Prediction of positive and negative elastic dilatancy in 2D and 3D liquid foams

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HAL Id: hal-00419999

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Submitted on 26 Sep 2009

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A tentative geometrical description of static dilatancy in liquid foams: ordered 2D and 3D foams

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(Dated: September 25, 2009)
Abstract

Liquid foams have been observed to behave like immersed granular materials in at least one re-
spect: deformation tends to raise their liquid contents, a phenomenon called dilatancy. We present
a geometrical interpretation thereof in foams squeezed between two solid plates (2D GG foams),
which contain pseudo Plateau borders along the plates, and in 3D foams. While experimental ob-
servations evidenced the effect of a continuous deformation rate (dynamic dilatancy), the present
argument applies primarily to elastic deformation (static dilatancy). We show that the negative
dilatancy predicted by Weaire and Hutzler (Phil. Mag. 83 (2003) 2747) at very low liquid fractions
is specific to ideal 2D foams and should not be observed in the dry limit of real 2D foams.

PACS numbers: 83.80.Iz Emulsions and foams – 83.60.Hc Normal stress differences and their effects –
45.90.+t Other topics in classical mechanics of discrete systems

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I. DILATANCY IN FOAMS AND GRAINS

Liquid foams \([1]\) and granular materials are common examples of materials whose mechanical behaviours exhibit complex features, such as non-homogeneous flows: shear-banding, fracture or jamming. In the present work, we are interested in yet another such complex behaviour, called “dilatancy”. Dilatancy was described by Bagnolds \([2]\) in the context of granular materials: upon deformation, because grains are forced to move while avoiding each other, the medium swells to some extent. In other words, the fluid volume fraction \(\phi\) is increased. This effect can remain unnoticed in air. By contrast, a spectacular absorption of liquid \([2]\) is obtained upon deformation of an immersed granular sample.

In liquid foams, it is not \textit{a priori} so obvious whether dilatancy should be expected or not. Because bubbles are deformable individually, a foam should be able to deform substantially without altering its liquid fraction.

In 3D foams, \textit{dynamic} dilatancy was indeed observed under stationary shear \([3]\): in a device where the shear rate localizes, the liquid volume fraction was observed, both visually and through electrical conductimetry, to stabilize, within seconds, at a higher value in the region being sheared continuously, while it remained lower in both statically deformed regions (and when shearing was stopped, the liquid fraction was observed to become uniform again within seconds). Dynamic dilatancy, whose microscopic origin is yet uncertain \([18]\), thus constitutes one out of many mechanisms that could favour shear banding, a family of rheological behaviours commonly observed in many other complex fluids such as giant micelle solutions, multilamellar vesicles or onions, granular pastes, etc. The stabilizing mechanism is as follows. As both the foam shear modulus and its yield strain are known to decrease as the liquid fraction increases \([1]\), a region that contains more liquid will host statistically more plastic events than other regions and, if dynamic dilatancy is indeed present, even more liquid will then permeate towards that region from less active regions. This process reaches a stationary regime (co-existing shear bands) when the dilatancy-induced pressure gradient is balanced by the osmotic gradient associated with fluid concentration gradients.

Whether \textit{static} dilatancy (resulting from a \textit{fixed deformation}) also exists in liquid foams has not been determined experimentally so far. It might constitute a destabilizing factor for a homogeneous flow and \textit{generate} shear banding in the first place.

A \textit{theory} of static dilatancy was derived from thermodynamical considerations a few
FIG. 1: Four types of 2D foams. (a) Top-view of an ideal, truly two-dimensional foam: either the third dimension is purely absent, or the foam is invariant with respect to the third dimension. (b) Side-view of a monolayer of bubbles between two glass plates (“GG” foam). (c) Monolayer of bubbles floating on a liquid bath below a glass plate (“LG” foam). (d) Monolayer of bubbles floating on the free surface of the liquid bath, in contact with air (“LA” foam). Dilatancy is no real issue for LG or LA foams because of the underlying reservoir of liquid. By contrast, the GG foam holds liquid not only in the Plateau borders just like an ideal 2D foam (second term of Eq. (1)), but also in the pseudo Plateau borders which run along the edge between two bubbles and touch one solid plate each (first term of Eq. (1)).

years ago [4, 5] for an ideal, truly two-dimensional foam (Fig. 1a), with predictions based on empirical or numerical results concerning the osmotic pressure and the elastic modulus. The model leads to the surprising prediction that the sign of the dilatancy effect should be negative for very low liquid fractions, as a consequence of the fact that the elastic modulus does not depend on the liquid fraction in the dry limit (Decoration Theorem [1, 6]).

In this paper, we provide a simple geometrical description of static dilatancy in a dry liquid foam. An elastic deformation of such a material induces an elongation of the bubbles while conserving their volume constant, thereby increasing the total Plateau border length per bubble, or per unit volume. Since most of the liquid is located in Plateau borders, the liquid fraction is thus increased.

According to this description, we calculate the magnitude of the dilatancy effect analytically for a crystalline, hexagonal GG foam (Fig. 1b) as well as for a 3D Kelvin foam, and we discuss negative dilatancy [4, 5].
II. STATIC DILATANCY IN A DRY 2D GLASS-GLASS FOAM

Here, we introduce a very simple geometrical description of static dilatancy. In a “GG” foam, there are two contributions to the liquid content, which are presented on Fig. 1:

\[ \Omega_{\text{liq}} \simeq \frac{4-\pi}{2} PR_{\text{ps}}^2 + (2\sqrt{3}-\pi) R^2 H \] (1)

where \( \Omega_{\text{liq}} \) is the volume of liquid per bubble. The first term corresponds to the pseudo Plateau borders (with the corresponding radius of curvature \( R_{\text{ps}} \) and the bubble perimeter \( P \)) while the second term corresponds to the genuine Plateau borders (radius \( R \) and gap \( H \) between the solid plates).

In the present work, we will assume both \( P \gg R \) (i.e., \( \phi \ll 1 \)) and \( R_{\text{ps}} = R \ll H \) (floor tile regime). In other words, we concentrate on regimes E, F and G of Fig. 4. Regime E corresponds to a floor tile GG-foam, while regime G is the truly 2D limit of refs. [4, 5]. We here focus on floor tile “GG” foams (regime E of Fig. 4), i.e., with not only a low liquid volume fraction (\( \phi \ll 1 \), achieved when \( R \ll P \)) but also small pseudo Plateau borders compared to the gap between the solid plates (\( R_{\text{ps}} \ll H \)), in which case one has \( R_{\text{ps}} \simeq R \). The opposite limit of “pancakes” (\( R_{\text{ps}} \simeq H/2 \ll R \), regime ABCD of Fig. 4) will be discussed elsewhere [7].

Let us consider such a 2D glass-glass foam free of any significant anisotropic (i.e., deviatoric) in-plane stress. If the foam is now deformed elastically, the average bubble perimeter \( P \) increases as the typical bubble elongates while conserving its volume constant. Since the total amount of liquid per bubble remains constant on short time scales, the increase in perimeter causes the pseudo Plateau borders to shrink accordingly (see Fig. 2). Later on, as permeation takes place, the pseudo Plateau border radius \( R_{\text{ps}} \) may return to its original value, and the part of the liquid that is located in the pseudo Plateau borders increases accordingly.

Thus, soon after deformation, the pressure difference between the gas and the liquid must increase as the pseudo Plateau borders (and Plateau borders) shrink. Later, after permeation has taken place, the liquid content of the foam has increased as compared to its value when the foam was at rest. In other words, there is a positive dilatancy effect as soon as the perimeter increases.

In order to evaluate the change in bubble perimeter, let us consider a dry, crystalline foam subjected to an arbitrary elastic, homogeneous deformation. Up to a global rotation,
FIG. 2: Pseudo Plateau borders between two bubbles squeezed between two glass plates. Original configuration, with top-view (a1) and perspective (a2). When a bubble which is initially at rest is suddenly deformed, its perimeter, \( i.e., \) the total length of its pseudo Plateau borders, is increased. On short time scales, the amount of liquid that surrounds the bubble remains constant, hence the quantity of liquid per unit length of the perimeter decreases (b1). Thus, the pseudo Plateau borders shrink, and the corresponding radius of curvature \( R_{ps} \) is reduced (b2). This implies that the pressure in the liquid is lowered. As a result, liquid from less deformed regions of the foam, where the liquid pressure is not as low, can soon permeate and compensate for the Plateau border shrinking (c).

This deformation is equivalent to an elongation by some factor \( \lambda \) in one direction and to a compression by the same factor in the perpendicular direction. In such a hexagonal foam, in the dry limit, the bubble perimeter increases in the following way:

\[
\frac{P^{2D}(\lambda)}{P^{2D}_0} = \left[ \frac{\lambda}{2} + \frac{1}{2\lambda} \right] \quad \frac{\delta P}{P^{2D}_0} = \frac{(\lambda - 1)^2}{2\lambda}
\]  

(2)

The perimeter thus increases when the foam is deformed \( (\lambda \neq 1) \), as can be seen on Fig. 3. This law, which is exact in the dry limit, holds as long as no T1 process occurs and does not depend on the orientation of the crystalline network with respect to the direction of elongation. In fact, it even holds for polydisperse hexagonal foams [8]. The elongation at which T1 processes occur depends both on orientation or on liquid volume fraction. In the dry limit, it is \( \lambda = \sqrt{3} \) when elongation is perpendicular to some facets and larger for other
elongations. This yield deformation is lowered as the foam becomes wetter as described already long ago [1]. The expected perimeter increase for \( \lambda = \sqrt{3} \) is of order 15\%, which means that gas-liquid pressure difference is expected to increase by 7\% on short time scales, while the foam liquid fraction is expected to increase by 15\% after permeation.

III. DILATANCY IN THREE DIMENSIONS

In a dry, three-dimensional foam, most of the liquid is located in the Plateau borders. Following the same line of thought as above, static dilatancy is expected if deformation induces an increase in the total Plateau border length per bubble (or per unit volume of foam).

To test this in a simple manner, we took an approximate version of Kelvin’s bcc bubble packing. Kelvin’s packing is one of the most commonly encountered in monodisperse 3D foams, even though it does not have the lowest energy [9]. In a real Kelvin foam, square faces have edges bending slightly outwards, while hexagonal faces are slightly non-planar. Here, each bubble is assumed to be a truncated octahedron, called tetrakaidecahedron, with six planar, square faces and eight regular hexagonal, planar faces. This approximation was used and discussed by Reinelt and Kraynik [10] to express the stress and the energy under large elastic deformations. Here, we present the changes in the total Plateau border perimeter.

Computing the bubble perimeter (sum of all edge lengths) as a function of elongation has been performed for three orientations of the elongation (with a compression by \( 1/\sqrt{\lambda} \) in the perpendicular directions. As can be seen on Fig. 3, the result is very similar to the two-dimensional case, and indicates that significant dilatancy is to be expected in dry 3D foams, too. Incidentally, Fig. 3 indicates that for the present approximation of a Kelvin foam, the response is independent of the direction of elongation:

\[
\frac{P_{3D}^A(\lambda)}{P_{3D}^0} = \frac{P_{3D}^B(\lambda)}{P_{3D}^0} = \frac{P_{3D}^C(\lambda)}{P_{3D}^0} = \frac{1}{3} \left[ \lambda + \frac{2}{\sqrt{\lambda}} \right]
\]

IV. OSMOTIC PRESSURE AND DILATANCY

The dilatancy coefficient is defined [4] as the (second) derivative of the osmotic pressure with respect to the deformation \( \epsilon \): \( \chi = \partial^2 \pi_{osm} / \partial \epsilon^2 \). With the variations of the average
FIG. 3: Sum of all Plateau border lengths in a 2D (hexagonal, Eq. (4)) foam or 3D (Kelvin, Eq. (3)) foam, as a function of the elongation $\lambda$. In 3D case, elongation is performed in direction A (normal to a square facette), B (normal to a hexagonal facette) or C (one of the edges of Kelvin’s cell) with compression $1/\sqrt{\lambda}$ in both perpendicular directions.

bubble perimeter in mind, let us now derive the variation of the osmotic pressure in a floor tile GG-foam. From Eq. (2), expressed in terms of $\lambda = 1 + \epsilon$, one has $\delta P/P \simeq \epsilon^2/2$. Hence, $\chi = \delta \pi_{\text{osm}}/(\delta P/P)$, where the variation $\delta \pi_{\text{osm}}$ is taken at constant liquid volume fraction $\phi$.

When the foam is confined in a container, the osmotic pressure $\pi_{\text{osm}}$ corresponds to the force that must be applied externally to one of the confining walls if the latter is permeable to the liquid but not to the bubbles. The osmotic pressure (which is in fact a symmetric tensor and not just a scalar quantity [4, 5]) is thus the difference between the stress in the foam and the pressure applied by the pure liquid on the other side of the semi-permeable wall: $\pi_{\text{osm}} = -p_l - \sigma$ (where tensile stresses and pressures are both counted positively). The stress in the foam includes a pressure contribution from the liquid ($p_l$) and from the gas ($p_g$), as well as a tensile contribution from the interfaces: $\sigma = -\phi p_l I - (1 - \phi) p_g I + \sigma^{\text{interf}}$. In the floor tile limit ($R \ll H$), the interface stress, averaged over in-plane orientations and over the sample thickness, is related to the bubble perimeter [4]: $< \sigma^{\text{interf}} >_{2D} \simeq (1 - \phi) \left[ \frac{\gamma PH}{2H} \right] + \frac{2\gamma}{\Omega}$, where $\gamma$ is the surface tension and $\Omega$ is the bubble volume. Hence, osmotic pressure (now expressed as a scalar quantity) can be written as: $< \pi_{\text{osm}} >_{2D} = (1 - \phi) \left[ \Delta p - \frac{\gamma PH}{2H} \right] - \frac{2\gamma}{\Omega}$, where $\Delta p = p_g - p_l = \frac{\gamma}{R_{ps}} = \frac{\gamma}{R}$ is the pressure difference between the gas and the liquid.
FIG. 4: Regimes of a GG-foam with low liquid fraction ($\phi \ll 1$), in terms of the bubble perimeter $P$ (measured on top view), the Plateau border radius $R$ (as seen from above too) and the cell height $H$. Real GG-foams ($P > H$) can be found in two main configurations: the “pancake” regime $\text{ABCD}$ ($H - 2R_{ps} \ll H \ll R$, region ABCD) and the “floor tile” regime E ($R = R_{ps} \ll H$). Regimes F and G do not correspond to real foams (which cannot sustain the condition $H \gg P$ without destabilizing into 3D foams), but regime G is the limit of truly two-dimensional foams ($H \to \infty$) addressed in earlier works on dilatancy [4, 5].

The dilatancy then results from Eqs. (1) and (2):

$$\chi = \frac{\delta < \pi_{\text{o sm}} >_{2D}}{6P/P} \approx \frac{\gamma/(2R)}{1 + \frac{\sqrt{3-\frac{\pi}{2}}}{1-\frac{\pi}{4}} H/P} - \frac{\gamma P}{2A}$$

(4)

The last term, in which $A = \Omega/H$, originates in the interface stress contribution $\sigma_{\text{interf}}$ and is equal to one half of the interfacial energy per unit height ($\gamma P$ per bubble).

V. NEGATIVE DILATANCY?

For a GG-foam (regime E with $P \gg H$, see Fig. [4]), one has $\chi \simeq \Delta p/2 > 0$. By contrast, when $H \gg P$ (regime G), one gets the limit of a truly two-dimensional foam since the pseudo Plateau borders (total length $P$ per bubble) are too small compared to the Plateau borders (length $H$) to contribute to dilatancy significantly. Hence only the last term remains in Eq (4), and dilatancy is negative: $\chi \simeq -\frac{\gamma P}{2A}$. We thus recover exactly the result $\chi \simeq -2G$. 

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by Weaire and Hutzler \[4\] in the dry limit \[19\]. For instance, in the case of a crystalline, hexagonal 2D foam: \(\chi \simeq -4\sqrt{3}\gamma/P\).

**VI. DISCUSSION**

In the present work, we have shown that positive static dilatancy in liquid foams might result from the overall increase in Plateau border lengths when the foam is being deformed elastically. In 2D GG foams, this would be in fact the pseudo Plateau borders (those along the solid walls). The present study was restricted to crystalline foams, whether 2D GG hexagonal foams or 3D Kelvin foams. A weaker (or even reversed) effect cannot be excluded, at this stage, in the case of disordered or polydisperse foams, or even simply for different 3D crystalline structures. Indeed, in 3D at least, there seems to be no trustworthy reason why the minimum in the total facet surface area (which is the definition of a foam at rest, in the dry limit) should coincide with the minimum in the sum of all Plateau border lengths. For a GG foam in a somewhat less dry configuration (regimes A-D of Fig. 4), the expected dilatancy effect can be shown to be less intense \[7\].

The negative dilatancy predicted \[4, 5\] in the dry limit in truly 2D foams (invariant along the third dimension of space) coincides with one particular limit of our study, which indeed corresponds to a truly 2D foam. It is then related to the interfacial (and deviatoric) part of the stress. By contrast, in the other regimes, we obtain a positive dilatancy, related to the pressure in both fluid phases. Let us mention that negative dilatancy is also possibly expected in the wet limit of the pancake regime \[7\].

If static dilatancy is confirmed experimentally, it should be coupled with drainage models (which describe permeation) and incorporated into rheological models of foams \[11, 12, 13, 14, 15\].

**Acknowledgements**

We gratefully acknowledge fruitful discussions with Benjamin Dollet, with Reinhard Höhler, and with other participants of GDR 2983 Mousses (CNRS) and of the Informal Workshop on Foam mechanics (Grenoble, 2008). This work was supported by Agence Na-
tionale de la Recherche (ANR-05-BLAN-0105-01).

[18] It might originate in the slippage of Plateau borders along inter-bubble films, and thus be related to the slippage of pseudo Plateau Borders along solid walls. Experiments have been carried out in this direction recently [18, 17].
[19] The shear modulus $G$ can be computed as follows. The change in energy for a shear strain $\Gamma$ is $\delta \mathcal{E} = \frac{1}{2} G \Gamma^2$ and is also related to the change in the bubble perimeter: $\delta \mathcal{E} = \gamma \delta P/A$. Now, in the case of a pure shear $\Gamma$, at least for a hexagonal, crystalline foam, one can show that Eq. (3) yields $\delta P/P = \Gamma^2/8$. Hence, $G = \frac{1}{4} \gamma P/A$ and $\chi \simeq -\frac{\gamma P}{2A} = -2G$. 

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