Actin expression in germinating seeds of *Phaseolus vulgaris* L.

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Abstract. Actin was present at very low levels in the seeds of common bean (*Phaseolus vulgaris* L.) compared with those from other species, and was observed mostly in the embryo. A time-course of actin expression in germinating bean seeds revealed an induced expression of both the mRNA and protein. Initially, the actin mRNA in seeds was barely detectable by northern blot analysis. However, there was a substantial increase in the expression of the actin mRNA at 24, 48 and 72 h after imbibition, compared with an internal control consisting of a late-embryogenesis-abundant (LEA) type IV gene from *P. vulgaris*. An increase in the amount of actin in total seed extracts that paralleled that of the mRNA was detected by western blotting starting at 24 h after imbibition. This increase was more apparent when the embryo alone was analyzed. Two-dimensional western blots initially revealed three actin isoforms with isoelectric points (pl's) of approximately 5.6, 5.7 and 5.8, the amounts of which increased within a 48-h period, when a new minor isoform of pl approximately 5.5 appeared; however, after 72 h, the pl-5.8 isoform had almost disappeared and the pl-5.5 isoform had disappeared completely, indicating that these two minor isoforms are expressed transiently. These results indicate that actin is at very low levels in the dry seed but undergoes an increased and differential expression during imbibition, an event probably required to carry out all the necessary functions for germination.

Key words: Actin – Cytoskeleton – Gene expression (actin) – Germination – Imbibition – *Phaseolus* (seed actin)

Introduction

The cytoskeleton is a fundamental organelle that is involved in most essential cellular functions of eukaryotes. One of its components, actin is a ubiquitous protein that is highly regulated at various levels, including transcriptionally (Lloyd and Gunning 1993) and post-translationally (Aktories and Wegner 1992; Howard et al. 1993; Jungbluth et al. 1995). In addition, actin can self-assemble into microfilaments, and the equilibrium between polymeric and monomeric actin is also under tight cellular control by pH, salts, and a subset of proteins that can interact with both the polymer and the monomer (Pollard and Cooper 1986; Sheterline and Sparrow 1994). All these control points are necessary since small variations in the actin monomer/polymer equilibrium can dramatically alter cell shape and function. For example, the capability of actin to polymerize can be affected by specific phosphorylation at tyrosine residues in *Dictyostelium discoideum* cells as a response to stress (Jungbluth et al. 1995). Consequently, this increased actin phosphorylation causes pseudopodium loss, rounding-up and reduced adhesion of *D. discoideum* cells to the substratum (Howard et al. 1993). Similarly, over-expression of an actin-binding protein such as profilin, which is presumed to induce actin depolymerization, produces slow growth, swelling and lysis of *Saccharomyces cerevisiae* cells, as well as an altered actin and chitin distribution compared with the wild type (Haarer et al. 1990).

While most of the data on cytoskeletal function and regulation have been obtained from studies on eukaryotes from the animal kingdom, the cytoskeleton in plant cells has also been reported to contain the main primary protein components such as actin (Vahey and Scordilis 1980; McLean et al. 1990b; Villanueva et al. 1990; Liu and Yen 1992), tubulin (Lloyd et al. 1979; Yadav and Filner 1983) and intermediate filament-like proteins (Beven et al. 1991). Among these proteins, tubulin (Yadav and Filner 1983) and actin (Andersland et al. 1992; Liu and Yen 1992; Ren et al. 1997) have been shown to self-associate, in vitro, into microtubules and
microfilaments, respectively. Although there is a notable lack of information on the particular roles and regulation of actin in plants, there is increasing substantial evidence on the involvement of actin in many plant cellular functions. Plant actin has been observed to be an important component that works in concert with tubulin during cell division (Schmit and Lambert 1990), and the disruption of the actin network leads to a number of effects including cessation of cytoplasmic streaming, cell division, organelle movement and tip growth (McCurdy and Williamson 1991). In addition, microinjection of the actin-binding protein profilin into Tradescantia stamen hair cells disrupts microfilament organization and the cells stop cytoplasmic streaming (Staiger et al. 1994). Thus, actin is a fundamental protein in the plant kingdom as well.

Due to the variety of roles of actin in many different plant cellular functions, it is also expected that plant actin is tightly regulated wherever cytoskeletal function occurs. For example, it has been reported that actin displayed a differential expression of isoforms during nodule ontogeny in *P. vulgaris* (Pérez et al. 1994). This differential expression of isoforms might be induced as early as nodule ontogeny as the initial infection steps, during the rearrangement of the actin network of *P. vulgaris* root hair cells that is induced by Nod factors derived from *Rhizobium etli* (Cárdenas et al. 1998). In soybean, actin was reported to consist of highly divergent isovariants with possible roles in the differential regulation necessary for the plasticity of cytoskeletal function (McLean et al. 1990b). Furthermore, antibodies generated against the specific λ-, κ- and ι-actin isoforms, showed a preferential distribution of the isoforms themselves. These studies also suggest that the expression of actin is not as constitutive during cell division, organelle movement and tip growth (McCurdy and Williamson 1991). Antisera on the involvement of actin in many plant cellular functions, it is also expected that plant actin is tightly regulated wherever cytoskeletal function occurs.

Materials and methods

**Plant material.** Seeds from *Phaseolus vulgaris* L. were used at all times and were obtained from the local supermarket. Seeds were surface-sterilized in 10% (v/v) commercial bleach, rinsed in running tap water, sown on water-saturated towels and germinated in the dark at 27 ± 1 °C and 100% relative humidity. Whole seedlings and embryo axes were harvested at different times as indicated and stored at −70 °C until used.

### Antibodies and chemicals

Anti-(calf thymus actin) polyclonal antibodies were a kind gift from Dr. John L. Wang, Michigan State University (East Lansing, USA) and have been shown to cross-react with plant actin (Villanueva et al. 1990). Monoclonal anti-actin antibody N350 was purchased from Amersham (Arlington Heights, Ill., USA) and has also been shown to cross-react with plant actin (McLean et al. 1990b). Alkaline-phosphatase-conjugated anti-rabbit antibodies and substrates were from Boehringer-Mannheim (Indianapolis, Ind., USA). Ampholytes were purchased from Sigma (St. Louis, Mo., USA). All other chemicals were reagent grade.

### Preparation of material for protein and RNA extracts

The surface-sterile seeds and embryo axes were used to prepare a powder by grinding them in liquid nitrogen and then in a mill. The frozen powders were kept at −80 °C until further analysis. Alternatively, a flour was also prepared by grinding the dry clean seeds or the cotyledons and embryo axes separated from them in a coffee mill.

### Extraction of RNA northern blot analysis and reverse transcriptase-polymerase chain reaction (RT-PCR)

Total RNA was prepared from the frozen powdered material by the hot-phenol method, following the procedure of de Vries et al. (1991). A 5-µg sample of total RNA was electrophoresed on a 1% agarose gel in the presence of 2.2 M formaldehyde and transferred onto nylon membranes (Sambrook et al. 1989). Blots were hybridized with *P. vulgaris* late-embryogenesis-abundant (LEA) class-IV (lea4-25) cDNA (Colmenero-Flores et al. 1997) and actin cDNA from leaves (*PLACT-6)*, (unpublished). Hybridization and washes were done at high stringency (Church and Gilbert 1984) and subjected to autoradiography. The RNA load was standardized by visualizing the ribosomal RNA in the gels after ethidium bromide staining. For RT-PCR, we used actin universal oligonucleotides from the carboxy terminus which is highly conserved (Sheterline and Sparrow 1994). The 5′ oligonucleotide encoding an adaptor sequence and a 5′ region (5′ GGAAATCCAGTGTCGGA- TTGGTG 3′), and the 3′ oligonucleotide (5′ GACAGTTT- CCTGTGAACTATTGATGGCCCAG 3′) were incubated in the presence of total cDNA synthesized by reverse transcriptase from total RNA extracted from different parts of the plant. The reaction was carried out using the Superscript II kit according to the instructions of the manufacturer (Gibco BRL, Gaithersburg, Md., USA) from 0 to 25 cycles, and the amplified cDNA was analyzed on 2% agarose gels. In some cases, the amplified DNA bands were quantified by densitometry by the same method used for protein bands (see below).

### Sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE), two-dimensional SDS-PAGE (2-D PAGE) and immunoblotting

For whole-seed flour, the powdered material was suspended in 1% SDS and boiled immediately for 8 min. Alternatively, the same material was suspended in Laemmli’s sample buffer and processed the same way. The supernatants were recovered by centrifugation at 14,000 g and analyzed by SDS-PAGE according to Laemmli (1977). Similarly, 0.1 g of frozen powdered embryo was suspended in Laemmli’s sample buffer and boiled immediately for 8 min. For analysis by 2-D PAGE, the embryo extracts in Laemmli’s sample buffer were diluted at a 1:1 ratio in O’Farrell’s (O’Farrell 1975) lysine buffer (9.5 M urea, 5% β-mercaptoethanol, 2% nonidet P-40, 1.6% ampholytes pH 5–7, and 0.4% ampholytes pH 3.5–10) (Sinclair and Rickwood 1981). The samples were then analyzed by isoelectric focusing in the first dimension and SDS-PAGE in the second dimension according to O’Farrell (1975). For immunoblotting, the gels were electrotransferred to nitrocellulose at 400 mA overnight (Towbin et al. 1979) and blocked with 3% bovine serum albumin in phosphate-buffered saline (PBS; 10 mM Na-phosphate; 150 mM NaCl, pH 7.4) for 1 h at 50 °C. The membranes were then incubated overnight at 4 °C in either a 1/1000 dilution of monoclonal anti-actin antibodies, or a 1/2000 dilution of an anti-(calf thymus actin) polyclonal antibody in

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0.05% Triton X-100 in PBS (PBST). After three washes of 15 min in PBST, the membranes were incubated with a 1/5000 dilution of the appropriate peroxidase- or alkaline-phosphatase-conjugated secondary antibodies for 1 h at 25 °C. The membranes were then washed three times for 15 min with PBST, rinsed briefly in PBS and the proteins were visualized by chemiluminescence or by incubating in a solution of nitro-blue tetrazolium and bromo-chloro indolyl phosphate according to the manufacturer (Boehringer-Mannheim).

Protein quantification and densitometry. The uniformity of protein loading in the gels was normalized by previously running the various extracts and staining with Coomassie blue. An approximation of the amount of protein loaded was determined according to Bradford (1976) using bovine serum albumin as a standard.

For densitometric scans, the images of the western blots or photographic films were captured by an Apple Color One Scanner using Adobe Photoshop software and the bands quantified according to their pixel value using NIH-Image software.

Results

Actin is present at low levels in P. vulgaris seeds and predominantly in the embryo. The presence of actin was detected by western blotting using an antibody to calf thymus actin that is known to cross-react with plant actin (Villanueva et al. 1990). When 40 µg of protein from several extracts from different dry seeds was analyzed, even with a 2-fold higher protein load for P. vulgaris extracts, the actin could not be detected (Fig. 1A, lane 1), whereas the 42-kDa band was readily detected in maize and oat seed extracts (Fig. 1B, lanes 2 and 3, respectively). Possible actin degradation products were also detected in a rice extract (Fig. 1B, lane 4). The facts that: (i) the amount of protein loaded was equivalent in lanes 2–4 (Fig. 1A); (ii) lane 1 where actin was not observed was actually about 2-fold overloaded with respect to the lanes where actin was detected (Fig. 1A); and (iii) we were unable to detect proteolytic activity using gelatin as substrate in seed extracts (not shown), indicated that in P. vulgaris seeds is present at very low levels and that this finding was not due to a differential extraction caused by omitting the β-mercaptoethanol, Tris and glycerol from the extraction buffer. In fact, an identical reaction was observed in lanes loaded with the same amount of protein extracted with either 1% SDS alone (Fig. 2A, lane 2) or Laemmli’s sample buffer (Fig. 2A, lane 3). In addition, the only way to observe the actin band in the P. vulgaris extracts was by overdeveloping the blot (Fig. 2A, lanes 2, 3; Fig. 3), but this caused a high background in the rest of the membrane (Fig. 3). A more detailed analysis of the different parts of the seed by western blotting using equal amounts of protein in each lane and after over-developing the blot, revealed that in P. vulgaris seeds is present mostly in the embryo (Fig. 2B, lane 2) since the equivalent extract from total seed (Fig. 2B, lane 1) or cotyledon (Fig. 2B, lane 3) showed a very faint reaction. These data indicated that most of the actin during germination arises from the embryo, and prompted us to check the time-course of its expression in both total seeds and embryo axes.

The actin product shows a substantial increase during imbibition. Western blot analysis and overdeveloping the blot for a longer time than used previously (Fig. 1) for whole seed extracts revealed that, initially, actin was hardly detectable (Fig. 3, 0 h) and that its level remained essentially unchanged during the first 24 h of imbibition (Fig. 3, 0–24 h). A substantial amount of the protein product was only detected after 48 h and prevailed from then until 72 h after imbibition (Fig. 3, 48–72 h).

The results confirmed our expectation that the majority of the actin was found in the embryo (Fig. 2B), since it is the seed part that elongates during germination to become a new plant. Therefore, we repeated the time-course of induction but analyzing only the embryo axis. The analysis was carried out with both a polyclonal (Villanueva et al. 1990), and a monoclonal anti-actin...
The positive by electrotransfer and immunoblotting. The blot shows actin in the amounts (SDS-PAGE sample buffer) and the supernatants loaded in equivalent amounts (calf thymus actin) polyclonal antibodies. The powdered material from seeds and seedlings at different time points was extracted with sample buffer. The seeds were separated into embryos and cotyledons, and extracts from each were made as for the whole seed. Equivalent amounts of protein (≈ 40 μg) were loaded in lanes 1–3. The 42-kDa actin band was not detected in whole seeds (lane 1) or cotyledons (lane 3) but it was readily detectable in embryos (lane 2) and the positive chicken-muscle control (lane 4). In both cases (A and B), the immunoblot had to be overdeveloped in order to visualize the 42-kDa polypeptide (arrows) in whole-seed and cotyledon extracts.

Fig. 2. A Western blot analysis with anti-(calf thymus actin) polyclonal antibodies of whole P. vulgaris seeds. The seeds were extracted with 1% SDS (lane 2) or sample buffer (lane 3) and equivalent amounts of protein were loaded in lanes 2 and 3 (≈ 50 μg). The actin polypeptide was barely detectable even after overdeveloping the blot (lanes 2, 3); however, the chicken-muscle actin control was readily detected (lane 1). B Western blot analysis with anti-(calf thymus actin) polyclonal antibodies of different components of P. vulgaris seeds extracted with sample buffer. The seeds were separated into embryos and cotyledons, and extracts from each were made as for the whole seed. Equivalent amounts of protein (≈ 40 μg) were loaded in lanes 1–3. The 42-kDa actin band was not detected in whole seeds (lane 1) or cotyledons (lane 3) but it was readily detectable in embryos (lane 2) and the positive chicken-muscle control (lane 4). In both cases (A and B), the immunoblot had to be overdeveloped in order to visualize the 42-kDa polypeptide (arrows) in whole-seed and cotyledon extracts.

Fig. 3. Time-course analysis of actin during germination of whole seeds of P. vulgaris as shown by western immunoblotting using anti-(calf thymus actin) polyclonal antibodies. The powdered material from seeds and seedlings at different time points was extracted with SDS-PAGE sample buffer and the supernatants loaded in equivalent amounts (≈ 50 μg) directly onto 10% polyacrylamide gels, followed by electrotransfer and immunoblotting. The blot shows actin in the dry seed at 0, 2, 4, 8, 12, 24, 48 and 72 h after imbibition. The positive chicken-muscle actin control (CA) is also shown.

(McLean et al. 1990b) antibody and the corresponding blots were analyzed by densitometry (see Materials and methods) to assess the relative increase in actin. In the case of the polyclonal antibody, we observed a certain difference in detection of the various RNAs was not (data not shown). These data indicate that actin is at low levels in the embryo axis of dry seeds and the polypeptide increases substantially after the first 24 h of germination. This is the reason for detecting the protein only after 48 h of germination in whole seeds.

The actin mRNA is induced during germination in seeds of P. vulgaris. Initially, when we analyzed 5 μg of total mRNA, we were surprised that actin mRNA could not be detected in dry seeds of common bean on northern blots probed with a P. vulgaris leaf cDNA (PLACT-6) (Fig. 5A, 0 h). Therefore, we carried out an analysis of the actin mRNA using the P. vulgaris leaf cDNA (PLACT-6) in whole seeds at various time points after imbibition. The actin mRNA was not detected during the first 0–12 h (Fig. 5A, PLACT-6, 0–12 h). The message was barely detectable even after 24 h of imbibition (Fig. 5A, PLACT-6, 24 h). However, its expression increased substantially after 48 h (Fig. 5A, PLACT-6, 48 h). In contrast, the message for a seed-specific gene, LEA-IV showed an initial high expression and then a decrease as the time of imbibition proceeded (Fig. 5A, lea4-25). In addition, analysis of the actin mRNA in embryo axes of P. vulgaris as shown by western immunoblotting using anti-(calf thymus actin) polyclonal antibodies. The powdered material from embryo axes at different time points was extracted with sample buffer and the supernatants loaded in equivalent amounts (≈ 45 μg) onto 10% polyacrylamide gels followed by electrotransfer and immunoblotting. The blot shows the presence of actin at 0, 6, 12, 24, 48 and 72 h after imbibition.
due to differential loading of the gel, the RNA amount loaded in all lanes was standardized using the ribosomal RNAs as a reference (Fig. 5A, B, rRNA). The dry-seed actin mRNA could not be detected by northern blot analysis; however, we were able to observe the presence of seed actin mRNA by RT-PCR with 5′ and 3′ conserved actin oligonucleotides from the carboxy region, after 25 cycles of amplification (Fig. 5C, seed). The results were consistent with the fact that the actin mRNA in seeds is present in low amounts since the amount detected in this tissue after 25 cycles of amplification was about one-half lower than that of flowers, leaves and nodules (Fig. 5C, densitometric scan). This indicated that the initial amount of seed actin mRNA is not enough to be observed by a regular northern blot analysis under our current detection parameters. These data also suggest that the increase in protein detected in whole seeds during germination is most likely due to an increase in the expression of the actin mRNA that takes place in the embryo.

The induction of the protein is due to an increase in the amount of the two main actin isoforms. To analyze with more detail the characteristics of the actin induction we ran the embryo extracts on 2-D PAGE gels followed by western immunoblotting to detect the isovariants. To ensure that both monoclonal and polyclonal antibodies recognized the same isoforms, the blots initially probed with the monoclonal antibody (Fig. 6A) were washed with SDS-urea to strip off the bound antibodies, re-probed with the immunopurified polyclonal anti-actin antibody and developed by chemiluminescence (Fig. 6B). The results obtained with either antibody were identical (Fig. 6A, B), indicating that both antibodies recognize the same actin isoforms. We observed that in the dry embryo three isoforms of actin already exist, two major isoforms migrate at isoelectric point (pIs) of approx. 5.6 and 5.7, respectively, and a minor isoform at pI 5.8 (Fig. 6, 0 h). The two major isoforms increase in amount as detected after 24, 48 and 72 h of imbibition (Fig. 6, 24–72 h). Additionally, a new minor isoform of pI ≈ 5.5 appears after 48 h (Fig. 6, 48 h); however, after 72 h of imbibition this new actin isoform is no longer detected (Fig. 6, 72 h). There was some variability in the stretch of the isoelectric focusing gel on which the isovariants migrated; however, the monoclonal antibody cross-reacts with a higher-molecular-weight spot (Fig. 6A, arrow) which has also been observed in soybean (McLean et al. 1990a). This spot was used as an internal reference which allowed us to identify the two major actin isoforms since it was always observed above and between the area where these two major isoforms were focused (Fig. 6A, arrow).

Discussion

When we tried to detect actin in dry seed extracts of *P. vulgaris*, we were unable to obtain a detectable signal; in contrast, profilin can be readily detected in whole-seed extracts (data not shown). When we compared extracts from different seeds, we observed that the 42-kDa polypeptide corresponding to actin was readily detectable in maize and oats, and possible degradation products from it in rice, but at the same or a higher protein concentration, common-bean actin was still undetectable (Fig. 1). The absence of actin was not
due to degradation during the extraction since common-
bean seeds have no detectable proteolytic activity (not
shown). Further analysis of separate components of the
seed revealed that a 42-kDa polypeptide corresponding
to actin was indeed present in the dry seed but it
predominated in the embryo (Fig. 2B) and it could be
observed in whole seeds only after developing the
western blot for a long time (Fig. 3). These data are
noteworthy in the context that the embryo is the part of
the seed that will become a new plant and it requires the
necessary machinery for the dynamic processes that will
occur at the onset of germination. The cytoskeleton is
certainly an essential component that will be required
during this process. Consistent with these data is the fact
that profilin is also present predominantly in the embryo
from *P. vulgaris* seeds (data not shown).

Actin has been previously used as an internal control
for constitutive genes (Cook et al. 1995). In vegetative
tissues this may work properly as a control for baseline
gene expression but it may ultimately depend on the
particular specificity of the probe and the tissue tested.
In fact, it was recently reported that an *Arabidopsis* actin
gene, *ACT7*, has a preferential expression in rapidly
growing zones of this plant (McDowell et al. 1996). The
finding of extremely low levels of actin in the dry seed of
*P. vulgaris* suggested the possibility that, in common
bean, there was also a differential actin expression
during the germination process. When we performed the
analysis of protein expression at different time points of
germination, we found that the protein was indeed
induced slowly at first (0–24 h) and then more strongly
after 48 h of imbibition (Fig. 3). We found a 2- to 5-fold
increase in the amount of the protein which was
consistent with the level of expression of the corre-
sponding mRNA. The times of induction were essen-
tially the same when either the whole seed or the embryo
axes alone were analyzed. Although the actin mRNA
could not be detected at time = 0 of imbibition (whole

**Fig. 6A,B.** Two-dimensional gel electrophoretic analysis of actin isoform expression at different time points after imbibition of *P. vulgaris* seeds. Embryo axes were excised and extracted in SDS-PAGE sample buffer and analyzed by 2D-PAGE according to O’Farrell (1975). Equal amounts of protein for all time points (≈ 40 μg) were loaded in the first dimension. **A** Immunodetection of the actin isoforms with Amersham N350 monoclonal antibody and alkaline-phosphatase-conjugated secondary antibodies at 0, 24, 48 and 72 h after imbibition. The arrow points toward a cross-reactive protein that was used as a reference point. **B** The bound antibodies were stripped off the blots and re-probed with immunopurified anti-calf thymus actin polyclonal antibodies followed by peroxidase-labeled secondary antibodies, and developed by chemiluminescence. The pI values of the corresponding isoforms are shown in the bottom panels. The developmental stage of the seedling is schematized along with the time point at the right of the figure.
dry seed) by northern blot analysis, we were able to amplify actin cDNA from seed mRNA by RT-PCR, indicating that the message is nevertheless present in the dry seed. The amount of the amplified cDNA detected in this plant component after 25 cycles was approximately 50% less than that obtained from flower, leaf and nodule mRNA (Fig. 5C), which supports the fact that initial actin mRNA in the seed is present in low quantity.

One possibility to account for the increase in the expression of actin was that new genes were being induced or their products modified during the process. Analysis of the extracts at the different time points by 2D-PAGE indicated that: (i) actin is present in the embryo of the common-bean seed as one minor and two major isoforms; (ii) that as many as four isoforms are detected after 48 h of imbibition but the appearance of the new minor isoform seems to be transient; and (iii) the increase in the protein product is mostly due to the increase in the quantity of the two main isoforms initially present in the embryo instead of a shift of isoform expression (Fig. 6). It is also noteworthy that the PI's determined for the actin isoforms in the common bean correspond closely to actin isoforms already reported from soybean (McLean et al. 1990b), which would suggest an equivalent expression in both species during the vegetative stage. We have determined that there are five to six actin genes present in the P. vulgaris genome (data not shown) so that each one of the proteins detected could account for one particular gene product. There is a possibility though, that they could arise from a few genes and be modified post-translationally. Such post-translational modifications of actin have been documented in the literature previously (Aktories and Wegner 1992). As was expected from the protein data, the lack of detection of the actin mRNA in the primary stages of germination did not mean that it was not present; indeed, RT-PCR analysis using universal and plant-actin-specific oligonucleotides showed that the seed actin mRNA could be amplified to a detectable level (Fig. 5c, seed).

It is clear that actin is not always a constitutively expressed protein as previously thought. High levels of regulation of actin have now been encountered at the transcriptional (Lloyd and Gunning 1993), translational and post-translational (Aktories and Wegner 1992; Howard et al. 1993) levels. It remains to be determined if one or both of the main actin isoforms that increase during germination in common bean are products arising from a homolog of the ACT7 gene (McDowell et al. 1996). This gene was observed to be highly expressed in rapidly expanding tissues of Arabidopsis, including a strong expression in the seedlings immediately after imbibition. Its expression was also observed to increase upon treatment with external stimuli such as auxin, light and wounding (McDowell et al. 1996). Thus, it would not be surprising to find an analogy in the expression of actin genes during the germinative process of common bean.

Finally, we could envision two possible scenarios for the existence of such a small amount of actin in cotyledons and the majority of the actin in the embryo.

One would be that actin is not required for the initial triggering of germination. The other possibility is that the basal-level amount of actin present in the seed is enough to allow the embryo to carry out the baseline functions required to trigger the germination event. Once this has occurred, actin expression is induced to continue with the development of the new plant.

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