THE SHELL MAP *

The structure of froths through a dynamical map

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1. Introduction

The shell map is a very simple representation of the structure of foams, combining the geometrical (random tiling) and dynamical (loss of information from an arbitrary cell out) aspects of disorder. We will illustrate it and give several examples, including a few arising from discussions in Cargese. This chapter is written by following the main lines of two previously published papers [1, 2].

In Nature, space-filling disordered patterns and cellular structures are widespread [3, 4]. These structures (froths) are partitions of $D$-dimensional space by convex cells. Disorder imposes that each vertex has minimal number of incident edges, faces and cells ($D + 1$ edges incident on a vertex, $D$ faces incident on a edge, $D - 1$ cells incident on a face, in $D$-dimensions, Fig.1). In this respect, a froth is a regular graph, but the number of edges bounding each face, the number of faces bounding a polyhedral cell, etc., are random variables [5]. Minimal incidences implies also that the topological dual of a froth is a triangulation (Fig.1), a useful representation of packings. Indeed, for any given packing, or point set, one can construct the Voronoï tessellation [6], which is a space-filling assembly of polyhedral cells. When the starting points are disordered (no special symmetries) the Voronoï tessellation is a froth. The space filled by the froth can be curved (Fig.1). This is for instance the case in amphiphilic membranes [7], fullerenes, the basal layer of the epidermis of mammals [8] or the ideal structure of amorphous materials [9, 10]. Disorder does not necessarily imply inhomogeneity. On the contrary, in many cases, disordered froths are very homogeneous (cells with very similar sizes and regular shapes). This is -for instance- the case in the epidermis, where the biological

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cells have homogeneous sizes and isotropic shapes but the structure is disordered. Indeed, the disordered arrangement is the one which best guarantee both the partition of the curved space into similar cells and the invariance under mitosis and detachment.

The interplay between disorder and curvature is illustrated in this chapter by representing the froth as organized in concentric layers of cells around an arbitrary central cell [1] (Figs. 2 and 2.2). The structure is built from the central cell outward like an ever expanding jigsaw puzzle without boundary. The radial map, from one spherical layer of cells to the next, is the logistic map [11], and the geometrical tiling is expressed mathematically as a dynamical system [12]. The isotropy of the disordered structure is expressed locally by averaging over each layer. The over-all translational invariance is manifest in the independence of the structure and properties on the choice of the central cell. The radial map from one layer to the next includes both effects of disorder and of space curvature.

In a two dimensional froth, given a cell with $n$ edges, one can define its topological charge as $q = 6 - n$. The total charge of a froth with $N$ cells is the sum over the charges of each cell: $Q_T = \sum_i q_i = (6 - \langle n \rangle)N = 6\chi$, with $\chi$ the Euler-Poincaré characteristic of the manifold tiled by the froth [13, 14]. The charge is a topological invariant, it cannot be generated or destroyed and the local topological transformations in the froth redistribute it between adjacent cells. The total charge is equal to zero in Euclidean froths, it is 12 for froths on the surface of a sphere.
and it is negative in spaces with negative Gaussian curvature. In two dimensions, it is therefore possible to define the curvature of a surface tiled by a froth by analyzing the local topological configuration of its tiles. In three dimensions, for $N \to \infty$ or in closed elliptic froths, the Euler’s relation is homogeneous ($\chi = 0$, independently on the space-curvature) and from the Gauss-Bonnet formula [14] it is not possible to distinguish the global curvature of a tiled manifold from the local average properties of its tiles. On the other hand, we show that, also in three dimensions, the map makes possible to define the curvature of the space from local topological configurations [1]. This is done by simply computing the number of cells in successive layers.

In this chapter, we describe the map which gives the number of cells $K_j$ in a layer distant $j$ from the central cell as function of the average topological properties of the cells in the previous layers (section 2) [1]. We discuss the link between the map and the space curvature giving examples in two and three dimensions (section 3) [1, 2]. By using the map we exploit the freedom of constructing a froth with different local topological configurations and determine the three dimensional Euclidean structures which maximize such freedom (section 4) [1, 2].

2. From a cell to the whole froth, a topological map

All froths can be studied as structured in concentric layers of cells which are at the same topological distance from a given central cell. The topological distance between two cells is the minimum number of edges crossed by a path from one cell to the other [1]. The layers are closed rings of irregular polygons in two dimensions and spherical caps of irregular polyhedra in three dimensions. The cells making the layer $j$
can be distinguished in two categories. Some cells have simultaneously neighbours in the layers $j - 1$ and $j + 1$, these cells make themselves closed layers and constitute the skeleton of the shell-structure. Other cells (or clusters of cells) are local inclusions (topological defects) between the layers of the shell-skeleton (they don’t have neighbours in the layer $j + 1$). The shell-skeleton is itself a space-filling froth, hierarchically organized around the germ cell. Once the germ cell is chosen, the shell-structure and its skeleton are univocally defined, but different germ cells generate different skeletons. We call shell-structured-inflatable (SSI) a froth free of topological defects. In this case, the shell-structure and its skeleton coincide. An example of a two dimensional SSI disordered cellular structure is given in Fig.(2). A three-dimensional regular SSI structure (the Kelvin froth [15]) is given in Fig. (2.1). In this paragraph, we first study SSI froths and then generalize the results to generic froths.

2.1. A topological map for SSI froths

For SSI structures, a recursive equation gives the number of cells $K_j$ in a layer at a distance $j$ from the germ cell, as a function of the number of cells in the previous two layers. In two dimensions, one can easily verify that [1, 12, 16, 17] (see Fig.2)

$$K_{j+1} = s_j K_j - K_{j-1} \quad (\text{for } j \geq 1),$$

(1)
Figure 4. (a) An example of 3D cellular system: the “Kelvin froth”. A part of its shell-network is shown in (b), it is the 2D elliptic set of facets which are bounding externally a layer. (c) Construction of the 3D “Kelvin cell” from the 2D shell network. (d) The “twisted Kelvin cell”.

where the inflation parameter $s_j$ is related to the average topological properties of the cells in layer $j$,

$$s_j = \langle n \rangle_j - 4,$$

with $\langle n \rangle_j$ the average number of neighbours per cell in layer $j$. The map starts with $K_0 = 0$ and $K_1 = n = \text{number of neighbours of the central cell}$. For example in the hexagonal lattice one has $n = 6$, $\langle n \rangle_j = 6$ and therefore $s_j = 2$. In this case the map (1) correctly gives $K_j = 6j$.

In three dimensions, one can write a recursive equation similar to Eq.(1) [1, 2]

$$K_{j+1} = s_j K_j - b_j K_{j-1} + c_j \quad \text{(for } j \geq 1)$$

with

$$s_j = \frac{1}{2}(\langle f \rangle_j - 6)(\langle n^N \rangle_j - 4) - b_j - 1$$

$$b_j = \frac{\langle n^N \rangle_j - 4}{\langle n^N \rangle_{j-1} - 4}$$

$$c_j = 2\left(b_j - s_j + 6(\langle n^N \rangle_j - 4) + 1\right).$$

Here $\langle f \rangle_j$ is the average number of neighbours of the cells in layer $j$. The quantity $\langle n^N \rangle_j$ is the average number of edges per face in the 2D elliptic set of facets which are bounding externally layer $j$ (the
shell-network, see Fig.2.1(a,b) and [1, 2] for details). The map starts with \( K_0 = 0 \) and \( K_1 = f = \text{number of neighbours of the central cell} \). For example in the Kelvin structure one has \( f = 14, \langle f \rangle_j = 14 \) and \( \langle n^N \rangle_j = \frac{5+(24j^2+24j+7)}{1} \). By using the map (3) we obtain \( K_2 = 50, K_3 = 110 \) and in general \( K_j = 12j^2 + 2 \), which is the correct answer [18].

Equations (1) and (3) take a much simpler form if the average topological properties of the cells in layer \( j \) are independent on the layer number (i.e. \( \langle n \rangle_j = \langle n \rangle, \langle f \rangle_j = \langle f \rangle \) and \( \langle n^N \rangle_j = \langle n^N \rangle \)). In general, these quantities can vary as one goes from one layer to the next (as for the Kelvin case). However, in the limit \( j \to \infty \) they must converge towards the averages over the whole structure (in the Kelvin case \( \langle n^N \rangle_j \to 5 \)). Moreover, since the choice of the central cell is arbitrary, in disordered systems this asymptotic behaviour is reached much faster if one averages over the central cell first. Finally, the relations (1) and (3) are also valid when the “central cell” is a cluster or even an infinite set of cells. For instance the starting configuration (the cells with \( j = 0 \)) can be an infinite linear strip of adjacent faces in 2D or a planar layer of adjacent bubbles in 3D. In this case the asymptotic behaviour and the average over the system can be set from the beginning.

When the average topological properties of the cells in layer \( j \) are independent on the layer number, Eqs.(1) and (3), give [1, 12]

\[
\begin{pmatrix}
\tilde{K}_{j+1} \\
\tilde{K}_j
\end{pmatrix} =
\begin{pmatrix}
s -1 \\
1 & 0
\end{pmatrix}
\begin{pmatrix}
\tilde{K}_j \\
\tilde{K}_{j-1}
\end{pmatrix}.
\]

(5)

This equation is the logistic map [11]. In two dimensions, \( \tilde{K}_j = K_j \) is the number of polygonal cells which are making the layer \( j \). In three dimensions, for \( s \neq 2 \), \( \tilde{K}_j \) is the number of polyhedral cells making the layer \( j \) plus the additive constant \( c/(2-s) \). When \( s = 2 \) Eq.(5) cannot be used and Eq.(3) must be used instead.

2.2. Topological defects and non-SSI froths

In almost all disordered froths, non-SSI inclusions (topological defects) are present. An example of non-SSI froth is given in Fig.2.2. A defect in layer \( j \) is a cell (or a cluster of cells) which have no neighbours in layer \( j+1 \). Examples of 2D topological defects are given in Fig.2.2. The fraction \( \delta \) of defects in layer \( j \) respect to the total number of cells, tends to a constant as \( j \) tends to infinity. Typically, in 2D physical froths one finds \( \delta \simeq 0.1 \) [19], whereas very disordered computer simulated froths have \( 0.15 < \delta < 0.6 \) [20].

Maps (1) and (3) apply to SSI froths only. But, for any froth, given a central cell, one can always single out the topological defects. Then
Figure 5. Example of non SSI froths (left). Defects, in layer $j$, are cells which have no neighbours in layer $j + 1$. They are topological inclusions between the layers that can be considered as vertex- or edge-renormalizations (right). The topological map doesn’t apply to such configurations, but it applies on the shell-skeleton, which is the SSI structure obtained by eliminating the defects.

the reduction to the shell-skeleton is made by eliminating the defects. The shell-skeleton is itself a space-filling froth hierarchically organized around the germ cell, and it is SSI by construction. Maps (1) and (3) are therefore applicable on this structure.

3. Space curvature from the map

Let start from a given cell of the froth and count its number of neighbours $K_j$ at a distance $j$. We expect different behaviours for $K_j$ v.s. the distance $j$ depending on the curvature of the manifold tiled by the froth, as schematically shown in Fig.3. Mathematically, these different behaviours can be easily studied from Eq.(5) when $s_j = s$ independent of $j$. In this case, one has the following solution of the map (5) [1],

$$
\tilde{K}_j \propto \left( \exp(\varphi_j) - \exp(-\varphi_j) \right).
$$

(6)

The parameter $s$ separates the solutions into classes. The region $|s| < 2$ is associated with $\varphi = i \cos^{-1}(s/2)$. It has bounded, finite trajectories in the $(j, K_j)$ parameter space. This is the characteristic behaviour of cellular tilings of elliptic manifolds (see Fig.(3c)). For $|s| > 2$ one has $\varphi = \cosh^{-1}(s/2)$, and unbounded exponential trajectories (typical of hyperbolic tilings, Fig.(3a)). The point $s = 2$ separates the elliptic
from the hyperbolic regions. It corresponds to Euclidean tilings and the solution of Eqs.(1) and (3) is $K_j \propto j^{D-1}$, with $D$ the space-dimension (this is the behaviour expected from elementary geometry, Fig.(3b)). Note that this connection between the space curvature and the inflation parameter $s$ applies both in two and three dimensions.

In two dimensions, equation (6) is the topological version of the relation (attributed to Gauss) between the circumference $C$ of a circle and its radius $\rho$ measured on a manifold of constant curvature $R$ (where $R = \text{Ricci curvature scalar} = 2 \times \text{Gaussian curvature}$),

$$C = \frac{\pi}{\sqrt{-\frac{R}{2}}} \left( \exp(\rho \sqrt{-\frac{R}{2}}) - \exp(-\rho \sqrt{-\frac{R}{2}}) \right). \quad (7)$$

In our case, $C$ represents the topological length ($K_j$) of the boundary of the cluster with topological radius $j$ and $\sqrt{-\frac{R}{2}}$ is the parameter $\varphi$ of Eq.(6), given here in terms of the curvature.
Here are some examples of regular 2D froths with different $s$. To $s = -1$ corresponds an elliptic froth made with four triangles, i.e. the surface of a tetrahedron. The value $s = 0$ corresponds to an elliptic froth made with six squares, the surface of a cube. To $s = 1$ is associated an elliptic froth with pentagonal faces, the surface of a dodecahedron. It is known that the hexagonal lattice ($s = 2$) is the only regular froth which tiles the Euclidean plane. Froths with heptagonal ($s = 3$) or octagonal faces ($s = 4$) tile hyperbolic surfaces. Two-dimensional disordered tilings can be generated by Voronoï [6] tessellations around points at random on a surface with arbitrary curvature. Here the map must be studied experimentally case by case. In the Euclidean plane, the linear behaviour of $K_j$ v.s. $j$ predicted by our map, has been experimentally confirmed by many measurements on natural and computer generated froths [20, 19, 21]. The behaviour of $K_j$ in random Voronoï networks constructed on curved surfaces is under investigation [22].

In two dimensions, the classification by the map (5) is identical to that provided by the combination of the Gauss-Bonnet theorem [14] and Euler’s equation:

$$\int \frac{R}{2} da = 2\pi \chi = \frac{\pi}{3} Q_T = \frac{\pi}{3} (6 - \langle n \rangle) N = \frac{\pi}{3} (2 - s) N , \quad (8)$$

where $N$ is the total number of cells in the system. Equation (8) shows clearly that the parameter $s < 2$ corresponds to global positive Gaussian curvature, $s > 2$ to negative curvature and $s = 2$ to the Euclidean limit. In 2D, the determination of the curvature by the map (5) or by the Gauss-Bonnet theorem are identical, but the two methods are independent. In three dimensions, the Euler’s relation is homogeneous and from the Gauss-Bonnet formula it is not possible to distinguish the global curvature of a tiled manifold from the local average properties of its tiles. However, the map (5) is still applicable. It provides therefore a general means of describing the curvature from topological considerations, also in 3D.

Here are two example of 3D froths with different inflation parameter $s$. The regular froth made by packing dodecahedras has $\langle f \rangle = 12$ and $\langle n^N \rangle = 5$, which substituted into Eq.(4) leads to $s = 1$, indicating therefore that this froth tiles a positively curved space. Indeed, it is known that it is a closed structure made with 120 dodecahedras. The Kelvin froth of Fig.2.1 has $\langle f \rangle = 14$ and asymptotically $\langle n^N \rangle = 5$, corresponding to $s = 2$, which correctly indicate that the Kelvin’s cells fill the Euclidean 3D space.
Figure 7. Three dimensional froths generated by using the map starting from the shell network. The shell network is always the result of a superposition of two three connected networks, in the examples (a), (d) it is the superposition of two hexagonal lattices.

4. Construction of three dimensional disordered structures from the map

Three-dimensional space-filling cellular systems are highly correlated structures. Each cell has a different shape and there exist in general very few configurations where all cells pack together and fill space without gaps or overlaps. The space-filling condition is a very important constraint which strongly determines the properties of disordered structures.

In froths, the incidence numbers (number of edges on a vertex, of faces on an edge, etc.) are fixed at their minimal value. By contrast, the coordination number (number of faces of a 3D cell, number of edges of a 2D face) is a random variable and its average characterize the structure topologically [5]. In two dimensions, the Euler relation fixes the average number of edges per face (see Eq.(8)). In three dimensions, the Euler relation associates the average number of edges per face $\langle n \rangle$ with the average number of faces per cell $\langle f \rangle$

$$\langle f \rangle = \frac{12}{6 - \langle n \rangle}.$$  \hspace{1cm} (9)

So that the two average coordination numbers are related, and one of these two quantities is not constrained.
The representation of the cellular system in concentric layer around a given central germ cell can be also regarded as an iterative way of constructing disordered packings. In the Euclidean space (and in the asymptotic limit when \( \langle f \rangle_j \) and \( \langle n^N \rangle_j \) are independent of \( j \)), the inflation parameter of the map is fixed at \( s = 2 \). For this value of \( s \), Eq.(4) gives

\[
\langle f \rangle = 6 + \frac{8}{\langle n^N \rangle - 4}.
\] (10)

This equation relates the average coordination \( \langle n^N \rangle \) of a two dimensional structure (the shell-network) with the average coordination \( \langle f \rangle \) of the three dimensional cellular system. The shell-network is the result of a superposition of two elliptic 3-connected networks, the “incoming” and the “outgoing” froths [1]. The pattern of edges constituting the shell-network sets the value of \( \langle n^N \rangle \). Therefore, Eq.(10) allows us to construct systematically 3D Euclidean SSI froths starting from 2D shell-networks.

The simplest 2D froth is the hexagonal lattice. The examples displayed in Fig.2.1 and Fig.4(a-e) illustrate the construction of ordered, monotiled 3D froths from a shell-network generated by different superpositions of two hexagonal lattices (see also [1, 23]). The cell in Fig.2.1(c) is topologically equivalent to Kelvin’s \( \alpha \)-tetrakaidecahedron (it builds up the Kelvin froth shown in Fig.2.1(a)), and the cell 2.1 (d), to its twisted variant [24, 25]. Both structures have \( \langle f \rangle = 14 \) and asymptotically \( \langle n^N \rangle = 5 \), and fill the Euclidean space. In Fig.4(a) is shown part of a shell-networks with 5-sided faces, generated by the superposition of two “squeezed” hexagonal lattices and its unit cell (b) (see also [1, 23]). This structure has again \( \langle n^N \rangle = 5 \), \( \langle f \rangle = 14 \) and is an Euclidean \( (s = 2) \) space-filler. With another intersection of the two “squeezed” hexagonal lattices, one generates the shell-network shown in Fig.4 (c), it has \( \langle n^N \rangle = 4.8 \) a 3D unit cell (4 (d)) with \( \langle f \rangle = 16 \) (8 quadrilaterals, 6 hexagons and 2 octagons). This unit cell is a monotile Euclidean space-filler that, as far as we know, was reported for the first time in [1]. Recently, this tiling has been proposed [26] as the structure of the ternary crystal \( ThCr_3Si_4 \). Fig.4 (e),(f) show an example of an Euclidean shell-structured-inflatable froth made of two different cells. The shell-network in Fig.4 (e) has also two different tiles with \( \langle n^N \rangle = 32/6 = 5.33... \). The associated 3D unit cell (4 (f)) has \( \langle f \rangle = 12 \) and \( \langle n \rangle = 5 \). This structure is an Euclidean space-filler with \( s = 2 \). One can in general show that any Euclidean shell-structured-inflatable froth made with topologically identical cells can be constructed from a shell-network generated by the superposition of two hexagonal lattices.

We can restrict our attention to a special class of structures where the packed polyhedras have only pentagonal and hexagonal faces. In
In the Z and A15 shell networks there are respectively 12 and 14 pentagons per each hexagon. Therefore, \( \langle n^N \rangle \) result 66/12 and 76/15 for Z and A15 respectively. In the Z phase, successive shells have alternating configurations \((j)\) and \((j+1)\).

this case, the range of variability of \( \langle f \rangle \) is restricted between 12 and 14 by Eq.(9) and (10). The lower value corresponds to a structure with only pentagonal faces: a packing of dodecahedras, and it cannot be realized in Euclidean spaces (it is an elliptic system of 120 dodecahedras). Whether the upper value is realizable or not in Euclidean spaces is still an open question. (The only known structure with \( \langle f \rangle = 14 \) made only with hexagon and pentagons is the Goldberg froth [27], but this packing is hyperbolic [1]). Two structures with coordinations number equal to 13.2 and 13.333... have been obtained by decurving iteratively the curved structure of 120 dodecahedras by introducing hexagonal faces [9, 28]. A natural class of ordered structures made only with pentagons and hexagons are the 24 known TCP. They are Euclidean and have 13.333... \( \leq \langle f \rangle \leq 13.5 \) [30, 31, 32, 33, 34]. The TCP A15, with \( \langle f \rangle = 13.5 \), is the known packing of polyhedral cells of equal volumes with minimal interfacial area per cell [35]. In a previous paper [1], we proved that all TCP structures could be described topologically by the map (5). In figure 4 are given two examples of shell networks generated by combinations of pentagonal and hexagonal tiles only. From these networks the structure of the Z phase and of the A15 phase can be constructed. They have coordination \( \langle n^N \rangle = 66/13 = 5.076... \) (Z) and \( \langle n^N \rangle = 76/15 = 5.066... \) (A15), that substituted into Eq.(10) give correctly \( \langle f \rangle = 13.42... \) for Z and \( \langle f \rangle = 13.5 \) for A15.

Figure 4 summarizes in a plot some of the previous considerations and results. The two lines in Fig.(4) are plots of Eq.(9) and (10). As one can clearly see, in a given structure, \( \langle n \rangle \) and \( \langle n^N \rangle \) are different in general. The packing is organized so that the faces of the shell-network have an average number of sides \( \langle (n^N) \rangle \) different from that of the whole.
structure ($\langle n \rangle$). But there is a point [1] where these two values coincide:

$$\langle n^N \rangle = \langle n \rangle = \langle n \rangle^* = \frac{10 + 2\sqrt{7}}{3}$$  \hspace{1cm} (11)$$

and, from (9) or (10)

$$\langle f \rangle^* = 8 + 2\sqrt{7} = 13.29\ldots$$ \hspace{1cm} (12)$$

When $\langle f \rangle = \langle f \rangle^* = 13.29\ldots$, an arbitrary cell has the freedom to adhere to a layer by any subset of its faces, without adjustment. This freedom gives therefore many more local possibilities for building up the structure. A coordination number $\langle f \rangle = 13.29\ldots$ characterizes therefore an hypothetical SSI topological packing which fills the Euclidean three-dimensional space with the largest possible number of local arrangements, that is with maximum entropy. Coordination 13.29\ldots lies in between the two froth obtained by decurving dodecahedras and it is inside the interval where $\langle f \rangle$ is ranging in natural structures and computer simulated froths [25, 29, 30, 31, 32, 34, 36, 37].
Note that, this structure has a coordination number which is irrational. It cannot be therefore a periodic crystalline structure. It can only be approximated by disordered froths or quasi-crystalline structures.

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