Two Class XI Myosins Function in Organelle Trafficking and Root Hair Development in Arabidopsis

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Multigene families encoding class XI myosins are conserved in higher plants, however, little information is available on specific functions of these ubiquitous molecular motors. We isolated gene knockout mutants for all 13 class XI myosins present in Arabidopsis (Arabidopsis thaliana) genome. Inactivation of 11 myosin genes resulted in no discernible phenotypes under the normal growth conditions. In contrast, the knockouts of the remaining two myosin genes, XI-2 (formerly MYA2) and XI-K, exhibited similar defects in root hair elongation suggesting that the myosin-driven motility plays a significant role in a polar tip growth. Strikingly, inactivation of each of these myosins also reduced trafficking of Golgi stacks, peroxisomes, and mitochondria in root hairs and in leaf epidermal cells. These results indicate that myosins XI-K and XI-2 play major and overlapping roles in the cell dynamics in Arabidopsis and highlight the redundant nature of myosin function in plants.

Myosins are signature molecular motors of eukaryotes that are involved in a broad spectrum of actin cytoskeleton-associated types of cellular dynamics (Vale, 2003). Comparative genomics revealed that myosins are conserved throughout the eukaryotic domain of life (Richards and Cavalier-Smith, 2005; Foth et al., 2006). Land plants possess two myosin classes, XI and VIII, each of which is evolutionary, related to animal and fungal class V myosins (Desnos et al., 2007), suggesting their origin was from a common ancestor that antedated the divergence of Plantae and Opisthokonts (Foth et al., 2006). Subsequent evolution of myosins was dominated by gene duplication and diversification that resulted in the presence of more than 10 myosin genes in all plant genomes sequenced so far (see accompanying article Avisar et al., 2008). In particular, Arabidopsis (Arabidopsis thaliana) encodes 13 class XI and four class VIII myosins (Reddy and Day, 2001). The studies that mostly involved cytoskeletal inhibitors have demonstrated the principal role of actomyosin motility in plant cell dynamics including organelle trafficking, remodeling, and inheritance (Boevink et al., 1998; Nebenfuhr et al., 1999; Sheahan et al., 2004; Kim et al., 2005; Runions et al., 2006).

Some of the class XI myosins were found in association with organelles suggesting their involvement in organelle transport (Wang and Pesacreta, 2004; Hashimoto et al., 2005; Li and Nebenfuhr, 2007; Reisen and Hanson, 2007). However, information on functional profiles of individual myosin motors is very limited. Two recent publications have demonstrated the role of Arabidopsis myosin XI-K in root hair growth (Ojangu et al., 2007), and implicated rice (Oryza sativa) myosin XI-B in pollen development (Jiang et al., 2007).

Here we screen the gene knockouts of all 13 class XI myosins of Arabidopsis to show that, in addition to myosin XI-K, myosin XI-2 (MYA2) is also required for root hair development. Furthermore, we demonstrate that each of these two highly expressed myosins functions in the rapid movement of the Golgi stacks, peroxisomes, and mitochondria in roots and leaves. Interestingly, inactivation of the genes encoding the most closely related paralogs of myosins XI-K and XI-2 and myosins XI-1 (MYA1) and XI-B, respectively, did not impair root hair growth or organelle trafficking. These results indicate that evolution of myosins in plants combines opposing tendencies of functional specialization and functional redundancy.

RESULTS

Isolation of the Homozygous Knockout Lines

Sixteen homozygous lines in which each of the 13 class XI myosin genes of Arabidopsis was inactivated by T-DNA insertion were obtained and the exact localization of the insert was determined by sequencing (Fig. 1A; data not shown). Furthermore, inactivation of the target genes was demonstrated using semiquantitative reverse transcription (RT)-PCR (e.g. Fig. 1B). The corresponding mRNAs were undetectable by RT-PCR analysis, therefore confirming complete abolishment
of myosin expression for each of the 16 knockout lines. To ensure that the observed phenotype is attributed solely to the inactivation of the myosin XI-K locus, line SALK_067972 was further backcrossed twice to the wild-type plant, and the homozygous progeny was selected and used for the experiments described below. In addition, two independent lines for myosin XI-K and three for myosin XI-2 genes were selected for further experiments to ensure that the observed phenotypes were due to the T-DNA insertion in the corresponding locus rather than to secondary site mutations. For both of the obtained independent myosin XI-K gene knockout lines, the lack of an expressed protein was confirmed using a polyclonal antibody specific to this myosin (Fig. 1C).

**Inactivation of Myosins XI-2 and XI-K Induces Similar Defects in Root Hair Growth**

Inspection of the insertional lines in which each of the 13 class XI myosin genes present in Arabidopsis genome was inactivated (Fig. 1; data not shown) revealed no detectable developmental defects in the aerial organs of plants grown under normal conditions. Therefore, the seeds of each line were planted to vertical plates to screen for root morphology. Although the overall root sizes and shapes appeared normal in all knockout lines, four lines exhibited obvious defects in the elongation of the root hairs (Fig. 2A). These lines were knockouts of two class XI myosin genes, namely, XI-2 and XI-K. Quantification of the root hair length for each of these lines revealed a dramatic reduction in the mean root hair length that varied from 28% to 40% of that in the parental Columbia line (Fig. 2B). Comparative analysis showed that the differences in the root hair length between each of these four knockout lines and the wild-type line were statistically significant with \( P < 0.001 \) for all lines.

Interestingly, analysis of the light-regulated chloroplast relocation performed as described in an accompanying work by Avisar et al. (2008) revealed that none of the myosin gene knockouts exhibited observable defects in chloroplast movement (data not shown). This result suggested that several myosin motors may have overlapping functions in chloroplast movement. Alternatively, it is possible that although chloroplast movement requires intact actin cytoskeleton (Faves and Truve, 2007), it does not involve myosins.

To confirm the results of the gene knockout experiments by an independent approach and to examine the usefulness of the dominant negative inhibition of the myosin function that we used for *Nicotiana*.
benthamiana in an accompanying article (Avisar et al., 2008) for Arabidopsis, we generated several transgenic lines that expressed the cargo-binding, globular tale domain (GTD) of myosin XI-K. The hemagglutinin (HA)-epitope tagging of this domain was used to select transgenic lines that exhibited the highest levels of GTD expression (Fig. 2C). Because the mean root hair length in these selected lines was only 38% of that in the control (Fig. 2A, far right), very similar to that found for myosin XI-K knockout mutants, these data validated the use of the dominant negative inhibition approach in Arabidopsis.

Given that the principal function of myosin motors is physical translocation of various cargoes, our findings suggest that such translocation powered by myosins XI-2 and XI-K is required for the rapid polarized growth of the root hairs.

Myosins XI-2 and XI-K Are Required for the Rapid Organelle Transport in Root Hairs and Leaves

What are the cargoes transported by the myosins XI-2 and XI-K within the growing root hairs? To address this question, we studied trafficking of Golgi stacks and peroxisomes in the gene knockout lines that were transformed to express the fluorophore-tagged, organelle-specific reporters (Fig. 3, A–C). The mitochondrial transport was examined using a vital fluorescent dye Rhodamine 123 (Fig. 3E). Organelle trafficking in the live root hairs was visualized by confocal microscopy and the resulting movies were used for computer-assisted organelle tracking and velocity measurements.

As expected, the predominant pattern of Golgi movement was along the root hair longitudinal axis (Fig. 3A). It should be noted, however, that not infrequently the neighboring individual Golgi stacks were moving in the opposite directions or with the drastically distinct velocities (data not shown). This pattern and a mean Golgi velocity of approximately 1 μm/s were very similar to those recently described for Arabidopsis root hairs (Campanoni et al., 2007). It seems, therefore, that Golgi stacks move independently of each other rather than in a uniform stream-like pattern.

Strikingly, we found that inactivation of either myosin XI-2 or myosin XI-K resulted in an approximately 2-fold reduction in the mean velocity of the Golgi stacks and peroxisomes (Fig. 3D). In the case of mitochondria, the myosin XI-K knockout line exhibited more than 3-fold lower velocity than that in the wild type, whereas a less dramatic, approximately 1.5-fold velocity reduction was observed in the myosin XI-2 knockout line (Fig. 3F). In all cases, differences between the organelle velocities in knockout lines versus the parental Columbia line were statistically significant with \( P < 0.001 \). Taken together, these observations demonstrate that myosins XI-K and XI-2 each make a
significant contribution into the rapid transport of Golgi stacks, peroxisomes, and mitochondria in the root hairs. This, however, does not necessarily imply that the defects in root hair growth seen in the corresponding knockout lines can be directly attributed to the slower organelle movement.

To determine whether or not myosins XI-2 and XI-K are required for rapid organelle trafficking in organs other than root hairs, we examined motility of Golgi stacks and peroxisomes in the elongated epidermal cells found along the central vein on the leaf underside (Fig. 4B, central area). These elongated cells are most amenable to observing and measuring organelle movement in leaves. Interestingly, the mean velocity of these organelles in leaf cells was at least 30% greater than that in the root hairs (compare with Figs. 3D and 4D). Inactivation of either myosin XI-2 or myosin XI-K resulted in an approximately 3- to approximately 5-fold reduction of the mean velocity of Golgi stacks and peroxisomes (Fig. 4, A, C, and D), and also a stronger effect compared with that in the root hairs. The dominant negative inhibition of myosin XI-K closely mimicked the effect of the gene knockout, once again confirming the utility of this approach for the study of myosin function in plants (Fig. 4, A and D).

The mean velocity of mitochondria in the leaf cells was approximately 40% greater than that in the root hairs (compare with Figs. 3F and 4F). Interestingly, inactivation of the two myosin genes had distinct effects on the translocation of mitochondria: the myosin XI-2 gene knockout line showed only a moderate, although statistically significant ($P = 0.045$) reduction in the velocity of this organelle, whereas the myosin XI-K knockout line exhibited a drastic, 3.5-fold velocity reduction (Fig. 4, E and F).

Collectively, these results demonstrated that two class XI myosins, XI-2 and XI-K, make comparable contributions to rapid trafficking of Golgi stacks and peroxisomes in the Arabidopsis roots and leaves. It seems, however, that myosin XI-K plays a more significant role in the translocation of mitochondria than myosin XI-2 and that this difference is more pronounced in leaves than in root hairs.

**Paralogous Myosins XI-B and XI-1 Are Not Essential for Organelle Transport and Root Hair Growth**

Phylogenetic analysis of plant myosins presented in Figure 1A of an accompanying article (Avisar et al., 2008) shows that Arabidopsis myosins XI-2 and XI-K each possess closely related paralogs, myosins XI-B and XI-1, respectively, suggesting that the myosins in each paralogous pair may perform similar functions. To test this possibility, we examined root hair growth and trafficking of the mitochondria and peroxisomes in the knockout lines with inactivated myosin XI-B or XI-1. Our analyses revealed no defects in root hair growth in either of the mutant lines (Fig. 5A). Examination of mitochondria in the root hairs (Fig. 5, B and C) or leaf epidermal cells (Fig. 5, D and E) showed no significant changes in the mean organelle velocities due to mutational inactivation of either of two myosin

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**Figure 3.** Roles of myosins XI-2 and XI-K in organelle trafficking in root hairs. A, C, and E, Representative images of the indicated organelles (top rows) and paths of individual organelles plotted relative to a common origin (bottom rows; each axis is 100 µm). B, Images of the roots of parental and knockout lines transformed with the Golgi-specific GFP reporter. D, Mean velocities of the Golgi stacks and peroxisomes. F, Mean velocities of mitochondria.
paralogs. Only a modest reduction of the peroxisome velocity in leaf epidermal cells was observed in the myosin XI-1 knockout line compared with that in the parental Columbia line (Fig. 5, F and G).

It was also found that transgenic expression of the myosin XI-1/GTD (Fig. 5H) had only moderate negative effects on the root hair growth (Fig. 5A) or on peroxisome trafficking (Fig. 5, F and G). Taken together, these observations indicate that myosins XI-B and XI-1 might play only a relatively minor role in the root hair development and organelle translocation in the presence of intact paralogous myosins XI-2 and XI-K.

DISCUSSION

It is well established that actin cytoskeleton dynamics plays a paramount role in the polarized growth of the root hair cells (Hepler et al., 2001; Carol and Dolan,

Figure 4. Roles of myosins XI-2 and XI-K in organelle trafficking in leaves. A, C, and E, Representative images of the indicated organelles (top rows) and paths of individual organelles plotted relative to a common origin (bottom rows; each axis is 100 μm). B, Image of the leaf vein in the Columbia line transformed with the Golgi-specific GFP reporter showing a file of elongated epidermal cells used for organelle tracking. D, Mean velocities of the Golgi stacks and peroxisomes. F, Mean velocities of mitochondria.

Figure 5. Myosins XI-B and XI-1 do not play significant roles in root hair development and organelle movement. A, Analysis of the mean root hair length for the indicated gene knockout mutant lines and transgenic line that expresses XI-1/GTD. B, D and F, Paths of individual organelles plotted relative to a common origin (each axis is 100 μm). C and E, Mean velocities of mitochondria in root hairs and leaf epidermal cells, respectively. G, Mean velocities of the peroxisomes in leaf epidermal cells. H, Immunoblot analysis of the HA epitope-tagged XI-1/GTD expression using HA-specific monoclonal antibody. Samples were from a control plant (C, Columbia in the image) and plants representing three independent transgenic lines (1, 6, and 7). Positions of two size markers (molecular mass in kilodaltons) are shown by arrows.
pollen tubes. However, relatively low levels of myo-
involvement of myosins in the polarized growth of the
development of the root hairs and pollen (Hepler
myosin XI-K function.
proper cargo translocation in both knockout and dom-
both of these effects are likely due to the lack of the
transgenic expression of the cognate GTD mimics the
interference with cargo binding by myosin XI-K using

cell metabolism cannot be ruled out. However, because
more general processes required for maintenance of
motility of vesicles and organelles for the elongation
of the root hairs. In fact, only one class XI Arabidopsis
myosin, XI-K, was recently implicated in this process
(Ojangu et al., 2007), and even in this study, the
potential role of this myosin in organelle movement
has not been addressed.

Here we describe systematic screening of all 13 class
XI myosins of Arabidopsis for their potential functions
in the root hair development. Using gene knockout and
dominant negative inhibition approaches, we reveal
that two class XI myosins, XI-2 and XI-K, are essential
for the normal elongation of the root hairs. Inactivation
of each of these myosins in four distinct insertion lines
(Fig. 1) results in very similar phenotypes with a mean
root hair length of approximately one-third that in the
parental Columbia line (Fig. 2). In addition, we inves-
tigated the roles of myosins XI-2 and XI-K, and their
most closely related paralogs, myosins XI-B and XI-1, in
the trafficking of Golgi stacks, peroxisomes, and mito-
chondria in the root hairs. Conspiciously, we found
that myosins XI-2 and XI-K are each required for the
rapid movement of all three organelles (Fig. 3). Inactiva-
tion of the corresponding genes reduced the mean
velocity of organelles from 1.5- to 3-fold depending on the
organelle or myosin identity. In contrast, inactiva-
tion of the closely related, paralogous myosins XI-B and
XI-1, had no detectable effects on either root hair
growth or organelle motility (Fig. 5); additional work
is needed to identify functions of these myosins.

The observed correlation in the roles of myosins XI-2
and XI-K in root hair growth and organelle trafficking
suggests that the latter might be functionally required
for the former. Indeed, the rapid movement of the Golgi
stacks, peroxisomes, and mitochondria in a “reverse
fountain” manner (Hepler et al., 2001) may be needed
to enhance protein secretion and turnover of metabo-
lites and energy in the growing root hair tip. It also
seems possible that the vesicular trafficking that de-
livers building materials for the plasma membrane and
cell wall expansion in the tip is powered by myosins
XI-K and XI-F. Finally, the possibility that these myosins
are also involved in microfilament remodeling or even
more general processes required for maintenance of
cell metabolism cannot be ruled out. However, because
interference with cargo binding by myosin XI-K using
transgenic expression of the cognate GTD mimics the
phenotype of XI-K gene knockout in defective root hair
growth and slow organelle movement (Figs. 2 and 3),
both of these effects are likely due to the lack of the
proper cargo translocation in both knockout and dom-
inant negative inhibition approaches used to determine
myosin XI-K function.

Recent insight into the mechanistic parallelism in
the development of the root hairs and pollen (Hepler
et al., 2001; Cole and Fowler, 2006) implies potential
involvement of myosins in the polarized growth of the
pollen tubes. However, relatively low levels of myo-
sins XI-2 and XI-K in pollen (the myosin transcription
data were found at the Weigel World Web site; http://
www.weigelworld.org/resources) make these myo-
sins unlikely candidates for the major role in this
process. In contrast, myosins XI-A, XI-B, XI-C, XI-D,
and XI-J are almost exclusively expressed in pollen
and therefore are much better candidates for direct
involvement in pollen tube growth. Therefore, it seems
likely that the closest paralog of myosin XI-2, myosin
XI-B, has evolved to mediate organelle and vesicle
transport in the pollen tubes.

We found that myosins XI-2 and XI-K are required
for the rapid organelle trafficking not only in root hairs,
but also in the leaves (Fig. 4). This conclusion resonates
with the relatively high levels of these myosins in
leaves (http://www.weigelworld.org/resources). In
fact, myosins XI-2, XI-K, and XI-1 are the most abun-
dant myosins in the entire Arabidopsis plants. Surpris-
ingly, this abundance and major roles played by
myosins XI-2 and XI-K in organelle movement do not
translate into substantial defects in the development of
leaves, stems, or flowers in the corresponding gene
knockout lines. It appears that the reduction in the leaf
organelle velocity in these lines is not critical for leaf
development under normal growth conditions. There-
fore, perhaps due to their rapid elongation, root hairs
are a more sensitive indicator for the defects in organ-
elle trafficking than the leaf cells.

The lack of major developmental defects in two
myosin XI-2 knockout lines characterized here and in
an additional line described elsewhere (Hashimoto
et al., 2005) is in a stark contrast to the earlier article
that reported dwarf growth and flower sterility in a single
knockout line (Holweg and Nick, 2004). The reason for
this severe phenotype could be the second-
ary site mutation(s).

Rapid organelle trafficking is a hallmark of plant cell
physiology that is traditionally referred to as “cyto-
plasmic streaming” (Hepler et al., 2001; Smith and
Oppenheimer, 2005; Taiz and Ziegler, 2006; Shimmel,
2007). However, early studies on Golgi trafficking
(Boevink et al., 1998; Nebenfuhr et al., 1999) demon-
strated that individual stacks are moved independently
of each other by the actomyosin motility system rather
than being passively carried along by indiscriminate
cytosol flow (Nebenfuhr and Staehelin, 2001). Our
analyses of the trafficking of Golgi, peroxisomes, and
mitochondria in N. benthamiana (Avisar et al., 2008)
stressed the need to revise the concept of cytoplasmic
streaming. The patterns of organelle movement in
Arabidopsis (e.g. Fig. 4) are also incompatible with
passive “floating with the flow” and further support
introduction of a mechanistically more relevant con-
cept of active organelle translocation defined by the
actomyosin transport network.

On a broader scale, this work is relevant to the prob-
lem of the multigene families’ evolution in eukaryotes:
to what extent the lineage-specific expansion of these
families is due to adaptation as opposed to stochastic
gene birth and death processes (Lynch, 2007)? The myosin
gene families provide an excellent model to address this problem because they are nearly ubiquitous in eukaryotes. Our phylogenetic analysis (Avisar et al., 2008) suggested that the common ancestor of higher plants possessed five class XI myosins. This number has more than doubled in Arabidopsis, the plant that encodes more class XI myosins than the other three plants with completely sequenced genomes. The functional analysis of the Arabidopsis class XI myosins described here revealed a rather complex picture. The lack of major developmental defects in 11 myosin gene knockouts attests to a redundant nature of myosin functions in plants. The functional profiles of myosins XI-2 and XI-K that belong to distinct paralogous families largely overlap: both myosins are involved in the organelle transport and in root hair growth. The only functional specialization apparent among these Arabidopsis myosins is a somewhat larger contribution of myosin XI-K in the movement of mitochondria, an effect that is even more pronounced in N. benthamiana (Avisar et al., 2008). More extensive characterization of the myosin functions is required to determine how many and which myosins are essential for plant development.

MATERIALS AND METHODS

T-DNA Insertion Mutants

Seeds of Arabidopsis (Arabidopsis thaliana) ecotype Columbia T-DNA insertion lines

Seeds of Arabidopsis (Arabidopsis thaliana) ecotype Columbia T-DNA insertion lines SALK_055785 and SAIL_632_D12 (At5g43900; myosin XI-2), SALK_067972 and WiscDsLox17C12 (At5g20400; myosin XI-K), SALK_113062 (At1g04160; myosin XI-B), SALK_019031 (At1g17580; myosin XI-I), SALK_082070 (At1g29740; myosin XI-D), SALK_072023 (At1g54560; myosin XI-E), SALK_018032 (At2g20290; myosin XI-G), SAIL_365_D03 (At4g28710; myosin XI-H), SALK_082443 (At4g33200; myosin XI-I), and SALK_063159 (At5g38160; myosin XI-J) were acquired from the Arabidopsis Biological Resource Center (Alonso et al., 2003). Lines GABI_115C01 (At5g43900; myosin XI-2), GABI_622E02 (At1g04600; myosin XI-A), GABI_626B03 (At1g58700; myosin XI-C), and GABI_070F03 (At2g31900; myosin XI-F) were obtained from the European Arabidopsis Stock Center. All lines were selected and grown for 5 d on the same medium supplemented with 50 nM of Rhodamine 123 (Invitrogen). A series of eight consecutive images was acquired using a 510 Meta (Zeiss) confocal microscope and was used to measure the organelle velocities. Samples were excited using Argon laser at 488 nm, emission signal was collected through a band-pass 505–530-nm filter. The Golgi stacks and peroxisomes were observed using the following configurations of excitation and emission filters, respectively: 488 and 508 nm for GFP, 513 and 527 nm for yellow fluorescent protein, and 587 and 610 nm for mCherry. For time-lapse experiments, the consecutive images were taken at 1-s intervals for mitochondria or 2 s for Golgi and peroxisomes. For the root hair and leaf epidermis observations, more than 150 and 300 individual organelles, respectively, were traced. Tracking and measurements of velocities of individual organelles was performed using the Velocity3.7.0 Classification software (Improvision; Image Processing and Vision Company). Statistical analysis of the data was done using t test and Excel software. Additional details of organelle trafficking analyses are provided in an accompanying work (Avisar et al., 2008).

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