On the mechanical origins of waving, coiling and skewing in *Arabidopsis thaliana* roots

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We present theory and simulations of the well-known waving, coiling and skewing patterns exhibited by roots of *Arabidopsis thaliana* when grown on inclined agar substrates. By modelling growing slender organs as Cosserat rods that mechanically interact with the environment, our simulations confirm previous hypotheses of waving and coiling arising from the combination of active gravitropism and passive root-plane interactions. Skewing is instead related to intrinsic twist due to cell file rotation. Numerical investigations are outfitted with an analytical framework that consistently relates transitions between straight, waving, coiling and skewing patterns with tilt angle. Simulations are found to corroborate theory and recapitulate a host of reported experimental observations, thus providing a systematic approach for studying *in silico* interactions of plant organs with their environment.

The ability of roots to grow in soil is critical for plant health and crop yield, enabling the uptake of water and nutrients, providing anchorage and stability of eroding soil (1–3). This is no small feat. Indeed, soil is a highly heterogeneous environment, characterized by non-uniform concentrations of resources as well as obstacles such as rocks and compacted soil. In negotiating with the environment, plant roots combine passive physics and active growth-driven mechanisms, termed tropisms, whereby dedicated organs sense stimuli such as water (hydrotropism) or gravity (gravitropism), and redirect growth appropriately. While dynamics of tropisms are generally understood (4, 5), and aspects of root mechanics and their effect on growth have been described (6–8), a consistent framework to dissect the interplay between passive and active responses, heterogeneous environments and ensuing growth patterns, is still missing (9, 10). The essence of this complex interaction is on display in controlled experiments of roots growing on inclined agar gel substrates, whereby remarkably different behaviors are observed to emerge: roots grow straight on vertical planes, grow in waving patterns as tilt increases, skew in some cases, while growing on inclined agar gel substrates, whereby remarkably different behaviors are observed to emerge: roots grow straight on vertical planes, grow in waving patterns as tilt increases, skew in some cases, and eventually coil when tilt approaches the horizontal plane (Fig. 1a). These growth patterns are well-documented in genetically driven phenotypes of *Arabidopsis thaliana* (11–17). Although insightful, such genetic approaches are limited to given species, and do not formally address the fundamental role of root and environmental mechanics. Thus, while gravitropic responses and root-substrate mechanics clearly play a role (13), underlying mechanisms remain subject of debate. For instance, both circumnutation (12, 18, 19), the intrinsic circular movements of root tips, and thigmotropism (11, 20), the active response to touch, have been suggested to be at the basis of waving and skewing patterns. Here, we combine advances in the modeling of growing rod-like organs (21–30), and 3D numerical simulations (31) to gain broader insight.

**Results**

**Modeling of root mechanics and growth.** In modeling the waving and coiling experiments of *Arabidopsis thaliana*, we begin by assuming separation of timescales between slow growth-driven root responses and fast mechanical relaxation (Fig. 1d), allowing us to decouple the two processes in a quasi-static manner (22, 23).

Slender roots are represented as Cosserat rods (31), which are 1D elastic continuous elements able to undergo all modes of deformations — bending, twisting, stretching, shearing — and reconfigure in 3D space (Fig. 1b). We mathematically describe a slender rod by a centerline \(r(s,t) \in \mathbb{R}^3\) and a rotation matrix \(Q(s,t) = \{d_1, d_2, d_3\}\), providing a local material frame. When the rod is at rest, its length is \(L\) and the corresponding material coordinate is \(S \in [0,L]\), with \(t\) and \(s \in [0,l]\) denote the length and arc-length of the deformed (stretched) filament, and \(t\) is time. If the rod is unsheared, \(d_3\) points along the centerline tangent \(\partial d_3 / \partial s\), while \(d_1\) and \(d_2\) span the normal–binormal plane, i.e. the cross-section. Shearing and stretching shift \(d_3\) away from \(\partial d_3 / \partial s\), quantified by the shear vector \(\sigma = (\lambda_c \partial d_3 / \partial s - \lambda_c d_3)\), where \(\lambda_c = \partial S / \partial s\) is the local elastic stretch. The curvature vector \(\kappa = \kappa_1 d_1 + \kappa_2 d_2 + \kappa_3 d_3\) encodes the rotation rate of the local frame along the material coordinate so that \(\partial d_3 / \partial s = \kappa \times d_3\). We define the bending stiffness matrix \(B\) and shearing stiffness matrix \(S\) in the rest configuration. Then, the mechanical equilibrium of a rod-like root is described by

\[
\frac{\partial n}{\partial s} + f = 0 \quad (1)
\]

\[
\frac{\partial m}{\partial s} + \frac{\partial r}{\partial s} \times n + c = 0 \quad (2)
\]

where \(n = S(\sigma - \sigma^0)\) represents internal forces, related to shear and elastic stretch of the centerline, \(m = B(\kappa - \kappa^0)\) represents internal torques, related to bending and twisting, and \(\sigma^0\) and \(\kappa^0\) are intrinsic strains. As detailed in (31), incompressibility is incorporated by rescaling \(B\) and \(\kappa\) through \(\lambda_c\) such that \(B \rightarrow B / \lambda_c^2\), \(S \rightarrow S / \lambda_c\) and \(\kappa \rightarrow \kappa / \lambda_c\).
Lastly, external forces $f$ and couples $c$ (per unit length) capture overall environmental effects.

Next, we outfit this model with active gravitropic dynamics driven by differential growth (24, 35, 36). Axial growth is implemented via the additional stretch factor $\lambda_g$, defined with respect to an initial material coordinate $S_0 \in [0, L_0]$, such that $L_0$ is the initial length and $\lambda_g = \partial S / \partial S_0$ (Fig. 1d). By denoting time derivatives as $\frac{dx}{dt} = \dot{x}$, we introduce the average relative growth rate as $\dot{\lambda}_g = \lambda_g / \lambda_g$. Root growth is restricted to a finite growth zone of length $L_{gz}$ extending from the tip (Fig. 1e), reflecting experimental evidence (7, 37, 38).

Tropic movements are directed by environmental cues, which are generally perceived via dedicated sensory systems. Gravity in particular is sensed via specialized cells close to the tip (39), and translated into a redistribution of growth hormones across the root cross-section. This results in axial differential growth, which in turn leads to a change in the root’s curvature, redirecting the organ towards the stimulus. This machinery is mathematically captured via the local differential growth vector $\Delta(s, t)$ (24, 35, 36)

$$\Delta(s, t) = \beta \dot{g} - \gamma R \kappa(s, t) \hat{N}(s, t)$$

where the first term represents the gravity stimulus, with $\dot{g}$ the direction of gravity and $\beta$ the gravitropic gain or sensitivity (35). The second term represents the proprioceptive response. This can be thought of as a counter-curved response that balances the gravitropic one, based on the sensing of the organ’s own shape (35). Here $\gamma$ is the proprioceptive gain, $\kappa = \sqrt{\kappa_1^2 + \kappa_2^2}$ is the norm of the bending curvature, and $\hat{N}$ is the normal director of the center-line’s Frenet-Serret frame (Fig. 1c). Given the average growth rate $\dot{\lambda}_g$ and the organ radius $R$, we connect the change in local intrinsic curvature $\kappa^0(s, t)$ within $L_{gz}$, to the differential growth $\Delta(s, t)$ projected on the cross section (24, 35, 36)

$$\kappa^0 = \frac{\dot{\lambda}_g}{R} (\hat{d}_3 \times \Delta)$$

defined in SM 2. For simplicity, here we drop the explicit dependence on $(s, t)$. Thus, Eqs. 3 and 4 capture gravitropic dynamics, relating sensing $(\beta \dot{g})$ to differential growth $(\Delta)$, which in turn feeds into the root shape ($\kappa^0$). As described in the Methods section, $\Delta$ can be rewritten to include different types of internal and external cues, such as phototropism or circumnutations, as well as different sensing profiles (apical and local). Finally, to distill the essential mechanisms under-
lying the range of observed growth patterns, we neglect the effect of substrate deformations (13, 40), and the coupling between root growth, static friction, and stick-slip dynamics (38, 41, 42). These elements nonetheless deserve future attention.

Some Arabidopsis mutants (43–45) exhibit a cell profile which twists during growth (Fig. 4b). We model this additional effect by allowing the material frame in the growth zone $L_{\text{gz}}$ to twist around the centerline of the organ. This twist is described by an increasing register angle $\xi$, characterized by the angular velocity at the apex

$$\omega = \frac{\partial \xi}{\partial t} = \tau \dot{e}_g L_{\text{gz}}$$

where $\tau$ is a constant intrinsic twist line density (Fig. 4c).

Overall, Eqs. 1-5 define our model, which we numerically discretize and solve using the open-source software Elastica (46), demonstrated across a range of biophysical problems (47–52) entailing fiber-like structures (Methods and SM 3).

**Simulations recapitulate waving and coiling patterns.**

Based on waving and coiling experiments (11–17), we simulate Arabidopsis thaliana roots growing on a tilted plane for a range of angles $\alpha$ (from vertical to horizontal), as well as gravitropic and proprioceptive gains $\beta$ and $\gamma$. We use parameters and mechanical properties typical of Arabidopsis (Table 1 in Methods). For simplicity we assume frictionless growth while fixing the sessile (mature) zone to the substrate, emulating real roots attachment (13, 40) due to root hairs (Fig. 1a). At this stage, we neglect root twisting during growth, although we will revisit it in later sections.

As illustrated in Fig. 2a and supplementary movies 1 and 2, simulations capture observed root behavior. Indeed, simulated roots grow straight on a vertical plane ($\alpha = 0$), and transition to waving as the substrate tilt angle reaches $\alpha \approx 15^0$, consistent with experimental measurements (Fig. 1a). A further increase in tilt angle initiates a second phase transition, around $\alpha \approx 50^0$, and roots begin to coil, again consistent with experiments (Fig. 1a). Transition angles are also consistent with experimental values (40) (Fig. 3f), as we shall elaborate later.

To gain intuition about the waving and coiling process, as well as their transitions, we analyze the system from an energetic and topological perspective. Tip dynamics are tracked via the planar angle $\theta^{\text{tip}}$. This is the angle between the projections of gravity and the tangent of the apex on the tilted plane (Fig. 2a and c). The evolution of the stored energy in the growth zone is described through the approximate elastic bending energy $E_b = \int_{L_{\text{gz}}} m \cdot (\delta k_1 d_1 + \delta k_2 d_2) ds$ and elastic twist energy $E_t = \int_{L_{\text{gz}}} m \cdot \delta k_3 d_3 ds$, where $\delta k = k/\lambda - k^0$ and $m = B\delta k/\lambda_s^2$ (Fig. 2d).

Additionally, in order to concisely capture root morphology and its reconfigurations, we consider the topological quantities writhe (Wr) and twist (Tw) in the growth zone (Fig. 2e). Writhe is computed by treating the growth zone as an open loop (53), and is used here as a measure of out-of-plane bending of the growth zone’s centerline. If Wr = 0, then the centerline is in-plane, while positive or negative values of Wr correspond approximately to right- or left-handed out-of-plane bending, respectively (Fig. S2). Twist instead determines the rotation of the local frame around its centerline. Positive and negative values correspond to right-handed and left-handed rotations accordingly (Fig. S2).

In a closed or infinitely long rod, the geometric integral quantities Wr and Tw are related topologically, and their sum is constant and constrained by the number of formed loops (54). In our case their sum is not conserved, however their topological connection is useful to qualitatively understand the system dynamics.

As can be seen in Fig. 2a, the onset of waving and coiling follows a first turning event. Fig. 2b shows snapshots at different stages of a waving organ, animated in supplementary movie 1 and marked for reference in Figs 2c-e. Snapshot (i) occurs just before the first turning event, where the root grows in a plane defined by its initial tangent and the direction of gravity, as represented by the constant values of $\theta^{\text{tip}}$ (Fig. 2c), Wr and Tw (Fig. 2e). Due to gravitropism, the root tends to grow into the substrate (Fig. 1d), which in turn resists to growth, leading to an increase in bending energy $E_b$ (Fig. 2d). This energy is eventually released, breaking symmetry, in the form of an elastic instability - snapshot (ii) - whereby the root tip slips to one side, bending and twisting. This process is reflected by a sharp drop in $E_b$ accompanied by a jump in all other parameters (Fig. 2c-e).

We continue to follow the growth of the root during a subsequent turning event. Snapshot (iii) occurs right after the second turn where the tip angle $\theta^{\text{tip}}$ is at a maximum, and the bending energy $E_b$ is at a minimum. For further insight, it is useful to decompose $\Delta$ into components parallel and perpendicular to the substrate. The parallel component causes the root to reorient along the projection of gravity on the plane, leading to a decrease in the planar tip angle $\theta^{\text{tip}}$. The perpendicular component pushes the apex into the plane, leading again to an increase in $E_b$, similar to the initial symmetry breaking and in line with observations of substrate deformations (13). As new cells are produced at the tip, older cells stop growing and become part of the mature zone. Thus, accumulated twist leaves the growth zone, so that, at snapshot (iv), $E_t$ reaches a minimum while $E_b$ reaches a maximal value. Then, as for the first bending event, the organ releases bending energy by twisting, and $E_b$ is converted into $E_t$. This elastic relaxation is accompanied by a conversion of Wr into Tw, since these quantities are topologically related (53). Both $E_t$ and Tw reach a maximum value at snapshot (v). At this point Tw is converted out of the growth zone faster than it is generated, and $E_t$ and $E_b$ decrease. The organ is now back to stage (iii), and the process repeats itself. Since the gravitropic correction of curvature is slower than the elastic relaxation, $\theta^{\text{tip}}$ repeatedly overshoots in the direction of gravity, producing the observed waving patterns.

We note that our simulations show that neither thigmotropism nor circumnutation are strictly required in order to produce waving and coiling, unlike previously argued (11, 12, 18–20).
Scaling analysis. We now perform a scaling analysis of the growth dynamics described in Eqs. 3 and 4, and relate model parameters to experimentally measurable variables: wavelength \( \lambda \) and amplitude \( A \) of root waving patterns, as defined in Fig. 3d. Comparing predicted values to experimental measurements will allow to quantitatively corroborate our model. In order to derive relations for amplitude and wavelength, we focus on the curvature of the waving pattern. Simulations (Fig. 2) show that the elastic stresses associated with bending, due to the organ pushing into the substrate, are accumulated until they relax by twisting sideways, which rotates the curvature vector. This suggests that the maximal value of the in-plane curvature of the waving pattern \( \kappa \), obtained right after twisting, may be comparable to the maximal curvature \( \kappa^0 \) induced by the gravitropic response in a free organ tilted at the same angle \( \alpha \) of the substrate. Thus, we hypothesize that

\[
\max \kappa \approx \max \kappa^0
\]

In SM 6 we provide a detailed analysis supporting this assumption, decomposing the dynamics of a tilted organ into components parallel and perpendicular to the tilted plane.

This assumption is also supported by Fig. 2c, which compares the oscillating tip angle of a waving pattern of a root on substrate (blue) to the gravitropic bending of a corresponding free organ (pink). This comparison further suggests that the wavelength \( \lambda \) of the waving pattern is dictated by the damped oscillations inherent to the free organ’s gravitropic dynamics (57), while the amplitude \( A \) is modulated by the mechanical interaction with the substrate.

Therefore, in order to find \( \lambda \), we first evaluate the characteristic gravitropic turning period \( T_0 \), corresponding to the time a tilted free root takes to turn towards the direction of gravity (21, 35). We rewrite Eq. 4 in non-dimensional form, assuming \( \gamma = 0 \) for simplicity, and rescaling length and curvature by \( L_{gz} \), yielding

\[
\frac{d}{dt}(L_{gz}\kappa^0) = \varepsilon_g L_{gz} \beta (d_3 \times \hat{g})
\]

Values in parentheses are dimensionless, as well as the coefficient \( L_{gz} \beta \). Since \( d_3 \) and \( \hat{g} \) are unit vectors, we expect the turning time \( T_0 \), related to the rate of change of the normalized curvature \( \frac{d}{dt}(L_{gz}\kappa^0) \), to scale as a function of \( L_{gz} \beta \), with \( 1/\varepsilon_g \) the overall timescale of the problem. Based on
This final relation provides an estimate for \( \max \kappa_0 \), which together with Eq. 6 relates the waving curvature to the gravitropic curvature. In Fig. 3a we validate Eq. 6 for a range of tilt angles by plotting \( \max \kappa \), the maximal curvature measured from simulated waving patterns, against \( \max \kappa_0 \), the analytically predicted maximal curvature for a free tilted organ in Eq. 8, finding good quantitative agreement.

Having verified that the timescale of the waving pattern is related to the gravitropic bending, we can approximate the length \( \lambda/2 \) grown during a turning event (Fig. 3d)

\[
\lambda \approx 2A \approx \frac{L_{gz}}{R} \frac{v_g^{tip}}{T_0 \tan(\alpha)} \tag{9}
\]

where \( v_g^{tip} \) is the tip growth speed. Then, following simple geometric arguments, we relate the amplitude \( A \) to the wavelength. After a turn event, \( 2A \) and \( \lambda/2 \) form an orthogonal triangle (Fig. 3d). Based on Eq. 6, and since in the gravitropic response of a free organ the angle between the base and the tip is equal to the tilt angle \( \alpha \) (Fig. 1d), we can infer

\[
2A \approx \frac{L_{gz}}{R} \frac{v_g^{tip}}{T_0} \tan(\alpha) \tag{10}
\]

In Figs. 3b-c, we numerically test the scaling relations for both \( \lambda \) (Eq. 9) and \( A \) (Eq. 10) and compare with experiments (12, 40, 55, 56), assuming characteristic parameters from literature (Table 1 and SM 5). We assume local sensing throughout, and we report results for apical sensing in SM 7. The second column represents the configurations shown in Fig. 2a. Equivalent configuration spaces for high \( \gamma \) and apical sensing in SM 7. (f) Observed coiling transition. Experimental observations of coiling probability (40) agree with model prediction (represented by background colors as in (e) for \( \beta L_{gz}/R = 0.5 \)).

Transitions between straight, waving and coiling. Here we provide an intuitive rationale for the two critical tilt angles
at which the root transitions from straight to waving (αs→w) and from waving to coiling (αw→c), with the full derivation reported in SM 7.

Simulations of Fig. 2 suggest that waving patterns are initiated by an elastic instability, which occurs when it becomes energetically convenient to twist rather than to bend, i.e. \( E_t < E_b \). Again assuming that the intrinsic curvature is captured by the gravitropic response of the free organ \( \max \kappa_i^0 \) (Eq. 8), before the turn we can approximate \( E_t \propto \max \kappa_i^0 \). The accumulated angle during the gravitropic response is roughly equal to the tilt angle \( L_{gz} \max \kappa_i^0 \approx \alpha_t \), meaning that \( E_t \) increases with tilt angle. However \( E_t \) is constant, and at a critical angle \( \alpha_{s\rightarrow w} \) the two energies intersect \( E_f = E_b(\alpha_{s\rightarrow w}) \) and waving occurs. The critical angle (detailed in SM 7) follows

\[
\sin(\alpha_{s\rightarrow w}) = \sqrt{\frac{2}{3} \frac{2}{\pi} \frac{\beta L_{gz}}{R} - \frac{\bar{\gamma}}{3}}
\]

(11)

where \( \Delta \xi \) is a fitting parameter representing the change in the register angle \( \xi \) created by the integrated elastic twist (Fig. 1c).

The transition between waving and coiling is instead related to the ratio between (i) the time to turn towards gravity after a twist induced relaxation and (ii) passive orientation drift (21) – the rate at which the tip angle of a curved organ increases due to growth, maintaining the same curvature, thus reorienting growth uphill in this case. This is proportional to the amplitude of the waving pattern \( T_0 \tan(\alpha) \) in Eq. 10, and the latter follows \( T_{dr} \approx 1/(\xi L_{gz} \max \kappa) \) (21). While the turning time increases with \( \alpha \), passive orientation drift is constant. Therefore at a critical angle \( \alpha_{w\rightarrow c} \) orientation drift occurs faster than turning, \( T_{dr} < C T_0 \tan(\alpha) \), and coiling ensues. The coiling critical angle (detailed in SM 7) then follows

\[
\sin^2(\alpha_{w\rightarrow c}) = \frac{4}{C \pi^2} \left( \frac{\beta L_{gz}}{R} \right)^{1/2}
\]

(12)

where \( C \) is a fitting parameter.

We corroborate the predicted critical angles both numerically and experimentally. Fig. 3e shows the configuration space of simulated organs over a range of \( \alpha \) and effective gravitropic bending \( \beta L_{gz}/R \) (where for Arabidopsis roots we use 0.5, see Table 1). For each set of parameters we compare the conformation observed in simulations (■ symbols for straight, ◆ for waving, ● for coiling) to the predicted one (represented by background colors, separated by the critical angles), with \( \Delta \xi = \pi/8 \) and \( C = 0.5 \), finding good agreement. Configuration spaces for cases with \( \gamma = 1 \), and apical sensing, can be found in SM 7. In Fig. 3f we also compare our results to the coiling probability measured in vivo by Zhang et al. (40), again finding good quantitative agreement.

**Intrinsic twist yields skewing patterns.** Lastly, we focus on the skewed waving patterns observed in some Arabidopsis mutants (20, 33, 58, 59) (Fig. 1a). Observations suggest that the skewing angle is related to the twisting of root cell files (e.g. Fig. 4b), representing an intrinsic twist of the material frame around the centerline of the growth zone, as described in Eq. 5 and detailed in the Methods (Fig. 4c). When accounting for such rotations (\( \omega \) in Eq. 5), simulations recover observations of skewing (Fig. 4a and supplementary movies 3-5). We see how increasing the ratio between the bending time and the twisting velocity, \( \omega T_0 \), results in larger skewing angles and washes out the waving pattern. This is reflected in the evolution of energetic and topological variables: \( \omega T_0 = 1 \) (Fig. 4d-f) exhibits oscillatory behavior similar to that of regular waving patterns, while \( \omega T_0 = 8.5 \) (Fig. S10) transitions to a more monotonic behavior. Here the initial breaking of symmetry is not a mechanical instability but a smooth, twist-induced rotation. We find that circumnutations are not required, however they could have a secondary role, as we find that with no intrinsic twist circumnutations lead to disordered patterns and low skewing angles (SM 8), explaining observed coiling in agravitropic roots (12).

Next, we relate the average skewing angle \( \langle \theta \rangle \) (defined in Fig. 2a) to \( \omega \), for two limiting cases. When \( \omega T_0 \ll 1 \), the apical rotation due to intrinsic twist acts as a small perturbation to the waving mechanism, accelerating turns in the same direction, while slowing down opposite turns (Fig. 4d-f). Thus, the amplitude of the turn increases in one direction, \( A_+ \approx A(1 + \omega T_0) \), and decreases in the other, \( A_- \approx A(1 - \omega T_0) \). From trigonometric arguments \( \sin(\theta) \) is the horizontal displacement due to the difference in amplitude over two turns, divided by the wavelength \( \sin(\theta) \approx \Delta A/\lambda \), with \( \Delta A = 2A_+ - 2A_- = 4A_0 \omega T_0 \). Substituting the expressions for \( \lambda \) and \( A \) in Eqs. 9-10 yields

\[
\lim_{\omega T_0 \ll 1} \sin(\theta) \approx \omega T_0 \tan \alpha
\]

(13)

In the opposite limit \( \omega T_0 \gg 1 \), the twist-induced rotation dominates, and the waving pattern disappears altogether. Following simple geometric arguments (Fig. 1d), the skewing angle \( \langle \theta \rangle \) reduces the in-plane gravitropic component \( \beta_\parallel = \beta \cos \alpha \sin(\theta) \), however does not affect the perpendicular component directed into the plane, \( \beta_\perp = \beta \sin \alpha \), which causes the apex to drift uphill when twisted into the plane. The constant skewing angle suggests that these two competing processes balance each other, \( \beta_\perp = \beta_\parallel \), yielding

\[
\lim_{\omega T_0 \gg 1} \sin(\theta) \approx \tan \alpha
\]

(14)

We corroborate our skewing model, comparing simulations with a number of experimental observations of different mutants (20, 33, 58, 59), as well as the limiting relations predicted in Eqs. 13 and 14 (Fig. 4g). As can be seen, despite the anticipated experimental spread, our models do capture general trends.

**Conclusions**

We developed a numerical framework for simulating the interaction of plant roots with solid interfaces, integrating passive mechanics and active growth-driven mechanics. Our methods are shown to reproduce the range of growth patterns exhibited by Arabidopsis roots grown on a tilted plane,
a well documented biophysical plant model. Our simulations illustrate how the interplay between active gravitropic response and passive elasticity are the minimal requirements to generate straight, waving and coiling patterns, while the addition of intrinsic twist is responsible for skewing. We find that neither thigmotropism nor circumnutation are required to recover waving or skewing patterns, as previously argued (12, 18, 19, 28), however they could play a secondary role.

A scaling analysis reveals that the amplitude of waving patterns is modulated by the mechanical interaction with the plane, while the wavelength depends on oscillations inherent to the gravitropic dynamics of a free organ, regardless of the interaction with the plane. Based on this analysis we develop analytical expressions for the critical tilt angles for which the root behavior transitions from straight to waving and then coiling. Further, we elucidate the relation between skewing angles and intrinsic twist. We corroborate all these analytical insights by comparing model predictions to both simulations and experimental observations, obtaining good agreement.

This framework provides, effectively, an in silico laboratory to form and test hypotheses relative to the behaviors of plants and their mechano-sensory machinery in realistic environments. Future extensions will consider heterogeneous terrains via the inclusion of granular mechanics. Finally, we note that since our approach is general and agnostic to the underlying biological building blocks, it can be applied to growth-driven systems other than plant organs, such as neurons, fungal hyphae, and the new generation of growing robots (60–63).

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**Materials and methods**

Here we consider a plant organ as a slender rod, whose centerline is parameterized by $s$, with $s = 0$ at the base and $s = l$ at the tip, $l$ being the organ length (Fig. 1b). In order to describe its dynamics, we assume a separation of timescales between slow growth-driven processes and fast elastic relaxation (22, 23). At each time step of our integration scheme we first update the configuration of the organ according to the growth process alone, and then allow it to relax mechanically following Cosserat rod theory (31) (Fig. 1d). For the sake of clarity, the intermediate stress-free configuration is parameterized with $S$, in order to differentiate it from the final relaxed configuration described with $s$. The initial con-
figuration is marked with $S_0$. In what follows we bring the governing equations describing the growth dynamics and the mechanical relaxation.

**Shape.** At each point $s$ along the centerline we define the position vector $r(s,t)$, which provides the 3D description of the rod at time $t$ (Fig. 1b). We also define a local orthonormal material frame $\{d_1(s,t), d_2(s,t), d_3(s,t)\}$, where $d_1$ and $d_2$ span the cross-section of the organ, and in a shearless and inextensible system $d_3$ coincides with the tangent of the centerline. The local curvature vector $\kappa(s,t)$ (or Darboux vector) of the center-line is defined through the relation $\frac{\partial d_1(s,t)}{\partial s} = \kappa(s,t) \times d_1(s,t)$ (31). The components of curvature projected along the principal vectors of the material frame ($\kappa = \kappa_1 d_1 + \kappa_2 d_2 + \kappa_3 d_3$) coincide with bending ($\kappa_1, \kappa_2$) and twist ($\kappa_3$) strains in the material frame. Introducing elastic stretch may lead to a difference between the arc-length configuration $s$ and the stress-free configuration $S$, described by the local stretch (31) $\lambda_c(s,t) = \frac{\partial s}{\partial S}$. Shear may leads to an incongruity between $d_3$ and the tangent of the center-line, and the local deviation is described by the translation vector $\sigma(s,t) = \lambda_c(s,t) \frac{\partial r(s,t)}{\partial S} - d_3(s,t)$ (Fig. 1b).

**Growth.** Root growth occurs in the growth zone, an area of length $L_{gz}$ just below the root tip. Cells divide at the tip, and elongate within the growth zone, until stop elongating and reach the mature zone (Fig. 1e). The initial configuration then represents the material (or Lagrangian) coordinate which flows due to growth with velocity $v_g$, which is the integral of the mean axial relative growth rate $\dot{\varepsilon}_g \geq 0$

$$v_g(S,t) = \dot{S}(S_0,t) = \int_0^S \dot{\varepsilon}_g(u,t)du$$

Here the growth rate is the time derivative of the logarithmic growth strain $\dot{\varepsilon}_g = \ln \lambda_g$, where $\lambda_g \equiv \frac{\partial S}{\partial S_0}$ is the growth stretch. The increase in rest lengths enters the dynamics via the actual and intrinsic stretch vectors, defined as $\sigma = (\lambda_c \frac{\partial S}{\partial S} - \lambda_g d_3)$ and $\sigma^0 = -(\lambda_g - 1) d_3$ respectively, such that $\sigma - \sigma^0 = (\lambda_c \frac{\partial S}{\partial S} - \lambda_g d_3)$. As described in the main text, growth-driven movements are generally classified into tropic and nastic movements, where the former are due to external stimuli, such as gravitropism, and the latter are due to internal cues, such as the oscillatory movements of circumnutations. Changes in curvature are due to the anisotropic redistribution of growth hormones, leading to asymmetric growth along the cross-section captured by the differential growth vector $\Delta(s,t)$ (24). The dynamics resulting change in intrinsic curvature $\kappa^0$ for $L(t) - L_{gz} \leq s \leq L(t)$ within the growth zone follows (24)

$$\kappa^0 = \frac{\dot{\varepsilon}_g}{R}(d_3 \times \Delta)$$

Here $\kappa^0 = \frac{\partial \kappa^0}{\partial t} + v_g \frac{\partial \kappa^0}{\partial S} = \frac{D \kappa^0}{Dt}$ is a material derivative which accounts for the growth of the centerline. For simplicity we assume no intrinsic twist, $\kappa^0_3 = 0$. The cross product $d_3 \times \Delta$ represents a projection of $\Delta$ on the local cross-section of the organ (details in SM 2).

The tropic and nastic movements then contribute to the total differential growth vector $\Delta$. In the case of gravitropism we have

$$\Delta_g(s,t) = \beta R(s,t) \dot{\g} - \gamma Rx(s,t) \hat{N}(s,t)$$

(17)

The first term represents the gravity stimulus, with $\dot{\g}$ the direction of gravity and $\beta$ the gravitropic sensitivity or gain reflecting the variance in the magnitude of gravitropic responses across different species (35). The second term represents proprioception which can be thought of as a counter-curving response (35), where $\gamma$ is its gain, $R$ is the radius of the organ, $\kappa = \sqrt{\kappa_1^2 + \kappa_2^2}$ is the norm of the bending curvature, and $\hat{N}$ is the normal director of the Frenet-Serret frame. The matrix $R(s,t)$ in the gravitropic term represents the sensing mechanism: in the case of local sensing used here $R(s,t) = I$, and in the case of apical sensing where sensing occurs at the tip alone we set $R(s,t) = Q(s,t)Q^T(l(t),t)$. This rotation matrix takes vectors from the material frame at the apex, $s = l(t)$, to points along the growth zone $s$, and assures that the directional signal sensed at the apex is instantaneously transferred to every arc-length $s$. Results of simulations with apical sensing are described in SM 5 and SM 7. We model circumnutations as resulting from differential growth rotating around the center-line without introducing twist (24, 25, 27, 36)

$$\Delta_{CN} = \lambda(\cos(\Omega t)) \hat{\mathbf{m}}_1 + \sin(\Omega t) \hat{\mathbf{m}}_2,$$

(18)

where $\lambda$ is the circumnutation gain, $\Omega$ is its temporal angular frequency and $\hat{\mathbf{m}}_1$ and $\hat{\mathbf{m}}_2$ are unit vectors of a normal development of the centerline (24). We assume that the two processes are additive, allowing to take $\Delta = \Delta_{CN} + \Delta_g$.

**Adding a twisting material frame.** Various Arabidopsis mutants exhibiting skewing angles also seem to exhibit a twisting cell profile (43–45), which can be interpreted as a twisting material frame around the centerline of the organ, described by the relative angle $\xi = \arccos(\hat{\mathbf{m}}_1 \cdot \mathbf{d}_1)$ (Fig. 1c). Based on experimental observations, we chose a twist profile where the angle between the cell files and the root axis increases from the apex, where it is zero, to the mature zone where it reaches a constant value (Figs. 4b-c). This is implemented using a linear profile for the arc-length derivative of $\xi$

$$\frac{\partial \xi(S,t)}{\partial S} = \left\{ \begin{array}{ll} \frac{\tau}{L_{gz}} (S - (L(t) - L_{gz})) & S \in GZ \\ \frac{\tau}{L(t)} & S \in MZ \end{array} \right.$$ 

(19)

with GZ is the growth zone $L(t) - L_{gz} \leq S \leq L(t)$, and MZ is the mature zone $0 \leq S < L(t) - L_{gz}$, where $L(t)$ is the length of the organ and $L_{gz}$ is the length of the growth zone. Here $\tau$ represents twist, where a positive or negative value represents a right or left-handed twist respectively. The intrinsic angle between the cell files and the root axis is $\arctan\left( R \frac{\partial \xi}{\partial S} \right)$ with a maximal value in the mature zone of $\arctan(R\tau)$. The angular frequency of rotation of the tissue at the tip follows $\omega = \frac{\partial \xi(L(t),t)}{\partial \tau} = \tau \dot{\varepsilon}_g L_{gz}$ clockwise.
(looking from the base to the apex, see Fig. 4c). When incorporating twist into the dynamics of Eq. 4, only the component in the direction of \( d_3 \) is affected, reading

\[
\kappa \frac{d}{dt} \frac{d}{dS} \xi(S,t) = \frac{d}{dt} \frac{d}{dS} \xi(S,t).
\] (20)

See more details in SM 2.

**Mechanics.** Mechanical equilibrium is achieved at each time step when the time independent Cosserat rod equations are fulfilled along the centerline (31)

\[
\frac{\partial n}{\partial S} + f = 0 \tag{21}
\]

\[
\frac{\partial m}{\partial S} + \frac{\partial r}{\partial S} \times n + e = 0 \tag{22}
\]

where \( n \) are the internal contact forces, \( \tau \) are the internal torques or bending moments, \( f \) are the external forces per unit length and \( e \) are the external couples per unit length, all functions of \( s \) and \( t \). The full elastic relaxation dynamics of the second step are based on the Cosserat model with a dissipation mechanism, as described in (31) and SM 3B. We follow (31) and choose a constitutive model that assumes linear elasticity, namely the internal contact forces are linearly related to the shear and stretch of the centerline, and the internal torques are linearly related to the bending and twisting

\[
n = S (\sigma - \sigma^0) = S \left( \lambda_w \frac{\partial r}{\partial S} - d_3 \right) \tag{23}
\]

\[
m = B (\kappa - \kappa^0), \tag{24}
\]

where \( S \) and \( B \) are stiffness matrices which are related to the cross sectional area and the second moment of inertia and are therefore diagonal in the local material coordinates. We also assume volume conservation, varying the local radius of the organ in order to compensate for the elastic stretch \( R \rightarrow R/\sqrt{\lambda_w(s,t)} \).

**Simulating Arabidopsis thaliana roots.** We simulate classic wave experiments on *Arabidopsis thaliana*, placing a rod on a plane tilted at an angle \( \alpha \) with respect to the vertical (see Fig. 1d), allowing it to grow during 1500-3000 growth time-steps, which increases their length by a factor of 15-30. We adopt values for *Arabidopsis thaliana*, taking the radius \( R = 0.1 \text{mm} \) (37, 57), and an initial length \( L_0 = 1 \text{mm} \) (see Table 1). We assume that growth is restricted to a sub-apical growth zone of length \( L_{gz} \), such that \( \hat{\varepsilon}_g(s,t) = \varepsilon_{gz}^0 \) within the growth zone \( L(t) = L_{gz} \leq L(s) \leq L(t) \), and \( \hat{\varepsilon}_g(s,t) = 0 \) elsewhere (7) (see Fig. 1b and e). According to Eq. 15, the maximal growth velocity at the tip is \( v_{\text{tip}} = \varepsilon_{gz}^0 L_{gz} \), so that \( L(t) = L_0 + \varepsilon_{gz}^0 L_{gz} t \). In our simulations we take \( L_{gz} = 1 \text{mm} \) and \( \varepsilon_{gz}^0 = 0.2 \text{h}^{-1} \) following experimental data (37). We assume the roots are incompressible with an effective Poisson’s ratio of 0.5 and Young’s modulus of \( E = 50 \text{MPa} \) (7, 64), yielding the stiffness matrices \( S = \frac{E \pi R^4}{4} \cdot \text{diag}(8/9, 8/9, 1) \) and \( B = \frac{E \pi R^4}{4} \cdot \text{diag}(1/4, 1/4, 1/6) \), expressed in the local material frame (31). Values are summarized in Table 1. We assume the plane is frictionless. In order to emulate the stiff agar surface we assume the plane applies a normal restoring force with a spring constant of 100kg/s². The mature zone adheres to the substrate due to development of root hairs and lignification (7), which we emulate by using a restoring force with a spring constant of 1000kg/s² which fixes in place material that exists the growth zone. Thus only the growth zone is free to change its form, while the mature zone is only free to twist around its axis. We vary the gravitropic sensitivity between \( 0.05 \leq \beta \leq 1.0 \) following experimental observations (37), and vary proprioceptive sensitivity in the range \( 0 \leq \gamma \leq 10 \) (21), though its role in roots hasn’t been established as clearly as in shoots (57).

**Solver validation.** We validate our solver in SM 3 by comparing our simulations to two analytically solvable cases: (i) a clamped organ growing in the direction of an obstacle until it buckles, and (ii) the tropic response to a constant stimulus in the case of apical sensing. In SM 3 we also provide additional information regarding the numerical discretization of growth, and a numerical criterion for mechanical equilibrium. For more details about *Elastica* and the implementation of interactions between the rod and external obstacles see (31, 46).

**Estimation of intrinsic twist profile.** The intrinsic twist can be evaluated by counting the number epidermal cells that cross a line tangent to the root axis, yielding the parameter CFR (cell file rotation) such that \( \text{CFR} = \text{Number/length}. \) The average projected length of one epidermal cell on the root axis is therefore 1/CFR. Assuming that a cell’s width is \( w \approx 0.01 \text{mm} \) (45), we can estimate the angle of the cell file with respect to the root axis by

\[
\sin(\eta) = \text{CFR} \cdot w
\] (25)

This angle in the mature zone can be expressed using \( \eta = \arctan(\tau \cdot R) \), and we estimate \( \tau \) from the CFR following

\[
\tau = \frac{1}{R} \tan(\arcsin(\text{CFR} \cdot w)) = \frac{1}{R} \frac{\text{CFR} \cdot w}{\sqrt{1 - (\text{CFR} \cdot w)^2}}
\] (26)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Value used</th>
<th>Units</th>
</tr>
</thead>
<tbody>
<tr>
<td>( R )</td>
<td>100</td>
<td>( \mu m )</td>
</tr>
<tr>
<td>( L_{gz} )</td>
<td>1</td>
<td>( mm )</td>
</tr>
<tr>
<td>( \varepsilon_{gz}^0 )</td>
<td>0.2</td>
<td>( 1/h )</td>
</tr>
<tr>
<td>( E )</td>
<td>50</td>
<td>( \text{MPa} )</td>
</tr>
<tr>
<td>( \beta )</td>
<td>0.05-1.0</td>
<td></td>
</tr>
<tr>
<td>( \gamma )</td>
<td>0.0-10.0</td>
<td></td>
</tr>
<tr>
<td>( v_{\text{tip}} = \varepsilon_{gz}^0 L_{gz} )</td>
<td>0.2</td>
<td>( \text{mm/h} )</td>
</tr>
<tr>
<td>( \beta L_{gz}/R )</td>
<td>0.5</td>
<td></td>
</tr>
</tbody>
</table>

Table 1. Parameters used to emulate *Arabidopsis thaliana* roots. The values taken have the same order of magnitude as their measured values, as described in (7, 21, 37, 57, 64). The last two relations were used in order to fit experimental data in Figs. 3, 4.

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