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Tribute to an exemplary man: Yves Couder Morphogenesis, elasticity / *Morphogenesis, elasticity*

Tip growth in morpho-elasticity

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Abstract. Growth of living species generates stresses which ultimately design their shapes. As a consequence, complex shapes, that everybody can observe, remain difficult to predict, even when the growth biology is over-simplified. One way to tackle this question consists in limiting ourselves to quasi-planar objects like leaves in the spring. However, even in this case the diversity of shapes is really vast. Here, we focus on growing tips with the aim to compare their role in elastic growth to classical viscous fingering and dendritic growth. With the help of complex analysis, we show that a parabola under constant growth is free of stress while growing but any growth perturbation will strongly affect its final shape. Two models of finite elasticity are considered: the Neo-Hookean and the poro-elastic model with incompressibility.

Résumé. La croissance biologique génère des contraintes mécaniques qui contribuent à façonner la forme des tissus, des organes et des organismes vivants. En raison de l'extrême complexité des phénomènes de croissance biologique, il est en général impossible de prédire ces formes. Dans certains cas géométriquement simples, par exemple des tissus biologiques minces en croissance quasi-planaire tels que des feuilles, les lois de la mécanique contraignent les formes possibles. Toutefois, l'espace des formes atteignables reste particulièrement vaste. Dans ce compte-rendu, nous nous intéressons au cas particulier des pointes en croissance, que nous décrivons dans le cadre de la théorie de la morpho-élasticité et de la poro-élasticité non-linéaire, et qui partage des similarités frappantes avec deux sujets d'étude classiques en physique : la croissance dendritique et la digitation visqueuse. Les outils de l'analyse complexe sont mobilisés pour montrer qu'une parabole en croissance homogène est stable et ne développe pas de contrainte mécanique. En revanche, la forme de la pointe est fortement affectée par les perturbations du champ de croissance.

Keywords. Nonlinear elasticity, Biological growth, Instabilities, Morphogenesis, Meristem growth.

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1. Introduction

Morphogenesis is the biological process by which living organisms acquire their shape. While biological growth is mostly under the control of genetics, the set of possible shapes of biological objects is ultimately constrained by physical principles. At a cellular scale, growth and division can be described by local, time-dependent, laws. However, integrating such local laws in order to explain the (possibly evolving) shapes of macroscopic objects (leaves, flowers, trees, organs, tumors, biofilms, etc.) is a formidable task. If we limit ourselves to botanics, which is simpler in the sense that growth is less constrained by boundary conditions, and even if we further restrict ourselves to leaves because of their geometric simplicity, a broad diversity appears in nature that perhaps physics and mechanics may help to understand or at least to classify.

Upon observing the growing leaves of trees in spring, one may first notice that some leaves are mostly planar, in particular when the ribs are not too thick compared to the leaf thickness. When this is not the case, the leaf typically buckles between two successive ribs and the differential growth leads to sharp pointed tips, one being along the symmetry axis of the leaf with the others organized on both sides, as observed on holly leaves. But for thin leaves and tiny ribs, on the other hand, the membrane remains planar and the outer contour is smooth, with either regular or undulated boundaries depending on species.

Tip instabilities of growing inert matter are also frequently observed. Archetypal examples of "growing" systems exhibiting morphological instabilities in physics are dendritic (diffusive) growth and radial viscous fingering [1, 2], illustrated in Figure 1. Of course, unlike biological growth, there is no creation of matter in these classical experiments but rather a displacement of a material into an another one. Two broad classes of instabilities in these systems have been documented: tip splitting in viscous fingering and side-branching in diffusive out-of-equilibrium processes. Due to the nonlocal nature of the "growth" processes, ie the competition between surface tension and a diffusive or Laplacian field, understanding these patterns turned out to be a real challenge and both of these topics have been the subject of a considerable amount of experimental, numerical, mathematical and theoretical work in the past forty years.

A naive idea to explain the predominance of side-branching events in dendrite growth and of tip-splitting instabilities in viscous fingering lies on the difference between Laplacian and diffusional fields. Dendrite tips in diffusive systems preserve their size to maintain stabilization by surface tension while growing Laplacian fingers in open geometries keep growing and thus must divide in order to maintain stability by surface tension. This sharp separation of physical processes can be somewhat perturbed by the addition of a localized perturbation. As demonstrated by Yves Couder and his collaborators, trapping of a small bubble at the tip of a growing viscous finger results in a stable finger growing at constant velocity and emitting dendrite-like sidebranches [3–5].

The emergence of a theoretical framework coupling biomechanics to growth in the last years now offers a unique way to understand problems of morphogenesis and embryogenesis where the biology at small scales is strongly coupled to biomechanics, and elasticity, see in particular [6, 7]. By analogy with some of his favorite hydrodynamic instabilities, namely crystal growth and viscous fingering, Yves Couder has suggested some universality in growth patterns potentially dominated by leading growing tips, in particular in botany. More precisely, he suggested that growing tips, described within the formalism of morpho-elasticity, might share some characteristics with Laplacian or diffusive growth. The aim of this paper is thus to scrutinize carefully this hypothesis. Let us stress however, that it took fifty years to understand Laplacian [8] or dendritic growth [9] following the first pioneering works of respectively Saffman and Taylor [10] and Ivantsov [11] and unfortunately, the mathematics of finite or nonlinear elasticity is much more complex than that of diffusive or Laplacian growth. As a consequence, our modest contribution will be limited to the simple case of a parabola under: (i) constant volumetric growth, isotropic



Figure 1. On the left, viscous fingering in an open geometry. The anomalous dendritic finger is due to a pulsating tip. The image is reproduced from Couder, Y., Cardoso, O., Dupuy, D., Tavernier, P. and Thom, W. [3]. On the right, a dendrite of succinonitrile growing in an undercooled melt, exhibiting the characteristic paraboloidal tip and the secondary sidebranching behind the tip. The figure is reproduced from J. S. Langer [4].

or slightly anisotropic and (ii) homogeneous poro-elastic swelling and (iii) to the existence of undulating modes at the leave borders.

2. Finite-Elasticity in complex geometry

The basis of elasticity dates at least from the 17th-century with the first statement *De Potentia Restitutiva* of Robert Hooke (1678), but any application of the first principles remains challenging in bodies of non trivial shapes and of finite size [12]. Indeed, if we really want to analyze the field of deformations of a specific body under an external loading, we are faced to its geometrical description. It is why, in textbooks, examples are restricted to bars, spheres, cylindrical or spherical shells. This challenge is even increased in non-linear elasticity [13–15] since we need to consider both the initial and the final geometries or initial and final configurations. This explains why, only simple geometries have been considered since not only the shape is important but also the writing of equilibrium equations (density of forces and torques or Euler–Lagrange equations) in a more or less standard coordinate systems. As for living systems which to some extent, can be considered as soft elastic tissues, their shapes are much more diverse and also their elastic properties. In addition, it exists strong differences between living and inert matter, the most obvious one being the ability to grow with specific rules. This induces stresses that need to be combined to loading or to other constrains, as the boundary conditions for example.

Keeping the strategies that many have followed for viscous fingering and dendritic growth, that is to focus on growing tips, we consider this shape which has been completely discarded in the context of volumetric growth. Tips are present in nature as the ends of branches, leaves, blades. We restrict on planar or quasi-planar objects to maintain plane strain elasticity. The advantage of this geometry is a simple expression with holomorphic functions for borders, a simple way to construct a curvilinear system of coordinates where the initial shape turns out to correspond to one coordinate and finally a simple writing of the equations of non-linear elasticity with growth, as shown in the following. Adopting the technique of needle-crystals which have been assimilated to parabola or paraboloid, discarding side-branchings, we perform the elastic treatment in parabolic coordinates [16, 17].

2.1. Elasticity in parabolic coordinates

We consider a growing parabolic tip and write all the relevant equations of finite elasticity with growth [15] in the parabolic system of coordinates based on the conformal mapping of the (X, Y) plane:

$$Z = X + IY = -\frac{I}{2}((\mu + I\eta)^2 + 1) = \eta\mu + I/2(\eta^2 - \mu^2 - 1).$$
(1)

If the coordinate η is fixed to one, we derive the following relationships: $X = \mu$ and $Y = -\frac{1}{2}X^2$ which represent a parabola of radius of curvature equal to unity ($\Re = 1$), oriented symmetrically along the negative *Y* axis. Here after, we choose as unit of length this radius of curvature. μ and η are orthogonal coordinates with a scale factor h_{μ} equal to $|\partial_{\mu}\vec{R}|$, where $\vec{R} = X\vec{e}_X + Y\vec{e}_Y$, and the equivalent definition for h_{η} . In the following, we will adopt arbitrarily either the notation $\partial/\partial \alpha$ or the notation ∂_{α} for the partial derivative with respect to α . h_{η} and h_{μ} are the classical notations for changes of coordinate systems, see Morse and Feshbach's book [18]) and for the parabolic coordinate defined by (1), we easily derive $h_{\mu} = h_{\eta} = \sqrt{\eta^2 + \mu^2}$. Considering that *Z* is the complex coordinate in the configuration of reference and *z* in the current configuration, the geometric strain **F** and the elastic strain **F**_e (after conformal mapping transformation of coordinates) are then:

$$\mathbf{F} = \frac{1}{h_{\mu}^{2}} \begin{bmatrix} \partial_{\mu} x \ \partial_{\eta} x \\ \partial_{\mu} y \ \partial_{\eta} y \end{bmatrix} \quad \text{and} \quad \mathbf{F}_{e} = \frac{1}{h_{\mu}^{2}} \begin{bmatrix} \partial_{\mu} x/g_{\mu} \ \partial_{\eta} x/g_{\eta} \\ \partial_{\mu} y/g_{\mu} \ \partial_{\eta} y/g_{\eta} \end{bmatrix}$$
(2)

where g_{μ} and g_{η} are the eigenvalues of a growth tensor, diagonal in the parabolic system of coordinates, possibly anisotropic when $g_{\mu} \neq g_{\eta}$. To simplify, these eigenvalues will be chosen independent of space coordinates but they can be dependent on time if the typical time-scale of growth is long compared to any dissipation process. We choose the simplest model of finite elasticity, the Neo-Hookean modelling [13–15], and we assume incompressibility so:

Det
$$\mathbf{F}_e = 1$$
 and Det $\mathbf{F} = g_\mu g_\eta$ so $\mathscr{J} = \partial_\mu x \partial_\eta y - \partial_\eta x \partial_\mu y - h_\mu^2 g_\mu g_\eta = 0.$ (3)

The elastic energy of the initially growing parabola is then:

$$\mathscr{E} = \frac{E}{2\tau} \int_{\tilde{V}} d\mu \, d\eta \{ (\partial_{\mu} x)^2 + (\partial_{\mu} y)^2 + \tau^2 \{ (\partial_{\eta} x)^2 + (\partial_{\eta} y)^2 \} - 2h_{\mu}^2 g_{\mu}^2 - 2p \mathscr{J} \}$$
(4)

where *E* is the shear modulus and *p* is the Lagrange multiplier which allows to impose the incompressibility constraint. A priori, *p* is a function of both μ and η . The coefficient of anisotropy is given by $\tau = g_{\mu}/g_{\eta}$. We can eliminate the anisotropy coefficient by dilating the variable $\mu = \tau v$ but the price to pay is the change of the scaling factors h_{μ} . Indeed, we must reminder that *h* is a function of μ and η so only when *h* depends on a unique coordinate, the anisotropic case can be treated as the isotropic one (this is the case of the cylindrical or radial geometry). If not, we are really constrained. In this curvilinear coordinate system, the Euler–Lagrange equations read:

$$\begin{cases} \Delta_{\eta,\mu} x = \frac{\partial^2 x}{\partial \mu^2} + \tau^2 \frac{\partial^2 x}{\partial \eta^2} = \frac{\partial p}{\partial \mu} \frac{\partial y}{\partial \eta} - \frac{\partial p}{\partial \eta} \frac{\partial y}{\partial \mu} \\ \Delta_{\eta,\mu} y = \frac{\partial^2 y}{\partial \mu^2} + \tau^2 \frac{\partial^2 y}{\partial \eta^2} = -\frac{\partial p}{\partial \mu} \frac{\partial x}{\partial \eta} + \frac{\partial p}{\partial \eta} \frac{\partial x}{\partial \mu}. \end{cases}$$
(5)

One can notice that (5) are similar to the same equations for the growth of a sample in a stripe, written in cartesian coordinates. If the choice is made of holomorphic function for *x* and *y* and for isotropic growth, $\tau = 1$, the Lagrange parameter *p* is a constant. Because of the incompressibility constraint, our choice for the deformation will be in favor of a solution which restores the parabolic shape. So we hypothesize:

$$x = \lambda_x \eta \mu$$
 and $y = \frac{\lambda_y}{2} (\eta^2 - \mu^2 - 1).$ (6)

The Jacobian of the new configuration is then $J = \lambda_x \lambda_y \cdot (\eta^2 + \mu^2) = \lambda_x \lambda_y h_{\mu}^2$ so the incompressibility condition imposes $\lambda_x \lambda_y = g_{\eta}^2 \tau$, according to (3). But we one cannot verify the second Euler– Lagrange equation for y and new guess of possible solution is not so easy to find. It is why we choose first the isotropic case with $\tau = 1$, $\Delta_{\mu,\eta} x = \Delta_{\mu,\eta} y = 0$ and the pressure p constant and opt for a weak anisotropic perturbation. So, our exact result is limited to isotropic growth and weak perturbed elastic fields will be evaluated once driven by weakly anisotropic growth. Boundary conditions for $\eta = 1$ concern the cancellation of the two components of the nominal stress:

$$S_{\eta,\eta} = E\left(\tau^2 \frac{\partial y}{\partial \eta} - p \frac{\partial x}{\partial \mu}\right) \quad \text{and} \quad S_{\eta,\mu} = E\left(\tau^2 \frac{\partial x}{\partial \eta} + p \frac{\partial y}{\partial \mu}\right) \tag{7}$$

which gives: p = 1 and $\lambda_y = \lambda_x = g_\eta = g$. So finally a parabola which grows isotropically enlarges during development and its radius of curvature \mathscr{R} is simply the growth factor g. This solution differs from the initial configuration. If g > 1, the parabola seems smoother than the initial one while if g < 1, it appears more sharp-pointed. Since this constant homogeneous growth process does not generate any kind of stress, the parabolic shape is stable and no shape bifurcation is then expected, contrary to the growth of a layer [19, 20]. In the next section, we confirm that the parabola, with constant isotropic growth, is a robust shape and that deviation from this shape will require a change in the growth conditions.

3. Variations around the stress-free parabola

The stress-free parabola found previously is an exact solution of the elasto-static growth problem. To find exact solutions, even in the Neo-Hookean approach remains a challenge but it remains possible to slightly perturb the growth conditions. Considering anisotropic growth, the weakness is represented by the small coefficient $\epsilon = \tau - 1$ where τ intervenes in (4), (5), (7). We will use this parameter to expand linearly the main equations.

3.1. Weakly anisotropic growth

At linear order in ϵ , x, y, p and J are transformed into:

$$\begin{cases} x = g(\eta \mu + \epsilon u(\mu, \eta)) \\ y = \frac{g}{2}(\eta^2 - \mu^2 - 1) + \epsilon g v(\mu, \eta) \\ p = 1 + \epsilon \pi(\mu, \eta) \\ \mathscr{J} = g^2(\eta^2 + \mu^2) + \epsilon j(\mu, \eta) - g^2 \tau(\eta^2 + \mu^2). \end{cases}$$
(8)

For $j(\mu, \eta)$, it reads:

$$\mu \left(\frac{\partial u}{\partial \eta} - \frac{\partial v}{\partial \mu}\right) + \eta \left(\frac{\partial v}{\partial \eta} + \frac{\partial u}{\partial \mu}\right) = a_0(\eta^2 + \mu^2) \tag{9}$$

where a_0 is equal to 1 if $\tau \neq 1$ and is zero for the isotropic case, $\tau = 1$.

that we can transform into:

$$\begin{cases} \frac{\partial u}{\partial \eta} - \frac{\partial v}{\partial \mu} = \eta m(\mu, \eta) + a_0 \mu \\ \frac{\partial v}{\partial \eta} + \frac{\partial u}{\partial \mu} = -\mu m(\mu, \eta) + a_0 \eta \end{cases}$$
(10)

where $m(\mu, \eta)$ is an arbitrary function. Notice that m = 0 means that (u, v) are complex conjugates since they verify the Cauchy relations. Deriving the first equation of (10) with respect to η and the second equation with respect to μ and adding both results gives:

$$\Delta u = \eta \frac{\partial m}{\partial \eta} - \mu \frac{\partial m}{\partial \mu} \tag{11}$$

and the same operation consisting in taking first the derivation with respect to μ , then with respect to η gives after substraction:

$$\Delta v = -\eta \frac{\partial m}{\partial \mu} - \mu \frac{\partial m}{\partial \eta}.$$
(12)

These two relationships have to be compared to the linear version of (5) which gives:

$$\Delta u = \left(\eta \frac{\partial \pi}{\partial \mu} + \mu \frac{\partial \pi}{\partial \eta}\right) \tag{13}$$

$$\Delta \nu + 2a_0 = \left(-\mu \frac{\partial \pi}{\partial \mu} + \eta \frac{\partial \pi}{\partial \eta}\right). \tag{14}$$

So we relate π to m:

$$\frac{\partial \pi}{\partial \mu} = \frac{\partial m}{\partial \eta} - \frac{2a_0\mu}{\eta^2 + \mu^2} \quad \text{and} \quad \frac{\partial \pi}{\partial \eta} = -\frac{\partial m}{\partial \mu} + \frac{2a_0\eta}{\eta^2 + \mu^2}$$
(15)

 π satisfies the Poisson equation:

$$\Delta \pi = 4a_0 \frac{\mu^2 - \eta^2}{(\eta^2 + \mu^2)^2}.$$
(16)

The general solution for *u* and *v* can be found more easily if we combine both (41) and (42) and define U = u + Iv

$$\Delta U = -2I \frac{\mathrm{d}\pi}{\mathrm{d}\zeta} \zeta - 2Ia_0 \quad \text{where } \zeta = \mu + I\eta \tag{17}$$

 π is a real function which can be decomposed into an holomorphic one $P(\zeta)$ and a contribution due to the anisotropy. It is easy to show that

$$\pi = \frac{1}{2} \left(P(\zeta) + \bar{P}(\bar{\zeta}) - a_0 \left(\frac{\zeta}{\bar{\zeta}} + \frac{\bar{\zeta}}{\zeta} \right) \right)$$
(18)

and one easily find that

$$\frac{\partial^2 U}{\partial \zeta \partial \bar{\zeta}} = -I/4 \left(\frac{\mathrm{d}\bar{P}(\bar{\zeta})}{\mathrm{d}\bar{\zeta}} \zeta + a_0 \left(1 + \frac{\zeta^2}{\bar{\zeta}^2} \right) \right). \tag{19}$$

Before going further, let us remind the boundary conditions: first symmetry along the *X* axis: $x = U(\mu, \eta) = 0$ and $\partial_{\mu}x = 0 = \partial_{\mu}U = 0$ for $\eta = 0$, second cancellation of the stresses $S_{\eta,\mu}$ and $S_{\eta,\eta}$ for $\eta = 1$, see (7). Let us first consider linear perturbation of the stresses:

$$\Sigma(\mu,\eta) = S_{\mu,\eta} + IS_{\eta,\eta} = -2I \frac{\partial U}{\partial \bar{\zeta}} - \pi \zeta + 2a_0 \zeta = 0 \quad \text{for} \quad \eta = 1, \forall \mu.$$
⁽²⁰⁾

Integrating (19) with respect to ζ , we obtain:

$$\frac{\partial U}{\partial \bar{\zeta}} = -\frac{I}{8} \left(\frac{\mathrm{d}\bar{P}(\bar{\zeta})}{\mathrm{d}\bar{\zeta}} \zeta^2 + 2a_0 \left(\zeta + \frac{\zeta^3}{3\bar{\zeta}^2} \right) \right) + Q'(\bar{\zeta}) \tag{21}$$



Figure 2. On left, quasi parabolic shapes for anisotropic growth. In blue, the growing parabola with $\epsilon = 0$ (isotropic growth), in green the perturbed shape with $\epsilon = -0.1$, in orange similar result with $\epsilon = 0.1$. $\epsilon \ge 0$ means that the growth factor along μ is larger than along η . All lengths are scaled by the growth coefficient g. On right, $\sqrt{\sum_{i,j} S_{ij}^2}$ (with i < j) average stress density inside the parabolic shape. Notice the divergence of the stresses near the tip: $\mu \sim 0, \eta \sim 0$.

 $Q(\bar{\zeta})$ can be evaluated once introduced in the stress function, Equation (20), which vanishes for $\eta = 1$ so for $\zeta = \bar{\zeta} + 2I$. So we get:

$$Q(\bar{\zeta}) = \frac{1}{6} \left(-4/\bar{\zeta} + 12\bar{\zeta} - \frac{7I}{2}\bar{\zeta}^2 \right).$$
(22)

Finally, *U* will be fully determined by integrating (21) imposing $u = 0 = \partial_{\mu}u = 0$ for $\eta = 0$ and imposing also convergence when $\mu - > \pm \infty$. Then, the profile function becomes, under weakly anisotropic growth:

$$\begin{cases} x = g\eta\mu \left\{ 1 + \epsilon \left(-\frac{3}{2} + \frac{2}{3} \frac{\eta^2}{\mu^2 + \eta^2} \right) \right\} \\ y = \frac{g}{2} \left\{ \left(1 + \frac{3\epsilon}{2} \right) (\eta^2 - \mu^2 - 1) + \frac{34\epsilon}{3} - 2\epsilon\eta \left(4 + \eta - \frac{2}{3} \frac{\eta^3 - 2}{\eta^2 + \mu^2} \right) \right\} \end{cases}$$
(23)

where $\epsilon = \tau - 1 = g_{\mu}/g_{\eta}$. So anisotropic growth reduces the radius of curvature of the parabola when $g_{\mu} > g_{\eta}$ as shown in Figure 2 on left. One also notices on right that the stresses diverges at the tip which seems unphysical. This explains that the real tip requires a modification of the growth law or a change in structure like a rib. If we think about botany, it is clear that this model has not enough flexibility to represent a real leaf.

3.2. Free harmonic modes

We consider now free harmonic modes, modes which can be superposed to the stress-free parabola found in Section 2, without modification of the growth condition. These modes can be generated by a noisy environment and may play a crucial role in dynamics. For a growing tissue, such a noise may arise as a consequence of fluctuations in cell growth rates and/or mechanical properties. Interestingly, an interface that is linearly stable in absence of noise may roughen in response to small fluctuations [21, 22]. During this process called kinetic roughening, the interface will remain flat on average but its width (i.e. the typical distance between the crests and

the valleys of the interface) will grow over time. Such a behavior arise when modes of different wavelengths exhibit different dynamics of relaxation and have been observed in various inert or biological systems (for example [23,24]). Free harmonic modes are also observed during dendritic growth. They are considered as the result of a thermal noise mostly located at the tip, generating growing side-branching events on both sides, far from the tip [25]. In viscous fingering in a channel, on the contrary, the finger is especially stable and side-branching is observed only when a perturbation is introduced artificially [3, 5]. At high forcing, a tip-splitting event may occur, similar to the ones observed in radial geometry. Indeed, in this geometry and also in the wedge geometry, a cascade of tip-splitting events is observed as time goes on, mimicking a quasi fractal pattern at long time [26–28]. But this scenario can be inhibited and transformed into dendrites if anisotropy occurs [29]. Here we assume an harmonic mode for the pressure and examine the consequences for the shape of the parabolic growing elastic tip. The wavelength is not a priori specified and modes of any wavelength can be superposed but they do not interact at the linear approximation. In addition, they can be superposed to the previous solution given by (23). So we focus on one mode of wavenumber k which gives a symmetric shape. In addition, we require, as previously, that the asymptotic deformation (u, v) will not be larger than the initial shape so $|u| < \eta |\mu|$ and $|v| < |\mu^2 - \eta^2|$. Most of the results of Section 3.1 will remain valid but with $a_0 = 0$. In particular, the pressure π will be the real part of an holomorphic function as shown by (16), a necessary condition to check incompressibility. For deriving the free modes, we focus on the pressure which is the key quantity to solve the elasticity problem. A symmetric shape will require for the pressure π an even function of μ vanishing on the border. At the linear approximation, it reads

$$\pi = \sinh(k\eta - k)\cos(k\mu) = -Re[\sinh(k + Ik(\mu + I\eta))].$$
⁽²⁴⁾

Solving (19), we deduce on the interface $\eta = 1$

$$\frac{\partial U}{\partial \bar{\zeta}} = -\frac{I}{8} \left(\frac{\mathrm{d}\bar{P}(\bar{\zeta})}{\mathrm{d}\bar{\zeta}} \zeta^2 \right) + Q'(\bar{\zeta}) = Q'(\bar{\zeta}) + \frac{1}{8} k \zeta^2 \cosh(k - \mathrm{i}k\bar{\zeta})$$
(25)

which allows the determination of $Q(\bar{\zeta})$ for ζ being replaced by $\bar{\zeta} = \zeta + 2I$. Finally, integrating *U* and adding a holomorphic function of ζ to limit the growth of the result so that:

$$U = \frac{1}{8}k\zeta^{2}(\cosh(k - ik\bar{\zeta}) - \cosh(k - ik\zeta) + Q(\bar{\zeta}) - Q(\zeta)$$
(26)

we derive:

$$\begin{cases} u = -\frac{1}{2k^2} \sin(k\mu) \{2k \cosh(k) (k\eta \cosh(k\eta) - \sinh(k\eta)) \\ + \sinh(k) (-k\eta \cosh(k\eta) + (1 + 2k^2) \sinh(k\eta))\} \\ + \frac{\mu}{2k} \cos(k\mu) \{-k\eta \sinh(k) \cosh(k\eta) + \sinh(k\eta) ((\eta - 2)k \cosh(k) + \sinh(k))\} \end{cases}$$
(27)

and

$$\begin{cases} v = \frac{1}{2k^2} \cos(k\mu) \{2k \sinh(k)(-k\eta \cosh(k\eta) + \sinh(k\eta)) \\ -\cosh(k)(-k\eta \cosh(k\eta) + (1+2k^2) \sinh(k\eta))\} \\ + \frac{\mu}{2k} \sin(k\mu) \cos(k) \{k\eta \cosh(k\eta) - \sinh(k\eta) - (\eta - 2)k \sinh(k) \sinh(k\eta)\} \end{cases}$$
(28)



Figure 3. Superposition of the growing parabola and free holomorphic modes: on left in green, k = 3, $\epsilon = 0.12/\cosh^2(3)$, in orange, k = 3, $\epsilon = -0.12/\cosh^2(3)$, in blue the non-perturbed parabolic shape. A tip-splitting event accompanies the side-branching on the curve in green but not on the orange-profile. So the undulated mode depends on the tip-perturbation. On the right: zoom on the tip.

which gives for $\eta = 1$

$$\begin{cases} u = -\frac{\sin(k\mu)}{4k^2} \{(4k^2 + 1)\cosh(2k) - 3k\sinh(2k) - 1\} \\ -\mu \frac{\sinh(k)}{2k}\cos(k\mu)(2k\cosh(k) - \sinh(k)) \\ v = -\frac{\cos(k\mu)}{4k^2} \{(4k^2 + 1)\sinh(2k) + k - 3k\cosh(2k)\} \\ +\frac{\sin(k\mu)}{2k} \{k(\sinh^2(k) + \cosh(k)^2) - \sinh(k)\cosh(k)\}. \end{cases}$$
(29)

In Figure 3, we give several possible solutions for the new shapes at fixed wavenumber: k = 3 which means a wavelength close to the tip radius. At linear approximation, it is easy to obtain a tip-splitting event versus a dendrite but the two kinds of mode differ only at the tip which is a difference with viscous fingering. Concerning the shape of leaves, both undulating modes exist but the sharp tips seem to be the most common case. In the next section, we present another growth model triggered by diffusion: the poro-elastic model.

4. Nonlinear poro-elasticity in parabolic coordinates

Let us now extend the preceding analysis to the case of a simple nonlinear poro-elastic material. This model is well-suited to describe swelling gels but also, to some extent, the behavior of vegetal tissues as it incorporates both an elastic and a liquid phase. In response to pressure gradients, the liquid phase moves inside the solid phase in order to minimize the total energy of the system. At equilibrium, a nonlinear poro-elastic material is therefore equivalent to an elastic material with a space-dependent compressibility. As such it also provide a (highly simplifed) model of stress-modulated growth. Ignoring the mixing contributions arising from the interactions between the

solvent molecules and the polymer chains, the Helmoltz free energy is simply that of the highly compressible cross-linked polymer network [30]:

$$\mathcal{W} = \frac{E}{2} (\operatorname{Tr}(F^T F - 2) - 2 \log \det F)$$
(30)

so that the Grand potential to be minimized is simply:

$$\mathscr{E}[\vec{u}] = \frac{E}{2} \int_{\vec{V}} d\mu \, d\eta \left\{ (\partial_{\eta} x)^2 + (\partial_{\eta} y)^2 + (\partial_{\mu} x)^2 + (\partial_{\mu} y)^2 - 2h_{\mu}^2 \right\}$$
(31)

$$-2\log\left(\frac{\partial_{\mu}x\partial_{\eta}y - \partial_{\eta}x\partial_{\mu}y}{h_{\mu}^{2}}\right)\right\}$$
(32)

$$-\frac{\mu_0}{\nu}\int_{\tilde{V}} d\mu \,d\eta (\partial_\mu x \partial_\eta y - \partial_\eta x \partial_\mu y - h_\mu^2). \tag{33}$$

The first two terms in the equation above are just the Helmoltz free energy density integrated over the volume of the poro-elastic tip while the third integral enforces the molecular incompressibility constraint at the thermodynamic equilibrium. Here μ_0 is the chemical potential of solvent molecules in the particle reservoir in contact with the system and v is the volume per solvent molecule. In the context of plant sciences, μ_0 is sometimes referred to as the pore pressure. Similarly to the growing case, we may obtain the Euler–Lagrange equations by requesting the Gâteaux derivative of the functional $\mathscr{E}[\vec{u}]$ above to vanish and we obtain the following set of equations:

$$J\Delta_{\mu,\eta}x = \frac{\partial \log J}{\partial \mu} \frac{\partial y}{\partial \eta} - \frac{\partial \log J}{\partial \eta} \frac{\partial y}{\partial \mu}$$

$$J\Delta_{\mu,\eta}y = -\frac{\partial \log J}{\partial \mu} \frac{\partial x}{\partial \eta} + \frac{\partial \log J}{\partial \eta} \frac{\partial y}{\partial \mu}$$
(34)

where:

$$J = \frac{\partial_{\mu} x \partial_{\eta} x - \partial_{\eta} x \partial_{\mu} y}{h_{\mu}^2}.$$
(35)

As previously suggested [31, 32], a swelling body is, at equilibrium, equivalent to a compressible structure subjected to an hydrostatic pressure at its free surface. This is illustrated by writing the boundary condition at the free surface in the following form:

$$\mathbf{S}^T \vec{N} = \frac{\mu_0}{\nu} J_F \mathbf{F}^{-T} \vec{N}$$
(36)

where the definition for the nominal stress has been kept:

$$\mathbf{S} = \frac{\partial \mathcal{W}}{\partial \mathbf{F}} \tag{37}$$

but now care must be taken to interpret **S** as the *partial stress* associated with the solid phase of the swelling material. Recalling that $h_{\mu} = \sqrt{\eta^2 + \mu^2}$, it is not difficult to check that the following solution:

$$x = \alpha \eta \mu$$
 $y = \frac{\alpha}{2}(\mu^2 - \eta^2)$ and $J = \alpha^2$ (38)

with

$$\alpha = \frac{1}{\sqrt{1 - \frac{\mu_0}{F_H}}}\tag{39}$$

is a nonlinear symmetric solution of the equilibrium equations and boundary conditions. This solution correspond to an homogeneous swelling process. The one-dimensional swelling ratio α is comprised between 0 and ∞ and is directly controlled by the dimensionless chemical potential

of the solvent particles reservoir $\mu_0/(E\nu)$. Let us now perturb the fields *x*, *y*, and *J* using the following expansion:

$$\begin{cases} x = \alpha \eta \mu + \epsilon \alpha u(\mu, \eta) \\ y = \frac{\alpha}{2} (\mu^2 - \eta^2) + \epsilon \alpha v(\mu, \eta) \\ J = \alpha^2 + \epsilon \alpha^2 j(\mu, \eta). \end{cases}$$
(40)

Inserting the expansions above in the equilibrium equations (34), we obtain the following equations:

$$\Delta u = -\frac{1}{\alpha^2} \left(\eta \frac{\partial j}{\partial \mu} + \mu \frac{\partial j}{\partial \eta} \right) \tag{41}$$

$$\Delta \nu = -\frac{1}{\alpha^2} \left(\mu \frac{\partial j}{\partial \mu} - \eta \frac{\partial j}{\partial \eta} \right) \tag{42}$$

where *j* is given by:

$$j = \frac{1}{\eta^2 + \mu^2} \left\{ \eta \left(\frac{\partial u}{\partial \mu} - \frac{\partial v}{\partial \eta} \right) + \mu \left(\frac{\partial u}{\partial \eta} + \frac{\partial v}{\partial \mu} \right) \right\}.$$
(43)

As previously, let us introduce the complex displacement U = u + Iv and the complex coordinates $\zeta = \mu + I\eta$ to obtain the following equations:

$$\Delta U = -\frac{2I\overline{\zeta}}{\alpha^2} \frac{\partial j}{\partial \zeta} \tag{44}$$

$$j = \frac{I}{\zeta} \frac{\partial \overline{U}}{\partial \zeta} - \frac{I}{\zeta} \frac{\partial U}{\partial \overline{\zeta}}.$$
(45)

The previous equations can immediately be integrated to give, after using the symmetry condition with respect to μ :

$$U(\zeta,\overline{\zeta}) = \frac{1+2\alpha^2}{2+2\alpha^2} \left\{ \frac{F'(\zeta)}{2\zeta} (\overline{\zeta}^2 - \zeta^2) + F(\overline{\zeta}) - F(\zeta) \right\}$$
(46)

where *F* is an arbitrary holomorphic function with appropriate behavior at infinity ($|u| < \eta |\mu|$ and $|v < |\mu^2 - \eta^2|$) that must satisfy the linearized boundary condition (36):

$$\frac{\partial U}{\partial \zeta} + \frac{\partial U}{\partial \overline{\zeta}} + \overline{F}'(\zeta) = 0 \quad \text{for } \overline{\zeta} = \zeta - 2I \tag{47}$$

which cannot be satisfied for periodic holomorphic functions *F*, indicating that the swollen poroelastic tip is also linearly stable.

5. Conclusion

Motivated by several discussions with Yves Couder, we have begun to investigate in this paper the behavior of growing elastic tips. By analogy with the shape of several biological structures but also with dendritic tips, we have chosen to describe growing tips as parabolas. We have derived the Euler–Lagrange equations ruling the shape of growing hyperelastic tips as well as nonlinear poro-elastic swelling tips, as a simple example of stress-modulated growth. In the case of homogeneous growth, as well as at the thermodynamic equilibrium for the swelling case, the parabolic tips were found to be linearly stable with respect to infinitesimal perturbations. Free harmonic modes, which may trigger a roughening transition depending on their dynamic and on the level of noise in the system, were also analyzed. More realistic models, incorporating additional constraints arising from boundary conditions and non-homogeneous growth will be considered in a forthcoming publication. Besides the problem of plant morphogenesis studied here, an understanding of the behavior of growing tips might also contribute to the design of complex elastic structures using swelling gels [32, 33], inflatable structures [34] or electricallyactuated materials [35].

This work, suggested to both of us by Yves, is the outcome of our last conversations. We are convinced that he would have gently pushed us to explore more deeply the leading role of tips in biological growth processes. This is only a first contribution on his physical intuitions which have always combined universality and simplicity.

Conflicts of interest

The authors declare no competing financial interest.

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