Linear stability analysis of morphodynamics during tissue regeneration in plants

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Abstract

One of the key characteristics of multicellular organisms is the ability to establish and maintain shapes, or morphologies, under a variety of physical and chemical perturbations. A quantitative description of the underlying morphological dynamics is a critical step to fully understand the self-organising properties of multicellular systems. Although many powerful mathematical tools have been developed to analyse stochastic dynamics, rarely these are applied to experimental developmental biology.

Here, we take root tip regeneration in the plant model system Arabidopsis thaliana as an example of robust morphogenesis in living tissue, and present a novel approach to quantify and model the relaxation of the system to its unperturbed morphology. By generating and analysing time-lapse series of regenerating root tips captured with confocal microscopy, we are able to extract and model the dynamics of key morphological traits at cellular resolution. We present a linear stability analysis of its Markovian dynamics, with the stationary state representing the intact root in the space of morphological traits. This analysis suggests the intriguing co-existence of two distinct temporal scales during the process of root regeneration in Arabidopsis.

We discuss the possible biological implications of our specific results, and suggest future experiments to further probe the self-organising properties of living tissue.

Keywords: morphogenesis, Arabidopsis, regeneration, morphodynamics, plant roots, morphometrics

Supplementary material for this article is available online (Some figures may appear in colour only in the online journal)

1. Introduction

In most multi-cellular systems, the function of an organ or a tissue relies on its morphology and internal cellular organization. Not surprisingly, evolutionary processes tend to select mechanisms to at least partially restore an optimal tissue state that has been damaged, through controlled developmental processes known as wound repair or regeneration [1, 2]. From a more abstract point of view, we could refer to such a state as a stable equilibrium in the space of morphologies, and the process of regeneration as the relaxation of the system back to equilibrium after a small perturbation. Tissues exhibiting such stability to perturbation are sometimes referred to as morphologically robust [3].

Since the ground-breaking and influential theory of transformation by D’Arcy Thompson [4], quantitative approaches have been successfully applied to study the changes occurring during shape formation, or morphodynamics [5]. In plants, rigid cell walls suppress virtually all cell migration, making the process of tissue organization much simpler to understand. Perhaps for this reason, plants have been used extensively as model systems to study morphodynamics [6, 7]. The very
successful molecular genetics approach in developmental biology has generated a body of knowledge on the molecular mechanisms involved in pattern formation and generally in tissue organization. In plants, a number of detailed computational models have been proposed to capture the complexity of genetic networks controlling morphogenesis [8, 9] and to simulate the diversity of macroscopic forms [10]. Nevertheless, although theoretical concepts such as morphostate and ontogenetic trajectories have been explored from the point of view of dynamical systems [11], a broad mathematical framework to link experimental data on morphodynamics with tissue self-organization is essentially missing.

We believe that a quantitative analysis of tissue regeneration can provide a significant step towards the development of such a framework. Among plant organs, roots offer the advantage of strong geometrical symmetries and simple internal organization, for most part composed of just few concentric layers of cell types. The root tip still maintains an approximate rotational symmetry and is organized in a very stereotypical pattern, harbouring actively dividing cells and an apical stem cell niche [12]. The root tip of the genetic model system Arabidopsis thaliana offers the further advantage of being quite transparent, which makes it an ideal system for live microscopy [13].

Here, we take advantage of the process of root tip regeneration in Arabidopsis [14] and combine cellular morphometrics with the theory of stochastic processes, to propose a novel quantitative approach to study dynamical perturbations of stable morphologies at cellular scale. We use time-lapse imaging with confocal fluorescence microscopy and measure cellular morphological traits. We propose a novel mathematical framework to describe the resulting morphological dynamics, borrowing from field theory methods. Finally, we present a mathematical analysis of the robustness of regeneration at cellular resolution.

2. Results

2.1. Distributions of cellular morphological traits

We followed an established protocol [14] to collect median longitudinal optical sections of uncut as well as cut (i.e. with their tip fully excised) and thus regenerating Arabidopsis root tips. Each root was imaged once a day, for at least 8 d (figure 1).

We developed an original image processing routine based on the marker-controlled watershed segmentation method [15] to identify cells in each optical section and to measure its area, eccentricity and the orientation of its major axis. At each time-point, we measured these traits for all the cells in all root meristems that have been cut and imaged through regeneration, as well as in all uncut root meristems (figure 2).

2.2. A vector field as a representation of cell identity

In our study, cell morphology is described by three metrics, namely area, eccentricity and orientation of the major axis of the cell, encoded in the sine and cosine of the orientation angle prior to registration (see section 4.4 for detailed methods), and thus mapped in a four-dimensional (4D) space (morphospace). After normalising the area and the eccentricity to the interval [0, 1], each cell is represented by a point on the surface of a 4D hyper-cylinder that has a unit circle as base. We asked if all morphologies we identified could, in fact, be subdivided into only a few clusters, or morphological states (morphostates).

We collected area, eccentricity and orientation of all cells from a separate set of 31 uncut root tips at day 0 (3 d after germination) and calculated the Davis–Bouldin index (DBi) [16] to find the number κ of clusters present in the data (figure S1 (stacks.iop.org/JPhysD/52/084002/mmedia)). While the minimization of DBi is a guiding principle for choosing a plausible number of well-separated clusters in the data, there is a level of arbitrariness in this method as in any clustering method. Following the interpretation of Davies and Bouldin [16], figure S1 suggests that the present dataset is hierarchical and that each local minimum corresponds to a partition at a given scale. We noted that the local minimum at κ = 7 is the closest to capture the complexity of the stereotypical organization of the Arabidopsis root meristem, in first approximation composed by the cell types epidermis, cortex, endodermis, pericycle, stele, quiescent center, columella, lateral root cap, although a one-to-one correspondence between morphostates and cell-types should be avoided in our view, as discussed below.

We repeated the analysis using another set of uncut roots at days 0–8, and a third set composed of all the images we collected of uncut and regenerating roots, taken over various days after the cut. All sets identified κ = 7 as a local minimum of the DBi. Finally, we applied Lloyd’s [17] and k-means++ [18] algorithms to cluster the data points in the morphospace and to calculate the position of each cluster’s centroids, which is the point to which the sum of the squared Euclidean distances from all cells in the cluster was minimal. The resulting set of κ centroids in the 4D space represent κ morphostates and thus provide a map to associate a morphostate with every point within that space. The morphostate of a cell is given by the centroid closest (in terms of Euclidean distance) to the coordinates representing the cellular morphology. Each cell was assigned to a morphostate i ∈ {1, 2, . . . , κ} by finding the centroid closest to the cell’s position in the morphospace (a typical map of morphostates is shown in figure S2).

In order to maintain the information about the relative position of each cell across roots, we performed ad hoc image registration to spatially overlap all the roots (maintaining separated cut and uncut sets) at any given time-point (see section 4.4). For each set of registered roots at time t, we defined a probability P(⃗x, t) of finding a pixel at position ⃗x ∈ Rd in the image plane, belonging to a cell in morphostate i. Thus, P(⃗x, t) = (P1(⃗x, t), . . . , Pκ(⃗x, t)) is a vector field representing the distribution of morphostates in a typical root and its evolution in time is a quantitative representation of the morphological changes occurring at cellular level as the root grows (for uncut roots) or regenerates (for cut roots). An equivalent interpretation of the field P(⃗x, t) is that each cell of the typical root is a weighted superposition of morphostates, with the probability to be in a certain morphostate i given by the
Figure 1. Optical longitudinal median sections of representative Arabidopsis roots at various days after tip excision (Regenerating) or mock-treated (Uncut). Confocal microscopy with propidium iodide to counter-stain cell membranes. Scale bar, 50 µm.

Figure 2. Box-plot representation of the distribution of the measured cellular morphological traits (area, eccentricity and orientation) as a function of time, after root tip excision (Regenerating) or mock-treated (Uncut).
components $P_i$ of $\mathbf{P}$. The spatial average of the field $\mathbf{P}(\vec{x}, t)$ is a seven-dimensional vector $\mathbf{P}(t)$, which we call morphovector, whose component $i$ represents the probability of finding anywhere a cell in morphostate $i$, at time $t$. The morphological dynamics of growing or regenerating roots is described by the evolution of its components $\mathbf{P}_i(t)$ in time $t$ (figure 3).

The uncut roots represent the control set, capturing the background noise due to the natural variation among individuals. As expected, the corresponding morphovector shows little variation in time. In contrast, the morphovector of regenerating roots contains components with changes and trends more pronounced than the controls. In essence, it appears that during regeneration the average morphological identity of the cells varies visibly as time progresses and converges back to a morphostate characteristic of the uncut root. Perhaps this is not surprising, as it captures the whole concept of regeneration over all of them) so that $S(\vec{x}, t) = 0$ if a cell at position $\vec{x}$ and time $t$ belongs to a single morphostate (i.e. if $P_i = 1$ for $i = a$ and $P_i = 0$ for all $i \neq a$), whereas $S(\vec{x}, t) = \ln 7 = 1.94...$ if any cell at position $\vec{x}$ and time $t$ belongs to any of the morphostates with equal probability (i.e. if $P_i = 1/7$ for every $i = 1, ..., 7$) and $\mathbf{P}(\vec{x}, t)$ is a uniform superposition of all morphostates.

The scalar field $S(\vec{x}, t)$ contains information about the local variation of cell morphologies within a sample of comparable roots. A visual representation of the scalar field $S(\vec{x}, t)$ mapped on top of a typical root (figure 4) shows large regions of high (large $S$, light gray) and low (small $S$, dark gray) morphological variability. Interestingly, low-variability regions loosely correspond to well-characterized tissue types such as lateral root cap and epidermis (figure 4, top row). Moreover, regenerating roots show a progressive re-establishment of low-variability regions that were lost immediately after tip excision (figure 4, bottom row).

To further quantify the progress of morphological reorganization of regenerating roots, we took the spatial average $\bar{S}(\vec{x}, t)$ of $S(\vec{x}, t)$ and plotted it as a function of time (figure 5). Its error was calculated by the Jackknife procedure [20],

$$\text{error}_{\text{Jackknife}} = \left[ \frac{N - 1}{N} \sum_{i=1}^{N} \left( \bar{S}_i - \frac{1}{N} \sum_{j=1}^{N} \bar{S}_j \right)^2 \right]^{1/2}, \quad (2)$$

where $\bar{S}_i$ is the spatially averaged entropy when root $i$ was left out from the sample. Large values of the spatial average $S$ thus refer to a sample of roots where the cellular morphologies are highly variable, while small values of $S$ indicate small

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{figure3.png}
\caption{Dynamics of each of the components $\mathbf{P}_1(t), \ldots, \mathbf{P}_7(t)$ of the morphovector $\mathbf{P}(t)$, plotted as their relative contribution and as a function of time $t$ after root tip excision (Regenerating) or mock-treated (Uncut).}
\end{figure}
morphological variation within the sample. This could respectively be interpreted as a measure of a disordered, less robust pattern (high entropy) and an ordered, more robust one.

2.4. Linear approximation of morphological perturbation

The biology of root tip regeneration suggests that the uncut state is a stable equilibrium in the morphological space, that the tip excision is a perturbation away from equilibrium and that the process of regeneration is the relaxation of the system back to equilibrium. The formalism and measurements described above provide us with the quantitative observables to describe mathematically the dynamics of such perturbation.

In the following, we call \( \mathbf{P}_u(t) \) and \( \mathbf{P}_c(t) \) the morphovectors for uncut and cut roots, respectively, as measured on days \( t \in \{0, 1, \ldots, T_{uc} - 1\} \). Here \( T_u = 9 \) and \( T_c = 10 \) denote the numbers of days uncut and cut roots have been observed for.

In a mean-field approximation, we consider the evolution of the (spatially averaged) morphovector \( \mathbf{P}_c(t) \) as a (linear) continuous-time Markov process with master equation

\[
\frac{d}{dt} \mathbf{P}_c(t) = M \mathbf{P}_c(t),
\]

Figure 4. Local Shannon entropy \( S(\mathbf{x}, t) \) mapped on a typical longitudinal median section of Arabidopsis root tip, as a function of time after tip excision (Regenerating) or mock-treated (Uncut).

Figure 5. Spatial average of the Shannon entropy \( S(\mathbf{x}, t) \) as a function of time after tip excision (Regenerating) or mock-treated (Uncut). Error bars, Jackknife.

4 Here and in the following the subscript \( u \) and \( c \) refer to uncut and cut roots respectively.
where the matrix $M$ is the Markov matrix governing the dynamics of $\mathbf{P}_c(t)$, or transition matrix. The off-diagonal elements are thus non-negative Poissonian transition rates between morphostates of cells in the cut root. The diagonal elements, on the other hand, are determined by the constraint that columns have to sum to 0 as a matter of probability conservation, $(1, \ldots, 1)M = 0$. Equation (3) is a linearisation of a potentially more complicated evolution of $\mathbf{P}_c$, which, here, we demand to obey simple linear Markovian dynamics. What follows therefore amounts to a linear stability analysis of the morphodynamics.

Taylor-expanding equation (3) to first order gives

$$\mathbf{P}_c(t + 1) - \mathbf{P}_c(t) = M\mathbf{P}_c(t)$$

(4)

and thus seven (namely number of morphostates and thus components of any morphovector $\mathbf{P}$) equations for $t = 0, 1, \ldots, T_c - 2$, resulting in 63 equations in total, used to determine a $7 \times 7$ matrix. As the diagonal elements are determined by the columns of $M$ summing to 0 ‘only’ 42 entries are to be fitted by a weighted least square fit.

The resulting matrix $M$ can be used to predict the cell frequencies $\mathbf{P}_c(t + dt)$ at future time $t + dt$ from those at given time $t$. Distinguishing the theoretical expectation from the experimental measurements, we write

$$\mathbf{P}_c^{\text{theo}}(t + dt) = \mathbf{P}_c^{\text{theo}}(t) + M\mathbf{P}_c^{\text{theo}}(t)dt,$$

(5)

where $\mathbf{P}_c^{\text{theo}}(t + dt)$ is the theoretical prediction of the morphovector $\mathbf{P}_c(t + dt)$ at time $t + dt$ and $M$ is the matrix obtained above. To start the integration we set $\mathbf{P}_c^{\text{theo}}(0) = \mathbf{P}_c(0)$ (see also equations (8) and (9)).

To validate the transition matrix $M$, we compared the components of the predicted morphovector $\mathbf{P}_c^{\text{theo}}(t)$ with the experimental ones as a function of time (figure 6).

Remarkably, the theoretical morphovector $\mathbf{P}_c^{\text{theo}}(t)$ appear to capture most of features of the evolution of their experimental counterparts. This is quite astonishing, given the fact that the theoretical predictions are based on a linear approximation, which can be considered only a simplified model of the true underlying dynamics.

2.5. Stability analysis of the uncut morphology

Given the transition matrix $M$ describing linear perturbations around the uncut morphostate, we can now proceed with a standard analysis [21] and calculate its right eigenvectors $\mathbf{e}_i$ and eigenvalues $\lambda_i$,

$$M\mathbf{e}_i = \lambda_i \mathbf{e}_i.$$  

(6)

By construction, $M$ has one left eigenvector $(1, \ldots, 1)$ with vanishing eigenvalue $\lambda_0 = 0$, as $(1, \ldots, 1)M = 0$. The corresponding right eigenvector $\mathbf{e}_0$ with eigenvalue $\lambda_0$ represents the stationary distribution of the Markov process in equation (3), i.e. the state towards which the root is predicted to relax.

Comparing it to the temporal average of the morphostate of the uncut root,

$$\mathbf{P}_u = \frac{1}{T_u} \sum_{t=0}^{T_u-1} \mathbf{P}_u(t)$$

(7)

we found a good correspondence between the weight of the seven different morphostates. This suggests that the period of $T_c$ days, that fed into the analysis, suffices to produce a rough estimate of the asymptotic distribution of morphostates.

In general, all other eigenvectors $\mathbf{e}_i$ and eigenvalues $\lambda_i$ of $M$ are complex. As $M$ is real, their complex conjugates are linearly independent eigenvectors themselves. If $M$ has eigenvalues $\lambda_i$ and eigenvectors $\mathbf{e}_i$ with $i = 1, 2, \ldots, 6$ that span the entire subspace of $\mathbb{R}^7$ of normalised morphostate vectors, namely those orthogonal to the left eigenvector $(1, \ldots, 1)$,

$$\mathbf{P}_c(0) = \sum_{i=1}^{6} q_i \mathbf{e}_i,$$

(8)

equation (3) can be integrated to give

$$\mathbf{P}_c^{\text{theo}}(t) = \sum_{i=1}^{6} e^{\lambda_i t} q_i \mathbf{e}_i,$$

(9)

with characteristic relaxational time scales $-1/\Re(\lambda_i)$ derived from the real part $\Re(\lambda_i)$ of the eigenvalues $\lambda_i$ (table 1). They represent the time it takes for a perturbation to decay by a factor $1/e$. The imaginary part of the eigenvalues, on the other hand, give rise to a superimposed oscillatory behaviour with a period of $2\pi/\Im(\lambda_i)$. The eigenvalues come in three pairs of...
complex conjugates and any linear combination (8) that is real at time $t = 0$ will remain real under the evolution (9).

We find that all eigenvalues have negative real parts, indicating that every initial state $\mathbf{P}_0(0)$ will eventually relax to the stationary state. This is a property of the linear Markov dynamics imposed in equation (3). What is less trivial is that the relaxation time-scales derived from $\mathbf{M}$ are 0.3 to about 0.4 days and 1.2 to about 1.9 d, clearly indicating two types of relaxations: one rapid with characteristic temporal scale around eight hours, and a slower one with a scale of around one to two days. The oscillatory periods, on the other hand, are more than 66 d long. They all stretch well beyond the observation period, so that, given the exponential decay of the relative amplitude, it is not realistic to expect to observe the oscillation experimentally.

3. Discussion

We have taken an unconventional approach in analysing quantitative morphometric data extracted from regenerating plant tissue. We collected morphometric data at cellular resolution during the regeneration of cut root tips back to their uncut morphology, and proposed an innovative way to describe the morphological state of each cell, as a weighted superposition of fundamental states. We believe this is a representation that is quite effective and useful from a biological perspective, as it highlights the known fact that developmental and metabolic cellular states are fundamentally noisy. So it makes more sense to have a mathematical framework that associates each position in the tissue with a set of basic cell types, each weighted with an evolving probability of actually occurring, instead of an over-simplified view where each position is defined at any single time by a unique cell type.

From this point of view, our proposed morphovector is designed to capture the intrinsic morphological variability observed at cellular scale in any living tissue, regardless of its overall shape or other characteristics at larger scales. Although no information is provided about the tissue shape, we believe that the proposed analysis of the cell morphology dynamics underlines the changes observed at larger scales. In fact, the same approach could potentially be extended to morphologies at any scale. We believe this is one of the strengths of this approach, as it makes it potentially exportable to other organs and even other organisms.

Moreover, we could fit a theoretical transition matrix to describe a first-order approximation of the dynamics of such a morphovector, and showed that it can indeed be described as a Markovian relaxation of a morphological perturbation.

From the point of view of developmental biology, this can be interpreted as a robust morphology (stable fixed point) that resists physical damages (perturbations) by regenerating (relaxing back to) the original shape, regardless of the way it is cut, or in which direction it is pushed within the space of morphologies.

We can go further than that. Our quantitative analysis of the morphological robustness offers information about the characteristic dynamics of the tissue regeneration. For example, the real part of the eigenvalues gives us a temporal scale of the relaxation, with respect to the direction of the corresponding eigenvector. Each eigenvector can be interpreted as a special direction in the morphospace, so that when the tissue is perturbed along that direction, we can predict it will regenerate with a characteristic temporal scale given by the corresponding eigenvalue. Interestingly, our results indicate two classes of temporal scales in the specific case of Arabidopsis root tip regeneration: one of about eight hours and another of one to two days. That is a surprising and intriguing result, because it suggests the co-existence of two potentially distinct developmental processes, each characterised by a quite different temporal scale. At the moment we could only speculate what these processes might be, but in principle it should be possible to isolate genetic mutations where the two temporal scales are uncoupled. So, for example, one class of mutants could exhibit a regeneration dynamics characterized only by the short time scale, while another class would be characterized only by the slower process over a longer time scale. Crucially, we might also predict that a wild-type root could in principle be perturbed (by cuttings or other means) exactly along the eigenvector corresponding to a regeneration with a single eight hour or one to 2 d temporal scale. Finally, the imaginary components of the eigenvalues might suggest superimposed oscillatory trajectories during relaxation in the morphospace. Although in the system studied the oscillations would be on a very different time scale and are in effect not observable, we should not rule out the possibility that in other morphodynamic systems (organisms) these might become visible.

It should be noted that in this study we used only morphological data from root tips that actually regenerated and that we imposed Markovian dynamics. This means that we forced ourselves to explicitly study the morphodynamics in the basin of attraction of the single, stable fixed point representing the regenerated root tip. In fact, the probability of successful regeneration for this kind of physical damage in Arabidopsis roots is a function of the position of the incision and is always less than unity [14]. Analogous morphometric data could be collected for those roots that did not successfully regenerate. In morphospace this may correspond to a complete loss of a well-defined position if no meaningful morphometric data can be extracted from the roots or it may correspond to another fixed point. To model this, we would need to allow for a more complicated dynamics beyond simple (linear) Markovian evolution.
In the presence of another fixed point, what kind of perturbation would push the system out of the basin of attraction of the first fixed point and towards the second one? And what would be the characteristic temporal scales of the morphodynamics towards the second fixed point? At the moment, these remain open questions that we plan to address experimentally in future works.

To our knowledge, this is the first time that such a technique is applied to morphodynamics in the context of developmental biology, and we suggest it can be developed as a standard method to predict robustness of morphologies. We suggest that this approach could be applied to any kind of regenerating organism, where time-lapse images of morphological dynamics are available. In more general scenarios, where the eigenvalues take both negative and positive values, this method could be used to predict which perturbations (along which eigenvector in the morphospace) are more likely to produce a rapid regeneration (relaxation with large negative eigenvalues), slow regeneration (eigenvalues with small negative values) or no regeneration at all (positive eigenvalues).

4. Methods

4.1. Plant material

The plants used in this study were all wild-type Arabidopsis thaliana individuals of Columbia (Col-0) ecotype. Seed sterilization and synchronization were performed following standard procedure [14]. First, seeds were imbibed in water and stored at 4 °C for 2 d. The seeds were then sterilized in 50% household bleach (Sodium Hypochlorite, 5%) and 0.0005% Triton X-100 (Sigma, T8787) for 3 min and rinsed six times in sterile water. Under sterile conditions the seeds were transferred to a standard 0.8% agar solid MS medium (4.4 g l\(^{-1}\), or 1 × MS basal salts [Sigma, M5519], 0.5% sucrose, 0.05% MES hydrate [Sigma, M8250], adjusted to pH 5.7 with 1 M KOH before adding 0.8% agar [Sigma, A5040]) and germinated vertically in a growth chamber (23 °C, 120 µmol m\(^{-2}\)-s\(^{-1}\) light intensity on a 16 h/8 h light/dark cycle).

4.2. Root excision

The root tip excision was performed on primary roots 3 d post-germination, following an established protocol [14]. Plants were transferred to 5.0% agar solid medium (4.4 g l\(^{-1}\), or 1 × MS basal salts [Sigma, M5519], 0.5% sucrose, 0.05% MES hydrate [Sigma, M8250], adjusted to pH 5.7 with 1 M KOH before adding 5.0% agar [Sigma, A5040]) and the root tips were cut by hand with sterile 27G needles (Sterican) under a dissecting stereo-microscope (Nikon SMZ1000 at 4.3× magnification). The plants were then returned to 0.8% agar [Sigma, A5040], adjusted to pH 5.7 with 1 M KOH before adding 0.8% agar (Sigma, A5040) and the root tips were transferred from the microscope slide to the 0.8% agar solid medium and returned to the growth chamber.

4.3. Microscopy

Individual plants were imaged once a day for at least 9 d, following an established protocol [14]. Before each imaging session, each root was stained with filter-sterilized 10 µg ml\(^{-1}\) propidium iodide (Sigma, P4170) for 30 s to 3 min (shorter time for days closer to the root tip excision), briefly washed in sterile water and mounted in sterile water on sterile microscope slides. Stained root tips were imaged using a Leica SP5 laser scanning confocal microscope with a 63X water immersion objective, with excitation at 488 nm and emission between 579 nm and 698 nm. After imaging, the roots were transferred from the microscope slide to the 0.8% agar solid medium and returned to the growth chamber.

4.4. Image processing

All image processing routines were performed in MATLAB [22].

4.4.1. Segmentation. To identify single cells, each root was segmented in three steps: removing the background, minimizing noise and applying marker-controlled watershed segmentation (figure S3). The background was removed by hand, by drawing a mask on top of the root and setting all pixels outside of the mask to zero intensity. The noise was minimized by dividing the image by a blurred image of itself, obtained applying a filter with a kernel sized 101 × 101 pixels. The marker-controlled watershed segmentation is composed of five steps [15]: edge recognition through the Sobel method, identification of foreground markers (intensity maxima inside each cell) through opening-closing by reconstruction, identification of background markers (cell walls) by iteratively applying a global threshold, eliminating identified cells and increasing the threshold by a small step. In addition, the foreground marker list was updated with an eroded version of the cell walls found in the previous step. Finally, the foreground and the background markers were added to the image as regional minima, and a watershed-based segmentation was performed. The segmented objects were identified as cells only if the surface area of the cells was between 100 and 10 000 pixels.

4.4.2. Morphological traits. For each identified cell, three morphological traits were measured: area, eccentricity and orientation. The cell area was defined by the number of pixels this cell encapsulated. To derive a coordinate from it, the minimum cell size observed was mapped to 0 and the maximum to 1. The cell eccentricity was defined as the eccentricity of an ellipse with the same second-moments as the cell, and then normalised between 0 (minimum measured eccentricity) and 1 (maximum measured eccentricity).

The cell orientation was defined as the angle \( \theta \in [0, \pi] \) between the middle line of the root prior to registration and the major axis of the ellipse with the same second-moments as the cell (figure S4(a)).

The cell orientation ought to be continuous and periodic with a period of \( \pi \) (figure S4(b)). The angle \( \theta \) returned by the code as the orientation of a cell lies on the interval \([0, \pi]\). Two seemingly very different orientations of \( \theta = \epsilon \) and \( \theta = \pi - \epsilon \) are in fact identical in the limit \( \epsilon \to 0 \). For the clustering to consider these two orientations to be similar for small \( \epsilon \), a suitably periodic, continuous mapping of the angle

\[
\begin{align*}
\theta &\rightarrow \pi - \theta \\
\theta &\rightarrow \theta - \pi
\end{align*}
\]
is needed. In the present work, we have used the two coordinates $c = \cos(2\theta)$ and $s = \sin(2\theta)$. These two coordinates are located on a unit circle.

To account for the presumed symmetry of the root about its central axis, the orientation was recorded in conjunction with the relative position of the cell’s centroid in relation to the central axis. As far as the orientation enters into the determination of a cell’s morphostate, a cell located to the right of the central axis and having angle $\theta$ with it is considered to have the same orientation as a cell to the left with angle $\theta' = \pi - \theta$ (figure S4(c)). On the other hand, cell orientations of two cells on the same side with angles $\theta$ and $\theta' = \pi - \theta$ respectively should generally be considered as different. The mirror symmetry is thus accounted for by recording the original angle $\theta$ for cells to the right, but $-\theta'$ for cells on the left. If $\theta' = \pi - \theta$ (figure S4(c)) the angles recorded are therefore $\theta$ and $\theta - \pi$, which means that both are effectively identical, given the periodicity $\pi$ of the orientation.

Together, the area and the eccentricity both on unit intervals $[0, 1]$, and the coordinates $(c, s)$ on a unit circle, place the set of morphological traits of a cell on the surface of a hyper-cylinder with a unit circle as the base.

4.4.3. Registration. To create a coordinate reference, a middle line was drawn on each root by linking the very root tip to the QC and continuing proximally in the middle of the root and parallel to its main axis (figure S5(a)). The position of a cell was then defined with a radial coordinate $r$ given by the distance of the centroid of the cell to the root middle line just defined, and with a height coordinate $h$ given by the distance between the QC and the centroid’s projection on the middle line (positive when proximal and negative when distal). Radial coordinates $r$ were then normalized to $\bar{r} = \frac{r}{R}$ with $R$ the maximum radius of the root. Each root was rotated and translated such that the middle line was vertical (purple line in figure S5(a)) and that all QCs would overlap in the same position. The entire middle line (purple and green in figure S5(a)) was then drawn vertical and each centroid was translated accordingly, to maintain their original $(\bar{r}, h)$ coordinates. Finally, new virtual cells were created using Voronoi tessellation [23] around the centroids (figure S5(c)). This registration step was important for the calculation of the morphovector $\mathbf{p}(\bar{x}, t)$ described above, as each Voronoi cell in the registered image is associated with a centroid of a cell in the original image, so that each pixel $\bar{x}$ within a Voronoi cell can be associated with the morphological traits of that cell. These morphological traits are not distorted by the registration as they have been taken prior to it.

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