Light Promotion of Hypocotyl Gravitropism of a Starch-Deficient Tobacco Mutant Correlates with Plastid Enlargement and Sedimentation

Stanislav Vitha, Ming Yang², John Z. Kiss³, and Fred D. Sack*

The Ohio State University, Department of Plant Biology, 1735 Neil Avenue, Columbus, Ohio 43210

Dark-grown hypocotyls of a starch-deficient mutant (NS458) of tobacco (Nicotiana sylvestris) lack amyloplasts and plastid sedimentation, and have severely reduced gravitropism. However, gravitropism improved dramatically when NS458 seedlings were grown in the light. To determine the extent of this improvement and whether mutant hypocotyls contain sedimented amyloplasts, gravitropic sensitivity (induction time and intermittent stimulation) and plastid size and position in the endodermis were measured in seedlings grown for 8 d in the light. Light-grown NS458 hypocotyls were gravitropic but were less sensitive than the wild type (WT). Starch occupied 10% of the volume of NS458 plastids grown in both the light and the dark, whereas WT plastids were essentially filled with starch in both treatments. Light increased plastid size twice as much in the mutant as in the WT. Plastids in light-grown NS458 were sedimented, presumably because of their larger size and greater total starch content. The induction by light of plastid sedimentation in NS458 provides new evidence for the role of plastid mass and sedimentation in stem gravitropic sensing. Because the mutant is not as sensitive as the WT, NS458 plastids may not have sufficient mass to provide full gravitropic sensitivity.

The availability of starch-deficient mutants has provided new opportunities to test the starch-statolith hypothesis, i.e. the idea that gravity sensing relies on the mass of amyloplasts that sediment. In WT plants amyloplast sedimentation occurs in highly specific tissues such as the central rootcap (columella) and in the starch sheath (endodermis) in stems (Sack, 1987, 1991). In mutants with little or no starch, plastid sedimentation appears to be absent from these locations (Caspar and Pickard, 1989; Kiss and Sack, 1990). Roots and hypocotyls of starchless or starch-deficient mutants of Arabidopsis thaliana are gravitropic, but sensing is reduced, as measured by an increased threshold of stimulation and by a greater variability in organ orientation in response to prolonged stimulation (Caspar and Pickard, 1989; Kiss et al., 1989, 1996). Light-grown roots of a starch-deficient mutant of tobacco (Nicotiana sylvestris), NS458, are also gravitropic and have reduced sensitivity (Kiss and Sack, 1989).

The findings that five different mutants at four loci have no or decreased starch and reduced gravitropism support the conclusions that starch plays a role in sensing when present, and that at least moderate levels of starch are necessary for full gravitropic sensitivity (Poff et al., 1994; Kiss et al., 1996; Sack, 1997). These hypotheses are also supported by many other correlative data, including older experiments with maize plastid mutants and with the depletion of starch by experimental manipulation (Hertel et al., 1969; Sack, 1991, 1997). Data from NS458 hypocotyls provide striking support for the starch-statolith hypothesis, since the endodermis contains small, unsedimented plastids and these hypocotyls are severely deficient in gravitropism when grown in the dark (Kiss and Sack, 1990). However, preliminary experiments indicated that light-grown NS458 hypocotyls were strongly gravitropic without obvious increases in starch content, data that raised the possibility that light restored gravitropism independent of plastid mass or position. To analyze further the effects of light, we determined whether light-grown NS458 hypocotyls are as sensitive to gravity as WT hypocotyls, and whether light affects the amount of starch or plastid position compared with dark-grown mutant hypocotyls. This analysis shows that light induces plastid sedimentation in the NS458 endodermis, probably through an increase in plastid volume and in total starch content, and that light-grown mutant hypocotyls exhibit significant gravitropism but are less sensitive than the WT.

MATERIALS AND METHODS

Plant Material and Cultivation

Seeds used were WT tobacco (Nicotiana sylvestris Spec. et Comes) and the starch-deficient mutant NS458 (fourth generation with reselection for the phenotype after the first backcross). NS458 is deficient in the activity of plastidic phosphoglucomutase (Hanson and McHale, 1988).

Abbreviation: WT, wild type.
Seedlings were grown in square polystyrene Petri dishes (100 × 15 mm) on 1% (w/v) agar containing nutrients supplemented with 1% (w/v) Suc, as described by Kiss and Sack (1990). The dishes were sealed with Parafilm (American National Can, Greenwich, CT) and placed on the edge so that the surface of the agar was vertical, and were placed under continuous illumination (60–80 μmol m⁻² s⁻¹ from 40-W cool-white fluorescent lamps, General Electric). After 8 d of cultivation, the hypocotyls were about 1 to 1.5 mm long. For assessment of plastid sedimentation, dark-grown seedlings were also used and cultivated as described by Kiss and Sack (1990).

**Measurement of Gravitropic Curvature and Growth**

To measure curvature and growth, light-grown seedlings were photographed and then transferred to the dark and either turned to the horizontal (for measurement of curvature) or kept upright (for measurement of growth). Dark-grown seedlings were intermittently photographed using Kodak T-Max 400 ASA film and illumination with dim-green light (intensity of approximately 0.9 μmol m⁻² s⁻¹ at the level of the hypocotyl) provided by an incandescent lamp filtered through two layers of a Roscolux filter (no. 1090, Rosco Laboratories, Port Chester, NY) with a peak transmission of 526 nm and one-half bandwidth of 58 nm. Growth rates were measured during the first 6 h in darkness using digitally scanned photographic negatives and NIH Image program (National Institutes of Health, Bethesda, MD). Gravitropic curvature was measured from photographic prints as the increment over the initial angle (within 10° from the vertical, since seedlings were grown throughout. The hypocotyls were then washed in 50% ethanol, dehydrated, and embedded in paraffin or Steedman’s wax (Vitha et al., 1997). Longitudinal 10-μm sections were dewaxed and stained for starch with an IKI solution (2% [w/v] KI, 1% [w/v] I).

In WT stems the endodermis is obvious because of the presence of large, sedimented amyloplasts. In NS458 the endodermis can still be recognized by its location as the innermost layer of the cortex. Only endodermal cells within the most apical 0.5 mm (light-grown hypocotyls) or 3 mm (dark-grown hypocotyls) were analyzed for sedimentation and plastid size. Sedimentation was evaluated by counting the number of plastids in the top, middle, and bottom thirds of individual endodermal cells visualized in the microscope. Sections from five hypocotyls were assessed for each genotype and light treatment (WT versus NS458; light versus dark).

For measurement of plastid size, the sections were photographed using a Plan 100× oil-immersion lens (model NA 1.3, Zeiss), and negatives were digitally scanned. NIH Image program software was used to trace plastid outlines at a total magnification of 20,000 to determine the minimum and maximum diameters of each plastid. For NS458, individual starch grains within plastids were also measured. Optical sectioning of endodermal plastids in hypocotyls indicated that plastid shape could best be approximated as an ellipsoid in both genotypes. Thus, plastid volume, V, was calculated for an ellipsoid, where $V = [4/3 \times \pi \times (\text{length}/2) \times (\text{width}/2)^2]$. The same calculation was used for the volume of starch grains within the NS458 plastids.

Stokes’ law was used to calculate the theoretical velocity, $v$, of plastid sedimentation, where $v = 2/9 \times (d_1 - d_2) \times g \times r^2 / \eta$, and $(d_1 - d_2)$ is the difference between the densities of the plastid and the cytoplasm, g is the gravitational constant, $r$ is the plastid radius (calculated from plastid volume assuming a sphere), and $\eta$ is the viscosity of the cytoplasm (here assumed to be 0.3 Pa; Björkman, 1988). The density values used were 1.015 g cm⁻³ for cytoplasm, 1.42 for amyloplasts, and 1.45 for starch. For NS458 plastids, starch occupied 10% of plastid volume (see “Results”), and the density of the remaining volume was assumed to be 1.113 g cm⁻³ (approximated from the density of 30% Glc). The potential energy of the sedimenting particle was calculated as $(d_1 - d_2) \times V \times g \times s$, where $V$ is the plastid volume and $s$ is the distance displaced (here set at 10 μm).

Endodermal cell length was measured using an ocular micrometer from 4-μm sections of hypocotyls that were fixed in glutaraldehyde and embedded in Spurr’s resin as described by Kiss and Sack (1990). Cell-length measure-
ments were made only in regions also used to measure plastid size and sedimentation.

RESULTS

Light Significantly Improves NS458 Hypocotyl Gravitropism

NS458 hypocotyls grown in the dark for an extended period appeared almost agravitropic (Fig. 1E), whereas WT hypocotyls under similar conditions were upright (Fig. 1A). Light substantially improved the gravitropism of NS458 hypocotyls compared with those grown in the dark. (Note that gravitropism was allowed to develop in the dark, regardless of whether the plants had previously been grown in the light or the dark.) Whereas gravitropic curvature was about 10° (after 70 h horizontal) for dark-grown NS458 hypocotyls, it was almost 50° (after 50 h) for light-grown NS458 hypocotyls, a value close to that of the light-grown WT hypocotyls (Fig. 2). In addition, the onset of gravitropic curvature was much earlier and the rate of curvature was much greater in light- versus dark-grown NS458 hypocotyls (Fig. 2). This difference was not the result of growth rate, because dark-grown hypocotyls elongated twice as fast as light-grown hypocotyls (Table I). Also, within the same light treatment, WT and NS458 hypocotyls had the same growth rates (Table I). Thus, the 50% slower rate of gravitropic curvature of light-grown NS458 compared with WT hypocotyls (Fig. 2B) also cannot be explained by a difference in growth rate. Cultivation in the light had much less of an effect on the gravitropism of the WT hypocotyls (Fig. 2).

Light-Grown WT Hypocotyls Are More Sensitive

One measure of gravitropic sensitivity is the presentation time (also known as the induction time), which is the shortest single dose of gravistimulation that is a threshold to curvature. Regression lines were calculated from plots of stimulation time versus curvature (Fig. 3). The regression lines shown in Figure 3 were calculated from all stimulation time versus curvature (Fig. 3). The regression lines shown in Figure 3 were calculated from all stimulation time versus curvature (Fig. 3). The regression lines shown in Figure 3 were calculated from all stimulation time versus curvature (Fig. 3).

Light Increases Mutant Plastid Size but Not the Volume Fraction of Starch

Light increased the plastid size in the endodermis of both genotypes (Table III). Regardless of light treatment, WT plastids were larger than NS458 plastids (Fig. 1, B, D, F, and H). The volume of WT plastids consisted almost entirely of starch, whereas only about 10% of plastid volume was occupied by starch in both light- and dark-grown NS458 plastids (Table III). Thus, although plastids of the light-grown NS458 have more starch on an absolute basis than in the dark-grown mutant, the volume fraction of starch is the same in both the light and in the dark.

Light Induces Mutant Plastid Sedimentation

In both dark- and light-grown WT hypocotyls, the plastids in the endodermis (starch sheath) were amyloplasts that were filled with starch and consistently sedimented (Fig. 1, B, D, J, and K). Previous qualitative observations have shown that dark-grown NS458 hypocotyls lack obvious plastid sedimentation (Kiss and Sack, 1990). Quantification of plastid position confirms this conclusion because plastids were found with equal frequency in the apical, middle, and basal thirds of endodermal cells (Fig. 4). However, plastid sedimentation did occur in the endodermis of light-grown NS458 hypocotyls (Fig. 1, L, and M). Endodermal cells can be identified based on their position outside of the stele and inside of the other cortical layers, even in the absence of an obvious starch sheath (Fig. 1, I, L, and O). Quantification of plastid position in light-grown NS458 hypocotyls confirmed the presence of plastid sedimentation, even though it was not as complete as in the WT (Fig. 4). NS458 endodermal cells were shorter than those of the WT in light-grown hypocotyls (Table IV), a difference that might overestimate the extent of sedimentation in NS458 relative to the wild type because sedimentation was measured by dividing the cell into thirds rather than into cell segments of fixed lengths. Endodermal cell length was essentially unaffected by light treatment in the wild type, whereas these cells were much longer in dark- versus light-grown NS458 hypocotyls (Table IV).

DISCUSSION

The light promotion of hypocotyl gravitropism in NS458 correlates with the onset of plastid sedimentation, a finding consistent with the hypothesis that gravitropic sensing is plastid based. Whereas gravitropism in dark-grown NS458 hypocotyls is severely impaired (Fig. 2) (Kiss and Sack, 1990), mutant hypocotyls from seedlings grown in the light are strongly gravitropic. This light promotion cannot be attributed to effects on growth, because the growth rates of
Figure 1. Gravitropism and endodermal plastid sedimentation. Dark-grown WT plants (WT-D) are gravitropic (A) and contain large, sedimented amyloplasts (B) located approximately at the arrow in A. Dark-grown NS458 hypocotyls (M-D) are severely disoriented (E) and the endodermis (at the arrow in E) contains small plastids with small starch grains (F). Light-grown WT seedlings (WT-L, C) contain large amyloplasts that are sedimented in the endodermis (D, arrows in J and K). Light-grown NS458 seedlings (M-L, G) contain plastids that are larger and contain more starch (H) (compared with dark-grown mutants, F) that are sedimented in the endodermis (arrows in L and M). The position of the lower cell wall is indicated by an arrowhead in M. The endodermis in light-grown NS458 hypocotyls (arrows in I, L, M, and O) can be identified based on its position as the innermost layer of the cortex adjacent to the stele, even though it has less starch than the WT (N). All sections come from tissues that were fixed after being inverted for 1 h (see “Materials and Methods”) except for the section in I, which was kept horizontal for 1 h before fixation. All sections are longitudinal except for those in N and O, which are cross-sections. The gravity vector is toward the bottom of all figures, except for the cross-sections. All sections are of Steedman’s wax stained with IKI to localize starch (blue-black) except for the section in I, which is of Spurr’s resin stained with toluidine blue. Scale bars = 5 μm (B, D, F, and H), 50 μm (J–M), 100 μm (I, N, and O), or 10 mm (A, C, E, and G).
WT and NS458 hypocotyls are equal and because the growth of both genotypes is twice as fast in the dark as in the light.

Light, especially red light, has been shown to promote gravitropism in stems in other genera (Britz and Galston, 1982; Woitzik and Mohr, 1988), and light and phytochrome modulate stem gravitropism (Poppe et al., 1996; Robson and Smith, 1996). We did not investigate either the wavelength(s) or the duration of irradiation necessary for the light promotion of gravitropism in NS458 hypocotyls. But unlike in the studies cited above, we examined the effects of light on gravitropic sensitivity and on the size and sedimentation of endodermal plastids.

The finding that endodermal plastid sedimentation is absent in dark-grown NS458 hypocotyls but is largely present in light-grown mutant hypocotyls suggests that this sedimentation is responsible for the significant promotion by light of gravitropism. This is supported by data for the WT in which the extent of both amyloplast sedimentation and gravitropism are comparable in the light and in the dark. However, we cannot rule out the possibility that light promotes gravitropism independently of an effect on plastid size, e.g. via signal transduction.

Plastid sedimentation in light-grown NS458 probably results from an increase in plastid mass. Although the NS458 mutant has been described as starchless (Hanson and McHale, 1988; Sicher and Kremer, 1996), its plastids do not have amyloplast sedimentation. Table I. Growth rates (mean ± se) of vertically grown hypocotyls

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<thead>
<tr>
<th>Genotype</th>
<th>Growth Rate</th>
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<tr>
<td></td>
<td>Light</td>
</tr>
<tr>
<td>WT</td>
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<tr>
<td>NS458</td>
<td>12.5 ± 2.9</td>
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* Data from Kiss and Sack (1990).
Table II. Gravitropic curvature after intermittent stimulation

The seedlings were intermittently stimulated for 2 h (12 10-min cycles with 0.5–8 min of stationary, horizontal stimulation for each cycle) and then rotated continuously on a clinostat for 1 additional hour. The curvatures (degrees) were tested by paired Student’s t tests to determine whether the values were significantly different from 0. Each number represents the mean from one experiment with 33 to 50 seedlings.

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<td>1.3b</td>
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a Curvature is significantly greater than 0° (α = 0.01). b Curvature is significantly greater than 0° (α = 0.05).

Table III. Plastid and starch volumes in endodermal cells of hypocotyls

Table IV. Length of endodermal cells of WT and NS458 hypocotyls

Figure 4. Plastid sedimentation in light- and dark-grown hypocotyls. Shown is the percentage of plastids present in each third of endodermal cells of seedlings that were inverted for 1 h, fixed in this orientation, embedded in wax, sectioned, and stained. Plastids are sedimented, except in dark-grown NS458 hypocotyls. Bars = ±SE. A total of 39 to 61 endodermal cells from five different hypocotyls were scored for plastid sedimentation for each genotype and condition. Open bars, Light-grown WT; black bars, dark-grown WT; white bars with horizontal lines, light-grown NS458; and black bars with horizontal white lines, dark-grown NS458.

Table IV. Length of endodermal cells of WT and NS458 hypocotyls

Cell length (mean ± se) from the apical part of the hypocotyl (see text). n = 40 to 85 cells. All values are different (α < 0.01); analysis of variance, Newman-Keuls multiple comparisons.

Figure 4. Gravitropic curvature after intermittent stimulation

The seedlings were intermittently stimulated for 2 h (12 10-min cycles with 0.5–8 min of stationary, horizontal stimulation for each cycle) and then rotated continuously on a clinostat for 1 additional hour. The curvatures (degrees) were tested by paired Student’s t tests to determine whether the values were significantly different from 0. Each number represents the mean from one experiment with 33 to 50 seedlings.

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<table>
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<th>Genotype</th>
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<th>Dark</th>
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<tr>
<td>WT</td>
<td>97 ± 2.9</td>
<td>89 ± 5.0</td>
</tr>
<tr>
<td>NS458</td>
<td>66 ± 2.1</td>
<td>126 ± 6.6</td>
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</table>
be capable of discriminating a signal from noise at some threshold greater than thermal noise (Sack, 1997). If all gravitropic sensing is plastid based, then the potential energy in the plastids of dark-grown NS458 may be sufficient to barely trigger enough gravitropic sensing to produce the residual levels of gravitropism observed in these hypocotyls (Kiss and Sack, 1990). The estimated 4-fold greater potential energy of light-grown (compared with dark-grown) mutant plastids may be sufficiently higher than threshold levels to dramatically increase the orientational signal and thus gravitropism. The increased mass may also be important in inducing sedimentation, which places these plastids in a part of the endodermal cell that may contain or be enriched in mechanosensitive receptors that transduce plastid mass.

This reasoning might also explain why light-grown WT hypocotyls are at least twice as sensitive as NS458 hypocotyls, as estimated by the presentation time and by intermittent stimulation. The reduced sensitivity of the light-grown mutant compared with the WT might result both from the lower plastid mass and from the less-consistent sedimentation of mutant plastids. Based on data from roots of starch-deficient mutants of Arabidopsis, Kiss et al. (1996, 1997) estimated that mutants, the plastids of which contain \( \geq 60\% \) of the starch of the WT, exhibit full WT levels of gravitropic sensitivity. The starch content of NS458 plastids is significantly below this level.

The relative importance of plastid sedimentation to gravitropic sensitivity may differ in stems and roots. As in hypocotyls, light-grown roots of NS458 are strongly gravitropic but less sensitive than the WT, but mutant roots apparently lack sedimentation in columella cells (Kiss and Sack, 1989). These data show that plastid sedimentation is not necessary for significant gravitropism in roots, whereas there is a strong correlation for hypocotyls of the same mutant. This difference might be related to the much smaller size of columella versus endodermal cells.

Although light promotes gravitropism in NS458 hypocotyls, comparable data are not available for an Arabidopsis mutant that is also deficient in plastidic phosphoglucomutase, the pgm1 locus represented by two alleles, Tc7 and ACG21 (Caspar and Pickard, 1989; Kiss et al., 1996). pgm1 appears to be completely devoid of starch (Caspar, 1994). Dark-grown hypocotyls are severely disoriented, although they are capable of weak gravitropism (Caspar and Pickard, 1989; Kiss et al., 1997). Depending on the report, light-grown pgm1 hypocotyls either show WT levels of curvature (fig. 7 in Caspar and Pickard, 1989) or less gravitropic curvature than in mutant hypocotyls grown in the dark (figs. 5 and 6 in Kiss et al., 1997). Neither sensitivity nor plastid size and position were measured in these studies. Based on our results with NS458, we predict that light-grown pgm1 hypocotyls should be slightly more sensitive than dark-grown mutant hypocotyls; whereas light might enlarge pgm1 plastids, the increase in plastid mass should be much less than in NS458 because, unlike NS458 plastids, pgm1 plastids entirely lack starch.

In summary, our data are consistent with the conclusions that the light promotion of gravitropism in NS458 results from an increase in plastid mass that induces plastid sedimentation and greatly increases gravitropic sensitivity compared with the dark-grown mutant. These data support the importance of both plastid mass and plastid sedimentation in gravitropic sensing.

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**LITERATURE CITED**


Robson PRH, Smith H (1996) Genetic and transgenic evidence that phytochromes A and B act to modulate the gravitropic


