Computational Modelling of a Spanning Drop in a Wedge with Variable Angle

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Abstract
In this work, mathematical and computational techniques were used to model the phenomenon of a bubble or liquid drop as it slides to the narrowest side between two tilted plates. Practical applications of this phenomenon, such as the adjustment of liquid propellant tanks, are also discussed. Computations involved the use of Surface Evolver—a fully open, interactive program that allows the simulation of physical experiments in a virtual environment.

Keywords
Surface Evolver, Computational Modeling, Spanning Drop, Wedge

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INTRODUCTION

It is frequently easy to perform experiments with bubbles and soap films, which show intriguing behaviors. For instance, two co-axial parallel rings spanned by a soap film can produce a surface called a catenoid. This is a surface of revolution around the axis of the rings, and its name derives from catenary, which is the natural curve that a chain forms when its extremes are fixed at a certain height. The word chain is catena in Italian, and this explains the origin of the name. Besides the catenoid, two parallel rings can also be spanned by another continuous soap film, as illustrated by Isenberg (1978). However, this example is not entirely smooth due to its triple junctions.

A catenoid is also formed when two parallel plates are bridged by a soap film (Figure 1). A very curious fact is that, if one of the plates is tilted a small amount the soap film will slide to the edges of the plates that have the narrowest distance between them.

One practical application of this property of thin films in engineering is found in liquid propellant tanks. Unlike soap bubbles, a liquid drop can bridge non-parallel plates that form a small angle. Increasing this angle is equivalent to increasing the tilt of the plates. Eventually, as in the case of the soap film described above, the liquid drop slides to the edges of the plates that have the least distance between them. A fuel supplier in the shape of a wedge can be used to access small quantities of fuel as needed (Figure 2). By increasing the angle of the wedge, the fuel slides to the outlet of the supplier. Then the angle of the wedge can decrease again, so that another small portion of fuel is launched into the supplier, and then the supplier waits for the next access. This process is analogous to a pair of bellows that expels fuel instead of air.

Another factor that influences the way a liquid drop moves when the angle is varied is the so-called wetting property of each plate (Good, 1992). This can be described as a balance between how much the drop keeps its own cohesion and how much it adheres to a contact surface. Wetting and contact forces together produce an important effect called capillarity, which we mention in the next section.

The focus of the work discussed in this paper is using computational techniques to model the phenomenon of a bubble or liquid drop as it slides to the narrowest side between two tilted plates. In order to build our model we made use of a publicly available computer program called Surface Evolver. This program was created by Kenneth Brakke, professor at Susquehanna University, Department of Mathematical Sciences (USA). Surface Evolver allows the simulation of physical experiments in a virtual environment, and its source code is fully open (Brakke, 1994, susqu.edu/brake). In the next section we explain the strategies that were used to write the code of our simulator.1

METHODS

Surface Evolver reads data files that specify parameters, quantities, constraints, geometrical elements and scripts. In order to
write the data file that will simulate the phenomenon of interest for this work, the problem must be first described from a mathematical point of view.

This phenomenon can be described mathematically using a few geometrical properties. In doing so, some simplifying assumptions are made such as ignoring the thickness of the soap bubble, gravity applied to the liquid drop, etc. Even with these approximations, the essential ideal properties of the phenomenon can be understood.

The first requisite geometrical property is called mean curvature. Consider a point p in a smooth surface S. For each plane P that contains the normal line n to S through p, we have a planar curve S∩P.

Of course, the set of all such planes can be parameterized by q in the interval [0,p] with P(0) = P(p). The curvature of P(q)∩S at p is a continuous function k(q) that attains its minimum k₁ and maximum k₂ within the compact interval [0,p] (Figure 3). The mean curvature of S at p is defined as H(p) = (k₁+k₂)/2.

The second requisite geometrical property is the contact angle between the drop and the plates. Liquids are endowed with capillarity, a concept that is beyond the scope of this paper. However, mathematically speaking, this means that a drop that is rotationally invariant around an axis of symmetry makes a constant angle with its surface of contact. Rotational invariance does not apply to our configuration, but we will take constant contact angle as a simplification of our model. In practice, the contact angle is not constant. The (external) contact angle increases when we approach the apex of the wedge. This is the point that advances to the apex, and for this reason it is called advancing contact angle. It decreases when we approach the opposite point of contact, where we have the receding contact angle (Luo et al., 2014). Of course, this is what happens when the wetting property is uniform on the whole surface of contact. We could get a constant contact angle if the wetting property had a suitable variation over that surface.

Figure 4 shows the geometry of interest in this paper. We consider that the contact angle γ₁ between the drop S and the upper plate Π₁ has a constant value throughout the intersection S∩Π₁, and the same holds to the contact angle γ₂ between S and the lower plate Π₂. We call α the angle between the plates.

Of course, by neglecting gravity we should a priori have γ₁ = γ₂. But this is still not the case if we consider that the plates have different wetting properties. Hence, in our simulations their constant values are not necessarily equal.

It is important to notice that Figure 4 shows a profile not in accordance with Luo et al. (2014), who conclude that both γ₁ and γ₂ should be acute. However, soon we will present Theorem 1 of McCuan (1997), which states that under certain hypotheses we cannot have γ₁ + γ₂ ≤ π + α, and
consequently $\gamma_1 + \gamma_2$ cannot be less than $\pi$. This contradiction occurs because the theorem refers to a mathematically ideal model that includes some simplifications.

McCuan (1997) establishes $S$, $\Pi_1$, $\Pi_2$, $\gamma_1$, $\gamma_2$ and $\alpha$ as above. Moreover, he supposes that $\gamma_1$, $\gamma_2$ have constant values and that $S$ is embedded and has constant mean curvature. In our context embedded simply means “without self-intersections.”

The hypothesis of embedding has to do with the fact that, in nature, we do not typically find drops with knots. There are soap films with self-intersections. If a closed wire is dipped into a soap solution the resulting soap film is not generally embedded. However, this is a special case and not one that is relevant for this work.\(^2\)

The hypothesis of constant mean curvature (HCMC) is grounded in a physical experiment that was first performed by the French mathematician Jean Baptiste Meusnier in 1776. His experiment consisted of observing the behavior of an elastic membrane in the middle of a glass tube. He could inject and remove gas either above or below the membrane, so that each level was under different constant pressures $P_1$ and $P_2$. The membrane spanned a non-planar curve, as depicted in Figure 5.

![Figure 5. Elastic membrane in a glass tube.](image)

Meusnier observed that $P_2 - P_1 = \lambda H$, where $\lambda$ is a constant that depends only on the material and the thickness of the membrane. Namely, the surface that separates two ambients at pressures $P_1$ and $P_2$ has constant $H$. In the case of a drop the liquid is denser than the surrounding air. The effect is the same, and so the surface $S$ of the drop has constant $H$. For a soap film, however, $P_2 = P_1$ and what we have is a so-called minimal surface $S$. Notice that in this case $S$ is concave (see Figure 1) and that $k_1 = -k_2$. This characterizes saddle points (see Figure 3).

Because of these particularities, and using the term ring type surface to denote a topological annulus, McCuan (1997) assumes that $S$ is of ring type and also verifies the following hypotheses: it is an embedded surface of constant $H$ which spans a wedge of opening angle $\alpha$ and meets the planes $\Pi_1$ and $\Pi_2$ of the wedge at constant contact angles $\gamma_1$ and $\gamma_2$, respectively. With this setting he proves

**Theorem 1.** No embedded ring type spanner $S$ of constant $H$ and constant contact angles can exist if $\gamma_1 + \gamma_2 \leq \pi + \alpha$.

**Theorem 2.** A spherical spanner exists if and only if $\gamma_1 + \gamma_2 > \pi + \alpha$. Letting $A > 0$ denote either enclosed volume or mean curvature, the family of all spanning spheres may be indexed uniquely by the set of 4-tuples $(\gamma_1, \gamma_2, \alpha, A)$ for which this existence criterion is satisfied.

Of course, spherical rings $S$ clearly verify the inequality of Theorem 2. The strength of this theorem resides in its second statement, for it characterizes any spherical ring by means of a 4-tuple. Now, if $S$ has constant $H$ and is a compact embedded ring type spanner in a wedge, and also meets the planes of the wedge at constant angles along its boundary, then $S$ must be a spherical ring according to (Park, 2005).

In Theorem 2, notice that $A$ can be either $H$ or $\text{Vol}(S)$. In fact, either of these two quantities determines the other because $\text{Vol}(S) = -\pi/(3H^3)\left[\cos\gamma_1(2 + \sin^2\gamma_1) + \cos\gamma_2(2 + \sin^2\gamma_2)\right]$. This equation does not
depend on \( \alpha \), which is however necessary to determine the spherical ring.

As mentioned previously, we assume that gravity is negligible for both soap films and drops. Of these, liquid drops would be the most massive. A drop has a volume of about 50ml. If we assume it is made of mercury, one of the densest liquids (13.6kg/m\(^3\)), then it weighs just 0.68 mg. In order to see that it is negligible, just change lines 1 and 88 of our datafile as we will explain below.

Our datafile `sdrop.fe` begins as follows:

```
GRAVITY_CONSTANT 0.0
PARAMETER va   = pi/6
PARAMETER gam1 = pi/2+1
PARAMETER mcv  = 1
PARAMETER fa   = va
PARAMETER gam2 = pi+fa+0.5-gam1
#define av -pi/(3*mcv^3)*(cos(gam1)'*(2+(sin(gam1))^2)+cos(gam2)*(2+''(sin(gam2))^2))
```

These are initial values like \( \alpha \) (variable alpha), \( \tau \) (fixed alpha), \( \text{mcv} \) (mean curvature) and \( \text{av} \) (annulus volume). In Surface Evolver we usually call “parameter” a value that does not depend on a formula. The value of a defined macro is always computed through a formula. Interaction with the Surface Evolver is via command line at a prompt. Hence, at the Surface Evolver prompt you can invoke and change \( \text{va}, \text{fa}, \text{etc}. \) by typing `print mcv, gam1 := pi/1.8, etc`. The macro \( \text{av} \) is not recognized at the prompt but only internally by the program. The symbol `\'` means “line break”. In Surface Evolver you turn gravity on by replacing the above 0.0 with 1.0.

Next in our datafile we have:

```
constraint 1 /* over Pi 2 */
formula: z = 0
energy:
e1: cos(gam2)*y
e2: 0
e3: 0
```

This means that Evolver will try to construct a triangulated surface with volume as close as possible to \( \text{av} \). Namely, we will work with a triangulated surface and the more triangles we have, the more precise is the computed volume. Although fixed, whenever we change \( \text{mcv}, \text{gam1}, \text{etc}. \), \( \text{av} \) is re-calculated and gives a new fixed value for \( \text{fvol} \). This quantity will be computed with method facet_vector_integral as follows:

\[
\int_S (q_1, q_2, q_3) \cdot N dS = \int_V \text{div}(x, 0, 0) dV
= \int_V \left( \frac{\partial x}{\partial x} + 0 + 0 \right) dV = \int_V 1 dV = \text{av}.
\]

In applying Gauss’ Theorem in (1), the ring type spanner \( S \) can only comprise a closed volume if we add the circles \( C_1 \) and \( C_2 \) that correspond to \( S \cap \Pi_1 \) and \( S \cap \Pi_2 \), respectively. If added to the surface integral in (1), their integrand would be \((x, 0, 0) \cdot (0, -\sin \alpha, \cos \alpha) = 0 \) over \( C_1 \) and \((x, 0, 0) \cdot (0, 0, -1) = 0 \) over \( C_2 \).

Our choice of the integral field \( q_1, q_2, q_3 \) allows Surface Evolver to compute the volume by numerical integration over \( S \).

Next in our datafile we have:

```
constraint 1 /* over Pi 2 */
formula: z = 0
energy:
e1: cos(gam2)*y
e2: 0
e3: 0
```

The reason for these commands is the same as explained by Brakke (1994). But for the convenience of the reader we will detail it here. Notice that \( \Pi_2 \) is equated as \( z=0 \), and the surface tensions \( T_1, T_2 \) over \( C_1 \) and \( C_2 \), respectively contribute to the overall energy that Surface Evolver seeks to minimize. This energy acts on the closed surface \( S \cup C_1 \cup C_2 \), but in Section 3.3 of Brakke (1994) the author explains the convenience of omitting
additional parts like $C_1$ and $C_2$, and working with line integrals along their boundaries in order to get the same numerical values. Because of that we will compute the surface integrals over $C_1$ and $C_2$ by means of Stoke's Theorem. The surface tension that acts on $C_2$ corresponds to the force $(0,0,T_2)$, where $T_2 = -\cos \gamma_2$. Therefore,

$$\int_{C_2} (0, 0, T_2) \cdot d\vec{S} = \int_{\partial C_2} (-T_2 y, 0, 0) \cdot d\vec{r}. \quad (2)$$

Similarly, $\Pi_1$ is equated as $z=\tan(\alpha) \cdot y$ and over $C_1$ the surface tension corresponds to $(0,-T_1 \sin \alpha,T_1 \cos \alpha) = T_1 \vec{N}$, where $\vec{N}$ is the unitary normal and $T_1 = \cos \gamma_1$. Now we choose $(e_1, e_2, e_3)$ in such a way that the dot product of its rotational field with $\vec{N}$ will result in $T_1$:

$$\text{constraint 2} /* \text{over } \Pi_1 */$$

formula: $z = \tan(\alpha) \cdot y$

energy:

e1: $\cos(\gamma_1)/\sin(\alpha) \cdot z$
e2: 0ne3: 0

The final three constraints are purely geometrical:

$$\text{constraint 3}$$

formula: $x = 0$

$$\text{constraint 4}$$

formula: $x = 3$

$$\text{constraint 5}$$

formula: $y^2+z^2 = 9$

They mean that we will display both $\Pi_1$ and $\Pi_2$ as squares of edge length 3, and $\Pi_1$ can only vary inside the cylinder $y^2+z^2 = 9$.

Now we define a prototype of $S$: a polyhedral ring surface with 4 facets. It looks like a bottomless box without a lid. Surface Evolver displays it as in Figure 6. In order to define this prototype we begin by declaring its vertices.

vertices

1 1.0 1.0 0.0 constraint 1
2 2.0 1.0 0.0 constraint 1
3 2.0 2.0 0.0 constraint 1

The vertices that represent $\Pi_2$ by a 3×3 square must come in the sequel:

4 1.0 2.0 0.0 constraint 1
5 1.0 1.0 1.0 constraint 2
6 2.0 1.0 1.0 constraint 2
7 2.0 2.0 1.0 constraint 2
8 1.0 2.0 1.0 constraint 2

The vertices that represent $\Pi_2$ by a 3×3 square must come in the sequel:

9 0.0 0.0 0.0 constraints 1,2,3 fixed
10 3.0 0.0 0.0 constraints 1,2,4 fixed
11 3.0 3.0 0.0 constraints 1,4,5 fixed
12 0.0 3.0 0.0 constraints 1,3,5 fixed

Notice that vertices 9 and 10 are shared with the 3×3 square that represents $\Pi_1$. Therefore, we only need to add another two vertices.

13 3.0 3*\cos(\alpha) 3*\sin(\alpha) constraints 2,4,5 fixed
14 0.0 3*\cos(\alpha) 3*\sin(\alpha) constraints 2,3,5 fixed

After having defined the vertices we need to list the edges, faces and bodies (body is the term that Evolver uses to link facets as parts of the same object). For the sake of conciseness they will be omitted here.

At line 88 of our datafile sdrop.fe the reader can add density 0.0136 to the body. This corresponds to a mercury drop as commented beforehand. Now, by running the simulation with GRAVITY_CONSTANT 1.0 the reader will not see any significant
changes in volume, area or shape of the surface. That is why we can neglect gravity.

In the next section we explain some of our scripts, which will make the initial surface evolve to a spherical annulus S, vary the angle between $\Pi_1$ and $\Pi_2$, and show how this affects S in a dynamic way.

RESULTS

In the Methods section we discussed the beginning of our datafile, in which important parameters are defined, especially the mean curvature $m_{cv}$ and the annulus volume $av$. Notice that $m_{cv}$ is only used to define $av$ and does not appear elsewhere in the code. Thus, we could have fixed a numerical value for $av$ and the simulation would have given the same output independent of the HCMC.

Moreover, our simulation gives evidence that S must be spherical, a fact that was formally proved by Park (2005) but still under the HCMC. According to the results of our program, the HCMC may be unnecessary. However, many theoretical results rely on it, perhaps as a means of making progress in otherwise extremely difficult proofs. For instance, we cite:

1. Concus et al. (2001) deal with instability of tubular liquid bridges between parallel plates.

2. Finn (2002) lists and proves eight geometrical and physical properties regarding capillarity.

3. Luo and Heng (2014) describe the invention of a technique to separate oil from water by pressing plates on the mixture. The perfect separation relies on the difference of pressure inside the liquid drop. In our mathematical model we could add different constant mean curvatures to simulate the effect of these different pressures.

Such simplifications have already shown some contradiction with what happens in practice, as mentioned in the Methods section. However, Surface Evolver allows additional complexity beyond the simple example shown in this work. For professional purposes, we believe that Evolver achieves virtual simulations that can come much closer to what is observed in real experiments.

We mentioned the wetting property in the Introduction section and our program apparently neglects that important parameter. However, in the Methods section we explained that, although in practice the angles $\gamma_1$ and $\gamma_2$ are not really constant, the fact that we can take $\gamma_1$ different from $\gamma_2$ is a way of considering an effect of the wetting property. Hence, despite our simplifications we take that property into account by considering one of its effects.

Therefore, our code is very likely to improve both the theoretical and practical techniques that rely on the HCMC. Namely, from the numerical point of view, spherical surfaces of constant mean curvature automatically arise.

DISCUSSION

We have presented a simple physical experiment simulated with Surface Evolver. Such an experiment can be easily reproduced in a laboratory but perhaps only for the special case of soap bubbles. Drops are too small to work with, and in fact our practical example in engineering, given in the Introduction section, includes a narrow slot at the apex of the wedge. We will implement this slot in future work.

Moreover, notice that we cannot force prescribed angles $\gamma_1$ and $\gamma_2$ for a soap bubble without the addition of extra constraints as has been demonstrated using soap films and loops of thread. Each thread will assume the fixed shape of a circumference and so we cannot use our simulator to show what happens when $\alpha$ varies above the threshold. Namely, the exact case in which both circles $C_1$ and $C_2$ turn out to be tangent to each other.
We hope to have motivated students and professionals to use the Surface Evolver as a virtual environment for their physical experiments.

REFERENCES


1 Our code is named sdrop.fe and can be downloaded from our homepage https://sites.google.com/site/chrondon1992 but it needs installation of Evolver 2.70 to run, preferably in the operating system Linux Ubuntu 14.04 or higher. The user can install the older version 2.30 through the Ubuntu software center. Alternatively, just enter the command line sudo apt-get install evolver at the terminal prompt.

2 See susqu.edu/brakke/knots for some illustrations.

3 Here we show how to perform the simulation. Our main scripts are: gogo, olid and omre. Basically, each of them consists of a list of single-letter commands, of which the meanings are explained by Brakke (1994). Therefore, we will omit details here and focus on the functionality of our scripts.

Now you should install Evolver and download our file sdrop.fe, as explained at the Introduction. Open a terminal, go to the directory where sdrop.fe is and then type evolver sdrop at the prompt. You will get an interactive picture as illustrated in Figure 6. Now run the first script at the Evolver prompt, which is gogo. This script refines and smoothens our prototype until we see a virtual drop inside the wedge, as depicted in Figure 7.

The opening of the virtual wedge is \( \alpha = \pi/6 \). In the datafile sdrop.fe there are two variables that refer to \( \alpha \), namely \( va \) and \( fa \). Their names are a mnemonic to variable and fixed alpha, respectively. We made this distinction in order for our next scripts to open the wedge by increasing the value of \( va \). However, we keep \( fa \) fixed because the values of \( \gamma_1 \) and \( \gamma_2 \) are fixed and satisfy the equality \( \gamma_1 + \gamma_2 = 0.5 + \pi + \alpha \), for the initial value \( fa \) of \( \alpha \). In the datafile they are...
mcv = 1, gam1 = $\pi/2 + 1$ and gam2 = $\pi + fa + 0.5 - gam1$, respectively. With this setting we begin our simulation according to Theorem 2. The inequality $\gamma_1 + \gamma_2 > \pi + \alpha$ is equivalent to $va < fa + 0.5$, which is true at the beginning since $va = fa = \pi/6$.

Now run the script `olid`, which will “open the lid” $P_1$ at $va = \pi/5$ (Figure 8).

You can open it a bit more with `omre`, for which $va = \pi/4$ (Figure 9). The threshold $va = fa = \pi/6$ (Figure 10) is arrived at by running `thrs`.

The reader must have noticed that Fig. 7 is more refined than the subsequent ones. We did this on purpose because the scripts `olid`, `omre` and `thrs` aim at arriving at the threshold. Therefore, refining the picture for each of these steps would increase the computational cost. Moreover, it is possible to enhance and create an animation with them by means of a viewing program, for instance `Geomview`. The reader can access the animation file `sdrop.mp4` at our homepage.

Finally, according to Theorems 1 and 2 we can determine when the ring type spanner drop will cease to exist. Figure 11 shows the case $\alpha = va < fa + 0.5 = \gamma_1 + \gamma_2 - \pi$.
There the shaded quadrilateral is concave and the sum of the three internal angles $\pi - \gamma_1$, $\pi - \gamma_2$ and $\alpha$ is less than $\pi$.

As $\alpha$ approaches $\alpha + 0.5$ the concave quadrilateral converges to an infinitesimal triangle for which we finally have $\alpha = g_1 + g_2 - \pi$.

By increasing $\alpha$ further on the ring pops and assumes the shape of a half-moon. This can be seen by running the script `brst` of our simulator (Figure 12).

![Figure 12. Ring is replaced by half-moon.](image)

Finally, the scripts `aflt` and `flat` are two steps that show what happens to the half-moon when we open $\alpha$ until the extreme value $\pi$. This can be seen in the video `sdrop.mp4` linked from our homepage: [https://sites.google.com/site/chrondon1992](https://sites.google.com/site/chrondon1992).

We end up with a spherical cap that makes an internal angle $\gamma_2$ with the horizontal plane. The upper arc of the half-moon collapses to a single point on the line that separates $\Pi_1$ and $\Pi_2$.

4 An example can be seen in the video `Soap Film Loops.mp4` on our homepage [https://sites.google.com/site/chrondon1992](https://sites.google.com/site/chrondon1992), taken from the publicly available *Harvard Natural Sciences Lecture Demonstrations*. 