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Capillary Origami: Spontaneous Wrapping of a Droplet with an Elastic Sheet

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The interaction between elasticity and capillarity is used to produce three-dimensional structures through the wrapping of a liquid droplet by a planar sheet. The final encapsulated 3D shape is controlled by tailoring the initial geometry of the flat membrane. Balancing interfacial energy with elastic bending energy provides a critical length scale below which encapsulation cannot occur, which is verified experimentally. This length is found to depend on the thickness as $h^{3/2}$, a scaling favorable to miniaturization which suggests a new way of mass production of 3D micro- or nanoscale objects.

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Origami and haute couture design both rely on folding and assembling planar material to create elegant three-dimensional shapes whose variety and complexity are governed by the number, order, and orientation of folds. Folding is also a way to reduce the size of deployable structures in space industry (satellite pannels or sun sails [1]) as well as in nature (plant leaves folded in a bud [2]). In micro and nanofabrication, the folding of planar structures is a promising approach to build 3D objects, since most microfabrication technologies produce planar layers through surface etching [3].

While centimeter-sized objects may be folded using well disposed magnets [4], capillarity is a particularly relevant mechanism for microfabrication [5], as surface forces dominate over bulk forces at small scales. Indeed, the capillary attraction and stiction of wet slender structures may induce disastrous damage in micro and nanoscale devices [6–9], in addition to being involved in lung airway closure (neonatal respiratory distress syndrome) [10]. Capillary interactions have already been proposed as a way of assembling and orienting rigid objects in 2D at the surface of water [11] or in 3D by capillary bridges [12], while further studies have focused on the capillary induced deflection of elastic rods [13–15]. Recently, the multifolding of a flexible ribbon squeezed in a meniscus has also been described through an elegant experiment [16]. Here, we address the effect of capillary forces on a flat elastic membrane and we show how the spontaneous folding of a flexible sheet around a liquid droplet leads to a predetermined 3D shape.

Our experiments were conducted using polydimethylsiloxane (PDMS) membranes. The PDMS (Dow corning Sylgard 184, 10:1 polymer/curing agent mix) was spin coated at 24 °C on a glass microscope slide at rotation rates of 1000–2000 rpm. Once the PDMS was cured, this resulted in sheets with thickness in the range 80–40 μm.

During a typical experiment, a geometric shape is manually cut out from the PDMS layer and placed on a superhydrophobic surface. A drop of water, of volume 1–80 μL depending on membrane size, is deposited on the PDMS, while making sure that the water reaches all the corners of the flat sheet. The water is then allowed to evaporate at room temperature and a time lapse sequence of images is taken of the membrane-drop pair. As the water volume decreases from its initial value to complete evaporation, the surface tension of the liquid pulls the sheet around smaller volumes, thus increasingly curving it (Fig. 1). Sufficiently thin sheets eventually encapsulate the liquid with a shape that depends on the initial cut of the membrane.

In the simplest case, a drop is deposited on a square PDMS sheet, as shown in Fig. 1(a). This initially leads to the bending of the four corners towards the center of the drop. The final encapsulated 3D shape is controlled by tailoring the initial geometry of the flat membrane. Balancing interfacial energy with elastic bending energy provides a critical length scale below which encapsulation cannot occur, which is verified experimentally. This length is found to depend on the thickness as $h^{3/2}$, a scaling favorable to miniaturization which suggests a new way of mass production of 3D micro- or nanoscale objects.

![FIG. 1. Wrapping of a drop of water with (a) square and (b) triangular PDMS sheets. See movies Mov1 and Mov2 in Ref. [17].](image-url)
that we refer to as the elasto-capillary length [14]. Large deformations of the system are obtained when \( L \gg L_{EC} \). This scaling argument thus indicates that the critical size \( L_{crit} \) beyond which the sheet totally wraps the liquid droplet should be proportional to \( L_{EC} \), the only characteristic length in the problem.

The elasto-capillary length can be directly measured through a calibration experiment rather than by estimating each parameter in Eq. (1) which would produce large errors. The experiment consists in depositing a drop of wetting liquid onto a very long strip of PDMS that folds over into a loop with a self-contacting tail. We consider the late stage, when only a small meniscus still holds the loop together [Fig. 2(a)].

This equilibrium state is 2D, so the following analysis is understood to be per unit depth. The cross section follows planar rod elasticity coupled with surface tension. We note \( \theta(s) \) the angle made by the tangent to the rod \( t \) with respect to the horizontal, at curvilinear coordinate \( s \). We also note \( R \) the constant vectorial tension of the beam, so that Euler’s elastica equations for equilibrium may be expressed as [18]:

\[
B \frac{d^2 \theta}{ds^2} \mathbf{e}_z + t \times R = 0, \tag{2}
\]

where \( \mathbf{e}_z \) is the unit vector perpendicular to the plane of the section. The global shape of the loop is determined by elasticity, given the curvature at the contact point. In turn, this curvature is found from an equilibrium between surface tension and elasticity by considering a virtual displacement \( dx \) of the contact point for a loop of curvilinear length \( 2l \) [see Fig. 2(b)]. The elastic energy, \( E_{el} = \int \kappa \kappa^2 / 2 ds \), varies due to the change in the integral bounds, the integrand being stationary near elastic equilibrium. It may be written as \( dE_{el} = B \kappa^2 dx \), where \( \kappa \) is the curvature at the contact point. On the other hand, the variation of surface energy due to replacing a solid-gas interface (surface energy \( \gamma_{SG} \)) with a solid-liquid interface (\( \gamma_{SL} \)) is \( 2(\gamma_{SL} - \gamma_{SG})d \gamma = 2 \gamma \cos \alpha dx \), where \( \alpha \) is the contact angle of the liquid on the surface (here we use ethanol which has a zero contact angle on the PDMS).

Balancing the two energies yields the value of equilibrium curvature at the contact point: \( \kappa_c = \sqrt{2/L_{EC}} \).

The boundary value problem of Eq. (2) with the above boundary condition was solved numerically by enforcing the contact condition at the base of the loop, \( \int_0^\ell \sin(\theta)ds = 0 \). \( L_{EC} \) was then measured by scaling the numerical solution to fit the experimental shape. An excellent agreement is obtained [dashed line superimposed on experimental picture in Fig. 2(a)]. In practice the width of the loop (theoretical value \( 0.89L_{EC} \)) is sufficient to determine \( L_{EC} \).

The value of \( L_{EC} \) was measured for different membrane thicknesses and, for each thickness, the minimum length for encapsulation \( L_{crit} \) was determined for square and triangular membrane shapes (Fig. 3). The experimental

![FIG. 2. (a) Experimental image of a loop of PDMS held by a meniscus of ethanol and comparison with the theoretical shape (dashed curve). (b) Virtual displacement \( dx \) of the meniscus position.](image)
results are consistent with a linear relationship between $L_{\text{crit}}$ and $L_{\text{EC}}$ as expected from dimensional analysis [see discussion with Eq. (1)]. Based on this, a linear regression of the data leads to $L_{\text{crit}} = 7.0 L_{\text{EC}}$ for squares and $L_{\text{crit}} = 11.9 L_{\text{EC}}$ for triangles (Fig. 3).

Recovering the value of the critical length theoretically would require a full 3D description of the coupled thin plate and drop system. Here, we limit ourselves to a 2D model, which provides an understanding of the transition between the folding and reopening processes. Consider an elastic inextensible rod of length $L$ bent by pressure and surface tension forces at its ends (inset in Fig. 4). Equation (2) still holds but the vectorial tension $\mathbf{R}$ is not constant anymore due to the pressure forces; it follows

$$\frac{d\mathbf{R}}{ds} + \rho \mathbf{n} = 0,$$

(3)

where $\mathbf{n}(s)$ is the unit vector normal to the rod. By neglecting gravity, the pressure $\rho$ in the drop is uniform and the 2D shape of the liquid-air interface is a circular arc of radius $r = \gamma/\rho$.

Equations (2) and (3), with appropriate boundary conditions, are solved numerically and the distance $\delta$ between the rod tips is calculated as a function of drop section area $S$ for different rod lengths $L$, as shown in Fig. 4. Two trivial flat states ($\delta/L = 1$) exist in all cases for an infinitely large drop ($A$ and $A'$) or when no liquid is present ($B$ or $B'$). However, the evolution between these states is qualitatively different depending on $L/L_{\text{EC}}$.

For $L/L_{\text{EC}}$ slightly below the critical value [Fig. 4(a)], these two states are connected by a continuous family of solutions from $A$ to $B$: the drying of a large drop leads to transient limited bending but eventually to reopening as it is experimentally observed (Mov3 in EPAPS [17]). Closed states $C$ do exist in a restricted domain of the phase space, as the stable segment of a solution loop, while the unstable part of this loop corresponds to a drop which has depinned from the edges $D$. This bistability is also observed in the experiments: while small sheets never close spontaneously, they can be forced into a closed state if the liquid is pulled out with a syringe.

Above the critical ratio $L/L_{\text{EC}}$, the drying of a large drop $A'$ leads continuously to complete wrapping $C'$ in agreement with experimental observations (Fig. 1 and Mov1 in EPAPS [17]). Another branch links the open state $B'$ to unstable solutions $D'$. However, this branch is difficult to observe experimentally because it requires the spreading of a thin film of water on the flat membrane.

The 2D numerical simulation thus qualitatively reproduces the experimentally observed regimes and explains the closing-reopening transition. However, the crossing from diagrams (a) to (b) was found at $L_{\text{crit}} = 3.54 L_{\text{EC}}$, which is of the same order but below the value experimentally obtained for the square membranes. An improved model should include gravity and 3D effects.

Indeed, the drops considered in Fig. 3 are slightly flattened by gravity, since they are larger than the capillary length ($L_{\text{c}} = \sqrt{\gamma/\rho g}$, where $\rho$ is the density of the liquid and $g$ the acceleration of gravity). Moreover, the early stages are always 3D as seen in the initial folding of the corners of a square sheet. Only a full description of thin plate elasticity in 3D would account for the localization of the curvature around edges, which is reminiscent of the crumpling of thin sheets.

More generally, 3D wrapping of a droplet leads to a geometrical incompatibility described by Gauss’s theo-

![FIG. 3. Critical length for folding vs elasto-capillary length: black squares: square sheets, black triangles: triangular sheets, lines: linear regressions expected from dimensional analysis.](156103-3)
**FIG. 5.** Tuning of the initial flat shape to obtain (a) a spherical encapsulation, (b) a cubic encapsulation, or (c) a triangular mode-2 fold.

*remaegrium.* A classic example is that it is impossible to draw a planar map of the earth which conserves distances: the wrapping of a spherical object with a planar sheet must involve stretching as well as bending. For a thin membrane, the stretching energy scales as the thickness $h$ and is therefore much more expensive than the bending energy which scales as $h^3$. The unavoidable stretching is therefore localized in singular regions, leading to crumpling singularities [19,20]. We expect that for $L \gg L_{EC}$, the number of singularities will increase since surface tension becomes dominant over elastic forces, leading to quasispherical crumpled encapsulated shapes.

Precise engineering of the final closed state may be obtained by tailoring the initial sheet geometry. In this way, it is possible to approximate a smooth sphere by starting with a flower shape, as shown in Fig. 5(a). Figure 5(b) shows how to form a cube by starting from a cross shape. As in the case of the pyramid, evaporation after encapsulation takes place at a reduced rate and may buckle the sphere or the cube due to the negative pressure thus generated. However, one may also freeze (or polymerize) the encapsulated liquid at any point in this process, therefore fixing the desired shape in place. Finally, small perturbations of the initial shape may also be used to yield different final states. In particular, rounding off two opposite corners of a square sheet leads to the membrane closing along its diagonal [Fig. 5(c)] rather than parallel to the sides. These experiments suggest that a wide variety of final shapes may be achieved through careful tuning of the initial flat shape.

In summary, three-dimensional submillimetric objects are produced from flat membranes through the interaction of elasticity and capillarity. The limitation on the minimal size for folding is determined by the elasto-capillary length [Eq. (1)] which scales as the sheet thickness $h^{3/2}$. This scaling is favorable to miniaturization since thinner membranes lead to much smaller critical lengths. This opens the way to mass production of microscale 3D structures based on standard microfabrication methods since a wide variety of objects can be fabricated by tailoring the initial planar sheet.

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