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Dynamics of Plant Growth;
A Theory Based on Riemannian Geometry

by

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Abstract

In this work, a new model for macroscopic plant tissue growth based on dynamical Riemannian geometry is presented. We treat 1D and 2D tissues as continuous, deformable, growing geometries for sizes larger than 1mm. The dynamics of the growing tissue are described by a set of coupled tensor equations in non-Euclidean (curved) space. These coupled equations represent a novel feedback mechanism between growth and curvature dynamics.

For 1D growth, numerical simulations are compared to two measures of root growth. First, modular growth along the simulated root shows an elongation zone common to many species of plant roots. Second, the relative elemental growth rate (REGR) calculated in silico exhibits temporal dynamics recently characterized in high-resolution root growth studies but which thus far lack a biological hypothesis to explain them. Namely, the REGR can evolve from a single peak localized near the root tip to a double-peak structure. In our model, this is a direct consequence of considering growth as both a geometric reaction-diffusion process and expansion due to a distributed source of new materials.

In 2D, we study a circularly symmetric growing disk with emergent negative curvatures. These results are compared against thin disk experiments, which are a proxy model for plant leaves. These results also apply to the curvature evolution and the inhomogeneous growth pattern of the *Acetabularia* cap. Lastly, we extend the model to anisotropic disks and predict the growth dynamics for a 2D curved surface which develops an elongated shape with localized ruffling.

Our model also provides several measures of the dynamics of tissue growth. These include the time evolution of the metric and velocity field, which are dynamical variables in the model, as well as expansion, shear and rotation which are deformation tensors characterizing the growth of the tissue. The latter are physically measurable quantities that remain to be fully explored using modern tissue growth imagining techniques.
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Discovery, like surprise, favours the well-prepared mind. Discovery is in its essence a matter of rearranging or transforming evidence in such a way that one is enabled to go beyond the evidence so reassembled to new insights. It may well be that an additional fact or shred of evidence makes this larger transformation possible. But it is often not even dependent on new information.

- J. Bruner
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# List of Symbols, Abbreviations and Nomenclature

| Symbol     | Definition                                                                 |
|------------|---------------------------------------------------------------------------|
| 1D         | One Dimension                                                             |
| 2D         | Two Dimension                                                             |
| Manifold   | A topological space that is locally Euclidean.                            |
| KPZ equation | Kardar, Parisi and Zhang; three authors who described a stochastic model of crystal growth |
| ODE        | Ordinary Differential Equation                                            |
| PDE        | Partial Differential Equation                                             |
| Auxin      | A plant growth hormone.                                                   |
| PIN        | A protein found in plant cells that facilitates the transport of auxin from one cell to another. |
| Primordium | The precursor to a plant organ (leaf, branch, etc.)                       |
| Meristem   | A collection of undifferentiated cells at the tip of a growing shoot or root which generate plant organs. |
| Morphogenesis | The process by which shape develops in a plant or plant organ.           |
| REGR       | Relative elemental growth rate; a standard measure of growth for plant roots. |
Chapter 1

Introduction

Biological systems operate across an astounding breadth of physical scales, from quantum and molecular processes at nanometer scales \[56\] to architectures that grow to tens of meters in size. Over the past 60 years, the focus has been strongly biased toward molecular and cellular levels of understanding biology. However, important exceptions to this trend are the works of D’Arcy Thompson \[71\] and Alan Turing \[74\] who, in different ways, examined mathematical patterns on a whole-system scale.

The goal of the work presented here is to establish a macroscopic theory of plant leaf growth. We start with two very simple observations about leaves and petals. The first is that many types of leaves and petals are not flat; by this we mean these tissues have intrinsic curvature that cannot be laid flat without either tearing the tissue (as in the case of flattening an orange peel), or overlapping of the tissue (as for a ruffled lettuce leaf). The second observation is that leaves and petals grow from millimeter to centimeter or meter scales apparently continuously, both in space and time. As will be seen, these facts lead to remarkable implications for the mathematics and physics of the system. Furthermore, such a model of macroscopic growth leads us to the hypothesis that feedback between growth and curvature is essential to the development of plant structures with highly regular patterns of curvature.

1.1 Biological motivation

Several well studied, though relatively recent, biological arguments exist to motivate a macroscopic and continuum approach to plant leaf development. These are discussed below.
1.1.1 Mechanotransduction in tissues

Mounting evidence supports the idea that physical forces produce a measurable effect at all scales during tissue development. In experimental biology, this effect is known as *mechanotransduction*, which can be summarized as the cascade of reactions a tissue has to a physical force starting at the macroscopic scale down to the cellular, sub-cellular and genetic scales.

A striking example of this is given in [8], where the genetic response of plants to touch has been reviewed. In one study, two groups of the same plant were grown. One group was left to grow without any external mechanical stimuli. The other group was touched twice per day by the experimenter who gently brushed the top of the plants with the palm of their hand. The resulting plants were markedly different; the unstimulated plants grew tall, while the touched plants were significantly shorter. Furthermore, the stunted growth of the touched plants was correlated with the expression of an extended set of TOUCH genes, fully 2.5% of the plant’s genome.

At the sub-cellular level, responses to mechanical stress have also been seen in microtubule arrangement. Microtubules are micrometer long filaments inside the plant cell that, among other functions, help the cell keep its shape. The studies reviewed in [45] indicate that microtubule arrangement within plant cells is dynamically coordinated by the stresses experienced at the tissue scale. The key observation in these studies is that when an external stress is put on a portion of plant tissue, the microtubules within each cell will realign to be ‘parallel to the direction of maximal stress’.

Another candidate for a mechanosensing response can be seen in *Drosophila* (fruit fly) wing disks [33]. When the wing disk reaches maturity, cells will stop growing in unison throughout the disk. The lack of a time-dependent chemical signal (morphogen) and a nearly instantaneous response of all the wing disk cells strongly suggests a mechanically mediated response to growth induced stress.

Although the previous example involves animal cells, it is not unreasonable to think that
a similar process could occur in plants. Indeed, it has been observed in both plant and animal studies that organ size can be independent of the number of cells in the organ [69] [14]. In other words, there is some tissue-scale mechanism than governs growth in addition to cellular level processes. One possibility is that this regulation occurs as a result of mechanical stress.

In all of the above examples, the most important feature of how forces affect development is that an organism’s response to stress is direct and nearly instantaneous. That is to say, the response to a force is not mediated in the same way as other signaling pathways in a plant, namely the relatively slow processes of diffusion or active transport of a signaling molecule from one cell to another. Therefore, physics must play a direct and key role in the development of all kinds of tissues: embryos, organs, leaves, petals, roots. Indeed, biological systems live in a physical world, which means they must have mechanisms to respond and adapt to the physical constraints and stimuli around them. The study of these pathways is an active field of research in both plant and animal systems [26].

One example of this physics-based approach in biological modeling is the work of Green [24] [23]. To understand the patterning of primordia on the meristem (phyllocladus), Green analyses the mechanics of deformable, constrained disk and annulus geometries using a thin-disk approximation. The patterns of deformations, specifically peaks and troughs, allowed by these geometries are strikingly similar to the ones seen on the meristem, suggesting that the mechanics and geometry of plant structures play a key role in morphogenesis.

However, the exact nature of how mechanical signals propagate through tissues is not known. This is because plant tissues can be considered 'soft matter', in the same category as gels and films whose internal structure accommodates dynamics somewhere in between rigid matter and fluids. Indeed, it is the nonlinear viscoelasticity of soft matter that makes characterizing dynamics from first principles so difficult. Often, phenomenological approaches combined with experimental data are needed to understand these types of materials. The study of plant tissues from the perspective of soft matter is only beginning to take shape.
within the scientific community. [18] [15]

1.1.2 Plants as continuous structures

In this section, we aim to motivate the study of plant leaves at scales larger than 1mm as continuous thin sheets using well known facts about plant leaves. More discussion on established work using this paradigm will be found in Sections 1.3.1 and 1.3.2.

First, a simple comparison of the size of cells to the scale of curvature patterning suggests a continuum approximation of the leaf surface is appropriate. In the model plant *Arabidopsis*, leaf cells grow to about 1000µm² in size [19]. The resolution of the curvature patterns we seek to study (in particular, buckling and ruffling) is on the order of 1mm², or 10^6µm² [61]. Therefore, curvature patterns are collections of many cells, which indicates that a continuum approximation is an appropriate tool for analyzing the problem of macroscopic plant growth.

Another argument for a continuum approach is that the patterns of ruffling and buckling seen at the millimeter and larger scale are continuous. The surfaces of even highly curved leaves are locally smooth and connected. This suggests that leaves can be abstracted as differentiable manifolds, allowing for the use of powerful mathematical methods to provide a precise description of the system being studied. In particular, the tools from differential geometry will allow the intrinsic curvature of the tissues to be well understood and incorporated into the growth of the tissue over time.

Another useful fact in constructing our model is that macroscopic leaf growth is dominated by cell expansion, not cell division [25] [2]. Also of note is that cell expansion is not uniform across space or time. This process is not well understood and varies from species to species. Furthermore, plant cells have fixed neighbours and cannot move relative to one another, so each cell is like a point on an expanding geometry, with distances between points growing and deforming over time.

Lastly, cells in plant leaves are highly connected to each other via plasmodesmata. These 'windows' between cells facilitate fast information exchange between cells, including the
transfer of large molecules such as proteins, RNA and DNA. Each cell has many such links to its neighbours. This high interconnectedness suggests that tissue and system scale communication is efficient and evolutionarily favoured.

Indeed, all of these arguments fall in the purview of an organismal theory of plants, which the botanist Julius von Sachs summarizes as 'The plant forms cells, the cells do not form plants.'

1.1.3 Biological processes are feedback loops

In recent years, the view that biological processes are linear, with a clear cause-and-effect behaviour, has been replaced with a better understanding of the complexity of feedback mechanisms. These mechanisms operate at many different scales, and may interact across scales.

Gene regulatory networks are widely known to be complicated feedback loops that regulate gene expression in cells. They are represented visually as a network of mRNA and protein nodes with multiple links, but can also be described mathematically in terms of a set of coupled ordinary differential equations.

When coupled with chemical signaling across the organism, gene regulatory networks can be triggered in a way that supports pattern formation on the system scale. Information about the state of a cell’s neighbouring region can be transmitted by the diffusion of a chemical signal that can then activate a genetic pathway within the cell. The cell can pass on information about its new state using another chemical signal, thereby affecting the development of its neighbours.

A specific example that illustrates this point is the self-organization of PIN transporters in plant meristems. In order to grow, a plant cell must be exposed to the growth hormone auxin. Auxin is transported from cell to cell with the help of the transporter protein PIN. In silico modelization of the meristem shows that if PIN localization within the cell responds to the flux of auxin through the cell, PIN transporters will dynamically align (polarize) to
facilitate more auxin flow to an emerging organ on the meristem. This self-organization also leads to regular patterning of organ primordia on the meristem (phyllotaxy). Hence, the flow of auxin changes the structure of each cell in the meristem to cause a positive feedback loop that supports large-scale tissue growth and patterning.

At the system level, a very important feedback loop was hypothesized by Turing in 1952 [74] and recently confirmed experimentally [72]. Turing’s revolutionary idea was that large scale pattern formation in biological systems is a result of reaction-diffusion processes of chemicals called morphogens. Turing’s original paper was largely concerned with the dynamics of linear chemical reactions on a ring of discrete permeable cells. This model successfully predicted the classes of pattern formation seen in the recent experimental study.

The linear models are, however, only an approximation of the much more complex nonlinear dynamics seen in experiment. Indeed, Turing acknowledged that nonlinear couplings would be more realistic for modeling biological systems, but these would require numerical simulations that surpassed the computing power available at the time. Today, phase diagrams produced from linear and nonlinear models show just how different the dynamics are between the approximate model and a more realistic, nonlinear one [72].

Lastly, of particular interest in the context of forces and feedback loops, is recent experimental evidence that mechanotransduction pathways are evolutionarily ancient and shared across what are now distantly related species. The same mechanical mechanism for mesoderm specification has been shown to occur in both zebrafish and Drosophila, which last shared a common ancestor some 570 million years ago [9]. What mechanotransduction pathways exist in plants remains an open question [46] [51].

1.1.4 Patterning as a reaction-diffusion process

Reaction-diffusion dynamics occur in many natural systems including those in chemistry and biology. The resulting phenomena are identified as having, among other behaviours, self-organizing spatial and temporal patterns emerging from a random initial state. Mathe-
matically, the system’s dynamics must have both a diffusive and reactive term:

\[ \partial_t u = D \nabla^2 u + R(u), \]

where \( u \) is a \( N \)-dimensional dynamical variable in space and time, \( D \) is a matrix of diffusion coefficients and \( R \) is a real vector functional of \( u \) that may have many terms including nonlinearities and differential operators. We define Equation 1.1 to be quasi-linear if \( D = D(u) \) and \( R = R(u, \nabla u) \).

Turing’s work on morphogens falls into this class of equations, as do many studies of pattern formation in biology \[42\]. Models for coupling reaction-diffusion chemical dynamics with growth patterns on the plant meristem and other tip growth structures have been studied by Harrison and Holloway \[27\]. In the context of plant morphogenesis, the notion of reaction-diffusion dynamics is particularly relevant because of the highly regular patterns of curvature seen in many leaves and petals. Indeed, the structure of reaction-diffusion equations will guide the development of a set of equations describing plant growth (Chapter 2).

1.2 A note on modeling in biology

Biological disciplines today are dominated by observation, with theory playing a rather secondary role. In contrast, physics tends to place a more balanced emphasis on theory and experiment, so that the two drive each other to an ever more rigorous understanding of the world around us.

Often observations of biological systems are overwhelmingly rich and complicated, reflecting the inherent complexity of life. The value of constructing a model for such a system is therefore to focus our attention on the salient features of a certain problem. When we construct a framework that simplifies the system, we can then test the assumptions of that framework in the form of predictions.
Modeling a biological system can help guide experiments to be performed on a complicated system. Rather than starting with all the details in the hopes of sifting out the important parts, we can hypothesize what those important parts may be and then identify key processes occurring in vastly more complicated biological system.

A hugely successful example of this is Turing’s model of morphogens [74]. By postulating the existence of chemical messengers and how they interact, he constructed a theory that explained how large-scale patterns arise in biological systems. It was only decades later that his theory was fully verified by experiment [72].

In the model of plant growth presented here, we make the simple observations that plant leaves grow and exhibit intrinsic curvature (recall how neither an orange peel nor a lettuce leaf can be laid flat without tearing or folding). To understand these shapes in a mathematically rigorous way requires certain tools from differential geometry. As it happens, these are the same tools used to understand the nature of curved spacetime first described by Einstein in his theory of general relativity. Because of this, there will be parallels between the two systems - systems that are both geometric and dynamic. This was a huge shift in perspective for the science of cosmology a century ago, and hopefully dynamic geometry will shed some new light on plant growth as well.

1.3 Established work

1.3.1 Plant growth as a continuous process in space and time

One of the earliest works on treating growing tissues as continuous media was done by Silk and Erickson in 1979 [66]. In it they argue for the use of continuum mechanics in studying plant growth, noting that certain structures such as the meristem are defined by their geometry and position on the plant, rather than a fixed set of cells. They also use the fact that growth curves are time-continuous to motivate both temporal and spatial continuity at millimeter and larger length scales.
1.3.2 Material transport causes leaf growth

A very simplified but sufficient view of how leaf development occurs at scales larger than 1mm is that material from the stem is transported to the leaf, which enables cells in the leaf to expand. Material is also created locally in cells by transforming sunlight into carbohydrates (among other local processes) and can be transported to neighbouring cells in a variety of ways. In the continuum limit, this is enough to allow material transport in the leaf to be expressed as a conservation of mass equation with an associated source term.

Continuous material transport in the leaf also allows us to use the equations of viscous fluid flow to describe the velocity field of a growing leaf [66]. Just as the boundary of the leaf expands with some velocity, so too does every point inside the boundary. This is the velocity field of the leaf. The velocity field can be thought of as having two contributing effects: one is the diffusion of material between cells through a number of molecular transport pathways, and the other is the how material flow is altered by the expansion of the tissue itself.

1.3.3 Stress, strain and growth are tensor quantities

As early as 1984, Hejnowicz and Romberger discussed how tensors are the only mathematical objects to fully describe the physical deformations experienced by a growing tissue: expansion, rotation and shear [30] (see Figure 1.1). In addition, geometric effects such as anisotropy and inhomogeneity can also be considered as results of growth.

Indeed, tensors are used to describe the properties of both physical objects and geometry. Tensors can be as simple as a single number, like the temperature at a specific point in a room. These are zero-rank tensors which are also known as scalars. A tensor with information about magnitude and a single direction is a first-rank tensor, or vector. Higher rank tensors can express physical quantities with multi-directional properties like stress, strain and growth or geometric quantities like curvature. They are a very powerful way of encoding information about a set of points in a multidimensional space and are used in disciplines like fluid
mechanics, elasticity theory and general relativity.

One example where tensors have been used to better understand a biological system is given in [47]. Here, a time-independent growth tensor in curvilinear coordinates [29] is used to model the tip of a radish root. This information is then used to generate the principle directions of growth at each cell, which dictate how the cells will divide and grow. While it was long postulated that cells divide along principle directions of growth, it was unclear how the cells coordinate these directions [44]. By using a growth tensor, this model is able to build an algorithm that links a tissue-level property with cellular division and growth.

Another tensor based approach to biological growth is to consider elastic (reversible) deformations due to stress as separate from the plastic (irreversible) deformations of growth itself [53]. Both types of deformations are described using tensor quantities, but have very different time scales over which they act, allowing a multiplicative decomposition of the two effects. This approach has been successfully applied to simple growth modes such as constant isotropic growth, but the computational complexity of this algorithm for more realistic scenarios has prevented these models from developing further [35].

Figure 1.1: Some examples of rank-two tensor quantities.
1.3.4 Curved surfaces are a result of nonhomogeneous growth

More recently, there is also strong empirical evidence showing that nonhomogeneous growth leads to curvature.

In [41], a thin disk of gel is treated with a heat-sensitive monomer whose concentration varies with radius. When heat is applied to the disk, the disk contracts proportionally to how much monomer is present at a given radius. Thus, the disk can be manipulated to contract more along the edges than at the center, causing a dome shape, or the contraction can occur more in the center than at the edges, causing ruffles along the edges. The amount of contraction is a physical representation of the geometry of the disk, and thus the link between nonhomogeneous growth and intrinsic curvature can be studied. These results will play a key role in interpreting the 2D geometries of the model developed here (Chapter 5).

Other experiments on torn plastic sheets show similar results. When a plastic sheet is put under tension and then torn in the direction perpendicular to the direction of tension, the result is ruffled edges along the tear. If one then measures lengths along different directions on the sheet, it can be seen that the torn (ruffled) edge is longer than the flat edge [60]. Moreover, the ruffles form a fractal cascade over several orders of magnitude [61] [63] which can be understood in terms of minimizing the bending energy of the long edge under geometrical and elastic constraints [3]. A similar analysis for a disk geometry where anisotropic growth results in curvature is discussed in [16]. Intuitively, these models show that buckling will always occur in systems where there is an excess of material along one edge that, for geometric reasons, cannot spread out to be flat. For example, a lettuce leaf and a ballerina’s skirt both have this property; if one tries to flatten a portion of the leaf or skirt, the rest becomes more buckled. The excess material along the outer edges cannot relax to a flat geometry, so it folds up in a way that distributes the curvature as evenly as possible across the whole surface. How the buckling manifests itself, whether through short, long or fractal buckling, depends on the material itself, in particular its elasticity and thickness. Given two
sheets of equal thickness and equal excess edge length, but different elasticity (for example, one made of tissue paper and the other of cardboard), the more elastic one will be easier to bend, and so produce more short-wavelength, low amplitude buckling than the less elastic one. Similarly, the thickness of the sheet will influence how easy it is to bend the material; a thicker sheet is more difficult to bend, so it will tend to have more long-wavelength buckling with higher amplitudes.

The idea behind these experiments is to correlate the final shape of a manipulated system (thin sheets) to the shapes found in nature, in particular plant leaves and petals. Indeed, based on these experiments, it is obvious that intrinsic curvature is the result of nonhomogeneous growth processes.

Nonhomogeneous growth has been studied from a genetic perspective [11] [54] [48], and models for how genetically programmed growth rates affect tissue shape and curvature have been developed [40] [43]. These authors postulate that growth processes and the stresses they produce could influence the genetic expression of the tissue through mechanotransduction, however no model has incorporated this feedback loop yet. Additionally, these models of genetic control of curvature do not take into account how curvature effects react back onto the growth mechanism.

A relevant model organism for studying the development of curvature is the single-celled algae *Acetabularia*. Despite being single-celled, it grows to centimeter length scales and has a cap that changes its curvature as it grows (See Figure 1.2). Tracking the growth of the cap has also revealed that the cap grows mostly in an annular region around the main stalk, indicating highly nonhomogeneous growth processes at work [59]. These characteristics will be revisited in Chapter 5 as we seek to understand the dynamics of our simulated growth system.
1.4 What is needed for a better model?

The elements of continuity, material transport and nonhomogeneous growth are all important first steps in understanding plant growth dynamics, but are incomplete without considering the physical effects of curved geometries. Silk and Erickson consider only flat geometries for their continuum model, which we show omits very important dynamics found only in systems with curvature.

The idea of using tensors to describe the geometry, growth, velocity field and scalar density of the plant leaf is fully embraced in our model, and as will be seen in Chapter 2 provides the basis of a self-consistent physical model of plant growth. However, rather than being merely descriptive, tensor quantities in our model will be time dependent and evolve according to the structure of the dynamical equations of plant growth.
The thin-sheet experiments correlate the final shape of a plastic sheet with the shapes found in plant leaves and petals, but do not encompass the growth process itself. Indeed, models developed to describe thin sheet experiments do not address the feedback that a changing geometry imposes on the dynamics of the physical processes creating the curvature. In our model, this two-way feedback between growth and geometry is explicit and results in predictions on the large scale growth dynamics of plant tissues.

1.5 Chapter summary:

Geometry as the missing link

As seen in this chapter, there is strong evidence from both biology and thin-sheet physics that tissue curvature in plant leaves and petals is a rich interplay of many factors, as depicted in Figure 1.3. To model this curved, continuous system requires specialized tools in mathematics and physics, which will be discussed in Chapter 2. The numerical methods used to study the equations of plant growth will be discussed in Chapter 3 and results for 1D and 2D models will be presented in Chapters 4, 5, 6. Finally the implications of the numerical simulations and future directions of the work will be considered in Chapter 7.
Figure 1.3: A conceptual representation of the ingredients for a physical model of plant leaf growth.
Chapter 2

Developing a Physical Model

In this chapter, the physics and mathematics of our model are developed. Each assumption made about the physics of the plant leaf system provides a guide to the appropriate mathematical tools needed to study it. Certain crucial facts about the system also aid in this process. The goal is to create a minimal set of assumptions which will then generate a realistic model of leaf growth dynamics.

2.1 Tensors in physical systems

A fundamental property of tensors is that the quantities they relate through equations are coordinate independent, and they can be transformed between coordinate systems [17]. This is an important feature when constructing any physical theory. No matter what coordinates the system ‘lives in’, the relationships among the tensor quantities must remain the same. For the plant leaf model, this allows us the freedom to write down the tensor equations representing the physics of the system without worrying about how to represent the tensor components in the chosen space.

To illustrate the point, we can think of tracking the growth of the leaf by drawing small dots on the surface and measuring how each dot moves over time. The distance between dots tells us about the geometry of the leaf. The velocity of each dot (both speed and direction) gives information about how the leaf is growing. Clearly, where we draw the dots does not impact the dynamics of the leaf. The dots are merely a coordinate system that helps us observe growth. There may be very convenient arrangements of dots that immediately bring to light key aspects of the physics occurring in the leaf. Other coordinates may obscure our understanding.
A corollary of the fact that tensors can be transformed between coordinate systems is that tensor equations must equate only tensors of equal valence, and must have the same units. The valence \((p, q)\) denotes how many contravariant indices \(p\) and covariant indices \(q\) a tensor has. For example, the tensor \(A^i_j\) is a rank 2 tensor with valence \((1, 1)\); \(A_{ij}\) has valence \((0, 2)\); \(A^{ij}\) has valence \((2, 0)\).

Transforming tensors between coordinate systems follows well defined rules. The simplest transformation is for scalar (rank 0) quantities, which stay the same in different coordinate systems:

\[
\phi'(x'^1, x'^2, \ldots, x'^n) = \phi(x^1, x^2, \ldots, x^n). \tag{2.1}
\]

In an \(n\)-dimensional differentiable manifold, we can define a contravariant vector quantity \(A^i\) in the \(x^i\) coordinates \((i = 1, 2, \ldots, n)\). If we transform from the \(x^i\) coordinates to another coordinate system, say \(x'^i\), then the tensor \(A^i\) also transforms to \(A'^i\):

\[
A'^i = \sum_{j=1}^{n} \frac{\partial x'^i}{\partial x^j} A^j. \tag{2.2}
\]

The \(\partial x'^i/\partial x^j\) term forms the \(n \times n\) transformation matrix, or Jacobian that takes the \(x^i\) coordinates to the \(x'^i\) coordinates.

In practice, Einstein summation notation is used to reduce the clutter in such equations. Any repeated index is understood to indicate a sum over \(n\):

\[
A'^i = \frac{\partial x'^i}{\partial x^j} A^j = \frac{\partial x'^i}{\partial x^b} A^b. \tag{2.3}
\]

Higher rank tensors can be constructed out of covariant and contravariant tensors, and transformation rules can be written down for any tensor. Some relevant examples include:

Covariant tensor transformations:

\[
A'_i = \frac{\partial x'^i}{\partial x^i} A_j. \tag{2.4}
\]
Second rank tensor transformations:

\[ A'^{ij} = \frac{\partial x'^i}{\partial x^k} \frac{\partial x'^j}{\partial x^l} A^{kl}. \]  

\[ A'^{ij} = \frac{\partial x'^k}{\partial x^i} \frac{\partial x'^l}{\partial x^j} A_{kl}. \]  

Mixed rank tensor transformations:

\[ A'^i_j = \frac{\partial x'^k}{\partial x^i} \frac{\partial x'^j}{\partial x^l} A^l_k. \]  

One mathematical aspect that becomes very important is how to do differential calculus using tensor quantities. Let us first take a look at the partial derivative of a contravariant tensor in two different coordinate systems:

\[
\frac{\partial A'^i}{\partial x'^k} = \frac{\partial}{\partial x^k} \left( \frac{\partial x'^i}{\partial x^j} A^j \right) \\
= \frac{\partial x'^i}{\partial x^k} \frac{\partial}{\partial x^l} \left( \frac{\partial x'^i}{\partial x^j} A^j \right) \\
= \frac{\partial x'^i}{\partial x^k} \frac{\partial x'^j}{\partial x^l} \frac{\partial A^j}{\partial x^l} + \frac{\partial^2 x'^i}{\partial x^j \partial x^k} \frac{\partial x'^l}{\partial x^i} A^j 
\]

The standard transformation rule that worked before now yields a non-tensorial quantity, namely the second term on the right hand side of the equation. This presents a problem since partial derivatives form the basis of more complex differential operators, such as the gradient, divergence, etc. If partial derivatives of tensors cannot transform linearly between coordinate systems, then physics would act differently in different coordinate systems.

To remedy the situation, we can introduce a set of terms that compensate for the non-tensorial behaviour of partial derivatives in different coordinate systems called the *connection coefficients*, which are denoted by \( \Gamma^i_{jk} \). This allows for covariant derivatives to transform appropriately between coordinate systems. The details of this are beyond the scope of this...
short review, but can be found in [17]. We simply state that by using the connection coefficients, we can then define the \textit{covariant derivative}:

\begin{equation}
\nabla_k A^i = \partial_k A^i + \Gamma^i_{jk} A^j
\end{equation}

where $\partial_k A^i$ is short hand for $\partial A^i / \partial x^k$.

As will be seen in the next section, the connection coefficients are directly linked to geometry of the space in which the tensor exists.

Just as different tensors have different transformation rules, covariant derivatives act differently on different tensors.

For a scalar:

\begin{equation}
\nabla_k \phi = \partial_k \phi.
\end{equation}

For a covariant vector:

\begin{equation}
\nabla_k A_i = \partial_k A_i - \Gamma^j_{ik} A_j.
\end{equation}

2.2 Physical tensors: expansion, rotation and shear

As a first example of how tensors can describe physical quantities of a system, we can look at three tensors that are often used in continuum dynamics to describe transport phenomena: expansion, shear and rotation. Indeed, these formalize the notions introduced in Section [1.3.3] and will be useful later on when analyzing the stresses and deformations of simulated tissue growth. Also, tensor equations for these quantities are needed so that dynamical laws governing the physics of deformable systems remain coordinate independent.

A viscoelastic fluid flow can be described with a velocity field $v^a$ that varies over space and time. The same velocity field can be used to construct the expansion scalar which is the covariant divergence of the velocity field:

\begin{equation}
\Theta = \nabla_a v^a,
\end{equation}
the shear tensor:

\[ \sigma_{ab} = \frac{1}{2}(\nabla_a v_b + \nabla_b v_a), \]  \hspace{1cm} (2.13)

and the rotation tensor:

\[ \omega_{ab} = \frac{1}{2}(\nabla_a v_b - \nabla_b v_a). \]  \hspace{1cm} (2.14)

In each of these equations, \( \nabla_a \) is the covariant derivative as presented in the previous section.

Intuitively, these deformations are summarized in Figure 1.1. The expansion scalar measures the rate of relative increase in scale, the shear tensor measures an area preserving deformation of shape change when forces act to skew the shape of the element, and the rotation tensor measures how much an element is rotated during deformation.

In viscoelastic systems, a continuous medium is described using fluid elements. As the medium is deformed over time, shape of the fluid elements change as well, which can in turn be described using the velocity field of the medium. A full derivation of these equations can be found in \[67\].

2.3 Geometric tensors

We now move on to the introduce the roles of tensors in curved spaces, and hence their roles in plant tissues. Based on the discussion in Section 1.1, plant leaves can be viewed as continuous, curved and growing 2D surfaces. An appropriate mathematical tool to describe such a system is Riemannian geometry. A thorough treatment of this branch of mathematics is beyond the scope of this discussion, but the most salient points are presented here. Further details can be found in \[17\].

The fundamental object in Riemannian geometry is the metric tensor. The metric tensor encodes all the geometric information about a space. From it, basic quantities like the
distance between two points or the curvature at a given point can be calculated. We define
the geometry of a leaf’s surface using a 2D metric tensor.

In an \( n \)-dimensional geometry, we need \( n \) coordinates to describe any given point. This
can be written as \( x^i = (x^1, x^2, ..., x^n) \). If we wish to calculate the distance \( s \) between two
points in the geometry, we can do so by generalizing the Pythagorean theorem \( ds^2 = dx^2 + dy^2 \)
by defining an infinitesimal distance \( ds \) that is a function of the coordinate displacement \( dx^i \)
and the metric tensor \( g_{ik} \):

\[
ds^2 = g_{ik} dx^i dx^k
\]  \hspace{1cm} (2.15)

where \( g_{ik} \) is an \( n \times n \) tensor. Again, Einstein summation notation is used to denote a sum
over repeated indices.

Similarly, the metric defines an inner product between vectors or components of higher
rank tensors:

\[
C = g_{ik} A^i B^k.  \hspace{1cm} (2.16)
\]

It must be remembered that, in general, the coordinates themselves do not determine
distances. For example, in a plane polar coordinate system, the angular coordinate \( \theta \) is used
to distinguish between points on a circle of radius \( R \). The distance between \( \theta_2 \) and \( \theta_1 \) along
the circle is given by \( \Delta s = R \Delta \theta \) where \( \Delta \theta = \theta_2 - \theta_1 \). The factor multiplying the difference
in the coordinates is represented by a component of the metric tensor.

Extending this example to a flat circular geometry, we choose \( x^1 = r \) and \( x^2 = \theta \). The
distance between two infinitesimally separated points is then:

\[
ds^2 = dr^2 + r^2 d\theta^2.  \hspace{1cm} (2.17)
\]

The corresponding metric tensor for a flat 2D circular geometry in \((r, \theta)\) coordinates is
given by \( g_{11} = g_{rr} = 1, g_{12} = g_{21} = g_{r\theta} = 0, g_{22} = g_{\theta\theta} = r^2 \). This can also be written as a
2-dimensional array:

\[
g_{ik} = \begin{pmatrix}
1 & 0 \\
0 & r^2
\end{pmatrix}.
\] (2.18)

For non-flat 2D geometries such as those describing plant leaves, the metric tensor is more general and may involve arbitrary coordinates \((u, v)\):

\[
g_{ik} = \begin{pmatrix}
f(u,v) & h(u,v) \\
h(u,v) & g(u,v)
\end{pmatrix}.
\] (2.19)

In this case, the distance between two infinitesimally separated points is:

\[
ds^2 = f \, du^2 + 2h \, du \, dv + g \, dv^2.
\] (2.20)

Another interesting issue regarding the representation of curved surfaces the use of intrinsic rather than extrinsic geometry. The leaves we see around us are all comfortably embedded in flat 3D space. However, under our assumptions of the leaf as a thin sheet, the dynamics of the system only require a 2D space. If we study the system in a 3D space external to the leaf, we have chosen an extrinsic (Eulerian) approach. If we choose a 2D space defined on the leaf itself, such as dots drawn on leaf, then we have chosen an intrinsic (Lagrangian) approach.

Two arguments exist to support the intrinsic approach. First, Silk and Erickson noted that patterns of curvature are stationary in intrinsic coordinates [66], suggesting that the plant uses a local reference frame to guide its growth.

The second argument follows from the fact that curvature leads to forces (more details on this in the next section) and is an intrinsic property of a geometry. With clear evidence that plants respond to forces (Section 1.1), our hypothesis is that plants must also be able to sense forces arising from curvature. These could conceivably be sensed locally by each cell, primarily because curvature contributes to expansion and shear in a 2D manifold [36].
Because these are properties intrinsic to the space itself (i.e. they are independent of coordinate choice), a 2D coordinate system is sufficient to describe the physics associated with a curved leaf geometry.

The role of extrinsic coordinates, however, cannot be altogether discounted. External forces, including bending, undoubtedly affect plant tissue. Furthermore, embedding a curved 2D geometry in a flat 3D space provides a sort of symmetry breaking for the curved system, causing buckling cascades that minimize the elastic and bending energies of the tissue [3][62].

Since we are presently concerned with the effect of internal forces generated by nonhomogeneous growth of leaf tissue, we proceed with using a 2D metric and a 2D velocity field defined on the surface of the leaf, with curvature defined intrinsically on the surface of the manifold.

2.4 Curvature leads to forces

One of the great discoveries of 20th century science was that gravity is really an effect of curved spacetime. Indeed, general relativity uses the same concepts of Riemannian geometry, tensors, continuity and differential geometry to formalize how curvature affects motion at the largest scales.

But first, let us return to Newton’s Second Law:

\[ F = m a = m \frac{d\mathbf{v}}{dt} = m \frac{d^2\mathbf{x}}{dt^2}. \] (2.21)

Again note that a force is related to a change in an object’s velocity vector. To show how curvature leads to forces, let us look at two illustrative examples.

The first example is of two ants traveling on a surface. The ants walk in straight lines, which is locally defined as taking steps of equal length on the right and left with each stride forward [1]. Figure [2.1] shows three cases of such walks, also known as geodesics.
Figure 2.1: From left to right, geometries with flat, positive and negative curvatures and their corresponding 'straight' paths.

What is happening? The three shapes represent three possible geometries, as defined by Gauss’ *Theorema Egregium*. The flat disk can be defined by the fact that it’s circumference $C = 2\pi r$. The middle shape is defined as a positively curved surface in which there is a deficit in circumference so that $C < 2\pi r$ (the yellow area is removed from the flat disk). The last shape is a negatively curved surface in which there is an excess of circumference so that $C > 2\pi r$ (the red area is added to the flat disk). Because area measurements can be done without leaving the surface of the manifold, the curvature calculated this way is an intrinsic property of the space. There are also extrinsic measures of curvature, such as mean curvature, but these are not considered here.

In the flat geometry, two ants that start walking on parallel lines will remain the same distance from each other forever. On the positively curved geometry, the ants appear to attract each other. On the negatively curved geometry, they appear to repel each other. In each case, the ants still walk in locally straight lines (the black line is never redrawn), but the velocity vector of the ants is clearly altered by geometry. Moreover, this change in velocity will be interpreted by the ants as a force, just as we interpret curvature as the force of gravity. Again, these perceived forces are an intrinsic part of the curved manifold.

In relativity, this effect is formalized in the notion of geodesic deviation, in which the separation of infinitesimally close geodesics is proportional to the curvature of the space itself. It is also related to the phenomenon of gravitational lensing, one of the first predictions of
general relativity to be observed experimentally.

In biology, this effect could in principle impact any phenomenon that is described with a vector field. In particular, in Section [2.10] we will consider how the flow of materials in a curved surface is altered through geometry by incorporating the geodesic flow of matter in a curved space into the velocity field of the growing leaf.

Let us look at a second example of how curvature can lead to force. Now assume the ants carry little cubes that always have a face resting on the ground. Observe what happens to the cubes in Figure 2.2. In the flat geometry, we could swap the two cubes and they would still lie flat on the ground beneath them. But in the positive and negative curvature spaces, this cannot be done without rotating the cubes to make them perpendicular to the ground again. Locally, the ants obey the rule of keeping the cubes flat on the ground, but the intrinsic curvature of the space says that the orientation of the cubes changes from point to point.

This effect is formalized in differential geometry through connection coefficients $\Gamma^i_{jk}$. Connection coefficients quantify how much a vector or tensor changes when evaluated at different points on the manifold. In the previous example, the cubes represent vector or tensor quantities. Next, we show how connection coefficients are related to the metric of a curved space.

Given a curved space with metric $g_{ik}$, one can ask what is the natural motion of a particle in that space. In other words, what are the minimal paths governing the motion of particles
in a curved space?

Following Chapters 6.9 and 7.6 in [17], this problem can be solved using a Lagrangian functional of the form

\[ L = \sqrt{g_{ik}\dot{x}^i\dot{x}^k} \]  
(2.22)

and calculating the variation of

\[ \int_{P_1}^{P_2} L \, du = \int_{P_1}^{P_2} ds = s \]  
(2.23)

where \( u \) is a parameter along a smooth curve such that \( \dot{x}^i = dx^i/du \), and \( s \) is the interval between points \( P_1 \) and \( P_2 \) where, as before, \( ds^2 = g_{ik}dx^i dx^k \).

Solving the Euler-Lagrange equations under the condition that \( u = s \) gives us a second order differential equation for the extremal paths of particles traveling in a curved space, also called geodesics:

\[ \ddot{x}^i + \Gamma^i_{jk}\dot{x}^j\dot{x}^k = 0. \]  
(2.24)

The connection coefficients are the result of partial derivatives of the metric that appear in the Euler-Lagrange equations:

\[ \Gamma^i_{jk} = \frac{1}{2}g^{it}\left( \frac{\partial g_{jl}}{\partial x^k} + \frac{\partial g_{lk}}{\partial x^j} - \frac{\partial g_{jk}}{\partial x^l} \right). \]  
(2.25)

Hence, the connection coefficients are an intrinsic part of motion in a curved space. We will revisit this notion when considering transport of material in a curved and growing tissue later in this chapter.

Another property of this particular form of the connection coefficients is that \( \nabla_k g_{ik} = 0 \). Connection coefficients created from the Riemannian metric are often called Christoffel symbols or metric connections.
Recall from Section 2.1 that partial derivatives of tensors do not transform linearly between coordinate systems. A corollary of this statement is that tensor quantities are affected by their location on a manifold, since moving to a different point on the manifold is mathematically similar to a coordinate transformation. Now, instead of a coordinate transformation, we consider the spatial variation of a tensor $A^i$ in curved space. The connection coefficients act like the corrective term introduced before in the covariant derivative:

$$\nabla_k A^i = \partial_k A^i + \Gamma^i_{jk} A^j.$$  \hspace{1cm} (2.26)

Covariant derivatives introduce curvature effects that are not possible in a flat system, but that are crucial to a correct mathematical understanding of dynamics in a curved space. As will be seen, the covariant derivative will play an important role in modifying the continuity and transport equations.

Another powerful feature of Riemannian geometry is the ability to quantify curvature locally. In the research program started by Gauss and completed by his student Riemann in the mid-19th century, the search for methods to measure curvature without leaving the curved space itself were pursued in earnest. The result was a generalization of Euclidean geometry, where basic notions of distance, parallel lines and the circumference of a circle had to be redefined.

Not too long after, at the beginning of the 20th century, Einstein showed that curved geometries can produce the force of gravity. As we have seen, forces are described by the local effects they have on an object’s momentum or velocity. Curvature can be described locally as a property of the metric tensor. One such measure of curvature is the Riemann tensor:

$$R^a_{bcd} = \partial_c \Gamma^a_{bd} - \partial_d \Gamma^a_{bc} + \Gamma^e_{bd} \Gamma^a_{ec} - \Gamma^e_{bc} \Gamma^a_{ed}.$$  \hspace{1cm} (2.27)

A flat geometry is defined as one in which $R^a_{bcd}$ is identically zero.
The Riemann tensor also acts as a measure of non-commutativity of the covariant derivative:

\[ \nabla_a \nabla_b A_c - \nabla_b \nabla_a A_c = R^d_{\ abc} A_d. \]  \hfill (2.28)

Note that if we substitute in the expression for the connection coefficients from Equation 2.25, the Riemann tensor can be seen as a nonlinear function of the metric tensor, composed of second order derivatives and nonlinear first order derivatives of \( g_{ik} \).

Two other curvature tensors exist as well. The Ricci tensor:

\[ R_{ab} = R^c_{\ acb} = g^{cd} R_{acbd} = \partial_c \Gamma^c_{\ ab} - \partial_b \Gamma^c_{\ ac} + \Gamma^d_{\ ab} \Gamma^c_{\ dc} - \Gamma^d_{\ ac} \Gamma^c_{\ db}. \]  \hfill (2.29)

and the Ricci scalar:

\[ R = g^{ab} R_{ab}. \]  \hfill (2.30)

With a mathematical description of curvature in hand, we can now look at its effect on motion through geodesic deviation. As mentioned earlier in the section, curvature directly impacts the geodesics of a space. If we imagine a vector \( \eta^i \) that quantifies the separation between two geodesics, then the equation of geodesic deviation is:

\[ \frac{D^2 \eta^i}{D\tau^2} = -R^i_{\ jkl} v^j v^k \eta^l \]  \hfill (2.31)

where \( \tau \) is the proper time along the geodesics and \( v^i = dx^i/d\tau \). The consequence of this equation is exactly what was seen in Figure 2.1, where a shortest path between two points becomes warped by the geometry which the path must traverse.

In addition to quantifying the effect of curvature of a static geometry, dynamical systems can be constructed where the curvature changes over time. Indeed, general relativity is one of those systems. In it, matter and energy bend spacetime, while the geometry of spacetime dictates how matter and energy move. This is the essence of Einstein’s field equations for
general relativity:

\[ R_{ab} - \frac{1}{2} g_{ab} R = \frac{8\pi G}{c^4} T_{ab} \tag{2.32} \]

where \( T_{ab} \) is the energy-momentum tensor. This equation defines a tensor relationship for a 4D pseudo-Riemannian spacetime. The equations are hyperbolic in structure, which allows for wave dynamics and leads to the prediction of gravitational waves.

More abstract curvature flows can also be constructed. Another important though less well known dynamical equation is the Ricci flow:

\[ \partial_t g_{ab} = -\kappa R_{ab} \tag{2.33} \]

where \( \kappa \) is a real-valued constant and the metric is purely Riemannian. For pure Ricci flow as developed by Hamilton, \( \kappa = 2 \) in order to normalize the system such that a factor of \( 1/2 \) in the connection coefficients is removed.

The Ricci flow was developed by Hamilton in the 1980s in an attempt to solve the remaining parts of the Poincaré conjecture. Later, in the early 2000s, Perelman completed the work, thereby solving a 100-year old problem in geometry and winning a Fields medal for the solution. What the equation does is dissipate the curvature of a metric in a diffusive-like way, allowing a highly convoluted geometry to be gradually deformed to a smoothed geometry, allowing direct analysis of the topology of a space.  

\[ \text{[73]} \]

A common feature of curvature flows is their quasi-linear partial differential structure. This makes solving them analytically very difficult even in the simplest scenarios, and often requires numerical techniques to tackle anything more than toy models.

What do curvature and Riemannian geometry mean in the biological context? As discussed previously, stress, strain and growth are all tensor quantities. Therefore, these quantities will be affected by the local curvature of space. For example, a stress applied locally to a portion of negatively curved tissue will register differently than the same stress applied
to a portion of positively curved tissue. Since cells react to forces through mechanotransduction, it is reasonable to postulate that curvature will modulate a cell’s response to any applied force. A similar argument can be made for the perception of strain and growth in curved spaces. Indeed, such curvature effects were not considered by Silk and Erickson, who proposed that the growth tensor is a gradient of the velocity field in a flat geometry. Petals and leaves are, however, intrinsically curved and their curvature changes over time.

As we have seen, an important property of Riemannian geometry is the ability to describe calculus in curved spaces. This may seem like an abstract mathematical point, but it has profound implications on physical structures like plant leaves as well as more abstract notions like spacetime. This is because calculus provides a local description of physical forces, so anything that may change how those local interactions behave, such as a non-flat geometry, directly affects how the system experiences forces.

For our theory of leaf growth, the implication of considering curvature is that in addition to tensors representing strain, growth, and material transport, we must also include terms that reflect how the system is affected by curvature.

2.5 The physics of material transport and growth models

2.5.1 The continuity equation

As discussed in the previous chapter, Silk and Erickson argued that a continuum approach is appropriate when studying large-scale plant growth [66]. Hence, a relevant equation to study is that of mass continuity with a source term in 2D:

\[
\frac{\partial \rho}{\partial t} + \nabla \cdot (\rho \mathbf{v}) = S(x, t). \tag{2.34}
\]

In this equation, \(\rho = \rho(x, t)\) is the mass density of the material, \(t\) is time, \(x\) is the position vector, \(S\) is the source function, and \(\mathbf{v}\) is the velocity field of the material. The term \(\rho \mathbf{v}\) is also known as the mass current \(\mathbf{j}\).
Equation 2.34 is sufficient in flat geometries, but as seen in the previous sections, differential operators such as the gradient $\nabla$ are influenced by geometry if the space is curved. The curved-space version of this equation can be written in tensor notation, and indeed all the terms including the gradient can be generalized to their curvilinear equivalents [52]:

$$\frac{\partial \rho}{\partial t} + \nabla_i(\rho v^i) = S(x^i, t).$$

(2.35)

The major difference between the two notations is that now $\nabla_i$ is a covariant derivative rather than a partial derivative.

Intuitively, the continuity equation states that the density of the material at a point $x$ depends on the creation or destruction of material in a source or sink, and the flux of material into or out of a portion of space. The significance of this equation in a growing medium will be revisited in Section 2.9.

2.5.2 The transport equation

Material transport in a viscoelastic system can also be understood from the perspective of Newton’s Second Law, which gives a relation between forces and inertia:

$$\mathbf{F} = ma = m \frac{dv}{dt} = m \frac{d^2x}{dt^2}.$$  

(2.36)

Motion, therefore, is a local phenomenon formalized using the methods of calculus. Here, infinitesimal time and spatial changes govern how an object moves. A change in an object’s velocity or momentum vector is proportional to a force. This notion is just as applicable for macroscopic objects as it is for a fluid subdivided into infinitesimally small units of volume.

The ordinary derivative with respect to time in Equation 2.36 can be rewritten in terms of partial derivatives using the chain rule:
\[
\frac{dv}{dt} = \frac{\partial v}{\partial t} + \sum_{i=1}^{n} \frac{dx^i}{dt} \frac{\partial v}{\partial x^i}
= \frac{\partial v}{\partial t} + v \cdot \nabla v.
\]

(2.37)

Hence, the time evolution of the velocity field can be understood in terms of the forces per unit mass \( f = F/m \) and the advection \( v \cdot \nabla v \):

\[
\frac{\partial v}{\partial t} = -v \cdot \nabla v + f
\]

(2.38)

where the mass \( m \) can be normalized to unity assuming a constant density tissue.

What can be said about the forces acting in a continuous system like plant tissue? They can be thought of as the external and internal forces affecting the tissue. In this model, we assume that external forces such as gravity are negligible, and direct mechanical contact forces are not considered. Internally, we assume that the pressure inside each cell is a constant in space and time (no pressure gradients) and that no significant elastic restoring forces act at this scale. This allows for the construction of the simplest possible version of the transport equation, and allows the model to examine the effects of curvature as separate from other possible physical effects. However, should terms representing pressure gradients and elasticity be needed in order to extend the applicability of the model in different regimes, it would be possible to incorporate them into the transport equation.

We do note, however, that individual cells are known to undergo elastic and plastic deformations during growth, and measurements of externally induced tissue strain have been performed \[31\] \[58\]. These elastic restoring forces generally do not affect tissues at the scales we consider here. For instance, cutting a leaf or petal tissue does not cause it to relax to a different shape, nor does stretching the tissue result in deformations on the mm length scale. Indeed, the deformations are on the \( \mu m \) scale, and the relationship between cellular level elasticity and tissue elasticity is still poorly understood \[58\]. This is in contrast to
highly elastic materials such as rubber, which can deform reversibly at the macroscopic scales we consider. These assumptions also do not exclude the presence of stresses in the tissue due to growth, as discussed in Section 2.2 as will be seen, expansion, shear and rotation tensors can be calculated for the numerically simulated tissue growth models.

Yet another assumption is that the material inside the leaf is incompressible. Leaves are composed mostly of water, while other structures like the mature cell wall are quite rigid [70]. Growth of the cell walls does not occur by pure expansion of the cell wall material, but by the addition more material to the walls.

Incompressibility is also associated with notions of elasticity and pressure gradients. Elasticity is defined as $\epsilon = VdP/dV = [1/VdV/dt]^{-1}(dP/dt)$. Having assumed constant pressure in the tissue previously, this leads directly to zero elasticity of the material. Compressibility is quantified as the inverse of elasticity, so for zero elasticity, we necessarily have an incompressible medium. Intuitively, one can think of a synthetic tissue made up of many balloons. The tissue is not constrained at the outer boundary. As the balloons expand, they can also add material to their walls (just like a cell), thereby maintaining constant pressure inside each balloon while at the same time accommodating ever more material (air) inside of them.

The balloon analogy can be taken one step further when we notice that the balloons are also in local equilibrium with each other. Because of this, the elasticity of the individual balloons is not related to the elasticity of the whole tissue in a simple way. If the balloons are connected in a such a way that they maintain the same neighbours over time (as is the case in plant tissue), externally stretching the tissue means stretching many connected balloons. Therefore, the individual elasticity of each cell contributes to the elasticity of the tissue as a whole, but the effective elasticity of the tissue will be much lower. This has in fact been measured to be the case in iris leaves, where the elasticity of the cell walls is $29 - 100$Mpa while the elasticity of the leaf is several orders of magnitude higher, at $0.838 - 22.6$Gpa [49].

We do assume there is viscosity in the system, noting that as plant cells grow they keep
the same neighbours throughout late-stage development, which can be viewed as a sort of friction between the elements in the tissue. Materials traveling through a plant tissue also experience viscosity by way of diffusion and active transport processes. Transport happens across cell walls in a number of ways, including by way of carrier proteins, osmosis, or plasmodesmata. Transport of material also happens within the cell, again in a variety of chemical processes. We assume that the net macroscopic effect can be understood in terms of a diffusive process.

Viscosity is a standard term in the transport equation given by the Laplacian of the velocity field $\nabla^2 v$. Under the preceding assumptions, the velocity field evolves under advection and diffusion:

$$\frac{\partial v}{\partial t} = -v \cdot \nabla v + D \nabla^2 v$$

where $D$ is the diffusion coefficient, which in the simplest scenario is a constant throughout the material. In more complex scenarios, the diffusion term can itself be a spatially varying tensor.

Indeed, Equation 2.39 is a form of the Navier-Stokes equation for incompressible viscous flow in flat space. The curved-space version of this equation will be revisited in Section 2.10.

2.6 Local interactions in open, driven systems

In constructing this model of plant growth, two ideas have not come into play. Namely, the notions of closed and near-equilibrium systems. Having adopted the view that leaf growth is a result of material transport from the stem into the leaf that at large scales predominantly causes cell expansion, these two commonly used assumptions for many physical systems simply do not hold. The plant leaf is an open system because of the influx of new material. The system is far from equilibrium because it is driven by new material flowing into it. Moreover, cells manufacture materials and consume energy locally in cellular processes, from
photosynthesis which stores solar energy as sugar, to endoreduplication which produces multiple copies of a cell’s DNA without cell division (mitosis).

For an open and driven system in which interactions are still local, we follow the general prescription given by Kardar that the dynamical equations are the ‘fundamental object of interest’ \[37\]. Since the system still relies on local interactions, the general structure of its deterministic dynamical equations is:

\[
\partial_t h(x, t) = v(h(x, t), \nabla h(x, t), ...) \tag{2.40}
\]

where \( h \) is a generalized field and \( v \) is a velocity.

Such a structure can clearly lead to nonlinear interactions, where the field interacts with itself and its gradients as determined by the structure of the velocity functional. Since we are dealing with a pattern forming biological system, this result is also consistent with reaction-diffusion dynamics in biology.

The idea of local interactions governing global behaviour is also consistent with the methods of differential geometry. Local behaviour can be integrated over the entire structure to produce global topological characteristics. For a plant leaf, each cell grows locally but its growth must be consistent with global mass continuity and velocity flow.

### 2.7 Leaf dynamics as nonlinear coupled tensor equations

In considering what factors influence the time evolution of a leaf’s geometry, two factors have already come to light: growth and curvature. Combined with the notion that growth, curvature and the metric are all tensor quantities, we can begin to piece together the structure of our dynamical equations.

Namely, what is necessary to capture the feedback between geometry, growth and curvature is a set of nonlinear coupled tensor equations. In the following sections, the tensor formalism will bring together all the important biological and physical arguments put for-
ward so far (forces, growth, continuity, curvature, geometry), steering us toward a specific mathematical structure.

A significant impact of adopting a formalism based on nonlinear coupled tensor equations is that, in general, they do not have analytical solutions. There is no superposition principle for nonlinear systems that guarantees a complete set of basis functions from which a general solution can be constructed. Instead, we must turn to numerical simulations, particularly in realistic situations.

This approach is also fraught with difficulty. Nonlinear systems can be inherently unstable, which makes computing them not an easy task. Constructing a numerical simulation of our coupled tensor equations will require choosing a stable finite differencing scheme. This is discussed further in Chapter 3.

Based on the issues discussed above, we now present the specific equations that constitute the model.

2.8 Defining the domain and boundaries of 1D and 2D tissue models

Figure 2.3: 1D growth in Lagrangian coordinates. The effect of nonhomogeneous growth on initially equidistant coordinate positions is shown. The total length of the 1D manifold is the sum of all the distances between coordinate positions from the origin to the outer boundary.

In modeling the growth of tissue, we choose a Lagrangian coordinate system that is fixed to the tissue itself. As discussed previously in this chapter, the Lagrangian formalism allows us to model the dynamics of growth in a way that is intrinsic to the tissue itself, that is, without reference to external, fixed coordinates.

This approach is analogous to the landmark method used in experimental biology to
measure growth of tissues in a nondestructive way. Initially, points (such as ink dots) are placed equidistantly on the tissue. Over time, growth displaces the points, revealing where the tissue has grown. In general, biological growth is nonhomogeneous, which manifests itself as distances between landmarks increasing nonhomogeneously as the system evolves. This method has been used for nearly a century and continues to be utilized in more modern variations [4] [21] [77].

Our mathematical model requires three dynamical variables: the mass density, a velocity field and a metric tensor. These will be discussed in more detail in subsequent sections, but first we define the domain and boundary conditions underlying the model.

In our model, the tissue is defined on the domain $x = (0, x_{max})$ in 1D, and $r = (0, r_{max})$, $\theta = (0, 2\pi)$ in 2D. Further discussion on why a circular domain is chosen for 2D models is discussed in Chapter 5. The domain is subdivided into an integer number of fixed coordinate positions. As the tissue grows, the distances between coordinate positions also grow. This process is encoded in the metric tensor evolution discussed Section 2.11. A similar approach is taken when studying cosmology and the expansion of the universe. In that system, galaxies
mark the fixed coordinates of the universe, and expansion is quantified by the rate at which galaxies move away from each other.

The outer boundary of the tissue stays fixed at \( x_{\text{max}} \) in 1D and \( r_{\text{max}} \) in 2D. Growth occurs throughout the tissue inside of the boundary by increasing the distance between coordinate positions. The total length or radius of the tissue is then the sum of all the distances between the origin and the outer boundary.

Time, represented by \( t \), is not in principle bounded for these models. It has arbitrary units that can be adapted to the magnitude of the diffusion coefficients. The model treats time as a continuous variable.

The boundary conditions for the model are based on both mathematical and physical considerations. As has been discussed, the equations forming the model will be differential equations involving second spatial derivatives that represent diffusive processes. This necessarily requires the functions to be at least twice differentiable everywhere. The dynamics also depend on reaction terms that are first spatial derivatives of the dynamical variables.

For the velocity field, we also assume the outer boundary of the tissue cannot allow material to flow out of it; the outer boundary therefore represents a condition on the diffusion term (second spatial derivative). One plausible condition for the outer boundary that satisfies all of the above arguments is a Neumann condition. In 1D, for a velocity variable \( v \), this would be \( \partial_x v(t, x = x_{\text{max}}) = \phi(t) \) where the function \( \phi(t) \) is equivalent to \( \partial_x v \) evaluated at \( x = x_{\text{max}} - 1 \). In 2D, the equivalent statement would be \( \partial_r v(t, r = r_{\text{max}}, \theta) = \phi(t, \theta) \) where \( \phi(t, \theta) \) is equivalent to \( \partial_r v \) evaluated at \( r = r_{\text{max}} - 1 \). These conditions quantify the notion that diffusion does not occur across the outer boundary while allowing the dynamical variables to remain twice differentiable.

For the inner boundary, a similar consideration of dynamics leads to the boundary conditions for the velocity at the origin. Here, a Dirichlet condition is appropriate since we hold the origin fixed in place for both 1D and 2D models. Hence, the velocity is null at the origin.
for all time.

The metric can in principle expand anywhere in the domain, including the origin, so a Neumann condition is more appropriate for this dynamical variable. One possible constraint on the metric components (for simplicity denoted by $f$) that also provides numerical stability in simulations discussed in future chapters is: $\partial_x f(t, x = 0) = 0$, $\partial_x f(t, x = x_{max}) = 0$ (1D model); $\partial_r f(t, r = 0, \theta) = 0$, $\partial_r f(t, r = r_{max}, \theta) = 0$ (2D model).

For 2D simulations, the angular coordinate $\theta$ has a periodic boundary condition, reflecting the symmetry of the imposed geometry.

2.9 The mass density equation

The first dynamical variable is the scalar mass density of the tissue $\rho(x, t)$. At each point $x$, the density is represented by a number (scalar) value. Also, the density $\rho$ may change over time $t$ according to the continuity equation.

$$\frac{\partial \rho}{\partial t} + \nabla_i (\rho v^i) = S(x, t). \quad (2.41)$$

The two terms are the same as in Equation 2.34 for mass conservation, but generalized to curved geometries because $\nabla_i$ is a covariant derivative rather than a flat-space partial derivative. We also note that $S(x, t)$ is called the deposition rate in [65].

As a simplification, we assume the leaf has constant density in time and space. This is also consistent with the assumption of incompressible fluid flow discussed in Section 2.5.2 and [50]. Indeed, plant tissues are made mostly of water and cell walls are rigid structures that grow by deposition of new cell wall material (rather than pure elastic stretching), further justifying this assumption at large spatial scales.

Therefore we have:
\[
\frac{\partial \rho}{\partial t} = 0
\]
\[
\nabla_i \rho = 0.
\]
(2.42)

This means we assume the source term is in balance with the velocity field to create a constant density tissue. Biologically, this can be interpreted as the tissue drawing as many resources as are required to maintain its growth pattern; more active cells will draw more resources than quiescent or mature cells.

It is important to note the implication of these assumptions, summarized in the form:

\[
S(x, t) = \rho \nabla_i v^i = \rho \Theta.
\]
(2.43)

Recall from Section 2.2 that \(\nabla_i v^i\) is the expansion scalar \(\Theta\). Hence, the influx of material from the distributed source directly influences the macroscopic, physical expansion of the tissue. The other influence becomes the geometry of the manifold itself, since \(\nabla_i\) is a covariant derivative calculated based on the metric tensor of the curved 2D space in which the material is flowing.

2.10 The velocity field equation

The second dynamical variable in this model is the velocity field describing the transport of material throughout the leaf. Since we consider the system to be open and driven, we must construct this equation using suitable local dynamical terms.

The most elegant way to understand how the geometry affects the velocity field is by considering the Lagrangian of a particle moving in curved space, as was done in Section 2.4. Expanding on this, we can extend Equation 2.24 to include external forces:

\[
\ddot{x}^i + \Gamma^i_{jk} \dot{x}^j \dot{x}^k = f^i.
\]
(2.44)
Next, we replace $\dot{x}_j$ and $\dot{x}_k$ with the velocity vectors $v^j$ and $v^k$, and split the full time derivative $\ddot{x}^i$ into partial derivatives in space and time of the velocity vector $v^i$.

\[
\frac{\partial v^i}{\partial t} + v^k \nabla_k v^i + \Gamma^i_{jk} v^j v^k = f^i.
\] (2.45)

The forces $f^i$ are the same internal and external forces discussed in Section 2.5.2. If we again assume a viscosity term, an incompressible medium, and no elastic restorative forces, no velocity dependent damping forces and no other external forces and pressure gradients across the tissue, we obtain the following equation for the time evolution of the velocity field:

\[
\frac{\partial v^i}{\partial t} = -\Gamma^i_{jk} v^j v^k - v^k \nabla_k v^i + \epsilon g^{jk} \nabla_j \nabla_k v^i.
\] (2.46)

Comparing this equation to Equation 2.39, we see it is equivalent to the Navier-Stokes equation generalized to a curved geometry. Just as the flat space Navier-Stokes equation is a restatement of Newton’s Second Law, all the terms in this equation can be understood in terms of forces acting on the infinitesimal elements of viscoplastic tissue.

The first term contains the connection coefficient $\Gamma^i_{jk}$. When combined with the velocity field, $\Gamma^i_{jk} v^j v^k$ becomes the contribution that the curvature of the leaf makes to material transport. It can be thought of as a ‘geometric force’.

Another way to think of the $\Gamma^i_{jk} v^j v^k$ term is that it represents the shortest path a particle can take across a curved geometry. The direct impact for material transport is that, in a curved geometry, the shortest path is no longer ‘straight’ in the conventional (flat space) sense. In other words, the curvature of the leaf will influence how material flows across it.

It is also important to note that even if the velocities themselves are small, the effect of the connection coefficients can still produce a large change in a particle’s motion. This is because the connection coefficients are calculated based on spatial derivatives of the metric (Equation 2.25). If the spatial derivatives of the metric are large, this can influence the transport of material even at small velocities. Indeed this will be the case in many of the
simulations studied in later chapters.

The second term, representing advection, comes from the fact that both the outer boundary and every point inside the boundary experience a velocity as the leaf grows. This velocity field is a result of material entering the leaf (i.e. the distributed source term from Equation 2.41). As the material flows through the tissue, it is also affected by the expanding geometry of the tissue.

The last term represents diffusive processes in the tissue. These arise from a host of cellular transport processes, both passive and active, which then can be generalized to an effective, macroscopic material diffusion throughout the tissue. Transport processes are also used to grow the tissue and generate curvature, which will be explored further in the next section.

This modified form of material transport now includes nonlinear velocity terms. It can be thought of as a general reaction-diffusion equation for the velocity. Note too the presence of the covariant derivatives $\nabla_k$ which further couple the material transport to a curved geometry, allowing curvature to act as a force on the vector field of the material velocity.

### 2.11 The metric tensor equation

The third and final dynamical variable is the metric tensor. The metric tensor characterizes the geometry of the leaf. Since we assume the leaf is a thin sheet, the metric is two dimensional.

\[
\frac{\partial g_{ik}}{\partial t} = -\kappa R_{ik} + \kappa_1 [\nabla_i v_k + \nabla_k v_i] \tag{2.47}
\]

where $\kappa$ and $\kappa_1$ are coupling constants that at a fundamental level will depend on the system under study. These may be determined by the environment and/or the genetics of the organism.
Let us start with the last term, which is a symmetric growth tensor

\[ T_{ik} = \nabla_i v_k + \nabla_k v_i. \]  

(2.48)

The notion of a growth tensor of this form was originally introduced by Silk and Erickson [66]. Including this coupling provides feedback between the velocity field and the metric tensor. Since the metric tensor and the Ricci tensor are symmetric \((g_{ik} = g_{ki} \text{ and } R_{ik} = R_{ki})\), the growth tensor must also be constructed symmetrically.

Why is the growth tensor a gradient of the velocity field? If we think of two points on the leaf with the same velocity relative to each other (no gradient), they experience no change in the distance between them, so no growth has occurred. If, however, the points move at different speeds or in different directions (non-zero gradient), then growth or contraction must be occurring between the two points. This also clearly indicates that the growth tensor must affect the metric tensor since the metric tensor measures distances between points.

The growth tensor is also closely related to the expansion, shear and rotation tensors (Equations 2.12, 2.13 and 2.14). This can be understood in the context of treating growth as a deformation of the surface. It also indicates that growth causes strain in the tissue, an important concept that will be revisited later on. In fact, the growth tensor \(T_{ik}\) as defined in Equation 2.48 is twice the strain rate tensor commonly used in continuum mechanics.

The first term is the coupling of the metric to its own curvature. Including this term expresses the notion that curvature directly influences the time evolution of the leaf’s shape, as discussed in Section 1.1. Alternatively, this equation can be interpreted as stating that growth leads to both metric expansion and curvature, an effect clearly described in thin sheet experiments (Section 1.3). The negative sign is kept from Equation 2.33 and indicates that an increase in scale factor leads to negative curvature, which is consistent with experiments on torn plastic sheets [60].

The Ricci tensor components are also functions of first and second derivatives of the
metric components. In this way, the Ricci flow can be thought of as a reaction-diffusion term for the metric itself.

Readers familiar with differential geometry may wonder why the metric flow is coupled to the Ricci tensor rather than some other measure of the curvature like the Riemann tensor or the scalar curvature (lowered or raised as needed to become valence (0, 2) tensors). In fact, choosing any of these curvature couplings is equivalent in 2D. The Riemann tensor has only one unique component in 2D, so there is no information lost going from the Riemann tensor to the Ricci tensor \( R_{ik} \) or scalar curvature \( R \). Even the Gaussian curvature \( K \) is directly proportional to the scalar curvature, \( K = \frac{1}{2} R \). Using the Ricci tensor is convenient because it has the same valence as the metric and growth tensors.

Quite importantly, Ricci flow allows the growing tissue to dissipate curvature because of its reaction-diffusion type behaviour. This is important in a growing tissue, since a region of very rapid growth causes strain on the neighbouring tissue as well as a localized area of high curvature. The ability to dissipate this area of high curvature prevents the tissue from tearing and excessive bending. If cells in the tissue were to sense this area of high curvature and respond through a mechanotransduction pathway such that the curvature dissipates, then Ricci flow is merely another physical phenomenon of macroscopic leaf growth, much like the growth tensor.

Another interesting feature of coupling the time derivative of the metric to its own curvature is that even if the tissue is not growing (i.e. the growth tensor vanishes), the tissue still seeks the most uniform shape possible through curvature diffusion. This is a general property of the Ricci flow equation.

Explicitly coupling curvature and growth to the geometry of the leaf can only be done using rank two tensors of the same valence. Hence, our assumptions about the feedback between growth, curvature and geometry in plant leaves have led us directly to using a set of coupled nonlinear tensor equations. In particular, an expression emerges that allows
these three important elements to interact directly with each other, while at the same time excluding other couplings. For instance, $\partial_t g_{ik} = T_{ik}$ couples the growth tensor directly to the metric, but neglects the reaction-diffusion terms of the Ricci tensor that allow areas of quick growth to dissipate their curvature. Another variation that is incompatible with the dynamical nature of growth is $R_{ik} = T_{ik}$ which does not allow for the metric itself to grow over time, only change shape.

Indeed, the number of 2nd rank tensors that can be constructed out of dynamical vector quantities is quite limited. We can either use tensor products or derivatives, and the quantities thus created must still be relevant to the physics of the problem. Given that we have assumed a scalar mass density that is constant in both space and time, derivatives and tensor products of this quantity will either be null or constant. The covariant derivatives of the velocity field have already been used to construct the symmetrized growth tensor. A tensor product of the velocity field would yield terms related to an energy-momentum tensor that would have to obey a conservation law; as we have seen, the influx of material and energy into the leaf violates conservation laws, so the inclusion of a velocity tensor product would not reflect the physics of the system. Lastly, the metric tensor and its derivatives have already been included by way of the Ricci tensor. Hence, the equations already contain most, if not all the possible rank 2 tensors that can be constructed self-consistently from the dynamical variables that make up the model.

A further refinement of which terms to include when constructing a dynamical model is related to the principle of minimal coupling \[17\]. There are many tensors that could be made to fit the rank two, rank one and rank zero equations, simply by raising and lowering indices. However, these do not necessarily represent any new physics acting on the system. Each term included in the equations introduced in this section is based on physical or biological arguments. The resulting mathematical structure is found to be consistent with the dynamics of an open, driven system. Minimal coupling specifies that only the metric
The tensor can be coupled to curvature; other tensors like our velocity field and mass density should have no explicit curvature coupling terms of the type \( \partial_t v^i = \kappa R^{ij} v_j \).

The possibility of other tensor couplings is, however, not strictly out of the question. These could include physical forces that are not considered here, such as restoring forces of the type \(-k^{ij} x_j\) or retarding forces of the type \(-B^{ij} v_j\) [13]. Should it be found that key dynamics are not present in the model as it is presented here, different couplings can be explored that still satisfy the physical and biological assumptions of the model.

2.12 Chapter summary:

The interplay of geometry, growth and curvature

From these biological and physical considerations of mechanosensing, continuity, growth and curvature, we can now state the problems our theory seeks to explore.

First, let us summarize the system of equations introduced in this chapter that are the foundation of a tissue growth model based in dynamical Riemannian geometry:

\[
\frac{\partial \rho}{\partial t} = 0 \quad (2.49)
\]

\[
\frac{\partial v^i}{\partial t} = -\Gamma^i_{jk} v^j v^k - v^k \nabla_k v^i + c g^{jk} \nabla_j \nabla_k v^i \quad (2.50)
\]

\[
\frac{\partial g_{ik}}{\partial t} = -\kappa R_{ik} + \kappa_1 [\nabla_i v_k + \nabla_k v_i] \quad (2.51)
\]

This can be considered to be a 'minimal' model where the only forces considered are geometric and diffusive, and growth is driven by deposition of material from a distributed source.

The central theme of our theory is the feedback between growth and curvature at macroscopic scales. On the one hand, growth causes inhomogeneous and anisotropic changes in geometry that manifests itself as a deformation of the tissue; hence, growth leads to curva-
ture. On the other hand, curvature generates strain on the tissue, as real as any externally applied force. Through mechanotransduction, these forces caused by curvature could be sensed by the tissue, creating a self-regulating system. This is feedback is captured in Figure 2.5.

The details of mechanotransduction are not addressed by our theory as that is clearly the domain of experimental biology. What is addressed is a new and rigorous mathematical model of how the feedback between growth and curvature can occur given the assumption that leaves act like continuous, deformable, thin-sheet systems at some threshold scale.

What are the possible questions that can be asked with a mathematical model of plant growth?

For one, we would like to know if such a model results in shapes similar to those found in wild-type and mutant plants. In other words, the shapes and structures that result from such a model form a landscape which can be explored by varying the parameters and initial conditions in our set of coupled tensor equations. The extent to which a comparison might be done between model and real leaves will be discussed in Chapters 5 and 7.

We can also ask if our assumptions are sufficient to understand the feedback between geometry and growth. If the solutions to our equations do not produce the shapes found in plant leaves, then our basic understanding of the system is flawed.

Lastly, can our model predict some fundamental but as yet undiscovered part of plant growth? As mentioned in Section 1.1, the dynamics of cell expansion as the leaf matures at scales larger than 1mm are not well understood. Modeling this stage of leaf development mathematically may produce results that help guide experiments toward understanding the interplay of growth, geometry and mechanotransduction in both wild-type and mutant plants.
Figure 2.5: A diagram of how geometry, growth and curvature may be connected in plant growth. Purple arrows indicate novel concepts from modeling the leaf as a Riemannian geometry.
Chapter 3

Numerical Methods

In the previous chapter, three dynamical equations were developed to model the physics of a growing plant tissue: the mass density equation, the velocity field equation and the metric tensor equation. Though initially developed as coupled tensor equations, they can be rewritten component by component using continuous functions. In this way, the coupled tensor equations become coupled partial differential equations (PDEs). More details on the functions themselves will be discussed in Chapters 4, 5 and 6.

The study of numerical methods to solve coupled nonlinear PDEs is a branch of mathematics and computation in its own right. Here, we present an outline of the guiding principles and tests of the numerical methods used to study the dynamical equations in our model. All code was written in C using standard libraries and can be run on any computer with a UNIX platform. This was a programming choice that allows the code to be open-source and free of 'black box' solutions to sensitive numerical problems. Full source code is available from the author upon request.

3.1 Finite element versus finite differencing methods

In order to numerically model the dynamics of our system, several important factors must be considered. On the one hand, we seek to accommodate the features of large scale leaf tissues we know to be true. These include continuous patterns at mm and larger scales, time continuous growth, nonlinear viscoelastic behaviour of both cells and the tissue continuum, and a nonhomogeneously expanding geometry with large changes in scale and curvature (typically 1000-fold areal expansion from the end of cell proliferation to maturity \[19\]). On the other hand, we must choose appropriate numerical methods that will be optimal for
modeling this system. A priori, there is no obvious choice for how to best simulate this set of coupled partial differential equations.

Numerical simulations of PDEs in physical systems are generally handled either by finite element methods or finite differencing methods \[50\]. Other more specialized techniques exist as well, but we restrict our attention to these two broad categories as they are widely adopted in scientific and engineering applications.

In finite element methods, the physical space being modeled is subdivided into a grid of small areas or volumes. This approach mimics the plant system at the cellular level, where a continuous tissue is made up of cells that do not move relative to each other and have rigid walls.

However, several issues arise with adopting a finite element approach. Foremost is the question of growth. For a growing manifold, the geometry of the system changes with every time step and is coupled to the dynamics within the geometry. Hence, the elements of the grid itself would have to change at every time step in nontrivial ways such as location-dependent rescaling, the addition of more elements throughout the grid and the boundary, anisotropic growth of each element, and so on.

The use of a finite element method also requires that some quantity be conserved in the system being studied. In viscoelastic theories, this is mass and momentum. For problems in fluid mechanics, this makes intuitive sense both globally and locally. Globally, matter and energy cannot be created or destroyed. Locally, the influx of matter and energy across one boundary of an element must balance the efflux across the other boundaries. In a growing cell, however, this is not necessarily the case. An influx of material can cause growth rather than balanced efflux. It is not obvious what the conserved quantity should be for a tissue driven to grow by external source terms, and whether such a term can be expressed in integral form as required for a finite element method.

Finally, the case for finite element methods is further weakened by the inherent assump-
tion of an Eulerian coordinate system. This is in fundamental conflict with our previous assumption of a Lagrangian coordinate system for the plant leaf (Section 2.3). While not irreconcilable, a finite element method would require nontrivial calculations to translate between the two coordinate frameworks. More importantly, key features of plant growth that depend on Lagrangian coordinates, such as the stationary behaviour of the elongation zone in roots and the curvature of a cotyledon [21], would be more difficult to compare if calculated in an Eulerian coordinate system.

In the continuum limit, certain factors come into play that suggest a finite differencing approach is more appropriate. The equations are written in differential form, which is easily representable in a variety of finite differencing schemes. In this way, a finite differencing scheme is the most direct way of numerically interpreting the mathematics of the problem. The requirement for a conserved, integrable quantity is also lifted.

Furthermore, the central physical aspects of tissue growth can all be formalized with continuous functions. Modeling the dynamics of these continuous functions in a curved, dynamical, growing geometry is similar to the problems encountered in numerical general relativity where matter and geometry interact via nonlinear feedback dynamics. There, progress has been made using finite differencing methods so it is not unreasonable to try an approach based on similar numerical methods.

Lastly, a finite differencing scheme can more easily accommodate calculations in Lagrangian coordinates.

### 3.2 The Dufort-Frankel method

In light of the above arguments, we adopt the DuFort-Frankel finite difference method for the numerical analysis where the equations have terms that are diffusive in one spatial dimension. That is to say, this method is appropriate for 1D and constrained 2D geometries, but is not generally used where mixed second derivatives appear.
The Dufort-Frankel method is a semi-implicit differencing method that is particularly accurate for diffusive, heat-like systems \[50\]. As we will see in the next chapters, the Ricci flow term is functionally similar to a diffusive term when written out for each metric component. Hence the Dufort-Frankel method is chosen to accurately represent the dynamics of Ricci flow in the metric. The diffusion in the velocity field will also be treated using this method.

In general, finite differencing methods discretize the parameter space of continuous functions. In the formulas considered here, space and time become discrete parameters for the metric and velocity field components.

To represent this discretization, let us consider the continuous function \( u \) with a spatial variable \( x \) and time variable \( t \). We let the spatial and time variables be discretized in increments of size \( \Delta x \) and \( \Delta t \), and enumerate them using indices \( j \) and \( n \) such that

\[
  u(x, t) \to u(j\Delta x, n\Delta t) \to u^n_j. \tag{3.1}
\]
For spatial derivatives, we use a central differencing:

\[
\frac{\partial u}{\partial x} \bigg|_{nj} \to \frac{u_{n+1}^j - u_{n-1}^j}{2\Delta x}.
\]  (3.2)

Let us look at how to treat the standard diffusion equation

\[
\frac{\partial u}{\partial t} = D \frac{\partial^2 u}{\partial x^2}
\]  (3.3)

where \( D \) is the diffusion coefficient.

Using Equation 3.2, the standard finite difference scheme for a second-order spatial derivative yields:

\[
\frac{\partial^2 u}{\partial x^2} \bigg|_{nj} \sim D \frac{u_{n+1}^j - 2u_n^j + u_{n-1}^j}{(\Delta x)^2}.
\]  (3.4)

It is important to note that this standard differencing scheme uses terms evaluated all at the same time step \( t = n\Delta t \). Because of this, when expression 3.4 is coupled to a time derivative as in the diffusion equation 3.3, the method becomes unstable for \( \Delta t/(\Delta x)^2 \geq 1/2 \) [50]. To avoid such a numerical instability, the Dufort-Frankel method can be used, which uses terms from two time steps rather than just one.

If we apply a central differencing scheme for the time derivative and the Dufort-Frankel method for the second-order spatial derivative, the discretized diffusion equation becomes:

\[
\frac{u_{j}^{n+1} - u_{j}^{n-1}}{2\Delta t} \sim D \frac{u_{j+1}^{n} - 2u_{j}^{n} + u_{j-1}^{n}}{(\Delta x)^2}.
\]  (3.5)

In order to solve this equation numerically, we isolate \( u_{j}^{n+1} \) algebraically, thus solving for \( u \) at \( t = (n+1)\Delta t \) using the previously calculated values of \( u \) at \( t = n\Delta t \) and \( t = (n-1)\Delta t \). See Figure 3.1 for a pictorial representation.

Using values averaged over both space and time allows the Laplacian terms to be more stable than with simpler, fully explicit differencing methods such as the Euler method. However, where second derivatives in two different coordinates appear, in the full 2D anisotropic
equations, a time-averaged Laplacian is used in place of the Dufort-Frankel method because it is easier to implement.

\[
\frac{u_j^{n+1} - u_j^{n-1}}{2\Delta t} = D \frac{1}{2} \left( \frac{u_{j+1}^n - 2u_j^n + u_{j-1}^n}{(\Delta x)^2} + \frac{u_{j+1}^{n-1} - 2u_j^{n-1} + u_{j-1}^{n-1}}{(\Delta x)^2} \right)
\] (3.6)

The trade-off for ease of implementation of the time averaged Laplacian scheme is reduced stability. Simulations using this method have a smaller range of parameters that exhibit stable solutions as compared to the Dufort-Frankel method used in the constrained 2D models. Furthermore, using 2D arrays to store the 2D information of each tensor component increases the computing time several fold. Of the three geometries developed in this work, the time-averaged Laplacian models of 2D systems are by far the most computationally expensive.

For a full derivation of how the dynamical equations become discretized, see Appendix A.

### 3.3 Precision of the Dufort-Frankel method

To analyze the precision of the numerical methods used in these simulations, a convergence test can be performed which tests the behaviour of the system as the spatial and temporal increments are reduced. For a well-behaved system, the simulation should have convergent values as \( \Delta x \) and \( \Delta t \) approach zero. It is important to note that using a standard analytical error analysis involving Taylor expansions of the derivative terms in the equations would not easily capture the nonlinear feedback effects from first-order terms in the equations.

The dynamical equations used in these convergence tests can be found in Chapters 4 and 5. They are not included here for brevity.

In Figures 3.2 and 3.3, convergence tests for 1D and 2D simulations using the Dufort-Frankel method model are shown. The graphs were generated by running the same simulation for increasingly precise time and spatial steps while maintaining the condition \( \Delta t = (\Delta x)^2 \).
The values used to calculate absolute error were all taken at $r = r_{max}/2$ so that error values can be compared consistently across rescaled spatial grids. The error was calculated relative to the highest precision values, and the correlation between error and precision was determined using a linear fit of the logarithms of the data sets.

In each case, the trend is for functional values to rapidly approach the highest precision value. This is a good indicator that numerical simulations using the Dufort-Frankel method are convergent.

Moreover, these data can be used to determine appropriate spatial and temporal resolutions when running simulations. This is necessary because a trade-off must be made between precision and computing time; with more computational power, the simulations can be more precise for the same amount of computing time. Hence, a balance must be found between reasonable computing time and the precision of the simulation. In the 1D simulations, $\Delta t = 10^{-5}$ and $\Delta x = 0.05$ were used. Throughout the 2D simulations, $\Delta t = 10^{-4}$ and $\Delta r = 0.1$ were used. Given the hardware available for running the simulations, this meant run times could be kept between a few minutes to a half hour for 1D simulations of several
Figure 3.3: Convergence of a 2D simulation. Here, $\epsilon \sim (\Delta x)^{2.2}$.

million time steps, and 2D simulations of 750000 time steps.

The coupling constants and initial conditions used in the convergence tests were chosen to be representative of those in the simulations presented in Chapters 4, 5 and 6. Thus, for the 1D tests:

- Geometric reaction-diffusion coupling $\kappa = 20.0$
- Growth tensor coupling $\kappa_1 = 10.0$
- Velocity diffusion coupling $c = 10.0$

\[
f(t = 0, x) = 1.0
\]

\[
v(t = 0, x) = \left[1.0 + \exp(-0.2(x - \frac{1}{2}x_{max}))\right]^{-1}
\]

For the 2D convergence tests:
Ricci flow coupling \( \kappa = 0.75 \)

Growth tensor coupling \( \kappa_1 = 0.75 \)

Velocity diffusion coupling \( c = 0.01 \)

\[
\begin{align*}
  f(t = 0, r) &= 5.0 \exp(-3.0(r - \frac{1}{2}r_{max})^2) + 2.0 \\
  g(t = 0, r) &= 2.0 \exp(-3.0(r - \frac{1}{2}r_{max})^2) + 2.0 \\
  v^1(t = 0, r) &= \frac{0.01}{[1.0 + \exp(-5.0(r - \frac{1}{2}r_{max})]} - \frac{0.01}{[1.0 + \exp(\frac{5}{2}r_{max})]}
\end{align*}
\]

The form of the initial velocity field is chosen based on data from root growth experiments. The details of this are discussed further in Section 4.2.

Performing convergence tests for the time averaged Laplacian method was not practical because of the long run times of these simulations even at \(10^5\) time steps. Instead, knowing that the Dufort-Frankel method is convergent, simulations using the time-averaged Laplacian method were compared against simulations done using the Dufort-Frankel method using the same initial conditions. That is to say, the time averaged Laplacian simulations had the same results as Dufort-Frankel simulations for the same initial conditions.

### 3.4 Accuracy of the Dufort-Frankel method:

**Comparison to an exact solution**

Comparing a numerical solution against an exact solution is possibly the best test of the accuracy of a given numerical method. Central to the simulations is the question of Ricci flow, however the problem of numerical Ricci flow is not widely studied, so a set of exact solutions had to be found first.

For a 2D conformally flat coordinate system in Cartesian coordinates, the line element is

\[
ds^2 = f(x, y)(dx^2 + dy^2)
\]
and the Ricci tensor is:

\[ R_{11} = R_{22} = \frac{1}{2f^2} \left[ f \frac{\partial^2 f}{\partial x^2} + f \frac{\partial^2 f}{\partial y^2} - \left( \frac{\partial f}{\partial x} \right)^2 - \left( \frac{\partial f}{\partial y} \right)^2 \right] \]

For Ricci flow, we seek a solution to the equation

\[ \frac{\partial g_{ik}}{\partial t} = \kappa R_{ik}. \]  

(3.7)

The sign convention used here is different than in Equation 2.33, but is still mathematically consistent with the equations studied in later chapters.

If we impose a constraint for circular symmetry and transform to polar coordinates where \( x^2 + y^2 = r^2 \), the solution for the metric satisfying Ricci flow is

\[ f(r, \theta) = \frac{1}{r^2 + e^{nt}}. \]

where \( n \) is a real number.

The Ricci tensor for this metric is

\[ R_{ik} = -\frac{2e^{nt}}{[r^2 + e^{nt}]^2}, \]

and the time derivative of the metric is

\[ \frac{\partial g_{ik}}{\partial t} = -\frac{n e^{nt}}{[r^2 + e^{nt}]^2}. \]

Therefore

\[ \frac{\partial g_{ik}}{\partial t} = \frac{n}{2} R_{ik}. \]

To test the numerical solution, we write the Ricci tensor in polar coordinates and use the Dufort-Frankel method on the diffusive term:

\[ R_{ik} = \frac{1}{2r^2 f^2} \left[ \left( f \frac{\partial^2 f}{\partial r^2} - \left( \frac{\partial f}{\partial r} \right)^2 \right) r^2 + r f \frac{\partial f}{\partial r} \right]. \]
The accuracy itself is tested by decreasing the spatial step size $\Delta r$. The results of these tests are shown in Figure 3.4. Here, $\epsilon \sim (\Delta r)^{1.9}$ indicating that the Dufort-Frankel method accurately represents the diffusive dynamics of Ricci flow.

![Figure 3.4: Convergence of numerical Ricci flow solutions to an exact solution using the Dufort-Frankel method. Here, $\epsilon \sim (\Delta r)^{1.9}$.](image)

3.5 Coordinate singularities

In the 2D models of tissue growth, a plane polar coordinate system will be used. The details of this are discussed in Chapter 5, but we note here that a significant difference between the 1D and 2D equations is the presence of $1/r$ and $1/r^2$ terms in the 2D case. This is a direct result of using polar coordinates and causes discretized operators with this behaviour to become singular at the origin. These types of singularities can, for the most part, be managed within the numerical simulations since their behaviour is known and can be anticipated. This includes imposing appropriate boundary conditions on the dynamical terms and any other terms experiencing this behaviour (growth tensor components, connection coefficients, Ricci tensor components, etc), and studying the region near the origin using Taylor series...
expansions to ensure that the solutions are in fact regular.

3.6 Boundary conditions and initial values

Nonlinear systems like the one constructed here for plant growth generally have large sensitivities to both boundary conditions and initial values.

Ideally, both boundary and initial conditions would be set using biological data. However, no data exists to constrain the boundary behaviour, and very little information exists on specific initial data for the geometric variables.

Boundary conditions were kept consistent across geometries. For the metric tensor, the first derivative of the continuous functions defining its components was set to zero (Neumann condition) at the inner and outer boundary. For the velocity field the inner boundary was always set to zero based on the argument that the base of the tissue should have a vanishing velocity (Dirichlet condition). At the outer boundary, the velocity field was defined to have a constant first derivative. Specifying derivative information rather than functional values at the boundary allows the absolute values of the functions to change over time in a way that can accommodate a wider range of growth modes. For instance, specifying a constant velocity field value at the outer boundary would necessarily cause the boundary to grow outward at a prescribed rate, rather than be modulated by the dynamics of the whole tissue. Likewise, specifying the values of the metric components at the boundary would not allow the tissue to grow at the boundary based on the physics inside the boundary.

Initial values were chosen based on the available data on velocity fields in 1D plant tissues in [21]. Though not strictly an initial velocity field for a growing root, the data show a monotonically increasing logistic function for the velocity of cells in a root at some intermediate time of development. This general behaviour was then extended to the velocity fields in our 2D models. Initial data for the metric is not available from experiments. For simplicity, our models start with flat initial metrics for growth models. In models that test
only the metric’s response to curvature, we start with Gaussian perturbations superimposed onto a flat metric.

3.7 Chapter summary:

The Dufort-Frankel method for parabolic PDEs

In choosing the numerical methods required to solve the dynamical equations presented in the previous chapter, the first aspects considered were the differential nature of the equations, and whether or not they have conserved quantities. Given that the system is most easily expressed in differential form due to the assumption of local interactions, and has no obvious conserved quantities because it is an open system, finite differencing schemes are favoured over finite element methods.

Next, we saw that the dynamical equations have parabolic-type terms due to Ricci flow in the metric and diffusion in the velocity field. A strong candidate for the finite differencing scheme was the Dufort-Frankel method, which is semi-implicit scheme that uses two previous times steps to calculate the next time step. Comparing numerical and exact solutions for Ricci flow on a circularly symmetric disk confirmed that the Dufort-Frankel method is accurate, and the absolute error follows $\epsilon \sim (\Delta r)^2$. Furthermore, convergence tests showed that the numerical solutions are also well-behaved with increasing precision in the time and spatial steps.

A number of other methodological questions were considered as well, including the effects of coordinate singularities and boundary conditions.

In the next three chapters, numerical solutions to the equations of plant growth will be developed based on the discussion and results presented here. Numerical Ricci flow will provide the basis of the results concerning how the metric behaves over time; these are novel results in their own right given how little is known about Ricci flow in 2D disk-type geometries. The dynamics of disk growth will be further augmented by coupling the velocity
field to the metric. This coupled, nonlinear system of equations will yield different modes of growth that can be controlled by selecting appropriate coupling parameters.
Chapter 4

1D Simulations

We now present results from numerical simulations of 1D growth. This model is a simplification of the full 2D model discussed up until now. Assuming that organs such as roots and stems can be considered one-dimensional objects, with the other two dimensions being negligible, the simulations discussed in this chapter will be applicable to such elongated structures in the macroscopic limit (i.e. length scales greater than 1mm).

4.1 Dynamical equations of growth in 1D

Studying the 1D model is a natural first step in testing the numerical techniques required to model the system and for building intuition about the dynamics of the system. It is not, however, fully analogous to the 2D system. The main difference is that Riemannian curvature does not exist for 1D systems, which fundamentally alters the mathematics of the system. Without curvature coupling to the metric tensor, the nonlinear behaviour in the metric tensor equation disappears.

To retain the nonlinear self-coupling term for the metric evolution, we introduce a term similar in functional form to 2D Ricci flow. This is consistent with Equation 2.40 and recovers the reaction-diffusion behaviour that allows the metric to interact with itself and its derivatives. In fact, the 1D nonlinear term is similar in structure to a model of interface growth introduced by Kardar, Parisi and Zhang \[38\]. The main difference between the two growth models is that the 1D interface growth of a height field accumulates mass by stochastic surface deposition, while in the 1D root model, growth in length can in principle occur anywhere along the manifold. Since the mathematical form of this one dimensional term has self-coupling to the metric and quadratic gradients in the reaction-diffusion term,
we call it the geometric reaction-diffusion term $R$. Including this term reflects the idea that plant tissues can grow by depositing new material anywhere throughout their tissue.

In 1D, the reduced metric tensor has only one component $g_{11}$ which can be written as a real-valued function $f$ called the scale factor:

$$g_{11} = f(x, t) \tag{4.1}$$

where $x$ is the coordinate position relative to the base of the root. The velocity field $v$ is also a function of position $x$ and time $t$ and has only one component $v^i = v^1(x, t)$.

Assuming that the root material remains at a constant density throughout its growth, the tensor equations for the metric and velocity field evolution (Equations 2.47 and 2.46) become two coupled PDEs for the scale factor and the material velocity. The parameter $x$ is a co-moving coordinate that has a constant coordinate value. This is related directly to the way the spatial growth measurements are made. Usually the immature organism is marked with a grid of ink spots or pin pricks that are assumed to have no effect on the growth. This is called the landmark method and has been used since the 1930s to measure plant growth (initial experiments measured tobacco leaves [4]). More recently, ink jet printing on leaves [77] and advanced optical techniques for roots [76] [10] [5] have allowed for precise measurements of growth at micrometer scales.

As the organism grows, the physical distance between the markings expands. This is generally a nonhomogeneous process. The marks are fixed in number (let us say $M$) and their coordinate values are also fixed at position $x^i$ where the index $i = 1, 2, ..., M$. This means that the coordinate separation between the marked positions are fixed as well ($\Delta x^i = x^{i+1} - x^i = h^i$) where $h^i$ is a constant numerical separation value that depends on how the markings are laid out and enumerated. The growth is then measured by the change in the
scale factor such that the physical distance of separation is given as

\[ L(t) = \int_{x_1}^{x_2} \sqrt{g_{11}(x,t)} dx = \int_{x_1}^{x_2} \sqrt{f(x,t)} dx. \] (4.2)

This is exactly the role that the metric is supposed to play for a Riemannian geometry. It acts as the mathematical object that turns coordinate separations into physical length separations.

In a 1D geometry, we can express all the geometrical quantities in terms of the scale factor \( f(x,t) \). The connection coefficient is:

\[ \Gamma^1_{11} = \frac{1}{2f} \frac{\partial f}{\partial x}. \] (4.3)

The covariant derivative of the velocity is:

\[
\nabla_1 v^1 = \frac{\partial v^1}{\partial x} + \Gamma^1_{11} v^1 \\
= \frac{\partial v^1}{\partial x} + \frac{1}{2f} \frac{\partial f}{\partial x} v^1.
\] (4.4)

The covariant derivative can be treated as a mixed second rank tensor in order to determine the curved-space Laplacian, which is needed for the velocity diffusion term:

\[
g^{11} \nabla_1 \nabla_1 v^1 = \frac{1}{f} \frac{\partial^2 v^1}{\partial x^2} + \frac{1}{2f^2} \left( \frac{\partial^2 f}{\partial x^2} v^1 + \frac{\partial f \partial v^1}{\partial x \partial x} - \frac{1}{f} \left( \frac{\partial f}{\partial x} \right)^2 v^1 \right). \] (4.5)

From the covariant derivative, the growth tensor can be calculated as:

\[
T_{11} = \nabla_1 v_1 + \nabla_1 v_1 \\
= 2g_{11} v^1 \\
= 2f \frac{\partial v^1}{\partial x} + \frac{\partial f}{\partial x} v^1.
\] (4.6)
Finally, we define the geometric reaction-diffusion term $R$ which replaces the Riemann tensor $R_{ik}$:

$$R = \frac{1}{2f} \left[ \frac{\partial^2 f}{\partial x^2} - \frac{1}{f} \left( \frac{\partial f}{\partial x} \right)^2 \right] . \hspace{1cm} (4.7)$$

All the contributions to the metric and velocity field time evolution can now be combined for the 1D system, as stated in Section 2.12. Once again, assuming the density is constant over the time scale during which the evolution takes place, the equations governing the 1D dynamical system are given by:

$$\frac{\partial f}{\partial t} = \frac{\kappa}{2f} \left[ \frac{\partial^2 f}{\partial x^2} - \frac{1}{f} \left( \frac{\partial f}{\partial x} \right)^2 \right] + \kappa_1 \left( 2f \frac{\partial v}{\partial x} + v \frac{\partial f}{\partial x} \right) \hspace{1cm} (4.8)$$

$$\frac{\partial v}{\partial t} = -\frac{1}{2f} \left( \frac{\partial f}{\partial x} \right) v^2 - v \left[ \frac{\partial v}{\partial x} + \frac{1}{2f} \left( \frac{\partial f}{\partial x} \right) v \right]$$

$$+ c \left[ \frac{1}{f} \frac{\partial^2 v}{\partial x^2} + \frac{1}{2f^2} \left( \frac{\partial^2 f}{\partial x^2} v + \frac{\partial f}{\partial x} \frac{\partial v}{\partial x} - \frac{1}{f} \left( \frac{\partial f}{\partial x} \right)^2 v \right) \right] \hspace{1cm} (4.9)$$

where we have replaced $v^1$ with $v$.

Before discussing the results obtained from numerical simulations based upon these equations, it should be noted that the 1D equations under certain circumstances reduce to other well known equations used to explore uni-dimensional systems.

First the velocity equation in the case of flat Euclidean space ($f(x) = 1$) reduces to the viscous Burgers equation:

$$\frac{\partial v}{\partial t} = -v \frac{\partial v}{\partial x} + c \frac{\partial^2 v}{\partial x^2} , \hspace{1cm} (4.10)$$

or if $c = 0$, the inviscid Burgers equation. These equations have been used to model different flow patterns arising in systems ranging from turbulent gas dynamics, to urban traffic flow, to large scale cosmological structures.
Equation 4.9, the velocity field evolution, is also a generalized reaction-diffusion equation of the form:

\[ \frac{\partial v}{\partial t} = -\frac{1}{f} \left( \frac{\partial f}{\partial x} \right) v^2 - \frac{c}{f} \frac{\partial^2 v}{\partial x^2} + \cdots \]

The first term is the reaction component, given by nonlinear geodesic flow which is proportional to the connection coefficient. Note that the effect of the covariant derivative in the advection term of Equation 4.9 is to double the geodesic flow term. The second term is diffusive flow. The remaining terms in Equation 4.9 are linear in the velocity or its derivative with respect to \( x \).

Equation 4.8, the evolution of the scale factor, reduces to an equation developed by Kardar, Parisi and Zhang [38] which, as discussed in Section 2.6, takes the form:

\[ \frac{\partial h(t, x)}{\partial t} = \nu \nabla^2 h + \frac{\lambda}{2} (\nabla h)^2 + \cdots \]

where \( h \) is the dynamical 'height function' that changes due to deposition of material on a two dimensional surface. Comparing with Equation 4.8, we identify \( h(t, x) \) with \( f(t, x) \), \( \nu \) with \( \kappa / 2f \) and \( \lambda / 2 \) with \( -\kappa / f^2 \). Recently, the KPZ equation has sparked interest among mathematicians studying nonlinear PDEs, including a 2014 Fields medal won by Martin Hairer for his work in the area.

This equation describes the growth of a surface through deposition of material where \( h(t, x) \) is a height field measured from some flat horizontal reference surface. Examples include the build up of a snowpack during a snow storm, the formation of sand dunes by wind deposition, etc. In our case the height function is replaced by a scale factor that can evolve as a result of a build up of material occurring throughout the system.

The fact that the equations reduce to those discussed above, all of which have been studied extensively, provides us with a number of comparisons that can be made with well known results in order to verify that the numerical methods we have chosen are valid.
4.2 Initial conditions, boundary conditions and coupling parameters for a growing root

In the numerical simulations, we evolve the metric tensor and velocity field over time for $5 \times 10^5$ time steps, subject to coupling parameters:

- **Geometric reaction-diffusion coupling** $\kappa = 20.0$
- **Growth tensor coupling** $\kappa_1 = (0.0, 10.0, 20.0, 30.0)$
- **Velocity diffusion coupling** $c = 10.0$

where the variation in $\kappa_1$ represents different growth tensor couplings.

The initial conditions are:

\[
\begin{align*}
f(t = 0, x) &= 1.0 \\
v(t = 0, x) &= [1.0 + \exp(-0.2(x - \frac{1}{2}x_{max}))]^{-1}
\end{align*}
\]

The boundary conditions in 1D are:

\[
\begin{align*}
\partial_x f(t, x = 0) &= 0 \\
\partial_x f(t, x = x_{max}) &= 0 \\
v(t, x = 0) &= v(t = 0, x = 0) \sim 0 \\
\partial v(t, x = x_{max})/\partial x &= \phi(t)
\end{align*}
\]

The function $\phi(t)$ is equivalent to $\partial v^1/\partial x$ evaluated at $x = x_{max-1}$.

The initial condition on the velocity field was chosen based on data available on relative elemental growth rates of corn roots, which indicates that the velocity field has a logistic shape [21] [64]. We approximate the velocity field of the root as a logistic function. The functional fit is not a true regression, but is consistent with the form of measured velocity fields in roots across many species [75]. The magnitude of the velocity field is scaled so
that the maximum speed is 1.0 (distance unit)/(time unit). Since distance and time units in this model are arbitrary, the magnitude of the velocity field can be rescaled without loss of generality. We preserve the convention found in most root growth literature that the tip is the origin of the coordinate system, i.e. $x = 0$ corresponds to the tip of the growing root with the base of the root appearing to grow away from the tip.

For the initial condition on the metric tensor, we assumed a simple flat geometry. This is analogous to inscribing a pattern of equidistant markers on the root at the beginning of a growth experiment.

4.3 What do the data look like?

In presenting the information, we choose the total plant length and sectional growth as data most easily comparable to biological data on root growth (using Equation 4.2).

The sectional growth in our model is the equivalent of the landmark method discussed earlier in the chapter. This form of presenting the data allows us to understand where growth occurs in a tissue. In plant roots, growth does not occur homogeneously throughout the tissue. The portion of root that grows the most is called the elongation zone, and is a common feature in many species found just behind the root tip extending for several mm, and ending in the mature root tissue that experiences little to no growth once it leaves the elongation zone [70]. The markings placed on the root tissue are, in our model, represented by initially equidistant coordinate positions. As the dynamics of the system evolve, growth occurs nonhomogeneously which becomes evident as different segments come to have different lengths.

We also present the metric tensor, material velocity, geometric reaction-diffusion term and growth tensor which reflect the dynamics of the system itself.

An additional interpretation of the velocity data can be made by plotting the velocity field as a function of proper radial distance $r'$ from the origin (tip), where $dr' = \sqrt{g_{11}} dr$. This
format will be called a *expansion diagram* because it shows how the velocity is distributed over the increasing length of the growing root. For the 2D plant leaf, it is a representation of the velocity of points on the leaf relative to an observer at the origin of the leaf \((x_0 = 0)\) where the local velocity vanishes. From this point of view the expansion diagram is analogous to a Hubble diagram that in cosmology plots the recessional velocity of galaxies with respect to the observer. It is interesting to note the analogy between plant growth and the expansion of the universe is rather close. The markers used in cosmology are the positions of the galaxies. Each galaxy can have its own proper motion due to a combination of interactions with other galaxies and its initial conditions. These proper motions are 'dragged' along with the expansion of the universe, so one of the main problems in cosmology is to distinguish between the motion strictly due to expansion and the motion driven by local interactions. However, since the standard model of our universe is homogeneous and isotropic, the scale factor that measures the physical distances between the galaxies depends only on time. Also, both in cosmology and a growing plant tissue, local expansion contributes to the overall structure on the largest scales.

Lastly, we can also use the velocity field to construct the expansion scalar \(\Theta\). As discussed in Sections 1.3.3 and 2.2, growth is a tensor quantity that is closely related to viscoelastic deformations (expansion, shear and rotation). In the 1D system, given the definitions in Section 2.2, shear is directly proportional to the growth tensor component \(T_{11}\), rotation vanishes, and the expansion scalar becomes:

\[
\Theta = \nabla_1 v^1 = \frac{T_{11}}{2f}.
\] (4.11)

Recall from Section 2.9 that the assumption of constant tissue density in space and time leads to the source field \(S(x, t)\) being directly proportional to the expansion scalar \(\Theta\). Hence, mapping out the expansion scalar will also give us information on how the source of material is distributed through the tissue.
4.4 Results for 1D simulations

Figure 4.1 shows that the total root length increases in a nearly linear fashion (slightly superlinear). Depending on how strongly the growth tensor is coupled to the evolution of the metric, the root may not grow at all (null coupling), or more than quadruple in length (highest coupling).

Figure 4.2 shows the sectional growth pattern of the root simulation. Initially, the 'landmarks' are equally spaced on the spatial domain. Over time, growth is localized near the origin, as can be seen by the landmarks near the origin spreading apart from each other, while landmarks further away from the origin experience little growth relative to their neighbours. This qualitatively resembles the general pattern of root growth [70], with a clearly recognizable elongation zone behind the growing tip. For a direct comparison with biological data, see Figure 3 in [64], and Figures 4 and 7 in [21].

Figures 4.3, 4.4 and 4.5 show that the scale factor, growth tensor and material velocity are self-consistent with the results above. The metric tensor and growth tensor show that
Figure 4.2: Modular growth in 1D for growth tensor coupling $\kappa_1 = 30.0$. Note that the tip is designated as the origin of the coordinate system.

Distances between neighbouring points increase the most in a region that corresponds with the elongation zone seen in Figure 4.2.

The velocity field undergoes nontrivial dynamics due to both viscous Burgers’ type behaviour and contributions from the position-dependent expansion of the root. The steepening of the velocity field is typical of what happens in Burger’s equation, but the diffusive term prevents the flow from forming a discontinuity (shock) as happens in the inviscid Burger’s equation. This type of shock in the root model would in fact be physically impossible to sustain, since the steepening of the velocity field would eventually lead to a step function, indicating that the material would have two different and discontinuous velocities in adjoining regions of the tissue. An alternate mathematical technique to avoid these types of shocks would be to average the velocity field over neighbouring spatial regions. Physically, it is unclear how this would happen in the tissue; a diffusive mechanism provides a more intuitive way for the tissue to avoid such shocks.

The functional form of the initial velocity field is an important factor in the model.
Recalling from earlier in the chapter, the initial velocity field is based on experimental root growth data which shows that roots have a monotonically increasing logistic velocity field. Most significantly, the gradients of the velocity field determine where growth occurs initially. As determined earlier in the chapter, the growth tensor in the 1D geometry is:

$$ T_{11} = 2f \frac{\partial v^1}{\partial x} + \frac{\partial f}{\partial x}v^1. \tag{4.12} $$

Hence, at early times when the metric is flat or nearly flat, the region with the steepest velocity gradient will experience the highest growth rates, causing the scale factor in that region to expand significantly more than in neighbouring regions. However at late times, the feedback between the scale factor and the growth tensor becomes dominant. This late-time behaviour is interesting in that the dominant term in the growth tensor is no longer the spatial derivative of the velocity field, but rather the spatial derivative of the scale factor (metric). This can be seen in Figures 4.3, 4.4 and 4.5 where the peak of the growth tensor is initially small and correlates to the steepest part of the velocity’s logistic shape, but in late times the peak of the growth tensor shifts closer to the origin and is correlated with the
highest values of the scale factor’s spatial derivatives. Hence, the geometry of the root has a significant impact on where growth occurs in the tissue.

Figure 4.6 shows how the velocity field evolves as the length of the root increases. This plot combines information about the root length as it would be measured in an experiment (the proper length) with the velocity field of the material that makes up the root. True to the general model of root growth, the base of the root does not contribute much to the growth, whereas the region behind the elongation zone continues to grow, pushing the root forward and allowing the tip itself to maintain a nearly constant velocity relative to the base.

Figure 4.7 shows how the expansion scalar evolves over time. As Equation 4.11 shows, it is closely related to the growth tensor $T_{11}$, and from Section 2.9 we also know the expansion scalar is directly proportional to the distributed source field $S(x,t)$. We can see that expansion decreases over time and moves in a wave-like fashion toward the root tip. Given how large the scale factor $f$ gets over time (Figure 4.3), this plot essentially shows that growth has a positive feedback quality to it; at late times, even a small expansion leads to large overall growth because the geometric space itself has expanded. This plot is also consistent with
the dynamics of the growth tensor discussed above, where at early times $T_{11}$ is dominated by velocity field gradients, while at late times is dominated by scale factor gradients.

Another interesting aspect of the expansion scalar (or equivalently in our model, the source term) is that it is related to the geometric expansion of the root. For an initially constant scale factor, Equation 4.8 reduces to

$$\frac{\partial f}{\partial t} \sim \kappa_1 T_{11}. \tag{4.13}$$

If we divide both sides by $2f$ and use Equation 4.11

$$\frac{1}{2f} \frac{\partial f}{\partial t} \sim \frac{\kappa_1 T_{11}}{2f} = \kappa_1 \Theta. \tag{4.14}$$

Hence, the expansion scalar gives us some information on where the scale factor, and hence the geometry, is expanding.

Interestingly, the relation in Equation 4.14 is often used in tissue growth literature to
express the relative elemental growth rate (see for example [10]) such that

\[ REGR = \frac{1}{f} \frac{\partial f}{\partial t}. \]  \hfill (4.15)

Clearly, the REGR is proportional to the expansion scalar $\Theta$ only in the absence of a geometric reaction-diffusion term $\mathcal{R}$. Indeed, the standard definition of REGR in plant growth dynamics literature equates REGR with the expansion scalar. In the model developed here, the time evolution of the scale factor also depends on the diffusive behaviour of the geometric reaction-diffusion term. Hence, in a model of plant growth that includes diffusive coupling to the geometry, the REGR and expansion scalar are not equivalent. The REGR can be obtained by dividing Equation 4.8 by the scale factor $f$ and using the definition from Equation 4.11:

\[ REGR = k \frac{R}{f} + 2k_1 \Theta. \]  \hfill (4.16)

Recalling that $\mathcal{R}$ is similar to the deposition term in the KPZ equation (Section 4.1),
this result states that growth is due to both a deposition of material throughout the tissue as well as expansion of the tissue, which is also consistent with the distributed source term introduced in the continuity equation (Section 2.9). Furthermore, REGR can be understood as a true geometric measure of growth when calculated from the metric tensor as in Equation 4.16. In fact, the same measure of geometric growth is used in cosmology to describe the expansion of the universe; in that context it is known as the Hubble parameter $H = \dot{S}/S$ where $S(t)$ is a scale factor representing the ‘radius’ of the universe [17].

The time evolution of the REGR using proper distances is shown in Figure 4.8 in a format that allows for direct comparison to the results of Figure 7 in [76] and Figure 4 in [5], where the dissipative, double peaked behaviour of the REGR near the root tip is consistent with the results of our simulation. If we had assumed that only the gradient of the velocity field contributes to the REGR, then Figures 4.5 and 4.6 would imply a very different behaviour than is seen in Figure 4.8. The difference is due to the effects of the 1D geometry, both in the geometric reaction-diffusion term, and the covariant derivative used to calculate the expansion scalar. In the case of this simulation, the REGR has nearly equal contributions

Figure 4.7: Expansion scalar time evolution for growth tensor coupling $\kappa_1 = 30.0$. 

![Graph showing expansion scalar time evolution](image)
from the expansion scalar and geometric reaction-diffusion term, but with opposing signs and slightly misaligned peak values, leading to the double-peaked behaviour. It is interesting to note that new, high-resolution imaging techniques of root growth have indeed shown that the REGR is not merely the gradient of the velocity, but that it exhibits a double-peaked, time-varying behaviour not fully correlated with just the velocity field structure (Figure 5 in [76] and Figure 4 in [5]). To our knowledge, a hypothesis explaining these results is still lacking, however, our theory provides a mechanism to extend the definition of the REGR that could in principle be used to explain time-varying REGRs with behaviours not solely linked to the gradient of the velocity field.

Let us now look at the time evolution of the geometric reaction-diffusion term $\mathcal{R}$ in Figure 4.9. The most important feature is the emergence of domains in the geometry that alternate between positive and negative values of the geometric reaction-diffusion term. This hints at the possibility that in 2D, domains of positive and negative curvature can emerge from growth dynamics, yielding the ruffled and buckled geometries characteristic of non-flat leaves.
Figure 4.9: Time evolution of the geometric reaction-diffusion term for growth tensor coupling $\kappa_1 = 30.0$. Note that the $5 \times 10^5$ timestep is omitted so that earlier time steps may be more clearly discerned.

Another feature of the dynamics of the geometric reaction-diffusion term and other variables is that there is a wavelike motion of the peaks and troughs along the length of the organism, including an increase in the magnitude of the peaks and troughs over time. This behaviour occurs in many other systems whose dynamics is governed by reaction-diffusion equations where patterns in the field variables form and shift over time.

The results shown above are a small subset of the parameter sets tested overall. They are, however, representative of the main features that came to light. Namely, the coupling to the growth tensor $\kappa_1$ dominates the dynamics of the simulation. In terms of the coupling parameters in the equations, we find that the same variance in geometric reaction-diffusion coupling $\kappa$ or velocity diffusion $c$ (i.e. $0.0 - 30.0$ in coupling strength) does not produce as dramatic differences in final root length and the associated dynamics in the metric tensor and velocity field.

Another important aspect of the parameter space is that in the absence of geometric
reaction-diffusion coupling ($\kappa = 0$) the system is unstable and the growth is unrestricted.

If we look at the evolution of an initially flat metric under this condition we have, from Equation 4.8:

$$\frac{\partial f}{\partial t} = 2\kappa_1 \frac{\partial v}{\partial x} f,$$

which has an exponential solution of the form:

$$f(x, t) \propto \exp \left[ \int \left( 2\kappa_1 \frac{\partial v(x, t)}{\partial x} \right) dt \right].$$

Moreover, as $f$ increases exponentially, the velocity field is driven even more quickly to a shock front. This can be seen in Equation 4.9 where velocity diffusion is inversely proportional to $f^3$. Hence, as $f$ increases, the velocity field is less and less capable of dissipating.

Thus, in the absence of a geometric reaction-diffusion term, the growth of the metric is initially exponential, with the fastest growth occurring where the spatial derivative of $v$ is largest. At the same time, the exponential growth of the metric causes a shock front in the velocity field by shutting down the diffusion term where it is needed most. This seems to imply that a geometric dissipative term is essential in distributing the energy associated with growth between elongation and buckling of the metric.

4.5 Conclusions on 1D results

The 1D results presented here serve as a proof of concept for the dynamical equations of plant growth. The results show qualitative similarities to real biological systems (roots) and demonstrate that the dynamics of a model based on nonlinear coupled tensor equations can produce biologically relevant global properties. For a comparison to biological data, see Figure 3 in [64] which shows the modular growth of a corn root; Figures 7 and 8 in [21] showing time-continuous root growth data and the corresponding root velocity field.
and REGR data; Figures 4, 5 and 7 showing the REGR distribution in three different root specimens, the time variation of REGR at short time scales, and the REGR time evolution in a root over 120min in [76], Figure 4 in [5] showing a double-peaked REGR time evolution over 5 hours in two different plant species.

In the 1D model, growth is initiated by gradients in the velocity field, which is initially a logistic function mimicking the velocity field of the primary root in many plant species. Gradients in the velocity field provide the growth tensor with its initial form. The growth tensor then causes significant changes in the metric tensor. This in turn affects the transport of material through the connection coefficients found in the velocity equation. As seen in analyzing the REGR, growth occurs both through material expansion via a distributed source of material, and the deposition of material throughout the tissue.

To our knowledge, no other root growth model has predicted the temporal behaviour of the velocity field, metric tensor or any of the other tensor quantities presented here. Furthermore, our theory provides a mechanism to extend the definition of the REGR that could in principle be used to explain time-varying REGRs with behaviours not solely linked to the gradient of the velocity field, as observed in [76] and [5]. This is because the REGR in our model includes a geometric reaction-diffusion (deposition) term in addition to a covariantly defined expansion scalar in the REGR.

One major limitation in producing these simulations is that how the velocity field of a primary root changes over long periods of time (i.e. over a quadrupling in length scale) is not well documented in biological literature. The measurement is difficult with such large root tip displacements, and corresponds to time scales on the order of several days [6]. Indeed, the initial velocity field used here represents some intermediate stage of growth and would presumably be different during different stages of development (which appears as a prediction of velocity field dynamics in our model). Also, the physiology of a plant changes with age, which in this model could be represented by time-varying coupling constants that
make the plant more or less susceptible to growth.

Lacking sufficient experimental data to constrain all these parameters in our dynamical model, the initial conditions for the metric tensor and velocity field are based on available static and short-time data that shows a logistic velocity profile in the primary root of many species [21] [64] [75]. The coupling constants for the geometric reaction-diffusion term, growth tensor and velocity diffusion are currently static and arbitrary.

In the next chapters, 2D models representing simplified leaf shapes will be developed. Modeling of the 2D system is necessary to encompass the effect of curvature flow on the metric tensor. This needs to be built up, first from 2D systems exhibiting symmetries like circular symmetry that constrain and simplify the system. A thorough investigation of the parameter space of the system is also needed, including the space of geometries produced by the dynamics.

Comparison of 2D systems to experiment can begin to be more quantitative. The metric tensor from a 2D numerical simulation of our model can be compared against the metric from plastic thin-sheet experiments [41] and models [3]. Comparing the metric of a mature plant leaf is also possible. The more difficult problem will be to compare the growth dynamics between model and experiment since this would require a non-destructive way of probing leaf tissue geometry as it grows over time.
Chapter 5

2D Simulations with Circular Symmetry

5.1 Why circular symmetry?

The next step we take is to build a 2D growth model that represents growing disks. A disk with circular symmetry reduces the complexity of the tensor equations and boundary conditions required to model its growth, thereby simplifying the numerical simulations as well. It is a toy model because of its simple boundary shape and angular symmetry, but can still exhibit the non-zero curvatures found in plant leaves. Also, a circularly symmetric 2D geometry can be thought of as a 1D model generalized through rotational symmetry to two dimensions.

Geometries with circular symmetries do mimic simple plant leaf shapes like the lotus leaf, nasturtium leaf, lily pad and the cap of the algae *Acetabularia*. They can also be extended to other shapes parameterized in \((r, \theta)\) coordinates such as ellipses, lobes and cusps.

Another significant advantage of modeling growing disks is the ability to compare numerical simulations against experimental data on deformations of thin gel disks with circular symmetry. The physics of thin disk deformations have been studied both experimentally \[41\] and theoretically \[3\], in large part to understand the physical and geometric basis of biological shapes like plant leaves. However these studies concentrate on objects whose total mass is fixed so that the deformations result from externally impose forces rather than the addition of material.

In building these first 2D models, we will examine two cases in particular. The first will be to study Ricci flow in disk geometries without the influence of a velocity field. This is important because Ricci flows have not been studied widely using numerical simulations or in 2D disks. This will also give us an idea of how the metric behaves when coupled only to

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its own curvature, thus separating out the influence of geometry in our dynamical model.

The second case that will be studied is that of a growing disk. This will involve the full dynamical equations developed in Chapter \[2\]. The results of both the Ricci flow and dynamical growth simulations will be compared and discussed in the context of thin disk models of plant tissues.

5.2 Dynamical equations of growth with circular symmetry

The critical step in deriving the dynamical equations for a growing disk is to cast the components of the metric tensor and velocity field as continuous functions of the radius \( r \). Indeed, a similar approach was taken in the 1D case, and stems directly from the principles of Riemannian geometry \[28\]. It allows the coupled tensor equations for plant growth to be restated as a set of nonlinear coupled PDEs, which can then be simulated numerically.

Imposing circular symmetry necessarily removes any angular dependence, hence the metric components and velocity field vary only in \( r \). Symmetry also requires the off diagonal terms to vanish since the condition \( d\theta = -d\theta \) is required by isotropy. Hence \( g_{12} = g_{21} = 0 \) so that distances of the form \( g_{12}drd\theta \) vanish. However, \( g_{22} \) does not vanish since purely angular lengths are of the form \( g_{22}d\theta^2 \), thus allowing \((d\theta)^2 = (-d\theta)^2\).

The metric components are therefore:

\[
g_{ik} = \begin{pmatrix} f(r) & 0 \\ 0 & r^2g(r) \end{pmatrix}. \tag{5.1}
\]

Equivalently,
\[ g_{11} = g_{rr} = f(r) \]
\[ g_{22} = g_{\theta \theta} = r^2 g(r) \]
\[ g_{12} = g_{21} = g_{r\theta} = 0 \]

The velocity field components reduce to

\[ v^1 = v(r) \]
\[ v^2 = 0. \]

This is equivalent to stating that the velocity vector \( v^i \) has no \( \theta \) component, and is consistent with isotropy imposing the \( d\theta = -d\theta \) condition \( (v^2 = -v^2 = 0) \).

As discussed in Chapter 2, the full tensor equations for plant growth under the assumptions of constant tissue density are:

\[ \frac{\partial g_{ik}}{\partial t} = -\kappa R_{ik} + \kappa_1 T_{ik} \tag{5.4} \]
\[ \frac{\partial v^i}{\partial t} = -\Gamma^i_{jk} v^j v^k - v^k \nabla_k v^i + c g^{ik} \nabla_j \nabla_k v^i \tag{5.5} \]

where \( T_{ik} \) is the growth tensor.

Writing the metric and velocity field components as continuous functions of \( r \) allows all the parts of the dynamical equations to also be cast as functions of \( r \).
The connection coefficients are:

\[ \Gamma^1_{11} = \frac{f_r}{2f} \]
\[ \Gamma^2_{11} = 0 \]
\[ \Gamma^1_{12} = 0 \]
\[ \Gamma^2_{12} = \frac{g_{r,r} + 2g}{2gr} \]
\[ \Gamma^1_{22} = -\frac{g_{r,r}^2 + 2gr}{2f} \]
\[ \Gamma^2_{22} = 0. \]  

(5.6)

where \( f_r = \partial f / \partial r \).

The scalar curvature function (also known as the Ricci scalar) is:

\[ R = g^{lm} R_{lm} = -\frac{1}{4d^2} \left( 2g_{r,r} f - f(g_{r})^2 - f_r g g_{r,r} + \frac{1}{r} (4 f g g_{r,r} - 2 f_r g^2) \right) \]  

(5.7)

where \( d = -fg \).

In 2D, the Ricci tensor can be expressed as

\[ R_{ik} = R g_{ik}, \]  

(5.8)

so the metric evolution becomes:

\[ \frac{\partial g_{11}}{\partial t} \rightarrow \frac{\partial f}{\partial t} = -\kappa R f + \kappa_1 T_{11} \]  

(5.9)

\[ \frac{\partial g_{22}}{\partial t} \rightarrow \frac{\partial g}{\partial t} = -\kappa R g + \kappa_1 \frac{T_{22}}{r^2} \]  

(5.10)

The general expression for the growth tensor is:

\[ T_{ik} = \nabla_i v_k + \nabla_k v_i \]
\[ = g_{ki} \nabla_i v^i + g_{kp} \nabla_k v^p \]  

(5.11)
The specific expressions are:

\[ T_{11} = 2f(v^1_r + \Gamma_{11}^1 v^1) \]
\[ T_{22} = 2gr^2 \Gamma_{12}^2 v^1 \]
\[ T_{12} = 0. \]  
(5.12)

The velocity equations become:

\[
\frac{\partial v^1}{\partial t} = -\Gamma_{11}^1 v^1 v^1 - v^1[v^1_r + \Gamma_{11}^1 v^1] + cg^{11} \left( \frac{\partial^2 v^1}{\partial r^2} + \Gamma_{11}^1 v^1 + \Gamma_{11}^1 v^1_r \right) \\
+cg^{22} \left( -\Gamma_{22}^1 v^1_r + \Gamma_{22}^1 (\Gamma_{12}^2 - \Gamma_{11}^1) v^1 \right) \]

(5.13)

\[
\frac{\partial v^2}{\partial t} = 0. 
\]
(5.14)

Hence, by introducing continuous functions of \( r \) to represent the metric and velocity field components, we can view the coupled tensor equations as nonlinear coupled PDEs. The additional constraint of null \( g_{12} \) and \( g_{21} \) terms means that the dynamics are not affected by any mixed-derivative terms. Meanwhile, the imposed angular symmetry removes any derivatives with respect to \( \theta \), leaving the equations dependent only on \( r \) and derivatives in the radial direction.

5.3 Numerical methods

The set of coupled PDEs we are left to consider are Equations 5.9, 5.10 and 5.13. Most importantly, the time evolution of \( g_{22} \) and \( v^1 \) contain parabolic (diffusive) terms that can be stabilized by implementing the Dufort-Frankel method, similarly to how the parabolic terms in the 1D equations were handled.

Having more nonlinear terms than the 1D equations, the 2D evolution equations are expected to be less numerically stable than their 1D counterparts. This manifests itself in requiring smaller time steps and smaller coupling coefficients in the 2D models, which effectively slows down the simulation. Simulations of the 2D dynamics will be more computationally expensive than the 1D dynamics.
The boundary conditions in the isotropic 2D simulations are:

\[
\begin{align*}
\partial_r f(t, r = 0) &= 0 \\
\partial_r f(t, r = r_{max}) &= 0 \\
\partial_r g(t, r = 0) &= 0 \\
\partial_r g(t, r = r_{max}) &= 0 \\
v^1(t, r = 0) &= 0 \\
\partial v^1(t, r = r_{max})/\partial r &= \phi(t)
\end{align*}
\]

The function \(\phi(t)\) is equivalent to \(\partial v^1/\partial r\) evaluated at \(r = r_{max-1}\).

5.4 What do the data look like?

The data generated by the 2D simulations is not unlike the 1D data set. Information about the state of the metric, velocity field, growth tensor, connection coefficients and scalar curvature is gathered at prescribed time slices. The bigger problem is how to represent this data in a meaningful graphical way.

Ideally, the data would be represented visually as a growing disk with ruffling and buckling developing over time, in a way that strongly resembles a time-lapse video of a plant leaf growing over time. However, the problem of interpreting geometric data in this way is a mathematically difficult problem. It is called the embedding problem, which can be summarized as the problem of representing a \(N\) dimensional geometry in a \(M\) dimensional space where \(N < M\). In the case of plant growth considered here, we would seek a solution of the embedding problem of a non-flat 2D manifold embedded in a flat 3D space. However, the general solution to the embedding problem when \(M = N + 1\) remains an open problem in geometry. Moreover, no general solution exists for embedding a 2D surface of non-constant curvature in a flat 3D space \[55\], this despite the fact that many leaves found in nature have negatively curved surfaces and remain comfortably embedded in 3D space. Computational
methods do exist for visualizing these types of surfaces, however the techniques remain non-trivial and would generally require a finite element representation of the data rather than the finite differencing data used here \[40\]\[51\].

Lacking a general method to visualize the data generated by the simulations, what are other possible ways to interpret the data? One option is 3D printing each time slice, for which there are a number of methods to explore: hinged plastic components \[57\], programmed thin disks \[41\] and even crochet \[32\]. The advantages of physically representing the data in this way is that it is highly realistic and intuitive. The disadvantage is that these representations would be sparse in time and miss the dynamical aspect of the simulations.

Another interesting consideration is how similar the geometric data will be to the actual shape of the plant tissue. As a soft material conforms to a non-flat metric, the material must bend and stretch to accommodate its new shape. Studies of non-Euclidean plates have indicated that it is the interplay of a target metric with the properties of the material (such as elasticity and plasticity) that determine its final shape \[62\]. It is hypothesized that growing plant tissues experience these effects, but a full understanding of the interplay of growth and embedding remains an open question \[20\].

In the current work, data representing the dynamics of the system is presented in a relatively raw format. For the circularly symmetric simulations, the metric components, velocity field, expansion diagram, growth tensor components and modular growth are graphed as functions of the coordinate distance \(r\) along the \(\theta = 0\) direction, so are similar in format to the 1D data.

We can also determine the deformation tensors for expansion, shear and rotation. In the 2D isotropic case, the expansion is:

\[
\Theta = \frac{T_{11}}{2f} + \frac{T_{22}}{2gr^2}
\]  (5.15)

where \(T_{11}\) and \(T_{22}\) are the growth tensor components from Equation 5.12. The shear tensor
can be defined in its trace-free form:

$$\sigma_{ab} = \frac{1}{2}(\nabla_a v_b + \nabla_b v_a) - g_{ab}\Theta. \quad (5.16)$$

This form of the shear tensor is useful given that we are dealing with expanding surfaces; subtracting the trace eliminates the effect of expansion on the shear. The components of the trace-free shear tensor are then:

$$\sigma_{11} = -f \frac{T_{22}}{2gr^2}$$
$$\sigma_{22} = -g \frac{T_{11}}{2f} \quad (5.17)$$

and the rotation is null.

Calculating the deformation tensors is useful in that these represent the physical changes the tissue goes through as it grows. These quantities could be measured using techniques similar to those in [77] where a regular grid of dots is printed on a growing leaf. Over time, the grid becomes deformed, and could presumably be analyzed to yield the deformations that have resulted from growth. Also, under the assumption of constant tissue density in space and time, the expansion scalar is directly proportional to the distributed source field (Section 2.9), providing yet another way to characterize tissue growth empirically.

Figure 5.1: From left to right, geometries with flat, positive and negative curvatures.

Data specific to 2D simulations also exists and requires additional plots. The perimeter $C$ of the growing disk can be plotted relative to the proper radial distance $r'$. This is entirely
analogous to the plots in [41] and is chosen so that comparisons to existing experiments on expanding thin disks can be readily made. Indeed, plotting the perimeter as a function of radial distance is a fairly intuitive way of knowing whether a surface is positively or negatively curved. A basic result of Gauss’ Theorema Egregium is that surfaces of positive curvature will have a perimeter \( C < 2\pi r' \) while surfaces of negative curvature will have \( C > 2\pi r' \). In Figure 5.1, this is represented by a positive curvature disk that has a deficit perimeter when wedges of the disk are removed, and the negative curvature disk that has excess perimeter when wedges are added.

An analogy of the relative elemental growth rate (REGR) can also be defined in 2D. In the 1D case, the REGR was the rate of change of the scale factor normalized by the local length scale. In 2D, we must work with area instead. One way to define area is using the product of the two scale factors \( f \) and \( g \)

\[
dA = \sqrt{f}dr \times \sqrt{g}rd\theta = \sqrt{fg}rdrd\theta.
\] (5.18)

The corresponding measure of elemental growth is then

\[
REGR = \frac{1}{A} \frac{\partial A}{\partial t}.
\] (5.19)

Using the expressions for the time evolution of the metric components (Equation 5.9 and 5.10), we obtain

\[
REGR = -\kappa R + \kappa_1 \Theta
\] (5.20)

where \( R \) is the scalar curvature (Equation 5.7) and \( \Theta \) is the expansion scalar (Equation 5.15). Hence, the REGR equation shows that growth of a disk is not just the expansion of the space, but also a result of curvature dissipation. When we consider that the scalar curvature is a KPZ-type function generalized to 2D space, the notion that growth is both expansion and deposition again emerges, in analogy to the results of Chapter 4.
Measurements of the REGR for a 2D leaf or petal could be made using existing experimental methods \cite{77}. By imprinting a growing tissue with a grid of dots (the landmark method from Chapter 4), the area inside each segment of tissue could be measured over time, thereby allowing the REGR to be calculated.

Finally, the curvature evolution of the disk is represented as a 2D heat map. Regions of positive and negative curvature are mapped to the disk. Regions of positive curvature, when embedded in flat 3D space, will generally look like portions of a sphere, while areas of negative curvature will be buckled and ruffled. Two versions of the heat map will be presented. One is a disk that changes size over time, which has the radial growth information overlayed with curvature information. Another format is the mapping the curvature data onto the unit disk. This does not mean that the disk does not grow; rather, it is the curvature information separated from growth information. In these cases, the radial growth graphs are shown following the curvature maps.

5.5 Numerical Simulation: Ricci flow on the disk

The first 2D simulation we can look at is the evolution of the metric in response to its own curvature, i.e. pure Ricci flow. The equation using the Ricci tensor $R_{ik}$ is:

$$\frac{\partial g_{ik}}{\partial t} = -\kappa R_{ik}. \tag{5.21}$$

Just as before, we can replace the metric $g_{ik}$ with its components $g_{11} = f(r)$ and $g_{22} = r^2 g(r)$, and the Ricci tensor with $R_{g_{ik}}$, where $R$ is given by \ref{5.7}. The evolution equation becomes:

$$\frac{\partial g_{11}}{\partial t} \rightarrow \frac{\partial f}{\partial t} = -\kappa R f \tag{5.22}$$

$$\frac{\partial g_{22}}{\partial t} \rightarrow \frac{\partial g}{\partial t} = -\kappa R g. \tag{5.23}$$
We choose the coupling constant and initial conditions as:

\[
\begin{align*}
\text{Ricci flow coupling} & \quad \kappa = 0.5 \\
 f(t = 0, r) & = 1.0 \\
 g(t = 0, r) & = 0.5 \exp(-2.0(r - \frac{1}{2}r_{\text{max}})^2) + 1.0
\end{align*}
\]

and the results are shown in Figures 5.2 and 5.3.

Since Ricci flow on a disk has not been widely studied, it is important to understand the dynamics of this curvature coupling to the metric before introducing coupling to the velocity field. There are several important points to make about these results.

First, the radius of the disk increases over time, indicating that growth occurs due to Ricci flow alone as anticipated by Equation 5.20. The stagnation of growth is related to curvature values dropping to nearly zero at late times, which itself is due to the dissipation term in \( \partial g_{22}/\partial t \). This result also highlights the numerical stability of the code, which does not pick up oscillations when curvature values hover near zero.

Second, it is important to note that the curvature tends toward zero at late times, but the components of the metric are not trivial. A flat metric can be easily achieved if the components are constant (i.e. do not vary with radius). However, this is not the case at late times in this simulation where both \( g_{11} \) and \( g_{22} \) reach asymptotic solutions that clearly vary with radius.

5.6 Numerical Simulation: The growing disk

We now move on to study an example of growth driven by the velocity field and growth tensor coupling. Coupling to the Ricci tensor remains, and as will be discussed, is a key part of the dynamics of growth.

Starting with a flat disk and a radial logistic velocity field (a generalization of the one used in 1D simulations), the disk evolves under Equations 5.9, 5.10 and 5.13. The coupling
constants and initial conditions are:

| Coupling                  | Value       |
|---------------------------|-------------|
| Ricci flow coupling       | κ = 7.0     |
| Growth tensor coupling    | κ₁ = 5.0    |
| Velocity diffusion coupling| c = 0.05   |

\[
\begin{align*}
    f(t = 0, r) &= 1.0 \\
    g(t = 0, r) &= 1.0 \\
    v^1(t = 0, r) &= 0.01/[1.0 + \exp(-5.0(r - \frac{1}{2} r_{max}))] \\
                   &-0.01/[1.0 + \exp(\frac{5}{2} r_{max})]
\end{align*}
\]

and the results are shown in Figures 5.4 to 5.12

The disk evolves in several notable ways. First, the radius of the disk roughly doubles (Figures 5.5 and 5.6), demonstrating significant growth over what is physiologically a relatively short span of time. The disk growth is approximately linear in time, so it is likely only a portion of the larger sigmoidal pattern of growth that a real leaf would experience over its entire development.

The graphs of perimeter (Figure 5.5) and curvature (Figure 5.4) show that ruffling on the edges will occur at late times for this growth pattern due to the globally negative curvature values and an excess of perimeter compared to a flat disk. Correspondingly, Figures 5.6 and 5.7 all show that the disk grows more at the outer edge than at the center, again indicating the emergence of negative curvature.

Hence, the data indicate that under the conditions of this simulation, a growing, buckled surface will develop from an initially flat disk.

5.7 Numerical Simulation: Modeling the growing cap of *Acetabularia*

A very interesting organism on which to test the assumptions of our model is the algae *Acetabularia* (Figure 1.2). The algae is single-celled, but grows to centimeter length scales.
Moreover, the algae has a cap structure it uses in its reproductive cycle that starts as positively curved, then grows to a flat circular shape, and finally becomes negatively curved. Along with this, it is known that growth occurs in an annulus around the central stem. [59]

*Acetabularia* has a property that is rare in our understanding of dynamical Riemannian geometries. The growth of its cap begins from the top of the stem with a positive curvature everywhere. As it matures its curvature lessens and eventually becomes flat. However it then continues to evolve until the cap’s curvature is completely negative. This is unusual since flat space it often a stable fixed point in for dynamical Riemannian curvatures. For example in pure Ricci flow, in two-dimensions, the Ricci tensor vanishes only if the Riemann tensor vanishes therefore when the manifold is flat $R_{ik} = 0$ implies $\partial g_{ik}/\partial t = 0$. If such a fixed point is reached asymptotically then the curvature cannot not change its sign. Similarly if the energy momentum tensor of general relativity obeys certain “positive energy conditions” (generally the energy density of matter and fields in the space-time is greater than the momentum density which is itself positive), the signature of constant curvature space times cannot change without becoming singular in its evolution [7].

Of course one can engineer a flat thin disk into a positively or negatively curved surface with the appropriate tools, or change the topology of the universe from positive curvature to flat by adding negative energy (Dark Energy), but it would seem that some how some plants are able to do this quite naturally without requiring external manipulation or bizarre physical conditions. Thus *Acetabularia* presents a challenge to any biological models to explain how its cap changes the signature of its curvature through the growth process.

As a first attempt one can ask if the coupling of Ricci flow to the growth tensor is capable of changing the sign of the curvature if the surface has a spatially constant but time dependent curvature. One can then ask what the necessary conditions are on the velocity field to drive the surface past zero curvature. As will be proven below, a constant curvature space cannot change signature, even when driven by a growth tensor.
The simplest non-flat surfaces to describe are those with constant curvature. If the curvature is positive the surface is that of a sphere, if it is zero it is a flat plane and if it is negative the surface can be described by a hyperbolic paraboloid. Constant curvature surfaces have constant Gaussian curvature

\[ K = \frac{1}{\rho_1 \rho_2}. \]

If \( \rho_1 \) and \( \rho_2 \) are the radii of curvatures of the principle directions of the surface, the surface has positive curvature of the radii are measured form the same side of the surface, if one measures the radii from two different sides then the curvature is negative and if one of the radii is infinite then the surface has zero curvature and is flat.

A space of constant Riemannian curvature is isotropic and homogeneous and the line element in two dimensions can be written as:

\[ ds^2 = \frac{1}{1 - Kr^2} dr^2 + r^2 d\theta^2. \] (5.24)

A space of constant curvature has the following relation between the metric and the Riemann tensor:

\[ R_{1212} = K(g_{11}g_{22} - (g_{12})^2) \]

and therefore

\[ R_{ik} = Kg_{ik}. \]

Suppose it is assumed that the curvature is a spatial constant but can evolve over time. Then \( K = K(t) \).

Given the line element above we can identify the metric coefficients;

\[ f = \frac{1}{1 - Kr^2} \quad \text{and} \quad g = 1. \]

Therefore

\[ R_{11} = \frac{K}{1 - Kr^2} \quad \text{and} \quad R_{22} = Kr^2. \]
With a time dependent $K$ the time derivatives of the metric coefficients are
\[
\frac{\partial f}{\partial t} = \frac{\dot{K}r^2}{(1 - K r^2)^2} \quad \text{and} \quad \frac{\partial g}{\partial t} = 0.
\]

Given the metric above connection coefficients are:
\[
\Gamma^1_{11} = \frac{K r}{1 - K r^2}, \quad \Gamma^2_{12} = \frac{1}{r}, \quad \Gamma^1_{22} = r(K r^2 - 1)
\]
and these lead to:
\[
T_{11} = 2f\left(\frac{\partial v}{\partial r} + \Gamma^1_{11} v^1\right)
\]
and
\[
T_{22} = 2r^2\Gamma^2_{12} v^1.
\]

Since the time derivative of $g$ vanishes, the metric evolution equation leads to an expression for $v^1$ in terms of the radial coordinate:
\[
\frac{\partial g}{\partial t} = 0 = -\kappa R_{22} + \kappa_1 T_{22} = -\kappa K r^2 + 2\kappa_1 r^2 \left(\frac{1}{r}\right) v^1
\]
and this leads to :
\[
v^1 = \frac{\kappa}{2\kappa_1} K r.
\]

Therefore the velocity field grows linearly with the radius $r$. The other metric evolution equation is:
\[
\frac{\partial f}{\partial t} = -\kappa R_{11} + \kappa_1 T_{11} = -\kappa \left(\frac{K}{1 - K r^2}\right) + \kappa_1 \left(\frac{2}{(1 - K r^2)}\right) \left[ + \frac{\kappa}{2\kappa_1} K + \frac{K r}{1 - K r^2} \left(-\frac{\kappa}{2\kappa_1} K r\right) \right]
\]
The first two terms cancel exactly and writing the explicit expression for $\partial f/\partial t$ leads to
\[
\frac{\dot{K}r^2}{(1 - K r^2)^2} = \kappa \left(\frac{K^2 r^2}{(1 - K r^2)^2}\right)
\]
which reduces to:
\[
\frac{dK}{dt} = \kappa K^2.
\]

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Using separation of variables and assuming that the constant curvature at $t = 0$ is $K = K_0$, the explicit time dependence for $K(t)$ is

$$K(t) = \frac{K_0}{(1 - \kappa K_0 t)}.$$

Therefore the explicit velocity field required to maintain a time dependent spatially constant curvature should be

$$v^1 = \frac{1}{2} \frac{\kappa}{\kappa_1} \left( \frac{K_0}{1 - \kappa K_0 t} \right) r.$$

If $K$ is initially positive then a sign change cannot occur without the formation of a singularity. If $K < 0$ then the curvature becomes flat asymptotically and if $K = 0$ no evolution occurs. This result is similar to that found for cosmological spacetimes filled with positive energy.

From this discussion we can infer two things about the initial conditions on a model of *Acetabularia* development. The first is that if the sign of the curvature is to change, the idea of constant curvature must be abandoned and a more general approach needs to be taken. However the cap of the *Acetabularia* is close to constant curvature so we will chose as initial conditions for the metric a functional form for the coefficients that is close to that for the constant curvature case. Indeed, closer examination of the the cap structure at early times reveals a concentration of positive curvature near the stem, and a flatter but still positively curved surface toward the outer edge of the cap (Figure 7 in [59]). In the simulation, we choose $f(t = 0, r) = 1.0 + 0.25[1 + \exp(-(r - \frac{1}{2}r_{max})]^{-1}$ and $g(t = 0, r) = 1.0$, which allows the metric to have globally positive curvature, but also spatial variability such that the inner regions of the disk begin with higher positive curvature.

The second inference is that the velocity field cannot be linear if it is to drive a sign change from positive to negative curvature. Indeed, significantly more material needs to be transferred to the outer regions of the tissue in order to flatten out and then buckle the surface. Furthermore, the cap appears to have a localized annular area where most of the growth occurs (Figure 4 in [59]), which hints at a velocity field that is not linear but rather has a
steep gradient across part of the radius that drives the growth in this way. These behaviours were also seen in the previous simulation of a flat disk developing negative curvature over time. We therefore initiate the simulation with a monotonically increasing sigmoidal velocity field.

In summary, to simulate the cap changing curvature we use the following parameters:

|                           |               |
|---------------------------|---------------|
| Ricci flow coupling       | $\kappa = 1.0$ |
| Growth tensor coupling    | $\kappa_1 = 1.0$ |
| Velocity diffusion coupling| $c = 0.01$    |

\[
f(t = 0, r) = 1.0 + 0.25[1 + \exp(-r - \frac{1}{2}r_{\text{max}})]^{-1}
\]

\[
g(t = 0, r) = 1.0
\]

\[
v^1(t = 0, r) = 0.01/[1.0 + \exp(-5.0(r - \frac{1}{2}r_{\text{max}}))]
\]

\[
-0.01/[1.0 + \exp(\frac{5}{2}r_{\text{max}})]
\]

Compared to the flat disk simulation in the previous section, the coupling parameters are smaller but of the same magnitude, and rather than an initially flat metric, we choose a metric that is globally positive but not constant. The velocity field is the same as the previous simulation.

The results can be found in Figures 5.13 to 5.16. The curvature transitions from positive to negative values seamlessly. Also, the radial metric component increases in a localized ring around the center, while the angular metric component increases more at the outer boundary than near the center; both of these behaviours are consistent with biological observations. For brevity, we do not show the deformation tensors given the strong similarities between this simulation and the previous one for a growing disk that develops negative curvature.

5.8 Analysis of 2D simulations with circular symmetry

From the three simulations presented above, several key observations emerge.
First is the role of Ricci flow in disk growth. Ricci flow allows areas of high curvature to dissipate, which contributes to the growth of the disk as the metric expands (and in certain areas contracts) to accommodate a more uniform distribution of curvature. Another important observation is how the velocity field influences growth. As is clear from the two simulations, Ricci flow drives the growth of the disk for some time, but a non-constant velocity field is required to keep growth going.

With an initial logistic velocity field, the growth tensor generally has its highest values where the logistic function is steepest. Hence growth is localized, causing the metric to expand much more wherever the growth tensor has its largest values. This can be seen in the correspondence between the highest values of the growth tensor components (Figure 5.8) with the steepest parts of the velocity field (Figure 5.9). Without Ricci flow, these regions of fast growth lead to the development of a shock front in the metric where growth builds up in one area while neighbouring regions remain stagnant.

At the tissue level, such drastic differences in growth rates between neighbouring regions cause strain. In Figures 5.10 and 5.11 the values of the deformation tensors are shown over time. All the deformations are most pronounced in the area close to the steepest part of the velocity field, and hence the fastest growth. Here, expansion dissipates over time, as does the $\sigma_{22}$ component of the shear tensor, but the $\sigma_{11}$ shear component appears to keep increasing over time. Closer inspection of the same figures also shows that both shear components increase in magnitude at the outer boundary of the disk over time, even though the expansion scalar remains unchanged in this area at late times. These results suggest that although the dynamics are initially driven by the steep velocity field in the mid-region of the disk, the outer edge certainly begins to experience deformations due to the changing geometry of the disk. Note how this is also correlated with the metric tensor components steadily increasing over time at the outer boundary in Figure 5.7 and with the REGR becoming almost uniform everywhere on the disk over time (Figure 5.12), despite the
growth tensor components having localized maxima in the mid-radius regions.

The deformation tensors and REGR also exhibit wave-like motion of the peaks over time. This shows how the underlying reaction-diffusion dynamics affect the global properties of the tissue. Also, similarly to the 1D case, growth has a positive feedback behaviour, with the expansion scalar dissipating over time, yet generating large growth tensor values due to the geometric effects of an expanding metric. The REGR evolution also shows that the globally negative curvature of the disk allows growth to become almost equally distributed over the area of the disk rather than highly localized as the growth tensor dynamics alone would suggest.

The localization of deformations on the disk also suggests that if a disk or tissue experiences rapid growth, it must also have a mechanism to dissipate both its curvature and velocity field. In this phenomenological model, Ricci flow and velocity diffusion provide such a mechanism. When modeled numerically, this dependency manifests itself as a lower threshold on the Ricci flow and velocity diffusion coupling terms for a given value of growth tensor coupling. The lower threshold rises when growth tensor coupling is increased.

Recall from Section 2.9 that the assumption of constant tissue density in space and time leads to the source field \( S(r, \theta, t) \) being directly proportional to the expansion scalar \( \Theta \). Hence, plotting the expansion scalar as in Figure 5.11 will also give us information on how the source of material is distributed through the tissue. Throughout the simulation of disk growth and \textit{Acetabularia} development, the highest values of \( \Theta \) are localized in an annular region of the disk, again consistent with the region of highest growth rates observed in \textit{Acetabularia} measurements. A second feature of note in the simulation is that expansion at late times also influences the regions near the origin, indicating that the source of material becomes more uniformly distributed over time compared to the initial configuration. How these dynamics could be orchestrated by a single-celled organism is a truly intriguing question.

Lacking extensive data on the velocity field of a growing leaf at mm scales and larger,
a circularly symmetric logistic function was assumed as an initial condition. The logistic
function has certain characteristics that strongly influence the growth pattern of the disk.
Having a steep increase at half the radial distance to the edge means that growth will occur
mostly in a ring around the origin, as seen in the metric components of both growing disks
(Figures 5.7 and 5.14). Adopting this initial condition on the velocity field is also consistent
with the mathematical result in Section 5.7 that constant curvature metrics driven by linear
velocity fields cannot experience a change of curvature signature.

The logistic function increases monotonically with radius. In Equation 5.12, we see
that \( T_{22} \) (the angular component of the growth tensor) is proportional to the value of \( v^1 \).
Hence, it can be expected that a monotonically increasing velocity field will lead to \( g(r) \)
also having monotonically increasing values over time (Equation 5.10). This behaviour has
strong implications for the final shape of the disk because the circumference of the disk is
given by \( C = 2\pi\sqrt{g_{22}} = 2\pi r \sqrt{g(r)} \). If \( g(r) \) is greater than unity and increases with radius,
then \( C > 2\pi r \) for every value of \( r \), and must therefore represent a ruffled, negatively curved
surface (Figure 5.5).

An interesting question is what initial conditions might generate a growing disk with
positive curvature. One might start with \( 0 < g(r) < 1 \) to force \( C < 2\pi r \) for all values
of \( r \), however this is somewhat contrived. A monotonically decreasing velocity field would
produce the desired effect on \( T_{22} \), but would generate only shrinking solutions rather than
growing ones. Lastly, there may be an initial configuration of both metric components that
leads to a positive curvature solution.

5.9 Conclusions on 2D simulations with circular symmetry

The data presented here clearly indicate that under the conditions of this simulation, a
growing, ruffled surface will emerge from an initially flat disk due to the development of a
globally negative curvature on the disk. The monotonically increasing velocity field plays a
major role in this behaviour, as does curvature coupling of the metric to its Ricci tensor. Comparing Figure 5.5 with the results in [41] shows that the curvature patterns in our model of disk growth would be physically realizable shapes when embedded in flat 3D space.

A biological model for this type of growth can be found in *Acetabularia*. Two morphologically significant results emerge from our simulation. The first is that a positively curved disk can evolve to a negatively curved disk when driven by a logistic velocity field, thereby reproducing the curvature evolution of the *Acetabularia* cap. The second is that an annulus of growth is observed in the model, which is consistent with the pattern of growth seen in [59].

Building the simplest 2D model of disk growth is the bridge to the next level of complexity: simulations of 2D disks with $\theta$-dependent quantities. These are discussed in the next chapter.

Looking back on the similarities between the 1D and 2D dynamical equations, it is possible to draw comparisons between the two. Both react to an initial logistic velocity field by exhibiting an elongation zone where the velocity field is steepest, and hence the growth tensor is largest. Both require dissipation terms in the metric and velocity fields to prevent shocks from building up as a result of localized growth. Both grow as a result of deposition (metric self-coupling) and material expansion.

The differences between the 1D and 2D models ultimately stem from their different dimensionalities. Even under similar initial conditions, in particular the logistic velocity field, the two systems behave differently. The 2D model has more degrees of freedom in how the metric can evolve, and has two components of the growth tensor (as opposed to one in 1D). This causes the feedback between the velocity field and metric to occur differently than in the 1D case. For instance, the REGR in 2D does not evolve a double-peak behaviour, even though it has contributions from both the curvature and expansion.

These 2D simulations are also the first to have curvature coupling terms. The 1D simulations required a functionally similar term to mimic the effect of curvature since intrinsic
curvature cannot be defined with only one spatial dimension. As seen in the 2D isotropic simulations, it is possible to look at the effects of curvature coupling and velocity field coupling separately and together to determine their relative roles in the dynamics of growth. With the aid of physical deformation tensors, the REGR, perimeter and modular growth calculations, the dynamics of the geometry and velocity field can be interpreted in terms of measurable quantities. To our knowledge, these are the first such calculations made for a 2D disk growth model.
Figure 5.2: Time evolution of the scalar curvature under Ricci flow.
Figure 5.3: Dynamics of the metric under Ricci flow.
Figure 5.4: Time evolution of the scalar curvature of a growing disk.
Figure 5.5: Perimeter of a growing disk over time.

Figure 5.6: Modular radial growth of a growing disk.
Figure 5.7: Time evolution of the metric components of a growing disk.
Figure 5.8: Time evolution of the growth tensor components of a growing disk.
Figure 5.9: Time evolution of the velocity field of a growing disk.
Figure 5.10: Time evolution of the shear tensor components of a growing disk.
Figure 5.11: Time evolution of the expansion scalar of a growing disk.

Figure 5.12: Time evolution of the REGR for a growing disk.
Figure 5.13: Time evolution of the scalar curvature and perimeter of a disk with changing curvature signature.
Figure 5.14: Time evolution of the metric components of a disk with changing curvature signature.
Figure 5.15: Time evolution of the growth tensor components of a disk with changing curvature signature.
Figure 5.16: Time evolution of the velocity field of a disk with changing curvature signature.
Chapter 6

2D Simulations with Anisotropies

6.1 Breaking circular symmetry

In this chapter, simulations of growth involving angular anisotropies will be discussed. Circular symmetry can be broken by introducing $\theta$-dependent initial conditions on the metric or the velocity field.

As in the previous chapter on 2D circularly symmetric solutions, we will proceed in a similar progression of simulations. First, Ricci flow on the disk with angular anisotropies will be studied in order to understand the dynamics of the disk when it is coupled only to its own curvature. Building on these results, we will then introduce a circularly symmetric velocity field to the anisotropic disks. This will allow us to see the first effects of angular anisotropies in the full 2D dynamical equations. Lastly, an initially flat disk will be evolved under the influence of an anisotropic velocity field in a model that results in elongation of the disk along an axis as well as nontrivial curvature patterns.

6.2 Dynamical equations of growth in anisotropic polar coordinates

The equations for growth with angular anisotropy are derived in a similar way to the isotropic 2D equations. A fully general 2D metric in polar coordinates can be written as:

$$g_{ik} = \begin{pmatrix} f(r, \theta) & rh(r, \theta) \\ rh(r, \theta) & r^2 g(r, \theta) \end{pmatrix}$$

(6.1)
The connection coefficients are:

\[
\begin{align*}
\Gamma^1_{11} &= \frac{(2hh_r - f_r g)r + 2h^2 - f_\theta h}{2dr} \\
\Gamma^2_{11} &= \frac{- (2fh_r - f_r h) r + 2fh - ff_\theta}{2dr^2} \\
\Gamma^1_{12} &= \frac{g_r hr + 2gh - f_\theta g}{2d} \\
\Gamma^2_{12} &= \frac{- fg_r r - f_\theta h + 2fg}{2dr} \\
\Gamma^1_{22} &= \frac{gg_r r^2 + (-2gh_\theta + g_\theta h + 2g^2) r}{2d} \\
\Gamma^2_{22} &= \frac{- g_r hr - 2hh_\theta + 2gh + fg_\theta}{2d}
\end{align*}
\]

where \( d = h^2 - fg \).

The only unique Riemann tensor component is:

\[
R_{1212} = \frac{1}{4d}[(2g_r hh_r - 2g_r r - f(g_r)^2) - f_r gg_r) r^2 \\
+((4hh_\theta + 4gh + 2fg_\theta) h_r + 4dh_\theta + 2f_r gh_\theta - 6g_r h^2 \\
+(f_\theta g_r - f_r g_\theta) h + 4f gg_r - 2f_r g^2) r \\
+(2f_\theta h - 4fg) h_\theta - 2df_\theta + (2fg_\theta + 2f_\theta g) h - f_\theta g_\theta - f_\theta^2 g]
\]

(6.3)

The Riemann tensor in 2D has only one component and is related to the scalar curvature \( R \) by:

\[
R_{1212} = R[g_{11}g_{22} - (g_{12})^2].
\]

(6.4)

Since the Ricci tensor can be expressed as \( R_{ik} = Rg_{ik} \), the metric evolution becomes:

\[
\frac{\partial g_{11}}{\partial t} \rightarrow \frac{\partial f}{\partial t} = -\kappa R f + \kappa_1 T_{11} \\
\frac{\partial g_{12}}{\partial t} \rightarrow \frac{\partial h}{\partial t} = -\kappa R h + \kappa_1 \frac{T_{12}}{r} \\
\frac{\partial g_{22}}{\partial t} \rightarrow \frac{\partial g}{\partial t} = -\kappa R g + \kappa_1 \frac{T_{22}}{r^2}
\]

(6.5)
The general expression for the growth tensor is:

\[ T_{ik} = \nabla_i v_k + \nabla_k v_i \]

\[ = g_{kl} \nabla_i v^l + g_{lp} \nabla_k v^p \]  \hspace{1cm} (6.6)

since \( \nabla_i g_{kl} = 0 \).

The specific expressions are:

\[ T_{11} = 2f(v_{,r}^1 + \Gamma_{11}^1 v^1 + \Gamma_{12}^1 v^2) + 2hr(v_{,r}^2 + \Gamma_{11}^2 v^1 + \Gamma_{12}^2 v^2) \]
\[ T_{12} = hr(v_{,r}^1 + \Gamma_{11}^1 v^1 + \Gamma_{12}^1 v^2) + gr^2(v_{,\theta}^2 + \Gamma_{11}^2 v^1 + \Gamma_{12}^2 v^2) \]
\[ + f(v_{,\theta}^1 + \Gamma_{21}^1 v^1 + \Gamma_{22}^1 v^2) + hr(v_{,\theta}^2 + \Gamma_{21}^2 v^1 + \Gamma_{22}^2 v^2) \]
\[ T_{22} = 2hr(v_{,\theta}^1 + \Gamma_{21}^1 v^1 + \Gamma_{22}^1 v^2) + 2gr^2(v_{,\theta}^2 + \Gamma_{12}^2 v^1 + \Gamma_{22}^2 v^2) \]  \hspace{1cm} (6.7)

The velocity equations are:

\[ \frac{\partial v^1}{\partial t} = -\Gamma_{11}^1 v^1 - 2\Gamma_{12}^1 v^1 v^2 - \Gamma_{22}^1 v^2 v^2 - v^1 \nabla_1 v^1 - v^2 \nabla_2 v^1 + c(g_{11}^1 \nabla_1 v^1 + g_{22}^1 \nabla_2 v^1) \]
\[ + cg_{11}^1 \frac{\partial v^1}{\partial r} + \Gamma_{11}^1 v^1 + \Gamma_{12}^1 v^2 + \Gamma_{11}^2 v^1 + \Gamma_{12}^2 v^2 - v^2 [v_{,r}^1 + \Gamma_{11}^2 v^1 + \Gamma_{12}^2 v^2] \]
\[ + cg_{22}^2 \frac{\partial^2 v^1}{\partial \theta^2} + \Gamma_{12}^1 v^1 + \Gamma_{22}^1 v^2 + 2\Gamma_{12}^2 v^1 + 2\Gamma_{22}^2 v^2 - \Gamma_{22}^2 v_{,r}^1 - \Gamma_{22}^2 v_{,\theta}^1 \]
\[ + cg_{22}^2 (\Gamma_{12}^1 (\Gamma_{12}^1 - \Gamma_{22}^1) + \Gamma_{22}^1 (\Gamma_{12}^2 - \Gamma_{11}^1)) v^1 \]  \hspace{1cm} (6.8)

\[ \frac{\partial v^2}{\partial t} = -\Gamma_{11}^2 v^1 - 2\Gamma_{12}^2 v^1 v^2 - \Gamma_{22}^2 v^2 v^2 - v^1 \nabla_1 v^2 - v^2 \nabla_2 v^2 + c(g_{11}^2 \nabla_1 v^2 + g_{22}^2 \nabla_2 v^2) \]
\[ + cg_{11}^1 \frac{\partial v^2}{\partial r} + \Gamma_{11}^2 v^1 + \Gamma_{12}^2 v^2 + \Gamma_{11}^2 v^2 + \Gamma_{12}^2 v^1 - v^2 [v_{,r}^2 + \Gamma_{12}^2 v^1 + \Gamma_{22}^2 v^2] \]
\[ + cg_{11}^1 (\Gamma_{12}^1 (\Gamma_{12}^1 - \Gamma_{22}^2) + \Gamma_{22}^1 (\Gamma_{12}^2 - \Gamma_{11}^1)) v^2 \]
\[ + cg_{22}^2 \frac{\partial^2 v^2}{\partial \theta^2} + \Gamma_{12}^2 v^1 + \Gamma_{22}^2 v^2 + 2\Gamma_{12}^2 v^1 + 2\Gamma_{22}^2 v^2 - \Gamma_{22}^2 v_{,r}^2 - \Gamma_{22}^2 v_{,\theta}^2 \]
\[ + cg_{22}^2 (\Gamma_{12}^2 (\Gamma_{12}^2 - \Gamma_{22}^2) + \Gamma_{22}^1 (\Gamma_{12}^2 - \Gamma_{11}^1)) v^2 \]  \hspace{1cm} (6.9)
6.3 Numerical methods

Again, the coupled tensor equations are now written as a set of nonlinear coupled PDEs. We seek to model Equations 6.5, 6.8 and 6.9.

The equations now include more nonlinear terms due to the presence of the angular velocity, off-diagonal metric terms and the full set of connection coefficients. The equations now also contain diffusion terms in both $r$ and $\theta$. To handle the additional complexity of the $\theta$-dependent equations, we implement a simpler, fully explicit finite differencing scheme. For the second-derivative terms appearing in the scalar curvature, a time averaged scheme for the Laplacian terms is used as discussed in Section 3.2.

The boundary conditions in the anisotropic 2D simulations now require conditions on both the radial and angular coordinates. The most appropriate condition for angular coordinate is a periodic boundary since the solutions can be mapped to the unit circle.

\[
\begin{align*}
\partial_r f(t, r = 0, \theta) &= 0 \\
\partial_r f(t, r = r_{\text{max}}, \theta) &= 0 \\
\partial_r g(t, r = 0, \theta) &= 0 \\
\partial_r g(t, r = r_{\text{max}}, \theta) &= 0 \\
\partial_r h(t, r = 0, \theta) &= 0 \\
\partial_r h(t, r = r_{\text{max}}, \theta) &= 0 \\
f(t, r, \theta = 0) &= f(t, r, \theta = 2\pi) \\
g(t, r, \theta = 0) &= g(t, r, \theta = 2\pi) \\
h(t, r, \theta = 0) &= h(t, r, \theta = 2\pi) \\
v_1(t, r = 0, \theta) &= 0 \\
\partial v_1(t, r = r_{\text{max}}, \theta)/\partial r &= \phi(t, \theta) \\
v_1(t, r, \theta = 0) &= v_1(t, r, \theta = 2\pi)
\end{align*}
\]

The function $\phi(t, \theta)$ is equivalent to $\partial v_1/\partial r$ evaluated at $r = r_{\text{max}} - 1$. 

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6.4 Numerical Simulation: Ricci flow with angular anisotropies

To understand the effect of pure Ricci flow on a metric with angular anisotropies, we study a metric with 4 Gaussian perturbations arranged symmetrically on the metric. These are similar to the Gaussian used in the isotropic 2D case, but are now modulated by a Gaussian in the $\theta$ coordinate.

The 4-Gaussian simulation has parameters:

Ricci flow coupling $\kappa = 0.5$

$$f(t = 0, r) = 5.0 \exp(-2.0(r - \frac{1}{2}r_{\text{max}})^2)$$

$$\times \left[ \exp(-2.0(\theta - \frac{1}{8}\theta_{\text{max}})^2) + \exp(-2.0(\theta - \frac{3}{8}\theta_{\text{max}})^2) \right] + 2.0$$

$$g(t = 0, r) = 2.0 \exp(-2.0(r - \frac{1}{2}r_{\text{max}})^2)$$

$$\times \left[ \exp(-2.0(\theta - \frac{1}{8}\theta_{\text{max}})^2) + \exp(-2.0(\theta - \frac{3}{8}\theta_{\text{max}})^2) \right] + 2.0$$

$$h(t = 0, r) = 0.5 \exp(-2.0(r - \frac{1}{2}r_{\text{max}})^2)$$

$$\times \left[ 0.5 \exp(-2.0(\theta - \frac{1}{8}\theta_{\text{max}})^2) + 0.5 \exp(-2.0(\theta - \frac{3}{8}\theta_{\text{max}})^2) \right] + 1.0$$

Figures 6.1 to 6.4 show the results of these simulations.

Figure 6.2 indicates that the regions with high initial curvature will shrink radially over time, while regions that are initially relatively flat will grow radially. Just like for the isotropic 2D simulations, as the curvature dissipates, growth stagnates and the metric approaches a nontrivial asymptotic configuration, as shown in Figures 6.3 and 6.4.

An interesting difference in the curvature dynamics compared to the isotropic 2D simulations is the final configuration of the scalar curvature. In the isotropic simulations, the center of the disk had negative curvature and the edges were positive. In the anisotropic
simulations, the center is positive while the edges have negative curvature. The cause of this
difference is not known, but is clearly a result of introducing angular anisotropy into the
system.

6.5 Numerical Simulation: Anisotropic growth of a disk

We can also study an initially flat metric under the influence of an anisotropic velocity field.
Here, the velocity field is a logistic function just as in previous simulations, only this time
modulated by a $\theta$-dependent Gaussian function centered on $\theta = \pi$:

- **Ricci flow coupling**  $\kappa = 0.75$
- **Growth tensor coupling**  $\kappa_1 = 0.75$
- **Velocity diffusion coupling**  $c = 0.005$

\[
\begin{align*}
  f(t = 0, r) &= 1.0 \\
  g(t = 0, r) &= 1.0 \\
  h(t = 0, r) &= 0.0 \\
  v^1(t = 0, r) &= (0.01/[1.0 + \exp(-10.0(r - \frac{1}{2}r_{max}))] - 0.01/[1.0 + \exp(5.0r_{max})]) \\
                &\times \exp(-0.1(\theta - \frac{1}{2}\theta_{max})^2) \\
  v^2(t = 0, r) &= 0.0
\end{align*}
\]

Figures 6.5 to 6.16 show the results of these simulations.

The results of growth driven by a radially anisotropic velocity field show that regions of
buckling and ruffling can emerge from an initially flat disk. Highest growth rates at $\theta = \pi$
correspond to the largest values of the growth tensor and velocity field. These also become
the areas of highest curvature. Indeed, the growth tensor for this simulation not only drives
elongation of the disk, but also where and how much curvature the disk develops.
6.6 Analysis of 2D simulations with anisotropy

The results of the two simulations presented in this chapter can be interpreted both geometrically and physically.

The Ricci flow simulation shows that curvature can cause both expansion and contraction of the manifold depending on the local distribution of curvature (Figure 6.2). Globally, the curvature still dissipates over time, flattening the manifold (Figure 6.1).

The expansion, rotation and shear tensors, as well as the REGR, give insight into the physical development of the tissue. As before, the REGR gives a measure of local growth. In a general 2D manifold, the area is now

\[ A = \sqrt{\det(g_{ik})} dr d\theta = \sqrt{fg - h^2r} dr d\theta, \quad (6.10) \]

but the expression for REGR reduces to the same one as in the 2D isotropic case:

\[ \text{REGR} = -\kappa R + \kappa_1 \Theta. \quad (6.11) \]

As before, \( R \) is the scalar curvature and \( \Theta = \nabla_i v^i \) is the expansion scalar.

As can be seen in Figures 6.9 and 6.14, growth has contributions from both the expansion scalar and the curvature term, just as in the 2D isotropic case. The REGR at the outer is edge is clearly higher than at the center in both plots, indicating that a ruffled structure is emerging. Also, the magnitude of REGR along the outer edge at \( \theta = \pi \) is roughly twice that at \( \theta = \pi/2 \), showing that ruffling will be more pronounced along the \( \theta = \pi \) direction and will taper off in parts of the disk that are not growing as fast. This contribution is also clearly related to the anisotropic curvature of the disk, since the expansion scalars do not increase at the outer edge over time.

Unlike the isotropic case in the previous chapter, rotation is now possible. This is a direct consequence of having an angular velocity develop over time (Figure 6.13). Moreover, the angular velocity varies in both the radial and angular directions; indeed, the angular...
velocity and rotation tensor are null at $\theta = \pi$ but are clearly non-zero along $\theta = \pi/2$. This also implies that the shear tensor components will have more structure than in the isotropic case (Figures [6.10] and [6.15]).

The dynamics of the initially flat disk show that growth of the metric is driven by the gradients of the initial velocity field, a result consistent with the simulations in 1D and 2D isotropic cases studied in previous chapters. Overall, the deformations in the anisotropic case are richer than those of previous simulations, yet follow the same broad patterns. Deformations are all localized near the steepest gradient of the velocity field, and with the expansion of the metric, these effects tend to dissipate even as the manifold continues to grow. The expansion scalar (or equivalently in our model, the source field) has similar dynamics to the disk growth and Acetabularia models, indicating that growth will be localized in a deformed annulus around the center of the anisotropic disk.

The results for these simulations are limited due to the less stable numerical methods used to achieve them. However there is strong indication that, just as in the isotropic 2D simulations, growth and curvature feed back onto each other to produce non-trivial geometries. In these models, growth and buckling occur in regions where the velocity field and growth tensor are largest.

Similarly to the isotropic 2D simulations, a lower threshold for velocity diffusion and Ricci flow coupling exists for a given growth tensor coupling. Using the less stable finite differencing scheme in this version puts a strong upper limit on the growth tensor coupling. Again, dissipative terms are required to prevent large differences in growth rates between neighbouring regions that would otherwise excessively strain the tissue.

6.7 Conclusions on 2D simulations in anisotropic polar coordinates

The data presented in this chapter are a prediction of how anisotropic growth may occur in a leaf. Different radial distances along different values of $\theta$ indicate that the embedded
shape of such geometries would be elongated along the $\theta = \pi$ axis. Variations in scalar curvature suggest the tissue would be ruffled or buckled near the growing edge, but flat near the base. Under the influence of an anisotropic velocity field, the simulations show that the disk becomes elongated with nontrivial curvature patterns throughout the disk. To our knowledge, no experimental data in the plant biology or thin disk communities exist to verify these simulated results.
Figure 6.1: Time evolution of the scalar curvature under Ricci flow for a 4-gaussian initial metric.
Figure 6.2: Radial growth of a 4-gaussian initial metric under Ricci flow along different $\theta$ directions.
Figure 6.3: Dynamics for a 4-gaussian initial metric under Ricci flow at $\theta = \pi/4$. 
Figure 6.4: Dynamics for a 4-gaussian initial metric under Ricci flow at $\theta = \pi/2$. 
Figure 6.5: Time evolution of the scalar curvature for an initially flat metric coupled to an anisotropic velocity field.
Radial Growth of a Disk in Different $\theta$ Directions

radius along $\theta = 0$

radius along $\theta = \pi/2$

radius along $\theta = \pi$

Figure 6.6: Radial growth for an initially flat metric coupled to an anisotropic velocity field along different $\theta$ directions.
Figure 6.7: Dynamics of the metric at $\theta = \pi$ for an initially flat metric coupled to an anisotropic velocity field.
Figure 6.8: Growth tensor components and radial velocity field at $\theta = \pi$ for an initially flat metric coupled to an anisotropic velocity field.
Figure 6.9: Expansion scalar and REGR at $\theta = \pi$ for an initially flat metric coupled to an anisotropic velocity field.
Figure 6.10: Shear and rotation tensor components at $\theta = \pi$ for an initially flat metric coupled to an anisotropic velocity field.
Figure 6.11: Dynamics of the metric at $\theta = \pi/2$ for an initially flat metric coupled to an anisotropic velocity field.
Figure 6.12: Growth tensor components at $\theta = \pi/2$ for an initially flat metric coupled to an anisotropic velocity field.
Figure 6.13: Velocity field components at $\theta = \pi/2$ for an initially flat metric coupled to an anisotropic velocity field.
Figure 6.14: Expansion scalar and REGR at $\theta = \pi/2$ for an initially flat metric coupled to an anisotropic velocity field.
Figure 6.15: Shear tensor components at $\theta = \pi/2$ for an initially flat metric coupled to an anisotropic velocity field.
Figure 6.16: Rotation tensor component at $\theta = \pi/2$ for an initially flat metric coupled to an anisotropic velocity field.
Chapter 7

Conclusions and Future Work

7.1 Conclusions on 1D and 2D Simulations

The results presented for 1D and 2D simulations show that biologically relevant growth dynamics occur for a system where the metric, velocity field and mass density are coupled in a way that represents the biology and physics of macroscopic plant leaf growth.

In the 1D case, several features of primary root growth emerge from the mathematical model. Amongst these is a pronounced elongation zone in the root tip, and a double peaked REGR that emphasized the contribution of geometric growth in the model in addition to the effects of a logistic-type velocity field. It was also found that a minimum coupling to dissipation terms in the scale factor and velocity field terms is required to prevent shocks from occurring as neighbouring regions experience different rates of growth.

In the 2D simulations, growth occurs both due to Ricci flow and coupling to the growth tensor. Similarly to the 1D case, a minimum amount of coupling to the Ricci tensor and velocity diffusion terms is needed to counteract the effect of localized growth due to the growth tensor.

Additionally, the 2D simulations show that for a monotonically increasing logistic velocity field, the tissue can transition from globally positive to globally negative curvature. Simulations of a flat disk and an initially positively curved surface show that with a logistic velocity field, the perimeter of the leaf grows outward, becomes ruffled, and has an annular growth zone, just as in the curvature evolution of the Acetabularia cap. Comparing these results to thin disk experiments strongly suggests these are also physically realizable shapes. It is equally conceivable that the elongated and locally curved disks resulting from anisotropic velocity fields are also physically realizable shapes that correspond to leaf shapes found in
nature.

Using a logistic velocity field as an initial condition in the 2D models was an extension of 1D velocity field profile in roots. In the case of *Acetabularia*, this velocity field profile allows the annular growth pattern to arise, and is consistent with the mathematical finding that constant curvature metrics driven by linear velocity fields cannot change the sign of their curvature (Section 5.7).

An important feature of growth that emerges from this model is the dual contributions to the relative elemental growth rate (REGR) from a deposition term and an expansion term. As seen early on, the expansion term can be linked back to a distributed source function that drives the influx of material to the tissue. The deposition term, however, comes from a geometric KPZ reaction-diffusion term in one dimension, and the Ricci flow in two dimensions.

The correspondence of the KPZ term with Ricci flow is striking since they originate in very different contexts. The KPZ term has its origins in a surface growth model, which we generalize to a one dimensional geometric model where growth can occur not just at the surface, but throughout the tissue. Ricci flow is a process that allows a highly curved geometry to evolve continuously to a smoothed geometry. For a two dimensional tissue, it is what allows the tissue’s curvature to change over time. The mathematical similarity between the KPZ equation and Ricci flow resides in quadratic gradient terms appearing as reaction-diffusion processes in both one and two dimensions. Indeed, one can think of Ricci flow as a generalization of the KPZ equation to higher dimensions, under the constraints of being a tensor of the same valence and symmetry as the metric tensor. On the other hand, one can start with Ricci flow in two dimensions and recover a geometric reaction-diffusion term for a one dimensional system that is consistent with the assumptions of local dynamical interactions governing open, driven systems like the KPZ equations.

Characterizing the dynamics of these growth patterns is significant. Geometry is known to
be correlated with nonhomogeneous growth processes, but an understanding of the dynamic feedback between curvature and growth has been lacking. On the theoretical side, this required developing numerical techniques to study Ricci flow applied to disk geometries, then coupling the geometric flow to a velocity field driven by a source of material. Improving this macroscopic and dynamic model of plant growth will require input from the sophisticated plant imaging techniques coming to the fore in experimental biology [5] [76] [77], allowing us to expand our understanding of the biophysics happening at large scales.

### 7.2 Future Directions and Open Questions

The geometric information calculated from the simulations of 1D and 2D growth provides a unique understanding of the physics of a growing plant leaf. However, this information is not easy to interpret and there are many questions left to answer.

The next step in interpreting the data will be to embed the 2D data in flat 3D space. This will allow for a more intuitive way to visualize the curved structures resulting from growth. The mathematics to solve the embedding problem are not trivial, requiring solutions to another set of coupled nonlinear PDEs [55].

Another interesting problem would be to look at the geodesics of the metric as it grows. Are the geodesics linked to stress patterns and venation? A related question is how geodesics evolve under the influence of Ricci flow and growth. For this question, using the Raychaudhuri equations may be an interesting way forward, not least because these are questions usually only posed by cosmologists when studying spacetime dynamics [36].

For the simulations themselves, much more is left to explore. Symmetric, noncircular boundary shapes could be imposed, thereby simulating the shapes of budded plant species whose leaf shape is determined by space constraints within the bud [12].

Other aspects of the mathematics and physics of the model remain open questions. For example, can the full 2D equations be efficiently evolved in a conformally flat space? Such
a formulation of the equations would allow the use of more stable finite differencing models, like in the constrained 2D model. Another geometric question pertains to extending this model to 3D tissues such as those found in animal organs, or diseases like cancer. The mathematics of curvature flows in three spatial dimensions would be more complex than in the 1D and 2D models studied here, but the approach of coupling curvature and growth would still be a valid one to study. Just as in the lower dimensional models, nonhomogenous growth will again lead to stresses in the tissue as well as curved geometries that could alter cell behaviour through mechanotransduction.

Gathering more biological data will also be immensely useful for mapping out the solution space of the equations. Constraints on the parameters and initial conditions of the system that come directly from experiment would help to decide which numerical solutions are biologically relevant. For this, data on the velocity field evolution, macroscopic growth patterns (including measurements of deformation tensors and REGR in 2D tissues), and mechanical properties of leaf tissues would be required. Luckily, the last 5 to 10 years have seen a resurgence in collecting high-resolution digitized data on root and leaf growth that can be applied to many different species of plants 5, 76, 77.

As mentioned in Chapter 4, the shape of a physical non-Euclidean disk in a 3D space is governed by both a target metric (i.e. pure geometry) as well as the physical properties of the material that forms the disk. The question then arises: what is the interplay of geometry with the material properties of the plant tissue as it grows? How do tissue properties evolve over the lifetime of the plant? Does this influence how the geometry of the leaf is ‘interpreted’ in flat 3D space? Experiments on thin disks suggest this would indeed be an important aspect of how a plant leaf forms its curved shape since a disk that grows in a non-uniform way must minimize its stretching and bending energies when its target metric is not flat 62.

The question of uniqueness of individual plants also arises, especially since the equations presented in this work are deterministic. In the current model, individuation could occur by
selecting heterogeneous initial conditions or coupling constants. Since the system is nonlin-
ear, it is possible that this would be one way to generate significant variations in the final
shapes of the plant leaves. Another method to introduce variability in the simulations would
be to add a stochastic noise term to the dynamical equations themselves. This approach
is taken by Kardar, Parisi and Zhang \[37\] in their model of crystal growth. Again, the
nonlinearity of the equations combined with a noise term could allow the system to more
fully explore its solution space, allowing it to choose among several energetically equivalent,
yet morphologically distinct configurations.

Lastly, readers with a plant biology background may ask, where is the auxin? Or for
that matter, any of a number of biochemical processes that enable plant growth. In plant
systems, auxin is a hormone that causes growth wherever it is present, and indeed, this
phenomenological model does not explicitly address where growth comes from. However,
the velocity field, which largely dictates the form of the growth tensor, is an initial condition
that has a profound influence on where growth occurs in the system. Additionally, the
coupling to the growth tensor is a free parameter in this model, which also affects how
receptive the system is to growth; in a more complicated model, this coupling constant
could instead be a function of space and time as well. Hence, it is possible to incorporate
the effects of biochemical and genetic processes into this phenomenological model through
the growth tensor and its coupling strength.

Indeed, plant growth operates on multiple scales, from genetic encoding to tissue level
mechanics. As the authors who studied Acetabularia caps state: 'In sum, morphogenesis
in A. acetabulum may result partly from biophysical constraints arising indirectly from the
general growth process, rather than from the action of genes that code for specific shapes'
\[59\]. Integrating the levels of complexity in biological growth remains a vibrant field of study
that will require expertise from many disciplines of science over the years to come.
Appendix A

Discretization of the Dynamical Equations

In this section, the details of how the dynamical equations used in the models of plant growth become discretized are discussed.

In general, we begin with identifying the diffusive terms in order to apply the Dufort-Frankel method. The remaining differential terms are discretized using spatial central differencing on a single time step.

A.1 1D discretization

In the 1D case, we start with two equations, one for the scale factor time evolution \( g_{11} = f \) and the other for the velocity field time evolution \( v^i = v^1 = v \). These are the same as Equations 4.8 and 4.9 in Chapter 4.

\[
\frac{\partial f}{\partial t} = \frac{\kappa}{2f} \left[ \frac{\partial^2 f}{\partial x^2} - \frac{1}{f} \left( \frac{\partial f}{\partial x} \right)^2 \right] + \kappa_1 \left( 2f \frac{\partial v}{\partial x} + v \frac{\partial f}{\partial x} \right)
\]

\[
\frac{\partial v}{\partial t} = -\frac{1}{2f} \left( \frac{\partial f}{\partial x} \right) v^2 - v \left[ \frac{\partial v}{\partial x} + \frac{1}{2f} \left( \frac{\partial f}{\partial x} \right) v \right] + c \left[ \frac{1}{f} \frac{\partial^2 v}{\partial x^2} + \frac{1}{2f^2} \left( \frac{\partial^2 f}{\partial x^2} v + \frac{\partial f}{\partial x} \frac{\partial v}{\partial x} - \frac{1}{f} \left( \frac{\partial f}{\partial x} \right)^2 v \right) \right].
\]

Both equations have a diffusive term as well as a series of nonlinear differential terms \( N \) which can be summarized in the general form:

\[
\frac{\partial u}{\partial t} = D \frac{\partial^2 u}{\partial x^2} + N \left( u, \frac{\partial u}{\partial x}, w, \frac{\partial w}{\partial x} \right).
\]

In order to discretize this type of PDE, we take the following approach:

- Apply the Dufort-Frankel discretization to the second-order spatial derivative.
• Apply a central differencing to the first-order time derivatives.

• Apply a central differencing to any first-order spatial derivatives.

• Evaluate functional values at \((x = j\Delta x, t = n\Delta t)\).

For the scale factor, the discretized function will be:

\[
\frac{f_j^{n+1} - f_j^{n-1}}{2\Delta t} \sim \frac{\kappa}{2f_j^n} \left( \frac{f_{j+1}^n - f_{j+1}^{n+1} - f_{j-1}^{n+1} + f_{j-1}^n}{(\Delta x)^2} \right) - \frac{\kappa}{2(f_j^n)^2} \left( \frac{f_{j+1}^n - f_{j-1}^n}{2\Delta x} \right)^2 \\
+ \kappa_1 \left( 2f_j^n \frac{v_{j+1}^n - v_{j-1}^n}{2\Delta x} + v_j^n \frac{f_{j+1}^n - f_{j-1}^n}{2\Delta x} \right).
\] (A.4)

A similar discretization of the velocity equation can also be done. Straightforward algebra then allows for isolating \(f_j^{n+1}\) and \(v_j^{n+1}\), which are the values of the scale factor and velocity field at the future time step \(t = (n + 1)\Delta t\) and spatial step \(x = j\Delta x\).

When implementing this algorithm, the discretized functional values of \(f\) and \(v\) can be stored in arrays and a sequential calculation of \(f_j^{n+1}\) and \(v_j^{n+1}\) is then performed using the existing functional values at \(t = n\Delta t\) and \(t = (n - 1)\Delta t\).

A.2 2D isotropic discretization

For the 2D isotropic equations, the discretization method is entirely analogous to the one presented above for 1D systems.

The equations to solve are now:
where the scalar curvature $R$ is given by:

$$R = -\frac{1}{4d^2} \left( 2g_{rr}fg - f(g_r)^2 - f_rgg_r + \frac{1}{r} (4f gg_r - 2f_r g^2) \right)$$  (A.6)

As before, $g_r = \partial g / \partial r$ and $g_{rr} = \partial^2 g / \partial r^2$.

There are more terms than before, but the equations are still of the form [A.3] so we apply the same algorithm as for the 1D equations. In fact, the evolution equation for $g_{11}$ is no longer diffusive, since $R$ contains second spatial derivatives of $g_{22}$ only.

### A.3 2D anisotropic discretization

The full 2D equation set as presented in Chapter [6] require a slightly modified discretization method because of the mixed second derivatives found in the Riemann tensor:

$$R_{1212} = \frac{1}{4d^2} \left[ (2g_r h h, r - 2g_{rr} d - f(g_r)^2 - f_r gg_r) r^2 ight.$$  

$$+ \left( (-4h h, \theta + 4g h + 2f g, \theta) h, r + 4h, r \theta + 2f, r g h, \theta - 6g, h^2 ight) r^2 
+ (f, \theta g, r - f, r g, \theta) h + 4f gg_r - 2f_r g^2) r $$

$$+ (2f, h - 4f g) h, \theta - 2f, \theta \theta + (2f g, \theta + 2f, \theta g) h - f, \theta g, \theta - f^2 g \right].$$  (A.7)
Additionally, there are significantly more terms to calculate in this set of equations. To make the numerical simulation easier to implement, and therefore less prone to programmer error, we use a time-averaged Laplacian in place of the Dufort-Frankel discretization as discussed in Chapter 3:

\[
\frac{u_j^{n+1} - u_j^{n-1}}{2\Delta t} = D \frac{1}{2} \left( \frac{u_{j+1}^n - 2u_j^n + u_{j-1}^n}{(\Delta x)^2} + \frac{u_{j+1}^{n-1} - 2u_j^{n-1} + u_{j-1}^{n-1}}{(\Delta x)^2} \right). \tag{A.8}
\]

Aside from this change, the algorithm to discretize the equations is the same as before. The goal is to algebraically isolate the discretized differential equations for the future time step values \( f_j^{n+1}, h_j^{n+1}, g_j^{n+1} \) representing the metric components, and similarly for the two velocity components \( (v^1)_j^{n+1} \) and \( (v^2)_j^{n+1} \).

Another complicating factor in the full 2D simulations is a second spatial dimension, the angle \( \theta \), that must be discretized. In implementing this numerical simulation, we must therefore use a 2D array for each functional value, as well as for any calculated values such as connection coefficients and the curvature tensor. The use of 2D arrays instead of 1D arrays as before significantly increases computing time for these simulations.
Appendix B

Supplementary numerical simulations of 2D anisotropic growth

B.1 The interaction of an initially anisotropic metric with a circularly symmetric velocity field

Anisotropic growth can be studied by introducing a circularly symmetric velocity field to the anisotropic initial metrics studied in Chapter 6.

For a 2-Gaussian initial metric:

\begin{align*}
\text{Ricci flow coupling} & \quad \kappa = 0.75 \\
\text{Growth tensor coupling} & \quad \kappa_1 = 0.75 \\
\text{Velocity diffusion coupling} & \quad c = 0.01
\end{align*}

\begin{align*}
 f(t = 0, r) &= 5.0 \exp(-2.0(r - \frac{1}{2} r_{max})^2) \times [\exp(-2.0(\theta - \frac{1}{4} \theta_{max})^2) \\
 &\quad + \exp(-2.0(\theta - \frac{3}{4} \theta_{max})^2)] + 2.0 \\
 g(t = 0, r) &= 2.0 \exp(-2.0(r - \frac{1}{2} r_{max})^2) \times [\exp(-2.0(\theta - \frac{1}{4} \theta_{max})^2) \\
 &\quad + \exp(-2.0(\theta - \frac{3}{4} \theta_{max})^2)] + 2.0 \\
 h(t = 0, r) &= 0.5 \exp(-2.0(r - \frac{1}{2} r_{max})^2) \times [0.5 \exp(-2.0(\theta - \frac{1}{4} \theta_{max})^2) \\
 &\quad + 0.5 \exp(-2.0(\theta - \frac{3}{4} \theta_{max})^2)] + 1.0 \\
 v^1(t = 0, r) &= 0.01/[1.0 + \exp(-5.0(r - \frac{1}{2} r_{max})]) - 0.01/[1.0 + \exp(\frac{5}{2} r_{max})] \\
 v^2(t = 0, r) &= 0.0
\end{align*}

For a 4-Gaussian initial metric:
Ricci flow coupling \( \kappa = 0.75 \)

Growth tensor coupling \( \kappa_1 = 0.75 \)

Velocity diffusion coupling \( c = 0.01 \)

\[
\begin{align*}
  f(t=0, r) &= 5.0 \exp(-2.0(r - \frac{1}{2} r_{\text{max}})^2) \\
  &\quad \times \left[ \exp(-2.0(\theta - \frac{1}{8} \theta_{\text{max}})^2) + \exp(-2.0(\theta - \frac{3}{8} \theta_{\text{max}})^2) \right. \\
  &\quad + \left. \left( \exp(-2.0(\theta - \frac{5}{8} \theta_{\text{max}})^2) + \exp(-2.0(\theta - \frac{7}{8} \theta_{\text{max}})^2) \right) + 2.0 \right] \\
  g(t=0, r) &= 2.0 \exp(-2.0(r - \frac{1}{2} r_{\text{max}})^2) \\
  &\quad \times \left[ \exp(-2.0(\theta - \frac{1}{8} \theta_{\text{max}})^2) + \exp(-2.0(\theta - \frac{3}{8} \theta_{\text{max}})^2) \right. \\
  &\quad + \left. \left( \exp(-2.0(\theta - \frac{5}{8} \theta_{\text{max}})^2) + \exp(-2.0(\theta - \frac{7}{8} \theta_{\text{max}})^2) \right) + 2.0 \right] \\
  h(t=0, r) &= 0.5 \exp(-2.0(r - \frac{1}{2} r_{\text{max}})^2) \\
  &\quad \times \left[ 0.5 \exp(-2.0(\theta - \frac{1}{8} \theta_{\text{max}})^2) + 0.5 \exp(-2.0(\theta - \frac{3}{8} \theta_{\text{max}})^2) \right. \\
  &\quad + \left. \left( 0.5 \exp(-2.0(\theta - \frac{5}{8} \theta_{\text{max}})^2) + 0.5 \exp(-2.0(\theta - \frac{7}{8} \theta_{\text{max}})^2) \right) + 1.0 \right] \\
  v^1(t=0, r) &= 0.01/[1.0 + \exp(-5.0(r - \frac{1}{2} r_{\text{max}}))] - 0.01/[1.0 + \exp(\frac{5}{2} r_{\text{max}})] \\
  v^2(t=0, r) &= 0.0
\end{align*}
\]

Figures B.1 to B.14 show the results of these simulations.

When a circularly symmetric velocity field is coupled to the dynamics of an initially curved surface, the stagnation in radial growth seen in Section 6.4 is reversed. Even the shrinking solutions for the 4-Gaussian at \( \theta = \pi/4 \) are reversed dramatically (see Figure 6.2). Additionally, the introduction of a circularly symmetric velocity field seems to smooth out the curvature of the metric more than just the action of the Ricci flow by itself (compare Figure 6.1 with Figure B.8). This again indicates that a monotonically increasing velocity field drives growth beyond what Ricci flow alone can accomplish.
Another important feature of the data is that the angular velocity $v^2$ starts having behaviour. This is not possible in isotropic simulations, and shows the influence that the metric has on the velocity field.
Figure B.1: Time evolution of the scalar curvature for a 2-gaussian initial metric and a circularly symmetric velocity field.
Figure B.2: Radial growth of a 2-gaussian initial metric and a circularly symmetric velocity field along different $\theta$ directions.
Figure B.3: Modular growth along different $\theta$ directions for a 2-gaussian initial metric and a circularly symmetric velocity field.
Figure B.4: Dynamics of the metric and velocity field at $\theta = \pi/4$ for a 2-gaussian initial metric and a circularly symmetric velocity field.
Figure B.5: Dynamics of the metric and velocity field at $\theta = \pi/2$ for a 2-gaussian initial metric and a circularly symmetric velocity field.
Figure B.6: Deformation tensors at $\theta = \pi/4$ for a 2-gaussian initial metric and a circularly symmetric velocity field.
Figure B.7: Deformation tensors at $\theta = \pi/2$ for a 2-gaussian initial metric and a circularly symmetric velocity field.
Figure B.8: Time evolution of the scalar curvature for a 4-gaussian initial metric and a circularly symmetric velocity field.
Figure B.9: Radial growth of a 4-gaussian initial metric and a circularly symmetric velocity field along different θ directions.
Figure B.10: Modular growth at different $\theta$ directions for a 4-gaussian initial metric and a circularly symmetric velocity field.
Figure B.11: Dynamics of the metric and velocity field at $\theta = \pi/4$ for a 4-gaussian initial metric and a circularly symmetric velocity field.
Figure B.12: Dynamics of the metric and velocity field at $\theta = \pi/2$ for a 4-gaussian initial metric and a circularly symmetric velocity field.
Figure B.13: Deformation tensors at $\theta = \pi/4$ for a 4-gaussian initial metric and a circularly symmetric velocity field.
Figure B.14: Deformation tensors at $\theta = \pi/2$ for a 4-gaussian initial metric and a circularly symmetric velocity field.
Appendix C

Make your own curved geometry!
1. Cut out circle.
2. Cut into the circle.
3. Cut out wedge.
4. Glue grey parts together.
4. Glue grey parts together.
Bibliography

[1] Harold Abelson and Andrea DiSessa. *Turtle geometry: The computer as a medium for exploring mathematics*. MIT press, 1986.

[2] Megan Andriankaja, Stijn Dhondt, Stefanie De Bodt, Hannes Vanhaeren, Frederik Coppens, Liesbeth De Milde, Per Mühlenbock, Aleksandra Skirycz, Nathalie Gonzalez, Gerrit TS Beemster, et al. Exit from proliferation during leaf development in arabidopsis thaliana: A not-so-gradual process. *Developmental Cell*, 22(1):64–78, 2012.

[3] B Audoly and A Boudaoud. Self-similar structures near boundaries in strained systems. *Physical Review Letters*, 91(8), Aug 22 2003.

[4] George S Avery Jr. Structure and development of the tobacco leaf. *American Journal of Botany*, pages 565–592, 1933.

[5] Paramita Basu and Anupam Pal. A new tool for analysis of root growth in the spatio-temporal continuum. *New Phytologist*, 195(1):264–274, 2012.

[6] Gerrit TS Beemster and Tobias I Baskin. Analysis of cell division and elongation underlying the developmental acceleration of root growth in arabidopsis thaliana. *Plant Physiology*, 116(4):1515–1526, 1998.

[7] Peter G Bergmann and Venzo De Sabbata. *Topological properties and global structure of space-time*. Springer, 2013.

[8] Janet Braam. In touch: plant responses to mechanical stimuli. *New Phytologist*, 165(2):373–389, 2005.

[9] Thibaut Brunet, Adrien Bouclet, Padra Ahmadi, Démosthène Mitrossilis, Benjamin Driquez, Anne-Christine Brunet, Laurent Henry, Fanny Serman, Gaëlle Béalle, Chris-
tine Ménager, et al. Evolutionary conservation of early mesoderm specification by mechanotransduction in bilateria. *Nature Communications*, 4, 2013.

[10] Andrés Chavarría-Krauser, Kerstin A Nagel, Klaus Palme, Ulrich Schurr, Achim Walter, and Hanno Scharr. Spatio-temporal quantification of differential growth processes in root growth zones based on a novel combination of image sequence processing and refined concepts describing curvature production. *New Phytologist*, 177(3):811–821, 2008.

[11] E Coen, AG Rolland-Lagan, M Matthews, JA Bangham, and P Prusinkiewicz. The genetics of geometry. *Proceedings of the National Academy of Sciences*, 101(14):4728–4735, Apr 6 2004.

[12] Etienne Couturier, Sylvain Courrech Du Pont, and Stéphane Douady. A global regulation inducing the shape of growing folded leaves. *PloS One*, 4(11):e7968, 2009.

[13] Anirvan Dasgupta, Hemwati Nandan, and Sayan Kar. Kinematics of flows on curved, deformable media. *International Journal of Geometric Methods in Modern Physics*, 6(04):645–666, 2009.

[14] SJ Day and PA Lawrence. Measuring dimensions: the regulation of size and shape. *Development*, 127(14):2977–2987, Jul 2000.

[15] Julien Dervaux and Martine Ben Amar. Mechanical instabilities of gels. *Annu. Rev. Condens. Matter Phys.*, 3(1):311–332, 2012.

[16] Julien Dervaux and Martine Ben Amar. Morphogenesis of growing soft tissues. *Physical Review Letters*, 101(6), Aug 8 2008.

[17] Ray d’Inverno. *Introducing Einstein’s Relativity*. Clarendon Press, Oxford, 1992.

[18] Masao Doi. *Soft Matter Physics*. Oxford University Press, 2013.
[19] Petra M Donnelly, Dario Bonetta, Hirokazu Tsukaya, Ronald E Dengler, and Nancy G Dengler. Cell cycling and cell enlargement in developing leaves of arabidopsis. *Developmental Biology*, 215(2):407–419, 1999.

[20] Efi Efrati, Eran Sharon, and Raz Kupferman. The metric description of elasticity in residually stressed soft materials. *Soft Matter*, 9(34):8187–8197, 2013.

[21] Ralph O Erickson and Wendy Kuhn Silk. The kinematics of plant growth. *Scientific American*, 242(5):134–151, 1980.

[22] Ray F. Evert and Susan E. Eichhorn. *Raven Biology of Plants*. W.H. Freeman and Company, 2013.

[23] Paul B Green. Transductions to generate plant form and pattern: an essay on cause and effect. *Annals of botany*, 78(3):269–281, 1996.

[24] PB Green, CS Steele, and SC Rennich. Phyllotactic patterns: A biophysical mechanism for their origin. *ANNALS OF BOTANY*, 77(5):515–527, MAY 1996.

[25] Peter J Gregory. *Plant roots: growth, activity and interactions with the soil*. John Wiley & Sons, 2008.

[26] Olivier Hamant. Widespread mechanosensing controls the structure behind the architecture in plants. *Current opinion in plant biology*, 16(5):654–660, 2013.

[27] Lionel G Harrison, Richard J Adams, and David M Holloway. Dynamic regulation of growing domains for elongating and branching morphogenesis in plants. *Biosystems*, 109(3):488–497, 2012.

[28] Stephen Hawking. *God created the integers: The mathematical breakthroughs that changed history*. Running Press, 2007.
[29] Z Hejnowicz and J Karczewski. Modeling of meristematic growth of root apices in a natural coordinate system. *American Journal of Botany*, pages 309–315, 1993.

[30] Z Hejnowicz and John A Romberger. Growth tensor of plant organs. *Journal of Theoretical Biology*, 110(1):93–114, 1984.

[31] Z Hejnowicz and A Sievers. Tissue stresses in organs of herbaceous plants. i. *Journal of Experimental Botany*, 46(289):1035–1035, 1995.

[32] David W Henderson and Daina Taimina. Crocheting the hyperbolic plane. *The Mathematical Intelligencer*, 23(2):17–28, 2001.

[33] Lars Hufnagel, Aurelio A Teleman, Hervé Rouault, Stephen M Cohen, and Boris I Shraiman. On the mechanism of wing size determination in fly development. *Proceedings of the National Academy of Sciences*, 104(10):3835–3840, 2007.

[34] Johannes Jaeger, David Irons, and Nick Monk. Regulative feedback in pattern formation: towards a general relativistic theory of positional information. *Development*, 135(19):3175–3183, 2008.

[35] Gareth Wyn Jones and S. Jonathan Chapman. Modeling Growth in Biological Materials. *SIAM Review*, 54(1):52–118, 2012.

[36] Sayan Kar and Soumitra Sengupta. The raychaudhuri equations: A brief review. *Pramana*, 69(1):49–76, 2007.

[37] Mehran Kardar. *Statistical physics of fields*. Cambridge University Press, 2007.

[38] Mehran Kardar, Giorgio Parisi, and Yi-Cheng Zhang. Dynamic scaling of growing interfaces. *Physical Review Letters*, 56(9):889, 1986.

[39] Guy Karlebach and Ron Shamir. Modelling and analysis of gene regulatory networks. *Nature Reviews Molecular Cell Biology*, 9(10):770–780, 2008.
[40] Richard Kennaway, Enrico Coen, Amelia Green, and Andrew Bangham. Generation of diverse biological forms through combinatorial interactions between tissue polarity and growth. *PLoS Computational Biology*, 7(6):e1002071, 2011.

[41] Yael Klein, Efi Efrati, and Eran Sharon. Shaping of elastic sheets by prescription of non-Euclidean metrics. *Science*, 315(5815):1116–1120, Feb 23 2007.

[42] Shigeru Kondo and Takashi Miura. Reaction-diffusion model as a framework for understanding biological pattern formation. *Science*, 329(5999):1616–1620, 2010.

[43] Erika E Kuchen, Samantha Fox, Pierre Barbier de Reuille, Richard Kennaway, Sandra Bensmihen, Jerome Avondo, Grant M Calder, Paul Southam, Sarah Robinson, Andrew Bangham, et al. Generation of leaf shape through early patterns of growth and tissue polarity. *Science*, 335(6072):1092–1096, 2012.

[44] Dorota Kwiatkowska. Structural integration at the shoot apical meristem: models, measurements, and experiments. *American Journal of Botany*, 91(9):1277–1293, 2004.

[45] Benoît Landrein and Olivier Hamant. How mechanical stress controls microtubule behavior and morphogenesis in plants: history, experiments and revisited theories. *The Plant Journal*, 75(2):324–338, 2013.

[46] Gabriele B Monshausen and Simon Gilroy. Feeling green: mechanosensing in plants. *Trends in cell biology*, 19(5):228–235, 2009.

[47] Jerzy Nakielski. The tensor-based model for growth and cell divisions of the root apex. i. the significance of principal directions. *Planta*, 228(1):179–189, 2008.

[48] Utpal Nath, Brian CW Crawford, Rosemary Carpenter, and Enrico Coen. Genetic control of surface curvature. *Science*, 299(5611):1404–1407, 2003.

[49] Karl J Niklas. *Plant biomechanics: an engineering approach to plant form and function*. University of Chicago press, 1992.
[50] Richard H Pletcher, John C Tannehill, and Dale Anderson. *Computational Fluid Mechanics and Heat Transfer*. CRC Press, 2012.

[51] Przemyslaw Prusinkiewicz and Pierre Barbier de Reuille. Constraints of space in plant development. *Journal of Experimental Botany*, 61(8):2117–2129, May 2010.

[52] Kenneth Franklin Riley, Michael Paul Hobson, and Stephen John Bence. *Mathematical methods for physics and engineering: a comprehensive guide*. Cambridge University Press, 2006.

[53] Edward K Rodriguez, Anne Hoger, and Andrew D McCulloch. Stress-dependent finite growth in soft elastic tissues. *Journal of Biomechanics*, 27(4):455–467, 1994.

[54] Anne-Gaëlle Rolland-Lagan, J Andrew Bangham, and Enrico Coen. Growth dynamics underlying petal shape and asymmetry. *Nature*, 422(6928):161–163, 2003.

[55] Joseph D Romano and Richard H Price. Embedding initial data for black-hole collisions. *Classical and Quantum Gravity*, 12(3):875, 1995.

[56] Elisabet Romero, Ramunas Augulis, Vladimir I Novoderezhkin, Marco Ferretti, Jos Thieme, Donatas Zigmantas, and Rienk Van Grondelle. Quantum coherence in photosynthesis for efficient solar-energy conversion. *Nature Physics*, 10(9):676–682, 2014.

[57] Jessica Rosenkrantz and Jesse Louis-Rosenberg. Kinematics dress, 2014.

[58] Takami Saito, Kouichi Soga, Takayuki Hoson, and Ichiro Terashima. The bulk elastic modulus and the reversible properties of cell walls in developing quercus leaves. *Plant and cell physiology*, 47(6):715–725, 2006.

[59] Kyle A Serikawa and Dina F Mandoli. An analysis of morphogenesis of the reproductive whorl of acetabularia acetabulum. *Planta*, 207(1):96–104, 1998.
[60] E Sharon, M Marder, and HL Swinney. Leaves, flowers and garbage bags: Making waves. *American Scientist*, 92(3):254–261, May-Jun 2004.

[61] E Sharon, B Roman, M Marder, GS Shin, and HL Swinney. Mechanics: Buckling cascades in free sheets - Wavy leaves may not depend only on their genes to make their edges crinkle. *Nature*, 419(6907):579, Oct 10 2002.

[62] Eran Sharon and Efi Efrati. The mechanics of non-Euclidean plates. *Soft Matter*, 6(22):5693–5704, 2010.

[63] Eran Sharon, Benoit Roman, and Harry L. Swinney. Geometrically driven wrinkling observed in free plastic sheets and leaves. *Physical Review E*, 75(4, 2), Apr 2007.

[64] Robert E Sharp, Wendy Kuhn Silk, and Theodore C Hsiao. Growth of the maize primary root at low water potentials i. spatial distribution of expansive growth. *Plant Physiology*, 87(1):50–57, 1988.

[65] Wendy K Silk and Marie-Béatrice Bogeat-Triboulot. Deposition rates in growing tissue: Implications for physiology, molecular biology, and response to environmental variation. *Plant and Soil*, 374(1-2):1–17, 2014.

[66] Wendy Kuhn Silk and Ralph O Erickson. Kinematics of plant growth. *Journal of Theoretical Biology*, 76(4):481–501, 1979.

[67] Anthony James Merrill Spencer. *Continuum mechanics*. Courier Corporation, 2004.

[68] Szymon Stoma, Mikael Lucas, Jérôme Chopard, Marianne Schaedel, Jan Traas, and Christophe Godin. Flux-based transport enhancement as a plausible unifying mechanism for auxin transport in meristem development. *PLoS Computational Biology*, 4(10):e1000207, 2008.

[69] Keiko Sugimoto-Shirasu and Keith Roberts. “big it up”: endoreduplication and cell-size control in plants. *Current Opinion in Plant Biology*, 6(6):544–553, 2003.
[70] Lincoln Taiz, Eduardo Zeiger, Ian Max Möller, and Angus Murphy. *Plant Physiology and Development*. Sinauer Associates, Incorporated, 2015.

[71] D.W. Thompson. *On Growth and Form*. Cambridge University Press, 1917.

[72] Nathan Tompkins, Ning Li, Camille Girabawe, Michael Heymann, G Bard Ermentrout, Irving R Epstein, and Seth Fraden. Testing turing’s theory of morphogenesis in chemical cells. *Proceedings of the National Academy of Sciences*, 111(12):4397–4402, 2014.

[73] Peter Topping. *Lectures on the Ricci flow*, volume 325. Cambridge University Press, 2006.

[74] Alan Mathison Turing. The chemical basis of morphogenesis. *Philosophical Transactions of the Royal Society of London. Series B, Biological Sciences*, 237(641):37–72, 1952.

[75] Corine M van der Weele, Hai S Jiang, Krishnan K Palaniappan, Viktor B Ivanov, Kannappan Palaniappan, and Tobias I Baskin. A new algorithm for computational image analysis of deformable motion at high spatial and temporal resolution applied to root growth. roughly uniform elongation in the meristem and also, after an abrupt acceleration, in the elongation zone. *Plant Physiology*, 132(3):1138–1148, 2003.

[76] A Walter, H Spies, S Terjung, R Küsters, N Kirchgeßner, and U Schurr. Spatio-temporal dynamics of expansion growth in roots: automatic quantification of diurnal course and temperature response by digital image sequence processing. *Journal of Experimental Botany*, 53(369):689–698, 2002.

[77] Lisheng Wang, Simon T Beyer, Quentin CB Cronk, and Konrad Walus. Delivering high-resolution landmarks using inkjet micropatterning for spatial monitoring of leaf expansion. *Plant methods*, 7(1):1–10, 2011.