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VirtualLeaf: an Open Source framework for cell-based modeling of plant tissue growth and development

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Abstract

Plant organs, including leaves and roots, develop by means of a multi-level crosstalk between gene regulation, patterned cell division and cell expansion, and tissue mechanics. The multi-level regulatory mechanisms complicate classic molecular genetics or functional genomics approaches to biological development, because these methodologies implicitly assume a direct relation between genes and traits at the level of the whole plant or organ. Instead, understanding gene function requires insight into the role of gene products in regulatory networks, the conditions of gene expression, etc. This interplay is impossible to understand intuitively. Mathematical and computer modeling allows researchers to design new hypotheses and produce experimentally testable insights. However, the required mathematics and programming experience makes modeling poorly accessible to experimental biologists. Problem-solving environments provide biologically-intuitive in silico objects ("cells", "regulation networks") required for setting up a simulation, and present those to the user in terms of familiar, biological terminology. Here, we introduce the cell-based computer-modeling framework VirtualLeaf for plant tissue morphogenesis. The current version defines a set of biologically-intuitive C++ objects, including cells, cell walls, and diffusing and reacting chemicals, that provide useful abstractions for building biological simulations of developmental processes. We present a step-by-step introduction to building models with the VirtualLeaf, providing basic example models of leaf venation and meristem development. VirtualLeaf-based models provide a means for plant researchers to analyze the function of developmental genes in the context of the biophysics of growth and patterning. The VirtualLeaf is an ongoing Open Source software project (http://virtualleaf.googlecode.com) that runs on Windows, Mac and Linux.

Keywords: plant development; organ growth; cell division; cell growth; mathematical modeling; auxin
**Introduction**

Plant scientists have gathered a wealth of knowledge about plant growth and plant genetics. Yet, it is still impossible to predict by which mechanisms genetic changes affect plant growth and development. To unravel how genetic information determines the morphogenesis of plants, analyzing and reconstructing the dynamics of the genetic regulatory networks (Gonzalez et al., 2009) is only the first step (Merks and Glazier, 2005). To mechanistically predict morphogenesis from gene function, it is essential to have a detailed understanding not only of genetics, but also of a) how genetic networks regulate cell behaviors, b) how cell behaviors lead to tissue growth and patterning, and c) how the tissue level phenomena, including spatial patterns (Swarup et al. 2005, Jönsson et al., 2006; Bayer et al., 2009) and strains and stresses (Green, 1999; Hamant et al., 2008) induce responses at the molecular level. How these multilevel feedbacks between levels of organization produce biological function and form is perhaps the most central question in systems biology (Noble, 2006).

Traditionally, biologists are familiar with presenting and developing hypotheses using block-and-arrow diagrams, symbolically representing identified and hypothetical gene interactions. However, despite their frequent use, it is not possible to understand the behavior of gene networks just by reading such diagrams, except for very straightforward cases. Mathematical models and computational models can bring the gene networks represented by block-and-arrow diagrams to “life”, and can explain potential dynamic behaviors of the network even where precise quantitative information is unavailable. In this way, the biologist may discover that the hypothetical biochemical network cannot exhibit the expected behavior, or that it occurs only for biologically unfeasible values of the reaction rates or binding constants. The researcher can then propose new interactions or system components and add them to the model to check if the revised network better reproduces the observations the original diagram was attempting to explain. Subsequent experiments will then test the validity of the new elements in the hypothesis and can reveal a set of new observations that will be used to refine the mathematical model. This iterative approach of alternating experimental steps and model refinements is known as the systems biology cycle (Kitano, 2002). Mathematical and computational modeling support experimental biologists to test alternative hypotheses for internal “consistency” and helps them fine-tune their ideas.

A typical systems biology approach is taken in analyzing the de novo patterning of root hairs, and in analyzing the regular arrangement of trichomes in the leaf epidermis of *Arabidopsis*. These two processes are controlled by the same molecular regulators (Schnittger et al., 1999; Pesch and Hülskamp, 2004; Kang et al., 2009). Based on a series of knock-out experiments, an initial “block-and-arrow” model for the interaction of the transcription factors TTG, GL1 and TRY was
proposed (Schnittger et al., 1999). By applying a local activation and lateral inhibition model for pattern formation, it was predicted that TRY is activated by GL1 and/or TTG and inhibits GL1 and/or TTG, two predictions that subsequently could be confirmed experimentally. Newly identified components have been added to the initial model (see e.g. Pesch and Hülskamp, 2004), which was fine-tuned in a series of computer simulation studies (Benitez et al., 2007; Bouyer et al., 2008; Dupuy et al., 2008) that were validated experimentally (Kang et al., 2009). Ongoing cycles of computational modeling and experimental validation continue to fine-tune an awareness of epidermal patterning in Arabidopsis (Bouyer et al., 2008).

The analysis of epidermal patterning is one of several specific questions in plant morphogenesis that could be addressed with this approach, including gravitropism (Swarup et al., 2005), phyllotaxis (Jönsson et al., 2006; Smith et al., 2006), root tip specification (Grieneisen et al., 2007), leaf venation (Merks et al., 2007), mechanical signalling at the shoot apex (Hmant et al., 2008), lateral root initiation (Laskowski et al., 2008), Prusinkiewicz et al., 2009; Dun et al., 2009) and patterning of the cell wall (Tindemans et al., 2010). These projects demonstrate how plant biologists benefit from computational modeling by developing new ideas, insights and hypotheses and how it helps predict missing system components. To date, modeling typically requires close collaboration between wet-lab scientists and computational biologists or mathematicians, even for testing relatively straightforward ideas or variants of existing models for which no new modeling and simulation codes and techniques need to be developed. Developments in bio-informatics demonstrate that presumably this only represents a transient state in the development of modeling. Initially only computer scientists or computational biologists developed and applied sequence-based algorithms. Since then, experimental biologists have integrated several, now standard bioinformatics tools in their research, while computer scientists and mathematicians continue to develop new algorithms and methods. To perform for example a BLAST search, experimental biologists no longer require collaborations with an expert bioinformatician. We hope that experimental biologists will soon also have a standard set of tools available for testing simple dynamic hypotheses on morphogenesis (Merks et al., 2006a). Apart from developing their own dynamic models and testing those in the lab, it will also help experimental biologists develop an intuition of how computational models can help them advance their research and assist them in framing a set of specific questions to be answered using more advanced models in collaboration with computational biologists.

A number of simulation systems exist for modeling multicellular plant systems. A popular and mature system is L-studio (Karwowski and Prusinkiewicz, 2004). As one of its features L-studio implements vv-systems, a two-dimensional rewriting grammar to model cell division (Smith, 2006) derived from L-systems (Lindenmayer, 1968, 1968, 1975). L-studio and vv-systems have
been applied in a number of recent studies on plant development (Smith et al., 2006; Bayer et al., 2009; Prusinkiewicz et al., 2009). L-Studio has an intuitive user-interface and users can specify models using L-system or vv-system rules. In vv-systems a morphological transformation of the tissue as a whole is specified by the user or by a model variable (Smith et al., 2006). The cell division algorithm then partitions the resulting space. This is a powerful approach if one is primarily interested in how growth can drive pattern formation (i.e., by increasing the domain in which pattern formation takes place (Crampin et al., 2002), e.g., leaf development [Runions et al., 2005], phyllotaxis [Jönsson et al., 2006; Smith et al., 2006], or trichome initiation [Dupuy et al., 2008]), or if one is interested in how stable patterns can drive morphogenesis entirely downstream (see, e.g., the emerging primorida in Smiths’ phyllotaxis model [Smith et al., 2006]). In contrast to such morphostatic development (Salazar-Ciudad et al., 2003), in many cases morphogenesis is a closed feedback loop of pattern formation, growth and morphogenesis (Holloway, 2010); here cell divisions guided by spatiotemporal patterns lead to domain growth that in turn change pattern formation. In such morphodynamic systems (Salazar-Ciudad et al., 2003), the morphology emerges from a feedback-loop between pattern formation and simultaneous morphogenesis, while chemical patterns drive morphogenesis by locally affecting cell behavior.

Other popular multicellular modeling techniques include the cell-based models (Merks and Glazier, 2005). Cell-based models derive morphogenesis from the genetically-regulated behavior of individual cells, that can be modeled as individual particles or as collection of particles. Chaste (‘Cancer, heart and soft-tissue environment’; Pitt-Francis et al. 2009) provides a set of libraries for developing biological simulations of animal tissues, including a cell-based simulation system for modeling tissue growth. It represents cells by its centers and connects cells with virtual springs. A disadvantage of such single-particle methods is that they cannot represent any morphological, subcellular detail, including cell polarity or cell shape. An example of a multi-particle method is the Cellular Potts Model (CPM; Graner and Glazier, 1992; Glazier and Graner, 1993). The CPM was developed originally for simulating animal development (reviewed in Merks and Glazier [2005]) and was recently applied to plants (Grieneisen et al., 2007). In their model of root development, Grieneisen and coworkers (2007) propose that the auxin maximum at the root tip determines the location of the quiescent center in the root apical meristem. The auxin maximum results from a dynamic equilibrium between central auxin transport towards the root tip (acripetal transport), lateral auxin transport towards the shoot (basipetal transport), and auxin reflux from the lateral epidermis towards the stele at the root basal meristem. The same polar auxin transport mechanism also forms an auxin gradient along the root meristem, which is thought to regulate cell division, cell elongation and cell maturation in the growing root via the PLT1-4 genes (Galinha et al., 2007).

Although most of the work focuses on polar auxin transport on static cell lattices or cellular
configurations derived from microscopic images, the cellular Potts model shows that the mechanism can produce realistic root division patterns while maintaining a dynamically stable auxin maximum at the root apex.

Because Chaste and the CPM in their present form lack an accurate description of the physical properties of the cell wall, adjacent cells slide along one another in the cellular Potts models, making their use in modeling of growing plant tissues is relatively limited. Therefore most developmental phenomena simulated with Chaste and the CPM are examples of plastic morphogenesis, as it occurs in most animal tissues where in most tissues the cells can migrate and move relative to one another. Because plant cells cannot migrate and do not slide along each other\(^1\), plant morphogenesis depends exclusively on patterned cell division and cell expansion. This mode of development is often called symplastic growth (Priestley, 1930; Erickson, 1986).

Apart from ‘gluing’ plant cells to each other and maintaining the shape of plant cells, cell walls determine the mechanical properties of plant tissues. The directed orientation of cell wall microfibrils determines differential directionality of tensile strength in cell walls, allowing for anisotropic cell expansion that underlies organ morphogenesis (Green, 1980; Baskin et al., 2004). Moreover cells respond to forces in the tissue, e.g. by setting their division plane parallel to the strain direction in the shoot apical meristem (Hamant et al., 2008). Thus both the mechanical properties of the cell wall and the tight intercellular connections are crucial for understanding morphogenesis of plant tissues.

A number of cell-based modeling methodologies consider detailed cell wall mechanics. CellModeller\(^2\) (Rudge and Haseloff, 2005; Dupuy et al., 2008) is a two-dimensional environment for plant tissue simulation. It has explicit representations of the cells, the cell walls, and the relative interactions between the cells, and includes differential equation models of the dynamics of cellular properties, including biochemical networks. CellModeller describes cell walls as viscoelastic rods, which are maintained tensed by the turgor pressure. Cell wall dynamics is simulated with differential equations, while architectural, geometric models describe cell division and cell death. The model of Corson et al. (2009) also represents cell walls as viscoelastic beams, while using an energy minimization approach to simulate cell wall dynamics and cell growth.

A shortcoming in most of the current modeling approaches is that they require a high level of mathematic and computer programming skills, which often keeps biologists from using them. Therefore we introduce VirtualLeaf, a modeling environment that allows biologists to turn their hypotheses of multicellular development and patterning into a dynamic simulation model with little programming effort. For simplicity and computational efficiency the underlying computational

\(^1\) An exception is the intercalary growth of phloem fibers (Ageeva et al., 2005).

method describes cells in relatively little detail. Nevertheless the model explicitly considers the balance between turgor pressure and cell wall tension. Our method relates to CellModeller and the method by Corson et al. (2009) in its representation of the cell walls and its consideration of turgor pressure. We illustrate the use of our simulation environment with a number of example simulations, including simple growing tissues and systems in which chemical pattern formation interacts with local growth.

8 Model description

9 Cellular mechanics

Our computational method represents cell walls in a tissue in two dimensions (2D) using polygonal finite elements (Figure 1), an approach similar to that taken in other plant tissue models (Nagai and Honda, 2001; Rudge and Haseloff, 2005; Dupuy et al., 2008; Corson et al., 2009). A cell is bound by a set of elastic cell wall elements. To restrict relative cell movement, adjacent cells share a cell wall, where each cell wall consist of multiple cell wall elements, so that cell walls are flexible. A Monte-Carlo-based energy minimization algorithm straightforwardly describes cell behaviors, including expansion, division, and active shape change. This approach conceptually derives from the Cellular-Potts model (Graner and Glazier, 1992; Glazier and Graner, 1993), which is most suitable for plastic, animal development (reviewed in [Merks and Glazier, 2005]). We allow cell walls to move according to rules derived from the classic equations for turgor-driven cell expansion (Lockhart, 1965):

1. The intracellular turgor pressure exerts a uniform force on cell walls, attempting to enlarge cells.
2. The elastic cell walls counteract the turgor pressures.
3. Walls expand irreversibly if stretched over a threshold. Wall relaxation occurs under the influence of enzymes secreted into the wall that break specific bonds that crosslink cellulose microfibrils (Cosgrove, 2000; Robles P, 2009) and is also called cell wall yielding (Lockhart, 1965; Cosgrove, 1985).

Based on this set of rules, our algorithm defines for each cell $i$ a resting area $A_r(i)$ at which the cell’s turgor pressure balances the ambient pressure and a resting length $L_r(j)$ for each wall element $j$; this is the length a section of cell wall would assume in the absence of turgor pressure. The actual cell area is $a(i)$, and actual wall-element lengths are denoted, $l(j)$. We can describe the
balance between turgor pressure and cell wall resistance in terms of a generalized potential energy or Hamiltonian, $H$:

$$
H = \lambda_A \sum_i (a(i) - A_i(i))^2 + \lambda_M \sum_j (l(j) - L_T(j))^2
$$

where indices $i$ and $j$ sum over all cells and polygon edges, respectively, $\lambda_A$ is a parameter setting the cells’ resistance to compression or expansion, and $\lambda_M$ is a spring constant.

The simulation uses Metropolis dynamics (Metropolis et al., 1953) to minimize the potential energy $H$: the algorithm iteratively selects a random node and attempts to move it in a random direction $\vec{x}_{\text{new}} = \vec{x} + \xi \vec{r}$, where $\vec{r} = \{\rho, \theta\}$ is a random vector chosen uniformly within the unit circle (i.e., $\rho \in [0, 1]$ and $\theta \in [0, 2\pi]$) and $\xi$ is the step size, that is fixed for a given simulation. The algorithm calculates the energy difference $\Delta H$ associated with the attempt and always accepts the move if it leads to an energy drop. To prevent the system from getting stuck in local energy minima, we also accept moves producing a small energy increase, according to the Boltzmann probability function, $P(\Delta H) = e^{-\Delta H / T}$. $T$ is a parameter setting the amount of “noise” added in this way. To model wall relaxation (cell wall yielding) we introduce new nodes whenever a polygon edge’s length is stretched to the extent that it exceeds a threshold value $\vartheta_{\text{yield}}$, typically four times the original length of the wall element. After the algorithm has attempted to move each of the nodes in the simulation in random order (a Monte Carlo Step), it checks if the forces described in the Hamiltonian have balanced out sufficiently. Assuming shape relaxation is fast compared to biological changes, the algorithm repeats this procedure until turgor pressures, cell expansion and the resulting cell displacements have equilibrated sufficiently and the energy difference drops below a relaxation threshold ($\Delta H < \vartheta_H$). After this relaxation cycle has completed, the algorithm solves the biologically-motivated processes, including turgor pressure increases, cell divisions, active cell shape changes, and transport of chemical signals.

Thanks to the energy minimization formalism used for the model, we can include additional cellular properties in a straightforward way by describing them as energy constraints, e.g., body forces like gravity. For example, the length of cells (Merks et al., 2006b) can be constrained using:

$$
H' = H + \lambda_E \sum_j (e(j) - E(j))^2
$$

with $H'$ the extended Hamiltonian. $e(j)$ is the actual length of cell $j$, $E(j)$ is the resting length of the cell $j$, and $\lambda_E$ gives the strength of the length constraint and can be interpreted as a
deformability parameter. Assuming cell shapes are ellipsoid, we estimate the cell length from the

cells’ inertia tensors as \( e(j) = \sqrt{\lambda_b(j)/a(j)} \), with \( \lambda_b(j) \), the largest eigenvalue of the cell j’s
inertia tensor \( \mathbf{I} \):

\[
\mathbf{I} = \iint_D \begin{bmatrix}
(x-x_c)^2 & -(x-x_c)(y-y_c) \\
-(x-x_c)(y-y_c) & (y-y_c)^2
\end{bmatrix} \, dxdy
\] (3)

with \( \iint_D \) the area integral over the cell. Using Green’s Theorem, the inertia tensor derives from the
vertices of the polygon describing the cell. The algorithm reconstructs the inertia tensor from the
raw moments of the cells’ coordinates, quantities that can be efficiently updated using local
information after each move of the cell boundary. Thus this algorithm produces a computationally
efficient, instantaneous estimate of the cell lengths.

For cell division, we define a division axis and build a new cell wall (consisting of several
wall elements), shared by the two daughter cells. The nodes of the parent cell are assigned to the
two daughter cells. Typically, cell division is over the shortest axis, which is derived from the
eigenvectors of the cell’s inertia tensor (Eq. 3).

**Transport of chemical signals and metabolites**

Sets of ordinary differential equations in each cell describe the biochemical networks and genetic
regulatory networks. Additional differential equations model diffusive transport across the two cell
membranes and the cell wall separating adjacent cells. To model active transport by transport
proteins, e.g. PINs, we define an extra object “Wall” (Figure 1B), which spans several “Wall
Elements”. It stores the amounts of transport proteins present in the cell membranes at either side of
the walls. Thus the export rate of a chemical A from a cell 1 to an adjacent cell 2 would depend on
the concentration of A in cell 1 and the amount of A’s export protein in cell 1’s membrane adjacent
to cell 2. Although diffusion between apoplast compartments is not implemented yet, the wall
elements can also describe an extracellular space (apoplast). For simplicity, we do not consider
apoplasts in the examples described here. After each relaxation cycle (Figure 1A) we dynamically
construct a set of differential equations describing the intracellular reactions and transport of
chemicals (with one equation for each chemical per cell and one equation for each chemical per cell
wall), and numerically integrate them over a period \( \Delta t \). This setup implies that the set of
differential equation increases in size after each cell division, because division adds new cells and
new cell walls. As a numerical integrator, we use a fifth-order, adaptive step-size Runge-Kutta algorithm (Press et al., 1992).

Modeling framework

VirtualLeaf separates the conceptual biological model from its implementation in terms of the algorithms for cellular mechanics and chemical transport, and the visualization and steering of the simulations. The model definition itself is contained in a model plugin, a software module that is developed independently from the main framework. It is defined by combining software building blocks representing objects familiar to the experimental plant biologist, like cells, cell walls, molecules, transporter proteins, or signaling molecules. New model plugins are developed by adapting a small section of C++ code (see Supplementary Text S1). The model framework runs fast (minutes rather than hours, depending on model complexity), enabling quick interactive desktop model evaluation. The models can also be run without a user-interface, enabling high-throughput parameter screening on a computer cluster.

Results

Based on the developed computational VirtualLeaf framework it is possible to investigate various aspects of tissue growth. We will illustrate this with a number of example models that investigate emergent behavior of biological systems based on the characteristics attributed to individual component cells.

Example 1. Growth, margin stiffness, and cell shape

As a first study we test the behavior of a simple model of plant tissue growth (Figure 2A and Video S1). The simulation was initiated with a single cell, whose target area A (and hence its “turgor pressure” \( (A_i - a_i)^2 \) ) gradually increases at a constant rate. Cells divide over the short axis once their area \( a \) has doubled (see Supplementary Text S1, Tutorial 1 for implementation details). This basic simulation rule creates a rounded “callus” of plant tissue. In these simulations we assumed that cell walls at the leaf perimeter were thicker and stiffer than the internal walls (\( \lambda_m(\text{perimeter})=\zeta\lambda_m(\text{internal}) \)), with \( \zeta = 2 \) the stiffness ratio between internal and perimeter walls. Perimeter walls are marked with blue nodes in Figure 1B and Video S2. This assumption agrees with the observation that margin walls in plant leaves are thick relative to walls of internal cells.
With these assumptions our model rules produced a rounded aggregate of plant tissue. Interestingly, after reducing the stiffness of perimeter cell walls ($\zeta = 1; \lambda_M(\text{perimeter}) = \lambda_M(\text{internal})$) a more “tumorous” morphology develops (Figure 2B and Video S2). In the absence of a neighboring cell or a thick cell wall resisting turgor, the perimeter cells “bulge” outwards and expands faster than the internal tissue. Indeed in plant leaves the margin cells play a crucial role in morphogenesis (Reinhardt et al., 2007), which was proposed to be due to a biophysical function of the margin cells. In this simple model of plant tissue growth we have not included the elongated shape of margin cells and we neglect many other mechanical features of plant cells (e.g. microtubules) and of the whole organ (e.g. the venature). Nevertheless already in this simple model the mechanical properties of the margin can regulate the shape of the tissue as a whole, by constraining the expansion of the interior tissue. Figure 2C and Video S3 illustrate the effect of a cell shape constraint (Eq. 2). In this example, the cells elongate into a direction that depends both on the initial shape of the cells and on the forces the adjacent cells exert on them. Illustrating the shape constraint, this simulation is not intended to correspond to a real plant organ.

**Example 2: Signaling molecules regulating growth and differentiation**

Plant morphogenesis results from regulated cell divisions and pattern formation via polar or diffusive transport of chemical signals. Using the VirtualLeaf environment it is possible to investigate the interaction between a diffusive morphogen and growth (Figure 3 and Video S4). One of the cells produces a morphogen (e.g. cytokinin or auxin; concentrations indicated in shades of gray), which transports diffusively from cell to cell. If the concentration $c$ of the morphogen is higher than a threshold, $c > \theta_1$, with $\theta_1$ a concentration threshold, cells expand and divide once the area has doubled. At lower concentrations, $\theta_1 > c > \theta_2$, with $\theta_2 < \theta_1$ a second concentration threshold, cells only expand. At concentrations $c < \theta_2$, cells neither expand nor divide, i.e. they enter a maturation state. If the source cell divides, the source is assigned to one of the daughter cells at random. Note that in this illustration we do not intend to model a biological system in particular; but even in this simple model the cells naturally progress from a division phase, to an expansion phase and finally enter a maturation phase, analogous to what happens in e.g. a root tip (Beemster and Baskin, 1998). The model does not consider gravitropy; hence the growth tip does not correct for the accidental sharp bend in Figure 3D. A local growth stagnation causes the bend: the morphogen concentration in the large expanding remains too high for the cell to enter division. This example simulation could form a basis for more complicated, realistic models, focusing on the mechanisms responsible for morphogen gradient formation (Gieneisen et al., 2007) and on modeling the relation between cell division rate, cell expansion rate, and size of meristem and...
expansion zone (see e.g. Chavarria-Krauser and Schurr, 2004; Chavarria-Krauser et al., 2005). Tutorial 2 in the appendix shows how to construct such plant growth models involving signaling molecules with VirtualLeaf.

A more realistic and complicated simulation involving chemical reactions that illustrates the use of the VirtualLeaf framework for developing new hypotheses of biological development involves the studies of vascular patterning during Arabidopsis leaf development (Merks et al., 2007). The model predicts the existence of a travelling wave of auxin moving from the leaf tip to the leaf base. Auxin is concentrated into a peak by PIN1, which polarizes towards neighboring cells at a rate proportional to their concentration of auxin, a mechanism introduced by Jönsson et al. (2006) and Smith et al. (2006) for phyllotaxis. In our leaf venation model, auxin stimulates the production of auxin pumps. The pumps polarize towards the next cells and pump it onwards. In this way travelling waves form, which leave behind trails of polarized cells that we hypothesize to differentiate into the vascular system (Figure 4 and Video S5). The long-term goal of this continuing project is to reconstruct in the VirtualLeaf environment how leaves develop in the interplay between patterned cell division and cell expansion, the transport of auxin and other signaling molecules between cells and the physical characteristics of various cell types.

Example 3: Interplay between growth, cell division, and patterning

A next modeling example illustrates how tissue growth can interact with auxin-driven patterning. Here each cell implements Jönsson et al.’s upstream auxin transport model (Heisler and Jonsson, 2006; Jönsson et al., 2006; Sahlin et al., 2009), and lets local auxin concentrations drive cell expansion. For simplicity, we assume auxin is produced along the tissue’s perimeter. Cells divide along their shortest axis when they reach twice their original size. After a small cell cluster has formed, auxin accumulates in the cluster’s center and induces further growth. New auxin maxima appear at the tissue perimeter once sufficient tissue has formed, inducing further, localized growth. This feedback between growth and pattern formation eventually forms a bulbous morphology (Figure 5 and Video S6). Tutorials 3, 4, and 5 in the Appendix show in detail how such a model can be constructed in VirtualLeaf.

In the previous sections, we illustrated the use of VirtualLeaf to model the biomechanics of plant tissue growth and its use in modeling pattern formation and morphogen-regulated growth. These relatively simple models give us more insight into interactions that exist between pattern formation, tissue growth and tissue mechanics that drive the formation of plant growth. Supplementary Text S1 provides a step-by-step introduction to building models in VirtualLeaf.
Discussion

In this paper we introduced VirtualLeaf, a cell-based modeling framework for the simulation of symplastic growth, the typical mode of development for plants and some animal tissues in which adjacent cells cohere tightly and cannot slide relative to one another. The example simulations—ranging from simple tissue growth to models where growth and biochemical patterning interact dynamically—illustrate how cell-based modeling frameworks like VirtualLeaf can help developmental plant biologists to turn a hypothesis into a dynamic model, and illustrate which tissue-level patterns follow from the observed or hypothetical behavior of the individual cells (Merks and Glazier, 2005). Thus modeling can be used to check hypotheses for “internal consistency”, by showing whether a putative mechanism indeed produces the shapes and patterns it was meant to explain.

One of our motivations for developing VirtualLeaf was the observation that plant (symplastic) development seems fundamentally different from animal development, because the tight coherence of cell walls prohibits plant cells from moving relatively to each other. Would modeling symplastic development be impossible with a “plastic” cell-based simulation method like the Cellular Potts model or Chaste? Balter et al. (2007) propose that connecting cellular Potts cells with elastic springs, in an approach much like our “cell wall elements”, would prevent relative cell movement. However, doing so would 1) suppress the most powerful feature of the cellular Potts model—relative cell movement—at additional computational cost, while 2) the model would not include the interaction between turgor pressure and cell wall tension, one of the main features of plant development.

A further advantage of using an off-lattice method for modeling plant development is in the descriptions of chemical concentrations. In VirtualLeaf these are mathematically part of the cells and move along with them; in lattice-based formalisms, including the CPM from which VirtualLeaf was derived, the chemical concentrations are described as parts of the lattice instead. Hence, because cells move relative to the lattice, in lattice-based simulation formalisms the morphogen concentrations may “lag behind” if cells move fast relative to reaction or transport rates.

Currently, developing a new model for VirtualLeaf requires some C++ programming. Although models are defined using biological concepts and it is possible to modify existing models with only basic experience in programming, the required coding and compilation still are a hurdle for computationally untrained users. We are currently defining and implementing a provisional
domain-specific language based on the draft *Cell Behavior Ontology*\(^3\) (CBO), a well-defined set of
terms for describing the behavior of animal, plant, or bacterial cells. A biological modeling
language derived from the CBO would make it possible to define the model entirely in a conceptual
language familiar to biologists.

Apart from a new modeling language, future versions of *VirtualLeaf* will require several
improvements. Currently, cells only contain a single set of chemical concentrations, while many
simulation models of developmental processes depend on intracellular, non-homogeneous spatial
distributions of molecular signals (e.g. Grieneisen et al., 2007; Laskowski et al., 2008).

Intracellular, triangular grids will allow us to consider intracellular chemical distributions. We also
hope to develop a three-dimensional *VirtualLeaf* environment to allow the simulation of tissues
consisting of multiple cell layers; required algorithms are relatively straightforward, because
entities of our two-dimensional framework (polygons, line pieces, etc.) have one-on-one three-
dimensional counterparts (polyhedra, triangles, etc.). Further extensions of *VirtualLeaf* would
allow modeling of the physiological interactions of the growing plant organ with the rest of the
plant and its environment. Functional-structural plant models (FSPM; Godin and Sinoquet, 2005)
focus on the growth and functioning of the whole plant in interaction with its environment. A
number of functional-structural plant modeling frameworks, including *OpenAlea* (Pradal et al.,
2008) and *CrossTalk* (Draye and Pagès, 2008), integrate structural whole-plant models,
environmental models and tissue growth models to single multi-scale plant models. Interfacing
*VirtualLeaf* with *OpenAlea* or *CrossTalk* would involve 1) adding appropriate boundary conditions
to simulate in- and efflux of materials from the rest of the virtual plant, and 2) adding the control
handles to synchronize the *VirtualLeaf* simulations with the FSPM model.

We hope that future cell-based modeling frameworks will integrate off-lattice and lattice-
based cell-based methodology, and interface whole-plant and single-organ approaches, thus
enabling researchers to use the most suitable approach for the problem or even making it possible to
combine approaches. Crucial steps towards this goal include the development of advanced model
interfacing algorithms, generic, standard modeling languages for cell-based modeling, and
multiscale model integration platforms like *OpenAlea* or *CrossTalk*.

In its present form, the *VirtualLeaf* provides an accessible problem-solving environment (Merks et
al., 2006a; Cickovski et al., 2007) that allows biologists with minimal programming skills to turn
their hypotheses on the mechanisms of plant development and symplastic tissue growth into
dynamic simulations. *VirtualLeaf* is available as an Open Source project at
http://virtualleaf.googlecode.com. We hope *VirtualLeaf*’s open source model will encourage users

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\(^3\) http://bioportal.bioontology.org/ontologies/39336
to use VirtualLeaf for their modeling projects and to share new models and framework extensions with other VirtualLeaf users.

Supplemental material

Text S1. Modeling plant development with VirtualLeaf, a tutorial

Video S1. Symplastic cell-based plant tissue simulation, as in Figure 2A. Cells adhere tightly to each other and expand at fixed rate. Cells divide once their area $a_i$ has doubled. Walls at leaf perimeter have double stiffness relative to internal cell walls ($\lambda_m$(perimeter)$=2\lambda_m$(internal));

Video S2. Symplastic cell-based plant tissue simulation, as in Figure 2A. Cells adhere tightly to each other and expand at fixed rate. Cells divide once their area $a_i$ has doubled. If perimeter walls have “single stiffness” ($\lambda_m$(perimeter)$=\lambda_m$(internal)), perimeter cells bulge out generating a tumorous morphology.

Video S3. Symplastic cell-based plant tissue simulation, as in Figure 2A. Cells adhere tightly to each other and expand at fixed rate. Cells divide once their area $a_i$ has doubled. Growing tissue with elongating cells.

Video S4. Diffusive morphogen stimulates growth, as in Figure 3. One tip cell produces a morphogen. At concentrations $c > \theta_1$ cells expand and divide once the area has doubled, for $\theta_1 > c > \theta_2$ for cells only expand, while for $c < \theta_2$ cells neither expand nor divide, i.e. they go into a maturation state.

Video S5. A traveling-wave of auxin forming a polar auxin channel, as in Figure 4. Auxin (shown in green) flows in from the leaf tip at the purple cell walls. PIN (red) localizes preferentially at cell walls adjacent to cells with highest auxin concentration as in recent phyllotaxis models. A t=3h; auxin accumulates at the leaf edge, some auxin leaks into the lower cell layers. PIN1 production rises. B t=13h; auxin converges into a small peak, towards which the adjacent cells polarize. C t=20h; completed auxin channel, traced out by the traveling wave of auxin. See also Merks et al. (2007).
Video S6. Interaction between auxin accumulation and growth, as in Figure 5. Auxin accumulates according to Jönsson et al. (2006) model. Auxin induces cell expansion; cells divide over shortest axis after doubling in size.

Code S1. Source codes of the VirtualLeaf V1.0, written in C++ with Qt 4.6. More recent versions will be available from http://virtualleaf.googlecode.com. Users wishing to contribute to the further development of VirtualLeaf can use the googlecode site to share their additions and new model plugins.

Code S2. Installer for Windows. This distribution includes executables only for evaluating the VirtualLeaf. To develop your own models, download the source code distribution (Code S1).

Code S3. Installer for MacOSX. This distribution includes executables only for evaluating the VirtualLeaf. To develop your own models, download the source code distribution (Code S1).

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Figure legends

Figure 1: Cell-based modeling of plant tissue growth. Polygons represent cells, surrounded by cell walls, which consist of viscoelastic wall elements linked by nodes. A wall furthermore represents the membranes and transporter proteins on either side of the cell wall and the apoplast, i.e., the intercellular space. The algorithm iteratively displaces the nodes to balance the turgor pressure with the wall tension and possible extra constraints (e.g., a cell shape constraint).

A. Overview of simulation algorithm. The algorithm attempts to displace each node in random order; one such cycle is a Monte Carlo Step (MCS). MCSs are repeated until the energy fails to drop significantly. This iteration of MCSs is called a relaxation cycle. Once the system has relaxed, we simulate the biochemical networks and other cell behavior rules. The potentially affect cell properties (e.g., turgor pressure, wall extensibility, etc.) and a new relaxation cycle is required. B. Visualisation of the simulation algorithm. 1. This move of a cell wall node relaxes local turgor pressure and minimally strains the cell walls; the move will be kept. 2. This move compresses the right cell, and extends the left cell. The top wall element is overstrained, the bottom wall element is compressed. Because this potential move would be energetically unfavorable it will typically be rejected. 3. Each cell potentially contains a biochemical network, described by a set of differential equations; the chemical species often regulate cell and wall attributes, including turgor pressure and wall extensibility. 4. A molecular species (shown in purple) channels back and forth between the cytoplasm and the cell membrane, as described by sets of differential equations. 5. Transport of the blue molecular species from cell to cell; diffusion passively transports molecules from high to low concentrations, molecules at the cell membrane (e.g., PINs) actively transport molecules upstream or downstream.

Figure 2: Symplastic cell-based plant tissue modeling. Cells adhere tightly to each other and expand at fixed rate. Cells divide once their area $a_i$ has doubled. Simulation states are shown after 50, 100, 150, 200, 250, and 300 relaxation cycles, respectively. A. Walls at leaf perimeter have double stiffness relative to internal cell walls ($\lambda_m$ (perimeter) = 2 $\lambda_m$ (internal)); B. If perimeter walls have “single stiffness” ($\lambda_m$ (perimeter) = $\lambda_m$ (internal)), perimeter cells bulge out generating a tumorous morphology. C. Growing tissue with elongating cells.

Figure 3 Diffusive morphogen stimulates growth. One tip cell produces a morphogen. At concentrations $c > \theta_1$ cells expand and divide once the area has doubled, for $\theta_1 > c > \theta_2$ for cells...
only expand, while for $c < \theta_2$ cells neither expand nor divide, i.e. they go into a maturation state.

Simulation state after A 100, B 500, C 1000, D 1500, E 2000 relaxation cycles.

**Figure 4** A traveling wave of auxin forming a polar auxin channel. Auxin (shown in green) flows in from the leaf tip at the purple cell walls. PIN (red) localizes preferentially at cell walls adjacent to cells with highest auxin concentration as in recent phyllotaxis models. A $t=3\text{h}$; auxin accumulates at the leaf edge, some auxin leaks into the lower cell layers. PIN1 production rises. B $t=13\text{h}$; auxin converges into a small peak, towards which the adjacent cells polarize. C $t=20\text{h}$; completed auxin channel, traced out by the traveling wave of auxin. See also Merks et al. (2007).

**Figure 5.** Interaction between auxin accumulation and growth. Auxin accumulates according to Jönsson et al. (2006) model. Auxin induces cell expansion; cells divide over shortest axis after doubling in size. Simulation state after A) 10 B) 30 C) 50 D) 70) E) 90 F) 110 simulation cycles.
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