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P. Marmottant, C. Raufaste and F. Graner



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P. Marmottant^a, C. Raufaste^b, and F. Graner

Laboratoire de Spectrométrie Physique, UMR5588, CNRS-Université Grenoble I, B.P. 87, F-38402 St Martin d'Hères Cedex, France

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Abstract. The plastic flow of a foam results from bubble rearrangements. We study their occurrence in experiments where a foam is forced to flow in 2D: around an obstacle; through a narrow hole; or sheared between rotating disks. We describe their orientation and frequency using a topological matrix defined in the companion paper (F. Graner, B. Dollet, C. Raufaste, and P. Marmottant, this issue, **25** (2008) DOI 10.1140/epje/i2007-10298-8), which links them with continuous plasticity at large scale. We then suggest a phenomenological equation to predict the plastic strain rate: its orientation is determined from the foam's local elastic strain; and its rate is determined from the foam's local elongation rate. We obtain a good agreement with statistical measurements. This enables us to describe the foam as a continuous medium with fluid, elastic and plastic properties. We derive its constitutive equation, then test several of its terms and predictions.

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1 Introduction

A liquid foam, made of gas bubbles surrounded by liquid walls (Fig. 1), is elastic for small strain, plastic for large strain, and flows at large strain rates [1–3]. This complex mechanical behaviour is exploited in numerous applications, such as ore separation, drilling and extraction of oil, food or cosmetic industry [1]. It is not yet fully understood [3]. Existing models of complex fluids include Oldroyd's visco-elasticity or Bingham's visco-plasticity [4,5]. Complete models tend to unify elastic, plastic and fluid behaviour [6–12].

Most models describing the foam as a continuous material have to make an assumption to describe phenomenologically the elastic-to-plastic transition, and the plastic strain rate. Here, we want to link it directly with the observation of foam's individual components, the bubbles. In fact, bubbles play for foams the same role as microscopical components play for other complex fluids, but have the advantage of being easily observable. The individual plastic events [13] in foams are the topological rearrangements (neighbour swapping, also called "T1 processes" [1]). Predicting the occurrence and properties of T1's, and characterizing their impact at large scale, is thus a crucial step to describe the foam as a continuous material.

Section 2 lists the ingredients we use. We start from a scalar model of the foam plasticity and dynamics, which predicts correctly its rheological behaviour [9]. We then include robust statistical tools defined in the companion paper [14], which provides: i) a link between local (bubble) behaviour and global (foam) measurements of elastic, plastic and fluid behaviours; ii) matrix (rather than scalar) measurements of these quantities, that is, including direction and magnitude of anisotropy; iii) space dependence of these measurements (heterogeneous deformation instead of homogeneous).

Section 3 then presents a kinematic equation to describe the elastic-to-plastic transition, which is the main point of this paper. It is an analytical prediction of the plastic strain rate, based on the elastic strain and the total strain rate.

Section 4 tests its predictions on experiments in different foam flows. The relationship between experimental data agrees with our analytical prediction. Moreover, we correctly predict the frequency and spatial distribution of T1's, which differs much from the spatial distribution of both elastic strain and total strain rate.

Section 5 concludes by discussing these findings and possible applications to other systems. It proposes a dynamic constitutive equation for foam rheology, relating

^a e-mail: philippe.marmottant@ujf-grenoble.fr

^b Present address: Physics of Geological Processes, University of Oslo, Sem Selands vei 24 NO-0316 Oslo, Norway.



Fig. 1. Experimental 2D flows of a foam: top view, here flow from left to right. (a) Wet foam (between glass and water) flowing around an obstacle, picture from B. Dollet [15,16]; field of view 15 cm×10 cm. (b) Dry foam (between parallel glass plates) flowing around an obstacle, picture from C. Raufaste [14]; field of view 13.3 cm × 10 cm. (c) Moderately dry foam (between parallel plexiglass plates) flowing through a narrow hole, picture from M. Asipauskas [17]; field of view 7.5 cm × 10 cm. (d) Wet foam (between parallel glass plates) sheared between two concentric wheels with tooth-shaped boundaries to prevent slipping; the rotating inner wheel is visible at the bottom, the fixed outer wheel is visible at the top [18]. Arrows indicate the measured velocity field; field of view 12.5 cm × 11.5 cm. Liquid fractions are approximately estimated as: (a) $4 \cdot 10^{-2}$ [19], (b) $2 \cdot 10^{-4}$ [20], (c) 10^{-2} [17], (d) $5 \cdot 10^{-2}$ [18].

stresses to applied strains, with the introduction of a viscosity. This closes the complete set of equations describing a foam, leading to testable predictions.

2 Ingredients of model and tests

2.1 Continuous description of plasticity with scalar strains

This section summarizes the model presented in [9] for homogeneous strains, described by a simple scalar. This situation arises when considering the strain in a rheometer, where strain is induced by the displacement of a plate with respect to the other, and characterized by a scalar.

The total applied strain rate (or symmetrised velocity gradient) contributes in part to load the elastic strain (bubbles deform) and to the plastic strain rate (bubbles move relatively to each other). The total applied strain rate is the sum of an internal elastic strain rate and the irreversible plastic strain rate:

$$\dot{\varepsilon}_{\rm tot} = \frac{\mathrm{d}\varepsilon_{\rm el}}{\mathrm{d}t} + \dot{\varepsilon}_{\rm pl},\tag{1}$$

where we used a different notation for the time derivative of $\varepsilon_{\rm el}$, to emphasise that it is an internal state variable of the material.

A kinematic equation describes how the total strain rate is shared between change of elastic strain and plastic strain. For slowly sheared foams plasticity occurs if $\varepsilon_{\rm el}$ and $\dot{\varepsilon}_{\rm tot}$ have the same sign (loading of the material), and is zero otherwise (unloading) [9]:

if
$$\varepsilon_{\rm el}\dot{\varepsilon}_{\rm tot} > 0$$
, $\dot{\varepsilon}_{\rm pl} = h(\varepsilon_{\rm el})\dot{\varepsilon}_{\rm tot}$,
if $\varepsilon_{\rm el}\dot{\varepsilon}_{\rm tot} \le 0$, $\dot{\varepsilon}_{\rm pl} = 0$. (2)

The plasticity fraction, or *yield function*, h, is zero at zero strain (elasticity only, no plasticity), and reaches 1 (plasticity only, no elasticity) at the yield strain, which is a material-dependent parameter. A smooth onset of plasticity can be described by a plasticity function h which continuously interpolates between the value 0 (completely elastic) and 1 (completely plastic) [9].

Equation (2) closes the system of kinematic equations describing a foam. Indeed from equations (1, 2), we can write

$$\mathrm{d}\varepsilon_{\mathrm{el}}/\mathrm{d}t = \dot{\varepsilon}_{\mathrm{tot}} - h(\varepsilon_{\mathrm{el}})\dot{\varepsilon},\tag{3}$$

which can be integrated, knowing the applied strain rate, to calculate the internal elastic strain.

The scalar constitutive equation [9] proposed a total stress that is the sum of the elastic and viscous stress. It was simple to assume a linear relation

$$\sigma = G \varepsilon_{\rm el} + \eta \dot{\varepsilon},\tag{4}$$

with G an elastic modulus and with η a viscosity.

This model is thus based on three material-dependent parameters; an elastic coefficient, a viscous one, and a plastic yield strain. It successfully predicted foam dynamical properties. In oscillatory regime, it predicts storage and loss moduli G', G''. In stationary regime, it resembles a Bingham model. It also enables to predict transient behaviours and creep [9]. If needed, it is possible in principle to refine it, by introducing some non-linearities in



Fig. 2. Sequence of images of a T1 event in a measurement box: (a) shows the network of links ℓ (thin lines) joining neighbour bubble centers superimposed on snapshots extracted from a dry foam [14], and the disappearing link (ℓ_d , dashes), (b) the intermediate stage, and (c) the appearing link (ℓ_a , thick line).

equation (4); for instance by replacing $\dot{\varepsilon}$ with a power law, $\dot{\varepsilon}^n$ (in which case, in a stationary regime it looks like a Herschel-Buckley model); or by having G or η not constant.

2.2 Direct measurement of strain, rearrangements and flow

This section summarizes the parts of the companion paper [14] which are used in what follows. The matrices $\dot{\boldsymbol{\varepsilon}}_{tot}$, $\boldsymbol{\varepsilon}_{el}$, and $\dot{\boldsymbol{\varepsilon}}_{pl}$ generalise the total, elastic and plastic strains in equation (1), and are function of the position in the material. They are measured directly using the statistical strains based on the bubble network: **U**, **V** and **P**, respectively. They are all obtained from the texture tensor $\mathbf{M} \equiv \langle \boldsymbol{\ell} \otimes \boldsymbol{\ell} \rangle$, based on links $\boldsymbol{\ell}$ between each pair of neighbouring bubble centers (see Fig. 2). Coarse-graining procedures yield a continuous description of the foam.

The internal strain is defined from the relative deformation of bubbles with respect to the deformation at rest M_0 , as

$$\mathbf{U} \equiv \frac{1}{2} \left(\log \mathbf{M} - \log \mathbf{M}_0 \right). \tag{5}$$

The topological rearrangement rate is defined from the discontinuous deformation of the network after each rearrangements. It describes the time derivative of U due to T1 events

$$\mathbf{P} \equiv f \, \frac{1}{2} \left(\langle \boldsymbol{\ell}_d \otimes \boldsymbol{\ell}_d \rangle - \langle \boldsymbol{\ell}_a \otimes \boldsymbol{\ell}_a \rangle \right) \mathbf{M}^{-1} + sym$$
$$= f \delta \mathbf{U}_{\mathrm{T1}}, \tag{6}$$

f being the frequency of rearrangements in an observation box containing one link, and $\delta \mathbf{U}_{\text{T1}}$ is the sudden drop in internal strain associated with a T1. It is computed from the disappearing links ℓ_d and appearing links ℓ_a (see Fig. 2).

The rate of strain is defined from the *continuous* deformation of the network. It describes the time derivative of \mathbf{U} between T1 events as

$$\mathbf{V} \equiv \frac{\mathbf{W} + \mathbf{W}^t}{2}$$

where

$$\mathbf{W} = \mathbf{M}^{-1} \left\langle \boldsymbol{\ell} \otimes \frac{\mathrm{d}\boldsymbol{\ell}}{\mathrm{d}t} \right\rangle.$$
 (7)

2.3 Experimental data

Horizontal monolayers of bubbles (Fig. 1) offer a unique possibility to observe all bubbles: their deformations, rearrangements and velocity. They are quasi-2D foams, because they are not strictly 2D. However, their flow is truly 2D: it has no vertical component. A large set of detailed data is available; bubbles act as convenient tracers of elastic strain, rearrangements and velocity [14,17,21].

We reanalyse experimental data already published and courteously provided to us by the authors. For details of the materials and methods, see the original publications. In these four set-ups, coalescence and ageing are below detection level. We assume that the gas and liquid constituents of the foam move together, at the same velocity (no drainage).

In Figure 1a, the foam is confined between the surface of water and a horizontal plate of glass. Bubbles are rather round, due to the high effective liquid fraction [19]. Thus the region where T1's occur is larger, more widely distributed around the obstacle (compare Figs. 12 and 14 below). There are thus more regions of the flow where statistics are significant. This is why we use this experiment for the most detailed quantitative tests (Sect. 4.1). In Figures 1b-d, the foam is confined between two horizontal plates of glass, and drier liquid fractions can be obtained.

In Figures 1a-c, the channel (only partly visible) is horizontal, its length is 1 m, its width 10 cm, its thickness 3.5, 3 and 0.5 mm, respectively. It is filled with bubbles obtained by steadily blowing nitrogen in water with commercial dishwashing liquid. The bubbles are monodisperse and form a disordered monolayer which reaches the free exit at the end of the channel. The resulting steady plug flow, well in the quasistatic regime [15], is made hetereogeneous by inserting a 3 cm diameter obstacle (Fig. 1a,b) or a constriction [17] (Fig. 1c). Thus different regions simultaneously display different velocity gradients, elastic strains, and rearrangement rates, and allow to sample simultaneously many different conditions.

In Figure 1d the foam is in a 2D circular Couette geometry [18]. Briefly, the 2 mm thick foam monolayer is formed between two concentric disks (only partly visible) with semicircular teeth of radius 1.2 mm to match the bubble diameter, thus anchoring bubbles at the walls. The outer disk, of radius 122 mm, is fixed. The inner disk, of radius 71 mm, rotates at 0.25 mm s^{-1} , thus shearing the foam. The resulting velocity field decreases quickly with the distance to the inner disk. This experiment has stirred a debate about the existence and cause of velocity localization: for review, see for instance reference [8]. The experimental measurements we present here complement those of reference [21]; they are largely model-independent and might be used in the future to contribute to this debate.

2.4 Identification with continuous elastic, plastic and total strain rates

This section discusses an additional property of foams, whether 2D or 3D: they are a unique material where the statistical measurements (Sect. 2.2) can be identified with (usual continuous quantities.

2.4.1 Affine flow and total strain rate

As discussed in the companion paper [14], if the flow of a material is affine, then **W** measures the velocity gradient

$$\mathbf{W} \stackrel{\text{affine}}{\simeq} \nabla \mathbf{v}. \tag{8}$$

Figure 3 shows an example of our experimental tests of equation (8). We have measured the detailed components, including the rotational (asymmetric part), as well as eigenvalues and axes of the symmetric part. All these quantities are the same for \mathbf{W} and $\nabla \mathbf{v}$, within a few percents precision, with a correlation close to 1. The measurements of \mathbf{W} and $\nabla \mathbf{v}$ have a comparable precision, and suffer from similar imprecisions near the channel walls and obstacle. Both have a small trace (ellipses are nearly circular). Tests at smaller and larger scales, that is with different sizes of the representative volume element (RVE), yielded similar results (data not shown).

This agreement is unexpectedly good, given that with the dry foam chosen here the strain is large and its gradient is strong. At large scale, movements of individual objects within the same RVE can differ considerably; but even in this unfavorable case the affine assumption seems to hold, as shown in Figure 3. The reason seems to be that equation (8) is correct whenever **M** does not vary significantly within the chosen RVE [20]. Together, equations (7, 8) enable us to identify the rate of growth of links, **V**, with the total strain rate, $\dot{\boldsymbol{\varepsilon}}_{\text{tot}} = (\boldsymbol{\nabla} \mathbf{v} + \boldsymbol{\nabla} \mathbf{v}^t)/2$.

2.4.2 Elastic stress and strain

In a foam, the elastic energy is proportional to the bubble surfaces, so that the elastic strain directly stems from bubble deformation. It has been experimentally checked that \mathbf{U} (or at least its deviatoric part) actually determines the (deviatoric) elastic stress: see references [17,21], to which we refer for details of the measurement method. We check it here too (Fig. 4), using our driest example (Fig. 1b) in order to improve the measurement precision of the deviatoric elastic stress.

This indicates that the internal strain **U**, which is a state variable constructed from bubble deformations, measures well the reversible strain $\varepsilon_{\rm el}$ that gives rise to elastic stresses, although the flowing foam is clearly out of the elastic regime. In what follows, we thus call **U** the "elastic strain".

Up to a prefactor, namely the foam's shear modulus, $(U_d)^2$ represents the elastic energy stored (*e.g.* due to shear): that is, the difference between the energy of the current state and that of the local minimum. The latter varies at each T1.

2.4.3 Plastic strain rate

Disordered foams, which we consider here, are models for the plasticity of amorphous materials. Plastic events take



Fig. 3. Affine assumption: test of equation (8) on the foam of Figure 1b. (a) Comparison of maps of W (top) and $\nabla \mathbf{v}$ (bottom): ellipses, symmetric part; grey levels, antisymmetric part (bar: $5.6 \times 10^{-3} \text{ s}^{-1}$). (b) Quantitative comparison: top, $(W_{XX} - W_{YY})/2$ versus $(\nabla v_{XX} - \nabla v_{YY})/2$, and the same for all other components (superimposed): XX + YY, XY - YX, XY + YX; bottom, angle (in degrees) of ellipses plotted in (a). Each point comes from one RVE of the image.



Fig. 4. XY components of the elastic stress σ (in arbitrary units) versus XY components of U. Each point comes from one RVE of the image of Figure 1b. The slope of the cloud of points is the foam's 2D shear modulus; it is of order of the bubbles' line tension λ to diameter ratio. Data for the (XX - YY)/2components are superimposed: they have the same slope.



Fig. 5. Schematic impact of individual (discrete) rearrangements on the stored elastic strain U, for a constant loading rate $\dot{\varepsilon}$. Rearrangements relax exponentially the strain over a time τ_{relax} , with here $\dot{\varepsilon}\tau_{\text{relax}} = 0.02 \ll 1$. In the present continuous model, rearrangements are coarse-grained. Reprinted from reference [9].

the form of bubble rearrangements. When averaged over time or space, the effect of topological rearrangements is usually smooth, and the foam behaves as a continuous material (Fig. 5). This contrasts with ordered foams, where bubbles are arranged in a honeycomb lattice and topological rearrangements are highly correlated, which are models for the plasticity of crystals based on dislocation movement [23–25].

Each topological rearrangements is instantaneous (by definition of a topological change) and imply a change from a stable elastic branch to another. It is followed by a relaxation over a finite time τ_{relax} , determined by the ratio of dissipation to driving elastic force. The rate of T1's is determined by the shear rate, which we keep here slower than τ_{relax}^{-1} in what follows (for extension to higher shear rates, see [10]). In that case, the foam has time to relax towards a new local equilibrium state (Fig. 6), and the total energy dissipated is determined by the difference between the energy barrier and the new local energy minimum. Thus the foam stays in this configuration: the



Fig. 6. Shear of a 2D ordered foam [22], different liquid fraction: 9.3% (close-packing, lowest amplitude curve), 5.5% (onset of a negative slope unstable branch), 3% and 1%. Decreasing liquid leads to increasing maximum stress. The topological rearrangement occurs when the stress *versus* strain curve becomes unstable (dot-dashed line) and induces an irreversible strain. Three 1% foams states are represented (circles on curve). The stress σ is here normalised using surface tension γ and bubble radius *d*.

transformation is plastic. It is possible to come back, with hysteresis, to the state which preceded the T1: since it too is a local energy minimum, the succession of a T1 and its inverse T1 leads to exactly the same pattern [26,27].

Note that each relaxation following the T1 involves an *irreversible* dissipation, which measures the plastic dissipation rate (as seen in Fig. 6). Indeed, if the strain rather than the stress is imposed, the stress-strain curve of a 2D ordered foam with less that 5.5% of liquid has an unstable branch. The dissipation power of the plastic flow is proportional to the rate of T1's, thus to the shear rate: the stress is thus independent of the shear rate, which is characteristic of solid friction (plastic contribution to stress, see plateau of Fig. 5).

To summarize, a topological rearrangement is equivalent to a plastic strain. This is specific to dry foams: as discussed in the companion paper [14], this equivalence is not general to all materials. The irreversibility is associated with the relaxation after a rearrangement.

When a rearrangement occurs, the total strain is not changed locally. The elastic strain decreases by $\delta \varepsilon_{\rm el} = -\delta \mathbf{U}_{\rm T1}$, and the plastic strain thus increases by $\delta \varepsilon_{\rm pl} = \delta \mathbf{U}_{\rm T1}$. Hence, the matrix **P** measures well the plastic strain rate $\dot{\varepsilon}_{\rm pl} = \delta \varepsilon_{\rm pl} / \delta t$.

2.4.4 Complete identification

The preceding sections suggest that, in dry foams, it should be possible to achieve the complete identification between statistical measurements and continuous quantities:

$$\begin{array}{lll}
\mathbf{U} & \stackrel{\text{foams}}{\simeq} & \boldsymbol{\varepsilon}_{\text{el}}, \\
\mathbf{V} & \simeq & \dot{\boldsymbol{\varepsilon}}_{\text{tot}}, \\
\mathbf{P} & \simeq & \dot{\boldsymbol{\varepsilon}}_{\text{pl}}.
\end{array}$$
(9)

The statistical measurements are linked by

$$\mathbf{V} = \frac{\mathcal{D}\mathbf{U}}{\mathcal{D}t} + \mathbf{P},\tag{10}$$

with \mathcal{D} the total corotational derivative [14]: $\mathcal{D}\mathbf{U}/\mathcal{D}t = \partial \mathbf{U}/\partial t + (\mathbf{v}\nabla)\mathbf{U} + \boldsymbol{\Omega}\mathbf{U} - \mathbf{U}\boldsymbol{\Omega}$. Equation (10) is the matrix version of equation (1); here too, the total applied strain rate is the sum of the internal elastic strain rate and the irreversible plastic strain rate.

We conclude that the measurable quantities \mathbf{U} , \mathbf{V} and \mathbf{P} make the connection between the discrete pattern and its continuous mechanical behaviour. In the following, since we present experimental results, we stick to the statistical measurements of strains (l.h.s. of Eqs. (9)). They can of course be replaced with the usual notations from material science (r.h.s. of Eqs. (9)).

3 Model for foam plasticity

This section introduces a prediction of the plastic strain rate and topological rearrangements in foams.

3.1 Plastic strain rate

The deviatoric elastic strain (shear, without dilation) is defined as the traceless matrix

$$\mathbf{U}_d \equiv \mathbf{U} - \frac{1}{2} (\operatorname{Tr} \mathbf{U}) \mathbf{I} \,. \tag{11}$$

Its amplitude is defined as

$$U_d \equiv \left[\frac{\mathbf{U}_d : \mathbf{U}_d}{2}\right]^{1/2},\tag{12}$$

where we use the double contraction product, namely the scalar product of matrices, $\mathbf{A}:\mathbf{B} = \sum_{i,j} A_{ij}B_{ij}$. In a 2D configuration, U_d provides the absolute value of the eigenvalues of matrix \mathbf{U}_d . The matrix \mathbf{U}_d/U_d is then a directional matrix that writes diag(1, -1) in the eigenvector basis of elongation. For details of notations, see [14].

To generalize the scalar model, we need to specify not only the amplitude of the plastic strain rate matrix \mathbf{P} , which we take linear in the strain rate \mathbf{V} ; but also its direction, which we assume is aligned with the current deviatoric elastic strain \mathbf{U}_d . In other words, the plastic evolution is directed along the pre-existent elastic strain and occurs with a rate which is the projection of the total strain rate onto the elastic strain. As will become apparent below (Sect. 4.1), this amounts to a mean-field approximation. Our main assumption is thus that \mathbf{P} is determined by the "projection" of \mathbf{V} on \mathbf{U}_d , defined (in analogy with the projection of a vector on another) using the doublecontraction product:

$$\mathbf{V}_{\text{proj}} = \frac{(\mathbf{V} : \mathbf{U}_d)}{2U_d^2} \,\mathbf{U}_d.$$
 (13)

We thus generalise equation (2) using a scalar plasticity function, h, of the strain amplitude, U_d :

$$\mathbf{P} = h(U_d) \mathbf{V}_{\text{proj}}, \quad \text{if} \quad \mathbf{V} : \mathbf{U}_d \ge 0. \quad (14)$$

Equation (14) applies for the case where the total strain loads the internal strain ($\mathbf{V} : \mathbf{U}_d$ is positive). If \mathbf{V} is proportional to \mathbf{U}_d (same direction, same anisotropy), this equation reduces to $\mathbf{P} = h(U_d)\mathbf{V}$, equivalent to the scalar one, equation (2); moreover, if U_d has reached the yield value (h = 1), then $\mathbf{P} = \mathbf{V}$.

On the opposite, if the applied strain rate \mathbf{V} is in the direction opposed to internal strain ($\mathbf{V} : \mathbf{U}_d$ is negative), it contributes to unload it elastically. It thus does not induce many rearrangements [20]. We neglect them by setting \mathbf{P} to zero

$$\mathbf{P} = 0, \qquad \text{if} \qquad \mathbf{V} : \mathbf{U}_d < 0. \tag{15}$$

Note also that equation (14) would reduce to the classical Prandtl-Reuss model for a perfect plastic material that yields, when the elastic strain reaches the value U_Y [28], if the plasticity function was an Heaviside function, $h(U_d) = \mathcal{H}(U_d - U_Y)$, discontinuously jumping from the value 0 when $U_d < U_Y$ to 1 when $U_d \ge U_Y$.

3.2 Rearrangement frequency

Each rearrangement modifies the strain in its measurement box. We consider here that the strain is concentrated within the reference area attributed to one link $A_{\text{link}} = A/N_{\text{link}}$ (the perturbation in a continuous elastic space rapidly decays with the distance to the rearrangement location, see model [13]). If we assume that each plastic change has a constant amplitude ε_0 , we can write

$$\delta \mathbf{U}_{\mathrm{T1}} = \varepsilon_0 \; \frac{\mathbf{U}_d}{U_d} \,, \tag{16}$$

where we assumed that rearrangements are aligned with elasticity (following Eq. (14)). We have seen that the plasticity rate matrix can be written

$$\mathbf{P} = f \,\,\delta \mathbf{U}_{\mathrm{T1}},\tag{17}$$

where f is the frequency of rearrangements *per link*. When considering averages in larger counting boxes, containing N_{link} links (approximately 3 times the number of bubbles in the box [1]), equation (17) still holds and writes $\mathbf{P} = f_{\text{box}} \delta \mathbf{U}_{\text{box}}$. Indeed the frequency in the counting box is $f_{\text{box}} = N_{\text{link}}f$, and the impact of a T1 on a larger surface is diluted to the value $\delta \mathbf{U}_{\text{box}} = \delta \mathbf{U}_{\text{T1}}/N_{\text{link}}$.

376

Combining the plasticity equation (Eq. (14)) and the amplitude of strain relaxation (Eq. (16)), we obtain the frequency f of T1 events, per link:

if
$$\mathbf{V} : \mathbf{U}_d > 0$$
, $f = \frac{h(U_d)}{2\varepsilon_0 U_d} (\mathbf{V} : \mathbf{U}_d)$,
if $\mathbf{V} : \mathbf{U}_d < 0$, $f = 0$. (18)

It depends on the positive eigenvalue $\dot{\gamma}$ of the elongation rate, and on the relative angle θ between the eigenvectors of the elongation rate and strain. If $\cos(2\theta) < 0$, the frequency is zero; if $\cos(2\theta) > 0$, the order of magnitude of the frequency can be estimated: $f \propto \cos(2\theta)\dot{\gamma}$. This extends findings by [29]. It expresses that rearrangements are frequent where the total strain rate is strong, and when the elongation rate is parallel to the pre-existing strain thus loading it through the yield surface.

4 Experimental tests

4.1 Rearrangements: orientation and frequency

This section uses data of Figure 1a.

Figure 7 confirms the mean-field approximation of the model (Eq. (14)), namely that disappearing and appearing links are determined (in average) by the existing strain. More precisely, **P** makes an angle of $0 \pm 9^{\circ}$ with **M** or **U**: disappearing links ℓ_d are mainly in the elongation direction (Fig. 7, top). Their length is 1.2 ± 0.1 times larger than the average of existing links in that direction, $\ell_+ = \sqrt{2\lambda_1}$, where λ_1 is **M**'s largest eigenvalue (Fig. 7, middle). Conversely, links ℓ_a appear in the contracted direction of **M**, with a length 1.1 ± 0.1 times the average of existing links, $\ell_- = \sqrt{2\lambda_2}$ (Fig. 7, bottom).

The jump in elastic strain is therefore oriented along U. Its amplitude is approximately

$$\delta \mathbf{U}_{\mathrm{T1}} \approx \begin{pmatrix} \ell_d^2/\ell_+^2 & 0\\ 0 & -\ell_a^2/\ell_-^2 \end{pmatrix} \\ \approx \begin{pmatrix} 1.3 \pm 0.2 & 0\\ 0 & -1.2 \pm 0.2 \end{pmatrix},$$
(19)

using $\delta \mathbf{U}_{\mathrm{T1}} \approx \delta \mathbf{M} \times \mathbf{M}^{-1}/2$, from the differentiation of equation (5) [14]. We conclude from equation (16) that each rearrangement changes the elastic strain per link by a constant amount:

$$\varepsilon_0 \simeq 1.2 \pm 0.2. \tag{20}$$

Strain is decreased by slightly more than one average length in the elongation direction, and increased by slightly more than one average length in the orthogonal direction.

Rearrangement frequency is well predicted (Fig. 8) by equation (18). The main parameter required, namely U_Y , is directly read from measurements of U: here $U_Y = 0.15$, which is reasonable for a foam with 4% liquid fraction. Second, the shape of the elasto-plastic transition has been



Fig. 7. Histograms of measurements in all regions of the foam (Fig. 1a). Top: angle between topological events and elongation. Middle: length of disappearing links compared to average length in elongation direction ℓ_+ . Bottom: length of appearing links compared to average length in compressed direction ℓ_- .



Fig. 8. Frequency of rearrangements: observed versus predicted. Each point corresponds to a RVE of the foam, that is, one ellipse of Figure 12. Observations: frequency f_{measured} of rearrangements (per link and second) measured in Figure 1a. Predictions: f from equation (18), setting the yield function to $h = \min((U/U_Y)^2, 1)$ and the yield $U_Y = 0.15$, while $\varepsilon_0 = 1.2$ (Eq. (20)). Solid line: diagonal $f_{\text{measured}} = f$.

chosen as a quadratic h function, which is justified in Section 4.2.

The origin of the upstream/downstream asymmetry of plasticity can now be qualitatively explained, by following a bubble along its streamline (Fig. 9). The obstacle imposes a succession of opposite elongation rates: spanwise before the obstacle, and streamwise after it (see below, Fig. 12). Before the obstacle, the elastic strain and



Fig. 9. Foam flow. Top: superimposed images evidence the streamlines (here from Fig. 1b). Bottom: measured velocity field and streamlines. The thicker solid line highlights the streamline analysed in Figure 10.

the elongation rate are aligned, and the foam is yielding. After the obstacle, it takes some time until the elastic strain fully relaxes, then increases again in the new direction of elongation, orthogonal to the initial one (Fig. 10). Topological rearrangements are therefore concentrated in a smaller region.

4.2 Maps

This section presents the spatial distribution of measurements plotted as ellipse maps (matrix fields), which simultaneously display: position, orientation, anisotropy and frequency of rearrangements. Again, we predict plasticity from the measured elastic strain and total strain rate, using a yield strain U_Y directly read from the onset of a plateau in the plasticity fraction h, without adjustable parameter (see Fig. 11 for the different measurements). A smooth quadratic plasticity fraction h is chosen (as observed from rheometric measurements [9]), to account for possible plastic events below the yield. This *direct* prediction of **P** from h is robust: in fact, it is especially significant near the yield strain, where h is close to 1 (whatever the choice of h).

Conversely, the *inverse* estimate of h, from \mathbf{P} , is noisier. In fact, differences between various candidates for the h function are more significant far from the yield point, where measurements are both smaller and with less statistics. In practice, we suggest to estimate the amplitude of the deviatoric part \mathbf{P}_d of plastic strain with the amplitude $P_d = (\mathbf{P}_d : \mathbf{P}_d/2)^{1/2}$; and the amplitude of the deviatoric part of the total strain rate with $V_d = (\mathbf{V}_d : \mathbf{V}_d/2)^{1/2}$.



Fig. 10. Measurements along the streamline shown in Figure 9 bottom. Top: loading and unloading of elasticity, U_d . Bottom: T1 frequency along streamline, as in Figure 8; solid line: experimental f_{measured} ; dashed line: predicted f.



Fig. 11. Yield strain versus liquid fractions for the different setups of Figure 1: \circ , (a); \bullet , different experiments of (b); \Box , (c); \triangle , (d). Trend of data (b) can be qualitatively compared to a quadratic law [30]: $U_Y = C(\Phi_c - \Phi)^2$, with Φ_c corresponding to the rigidity limit liquid fraction ($\approx 30\%$ for setup (b) [14]) and C an empirical constant equal to 1.1.

If we assume that **P** and **V** are nearly parallel (which is not really the case with obstacles), equation (14) can be projected on the same axis: we obtain an estimate of the plasticity fraction as $h \simeq P_d/V_d$.

4.2.1 Flow around an obstacle: wet foam

For a wet foam, upon flowing around the obstacle (Fig. 12), we observe that the amplitude of U increases then decreases, while V changes orientation. U ellipses have a "capsule" shape before the obstacle, with a negative compression larger than the positive expansion. This is due to bubble compressibility: in this experiment bubbles can expand their height (in the water bath) and thus retract their horizontal area [15].



Fig. 12. Wet foam flowing around an obstacle (Fig. 1a). Measurements of **U** (a) and **V** (b) yield, through equation (14) with quadratic plasticity function h, the theoretical prediction of **P** (c), in good agreement with its measurement (d). Scale: for **U**, bar = 1 (no unit); for **V** and **P**: bar = 1 s⁻¹.

The agreement between prediction (Fig. 12c) and measurement (Fig. 12d) of \mathbf{P} is very good. In particular, we predict well the spatial distribution of T1 events: they occur mostly just before the obstacle, and in a narrower region after it. For horizontal positions just on the right of the center of the obstacle, the flow tends to decrease



Fig. 13. Estimate of h. Symbols: measurements from Figure 1a of $P_d/V_d \simeq h$. We represented the average (circle) and standard deviation (vertical error bar) after binning data along the horizontal axis on equal interval sizes (horizontal error bar). Dashdots: $h(U_d) = \min((U_d/U_Y)^2, 1)$, quadratic up to $U_Y = 0.15$.

the existing strain ($\mathbf{V} : \mathbf{U}_d < 0$): the predicted frequency vanishes.

We also predict well the direction of rearrangements, as well as their amplitude, represented by the direction of the coffee beans and their size, respectively. We do not observe in experiment purely elastic areas and purely plastic areas with a sharp transition line. This is why a discontinuous plasticity function h would be unsuitable, and we use a smooth one (Eq. (14)).

Conversely, statistics are just good enough that we can extract h from measurements. (Fig. 13). We observe that h increases, more or less like the proposed $(U/U_Y)^2$, up to $U_Y = 0.15$, then saturates. Interestingly, it plateaus at a value ~ 0.6 ± 0.1 significantly lower than 1. This is probably because after the obstacle V unloads U, which decreases before the foam enters the fully plastic regime.

4.2.2 Flow around an obstacle: dry foam

The same experiment with a dry foam (Fig. 1b) yields a qualitatively similar behaviour for \mathbf{U} , \mathbf{V} and \mathbf{P} (Fig. 14). Quantitatively, however, the maximum value of U_d is here 0.45, which is a reasonable value for a dry foam [22]. The spatial variation of \mathbf{U} , \mathbf{V} and \mathbf{P} is restricted to a much narrower range. Note that ellipses are more circular: foam is much less compressible than the previous dry case.

This means that we measure larger values but on much less points, resulting in poorer statistics. Still, the agreement between prediction and measurements of \mathbf{P} is qualitatively correct. Extracting h from the data is also qualitative, with apparently a plateau as low as 0.4 (Fig. 15).

4.2.3 Flow through a constriction

When a foam is forced to flow through a constriction (Fig. 1c), along any streamline U_d steadily increases. The



Fig. 14. Dry foam flowing around an obstacle (Fig. 1b). Same caption as Figure 12, except that for V and P bar = 0.063 s^{-1} . See similar figures in the companion paper [14].

constriction is here so narrow, comparable to the square root of a bubble area, that just at the constriction the continuous description (and thus our measurement method) breaks down. The influence of the constriction is visible far uphill: U_d is widely distributed, and we obtain good statistics (Fig. 16).



Fig. 15. Same figure as Figure 13, but for the dry foam of Figure 1b, $U_Y = 0.45$.



Fig. 16. Foam flowing through a constriction (Fig. 1c). Same caption as Figure 12, except that for U bar = 0.1, and for V and P bar = $0.25 \,\mathrm{s}^{-1}$. See similar figures in the companion paper [14].

Conversely, **V** is more localized near the orifice, and thus, as expected from equation (14), so is **P**. Plasticity is indeed oriented by the elastic strain, the angle between the main axis of **P** and of \mathbf{U}_d is $1 \pm 5^\circ$. Concerning the plasticity amplitude, we observe that P_d is much smaller than V_d when the foam enters in the field of view: P_d/V_d undergoes a 5-fold increase until it reaches 1 at the constriction (Fig. 17a), and U_d plateaus. We measure $U_Y \sim 0.32$; which is reasonable for a foam with $\sim 1\%$ liquid fraction.

The direct estimate of the plasticity fraction as $h \simeq P_d/V_d$ is obtained with reasonably good statistics (Fig. 17b). It plateaus close to 1, confirming that the sat-



Fig. 17. Constriction: analysis of Figure 1c. (a) P_d/V_d (open circles) and U_d (closed circles) versus the distance to the constriction. (b) Plasticity fraction h estimated as $h \simeq P_d/V_d$ versus strain U_d (data from (a)): same figure as Figures 13 and 15, here with $U_Y = 0.32$.

uration is reached. We obtain an extremely good quantitative agreement of the predicted plasticiy with the measurement (Fig. 16).

4.2.4 Couette flow

In a steady Couette flow (Fig. 18), as expected U respects the circular symmetry: it does not vary orthoradially. The advantage is that we can improve the measurements by averaging orthoradially. The drawback is that we have very few independent measurements (here 6), along the radial direction. U is significantly different from zero everywhere. Near the rotating (inner) disk, it means that U saturates. Near the fixed (outer) disk, it is probably a residual strain due to the foam preparation (there are not enough T1's to relax it). U_d is rather low (at most of order of 0.1), which is consistent with the high liquid fraction. All these findings confirm those of reference [21].

As expected, \mathbf{V} similarly does not vary orthoradially. It decreases quickly with the distance to the fixed disk, so that we have only two independent, non-zero measurements. It is thus impossible to perform the same analysis as in the above flows which truly vary with both space coordinates.



Fig. 18. Foam sheared between concentric disks (Fig. 1d). Same caption as Figure 12, except that for U bar = 0.1, and for V and P bar = 0.25 s^{-1} .

Still, we can predict \mathbf{P} from equation (14): this agrees quantitatively with the measurements. Concerning the orientation of \mathbf{P} , the angle between the main axis of \mathbf{P} and of \mathbf{U} is $-5 \pm 3^{\circ}$, they are indeed aligned. Since the flow is steady, \mathbf{U} is constant along a streamline, and we thus expect $\mathbf{P} = \mathbf{V}$: this agrees qualitatively with the measurements.

5 Conclusion

5.1 Summary

Using the formalism developed in the companion paper [14], we measure in different 2D foam flows the matrices which quantify the elastic strain, the total strain rate and the plastic rearrangements. We identify statistical measurements performed on the detailed structure of bubbles, with large-scale quantities describing the foam as a continuous material.

We then generalize to matrices a previous scalar model [9], and base it on local measurements on individual bubbles. We show that the plastic rearrangements arise from a combination of both the elastic strain and the total strain rate. As shown by the maps, they cannot be predicted from the elastic strain alone, nor from the total strain rate alone.

For instance, in the wet obstacle flow, the spatial symmetry with respect to the obstacle is very different in the three maps. In the dry foam obstacle, the elastic strain extends very far, while the velocity gradient has a narrower extension, but both are needed, and the total strain gives the orientation. In the constriction the elastic strain extends so much that it is mainly the total strain rate which determines where T1's occur.

In a first approximation, the plasticity is described mainly by the behaviour near yielding. The yield strain U_Y is the main relevant parameter. We determine it directly from image analysis and check that the obtained values are reasonable. We then statistically predict the position, orientation, anisotropy and frequency of topological rearrangements in a flowing foam, in good agreement with various experiments.

In a second step, to refine the description and improve the prediction, we introduce the proportion of plasticity, to account for the gradual appearance of plasticity instead of a sharp yield. It is a function $h(U_d)$ (also called "yield function") of the elastic strain, which interpolates between 0 (fully elastic) at small strain $h(0) \sim 0$, and 1 (fully plastic) near yielding, $h(U_Y) \sim 1$. We obtain here estimates of h.

5.2 Discussion

The model presented in Section 3 succesfully predicts the plasticity, and contributes to describe the foam as a continuous material. We now discuss its validity and some of its limitations.

5.2.1 Material

The general approach which links individual and collective descriptions is applicable to other complex materials (see companion paper [14]). The identification between statistical and continuous quantities is restricted to affine flows, which probably applies rather generally to foams and other cellular materials; it also requires that the elastic strain corresponds directly to individual objects, and the plastic strain rate to topological rearrangements: this is probably specific to dry foams and emulsions.

5.2.2 Shear rate

Foams coarsening, due to the diffusion of gas from one bubble to the other, couples to foam rheology at slow time scales [3]. Here, the present experiments are short enough to neglect this effect.

On the other hand, at very high shear rate, the rheology couples to the bubble's internal relaxation times, which are much shorter [10]. In a material like a foam, at such a high shear rate the bubbles' films and vertices do not relax towards the local mechanical equilibrium. Bubbles distort and the foam eventually breaks; in fact, at all velocities explored here, if the foam exists it means that the shear rate is slower than the internal relaxation times [20].

5.2.3 Dimension

Our formalism is written using matrices, so that it indifferently applies to 2D or 3D systems. In principle it could be tested in truly 2D foams (for instance in numerical simulations) as well as in truly 3D foams (detailed measurements are in progress). Here, for simplicity, we test it on bubble monolayers (quasi-2D foams) which flow horizontally (true 2D velocity field).

The friction due to plates of glass is an external force acting on bubbles, not to be confused with viscous stresses which are internal to the foam. It likely plays a role in the values of the quantities we measure above, but does not affect the *relations* between these quantities; that is, the equations we write are insensitive to this friction. This is in particular the case for the repartition of the total strain rate between its elastic and plastic contributions (Eq. (14)).

5.2.4 Disorder

We have assumed here that the foam is amorphous; that is, RVEs are homogeneous. This excludes foams with: crystalline order; avalanches of T1's; localization of the shear, and shear banding; or one dimension comparable to a bubble diameter.

Here we measure the effects of the function h, and thus indirectly trace back to its approximate expression. This is not precise, since apparently, the exact shape of h does not affect much the predictions. This implies that, reciprocally, once h is known, predictions based on it are very robust. We are thus currently trying to measure directly the function h for various foams, with different liquid fractions and disorders.

5.3 Perspectives: constitutive equation

5.3.1 Closing the set of equations

In principle we could study the fluctuations in time and space of the three matrices we measure: \mathbf{U} , \mathbf{P} , \mathbf{V} . However, we observe experimentally that their averages vary smoothly with space. As a first step, we want to use these average smooth fields to obtain a continuous description of 2D foam flows. For that purpose, we need a closed set of equations to relate these three matrices.

First, the kinematic equation (10) means that the total strain rate is shared between elastic and plastic contributions. It is equivalent to rewrite it as

$$\frac{\mathcal{D}\mathbf{U}}{\mathcal{D}t} = \mathbf{V} - \mathbf{P}.$$
 (21)

In other words, the elastic strain is loaded by the total applied strain rate, that is, the velocity gradient; this process is limited by the plastic strain rate, which saturates the elastic strain.

Second, the plasticity equation (14) describes how the total strain rate is shared between change of elastic strain and plastic strain rate. It involves the fraction of plasticity, h. Equation (14) can be combined with equation (15) into a single equation, using the Heaviside function

$$\mathbf{P} = h(U_d) \ \mathcal{H}(\mathbf{V} : \mathbf{U}_d) \ \frac{(\mathbf{V} : \mathbf{U}_d) \ \mathbf{U}_d}{2U_d^2} \ . \tag{22}$$

Third, \mathbf{V} is directly related to the symmetrised gradient of the velocity field. In fact, combining equations (8) and (7) yields

$$\mathbf{V} \stackrel{\text{affine}}{\simeq} \frac{\boldsymbol{\nabla} \mathbf{v} + \boldsymbol{\nabla} \mathbf{v}^t}{2} \,. \tag{23}$$

Fourth, the velocity field is determined by the fundamental equation of dynamics, that is, momentum conservation (Navier-Stokes):

$$\rho\left(\frac{\partial \mathbf{v}}{\partial t} + (\mathbf{v} \cdot \nabla)\mathbf{v}\right) = \operatorname{div} \cdot \boldsymbol{\sigma} + \mathbf{f}_{\mathrm{ext}}, \qquad (24)$$

where ρ is the foam density, σ the stress tensor, and \mathbf{f}_{ext} the external forces (such as the friction on the glass plates) not included in the stress.

Fifth, we now need the constitutive equation itself. That is, a dynamical equation which determines the elastic and viscous contributions to the stress, from the current elastic strain and strain rate. We suggest to generalize equation (4), now using matrices

$$\boldsymbol{\sigma} = 2G\mathbf{U}_d + K\operatorname{Tr}(\mathbf{U})\mathbf{I} + 2\eta\mathbf{V}.$$
 (25)

We have added here a compression modulus K; in practice, for foams, it is much larger than the shear modulus. We write the material-dependent parameters as scalar, which is correct for an isotropic disordered foam; in an anisotropic material they would be rank-four tensors. The drawback of the external friction on plates (mentioned in Sect. 5.2.3) is that it prevents us from measuring locally the viscous contribution to the stress in equation (25). We thus have assumed that its viscosity is linear, *i.e.* η constant. It is possible in principle that η depends on \mathbf{U}_d or \mathbf{V} , thus introducing some non-linearities.

In conclusion, this fully closed set of equations can be solved from the knowledge of two field variables: \mathbf{v} for transport and \mathbf{U} for internal strain.

5.3.2 Unified description of elastic, plastic and viscous behaviours

Together, equations (21–25) provide a fully closed system. It is a complete continuous constitutive equation for foams. It treats with an equal importance the elastic, plastic and viscous contributions in any of the three regimes: elasticity, plasticity and flow. It admits as limits the classical equations of elasticity or hydrodynamics, as well as elasto-plasticity, visco-elasticity and visco-plasticity. Extension to higher shear rates [10] is challenging, but not impossible in principle.

It should be enough to perform complete, testable predictions of \mathbf{U} , \mathbf{P} , \mathbf{V} , from the material's properties and the boundary conditions. This is much stronger than Sections 3 and 4, which predict \mathbf{P} from known \mathbf{U} and \mathbf{V} . Such analytical predictions, and the corresponding experimental tests, are in progress. They are outside of the scope of the present paper; still, we can make a few remarks regarding the foam dynamical properties.



Fig. 19. Storage and loss moduli versus strain amplitude for a monodisperse emulsion. Symbols: experimental G' (squares) and G'' (triangles) in a close-packed emulsion (Fig. 1 of Ref. [31]). Lines: model calculated with the complete set of equations (21–25). The method is the same as in reference [9]; the correspondence between scalars and matrices is obtained by replacing $(\sigma, U, \dot{\varepsilon})$ with $(\sigma_{xy}, 2U_{xy}, 2V_{xy})$. The factors 2 come from the tensorial generalization (Eq. (25)) of the scalar equation (Eq. (4)); this is why the yield strain is half of that used for the scalar case (Fig. 9 of Ref. [9]). The parameters are directly obtained from the experimental data: shear modulus $G = 1.7 \cdot 10^3$ Pa, yield strain $U_Y = 0.0223$, and viscosity $\eta = 30$ Pa s. The yield function used here is $h = (U/U_Y)^2$.

In oscillatory regime, the system of equations (21-25) predicts the storage and loss moduli G', G'' (Fig. 19). Since these are scalar quantities, this is very similar to predictions based on equation (4) [9].

In stationary regime, while we cannot yet measure locally the viscous contributions to the stress, we can indirectly measure its consequences at the global level. In fact, the force acting on a circular obstacle (Fig. 1a) results from the integral of the stress over the obstacle boundary: experimentally, it increases linearly with the foam velocity, with a non-zero intercept [15]. This agrees with equation (25). A consequence is that, in a stationary regime where the elastic contribution to stress is constant, the foam seems to behave as a visco-plastic (Bingham) fluid.

In any transient regime, however, the full visco-elastoplastic nature of the foam has to be taken into account. The internal variable \mathbf{U} , often overlooked, thus plays a role as important as that of \mathbf{P} or \mathbf{V} .

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