. We fix the area $V$ and select different values of perimeter $A$. For instance, we fix $V = \pi$ and choose $A$ to be $2.5V$, $4V$, and $5V$, respectively. For each value $A/V$ we run our code. We choose our computational domain to be $\Omega = (-3,3)^2$ and cover it by a uniform grid of $128 \times 128$ grid points. We set $\varepsilon = 0.1$, $\kappa = 1$, $M(\varepsilon) = 0.5$, and $\Delta t = 10^{-5}$. The initial phase-field function $u^{(0)} = u^{(0)}(x)$ is taken to represent a circle, ellipse, or “bean.”

Figures 1–3 display some of our computational results. In each row of these figures, the first picture is the initial configuration and the last picture is the steady-state solution. The two pictures in between are some intermediate states of $u$. If the red region $u$ is 1 and in the blue region $u$ is 0. We observe from these computational results a diverse family of minimum-energy configurations which depend on the initial configuration and increase in complexity as the perimeter-to-area ratio increases. In particular, we see clearly from Figs. 1 and 2 that the final equilibrium shapes with higher perimeter-to-area ratios have more small pieces that have larger total perimeters.

B. Three-dimensional (3D) results

In the 3D computations, we fix the constant area $A$ in the functional $E_2$ to be $4\pi r^2$, the area of unit sphere, and use the reduced volume $V_{red} = V/(4\pi r^2)$ as a new parameter, where $V$ the constant volume in the functional $E_2$ and $4\pi r^2$ is the volume of unit ball. We choose the computational domain $\Omega = (-1.5,1.5)^3$ and cover it by a uniform grid of $128 \times 128 \times 128$ grid points. Our other parameters are $\varepsilon = 0.2$, $\kappa = 1$, $M(\varepsilon) = 0.5$, and $\Delta t = 10^{-5}$.

We choose five different types of initial phase-field functions $u^{(0)} = u^{(0)}(x)$. The first four of them are in the form of

$$u^{(0)}(x) = 0.5 + 0.5 \tanh \left( \frac{r - \text{dis}(x)}{\varepsilon / 3} \right),$$

where the function $\text{dis}(x)$ and the parameter $r$ are given by the following:

1. Sphere:
   $$\text{dis}(x) = \sqrt{x_1^2 + x_2^2 + x_3^2} \quad \text{and} \quad r = 1,$$
   where $x = (x_1,x_2,x_3)$;
2. Prolate:
   $$\text{dis}(x) = \sqrt{\left(\frac{x_1}{2}\right)^2 + x_2^2 + x_3^2} \quad \text{and} \quad r = 1;$$
3. Oblate:
   $$\text{dis}(x) = \sqrt{x_1^2 + x_2^2 + (2x_3)^2} \quad \text{and} \quad r = 1;$$
4. Parachute:
   $$\text{dis}(x) = \sqrt{x_1^2 + x_2^2 + x_3^2} \quad \text{and} \quad r = 1 + \sin \theta,$$
   where
   $$\theta = \arctan \left( \frac{x_3}{x_1^2 + x_2^2 + 0.01} \right).$$

The last initial phase-field function is chosen to represent a cube.
5. Cube:
   $$u^{(0)}(x) = \begin{cases} 1 & \text{if } |x_1| < 1, |x_2| < 1, |x_3| < 1, \\ 0 & \text{otherwise.} \end{cases}$$
We first consider the axisymmetric 3D geometries. In Fig. 4 we present the final equilibrium configurations with different reduced volumes and oblate initials. Each row displays a sequence of four snapshots of numerical computations at different times, with the first being the initial shape and the last the steady-state configuration. The perimeter-to-area ratios are 2, 3.5, 4, 5.5, and 6.5, respectively, from top to bottom.

In Fig. 5 we start with a parachute initial. For $V_{\text{redu}} = 0.75$ and 0.60 (top two rows), the equilibriums are parachute shape. Note that a smaller reduced volume corresponds to a smaller open hole at the south pole. Once the reduced volume becomes even smaller (bottom row), the open hole at south pole merges and the equilibrium becomes a concentric sphere.
FIG. 6. (Color online) The bending energy vs reduced volume $V_{\text{redu}}$. The three branches of prolate, oblate, and parachute shapes are displayed. The two critical values of reduced volume from prolate to oblate and from oblate to parachute are $V_{\text{pr} \leftrightarrow \text{ob}} \approx 0.65$ and $V_{\text{ob} \leftrightarrow \text{pa}} \approx 0.597$, respectively. These results agree well with those obtained using the sharp-interface model [8].

Figure 6 shows a phase diagram of the axisymmetric 3D equilibriums. There are three curves that correspond to three different initial shapes: prolate, oblate, and parachute. Each curve is the bending energy (defined to be the final minimum energy divided by $8\pi \kappa$) versus the reduced volume. The 2D cross sections of three representative solutions correspond to the reduced volume $V_{\text{redu}} = 0.8, 0.6, 0.57$, respectively. We observe that, for $V_{\text{redu}} \geq V_{\text{pr} \leftrightarrow \text{ob}} \approx 0.65$, the prolate shape is energetically the most favorable one. For the reduced volume $V_{\text{redu}} \in (V_{\text{ob} \leftrightarrow \text{pa}}, V_{\text{pr} \leftrightarrow \text{ob}}) \approx (0.597, 0.65)$, the oblate shape has the lowest bending energy. If $V_{\text{redu}} \leq V_{\text{ob} \leftrightarrow \text{pa}} \approx 0.597$, the parachute shape is the most stable one. The two critical values of the reduced volume, $V_{\text{ob} \leftrightarrow \text{pa}}$ and $V_{\text{pr} \leftrightarrow \text{ob}}$ agree with the

FIG. 7. (Color online) Configurations with different reduced volumes, all started with a unit sphere. Each row displays a sequence of four snapshots of numerical computations at different times, with the first being an initial unit sphere and the last the steady-state configuration. The reduced volumes are 0.423, 0.411, 0.333, 0.250, and 0.167, respectively, from top to bottom.

FIG. 8. (Color online) Configurations with different reduced volumes, all started with a cube. Each row displays a sequence of four snapshots of numerical computations at different times. The reduced volumes are 0.45, 0.35, and 0.25, respectively, from top to bottom.

FIG. 9. (Color online) Configurations with different reduced volumes, all started with an ellipsoid. Each row displays a sequence of four snapshots of numerical computations at different times, with the first being an initial ellipsoid and the last the steady-state configuration. Small 2D images are cross sections of the corresponding 3D shapes. The reduced volumes are 0.600, 0.577, 0.429, 0.333, and 0.328, respectively, from top to bottom.
FIG. 10. (Color online) The dependence of the relaxed (local) minimum energy and the corresponding configurations on the reduced volume. Each initial is an ellipsoid. The reduced volumes of the representative equilibriums are 0.600, 0.577, 0.556, 0.536, 0.517, 0.500, 0.484, 0.469, 0.455, 0.441, 0.429, and 0.333, respectively, from left to right.

Note that analytical formulas can be obtained for the sharp-interface limit of some of the phase-field energy minimizers. For instance, if the energy minimizer is a torus, then the distance $R$ from the center of the tube to that of the torus and the radius $r$ of the tube are given by $R = \frac{3}{2\pi V_{\text{redu}}}$ and $r = 2V_{\text{redu}}/3$, respectively. If the energy minimizer is a concentric sphere, we can solve the system of equations $R^2 + r^2 = 1$ and $R^3 - r^3 = V_{\text{redu}}$ to determine the outer and inner radii $R$ and $r$, respectively.

We now explore more 3D patterns that are the (local) minimizers of our phase-field bending-energy functional with constrained volume and area. In Fig. 7 we start from a unit sphere and take five different values of the reduced volume. In each of the five cases, the sphere evolves into a small ball with six handles which become thinner and longer as the reduced volume decreases. In Fig. 8 we start from a cubic initial. With three different values of the reduced volume, we observe even more complicated equilibrium shapes. In Fig. 9 the initial shape is an ellipsoid. We observe the topological changes in the relaxation dynamics.

It is clear that the initial shapes determine the final shapes. For instance, final shapes with sphere initials tend to be more rounded or at least more symmetric; cf. Fig. 7. Final shapes with ellipsoid initials are more elongated in one direction, the same as in the initial ellipsoid; cf. Fig. 9. Moreover, we observe that for smaller reduced volumes the final equilibrium surfaces are in general smoother and have larger curvatures. These result from the balance of minimizing the bending energy and satisfying the surface-area and volume constraints. Breaking into more components of surface increases the surface area while keeping the same volume, and also smooths the surface, reducing the bending energy.

Figure 10 shows the dependence of the final equilibrium bending energy and the corresponding equilibrium shape on the reduced volume.

Note that analytical formulas can be obtained for the sharp-interface limit of some of the phase-field energy minimizers.

FIG. 11. Cross sections of an evolving surface. Initial profile is a sphere.

FIG. 12. Phase-field evolution of a surface. Initial profile is an ellipsoid (not shown here). Gray lines show the boundary of the interior regions and the shape of the back side of the configuration.
ent spontaneous curvature
medium values of reduced volume, since for such values
constraints. High-energy configurations often correspond to
minimization of bending energy and the surface-area and volume
smoothness result from the competition between the mini-
mum surface area while keeping the volume fixed. For a larger
configurations. Comparison with existing results indicates
reveal a wide range of different energy-minimizing
computation more stable. Our extensive numerical results
implemented a semi-implicit, finite-difference, Fourier
spectral method to solve numerically the gradient flow,
and study the resulting functional $E_\varepsilon$. Note that $\varepsilon |\nabla u|^2 \approx W(u)/\varepsilon$ for small $\varepsilon > 0$. This implies that the terms inside the
pair of brackets are approximations to $-\varepsilon \Delta u + W'(u)/\varepsilon - H_0 |\nabla u|$. The integral of the square of this sum, with an
effective parameter $H_0$, is exactly the bending energy with
the spontaneous curvature, proposed as in Eq. (2.1) by [35].
Figure 13 shows our phase-field computational results that
include the parameter of spontaneous curvature $H_0$.

We point out that our phase-field relaxation is a method
of steepest descent. Like any such a method, it can only
capture locally energy-minimizing states for which suitable
initial states are known. Consequently, many local minimizers
may not be discovered as the energy landscape can be very
complicated. In fact, for a given reduced volume, we have
seen from our computations that there are often many different
local minimizers of the bending energy functional. Moreover, a
method of steepest descent does not in general describe the real
dynamics. Therefore, it is often unable to sample all the states
to provide the “ensemble average” such as the statistically
weighted energy value.

Nevertheless, what we have presented is a first step toward
understanding a complicated system. In addition to exploring
different surface patterns as it is done in this work, we can
apply our relatively simple and reliable computational model
and method to study other problems, such as the stability
of vesicles [36,37]. In order for our phase-field approach to
be able to treat more realistic systems, however, we have to
develop new method to describe the surface fluctuations [38].
These are among the directions we plan to pursue.

V. CONCLUSIONS

We have constructed a phase-field bending-energy
functional with penalty terms enforcing the constraint of
surface area and volume. We have also designed and
implemented a semi-implicit, finite-difference, Fourier
spectral method to solve numerically the gradient flow,
minimizing the phase-field functional. The use of penalty
terms with dynamic parameters of penalty makes the
computation more stable. Our extensive numerical results
reveal a wide range of different energy-minimizing
configurations. Comparison with existing results indicates
that our models and methods are accurate and reliable.

In general, initial shapes determine the final, locally energy-
minimizing shapes. With a smaller reduced volume, the final
surface breaks up into several components to increase the
surface area while keeping the volume fixed. For a larger
reduced volume, the final surface often has one component
(no break up) but can have large curvatures. Break up and
smoothness result from the competition between the mini-
mization of bending energy and the surface-area and volume
constraints. High-energy configurations often correspond to
medium values of reduced volume, since for such values
no break up is needed to satisfy the constraints but more
complicated topologies appear.

We remark that the spontaneous curvature, an important
quantity in membrane modeling, can be included approx-
imately in our model and numerical implementation. For
instance, we can replace the bending energy term, the $\kappa$ term,
in (2.1) by [35]

$$E_\varepsilon(u) = \int_{\Omega} \frac{\kappa}{2\varepsilon} \left[ -\varepsilon \Delta u + \frac{1}{\varepsilon} W'(u) - \varepsilon H_0 |\nabla u| \right]^2 dx,$$

where $H_0$ is a constant representing the spontaneous curvature,
and study the resulting functional $E_\varepsilon$. Note that $\varepsilon |\nabla u|^2 \approx W(u)/\varepsilon$ for small $\varepsilon > 0$. This implies that the terms inside the
pair of brackets are approximations to $-\varepsilon \Delta u + W'(u)/\varepsilon - H_0 |\nabla u|$. The integral of the square of this sum, with an
effective parameter $H_0$, is exactly the bending energy with
the spontaneous curvature, proposed as in Eq. (2.1) in [35].

Finally, in Figs. 11 and 12 we display drawings that
semble our computational results to show how the surface is
evolves during the relaxation. Figure 11 shows a sequence of
cross sections of six three-dimensional configurations during
the time evolution of relaxation dynamics starting with a
sphere. Figure 12 shows a sequence of six configurations
during the time evolution of relaxation dynamics starting with
an ellipsoid. For a high reduced volume, the phase field usually
stops evolving at phase 2 or 3. For a low reduced volume,
the phase field progresses to phase 5 or 6. Again, with the
same volume, equilibrium shapes with lower reduced volumes
often have more branches to create larger surface areas. It is
interesting to see from Fig. 11 that an initial sphere evolves
finally to a sphere inside a concentric shell. All of these
configurations are spherically symmetric. But many of the
intermediate configurations are not spherically symmetric.
The existence of such particular intermediate states may depend
on the choice of numerical parameter $\varepsilon$ that characterizes the
width of diffuse surface and the dynamic parameters of penalty
$M_\lambda(\varepsilon,m)$ and $M_\nu(\varepsilon,m)$.

ACKNOWLEDGMENTS

This work was supported by the U.S. National Science
Foundation (NSF) through Grant No. DMS-1319731, the NSF
Center for Theoretical Biological Physics (CTBP) through
NSF Grant No. PHY-0822283, and the National Institutes
of Health through Grant No. R01GM096188. The authors
would like to thank the anonymous referees for their valuable
comments and suggestions.

(1976).
[34] B. Li and Y. Zhao, SIAM J. Appl. Math. 73, 1 (2013).