Cytoplasmic pH Dynamics in Maize Pulvinal Cells Induced by Gravity Vector Changes1[w]

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In maize (Zea mays) and other grasses, changes in orientation of stems are perceived by pulvinal tissue, which responds to the stimulus by differential growth resulting in upward bending of the stem. The amyloplast-containing bundle sheath cells are the sites of gravity perception, although the initial steps of gravity perception and transmission remain unclear. In columella cells of Arabidopsis roots, we previously found that cytoplasmic pH (pHc) is a mediator in early gravitropic signaling (A.C. Scott, N.S. Allen [1999] Plant Physiol 121: 1291–1298). The question arises whether pHc has a more general role in signaling gravity vector changes. Using confocal ratiometric imaging and the fluorescent pH indicator carboxy seminaphthorhodfluor acetoxymethyl ester acetate, we measured pHc in the cells composing the maize pulvinus. When stem slices were gravistimulated and imaged on a horizontally mounted confocal microscope, pHc changes were only apparent within the bundle sheath cells, and not in the parenchyma cells. After turning, cytoplasmic acidification was observed at the sides of the cells, whereas the cytoplasm at the base of the cells where plastids slowly accumulated became more basic. These changes were most apparent in cells exhibiting net amyloplast sedimentation. Parenchyma cells and isolated bundle sheath cells did not show any gravity-induced pHc changes although all cell types responded to external stimuli in the predicted way: Propionic acid and auxin treatments induced acidification, whereas raising the external pH caused alkalinization. The results suggest that pHc has an important role in the early signaling pathways of maize stem gravitropism.

The vector of the gravitational force is one of the main cues that determines the spatial orientation of plant organs (Masson, 1995; Tasaka et al., 1999; Kiss, 2000). A plant’s ability to respond to the direction of gravity, the process of gravitropism, ensures the correct positioning of the seedling after germination and also enables mature plants to correct their position after a forced reorientation, e.g. by strong wind. In most cases, gravity perception occurs in specialized cells that contain dense particles, such as starch-filled plastids (amyloplasts; Sack, 1997; Tasaka et al., 1999). These shift their location within the cell when the normal plant orientation gets disturbed, thereby generating a cellular signal that sets up a chemical gradient between top and bottom of the plant organ resulting in differential growth.

Whereas the gravity-induced growth response in plants is well documented, little is known about the mechanism of gravity perception and the nature of the early steps of the signaling cascade. Research has focused on gravity perception in roots that occurs in amyloplast-containing root cap cells. Some of the earliest measurable responses induced by gravistimulation occur in and around these cells, and include complex changes in cytosolic and apoplastic pH (Scott and Allen, 1999; Fasano et al., 2001), changes in plasma membrane potential (Sievers et al., 1995), and the induction of ion flux changes around the root cap (Behrens et al., 1985; Björkman and Leopold, 1987). It is intriguing that in Arabidopsis, cytoplasmic pH (pHc) changes occur only in the inner columella cells of the root cap (Scott and Allen, 1999; Fasano et al., 2001), which are the most competent in gravity perception (Blancaflor et al., 1998). Furthermore, artificial modification of pHc alters the gravitropic response (Scott and Allen, 1999; Fasano et al., 2001). This suggests that pHc changes play a key role in gravity-induced signaling.

Induced changes in pHc occur in response to a wide array of stimuli, apart from gravitropism, including phytohormones (Felle, 1988a; Gehring et al., 1990a, 1994; Beffagna et al., 1994), light (Felle and Berl, 1986; Okazaki et al., 1994), Nod factors (Allen et al., 1994; Felle et al., 1996), and other elicitors (Mathieu et al., 1996). Furthermore, protons are implicated as a mediator in plant signal transduction (Felle, 1989; Guern et al., 1992; Roos et al., 1998; Zhou et al., 2000). Whether the gravity-induced pHc changes measured in roots have a general role in gravity perception in plants is still unknown because pHc changes in shoots have only been linked to the later phases of the gravitropic response (Gehring et al., 1990b). The objective of this study was to elucidate whether gravity-induced changes in pHc occur

1 This work was supported by the National Aeronautics and Space Administration (NASA Specialized Center of Research and Training grant no. NAGW–4984).
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in shoot tissues that are specialized in gravity perception such as the maize (*Zea mays*) pulvinus. The stem pulvinus of maize is a disc-shaped tissue that is located above each stem node and has a specific role in both gravity perception and the bending response (Collings et al., 1998; for review, see Kaufman et al., 1987, 1995). When the stem is placed horizontally, amyloplast sedimentation occurs in bundle sheath cells, and a growth response follows characterized by cell elongation specifically within the pulvinal cells, causing the stem to bend upwards. Pulvinal cells retain the capacity to elongate in the presence of an appropriate stimulus even after the surrounding tissue has fully differentiated, allowing the normal growth responses to be spatially and temporally separated from those induced by changes in the gravity vector (Collings et al., 1998). These properties make the maize pulvinus an ideal system to investigate biochemical (Winter et al., 1997; Perera et al., 1999), structural, and physiological changes at the cellular level (Collings et al., 1998) during gravitropism.

In this study, we focused on early gravity-induced responses and monitored pHc changes in pulvinal cell regions after rotation on a horizontally mounted confocal microscope using the ratiometric pH indicator carboxyseminaphthodifluor acetoxymethyl ester acetate (SNARF-1 AM). We compared the responses of amyloplast-containing bundle sheath cells with those of parenchyma cells to find out whether pHc changes similar to those found in Arabidopsis root cells occur and whether they are associated with the cells that perceive the gravity stimulus. Furthermore, we dissected the pHc responses of base and side regions (relative to the gravity vector) of the stimulated cells and compared cells that exhibited amyloplast sedimentation with those that did not. That pHc changes were most pronounced in cells in which there was net amyloplast sedimentation, with these changes confined to specific sites within these cells, is discussed in light of current models for the mechanism of gravity perception and the rising importance of pHc as a messenger in cellular signaling.

**RESULTS**

**Specimen Preparation and Dye Loading**

The pulvinus is a region of cells found between the differentiated node and elongated cells of the internode. The unelongated pulvinal cells surrounding the vascular strands are of two types. Several layers of bundle sheath cells occur immediately adjacent to the vascular tissue. Potassium iodide staining confirms that these cells contain starch-filled amyloplasts, and that these plastids are effectively confined to the pulvinus (Fig. 1, A–C). Surrounding the bundle sheath are numerous files of ground parenchyma.

To measure cytosolic pH (pHc) in maize pulvinal cells, both before and after gravistimulation, we investigated several specimen preparation protocols. Two methods proved satisfactory for imaging on a sideways-mounted microscope. In the first method, mild enzymatic digestion of cell wall material resulted in the isolation of long files of healthy cells that showed vigorous cytoplasmic streaming. These files could either be parenchymal cells or amyloplast-
containing bundle sheath cells. When files of bundle sheath cells were incubated in a vertical position for 30 min and then gravistimulated, net amyloplast sedimentation occurred over the next several minutes (Fig. 2, A–C; for video sequence, see www.plantphysiol.org). In these cells, the average rate of plastid sedimentation was visibly slower than the streaming velocities for individual amyloplasts. However, amyloplast sedimentation in files of bundle sheath cells, as shown in Figure 2, A through C, was a relatively rare event, seen in only two out of 15 independent rotations. The second specimen preparation method involved taking 0.3- to 0.5-mm thick longitudinal sections through the pulvinus. Cells in these preparation also showed vigorous cytoplasmic streaming, but it is significant that sections were more likely to show amyloplast sedimentation following gravistimulation (Fig. 2, D–F; for video sequence, see www.plantphysiol.org), with this being seen in 67% of sections (n = 12).

To measure pHi, both before and after gravistimulation, we also investigated numerous ratiometric pH indicators. The most promising dye was carboxy seminaphthorhodafluor (SNARF-1) because of a range of technical issues, and because when excited at 514 nm, pulval cells showed little autofluorescence at the emission peaks of SNARF’s acidic and basic forms, recorded at 550 to 600 and 620 to 670 nm, respectively. After 1 h of incubation in the membrane permeant AM ester form of SNARF (10 μM), dye loaded predominantly into the cytoplasm and was largely excluded from the vacuole. However, SNARF fluorescence was also observed within amyloplasts. SNARF loaded rapidly into isolated file cells, whereas in tissue sections the dye loaded first into the vascular tissue then proceeded into bundle sheath cells and finally was found in the parenchyma. Attempts to load SNARF via the vascular bundles into excised pulvini without prior sectioning failed.

**pHi Measurements in Maize Pulvinal Cells: Effects of Weak Acids, External pH Changes, and Indole Acetic Acid (IAA)**

To determine the reliability of our confocal ratiometric imaging system, we measured pHi with SNARF-1 in non-gravistimulated file cells and tissue slices, observing both bundle sheath and parenchyma cells, and determined whether these cells responded to known stimuli in the normal way. Ratios of the two emission intensities (620–670 nm/550–600 nm) at 620–670 nm were measured.

![Figure 2. Dynamics of amyloplast sedimentation. A through C, Amyloplast sedimentation in isolated files of bundle sheath cells. Amyloplasts follow the tracks of intracellular particle movement as well as the path predicted by physical parameters such as plastid density and cytosolic viscosity leading to net sedimentation of plastids to the new bottom of the cell. A, Thirty seconds before rotation of cell files through 180°. QuickTime movie located at www.plantphysiol.org. B, Thirty seconds after rotation. C, Seven minutes after rotation. D through E, Plastid sedimentation in bundle sheath cells of maize. Ratio images E1 (620–670 nm)/E2 (550–600 nm) of SNARF-1 AM-loaded longitudinal pulval sections, excitation 514 nm. D, Before rotation; E, 2 min after rotation by 90°. QuickTime movie located at www.plantphysiol.org. F, 12 Minutes after rotation, bar = 50 μm; v, vascular tissue; b, bundle sheath cells; p, parenchyma cells.](image-url)
nm) were generally stable in non-stimulated cells for the duration of experiments. In vitro calibrations with dextran-linked SNARF (10 kD) were carried out after each experiment. In most cases, these in vitro calibrations gave calculated pH\textsubscript{c} values of about pH 6.6 (or lower) that were more acidic than the pH\textsubscript{c} values reported in earlier studies (Smith and Raven, 1979; Kurkdjian and Guern, 1989; Guern et al., 1991, 1992). However, the cells appeared healthy, showing normal intracellular particle movement, and could react to weak acid and other treatments with similar ratio shifts as those cells whose calibrations reported a resting pH\textsubscript{c} around 7. Thus, results are presented in this paper as changes in ratio rather than changes in pH\textsubscript{c}, with an increase in ratio reflecting alkalinization and a decrease in ratio reflecting acidification. Adjustments in the photomultiplier settings for the two emission windows led mainly to a parallel ratio shift in the calibration curve and only had a small effect on the slope of the pH dependence (Fig. 3). Therefore, the change in ratio obtained from data sets with different starting ratios can be compared even though the pH dependence is not linear. It should, however, be noted that the bulk of the data were obtained within a more narrow range of photomultiplier adjustments than those depicted in Figure 3.

In vivo calibrations with nigericin in the presence of high K\textsuperscript{+} (Vercesi et al., 1994) are not reported in this paper because such experiments did not lead to sustained ratio changes and required complex solution changing that could not be performed for the majority of measurements that were made with a sideways-mounted microscope.

To test whether SNARF-1 AM can be used reliably to monitor pH\textsubscript{c} changes in maize pulvinal cells, a range of conditions were applied that are known to increase or decrease pH\textsubscript{c} in a large variety of plant cells. When subjected to 0.5 mM propionic acid applied at an external pH of 5.5, bundle sheath cells from tissue sections responded with a drop in ratio by $\Delta r = -0.171 \pm 0.009$ ($n = 16$; acidification, Fig. 4A). Streaming remained visibly unaffected in these cells. After removal of the weak acid, the ratio returned to the resting value, and occasionally showed a transient overshoot to more positive values (alkalinization). The kinetics with which these ratio changes occurred are comparable to the weak acid-induced pH\textsubscript{c} changes measured with ion-selective microelectrodes (Felle, 1987; Frachisse et al., 1988) and $^{31}$P NMR spectroscopy (Guern et al., 1986) in a variety of other plant cells. Conditions that evoke alkalinizations were also tested. It is known from previous reports that changes in external pH lead to corresponding changes in pH\textsubscript{c} by about 0.1 pH\textsubscript{c} unit per pH unit change in external pH (for review, see Smith and Raven, 1979; Felle, 1988b). Figure 4B shows the response of pulvinal bundle sheath cells to a change in external pH from 5.5 to 9.0 that caused a rise in the ratio (alkalinization) by $\Delta r = 0.077 \pm 0.008$ ($n = 18$).

The phytohormone auxin (e.g., IAA), which plays an important role in the gravity-induced differential growth response (Evans, 1991; Estelle, 1996), also evoked ratio changes in maize pulvinal cells (Fig. 4C). Application of 0.1 mM IAA evoked a transient rise in the ratio (alkalinization) by $\Delta r = 0.027 \pm 0.004$ ($n = 12$), which was apparent in 72% of analyzed recordings. This was followed by a drop in $\Delta r$ (acidification) by $-0.175 \pm 0.015$ ($n = 18$). IAA concentrations of 10 $\mu$M IAA caused a rise in $\Delta r$ (alkalinization) by 0.040 $\pm$ 0.002 ($n = 3$, one data set), which returned to the resting level within about 10 min (data not shown). Although the acidification evoked by higher IAA concentrations might in part be attributable to a weak acid effect, the initial alkalinization could reflect a change in membrane transport activity. Previous studies concerning pH\textsubscript{c} changes in maize coleoptiles induced by auxins (1 $\mu$M) showed either an oscillatory pattern of pH\textsubscript{c} changes (Felle, 1988a) or a sustained acidification by 0.1 to 0.2 pH units (Gehring et al., 1990a).

Although the data shown in Figure 4 relate to inducible pH\textsubscript{c} changes in bundle sheath cells from tissue slices, we observed similar changes in the emission ratio in isolated files of bundle sheath cells (Fig. 5A), and in parenchyma cells in both slices and isolated files (data not shown).

**Figure 3.** In vitro calibrations with 50 $\mu$M SNARF-1 dextran (10 kD): pH dependence of emission intensity ratio $E_1$ (620–670 nm)/$E_0$ (550–600 nm) for different photomultiplier settings. An increase in the photomultiplier setting for $E_0$ leads to an approximately parallel shift in the ratio values with only a slight increase in slope.

**Figure 4.** Photomultiplier (pmt) $E_2/E_1$ versus pH dependence of different photomultiplier settings. An increase in the photomultiplier setting for $E_1$ leads to an approximately parallel shift in the ratio values with only a slight increase in slope.

**Gravistimulation Experiments Demonstrate pH\textsubscript{c} Changes in Bundle Sheath Cells**

Protocols for isolation of bundle sheath cell files were developed because these cells provide a much more favorable system for high resolution imaging than pulvinal slices. Although it was possible to measure pH\textsubscript{c} changes in this system in response to pro-
pionic acid (Fig. 5A), gravistimulation never led to any visible change in the ratio (Fig. 5B). Because plastid sedimentation was only rarely retained in isolated cell files, it might be argued that the lengthy isolation and dye loading procedure, the potential for cytoskeletal reorganization, or the loss of cell wall material and cell positioning information, led to disruption in cell function that might have affected early steps in the signaling cascade.

In longitudinal stem sections that were loaded with SNARF-1 AM, rotation by 90° resulted in significant ratio changes, indicative of pH_{c} changes, in cells that contained sedimentable amyloplasts, as depicted in Figure 6, B and C, and summarized in Table I. Parenchyma cells that lack amyloplasts provided an ideal control for these experiments because they did not exhibit any notable change in the emission ratio after turning (Fig. 6A) either at the base of the cell or at the side of the cell (Table I). In bundle sheath cells that showed amyloplast sedimentation, cytoplasmic regions at the sides of the cells often responded with a drop in ratio (acidification), whereas regions at the base of cells showed a rise in ratio (alkalinization; Figs. 6B and 7). When cytoplasmic areas of whole cells were measured, the acidification often prevailed. In about 12% of cases, a transient alkalinization was followed by an acidification at the sides of cells but not at the base (Fig. 6C). Rotation by 360° (no gravistimulation) did not elicit any significant changes in the pH_{c} of bundle sheath cells, suggesting that touch responses that might occur during rotation do not contribute to the ratio change (Fig. 6D).

In contrast to these observations of changes in pH_{c} in bundle sheath cells that showed amyloplast sedimentation, bundle sheath cells that failed to show amyloplast sedimentation within 30 min after rotation lacked pronounced changes in pH_{c}. The results are summarized in the histograms presented in Figure 7 and the statistics in Table I. In bundle sheath cells not exhibiting plastid sedimentation, a higher percentage of cells showed no pH_{c} response compared with cells that showed sedimentation; furthermore, alkalinization did not occur at the base of cells. The magnitude and rate of acidification measured in side regions was also markedly lower than that mea-

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**Figure 4.** Ratio changes E_{2} (620–670 nm)/E_{1} (550–600 nm) in maize pulvinal cells from tissue slices following: A, application of 0.5 mM propionic acid; B, a change in external pH from pH 5.5 to 9.0; and C, addition of 0.1 mM IAA. Representative measurements are shown and statistics are given in the text. Substances were applied in buffer C (pH 5.5). Increase in ratio reflects alkalinization, decrease in ratio reflects acidification.

**Figure 5.** Ratio changes E_{2} (620–670 nm)/E_{1} (550–600 nm) in isolated pulvinal cell files following: A, application of 1 mM propionic acid; and B, after rotation by 90°. Representative measurements are shown. Substances were applied in buffer B (pH 5.5). The average ratio change was: A, -0.214 ± 0.019 (n = 10) in response to propionic acid; and B, 0.003 ± 0.005 (n = 10) after rotation.
Figure 6. Ratio changes \( E_2 \) (620–670 nm)/\( E_1 \) (550–600 nm) in maize pulvinal cells from tissue slices before and after rotation by 90° (A–C) and 360° (D). A, Response of parenchyma cells (no amyloplasts); B, typical response of bundle sheath cells with amyloplast sedimentation base region of cells (▲) and side regions of cells (▼); C, transient response measured in side regions of bundle sheath cells showing amyloplast sedimentation, observed in 12% of recordings; D, response of bundle sheath cells to a full 360° rotation. Average values ± s.e. of representative data sets are shown and statistics are given in Table I.

sured in cells with sedimenting plastids (at the 90% confidence interval; Fig. 7B, Table I).

In Table I, the rate of ratio change after rotation was compared statistically (Student’s \( t \) test for two independent samples) for data sets obtained from side and base regions of parenchyma cells, and bundle sheath cells with and without plastid sedimentation. Bundle sheath cells showed a significantly different response than parenchyma cells (control), except for base regions of bundle sheath cells not exhibiting plastid sedimentation, which responded in a manner similar to the control. When cells with and without plastid sedimentation were compared the responses measured in the base regions were significantly different (95% confidence interval) between the two data sets, whereas those measured at the sides of the cells were different at the 90% confidence interval. Bundle sheath cells that were rotated 360° did not exhibit significant ratio changes (Fig. 6D) and their response was significantly different from bundle sheath cells that were rotated 90°.

DISCUSSION

\( \text{pH}_\text{r} \) Measurements in Pulvinal Cells with SNARF-1 AM

Lipophilic derivatives of ion sensitive fluorescent dyes such as SNARF-1 AM can permeate intracellular organelles and cause errors in the determination of cytoplasmic ion concentrations (Fricker et al., 1993; Williams et al., 1993; Roos, 2000). In pulvinal bundle sheath cells, dye accumulation into the central vacuole is minimal during the first 60 to 90 min following SNARF-1 AM incubation. However, a large amount of dye accumulates in amyloplasts, an observation previously noted in barley (Hordeum vulgare) pulvinal cells where amyloplasts accumulate fluorescence following esterase cleavage of fluorescein diacetate (Dayanandan et al., 1982). The result of this dye accumulation is frequent signal saturation from amyloplasts under the conditions used for our experiments. To minimize errors in the \( \text{pH}_\text{r} \) measurements arising from SNARF-1 AM-loaded amyloplasts, we used the following precautions. If possible, cytoplasmic regions free of amyloplasts were selected for measurement, and a thresholding method was used to determine the emission ratio eliminating most of the amyloplast-derived signal.

Quantification of Changes in the Emission Ratio

Because in vitro calibrations did not report reliable \( \text{pH}_r \) values, it was difficult to quantify the observed ratio changes. However, taking into account measurements that reported normal \( \text{pH}_r \) values when compared with in vitro calibration, and by comparing our results obtained with weak acids and external \( \text{pH} \) changes with those obtained by other investigators, we can estimate the range within which the
Table I. pH ratio changes in stem slices following gravistimulation

<table>
<thead>
<tr>
<th>Cell Type</th>
<th>Region</th>
<th>Cells</th>
<th>Average Rate Δt ± st</th>
<th>Significance Levels for Comparisons between Different Changes in Ratioa</th>
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<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Compared with parenchyma</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Base Side</td>
</tr>
<tr>
<td>Parenchyma</td>
<td>Base</td>
<td>25</td>
<td>-0.0004 ± 0.0003</td>
<td>$t = -0.845$</td>
</tr>
<tr>
<td></td>
<td>Side</td>
<td>27</td>
<td>-0.0007 ± 0.0003</td>
<td>$t = -0.845$</td>
</tr>
<tr>
<td>Bundle sheath</td>
<td>Base</td>
<td>42</td>
<td>0.0020 ± 0.0002</td>
<td>$t = 2.3691$</td>
</tr>
<tr>
<td>Bundle sheath, no amyloplast sedimentation</td>
<td>Side</td>
<td>95</td>
<td>-0.0049 ± 0.0004</td>
<td>nd</td>
</tr>
<tr>
<td>Bundle sheath turned 360°</td>
<td>Base</td>
<td>21</td>
<td>-0.0005 ± 0.0005</td>
<td>$t = -0.2289$</td>
</tr>
<tr>
<td></td>
<td>Side</td>
<td>31</td>
<td>0.0005 ± 0.0004</td>
<td>$t = 2.4002$</td>
</tr>
</tbody>
</table>

* Student’s t tests for two independent populations were performed to compare the responses. Significant values at $P < 0.05$ are shown in bold.

** Emission ratio changes in response to rotation by 90° (or 360° as indicated) measured in cytoplasmic regions at different locations within cells, presented as the rate of change in the ratio per minute. Negative values indicate acidification; positive values indicate alkalinization.

Gravity-Induced pHe Changes: An Early Step in Gravity Perception?

The present investigation shows that gravity induces pHe changes in the amyloplast-containing bundle sheath cells of the maize pulvinus, indicating that pHe changes might have a generally important role in early gravity-induced signaling. Average ratio changes evoked in bundle sheath cells (with plastid sedimentation) within 30 min following rotation were about $Δt = 0.06$ (alkalinization) in bottom regions and $Δt = -0.15$ (acidification) in side regions (see Table I and Fig. 6B). Taking into account the considerations discussed in the previous section, the
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many of the earliest measurable gravity-induced effects involve ionic fluxes in and around the root cap. These include membrane potential and ion flux measurements that showed gravity-induced hyperpolarizations and proton fluxes indicative of stimulation of the plasma membrane H⁺ ATPase (Behrens et al., 1985; Björkman and Leopold, 1987; Sievers et al., 1995). Fasano et al. (2001) also observed a uniform acidification of the Arabidopsis root cap apoplast from pH 5.5 to 4.5 within 2 min of gravistimulation. However, because the plasma membrane H⁺ pump can effect pHc changes as well as respond to them, it is difficult to tell which of these two early responses might come first.

Early Events in Gravisignaling and the Role of pHc

Three investigations have now shown that pHc changes are a key player in early gravity-induced signaling in root caps. First, Scott and Allen (1999) showed that microinjection of the pH-sensitive dye 1',7'-bis-(2-carboxyethyl)-5-(and)-6-carboxyfluorescein into gravity-perceiving root cap cells of Arabidopsis revealed a rapid alkalinization in tier 2 cells and a slower acidification in tier 3 cells (Scott and Allen, 1999), the sites with the greatest competence for gravity perception (Blancaflor et al., 1998). When pHc changes were induced solely within the root cap by various pHc modifiers, the gravitropic response was also altered, with acidification of the columella causing enhanced bending, whereas alkalinization resulted in inhibition of bending. These observations were expanded by Fasano et al. (2001), who demonstrated, using both dye 1',7'-bis-(2-carboxyethyl)-5-(and)-6-carboxyfluorescein-dextran microinjection and the constitutive expression of a pH-sensitive green fluorescent protein, that rapid alkalinization occurred in the Arabidopsis columella following gravistimulation. Furthermore, in a stanchless Arabidopsis mutant, where plastid sedimentation and bending were reduced, pHc changes were also markedly smaller (Fasano et al., 2001). Taken together with our observations in the pulvinus, these results indicate that pHc changes might have a generally important role in early gravity-induced signaling.

Concurrent with rapid pHc changes in the pulvinus, gravity also induces rapid changes in the inositol phosphate metabolism in pulvini of maize and oats (Avena sativa) beginning within minutes of gravistimulation and resulting in oscillations of phosphatidylinositol 4,5-bisphosphate and inositol 1,4,5-trisphosphate (IP₃; Perera et al., 1999, 2001). Increases in the second messenger IP₃ are known to open Ca²⁺ channels in the vacuolar membrane of plant cells (Alexandre et al., 1990; Allen et al., 1995), and suggest a role of cytosolic free Ca²⁺ ([Ca²⁺]ᵣ) in the gravity-induced signaling cascade. Although direct measurements of [Ca²⁺]ᵣ in gravity-perceiving cells in the

Figure 7. Histograms summarizing data represented in Figure 6B. The rate of ratio change after rotation was determined over a 15- to 20-min period using linear regression and least-square fit. Data obtained from bundle sheath cells showing amyloplast sedimentation (light shaded bars) were compared with those in which no amyloplast sedimentation occurred (dark shaded bars). A, Ratio changes in base regions; B, ratio changes in side regions. The maximum rates of ratio changes observed in data sets obtained from parenchyma cells (exemplified in Fig. 6A) fell within the range of Δr = ±0.0025 min⁻¹ and were designated as no response. The bin width was Δr = 0.005 min⁻¹; however, rates of acidification larger than Δr = −0.0125 min⁻¹ were included in the left column. Statistics are given in Table 1.

gravity-induced ratio changes are likely to reflect pHc changes in the range of 0.3 to 0.5 pH units. These changes are similar to stimulus-induced pHc changes reported in other plant systems where changes in pHc have a wide array of physiological effects in plant cells and can act as mediators in many signal transduction pathways (Felle, 1989; Guern et al., 1992; Putnam, 1998). Changes in pHc affect proton translocation by pumps and carriers (Felle and Johannes, 1991; Fortillo, 2000), and can modulate the activity of various anion (Schulz-Lessdorf et al., 1996; Johannes et al., 1998) and cation (Grabov and Blatt, 1997; Lacombe et al., 2000) channels.

The sequence of signaling events that follow gravistimulation, and the involvement of pHc, remain largely unknown. However, in root gravitropism,
root cap of Arabidopsis failed to reveal such changes in response to gravistimulation (Legué et al., 1997), a role for [Ca^{2+}], has long been implicated by indirect experimental evidence (for review, see Chen et al., 1999). It is conceivable that [Ca^{2+}] changes occur and act locally in cellular microdomains and that high resolution imaging techniques in conjunction with more sensitive Ca^{2+} dyes are required to make them detectable.

However, several significant questions remain unclear. First, how does amylloplast sedimentation cause changes in pH, phosphatidylinositol 4,5-bisphosphate/IP_3 levels, and possibly also [Ca^{2+}], and how are these changes linked? Second, how is this signal integrated from the cellular to the tissue level, such that pH changes visible in the majority of cells with amylloplast sedimentation can generate a chemical gradient between the top and bottom of the plant organ that results in differential growth and the bending response? And third, how does auxin, thought to be a key regulator of the bending response (Evans, 1991; Estelle, 1996), fit into this system? One possible integrating system that could satisfy all these questions would be the actin cytoskeleton.

Although subject to some debate (for review, see Baluška and Hasenstein, 1997), it has now been shown that the amyloplasts in the columella cells of the root cap from various plant species are surrounded by endoplasmic reticulum and actin filaments (Collings et al., 2001). In a similar manner, the dynamics of amyloplast streaming in the maize pulvinus (this study), and amyloplast motility in other gravitropic tissues (Sack and Leopold, 1985), indicate extensive interactions between amyloplasts and the actin cytoskeleton. Thus, it is possible that while moving to the new cell bottom, amyloplasts might interact with actin and/or membranes, thereby eliciting pH changes in cellular microdomains. The actin cytoskeleton has several features that make it suitable for regulating gravitropic signaling and response generation. Actin-binding proteins that regulate the equilibrium between F actin and G actin, such as actin-depolymerizing factor, are modulated by pH with depolymerization occurring at higher pH values (Gungabissoon et al., 1998; Kovar and Staiger, 2000). Various actin-binding proteins, including actin-depolymerizing factor and profilin (Gibbon and Staiger, 2000; Kovar and Staiger, 2000), also interact extensively with signaling by phospholipids. Furthermore, actin modulates auxin transport through its regulation of the auxin efflux carrier (Munday, 2000).

The current state of knowledge suggests that changes in membrane transport (Behrens et al., 1985; Björkman and Leopold, 1987; Sievers and Busch, 1992; Sievers et al., 1995), actin dynamics (Collings et al., 2001), and phosphoinositol metabolism (Perera et al., 1999, 2001) are early responses to a change in the gravity vector. It is likely that the observed pH changes have an integral role in the gravity-induced signaling cascade because pH alone or in conjunction with [Ca^{2+}], modulates the activity of membrane transport proteins (Felle and Johannes, 1991; Schulz-Lessdorf et al., 1996; Grabov and Blatt, 1997; Johannes et al., 1998; Lacombe et al., 2000; Fortillo, 2000), cytoskeletal polymerization (Gungabissoon et al., 1998; Kovar and Staiger, 2000), organization of the endoplasmic reticulum (Quader and Fast, 1990), and enzymatic reactions (Putnam, 1998; Paterson and Nimmo, 2000), all of which are likely to play a role in the initial steps of gravity-induced signal transduction. It will be a challenge for future investigators to elucidate how the gravity signal is integrated and to dissect the sequence of events leading from gravity perception to the gravitropic response. Progress in this field will be aided by the availability of mutants impaired in perception or transduction of the gravity signal (Sedbrook et al., 1999; Firm et al., 2000) and through the use of more refined techniques for visualization of ionic changes in cellular microdomains using pH-sensitive green fluorescent protein constructs that can be targeted to specific cellular compartments (Miesenböck et al., 1998).

MATERIALS AND METHODS

Plant Material

Maize (Zea mays) plants (line 3183; Pioneer, Des Moines, IA) were grown in greenhouses (22°C–27°C), three plants per 22-cm pot. Pulvinus 2 and 3 (the second and third pulvinus from the base of the stem) from 5- to 6-week-old plants were used for all experiments. These pulvinus exhibit the strongest response to gravitistimulation, and lie adjacent to internodes that have ceased elongation (Collings et al., 1998).

Isolation of Stem Slices

Longitudinal hand sections, approximately 0.2 to 0.5 mm thick, were made with a razor blade through the pulvinal regions of pulvinus 2 and 3. Replicate sections were stained to reveal and confirm the location of starch-containing amyloplasts with 0.2% (w/v) iodine in 5% (w/v) KI (10 min).

Isolation of Cell Files

Pulvinus 2 and 3 from one or several maize plants were harvested, chopped in 1- to 2-mm³ pieces, and incubated for 40 to 50 min in cell wall-digesting enzymes (1% [w/v] Cellulase YC, 0.1% [w/v] Pectolyase Y23, 0.5% [w/v] bovine serum albumin, and 0.3 mM mannitol in Murashige and Skoog medium, pH 5.3). Following enzyme digestion, the tissue was mechanically disrupted by gentle stirring and tapping at the tissue pieces with a glass rod to release files of cells. Buffer A [100 mM KCl, 0.285 mM mannitol, and 10 mM PIPES (piperazine-N,N'-bis(2-ethanesulfonic acid))/KOH, pH 6.8] was added and the suspension was allowed
to sediment for 30 to 45 min at 1g (4°C), and this fast-sedimenting debris removed. Files of cells were recovered by further sedimentation of the suspension at 1g (4°C) over the next several hours. The pelleted cell files then were resuspended in buffer A to remove residual enzyme and again pelleted at 1g (4°C) for several hours. Centrifugation to accelerate the recovery of cell files was not feasible because low-speed centrifugation (<80g) ruptured the amyloplast-containing bundle sheath cells.

Ester Loading of SNARF-1 AM and Mounting of Cells

Cell files were incubated for 60 min in 10 μM SNARF-1 AM (0.8 mL of buffer A and 1% [v/v] dimethyl sulfoxide, 22°C). Cell files settled at the bottom of the Eppendorf tube and were resuspended in the recording buffer B [5 mM KCl, 0.475 mM mannitol, 0.1 mM CaCl2, and 10 mM MES-[N-(morpholino) ethanesulfonic acid]/KOH, pH 5.5, standard recording medium for cell files] to remove external dye. After sedimentation at room temperature, cell files were embedded in a thin layer of low-melting-point agarose (1.3% [w/v] agarose VII, in buffer B, applied at about 40°C) on a prewarmed welled slide.

Freshly prepared longitudinal sections through the pulvinral region were rinsed and incubated for 60 min in 10 μM SNARF-1-AM (1% [v/v] dimethyl sulfoxide, 22°C). Following dye loading they were washed three times with buffer C (0.1 mM CaSO4, 0.2 mM K2SO4, 0.1 mM NaN3, 0.5 mM MgCl2, and 10 mM MES/Tris, pH 5.5, standard recording medium for tissue sections) to remove excess dye. The slices were then embedded in a thin agarose layer (1.3% [w/v] agarose VII, in buffer C, applied at about 40°C) on a prewarmed welled slide.

For gravity experiments, welled slides were sealed with a coverslip using melted valap (vaseline:lanolin:paraffin, 1:1:1, w/w), whereas for perfusion experiments, the slide well (approximate volume 0.3 mL) was left open on two sides and had a small reservoir on each side. This allowed rapid media exchange using filter paper.

Unless otherwise stated, chemicals were obtained from Sigma Chemical Co. (St. Louis) and fluorescent dyes were obtained from Molecular Probes Inc. (Eugene, OR).

Confocal Imaging and Ratiometric pH Measurements

Experiments were conducted with a confocal microscope system (Leica SP, Leica). The ratiometric pH indicator SNARF-1 AM was excited with an argon laser at 514 nm, and fluorescence emission windows were set to 550 to 600 nm (peak of acidic form) and 620 to 670 nm (peak of basic form). These wavelengths avoided autofluorescence from the amyloplasts present within the bundle sheath cells. Concurrent differential interference contrast images were also recorded. Images were acquired with a 20× numerical aperture 0.6 dry objective at 5 to 10-s intervals for up to 40 min. Ratiometric image analysis was performed with Metafluor 4.01 software (Universal Imaging) with cytoplasmic regions marked and thresholds set at 3 and 253 (8-bit image). Ratios are given as emission intensity at 620 to 670 nm divided by emission intensity at 550 to 600 nm. In vitro calibrations were carried out with 50 μM SNARF-1 dextran (10 kD) in 100 mM KCl, 1 mM ATP, 1 mM MgCl2, and 10 mM HEPES [4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid]/Tris adjusted to pH 6.0, 6.5, 7.0, 7.5, and 8.0 (Bibikova et al., 1998).

For gravity experiments, the confocal unit was attached to an “upright” microscope (Leica model DMRX-A) that was mounted sideways in a cradle to give a horizontal light path. Specimens were mounted on a rotating stage and kept in an upright position for at least 30 min before the start of an experiment. Specimens were gravistimulated by rotating the stage, and imaged with centerable lenses adjusted to give a constant field of view as the stage rotated. Images were recorded before and after gravistimulation, but images taken during and directly after rotation were often blurred and could not be used for ratio analysis, resulting in gaps in the recording of between 25 and 180 s.

Perfusion experiments were performed in the horizontal position on an inverted microscope (Leica model DM-IRB) to facilitate solution changes.

Data Analysis

Emission ratios were routinely measured at the sides and base of the turned cells. Throughout the manuscript, the terms “side” and “base” of cells refers to their position after rotation. Average ratio values were compiled from three to nine independent data sets (if not indicated otherwise) and are presented as average values ± se (n = no. of cells or regions measured). The rate of change in the ratio after gravistimulation was determined using linear regression and a least-square fit. Data sets obtained for different cell regions/types were compared statistically using Student’s t test for independent samples.

Received February 16, 2001; returned for revision April 23, 2001; accepted June 3, 2001.

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Gravity-Induced Cytoplasmic pH Dynamics in Maize Pulvinus


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