An isoform of *Arabidopsis* myosin XI interacts with small GTPases in its C-terminal tail region

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Abstract

Myosin XI, a class of myosins expressed in plants is believed to be responsible for cytoplasmic streaming and the translocation of organelles and vesicles. To gain further insight into the translocation of organelles and vesicles by myosin XI, an isoform of *Arabidopsis* myosin XI, MYA2 was chosen and its role in peroxisome targeting was examined. Using the yeast two-hybrid screening method, two small GTPases, AtRabD1 and AtRabC2a, were identified as factors that interact with the C-terminal tail region of MYA2. Both recombinant AtRabs tagged with His bound to the recombinant C-terminal tail region of MYA2 tagged with GST in a GTP-dependent manner. Furthermore, AtRabC2a was localized on peroxisomes, when its CFP-tagged form was expressed transiently in protoplasts prepared from *Arabidopsis* leaf tissue. It is suggested that MYA2 targets the peroxisome through an interaction with AtRabC2a.

Key words: AtRabC2a, AtRabD1, myosin XI, MYA2, peroxisome.

Introduction

Myosin is a molecular motor protein, which acts on actin filaments. It is composed of light and heavy chains which are divided into head, neck, and tail regions. The head region binds to actin filaments and moves along them, dependent upon ATP hydrolysis. On the basis of the primary structure of this region, myosin is classified into at least 24 classes (Foth et al., 2006). Light chains, in many cases calmodulin, bind to the neck region. The C-terminal tail region is diverse in size and primary structure among myosin classes, and is inferred to determine the physiological function of each class of myosin in cells.

Only three classes of myosin, VIII, XI, and XIII, have been identified in plant cells (Reichelt and Kendrick-Jones, 2000; Sellers, 2000). In *Arabidopsis thaliana*, 17 myosins have been identified by searching the database. Among them, four belong to class VIII and 13 to class XI (Reichelt and Kendrick-Jones, 2000; Reddy and Day, 2001). Pharmacological studies, using actin-depolymerizing drugs and myosin inhibitors, have demonstrated that most organelles, including endoplasmic reticulum (E Yokota, S Ueda, K Tamura, H Orii, S Uchi, S Sonobe, I Harashima, and T Shimmen, unpublished data), mitochondria (Logan and Leaver, 2000; Van Gestel et al., 2002), chloroplasts (Wada et al., 2003; Paves and Truve, 2007), plastids (Kwok and Hanson, 2003), peroxisomes (Jedd and Chua, 2002; Mano et al., 2002; Mathur et al., 2002; Collings et al., 2003), vacuoles (Higaki et al., 2006), and Golgi (Boevink et al., 1998; Nebenfuhr et al., 1999) move, or are translocated, using the actin–myosin system. Among these organelles, mitochondria and plastids in maize root cells (Wang and Pesacreta, 2004), mitochondria in tobacco pollen tubes (Romagnoli et al., 2007) and endoplasmic reticulum in tobacco cultured BY-2 cells (E Yokota, S Ueda, K Tamura, H Orii, S Uchi, S Sonobe,
I Hara-Nishimura, and T Shimmen, unpublished data), immunostained with an antibody against myosin XI. It has previously been shown that an isoform of myosin XI, MYA2, was co-localized with peroxisomes and, thus far, unidentified vesicles in Arabidopsis by immunostaining with a specific antibody against MYA2 tail region (Hashimoto et al., 2005). Furthermore, the transport and movement of Golgi, mitochondria, and peroxisomes were suppressed or modified by the knockout of certain movement of Golgi, mitochondria, and peroxisomes were (Reisen and Hanson, 2006), and myosin XI isoforms was expressed in tobacco, onion (Reins and Hanson, 2006), and Arabidopsis cells (Li and Nebenfuhr, 2007), Golgi and peroxisomes were tagged. In some cases, the movement of these organelles and mitochondria was reported to be suppressed or modified by the expression of the tail region (Avisar et al., 2008; Peremyslov et al., 2008). These results suggest that myosin XI is associated with organelles through its tail region. The tail regions of myosin XI isoforms are divided into two short subdomains, with organelles associating with either of them independently (Li and Nebenfuhr, 2007). Interestingly, a full-length tail or a long tail isoform, containing the coiled-coil domain of myosin XI, does not target specific organelles or bind to unknown organelles. Based on this evidence, it was suggested that the organelle targeting of each myosin XI isoform is selected and regulated in the cells, and it appears likely that myosin XI isoforms have a mechanism for recognizing and selecting organelles in their tails.

In animal and yeast cells, it is known that small GTPases, Rab proteins, interact with class V myosin, which is functionally and structurally similar to myosin XI, and are important in organellar and vesicular targeting. In budding yeast, a Rab protein, Ypt11p, functions in mitochondrial distribution into the developing bud through the interaction with the tail region of myosin V, Myo2p (Itoh et al., 2002). In mammalian cells, myosin Vb regulates the plasma membrane recycling system through its association with Rab11a and Rab11-family-interacting protein-2 (Rab11-FIP2; Lapierre et al., 2001; Hales et al., 2002). Furthermore, the formation of a complex between Rab27a, melanophilin and myosin Va, is required for normal melanosome transport in the melanosphere (Storm et al., 2002; Kuroda et al., 2003).

In the present study, small GTPase Rabs, AtRabD1 and AtRabC2a, were identified as interacting factors with the tail region of MYA2 by yeast two-hybrid screening. An in vitro binding assay showed that the tail region of MYA2 binds to these AtRab proteins in a GTP-dependent manner. Furthermore, AtRabC2a was localized on peroxisomes in protoplasts prepared from Arabidopsis leaf tissue. These results suggested that MYA2 is targeted to peroxisomes through an interaction with AtRabC2a.

Materials and methods

Yeast two-hybrid screening

Truncated MYA2 tails were amplified by RT-PCR with total RNA isolated from leaves of 4-week-old plants by a RNeasy Plant Kit (Qiagen, Valencia, CA, USA). The PCR amplification products were subcloned into pENT/D/TOPO vector (Invitrogen, Carlsbad, CA), and then integrated into a pBD-GAL4-GWRF (Stratagene, CA Lolla, CA) by LR clone (Invitrogen). A yeast AH109 strain (Clontech) was co-transfected with MYA2(aa1049–aa1505)/pBD-GAL4-GWRF and 300 μg of an Arabidopsis root cDNA plasmid library which was constructed in pAD-GAL4-GWRF, derived from a pAD-GAL4-2.1 vector (Stratagene), and plated on the selection media containing 5 mM 3-aminotriazole but lacking leucine, tryptophan, and histidine. After 2 weeks of growth, positive colonies were picked up and regrown on the selection media. Significant growth on the selection media indicated a positive interaction. The plasmids were then recovered from the colonies using a Zymoprep Kit (Zymo Research, Orange, CA) and introduced into an E. coli TOP10 strain (Invitrogen) with selection by ampicillin. A DNA sequencer (ABI PRISM 3000) was used for sequence analysis. For the examination of Rab binding sites in the MYA2 tail, the yeast AH109 strain was co-transfected with Rab/pAD-GAL4-GWRF and one of the various truncated MYA2/pBD-GAL4-GWRF, then plated on the control media. After 4 d growth, the colonies were regrown on the selection and control media. To confirm no positive interactions by self-activation, the co-transfection with MYA2 tail bait and empty prey vectors or Rab prey and empty bait vectors were performed. In addition, co-transfection with pBD-wt and pAD-wt (Stratagene) or empty bait and empty prey vectors were performed for the control experiments.

Isolation of recombinant proteins

MYA2 tail (aa1049–aa1505 or aa1049–aa1450) in a pENT/D-TOPO vector was amplified using a forward primer with EcoR1 and a reverse primer with Xho1 and cloned into a pGEX4T-1 vector (Amersham) using their restriction sites by a DNA ligation kit ver.2.1 (TaKaRa). The full-length cDNAs of AtRabD1 and AtRabC2a were also amplified by RT-PCR with total RNA isolated from leaves of 4-week-old plants using an RNeasy Plant Kit. PCR amplification products were subcloned into the pENT/D/TOPO vector (Invitrogen), and then transformed into a pDEST17 vector (Invitrogen). After 4 d growth, the colonies were regrown on the selection media containing 5 mM 3-aminotriazole but lacking leucine, tryptophan, and histidine. After 2 weeks of growth, positive colonies were picked up and regrown on the selection media. Significant growth on the selection media indicated a positive interaction. The plasmids were then recovered from the colonies using a Zymoprep Kit (Zymo Research, Orange, CA) and introduced into a TOP10 strain (Invitrogen) with selection by ampicillin. A DNA sequencer (ABI PRISM 3000) was used for sequence analysis. For the examination of Rab binding sites in the MYA2 tail, the yeast AH109 strain was co-transfected with Rab/pAD-GAL4-GWRF and one of the various truncated MYA2/pBD-GAL4-GWRF, then plated on the control media. After 4 d growth, the colonies were regrown on the selection and control media. To confirm no positive interactions by self-activation, the co-transfection with MYA2 tail bait and empty prey vectors or Rab prey and empty bait vectors were performed. In addition, co-transfection with pBD-wt and pAD-wt (Stratagene) or empty bait and empty prey vectors were performed for the control experiments.

In vitro binding assay

The same batch of beads coated with MYA2 tails was used to analyse the binding of each His6-taged Rab protein. 39 μl of each
Rab protein (30 µg ml⁻¹) and 1 µl of 80 mM GTP or deionized water was added to 40 µl of the suspension of beads (15 µl of void volume) in a microcentrifuge tube. The mixture was gently suspended at 3 min intervals for 20 min at room temperature. After centrifugation at 500 rpm at room temperature, the supernatant was carefully discarded, and 65 µl of NEMP solution was added to the pellet. During washing, 300 µl of NEMP solution with or without 1 mM GTP was added to the pellet. After centrifugation, one aliquot of pellet was pooled, while a second aliquot of pellet was washed with 300 µl of NEMP solution and then centrifuged. Each pellet was subjected to SDS-PAGE using a 12% gel by the method of Laemmli (1970). After staining with Coomassie brilliant blue (CBB), the intensity of each band of Rab proteins was measured using a densitometer (Densit-o-pattern analyser model no. EPA-3000, Cosmo Bio. Co. Ltd., Tokyo).

Leaf protoplast preparation

A transgenic Arabidopsis expressing the GFP-tagged peroxisomal targeting signal peptide 1 (GFP-PTS1; Mano et al., 2002) was used for examining positional and distributional relationships between peroxisomes and AtRab proteins. Arabidopsis seedlings were grown on Murashige and Skoog plates at 22 °C. Rosette leaves of 2-week-old plants were cut into 1–2 cm sections with a razor blade. Samples were digested for about 3 h in an enzyme solution [1.5% cellulase R-10 (Onozuka, Tokyo, Japan), 0.4% macerozyme R10 (Yakult, Tokyo, Japan), 0.4 M mannitol, 20 mM KCl, 10 mM CaCl₂, 0.1% BSA, and 20 mM MES (pH 5.7)] whilst being shaken at 40 rpm at 23 °C. The mixture was filtered with a 50 µm nylon mesh. After centrifugation at 100 g for 3 min, the pellet was suspended in a solution containing 0.4 M mannitol, 15 mM MgCl₂ and 4 mM MES (pH 5.7).

Transient expression in Arabidopsis leaf protoplasts

A full-length cDNA of AtRabC2a or AtRabD1 was fused to the C-terminus of CFP controlled by the 35S CaMV promoter by GATEWAY technology. The construct was introduced into protoplasts by a PEG-mediated transformation method (Sheen, 2002; http://genetics.mgh.harvard.edu/sheenweb/). Protoplasts expressing the fusion protein were observed under an Olympus BX60 fluorescence microscope equipped with filters for GFP (U-MNIBA, Olympus) and CFP fluorescence (U-MCPFHQ, Olympus). Images were taken with a digital camera (DP 70, Olympus) controlled by Aquacosmos software (Hamamatsu Photonics). Unmixed images with the two types of fluorescence were subsequently acquired by processing with U9677 Multiband Imaging software (Hamamatsu Photonics).

Results

Identification of AtRab proteins as interacting factors with the C-terminal tail region of MYA2

To investigate the function of the MYA2 tail, potential factors that interact with the C-terminal tail region of MYA2 were searched using a two-hybrid screening method. An Arabidopsis root cell cDNA library with the tail region of MYA2, aa1049–aa1505 (f1), lacking the coiled-coil domain, as shown schematically in Fig. 1A was screened. After screening 1.2×10⁶ clones, DNA fragments encoding AtRabD1 and AtRabC2a, members of the small GTPase Rab family, were identified. To determine the region in the MYA2 tail responsible for the interaction with each AtRab, their interaction with a series of deletion forms of the MYA2 tail, aa1049–aa1450 (f2 in Fig. 1A), aa900–aa1221 (f3 in Fig. 1A), in which the coiled-coil domain was included, and aa1195–aa1505 (f4 in Fig. 1A) was examined. The association of each AtRab to each truncation of the MYA2 tail is summarized in Fig. 1B. AtRabD1 interacted with a truncation of MYA2 tail, aa1049–aa1450 (f2) lacking 55 amino acids in the C-terminus, but not with aa900–aa1221 (f3) containing only the N-terminal region, and aa1195–aa1505 (f4), lacking the N-terminus. These results suggest that the boundary region between aa1195 and aa1221 in the MYA2 tail is important for the interaction with AtRabD1. By contrast, AtRabC2a interacted only with a full-length MYA2 tail (f1), but not with its truncated forms, where 55 amino acids from the C-terminus or 146 amino acids from N-terminus were deleted. Therefore, both tail termini appear to be necessary for the interaction with AtRabC2a.

Fig. 1. Interaction of AtRabD1 and AtRabC2a with truncated MYA2 tails in the yeast two-hybrid method. (A) A diagram of the tail regions of MYA2, fragment 1 to 4 (f1 to f4), tested for the interaction with AtRabD1 and AtRabC2a. (B) Interaction of AtRabD1 or AtRabC2a with each truncated form of the MYA2 tail (f1 to f4); + or – indicates a negative or positive interaction, respectively. (C) Photographs of yeast co-transfected with truncations of MYA2 tail (MYA2 f1 (c), MYA2 f2 (d) MYA2 f3 (e), MYA2 f4 (f)) as bait and AtRabD1 (upper) or AtRabC2a (bottom) as prey on the selection media. Negative or positive control interaction is (a) or (b), respectively. Significant growth indicated a positive interaction.
Interaction of recombinant MYA2 tail proteins with recombinant AtRabD1 and AtRabC2a in vitro

Like other regulatory GTPases, the Rab protein interactions between the active GTP-bound and inactive GDP-bound forms, each of which have different conformations from the other (Ali and Seabra, 2005). The GTP-bound form of the Rab protein interacts with effector proteins rendering it catalytically active. It was examined whether AtRabD1 and AtRabC2a interact with MYA2 tails at the protein level, and whether any interaction is GDP-dependent. AtRabD1 (lane a in Fig. 2A) and AtRabC2a (lane b in Fig. 2A) fused with His6, and MYA2 tails, aa1049–aa1505 (MYA2 tail 1; lane c in Fig. 2A) and aa1049–aa1450 (MYA2 tail 2; lane d in Fig. 2A), fused with GST, were expressed in *E. coli* and purified by affinity chromatography as described in the Materials and methods. Each MYA2-tail recombinant protein fused with GST was bound to glutathione Sepharose beads, and was used for an *in vitro* binding assay using a pull down assay. Beads without the MYA2 tail bound to them were used as a control.

First, the interaction of both recombinant AtRab proteins with the MYA2 tail 1 was examined. In the absence of GTP (lane b in Fig. 2B), the amount of both His6-AtRab proteins, which recovered in pellets with beads coated with the MYA2 tail 1, appeared to be same as that recovered with control beads (lane a in Fig. 2B). However, the amount of both AtRab proteins in pellets with MYA2 tail 1 coated beads was increased by the addition of GTP (lane c in Fig. 2B). Densitometric band analysis of AtRab protein in the pellet showed that the amount of AtRabD1 (white bars in Fig. 3A) and AtRabC2a protein (shaded bars in Fig. 3A) bound to the MYA2 tail 1 coated beads in the presence of GTP (+GTP in Fig. 3A) was 2.6-fold and 1.7-fold, respectively, larger than that in the absence of GTP (−GTP in Fig. 3A) or that bound to the control beads in the presence of GTP, indicating that both recombinant AtRab proteins had an ability to bind to the MYA2 tail 1 in a GTP-dependent manner and that the affinity of AtRabD1 for the MYA2 tail 1 was higher than that of AtRabC2a.

Next, the interaction of both AtRab proteins with the MYA2 tail 2 (Fig. 4A) was examined. The amount of AtRabD1 protein (white bars in Fig. 4A) bound to MYA2 tail 2 coated beads in the presence of GTP (+GTP in Fig. 4A) was 1.6-fold larger than that in the absence of GTP (−GTP in Fig. 4A) or that bound to the control beads in the presence of GTP, indicating that this AtRab protein was also able to associate with the MYA2 tail 2 in a GTP-dependent fashion. However, the amount of AtRabC2a protein (shaded bar in Fig. 4A) bound to the beads coated with the MYA2 tail 2 in the presence of GTP (+GTP in Fig. 4A) was similar to that in the absence of GTP (−GTP in Fig. 4A) or that bound to the control beads. Thus, AtRabC2a protein did not associate with the MYA2 tail 2 even in the presence of GTP, supporting the results of the two-hybrid analysis.

However, these recombinant AtRab proteins readily dissociated from the MYA2 tail 1 (Fig. 3B; white bars and shaded bars for AtRabD1 protein and AtRabC2a protein, respectively) or tail 2 (Fig. 4B) adsorbed to beads by washing with a solution containing GTP. These results indicate that AtRabD1 and AtRabC2a protein tagged with His6 interacted weakly with the MYA2 tails fused with GST in a GTP dependent manner.

Co-localization of transiently expressed AtRabC2a with peroxisomes

Next, the intracellular localization of each AtRab protein was examined in relation to peroxisomes in *Arabidopsis* cells, in which peroxisomal targeting signal 1 (PTS1) tagged with GFP was stably expressed (Mano et al., 2002). Since the primary structures of AtRab proteins show high levels of homology with isoforms in subgroups (A and B for AtRab isoforms belonging to C and D subgroups, respectively, in Fig. 5), it was considered that it was not feasible to make an antibody against AtRabC2a or AtRabD1. Hence, the localization of AtRabD1 or AtRabC2a protein was evaluated by the transient expression of their CFP-fused forms in leaf cell protoplasts. Dot signals of CFP-AtRabC2a (Fig. 6B) generally overlapped with GFP-tagged PTS1 signals, peroxisomes (Fig. 6A). When a protoplast not transformed with CFP-tagged AtRabC2a (Fig. 6C) was
observed through a CFP filter, few signals were detected (Fig. 6D). Conversely, diffuse signals of CFP-tagged AtRabD1 (Fig. 6F, H) were observed throughout the cytoplasm, and peroxisomes appeared to be embedded with these signals. In some cases, the GFP-labelled peroxisome signals did not overlap with those of AtRabD1 (white arrows in Fig. 6H). However, at present, it is not possible to determine the organelles targeted by AtRabD1.
In the present study, small GTPase Rabs, AtRabD1 and AtRabC2a, were identified as interacting factors with the MYA2 tail using the yeast two-hybrid screening method. Both recombinant AtRab proteins interacted with recombinant MYA2 tail protein in a GTP-dependent manner. However, the regions in the MYA2 tail important for the interaction with each AtRab were different between isoforms. Furthermore, it was found that CFP-tagged AtRabC2a expressed transiently in Arabidopsis leaf cells was co-localized with the peroxisome. Based on the evidence and results, it is suggested that the MYA2 is targeted to peroxisomes through the interaction with AtRabC2a.
In animal cells and yeast, Rab proteins play a role as an adaptor/receptor for myosin V, and are involved in targeting myosin V to the proper organelles and vesicles. For example, Myo2p of budding yeast (Itoh et al., 2002), myosin Vb (Lapierre et al., 2001; Hales et al., 2002), and myosin Va (Storm et al., 2002; Kuroda et al., 2003) of mammalian cells interact with the Rab protein, Ypt11p, Rab11 family proteins, and Rab27a, respectively, either directly or indirectly with other proteins to exert functions such as mitochondrial distribution, plasma membrane recycling, and normal melanosomal transport. Recently, ATM1, a plant specific myosin VIII isoform, was shown to be co-localized with vesicles decorated with GFP-tagged Ara6 (Golomb et al., 2008), which is a plant Rab5 orthologue and regulates endocytosis in Arabidopsis cells (Ueda et al., 2001, 2004). Although the direct interaction of ATM1 with Ara6 in vivo or in vitro has not been examined, it was attractive to consider that myosin VIII is also associated with the AtRab protein, Ara6, on endocytotic vesicles.

Each AtRab had different MYA2 tail regions necessary for their interactions; AtRabC2a for both N- and C-termini in the tail. In a similar way in which the MYA2 tail interacts with AtRabC2a, mammalian myosin Vb tails interact with Rab11 family proteins, such as Rab11a, Rab11b, and Rab25 (Lapierre et al., 2001). The tail of mammalian myosin Vb, in which the coiled-coil domain is not contained, is able to interact directly with Rab11 family proteins. Using the yeast two-hybrid screening method and its truncated forms, two interaction sites in the myosin Vb tail are indicated. One site is Leu-Glu-Lys-Asn-Glu (LEKNE; N-site) located in the N-terminus (aa1421–aa1425), and the other is Leu-Leu-Leu-Asp (LLLID; C-site) in the C-terminus (aa1849–aa1852). A similar sequence to the N-site was not found in the N-terminal region of the MYA2 tail (data not shown). Surprisingly, however, Phe-Lue-Lue-Asp (FLLD) corresponding to LLLD of the C-site was present within 55 amino acids in the C-terminus of not only the MYA2 tail, but also other Arabidopsis myosin XI isoforms (Fig. 7), with the
exception of myosin XI-G and XI-J. In the case of XI-G, the last amino acid Asp in the four amino acid sequence was substituted with Lys (K). This conserved sequence was considered to play an important role in the interaction of myosin XI with AtRab proteins. The sequence was not included in a dilute domain (arrow in Fig. 7), which had been deduced to be necessary for cargo binding of myosin V (Reck-Peterson et al., 2000), but whose exact role has not yet been clarified.

The importance of both N- and C-termini of the MYA2 tails for interacting with AtRabC2a has been indicated, and it is suggested that MYA2 is targeted to the peroxisome through this interaction. However, a subdomain 2GT2, a C-terminal half spanning from aa1283 to aa1505 in the MYA2 tail, possessed a binding ability to the peroxisome (Li and Nebenfuhr, 2007). This result was contradictory to our suggestion, because the deleted fragment f4 (aa1195–aa1505) of the MYA2 tail containing the 2GT2 subdomain did not interact with AtRabC2a in the yeast two-hybrid screening assay. At present, this discrepancy cannot be explained. The subdomain 2GT2 might be targeted to the peroxisome without interacting with AtRabC2a, but by other mechanisms. Li and Nebenfuhr (2007) reported that a full-length MYA2 tail (MYA2GT, aa1091–aa1505) containing both 2GT1, a N-terminal half, and 2GT2 subdomains could not be targeted to the peroxisome or unknown organelles, suggesting that targeting and binding to organelles of myosin XI is tightly regulated in the cells. Conversely, Reisen and Hanson (2006) found that a MYA2 (myosin XI-6) tail (no coil, aa1053–aa1505) is targeted to the peroxisome when it was expressed in tobacco leaves. It is suggested that the N-terminus, aa1049–aa1195 in the MYA2 tail is necessary for interacting with AtRabC2a. Non-targeted MYA2 tail (MYA2GT) to the peroxisome started from aa1091, while the targeted tail (no coil) started from aa1053. This raises the possibility that the interacting site in the N-terminus of the MYA2 tail with AtRabC2a locates between aa1053 and aa1091, and that MYA2 tails with the site are able to interact with the peroxisome.

In the pull-down assay, the GTP-dependent interaction of both AtRab proteins with MYA2 tails in vitro appeared to be weak. The AtRab proteins bound to MYA2 tails were gradually dissociated from them by washing. The binding intensity or affinity of myosin V to the Rab protein has not been evaluated, because the interaction was examined by the yeast two-hybrid screening or an immunoprecipitation assay using cell lysate. In the latter case, there was a possibility that the interaction of myosin V with Rab protein is modified by other factors. Therefore, the intensity of interaction of MYA2 tail with AtRab proteins could not be compared to that of the myosin V tail with Rab proteins. As described above, Li and Nebenfuhr (2007) demonstrated the complicated behaviour of the MYA2 tail and its fragments having different domain lengths and suggested the presence of a mechanism for selection of the organelles which MYA2 binds to in the cells. Other myosin XI isoforms, such as MYA1, myosin XI-I, and myosin XI-K, showed similar behavior to MYA2. Li and Nebenfuhr (2007) proposed the following model for explaining the selection mechanism of myosin XI. Subdomains, GT1 and GT2 can take on dynamically alternating conformations in the myosin XI tail. Initially, the tail binds to or interacts weakly with an organelle-specific adaptor, and selects one of these conformations, and then is locked effectively on to the organelle. According to this model, the weak interaction of the tail with the adaptor is one of the key events. Although it was not confirmed whether or not the weak interaction of MYA2 with AtRabC2a occurs in cells, our data might be consistent with and support the model. Alternatively, the possibility was considered that other factor(s), such as Rab11-FIP2 for the interaction of Rab11a with myosin Vb in mammalian cells (Hales et al., 2002), further regulates and modulates the interaction of AtRab proteins with the MYA2 tail. Hence, further study is needed to elucidate and confirm these points and possibilities, in addition to clarifying the location and function of AtRabD1.

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