The simulation model of growth and cell divisions for the root apex with an apical cell in application to *Azolla pinnata*

Anna Piekarska-Stachowiak · Jerzy Nakielski

Received: 19 March 2013 / Accepted: 16 August 2013 / Published online: 30 August 2013
© The Author(s) 2013. This article is published with open access at Springerlink.com

**Abstract** In contrast to seed plants, the roots of most ferns have a single apical cell which is the ultimate source of all cells in the root. The apical cell has a tetrahedral shape and divides asymmetrically. The root cap derives from the distal division face, while merophytes derived from three proximal division faces contribute to the root proper. The merophytes are produced sequentially forming three sectors along a helix around the root axis. During development, they divide and differentiate in a predictable pattern. Such growth causes cell pattern of the root apex to be remarkably regular and self-perpetuating. The nature of this regularity remains unknown. This paper shows the 2D simulation model for growth of the root apex with the apical cell in application to *Azolla pinnata*. The field of growth rates of the organ, prescribed by the model, is of a tensor type (symplastic growth) and cells divide taking principal growth directions into account. The simulations show how the cell pattern in a longitudinal section of the apex develops in time. The virtual root apex grows realistically and its cell pattern is similar to that observed in anatomical sections. The simulations indicate that the cell pattern regularity results from cell divisions which are oriented with respect to principal growth directions. Such divisions are essential for maintenance of peri-anticlinal arrangement of cell walls and coordinated growth of merophytes during the development. The highly specific division program that takes place in merophytes prior to differentiation seems to be regulated at the cellular level.

**Keywords** Apical cell · *Azolla* · Cell divisions · Principal directions · Root growth · Simulation model · Tensor approach

**Abbreviations**
- AC: Apical cell
- GT: Growth tensor
- PDG: Principal direction of growth
- R_{1}: Relative elemental rate of linear growth

**Introduction**

Roots and shoots of pteridophytes comprise apical cells which are “stem cells” in which they renew themselves and produce a derivative every time they divide (Gunning 1982; Barlow 1997). Each derivative becomes added to the apex as a new construction unit called merophyte, within which set sequences of cell division, expansion and differentiation occur. In roots of *Azolla* (Gunning et al. 1978) and *Equisetum* (Gifford and Kurth 1982), a single apical cell (AC) located at the pole of the root proper occurs. The apical cell has a tetrahedral shape; its three faces are oriented proximally, while the fourth one which is a part of the root/cap border is oriented distally. During ontogenesis, the AC divides always asymmetrically. The derivative cut off from the AC distal face contributes to the root cap, while the derivatives from the three proximal division planes give rise to the proper root. Divisions along proximal faces are sequential and successive merophytes overlap with one another forming a strict merophyte helix arranged into 120° sectors along the root axis. In each new
merophyte, formative divisions of the early longitudinal-tangential and longitudinal-radial types are observed in a definite sequence (Fig. 1). These formative divisions lead to determination of precursors of particular tissues which are defined starting from the outer tissues in the following order: epidermis and outer cortex, inner cortex and endodermis, pericycle and inner stele. Once the files are established, transverse divisions in the files take place and next differentiation occurs. As the AC divisions are predictable and the proliferation of cells in merophytes is highly specific (each cell type has its own program of divisions prior to terminal differentiation), the root apex has a very precise construction and its cell pattern is remarkably regular (Gunning et al. 1978). The nature of such regularity remains unknown.

The apical cell, though usually identified by morphological criteria, plays an important functional role (Bierhorst 1977). On the one hand, it is an ultimate source of all cells of the root apex and can be regarded as a single-cell meristem (Korn 1993). On the other hand, it works as a regulatory site and formative center in histogenesis and organogenesis. Positional signaling seems to play a role for such function (Gunning et al. 1978; Barlow 1991; Hou and Blancasflor 2009). However, how the AC specific division sequence is determined and maintained throughout the root ontogeny is still an open question.

The root apices, like other plant organs, grow symplastically (Priestley 1930; Erickson 1986). During such growth, considered typical for plants, cells are displaced in a coordinated way, preserving their mutual contacts. A description of the symplastic growth is based on the continuity condition (Gandar 1980; Silk 1984). It is assumed that at the organ level, there exists a field of the displacement velocity \( \mathbf{V} \) of points described mathematically by a continuous and differentiable function of position (Gandar and Chalabi 1989; Silk 2006).

The continuity of \( \mathbf{V} \) implies that values of the linear growth rate in different directions are coordinated and strongly depend on a direction (Erickson 1966). The relative elemental rate of growth in length at a given point along the direction \( \mathbf{e}_s \) is defined by the equation (Hejnowicz and Romberger 1984):

\[
R_l(s) = \left[ \nabla \mathbf{V} \cdot \mathbf{e}_s \right] \cdot \mathbf{e}_s
\]

where \( \mathbf{e}_s \) is the unit vector of the direction and dots represent scalar product. As that gradient acting on vector gives the second-rank operator (Spiegel 1959), the field of growth rates of the organ is of a tensor type (Silk and Erickson 1979; Kenaway et al. 2011). If \( \mathbf{V} \) is known, such field can be determined with the aid of the growth tensor (GT, Hejnowicz and Romberger 1984) calculated from \( \nabla \mathbf{V} \) (Supplementary material 1).

Modeling by use of the GT was used to study spatial and directional variation of growth rates in root apices (Hejnowicz 1989; Nakielski 1991; Hejnowicz and Karczewski 1993). It has appeared that in the root apex with the AC, there maximum volumetric growth rate occurs, which may relate to activity of the apical cell working as a single initial. Concerning directional \( R_l \) variation, locally three mutually orthogonal principal directions of growth (PDG) can be recognized unless growth is isotropic, as the directions in which \( R_l \) attains extreme values (maximal, minimal and of a saddle type). These PDGs, determined locally for many points, arrange themselves into three types of continuous lines called PDG trajectories (Hejnowicz 1984, 1989). Through every point, three such trajectories can be drawn. At the organ level, there is a pattern of PDG trajectories considered steady, if organ shape does not change in time. In roots (Nakielski 1987, 2008), similarly to other growing plant organs (Dumais and Kwiatkowska 2002; Kwiatkowska 2004), such pattern can be observed in the cell wall system. Two families of continuous and mutually orthogonal lines commonly used to describe cell walls seen in a longitudinal section, known as periclimes and anticlimes (Sachs 1887), have been postulated to represent PDG trajectories (Hejnowicz 1984). Hejnowicz (1989) hypothesized that cells divide with respect to PDGs, a division wall typically is

![Fig. 1 Schematic representation of formative divisions of A. pinnata root apex during twelve cell cycles durations (after Gunning 1982). In successive merophytes from 1 to 12 the longitudinal-tangential and longitudinal-radial divisions are shown; dashed lines represent new division in the sequence. The cell rows numbered in the merophyte 7 are considered in the text](image-url)
perpendicular to one of three PDG at the site of its forma-
tion. The question arises whether regularity of cell pattern
observed in roots with the apical cell as well as a highly
specific division program that takes place in merophytes
prior to differentiation may result from cell divisions ori-
tented according to this rule.

On the basis of the gt concept, the 2D simulation model
for growth in which cells divide with respect to PDGs was
worked-out (Nakielski 1999; Nakielski and Hejnowicz
2003). An application of the model to the root apex with
the quiescent center, exemplified by radish, gave interesting
results (Nakielski 2008). The virtual root apex grew realis-
tically, indicating that cell divisions with respect to PDGs,
are needed to generate cell pattern which is preserved
during growth.

The present paper shows a similar tensor-based model,
but for the root apex with the apical cell. The model, imple-
mented on the example of the Azolla pinnata, assumes that
growth field of the root apex is of the tensor type and cells
divide with respect to PDGs. Using simulation, one can
see how new merophytes are formed and cell pattern of the
root apex develops in time. The obtained results indicate a
role of principal growth directions in generation and main-
tenance of the cell pattern typical for the root apex with
the apical cell. The divisions with respect to PDGs appear
essential not only for coordinated growth of the individual
cells in merophytes but also to these merophytes in the
organ as a whole. Both the formative and proliferative di-
visions in merophytes are oriented taking PDGs into account
but their highly specific division program needs regulation
at the cellular level.

Materials and methods

Root material

Let us assume cell pattern in the axial section of A. pinnata
root apex adopted from Gunning (1982) as representative
for this species (Fig. 2a). In the cell pattern, all peculiar-
ties of growth organization typical for the root apex with
the single AC and merophytes can be seen. For our mod-
eling, regularity of the cell wall system is the most impor-
tant. Walls of cells located in different parts of the root
apex manifest either periclinal or anticlinal orientation.
Therefore, the cell pattern can be conveniently described
by periclines and anticlines. Two periclines correspond to
the root/cap junction, whereas borders between merophytes
are represented by anticlines. Notice that every two mutu-
ally orthogonal cell walls locally tangent to particular peri-
cline and anticline, though displaced as a result of continu-
of flow of cells from the distal portion of the root apex,
retain their orthogonal intersections in time. This is because
orthogonal intersections between both these clines are
preserved during growth (Sachs 1887; Hejnowicz 1989).

Orthogonal alignment of merophytes with respect to the
root cap border as well as division walls resulting from
longitudinal and transversal divisions within merophytes
is evident. This indicates that peri-anticlinal arrangement
of the cell wall system is especially important for A. pin-
nata root apex. If periclines and anticlines represent PDG
trajectories (Hejnowicz 1984, 1989), regularity of the cell
pattern may result from cell divisions oriented with respect
to PDGs.
The cell walls in the exemplary merophyte are shown in 3D in Fig. 2b. Let us assume that these walls are formed with respect to PDGs in such a way that the merophyte lies in the plane defined by $G_a$ and $G_l$, i.e. it is perpendicular to $G_p$, and two remaining walls resulting from longitudinal-tangential and longitudinal-radial divisions are perpendicular to $G_a$ and $G_l$, respectively. Notice that due to threefold symmetry of the root apex, in cell pattern, coming from the axial section, only two of three sectors can be seen. Moreover, the cell walls formed by the longitudinal-radial divisions (perpendicular to $G_l$) are not represented in it. Accordingly, in our simulation model showing development of cell pattern in the axial plane, only cell divisions perpendicular to $G_p$ (transversal, anticlinal in 2D) and $G_a$ (longitudinal-tangential, periclinal in 2D) will be generated. To simplify the description, division walls perpendicular to $G_p$ will be called anticlinal, whereas those perpendicular to $G_a$ will be periclinal.

In regard to input data for computer simulation, the AC with the youngest merophytes and contacting cells of the root cap was assumed. They were digitized on the basis of Fig. 2a taking the merophyte corresponding to the stage 7 in Fig. 1 as the oldest one. Orientation of cell walls with respect to the root axis was estimated using published sections (Gunning et al. 1978; Gunning 1982) and our own anatomical study of *A. pinnata* root apices. In this study, serial sections of Steedman’s wax-embedded root tips were stained by the periodic acid-Schiff reaction and observed by brightfield and epifluorescence microscopy.

The 2D tensor-based simulation model for growth of *Azolla pinnata* root apex.

To generate growth of *A. pinnata* root apex, the simulation model described previously (Nakielski 2008) was used. Briefly, the model is composed of three elements: the polygon meshwork representing cell pattern at the input, the GT field that generates growth, and the cell division algorithm according to which cells divide with respect to PDGs. The temporal sequences of the simulated growth are obtained by operational application of the GT field to the meshwork. During growth, the meshwork expands, deforms, and new cells are formed through divisions. The division occurs when the cell area assumed as critical is exceeded. Then the parent cell is replaced by two daughter cells separated by a division wall oriented with respect to one of PDGs. In the simulation plane that corresponds to the central longitudinal section there are two PDGs: $G_a$ and $G_p$. Previously, (Nakielski 2008) both these directions were considered, and finally, the PDG for which the division wall could be shorter was chosen (Fig. 3). In the present model, this rule is modified so that a choice of $G_a$ or $G_p$ depends on the cell state (will be described). After formation, the division wall is shortened slightly, with respect to its former length (Nakielski 2008), to have a more realistic three-way junction at each end of the new wall.

The meshwork

Individual cells were described by polygons and arranged into a meshwork (Nakielski 2008) in which two neighboring polygons had a common side, whereas three such polygons had a common vertex (three-way junction). At the start of the simulation, the meshwork consists of 19 cells (Fig. 4). It was an open structure in this sense that new walls, resulting from cell divisions, were added to it. At both ends of the division wall, new three-way junctions were formed. Due to shortening of the division wall, which was different depending on location, angles at each three-way junction were similar in that they were usually occurring in a given region of the root apex.
Fig. 4 The polygon meshwork representing cell pattern in apical part of the Azolla pinnata root apex and the growth tensor field (red) applied to it. In the meshwork, cells are described by polygons. The growth field, defined in the paraboloidal coordinates \((u, v, \varphi)\), is visualized by \(u\) and \(v\) lines which represent PDG trajectories. The field consists of two zones, I and II, separated by the line \(u_0\). Its application to the meshwork is such that assuming \(v_0 = 0.42\) as the root/cap border (dashed) the proximal faces of the apical cell are tangent to \(u_0 = 0.44\) (thick). Under such an application, the zone I generates growth of the apical cell and distal part of the root cap, whereas the zone II—of merophytes and lateral part of the root cap.

**Growth tensor (GT) field**

The displacement velocity field, \(\mathbf{V}\), defined by Hejnowicz (1989) was used. In the paraboloidal coordinate system \((u, v, \varphi)\), the \(\mathbf{V}\) vector is composed of three physical components: \(V_u\), \(V_v\), and \(V_\varphi\). Assuming the absence of rotation around the root axis during growth we have \(V_\varphi = 0\), while the two remaining components are the following:

\[
V_v = h_v \frac{dv}{dt}, \quad V_u = h_u \frac{du}{dt}
\]

where \(h_v = h_u = \sqrt{u^2 + v^2}\) are scale factors of the coordinate system. We deal with the axial section through the apex that corresponds to the plane \(\varphi = \text{const}\). For such a plane, the \(\mathbf{V}\) field is considered two-dimensional. It consists of two zones, I and II, separated by the line \(u_0\) (Fig. 4). These zones are responsible to generate growth in different parts of the root apex: zone I—the apical cell and central part of the root cap, zone II—merophytes with the remaining part of the root cap. At every point, including the line \(u_0\), the \(\mathbf{V}\) is continuous. Two components of the displacement velocity are given by the equations (Hejnowicz 1989):

**Zone I**

\[
\frac{du}{dt} = au \quad \frac{dv}{dt} = kv \left( u_0 - u \right)
\]

**Zone II**

\[
\frac{du}{dt} = au_0 + c \left( u - u_0 \right) \quad \frac{dv}{dt} = -bv \left( u - u_0 \right)
\]

where \(a, b, c, k\) are parameters. Values of these parameters were the following: \(a = 0.012, b = 0.005, c = 0.07, k = 0.035\). They were determined heuristically, by simulation, by taking cell pattern of the real root apex as reference. Let us take as the example the parameter \(c\), influencing a thickness of merophyte at different age. The thickness increases linearly with a value of \(c\). Different values of this parameter were tested and the best approximation of the rate was reached for \(c = 0.07\). Values of the other parameters were specified in a similar way.

By having \(\mathbf{V}\), the tensor field of growth rates of the root apex was able to be determined. In correspondence to \(\mathbf{V}\), the field consisted of two zones; I and II. Figure 4 shows application of the field to the meshwork. The line \(v_0 = 0.42\) corresponded to the root/cap border, and the proximal faces of the apical cell were tangent to \(u_0 = 0.44\). Under such an application, slightly modified in comparison to previous ones (Hejnowicz 1989; Nakielski 1991), PDG trajectories correspond to lines of the coordinate system (see Supplementary material 1), especially in the region of a formation and development of merophytes. It was assumed that a pattern of the coordinate lines represents PDG trajectories in the axial plane of the root apex.

The spatial and directional variation of the linear growth rates \((R_l)\) within the root apex is shown in Fig. 5. The highest \(R_l\) values are in the central region and decrease with increasing distance from the tip in both periclinal and anticlinal directions. In terms of PDGs, everywhere, the highest growth rate is along \(G_a\) (periclinal direction), the rates along two remaining PDGs are significantly smaller. Comparing the rates along \(G_\varphi\) (anticlinal direction) and \(G_l\) (latitudinal direction), it can be seen that they are similar at the root axis and reaching maximal rates in the central region. However, going from this region to the root tip, the rate in \(G_\varphi\) decreases (Fig. 5a), and the rate in \(G_l\) remains relatively high (Fig. 5b). For that reason, at the root peripheries, the rates in \(G_\varphi\) increase, whereas the rates in \(G_l\) decrease slightly with distance from the tip.

**Algorithm for cell divisions**

As mentioned earlier, cells divided with respect to PDGs and orientation of a division wall was determined by cell state. The apical cell could be in two states: 1 and 2. It divided always perpendicular to \(G_\varphi\) however, in state 1, a new merophyte was produced to the sector on the left, whereas, in state 2—to the sector on the right. In every case, the division was unequal and the newly formed wall was tangent to \(u_0\). After division, the cell state was changed from 1 to 2 and vice versa.

The cell initiating merophyte was in state 3, and cells formed within the merophytes were in states 4–7 (Fig. 6a). How cell state affected cell division is shown in Fig. 6b. If the cell was in one of the states from 3 to 6, its division was perpendicular to \(G_\varphi\) (longitudinal). The cells in states 3 and 6 divided into two equal parts, whereas division of cells in states 4 and 5 was unequal. The state 7 resulted in
division perpendicular to \( G_p \) (transversal) and both daughter cells were equal. This state was final in the sense that both daughter cells and their derivatives were in the same state as the parent cell.

Figure 6c shows changes in the state of cells during the merophyte development. The youngest merophyte (state 3) divides perpendicular to \( G_a \) producing cells in states 4 and 5. The cell at state 4, located closer to the root/cap border, divides perpendicular to \( G_a \) producing daughters in state 7. In further development, both daughters divide perpendicular to \( G_p \). The cell at state 5, in turn, divides unequally perpendicular to \( G_a \) giving two cells each in state 6. Their daughter cells reach state 7, which is the final state. Further divisions of these cells are perpendicular to \( G_p \). The root cap cells were once together in state 7 which means that they always divided transversally into two equal cells.

The cells of \( A. pinnata \) root apex differ a lot in their dimensions. For this reason, the area assumed as critical for cell division \( (A_{cr}) \) was not the same for all cells. In general, it was at the level of 150 % of the mean cell area \( (A_{av}) \). In estimation of \( A_{av} \), the apical cell, cells in merophytes and cells in the root cap were considered separately. Also, the shortening of division walls was not uniform at both ends of the new wall. For example, the wall cutting of new merophyte was diminished with respect to its former length about 1.2 % at the outer, and 7 % at the inner end. The shortening of new walls in merophytes depended on their orientation. Namely, the walls resulting from division perpendicular to \( G_a \) were shortened by 15 % and 8 % at the distal and proximal ends, whereas, divisions perpendicular to \( G_p \)—by 8 % and 2 % at inner outer ends, respectively. The cells of the root cap were shortened by 1.2 % at both ends.

Simulations

The simulation started from the meshwork is shown in Fig. 4. After ten steps, each with \( \Delta t = 0.02 \), the first loop \( (L_1) \) was finished, and the meshwork returned to the input. Then the next loop \( (L_2) \) was performed and the procedure was repeated. The whole simulation consisted of 16 loops during which the meshwork could be printed out at every step. The results presented in Figs. 7 and 8 show the meshwork in times corresponding to divisions of the apical cell, labeled with \( T_i \). In particular, the times \( T_{11} \) and \( T_{13} \) represent the meshwork after first and last division of the AC which occurred in \( L_1 \) and \( L_{13} \), respectively.
The GT field was steady by definition. At $T_0$, its application to the meshwork is as in Fig. 4, i.e. the root/cap border coincided with the line $v_0 = 0.42$. However, in the course of the simulation, the border was slightly displaced proximally with respect to this line (by a distance not greater than 1% of the apical cell height). It returned to the previous position at the start of each new loop.

Statistical analysis

Statistical analysis was performed to estimate differences in orientation of the newly formed walls between real and virtual root apices. Cell walls coming from nearly central sections of six roots (both sectors) were taken into account; four coming from our own studies and two published by Gunning (1982). For every root apex, the angle between the wall and the root axis was estimated by division into three classes of cell walls: A—cutting off new merophyte, B, C—resulting from longitudinal-tangential (B) and transversal (C) divisions of cells in merophyte. For each class, at first, differences in angles between particular roots were analyzed using the ANOVA test. Then the samples coming from real roots, taken together for a given class, were compared with corresponding data.
obtained for a single virtual root in course of the whole simulation. In this case, the t test for independent samples was applied.

Also, the width ratio of two neighboring merophytes in the same sector (older to younger, i.e. $M_{i+1}/M_i$ in Fig. 6a) was estimated in both sectors of real and virtual roots. The width was measured in the central part of a merophyte. Assuming that the corresponding merophytes $M_i$ of real and virtual roots make pairs, the t test for dependent samples was used. All tests were performed using STATISTICA, version 10, at the significance level 0.05.

**Results**

The computer sequence of growth and cell divisions in the root apex of *A. pinnata* are shown in Figs. 7, 8 and Supplementary material 2 (video file). The cell pattern assumed at the input expands from $T_0$ to $T_7$ in Fig. 7 and from $T_8$ to $T_{13}$ in Fig. 8. In the course of the whole simulation, the apical cell divided 13 times and the total number of cells in the meshwork increased from 19 to more than 170. At every time-step, the GT field operated in such a way that the apical cell remained under control of the zone I, whereas
merophytes developed in the zone II. Cell divisions were perpendicular to $G_a$ (longitudinal-tangent division, periclinal in 2D) and perpendicular to $G_b$ (transverse division, anticlinal in 2D).

New merophytes are cut off perpendicular to $G_p$, alternately to the left and right side of the root apex (Figs. 7, 8). The first new merophyte is formed at $T_1$ on the right side (light gray in Fig. 7). Let us consider development of this merophyte during the whole simulation. At $T_2$, the cell giving rise to the merophyte increases in area and at $T_3$, divides longitudinally (perpendicular to $G_a$). Two daughter cells are obtained; the inner lying closer to the root axis and the outer that keeps contact with the root/cap border. From $T_3$ to $T_4$, both cells expand and then divide longitudinally as the parent cell at $T_5$ and $T_6$. Later, at $T_8$ and $T_9$ (Fig. 8), the next longitudinal division occurs in two inner cells. At the same time, starting from $T_{10}$, the outer cells divide transversely (through division perpendicular to $G_p$) and such division is reached by the innermost cells of the merophytes at $T_{13}$. In reference to Gunning’s (1982) results, it is possible to indicate precursors of particular tissues. Using notation of the cell rows I–IV adopted from Fig. 1 (merophyte 7), they have been established as follows (Fig. 8): at $T_8$—for the epidermis and root hairs (row IV) and outer cortex (row III), at $T_{10}$—for the inner cortex and endodermis (row II), and at $T_{12}$—for the pericycle, xylem, sieve elements, phloem parenchyma (row I).

Notice that at the end of the simulation ($T_{13}$ in Fig. 8), the considered merophyte (light gray) appears to be similar to the younger merophyte marked as dark gray at $T_9$ in Fig. 7. Hence, it is possible to predict a sequence of divisions of cells of the dark gray merophyte. Namely, during a period from $T_9$ to $T_{13}$ (Fig. 8) cells of this merophyte will expand and new transverse divisions will occur in the outermost cell. Further divisions of cells of this merophyte will be of a proliferative type.

Both analyzed merophytes are situated on the left side of the root axis, but it is easy to notice that merophytes on the opposite side developed in a similar way. Now let us focus our attention on the root cap. There are a few cell divisions in this region occurring only in the inner cell layer. As assumed, all these divisions are perpendicular to $G_p$. Notice the cell at $T_0$ located in zone 1 (gray in Fig. 7). This cell expands periclinally and divides anticlinally at $T_5$ giving two daughter cells. Since the division wall happens to be near the border between the zones 1 and 2, these daughter cells find their way to different zones and their fates are different. The one controlled by zone 2 expands without division and is displaced proximally into the root periphery whereas the other one remaining in zone 1 grows and divides as its parent cell at $T_9$.

Cell patterns of both real and virtual root apices are compared in Fig. 9. It can be seen that the computer-generated cell pattern is generally similar to that observed in the real root apex, especially concerning longitudinal dimension of successive merophytes as well as orientation of formative and proliferative divisions in the merophytes. Small discrepancies between the real and modeled roots relate to number cells in merophytes and not perfectly anticlinal shape of the merophytes generated in the model.

The statistical analysis of orientation of cell walls with respect to the root axis gave the following results. In all considered classes (A, B, C), the ANOVA test provided the absence of significant differences in orientation of corresponding walls between considered real roots; for each class, the obtained $P$ values are much greater than 0.05 (A 0.99, B 0.88, and C 0.63). The $t$ test has shown the absence of statistically significant differences in orientation of cell walls between corresponding samples in real and virtual roots (Table 1). The ratio taken from width of neighboring merophytes decreased with the merophyte age in both real and virtual roots (Table 2). The $t$ test has shown the absence of statistically significant differences in ratio between corresponding samples of real and virtual roots (Table 3). This means that virtual and real roots are also statistically similar, at least taking considered characteristics into account.

In our model, fates of individual cells are determined by their position in GT field. Accordingly, PDG at a given position affect both cell extension and cell division. What happens when other rules for cell divisions (Besson and
Table 1 Orientation of newly formed walls in real and virtual root apices

| Merophytes | Real roots | Virtual root | t test |
|------------|------------|--------------|--------|
|            | $n$        | $x_{av} \pm SD$ | $n$        | $x_{av} \pm SD$ | $t$-value | $P$    |
| A          | 36         | 58.06 ± 5.09  | 12        | 60.17 ± 2.08  | 1.39     | 0.17   |
| B          | 36         | 16.31 ± 6.07  | 57        | 14.11 ± 6.30  | 1.66     | 0.10   |
| C          | 36         | 81.36 ± 7.99  | 48        | 78.71 ± 7.26  | 1.59     | 0.12   |

The walls cutting off new merophyte. The walls resulting from formative (B) and proliferative (C) divisions in merophytes. The angle between the new wall and the root axis was measured in degrees.

$n$ number of walls, $x_{av}$ mean value, SD standard deviation

Table 2 The ratio of width of two neighboring merophytes obtained for virtual (second column) and real (third column) $Azolla$ root apices

| Merophytes | Virtual root | Real root |
|------------|--------------|-----------|
| 2:1        | 1.27         | 1.28      |
| 3:2        | 1.29         | 1.30      |
| 4:3        | 1.29         | 1.28      |
| 5:4        | 1.21         | 1.23      |
| 6:5        | 1.22         | 1.20      |
| 7:6        | 1.16         | 1.18      |
| 8:7        | 1.14         | 1.16      |
| 9:8        | 1.14         | 1.12      |

The first column indicates from which merophytes the ratio was taken. The third column shows mean values.

Table 3 Comparison of the ratio width of two neighboring merophytes in the real and virtual $Azolla$ root apices

| Merophytes | Real root | Virtual root | t test |
|------------|-----------|--------------|--------|
|            | $n$        | $x_{av} \pm SD$ | $n$        | $x_{av} \pm SD$ | $t$-value | $P$    |
| Ratio      | 31         | 1.25 ± 0.12  | 31        | 1.21 ± 0.07  | 1.81     | 0.07   |

The samples were dependent and every ratio of the virtual apex was paired with corresponding ratio in the real apex.

$n$ number of walls, $x_{av}$ mean value, SD standard deviation

Discussion

This paper presents a 2D model of growth and cell division in the root apex with a single apical cell. The model shows development of cell pattern in the central longitudinal section of the root apex, assuming that growth field of the organ is of a tensor type (Hejnowicz and Romberger 1984) and cells divide with respect to principal directions of growth (PDGs). How PDGs affect cellular pattern of the root apex with the quiescent center was described previously (Nakielski and Hejnowicz 2003; Nakielski 2008). The present paper is the first in which a tensor field of growth rates has been used to generate growth together with cell divisions in the root apex with the apical cell and merophytes.

The root apex of $Azolla pinnata$ root is predisposed to such modeling. It has relatively simple but precise construction (Gunning et al. 1978; Gunning 1981) and successive divisions of the AC and its derivatives are the most completely documented among ferns. Moreover, cell pattern of the root apex derives from activity of one cell working as a single initial. A remarkable regularity of the cell pattern may suggest some special properties of the AC (Gifford 1983; Barlow 1994). However, our simulations indicate an important role of principal growth directions; the cell pattern is regular and becomes steady due to cell divisions oriented with respect to PDGs. New merophytes are formed perpendicularly to the $G_p$, whereas division walls in merophytes are formed perpendicularly to $G_e$ and $G_t$ (longitudinal-tangential and transversal divisions, respectively). Adopting growth rate anisotropy is shown in Fig. 5, the transversal divisions are perpendicular, whereas longitudinal ones are tangent to the direction of maximal $R_t$. The realistic development of cell pattern of the virtual root apex may suggest that the apical cell and its derivatives are able to detect PDGs at a site of their location and orient division walls with respect to them.

The PDGs are an integral part of a growth field (Hejnowicz and Romberger 1984). In our model they are considered locally and at the organ level through the PDG field. Locally they affect cell expansion and cell divisions (orientation of division wall), whereas at the organ level, the stability of the root shape and coordinated development of merophytes within the apex (cell fates depend on PDGs via position in GT field) are affected. As the PDG pattern determines the orientation of volume increase and is predetermined by the existing structure of the growing body (Hejnowicz and Hejnowicz 1991), there is a feedback...
between growth field and the cell wall system. When the volume increase is accompanied with proliferation, cell divisions are involved in mechanism governing growth and formation of division wall with respect to PDgs, which is natural. In the broader meaning, PDgs are needed for coor-
dinated growth of individual cells and the organ as a whole.

Concerning orientation of the division wall, several hypotheses have been proposed (reviewed by Smith 2001; Müller 2012; Minc and Piel 2012). It was postulated that cells divide: perpendicular to the main axis of growth (Hofmeister 1863), nearly perpendicular to the existing walls (Sachs 1878, 1887) and along the shortest path, dividing the cell into two equally sized daughters (Errera 1888). These geometrical hypotheses emphasize local properties of dividing cells, not taking organ level and directional signals into account. Besides, they are useful mostly for isotropic growth (Sahlin and Jönsson 2010) and geometrically uniform cells (Besson and Dumais 2011; Alim et al. 2012). Asymmetric divisions have been recently used to model stomata lineages in leaf epidermis through a polarity-switching mechanism (Robinson et al. 2011). Also mechanical stress has been suggested to affect ori-
entation of division plane (Green 1980; Lintilhac and Vesecky 1984), receiving support in empirical observations (Lynch and Lintilhac 1997; Hamant et al. 2008; Potocka et al. 2011). The Hejnowicz’s hypothesis, according to which cells divide with respect to PDGs, combines geometri-
cal and mechanical approaches. On the one hand, it can be seen as generalization of the cell geometry based rules (Hejnowicz and Hejnowicz 1991). On the other hand, its relation to mechanical stress is clear because from the point of view of mechanics, growth is an irreversible deformation that results from stresses in the cell wall system (Nakiel-
ski and Hejnowicz 2003). Both the growth rate and the mechanical stress are tensor quantities and one can expect a relationship between their principal directions. If mecha-
nical cues are sensed and interpreted by ‘highly dynamic and regulated microtubule cytoskeleton’ (Mirabet et al. 2011) they may give rise to PDG oriented cell divisions via the stress. The mentioned relationship needs investigations, however, empirical data suggest (Alim et al. 2012) that cell division is parallel to the direction of maximal stress. By analogy of previous modeling (Nakielksi 2008), such situation may take place in the case of proliferative divisions in merophytes.

The rule according to which cells divide with respect to PDGs is primary for the present model. The majority of remaining rules are used mainly to ‘translate’ 3D cellular architecture and cell division sequence which occurred in the real root apex (under threefold symmetry about central longitudinal axis), to our 2D model in which merophytes are produced only to two sectors and to the left and right. In addition, to be as close as possible to empirical data, these remaining rules are numerous and deal with details. This creates the impression of “hard-wiring” a correct division pattern by which the primary role of PDgs is obscure. In this context, it is worth noticing that if divisions of the api-
cal cell are considered in 3D, the rule about the ac state becomes unnecessary because all merophytes are formed exactly in the same way as perpendicular to $G_p$ at the site of its formation. The helical formation of successive merophytes may be a consequence of geometrical factors result-
ing from the division parallel to proximal face giving rise to the first merophyte of the new root. Whether they are produced clockwise or counterclockwise may depend on random factors as that both are similarly frequent either in root (Guning 1982; Korn 1993) or shoot (Imaichi 1988) apices.

The computer-generated cell pattern was realistic but also some discrepancies between real and virtual roots,

\begin{figure}[h]
\centering
\includegraphics[width=\textwidth]{fig10.png}
\caption{The computer-generated cell pattern in two merophytes obtained by use of different division rules. \textbf{a} Cell divides with respect to PDGs with the algorithm presented in this paper. \textbf{b} As in (a) but not taking cell states into account. \textbf{c} Cell divides perpendicular to the nearest wall (Sach’s hypothesis). \textbf{d} Cell divides by the shortest path (Errera’s hypothesis). The merophytes at the bottom and top correspond to the oldest ones at $T_2$ and $T_6$ in Fig. 7, respectively. In \textbf{b-d}, the division walls with orientation different than in (a) are indicated (red).}
\end{figure}
related to number of cells and not perfectly anticlinal alignment of merophytes were observed (Fig. 9). Such discrepancies seem to be inevitable when a highly specific cell division program designated for three sectors has been applied to two sectors and under limitation to the longitudinal section. In addition, our attention was focused rather on the order of formative and proliferative divisions in merophytes, not cell pattern in particular sectors of the real apex. Also the simplification according to which cell walls are represented by straight lines is important; in the case of the AC, it is responsible for not exactly anticlinal alignment of merophytes. On the other hand, even working with the microscopic section regarded as the best in the literature, it is worth noting that the section is not perfectly central (Gunning 1982) and two sectors, due to threefold symmetry of the root apex, are not equally represented in it.

The *Azolla pinnata* root growth is determinate (Gunning et al. 1978). The apical cell divides in sum 50–55 times but after approximately 35 divisions its mitotic activity is successively reduced and finally ceases when the root reaches the length of about 8–10 mm (Gunning et al. 1978). A similar reduction is also observed in roots of other species (Chiang and Gifford 1971; Nitayangkura et al. 1980; Gifford 1983). In our simulation, the AC divided 6–7 times per sector which corresponds to about 2/5 of the total number of its divisions. Such time period was sufficient to analyze the cell pattern maintenance, but too short to assume changes in a cell division rate. Such changes can be taken into account in future work focused on a relationship between the rate of growth and the rate of cell divisions during whole period of mitotic activity of the AC.

In the present modeling the mature root apex is considered. The question arises how such cell pattern is formed. The problem has been already studied by use of the growth tensor (Hejnowicz and Hejnowicz 1991). The GT field, similar to the one considered here but of an unstable type, was applied to the non-differentiated orthogonal grid of points. Under this field, the grid extended, deformed and a clear protrusion was formed. Adjusting by hand the apical cell, it was possible to observe deformation of this cell and provide for its divisions with respect to PDGs. Three routes of development, differing in orientation of the first division, were considered. They lead optionally to the root apex with a single and two apical cells (two routes). Adopting these results to the *A. pinnata* root apex, one can conclude that the first division of the apical cell (parallel to the distal face) is perpendicular to \( G_a \), whereas all next divisions parallel to each of three proximal faces, are perpendicular to \( G_p \).

Roots with a single apical cell and roots with a quiescent center differ in growth organization and in cellular architecture but are similar in PDG pattern (Hejnowicz and Karczewski 1993). This suggests that the tensor-based control related to PDGs is needed regardless of cellular organization of the root apex (quiescent center or the apical cell). It is worth mentioning that directional information included in PDGs is essential in the present paper to describe cell pattern maintenance during steady growth, and is also of importance for unsteady growth which takes place, for example, during the lateral root formation (Szymanowska-Pułka and Nakielski 2010).

The rule governing the cell state change is similar to ‘production rules’ commonly used in L-systems (Prusinkiewicz 1998, 2004). In the 2D modeling of growth of *Microsorum linguiforme* gametophyte (de Boer and de Does 1990) cell divisions oriented periclinally and anticlinally were generated. However, such orientation did not result from PDGs and the tensor basis of the symplastic growth but only from the gametophyte geometry; periclinal division was tangent, whereas anticlinal-perpendicular to the organ surface. This cannot be surprising, because the plane tangent to the surface is defined by two PDGs: \( G_a \) and \( G_p \). Hence, periclinal and anticlinal divisions seen in a longitudinal section through the organ are in 3D perpendicular to \( G_a \) and \( G_p \) respectively, i.e. exactly as in the model presented here.

The L-systems appeared useful first of all to analyze genealogy of cells (Lück et al. 1988, 1994a, b; Barlow 1991; Lück and Lück 1993). In this aspect both shoot and root apices possessing one or more apical cells were investigated (Barlow et al. 2000, 2001). Such analysis does not need simulation performed at the organ level. Therefore, the present paper is the first in which development of cell pattern in the section of the whole root apex with the apical cell has been shown. It is also the first in which growth and divisions of individual cell are controlled on the one hand at the organ level, by growth field with PDGs, and on the other hand locally by factors influenced the cell state, type of cell division (perpendicular to \( G_a \) or \( G_p \), equal or not) and shortening of a division wall. Moreover, the present approach, though restricted to the case of a single apical cell, can be relatively easily adapted to describe growth in roots or shoots of *Azolla filiculoides* (Gifford and Polito 1981; Nitayangkura et al. 1980), *Equisetum* (Gifford and Kurth 1982; Gifford 1993) and *Ceratopteris richardii* (Hou and Hill 2002, 2004). The case with more than one apical cell requires deeper modifications.

**Acknowledgments** We thank Jacek Pietrakowski for his help in preparing the video supporting material, as well as Dorota Kwaitkowska and Joanna Szymanowska-Pułka for discussion and critical reading of the manuscript.

**Open Access** This article is distributed under the terms of the Creative Commons Attribution License which permits any use, distribution, and reproduction in any medium, provided the original author(s) and the source are credited.
References

Alim K, Hamant O, Boudaoud A (2012) Regulatory role of cell division rules on tissue growth heterogeneity. Front Plant Sci 3:174. doi:10.3389/fpls.2012.00174

Barlow PW (1991) From cell wall networks to algorithms: the simulation and cytokinetics of cell division patterns in plants. Protoplasma 162:69–85

Barlow PW (1994) Structure and function at the root apex—phylogenetic and ontogenetic perspectives on apical cells and quiescent centres. Plant Soil 167:1–16

Barlow PW (1997) Stem cells and founder zones in plants, particularly their roots. In: Potten CS (ed) Stem cells. Academic Press, London, pp 29–57

Barlow PW, Lück HB, Lück J (2000) Organogenetic cellular patterning in plants. In: Carbone A, Gromov M, Prusinkiewicz P (eds) Pattern formation in biology, vision and dynamics. World Scientific Publishing, Singapore, pp 163–198

Barlow PW, Lück HB, Lück J (2001) Autoreproductive cells and plant meristem construction: the case of the tomato cap meristem. Protoplasma 215:50–63

Besson S, Dumais J (2011) Universal rule for the symmetric division of plant cells. Proc Natl Acad Sci USA 108:6294–6299

Bierhorst DW (1977) Owen the stem apex, leaf initiation and early leaf ontogeny in filicalean ferns. Amer J Bot 64:125–152

Chiang S-H, Gifford EM (1971) Development of the root of Ceratopteris thalictroides with special reference to apical segmentation. J Indian Bot Soc 50A:96–106

De Boer MJM, De Does M (1990) The relationship between cell division pattern and global shape of young fern gametophytes. I. A model study. Bot Gaz 151:423–434

Dumais J, Kwiatkowska D (2002) Analysis of surface growth in the shoot apices. Plant J 31:229–241

Erickson RO (1966) Relative elemental growth rates and anisotropy of growth in area: a computer program. J Exp Bot 17:390–403

Erickson RO (1986) Symplastic growth and symplasmic transport. Plant Physiol 82(4):1153

Erella L (1988) Über Zellformen und Seifenblasen. Bot Centralblatt 34:395–399

Gandar PW (1980) The analysis of growth and cell production in root apices. Bot Gaz 141:131–138

Gandar PW, Chalabi S (1989) Estimation of velocity fields from cell number densities in non-steady, one dimensional growth. Bot Gaz 150:93–100

Gifford EM (1983) Concept of apical cells in bryophytes and pteridophytes. Annu Rev Plant Physiol 34:419–440

Gifford EM (1993) The root apical meristem of Equisetum diffusum: structure and development. Am J Bot 80:468–473

Gifford EM, Kurth E (1982) Quantitative studies of the root apical meristem of Equisetum scirpoides. Am J Bot 69:464–473

Gifford EM, Polito VS (1981) Growth of Azolla filiculoides. Bio-Sciences 31:526–528

Green PB (1980) Organogenesis—a biophysical view. Annu Rev Plant Physiol 31:51–82

Gunning BES (1981) Microtubules and cytomorphogenesis in developing organ: the root primordium of Azolla pinnata. In: Kiermayer O (ed) Cell biology monographs, vol 8. Springer, New York, pp 301–325

Gunning BES (1982) The root of the water fern Azolla: Cellular basis of development and multiple roles for cortical microtubules. In: Shtehly A, Green PB (eds) Developmental order: its origin and regulation. Alan R. Liss, New York, pp 379–421

Gunning BES, Hughes JE, Hardham AR (1978) Formative and proliferative cell divisions, cell differentiation, and developmental changes in the meristem of Azolla roots. Planta 143:121–144

Hamant O, Heisler MG, Jönsson H, Krupinski P, Uyttewaal M, Bokov P, Corson F, Sahlén P, Boudaoud A, Meyerowitz EM, Couder Y, Traas J (2008) Developmental patterning by mechanical signal in Arabidopsis. Science 322:1650–1655

Hejnowicz Z (1984) Trajectories of principal directions of growth, natural coordinate system in growing plant organ. Acta Soc Bot Pol 53:29–42

Hejnowicz Z (1989) Differential growth resulting in the specification of different types of cellular architecture in root meristems. Environ Exp Bot 29:85–93

Hejnowicz Z, Hejnowicz K (1991) Modeling the formation of root apices. Planta 184:1–7

Hejnowicz Z, Karczewski J (1993) Modeling of meristematic growth of root apices in a natural coordinate system. Am J Bot 80:309–315

Hejnowicz Z, Romberger JA (1984) Growth tensor of plant organs. J Theor Biol 110:93–114

Hofmeister W (1863) Vorsätze und Berichtigungen zu den 1851 veröffentlichten Untersuchungen der Entwicklung höherer Kryptogamen. Jahrb Wiss Bot 3:259–293

Hou G, Blanchaer EB (2009) Fern root development. In: Beeckman T (ed) Root development. Annu Plant Rev, vol 37, pp 192–208

Hou GC, Hill JP (2002) Heteroblastic root development in Ceratopteris richardi (Parkeriaceae). Int J Plant Sci 163:341–351

Hou GC, Hill JP (2004) Developmental anatomy of the fifth shoot-born root in young sporophytes of Ceratopteris richardi. Planta 219:212–220

Imaichi R (1988) Developmental anatomy of the shoot apex of leptosporangiate ferns. II. Leaf ontology of Adiantum capillus-veneris (Adiantaceae). Can J Bot 66:1729–1733

Kennaway R, Coen E, Green A, Bangham A (2011) Generation of diverse biological forms through combinatorial interactions between tissue polarity and growth. PLoS Comput Biol 7:e1002071

Korn RW (1993) Apical cells as meristems. Acta Biotheor 41:175–189

Kwiatkowska D (2004) Structural integration at the shoot apical meristem: model, measurements, and experiments. Am J Bot 91:1277–1293

Lintlilac PM, Vesecky TB (1984) Stress-induced alignment of division plane in plant-tissues grown in vitro. Nature 307:363–364

Lück J, Lück HB (1993) Parallel rewriting -cellwork L-systems for plant development. In: Demongeot J, Capasso V (eds) Mathematics applied to biology and medicine. Wuerzburg Publishing LTD, pp 461–466

Lück J, Lindemayer A, Lück HB (1988) Model for cell tetrads and clones in meristematic cell layers. Bot Gaz 149:127–141

Lück J, Barlow P, Lück HB (1994a) Deterministic patterns of cellular growth and division within a meristem. Ann Bot 73:1–11

Lück J, Barlow PW, Lück HB (1994b) Cell genealogies in a plant meristem deduced with the aid of a ’bootstrap’ L-system. Cell Prolif 27:1–21

Lynch TM, Lintlilac PM (1997) Mechanical signals in plant development: a new method for single cell studies. Dev Biol 181:246–256

Minc N, Piel M (2012) Predicting division plane position and orientation. Trends Cell Biol 22:193–200

Mirabet V, Das P, Bouadaoud A, Hamant O (2011) The role of mechanical forces in plant morphogenesis. Annu Rev Plant Biol 62:365–385

Müller S (2012) Universal rules for division plane selection in plants. Protoplasma 249:239–253

Nakielski J (1987) Variation of growth in shoot apical domes of spruce seedling: a study using the growth tensor. Acta Soc Bot Pol 56:625–643
Nakielski J (1991) Distribution of linear growth rates in different directions in root apical meristems. Acta Soc Bot Pol 60:77–86
Nakielski J (1999) Tensorial model for growth and cell division in the shoot apex. In: Carbone A, Gromov M, Prusinkiewicz P (eds) Pattern formation in biology, vision and dynamics. World Scientific, Singapore, pp 252–286
Nakielski J (2008) The tensor-based model for growth and cell divisions of the root apex. I. The significance of principal directions. Planta 228:179–189
Nakielski J, Hejnowicz Z (2003) The description of growth of plant organs: a continuous approach based on the growth tensor. In: Nation J, Trofimova I, Rand JD, Sulis W (eds) Formal description of developing systems. Kluwer Academic Publishers, Dordrecht, NATO science series II, vol 121, pp 119–136
Nitayangkura S, Gifford EM, Rost TL (1980) Mitotic activity in the root apical meristem of Azolla filiculoides Lam., with special reference to the apical cell. Am J Bot 67:1484–1492
Potocka I, Szymanowska-Pułka J, Karczewski J, Nakielski J (2011) Effect of mechanical stress on Zea root apex. I. Mechanical stress leads to the switch from closed to open meristem organisation. J Exp Bot 62:4583–4593
Priestley JH (1930) Studies in the physiology of cambial activity. II. The concept of sliding growth. New Phytol 29:96–140
Prusinkiewicz P (1998) Modeling of spatial structure and development of plants: a review. Sci Hortic 74:113–149
Prusinkiewicz P (2004) Modeling plant growth and development. Curr Opin Plant Biol 7:79–83
Robinson S, Barbier de Reuille P, Chan J, Bergmann D, Prusinkiewicz P, Coen E (2011) Generation of spatial patterns through cell polarity switching. Science 333:1436–1440
Sachs J (1878) Über die Anordnung der Zellen in jüngsten Pflanzentheilen. Arb Bot Inst Würzburg 2:46–104
Sachs J (1887) Lecture XXVII. Relations between growth and cell division in the embryonic tissues. In: Lectures in plant physiology. Clarendon Press, Oxford, pp 431–459
Sahlin P, Jönsson H (2010) A modelling study on how cell division affects properties of epithelial tissues under isometric growth. PLoS One 5:e11750
Silk WK (1984) Quantitative descriptions of development. Annu Rev Plant Physiol 35:479–518
Silk WK (2006) Moving with the flow: what transport laws reveal about cell division and expansion. J Plant Res 119:23–29
Silk WK, Erickson RO (1979) Kinematics of plant growth. J Theor Biol 76:481–501
Smith LG (2001) Plant cell division: building walls in the right places. Nat Rev Mol Cell Biol 2:33–39
Spiegel MR (1959) Theory and problems of vector analysis and an introduction to tensor analysis. McGraw-Hill, New York
Szymanowska-Pułka J, Nakielski J (2010) The tensor-based model for growth and cell divisions of the root apex. II. Lateral root formation. Planta 232:1207–1218