

Analysis of the Small GTPase Gene Superfamily of Arabidopsis¹

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Small GTP-binding proteins regulate diverse processes in eukaryotic cells such as signal transduction, cell proliferation, cytoskeletal organization, and intracellular membrane trafficking. These proteins function as molecular switches that cycle between "active" and "inactive" states, and this cycle is linked to the binding and hydrolysis of GTP. The Arabidopsis genome contains 93 genes that encode small GTP-binding protein homologs. Phylogenetic analysis of these genes shows that plants contain Rab, Rho, Arf, and Ran GTPases, but no Ras GTPases. We have assembled complete lists of these small GTPases families, as well as accessory proteins that control their activity, and review what is known of the functions of individual members of these families in Arabidopsis. We also discuss the possible roles of these GTPases in relation to their similarity to orthologs with known functions and localizations in yeast and/or animal systems.

Small GTP-binding proteins are molecular switches that are "activated" by GTP and "inactivated" by the hydrolysis of GTP to GDP. The resulting cycles of binding and hydrolysis of GTP by small GTP-binding proteins represents a ubiquitous regulatory mechanism in eukaryotic cells. Members of this class of proteins are among the largest families of signaling proteins in eukaryotic cells. Their importance in cellular signaling processes is underscored by their conservation throughout evolution of eukaryotic organisms and by the presence of homologs that perform related functions in cells of yeasts, humans, and plants. Although the GTP-hydrolysis "core" of this class of regulatory molecules is highly conserved, the surrounding domains are highly variable and undergo conformational changes as these proteins switch from GTP-associated to GDP-associated states. Eukaryotes have harnessed this diversity of protein conformations, linked to the nucleotide-associated state of the GTP-binding domain, to regulate a myriad of cellular processes (for review, see Takai et al., 2001). Small GTP-binding proteins are involved in regulation of diverse eukaryotic cellular processes, such as cell proliferation, cytoskeletal assembly and organization, and intracellular membrane trafficking (for reviews, see Barbacid, 1987; Boguski and McCormick, 1993; Takai et al., 2001). Physiological control of these GTPase "switches" occurs through association of the

GTPase with accessory proteins, termed guanine nucleotide exchange factors (GEFs), that catalyze the conversion of the small GTP-binding protein to their GTP-bound "active" conformation. In their "active" state, small GTPases interact with various downstream "effector" proteins that perform the diverse cellular functions controlled by this class of regulatory molecules. Inactivation occurs through either the intrinsic ability of the small GTP-binding protein to hydrolyze GTP to GDP+Pi, or through association with another set of accessory proteins, GTPase-activating proteins (GAPs), which stimulate this hydrolytic activity. Upon hydrolysis of GTP, the small GTP-binding protein is returned to the "inactive" state and is ready to begin the cycle again (supplemental data can be viewed at www.plantphysiol.org).

Structural and functional similarities between different members of this large superfamily has led to establishment of five distinct families: Ras, Rab, Rho, Arf, and Ran (Kahn et al., 1992). Ras GTPases regulate cell proliferation in yeast and mammalian systems. Members of the Rho GTPase family control actin reorganization and signal transduction pathways associated with MAP kinases. The Rab and Arf GTPase families function in distinct steps of membrane trafficking, whereas Ras-related nuclear protein (Ran) GTPases regulate transport of proteins and RNA across the nuclear envelope. Individual members of these families share higher overall sequence conservation with one another than with any other small GTPase families (Fig. 1). Analysis of the genomes of *Saccharomyces cerevisiae*, fruitfly (*Drosophila melanogaster*), *Caenorhabditis elegans*, Arabidopsis, and human (*Homo sapiens*) has underscored the conservation of these classes of regulatory molecules and has provided interesting glimpses into the ways in which the

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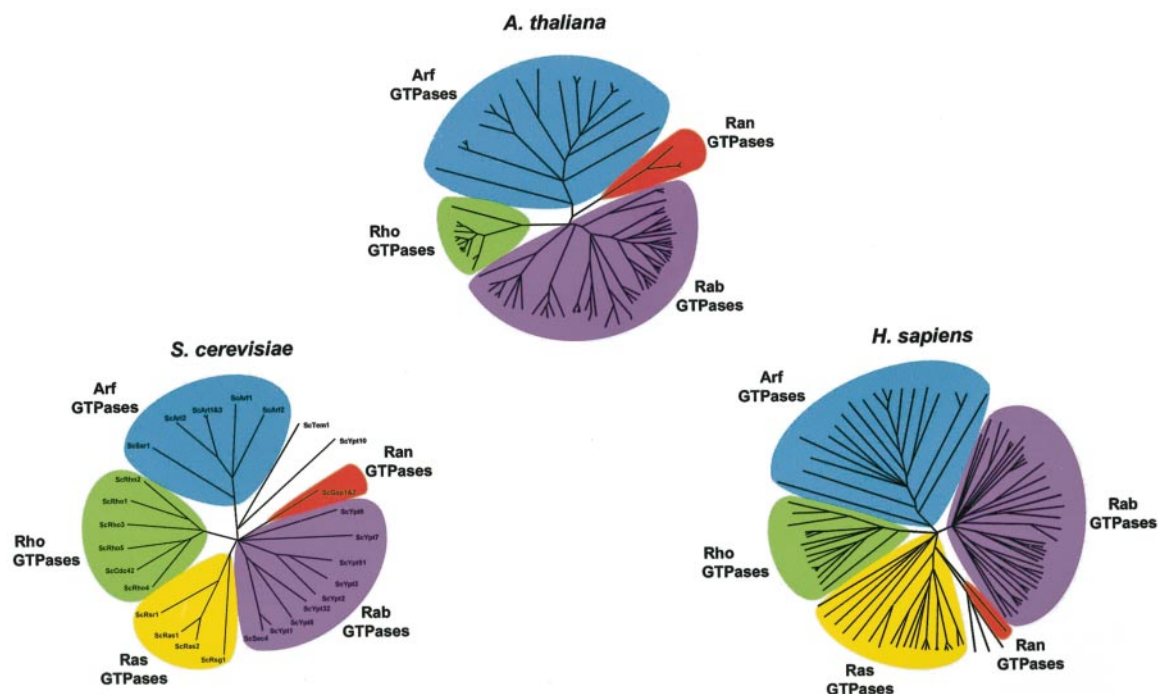


Figure 1. Phylogenetic star diagrams of small GTPases of *A. thaliana*, *S. cerevisiae*, and *H. sapiens*. Unrooted star diagrams were obtained using the program, ClustalW (Thompson et al., 1994). Sequences of small GTPases were obtained from the sequenced genomes of the three indicated organisms using BLAST (Altschul et al., 1997) sequence similarity searches against characterized GTPases. In each organism, distinct families (Ras, Rab, Rho, Arf, and Ran) could be distinguished, except for Arabidopsis in which no apparent Ras GTPase family could be identified.

small GTPases of these families have evolved and proliferated. Here, we describe the identification and classification of 93 small GTP-binding proteins in Arabidopsis. These GTPases were classified within four of the five small GTPase families: with 57 Rab GTPases; 21 Arf GTPases; 11 Rho GTPases; and 4 Ran GTPases. Interestingly, Arabidopsis does not contain any Ras GTPases that can be identified based on phylogenetic analysis, perhaps reflecting unique mechanisms for control of cell signaling during development in plants (Meyerowitz, 1999, 2002).

RESULTS

Identification and Classification of Small GTPase Genes in Arabidopsis

To identify small GTPase genes in the Arabidopsis genome, we first compiled sequences previously documented in *S. cerevisiae* and humans as members of the small GTP-binding protein superfamily of regulatory proteins. These genes were classified into the five accepted families of small GTPases (Kahn et al., 1992) based on their functions and were then used to generate a template phylogenetic tree for future grouping of Arabidopsis sequences. We then carried out BLASTP (Altschul et al., 1997) searches on the theoretical protein complement of the Arabidopsis Genome Initiative (AGI) database (Arabidopsis Ge-

nome Initiative, 2000) with representative sequences of each of the five families of small GTPases (Kahn et al., 1992). After individual protein sequences were identified, we ensured that we had eliminated duplications and contaminating sequences through careful comparison with theoretical cDNA and genomic DNA sequences in the AGI genomic database. In this manner, we identified a total of 93 small GTPase genes in the Arabidopsis genome.

To classify Arabidopsis small GTPase genes into Ras, Rho, Rab, Arf, and Ran families, theoretical protein sequences from these genes were aligned to previously compiled lists of small GTPases of *S. cerevisiae* and humans using ClustalW (Thompson et al., 1994) and then edited manually to restrict phylogenetic analysis to conserved "core" domains of these protein sequences. Phylogenetic trees were constructed using the neighbor-joining method as implemented in PAUP* with 1,000 bootstrap replicates (Swofford et al., 2001). Branches with bootstrap values of less than 80% were collapsed to simplify tree structures. After this phylogenetic analysis, we determined that Arabidopsis contained members of the Rab (57 members), Arf (21 members), Rho (11 members), and Ran (4 members) families, but no small GTPase genes could be determined to cosegregate with yeast or human members of the Ras GTPase family (Tables I–IV; Fig. 1).

Table 1. *Arabidopsis* RAB GTPase genes

a, Nomenclature used in Periera-leal and Seabra (2001). b, Nomenclature used in Bischoff et al. (1999). c, Nomenclatures used in: i, Anai et al. (1991) and Ueda et al. (1996); ii, Yi and Gueriot (1994); iii, Biermann et al. (1996); iv, Ueda et al. (2001); and v, Terryn et al. (1993).

Genes Name			AGI Gene ^a	Accession No. ^b	Expression/Localization ^c
a	b	c			
<i>AtRABA1a</i>	<i>AtRab11E</i>	<i>Ara-2</i> ⁱ	At1g06400	114086	
<i>AtRABA1b</i>			At1g16920	3024526	
<i>AtRABA1c</i>			At5g45750	9758678	
<i>AtRABA1d</i>	<i>AtRab11B</i>	<i>AthSGBP</i> ⁱⁱ	At4g18800	7438436	Highly expressed in roots ¹
<i>AtRABA1e</i>			At4g18430	7438435	
<i>AtRABA1f</i>			At5g60860	10176913	
<i>AtRABA1g</i>			At3g15060	8777489	
<i>AtRABA1h</i>			At2g33870	1707014	
<i>AtRABA1i</i>			At1g28550	6560749	
<i>AtRABA2a</i>	<i>AtRab11C</i>		At1g09630	3024516	
<i>AtRABA2b</i>			At1g07410	8778562	
<i>AtRABA2c</i>	<i>AtRab11A</i>		At3g46830	3915842	
<i>AtRABA2d</i>			At5g59150	9759237	
<i>AtRABA3</i>			At1g01200	9665141	
<i>AtRABA4a</i>			At5g65270	10178189	
<i>AtRABA4b</i>	<i>AtRab11G</i>	<i>AtGB3</i> ⁱⁱⁱ	At4g39990	7438426	
<i>AtRABA4c</i>			At5g47960	9758522	
<i>AtRABA4d</i>			At3g12160	9294115	
<i>AtRABA4e</i>			At2g22390	7438435	
<i>AtRABA5a</i>			At5g47520	9758780	
<i>AtRABA5b</i>			At3g07410	6041856	
<i>AtRABA5c</i>	<i>AtRab11F</i>	<i>Ara-4</i> ⁱ	At2g43130	114089	Ubiquitous expression/Golgi-derived vesicles ²
<i>AtRABA5d</i>			At2g31680	4582469	
<i>AtRABA5e</i>	<i>AtRab11D</i>	<i>Ara-1</i> ⁱ	At1g05810	114085	
<i>AtRABA6a</i>			At1g73640	6692728	
<i>AtRABA6b</i>			At1g18200	9719734	
<i>AtRABB1a</i>	<i>AtRab2B</i>		At4g17160	7438380	
<i>AtRABB1b</i>	<i>AtRab2C</i>	<i>AtGB2</i> ⁱⁱⁱ	At4g35860	2129702	
<i>AtRABB1c</i>	<i>AtRab2A</i>		At4g17170	1765895	High expression in cotyledons, fruits, and pollen ³
<i>AtRABC1</i>	<i>AtRab18</i>		At1g43890	2231312	Stems and roots ⁴
<i>AtRABC2a</i>	<i>AtRab18B</i>		At5g03530	11274537	
<i>AtRABC2b</i>	<i>AtRab18C</i>		At3g09910	6681328	
<i>AtRABD1</i>			At3g11730	4097557	
<i>AtRABD2a</i>	<i>AtRab1B</i>	<i>Ara-5</i> ⁱ	At1g02130	5902803	Ubiquitous expression ²
<i>AtRABD2b</i>	<i>AtRab1A</i>		At5g47200	2245111	
<i>AtRABD2c</i>	<i>AtRab1C</i>		At4g17530	7268505	
<i>AtRABE1a</i>	<i>AtRab8B</i>		At3g53610	11274528	
<i>AtRABE1b</i>	<i>AtRab8D</i>		At4g20360	10172744	
<i>AtRABE1c</i>	<i>AtRab8A</i>	<i>Ara-3</i> ⁱ	At3g46060	114088	
<i>AtRABE1d</i>	<i>AtRab8C</i>		At5g03520	11274535	
<i>AtRABE1e</i>	<i>AtRab8E</i>		At3g09900	6681329	
<i>AtRABF1</i>	<i>AtRab5C</i>	<i>Ara-6</i> ^{iv}	At3g54840	13160603	
<i>AtRABF2a</i>	<i>AtRab5A</i>	<i>Rha1</i> ^v	At5g45130	400976	Stomatal guard cells, stipules, and root tips ⁵
<i>AtRABF2b</i>	<i>AtRab5B</i>	<i>Ara-7</i> ^v	At4g19640	7438427	
<i>AtRABG1</i>			At5g39620	9758338	
<i>AtRABG2</i>	<i>AtRab7A</i>		At2g21880	4417298	
<i>AtRABG3a</i>			At4g09720	7438424	
<i>AtRABG3b</i>			At1g22740	3914521	
<i>AtRABG3c</i>	<i>AtRab7D</i>		At3g16100	9294457	
<i>AtRABG3d</i>			At1g52280	18389228	
<i>AtRABG3e</i>			At1g49300	5430767	
<i>AtRABG3f</i>	<i>AtRab7B</i>		At3g18820	9293907	
<i>AtRABH1a</i>			At5g64990	8843763	
<i>AtRABH1b</i>	<i>AtRab6A</i>		At2g44610	7438386	Highest expression in liquid root culture ⁶
<i>AtRABH1c</i>			At4g39890	7438387	
<i>AtRABH1d</i>			At2g22290	4567198	
<i>AtRABH1e</i>			At5g10260	11274352	

^aAGI gene nomenclature. ^bGenbank protein accession no. (GI) for a translation of the corresponding gene. ^cNumbers in superscript refer to articles where expression and localization data can be found: 1, Yi and Gueriot (1994); 2, Anai et al. (1994) and Ueda et al. (1994); 3, Moore et al. (1997); 4, Mikami et al. (1997); 5, Terryn et al. (1993); 6, Bednarek et al. (1994).

Table II. *Arabidopsis ROP GTPase genes*

a, Nomenclatures used in Li et al. (1998, 1999, 2001), Bischoff et al. (2000), and Molendijk et al. (2000). b, Nomenclature used in Winge et al. (1997, 2000). c, Nomenclature used in Kost et al. (1999) and Lemichez et al. (2001).

Gene Name			AGI Gene ^a	Accession No. ^b	Expression/Localization ^c
a	b	c			
<i>AtROP1</i>	<i>Arac11/AtRAC11</i>		At3g51300	2558666	Pollen and flowers only ¹ /specific domains of the PM ⁴
<i>AtROP2</i>	<i>Arac4/AtRAC4</i>		At1g20090	1777764	Vegetative tissues and inflorescence ^{1,2} /specific domains of the PM ⁵
<i>AtROP3</i> ^d	<i>Arac1/AtRAC1</i>		At2g17800	7211191	Vegetative tissues, inflorescence, and pollen ^{1,2}
<i>AtROP4</i> ^e	<i>Arac5/AtRAC5</i>		At1g75840	7211204	Vegetative tissues and inflorescence ^{1,2} /perinuclear organelle ⁶
<i>AtROP5</i> ^f	<i>Arac6/AtRAC6</i>	<i>AtRac2</i>	<i>At4g35950</i>	7211206	Leaves, stems, flowers, and pollen; low in roots ¹
<i>AtROP6</i> ^g	<i>Arac3/AtRAC3</i>	<i>AtRac1</i>	<i>At4g35020</i>	7211200	Vegetative tissues and inflorescence (open flowers only) ^{1,2} /entire PM ⁶
<i>AtROP7</i>	<i>Arac2/AtRAC2</i>		At5g45970	7211198	Roots and stems only ⁷
<i>AtROP8</i>	<i>Arac9/AtRAC9</i>		At2g44690	5381420	ND
<i>AtROP9</i>	<i>Arac7/AtRAC7</i>		At4g28950	7211208	Entire PM ³
<i>AtROP10</i>	<i>Arac8/AtRAC8</i>		At3g48040	7211210 ^h	Entire PM ³
<i>AtROP11</i>	<i>Arac10/AtRAC10</i> ⁱ		At5g62880	10177466	ND

^aAGI gene nomenclature. ^bGenebank protein accession no. (GI:) for a translation of the corresponding gene. ^cPM, Plasma membrane; ND, not determined. Numbers in superscript refer to articles where expression and localization data can be found: 1, Li et al. (1998); 2, Winge et al. (1997); 3, Zheng et al. (2002); 4, Li et al. (1999); 5, Jones et al. (2002) and Fu et al. (2002); 6, Bischoff et al. (2000); 7, Z.L. Zheng and Z. Yang, unpublished data. ^d*AtROP3* is the same as *ATGP2* (GI: 4097563). ^e*AtROP4* is the same as *ATGP3* (GI: 4097565). The accession number GI: 2654009 ("RHO-like GTP binding protein") refers also to *AtROP4*. ^fGI: 2654011 corresponds also to *AtROP5*, but the sequence is incorrect; the last 100 amino acid residues are missing. ^g*AtROP6* is the same as the clone At43 published by Xia et al. (1996) and defined in the database as "Arabidopsis Rho1Ps homolog" (GI: 1732519). *AtROP6* is also found in the database under the accession number GI: 2645643. ^hThe accession no. GI: 4678324 refers also to *AtROP10/Arac8*, but the splicing suggested in the database is inaccurate; the last exon is missing. ⁱThe first intron of *AtROP11/Arac10* gene contains a cryptic exon of unknown function (GI: 7211195).

Table III. *Arabidopsis ARF GTPase genes*

a, Nomenclature used in this manuscript. b, Nomenclature used in Bischoff et al. (1999). c, Nomenclatures used in Regad et al. (1993).

Genes Name			AGI Gene ^a	Accession No. ^b
a	b	c		
<i>AtARFA1a</i>	<i>AtArf1</i>	<i>AtArf</i>	At1g23490	8778579
<i>AtARFA1b</i>			At5g14670	7573316
<i>AtARFA1c</i>			At2g47170	166586
<i>AtARFA1d</i>			At1g70490	6175141
<i>AtARFA1e</i>			At3g62290	6899939
<i>AtARFA1f</i>			At1g10630	6573751
<i>AtARFB1a</i>			At2g15310	4662630
<i>AtARFB1b</i>			At5g17060	9755707
<i>AtARFB1c</i>			At3g03120	6714424
<i>AtARFC1</i>			At3g22950	11994726
<i>AtARFD1a</i>			At1g02440	9972392
<i>AtARFD1b</i>			At1g02430	9857537
<i>AtARLA1a</i>			At5g37680	9757978
<i>AtARLA1b</i>			At3g49860	6723431
<i>AtARLA1c</i>			At3g49870	6723432
<i>AtARLA1d</i>			At5g67560	9757877
<i>AtARLB1</i>			At5g52210	1184981
<i>AtARLC1</i>			At2g18390	4309728
<i>AtSARA1a</i>	<i>AtSar1</i>		At1g09180	3249104
<i>AtSARA1b</i>			At1g56330	12321751
<i>AtSARA1c</i>	<i>AtSar2</i>		At4g02080	1314860

^aAGI gene nomenclature. ^bGenebank protein accession no. (GI:) for a translation of the corresponding gene.

Table IV. *Arabidopsis* RAN GTPase genes

Gene Name	AGI Gene ^a	Accession No. ^b	Expression/Localization ^c
<i>AtRAN1</i> ^d	At5g20010	2058278	Ubiquitous, highest in meristematic tissue/cytoplasm and nucleus
<i>AtRAN2</i> ^d	At5g20020	1668706	Same as <i>AtRAN1</i>
<i>AtRAN3</i> ^d	At5g55190	9758116 ^e	Same as <i>AtRAN1</i>
<i>AtRAN4</i> ^f	At5g55080	9758105	ND

^aAGI gene nomenclature. ^bGenbank protein accession no. (GI:) for a translation of the corresponding gene. ^cND, Not determined. Expression and localization data for *AtRAN1*, -2, and -3 have been published by Haizel et al. (1997). ^dNomenclatures used in Haizel et al. (1997). ^eIn the database, several GI numbers refer to the same *AtRAN3* sequence (GI: 9758116, GI: 2058280, and GI: 2801433). ^f*AtRAN4* has been described in the database as “salt stress-inducible small GTP-binding protein Ran-1-like protein.”

Rab GTPases

Rab GTPases make up the largest family of the small GTP-binding protein superfamily. Both in vivo and in vitro experiments have demonstrated the roles of this class of proteins in intracellular membrane trafficking. Furthermore, given the specific distributions of Rab GTPases to different cellular membranes, it was hypothesized that Rab GTPases would, in conjunction with SNARE proteins, provide specificity for membrane fusion events (for reviews, see

Stenmark and Olkkonen, 2001; Zerial and McBride, 2001).

Our analysis of the *Arabidopsis* genome (*Arabidopsis* Genome Initiative, 2000) identified 57 Rab GTPase isoforms (Table I; Fig. 2), which we have named *AtRAB* GTPases. This corresponded with the number recently determined by others (Pereira-Leal and Seabra, 2001). With a few notable exceptions, little is known of the function of *AtRAB* GTPases. As a result, our discussion of this gene family will focus

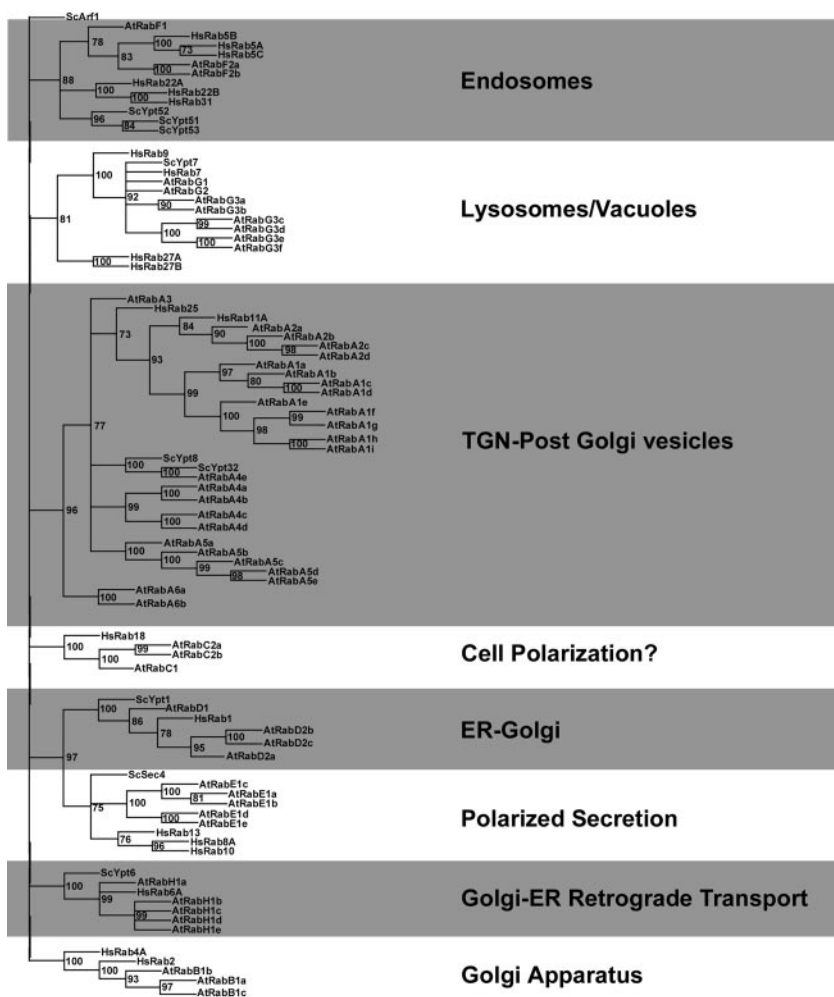


Figure 2. The Rab GTPase family of *Arabidopsis*. A neighbor-joining tree of *A. thaliana* Rab GTPases including representative Rab GTPase sequences of *S. cerevisiae* and *H. sapiens* was generated using ClustalW and scoring for amino acid differences (Thompson et al., 1994). The tree was rooted with a *S. cerevisiae* Arf1 sequence, and branches with percentage bootstrap values of less than 70% (of 1,000) were collapsed to simplify the tree. Labels to the right of the Rab GTPase subfamilies reflect functions determined for distinct Rab GTPases in that subfamily.

on organization and classification of members of the Rab GTPase family from Arabidopsis. Examination of the phylogenetic trees generated with protein sequences from human, yeast, and Arabidopsis Rab GTPases indicated that this GTPase family could be further divided into eight subfamilies based on sequence similarity and segregation with yeast and mammalian orthologs (see Fig. 2). A number of plant Rab GTPases have been cloned and named according to various schemes. To simplify and synchronize the nomenclature of the Rab GTPase family in Arabidopsis, we have followed the naming and classification scheme described in recent papers (Pereira-Leal and Seabra, 2000, 2001), and have named them according to their presence in one of the eight subfamilies that arose from our analysis (AtRABA-AtRABH; Fig. 2).

Correlation of Sequence Similarity with Localization in Different Species

Rab GTPase functions have been studied extensively in yeast and mammalian systems (for review, see Stenmark and Olkkonen, 2001). Individual members of the Rab GTPase family localize to different intracellular compartments where they regulate vesicle trafficking events (for review, see Simons and Zerial, 1993). *S. cerevisiae* only contains 11 Rab GTPases. Because of the cellular similarities between eukaryotic cells, we used the known distribution and function of these Rab GTPases as a basis for classification of plant Rab GTPases. *S. cerevisiae* Rab GTPases, termed yeast protein transport (YPT) proteins, are localized to the endoplasmic reticulum (ER), the Golgi apparatus and trans-Golgi network (TGN), the endosomal/prevacuolar compartments, and vacuoles in *S. cerevisiae*. They also regulate retrograde trafficking of proteins from Golgi to ER and polarized secretion (for review, see Lazar et al., 1997).

The correlation between sequence similarity and regulation of membrane trafficking through related compartments appears to be a conserved feature in the Rab GTPase family (Fig. 2). When functions of mammalian Rab GTPases sharing significant similarity to yeast counterparts have been studied, they regulate membrane trafficking through compartments with related function. For example, the human Rab8 GTPase is targeted to the protein secretion pathway, and displays high sequence similarity to Sec4p/Ypt2p, which regulates protein secretion in *S. cerevisiae*. Rab8 complements the *ypt2* mutant (Sec4p homolog) in fission yeast (*Schizosaccharomyces pombe*; Huber et al., 1993a, 1993b). Rab8 and Sec4p cosegregate in the same subfamily in our phylogenetic analysis (Fig. 2). Functions for most AtRAB GTPases have not yet been established. However, when plant Rab GTPase function is known, these isoforms also cosegregate in subfamilies containing their mammalian and yeast counterparts (discussed below). In particular, studies of Rab GTPases that are members of five

of the eight subfamilies highlighted by our phylogenetic analysis have been performed and published (AtRABA, Inaba et al., 2002; AtRABB, Cheung et al., 2002; AtRABD, Batoko et al., 2000; AtRABF, Ueda et al., 2001; AtRABH, Bednarek et al., 1994). In every case, their localization and/or function in membrane trafficking correlate with their segregation in subfamilies for yeast and mammalian Rab GTPases whose functions not only are the same, but also are well established. We have therefore indicated the compartments to which mammalian and yeast homologs are localized, and suggest that this may prove a useful initial framework for classification of AtRAB GTPases (Fig. 2).

Rab GTPases of the Endocytic Trafficking Pathway

Endocytosis is the primary means by which proteins and macromolecules too large to pass through the plasma membrane enter the cell. During endocytosis, regions of the plasma membrane with associated cargo molecules are invaginated and pinch off into transport vesicles, which then fuse with endocytic compartments. In addition, many proteins targeted to lysosomal and/or vacuolar compartments are sorted in endosomal compartments.

When compared with other eukaryotic Rab GTPases, three members of the AtRABF subfamily show highest similarity to human Rab5 and yeast Ypt51p family members (Fig. 2). In mammals, Rab5 GTPases localize to early endosomes and regulate fusion of clathrin-coated vesicles to early endosomes and fusion between early endosomes (Bucci et al., 1992). In yeast, Ypt51p family members similarly regulate membrane trafficking through prevacuolar compartments (Horazdovsky et al., 1994; Singer-Krüger et al., 1995). Both AtRABF2a (E. Nielsen, unpublished data) and AtRABF2b (Ueda et al., 2001) localize to compartments observed upon uptake of the fluorescent styryl dye, FM 4-64. Interestingly, the third Arabidopsis member of this subfamily, AtRABF1, appears to be a novel, plant-specific Rab GTPase (Bolte et al., 2000; Ueda et al., 2001). AtRABF1 lacks the traditional carboxy-terminal-CAAX motif for posttranslational isoprenylation but is myristoylated at the amino terminus (Ueda et al., 2001). Despite these differences, AtRABF1 colocalizes with AtRABF2b on the plant endosomal compartment where it may provide novel regulatory functions (Ueda et al., 2001).

The AtRABG subfamily contains eight members with significant similarity to human Rab7 and Ypt7p (Fig. 2). In mammals, Rab7 regulates transport of cargo from early endosomes to late endosomes and lysosomes (Feng et al., 1995; Mukhopadhyay et al., 1997). In yeast, Ypt7p localizes to vacuoles and regulates homotypic fusion between vacuolar compartments (Price et al., 2000). In plants, vacuoles are used as storage organelles in addition to having lytic roles, and single cells can contain multiple vacuole types

(Paris et al., 1996; Vitale and Raikhel, 1999). This increased complexity appears to be reflected in the large number of members in this Rab GTPase subfamily in plants.

Rab GTPases of the Biosynthetic Trafficking Pathway

Newly synthesized proteins enter the secretory pathway by translocation through ER membranes. Subsequent transport from the ER to Golgi complexes involves recruitment of cargo proteins into vesicle transport intermediates. The AtRABD subfamily contains five members that display significant similarity with mammalian Rab1 and Ypt1p Rab GTPases (Fig. 2). In mammals, Rab1 GTPase isoforms localize to ER, ER Golgi intermediate compartment, and Golgi compartments and regulate ER-to-Golgi membrane trafficking steps (Tisdale et al., 1992; Nuoffer et al., 1994). In yeast, *YPT1* is an essential gene required for ER-to-Golgi trafficking events (Segev et al., 1988). In plants, transient expression of a dominant-negative mutant of AtRABD2a resulted in accumulation of a secreted GFP marker in an intracellular compartment reminiscent of the ER and inhibited movement of Golgi complexes along cytoskeletal elements (Batoko et al., 2000). Arabidopsis contains three AtRABB subfamily members related to the human Rab2 GTPase. Rab2 GTPases are found associated with ER Golgi intermediate compartment membranes and COP-I transport vesicles in mammalian cells (Tisdale et al., 1992). The function and localization of the AtRABB subfamily members as yet remains uncharacterized in plants.

Recycling of ER-resident proteins, such as the KDEL-receptor Erd2p and Sec23p/Sec24p cargo receptor proteins requires the specific enrichment of these proteins in transport vesicles and subsequent delivery back to ER membranes (for review, see Kirchhausen, 2000). In mammals, Rab6 GTPases regulate this process (Martinez et al., 1997; White et al., 1999). Five isoforms of this subfamily of Rab GTPases (AtRABH) are present in the Arabidopsis genome. Of these, the AtRABH1b homolog, previously identified as AtRAB6, was shown to complement a yeast *ypt6* mutation, although the function of this Rab GTPase in plants was not determined (Bednarek et al., 1994).

Rab GTPases Involved in Polarized Secretion

Arabidopsis contains five Rab GTPases that cosegregate with yeast and mammalian isoforms that regulate polarized secretion (AtRABE subfamily; Fig. 2). In yeast, Sec4p regulates membrane trafficking to the daughter cell bud site (Salminen and Novick, 1987; Goud et al., 1988). In plants, members of this Rab GTPase subfamily may have roles in cellular responses to bacterial pathogens. In tomato (*Lycopersicon esculentum*), yeast two-hybrid screening for plant proteins that interact with the *Pseudomonas* sp. aviru-

lence factor, *avrPto*, identified a RabE subfamily member with significant similarity to mammalian Rab8 (Bogdanove and Martin, 2000). This interaction only occurred in absence of the tomato resistance protein, Pto (Bogdanove and Martin, 2000), raising the possibility that in susceptible plants, *avrPto* may interfere with membrane trafficking pathways regulated by this RabE homolog. In an intriguing possibility, the authors suggest this Rab GTPase might regulate polarized secretion of antimicrobial compounds and/or components involved in mounting cellular responses to attack by bacterial pathogens.

A Plethora of Post-Golgi Rab GTPases in Plants

One of the most striking features evident upon examination of the AtRAB GTPases is the large number of AtRABA subfamily members (Fig. 2). This subfamily contains Rab GTPases from mammals and yeast that regulate TGN membrane trafficking pathways. In mammals, human Rab11A and Rab11B isoforms localize to recycling endosomes (Ullrich et al., 1996), and in epithelial cells, mammalian Rab11A is critical for exit of internalized proteins from apical recycling endosomes (Calhoun et al., 1998; Duman et al., 1999). Yeast Ypt31p and Ypt32p Rab GTPases appear to function during exit of membrane traffic from trans-Golgi cisternae (Benli et al., 1996; Jedd et al., 1997).

The Arabidopsis genome contains 26 distinct AtRABA subfamily members. Whether this multitude of genes provides distinct functions or represents functionally redundant gene families remains to be determined. Antisense inhibition of RABA subfamily GTPases in tomato results in complex developmental abnormalities and delayed fruit ripening (Lu et al., 2001). In pea (*Pisum sativum*), expression of two RABA GTPases, *PRA2* and *PRA3*, was examined (Nagano et al., 1995). Whereas *PRA3* was constitutively expressed, *PRA2* was specifically up-regulated in the stem-elongation zone of dark-grown seedlings, a region in which rapid plant cell expansion is observed (Nagano et al., 1995). More recent experiments indicated that *Pra2* GTPase localizes predominantly to Golgi and possibly endosomal compartments, whereas *Pra3* appears on TGN and/or prevacuolar compartments (Inaba et al., 2002). These RABA isoforms were suggested to play a role in delivery of new cell wall components to the plasma membrane. Interestingly, identification of effector proteins for RABA GTPases may further support roles for this class of GTPases in stem elongation in dark-grown seedlings because one effector protein for this Rab GTPase was identified as a cytochrome p450 involved in biosynthesis of brassinosteroids (Kang et al., 2001). Plant cells display a high degree of polarized deposition of primary and secondary cell wall components, in some cases, adjacent walls of a cell display differential localization of antigenic determinants (Lynch and Staehelin, 1992). The delivery of hemicelluloses, integral cell wall proteins,

and probably even delivery of the cellulose synthase complex to the plasma membrane likely occur via post-Golgi membrane trafficking pathways. Could the large numbers of Rab GTPases in the AtRABA subfamily reflect the increased complexity associated with the highly polarized deposition of cell wall material observed in plants?

Rab-Interacting Proteins

Rab GTPases cycle between an inactive GDP-bound form located in the cytosol and an active GTP-bound form that is membrane associated. For most Rab GTPases, membrane association is promoted by posttranslational lipid modification. Stabilization of the lipid-modified Rab GTPase in cytosol occurs through association with Rab GDP-dissociation inhibitor proteins (RabGDI). Three RabGDI homologs (*AtRabGDI1-AtRabGDI3*; see supplemental data) are present in Arabidopsis. Two of these, *AtRabGDI1* and *AtRabGDI2* were identified by complementation of yeast *sec19* mutants, indicating functional conservation of activity between yeast and plants. Although *AtRabGDI1* was ubiquitously expressed, *AtRabGDI2* displayed higher expression levels in suspension cultures cells and in roots (Ueda et al., 1996, 1998; Zarsky et al., 1997). *AtRabGDI1-AtRabGDI3* homologs all share significant similarity (more than 77% identity). Interestingly, two RabGDI transcripts were determined to be up-regulated during early stages of fungal infection in rice (*Oryza sativa*; Kim et al., 1999), however the exact roles of these proteins during early plant defense responses to fungal infection remain unknown. A fourth related protein, *AtREP1* displays similarity to the *AtRabGDI* homologs, but appears somewhat diverged (27% identity, 44% similarity), instead displaying higher similarity to mammalian REP-1 (Alexandrov et al., 1994). In mammals, REP-1 associates with newly synthesized Rab GTPases and presents them for post-translational lipid modification.

GTP/GDP exchange is essential for Rab GTPase function. Upon binding to its target membrane, the Rab GTPase is converted from Rab:GDP to Rab:GTP through the action of RabGEF proteins. RabGEFs have been identified for several Rab GTPases in mammals and yeast (Horiuchi et al., 1997; Wada et al., 1997; Walch-Solimena et al., 1997; Hama et al., 1999; Iwasaki and Toyonaga, 2000; Siniosoglou et al., 2000). In general, significant levels of sequence similarity have not been observed between RabGEFs for different Rab GTPases. This suggests that despite having similar guanine exchange functions, different RabGEF proteins may have diverse evolutionary origins. However, in one notable exception to this trend, Vps9p, the RabGEF for the yeast Rab GTPase, Ypt51p, and Rabex-5, which catalyzes nucleotide ex-

change on mammalian Rab5, shows significant sequence similarity (Horiuchi et al., 1997; Hama et al., 1999). The Arabidopsis genome contained two sequences with significant similarity to the Vps9p and Rabex-5 RabGEF sequences. We designated these RabGEFs *AtVPS9A* and *AtVPS9B* (supplemental data). No Arabidopsis sequences with significant primary amino acid similarities could be detected for non-Vps9p-like RabGEFs.

As opposed to RabGEFs, which activate Rab GTPases, RabGAPs inactivate Rab GTPases by accelerating the slow intrinsic Rab GTPase activity. The first Rab-specific GAP proteins, Gyp6p, Gyp7p, and Gyp1p, were identified in yeast (Strom et al., 1993; Vollmer and Gallwitz, 1995; Vollmer et al., 1999). These proteins all contained six conserved sequence motifs within their catalytic domains (Albert et al., 1999). In other proteins, such as RN-tre and GAPCenA proteins, presence of these motifs successfully predicted their RabGAP activity (Cuif et al., 1999; Lanzetti et al., 2000). Analysis of the Arabidopsis genome revealed 20 proteins with all or most of the RabGAP "catalytic core" motifs, which we have named *AtGYP* proteins (supplemental data). All *AtGYP* proteins contained a conserved Arg residue critical for RabGAP activity (Albert et al., 1999) and at least five of the six "catalytic core" motifs. Although additional sequences containing portions of the RabGAP "catalytic core" were detected, these were not included because they (a) were missing two or more conserved motifs, and (b) did not contain essential conserved residues. Apart from the RabGAP motif, *AtGYP* family members contain no other detectable protein domains and displayed significant sequence diversity. In animals, two RabGAP proteins have been found associated with noncatalytic partner proteins (Nagano et al., 1998; Lanzetti et al., 2000). It seems likely that other RabGAP proteins may also be found in association with partner proteins.

In the last several years, many Rab effector proteins have been characterized. Because Rab GTPases are highly conserved and function similarly in different organisms, one might expect that Rab effector proteins also be conserved. But this is not apparently the case (Zerial and McBride, 2001). The structural heterogeneity of Rab effectors make it unlikely that plant Rab effectors can be identified by sequence similarity alone. However, some established Rab effectors, such as lipid kinases (Christoforidis et al., 1999), are present in Arabidopsis. In addition, in several cases, zinc-finger domains are associated with Rab effector protein functions (Christoforidis et al., 1999; Simonsen et al., 1998), and proteins with these domains are present in Arabidopsis (Jensen et al., 2001; Heras and Drobak, 2002). Identification of plant Rab effector proteins clearly is likely to uncover specialized proteins whose activities have been exclusively tailored for plant-specific transport systems.

The expression and localization patterns of the 11 AtROP GTPases (Table II) is consistent with their involvement in complex signaling networks in plants (Yang, 2002). Given the conspicuous absence of Ras-family GTPases and other RHO, RAC, and CDC42 subfamily members, it was hypothesized that the ROPs might replace these GTPases in plant signaling (Li et al., 1998; Winge et al., 2000). Overexpression of wild-type ROP GTPase genes and expression of mutants either deficient for GTPase activity (constitutively active; CA) or that can only bind GDP (dominant negative [DN]) have provided evidence for the above hypothesis (for reviews see, Yang, 2002; Zheng and Yang, 2000a).

Mammalian and fungal Rho GTPases regulate the establishment of cell polarity and influence cell morphogenesis (for review, see Arellano et al., 1999). Plant ROP GTPases appear to have retained this function. Three *AtROP* genes, *AtROP1*, *AtROP3*, and *AtROP5*, are expressed in pollen and may be functionally redundant during pollen tube growth. *AtROP1* and *AtROP5* are preferentially localized to the plasma membrane in apical regions of the pollen tube and control tip growth (Kost et al., 1999; Li et al., 1999). *AtROP2* and *AtROP4* localize to tips of elongating root hairs (Molendijk et al., 2001; Jones, 2002). CA mutants of *AtROP2*, *AtROP4*, and *AtROP6* either caused isotropic growth or increased length in root hairs of Arabidopsis, whereas DN-*AtROP2* mutants inhibited root hair elongation, indicating that *AtROP* GTPases also control tip growth during root hair development (Molendijk et al., 2001; Jones, 2002). In both root hairs and pollen tubes, *AtROP* GTPases control tip growth by modulating the formation of both the dynamic fine tip F-actin and a tip-focused cytosolic calcium gradient (Li et al., 1999; Fu et al., 2001; Fu and Yang, 2001; Molendijk et al., 2001; Jones, 2002). *AtROP* GTPases also regulate formation of cell shape by controlling assembly of dynamic cortical F-actin in cells that do not undergo tip-based growth such as epidermal cells (Fu and Yang, 2002).

In mammals, RAC GTPases control production of reactive oxygen species by directly associating with and regulating the activity of plasma membrane-associated NADPH oxidase complexes (for review, see Ridley, 1995). In Arabidopsis, oxygen deprivation rapidly and transiently activates ROP GTPases, resulting in H₂O₂ accumulation and alcohol dehydrogenase gene expression (Baxter-Burrell et al., 2002). ROP GTPase-dependent H₂O₂ production is blocked by treatment with DPI, an inhibitor of the plasma membrane NADPH oxidase (Kawasaki et al., 1999; Baxter-Burrell et al., 2002). These results are consistent with the suggestion that plant ROP GTPases may regulate the plasma membrane NADPH oxidase complex.

AtROP GTPases are also involved in signal transduction pathways mediated by the plant hormone, abscisic acid (ABA). ABA promotion of seed dormancy was enhanced and inhibited respectively by

DN-*Atrop2* and CA-*Atrop2* expression in Arabidopsis (Li et al., 2001). Lemichez et al. (2001) showed that *AtROP6* was implicated during the negative regulation of stomatal closure, and studies using loss of function mutants have revealed that *AtROP9* and *AtROP10*, two putative ERA1 targets, act as general negative regulators of ABA responses (Zheng et al., 2002). ERA1 encodes a β -subunit of protein farnesyltransferase involved in the negative regulation of ABA responses both in guard cell movement and seed dormancy.

Plant ROP GTPases may also regulate the accumulation of and/or responses to brassinolide and auxin. CA-*AtROP2* plants exhibit diverse morphologies that resemble auxin- or brassinolide-overproduction mutants, whereas DN-*AtROP2* plants show opposite phenotypes (Li et al., 2001). Readers are referred to several recent reviews for more detailed discussion of the function of the ROP family GTPases in these processes (Valster et al., 2000; Zheng and Yang, 2000a, 2000b; Fu and Yang, 2001; Yang, 2002).

Rop-Interacting Proteins

Several types of Rho GTPase regulators are known in animals: GAPs and GDIs (guanine nucleotide dissociation factors) that act as negative regulators of Rho GTPase function, and GEFs, which activate these GTPases through exchange of GDP for GTP (see introduction and "RAB GTPases" section).

In most fungal and animal RhoGEFs, GDP-GTP exchange activity typically is localized in Dbl-homology domains (Cerione and Zheng, 1996). Surprisingly, plants lack proteins containing obvious Dbl-homology domains. This suggests that plants may have evolved different mechanisms to activate the GDP-bound ROP GTPases. One possible mechanism of activation may be through direct association of ROP GTPases with plant receptor-like kinases (RLKs). A plant ROP GTPase was co-immunoprecipitated with the plant RLK, CLAVATA1 (Trochoud et al., 1999). Therefore, plant ROP GTPases could be activated by RLKs directly, although the functional significance of ROP-RLK association remains to be determined.

Three RhoGDI homologs are present in Arabidopsis. *AtRhoGDI1* is expressed in all tissues examined and has been shown to interact specifically with *AtROP* GTPases (Bischoff et al., 2000), but so far, no functional data is available. Using the yeast two-hybrid method, Wu et al. (2000) identified a family of Rop-specific, Rho GAPs called RopGAPs. Arabidopsis contains six RopGAPs, all with an N-terminal Cdc42/Rac-interactive binding (CRIB) motif located adjacent to a conserved RhoGAP catalytic domain. In yeast and animals, CRIB domains mediate GTP-dependent interaction between Rho GTPases and their effector proteins, but are not found in conjunction with RhoGAP domains. In plants, presence of

the CRIB domain is critical for the Rop-specific regulation of GAP activity (Wu et al., 2000). RopGAP1 localizes in the apical PM region in pollen tube and acts as a negative regulator of AtROP1 function during pollen tube growth (G. Wu and Z. Yang, unpublished data). A role for RopGAP4 in Arabidopsis responses to oxygen deprivation has been revealed using a *ropgap4* knockout mutant (Baxter-Burrell et al., 2002).

No obvious homologs of animal and yeast Rho effector proteins, except for phosphatidylinositol-phosphate kinases, are present in the Arabidopsis genome. Because the CRIB motif is a hallmark structure of many CDC42/RAC effectors, plant proteins containing these domains make attractive Rop GTPase effector protein candidates. Arabidopsis contains a novel plant-specific family of 11 putative ROP GTPase effector proteins, Rop-interactive CRIB motif-containing proteins (RICs; Wu et al., 2001). RIC1 associates with AtROP1 in a nucleotide-dependent manner, and this association is dependent upon the CRIB domain. Because RIC family members display little sequence similarity with other proteins or even with one another (apart from the CRIB domain), it seems unlikely that the variable regions of RICs function as enzymes to regulate downstream events. RICs could act as adaptor proteins linking Rop GTPases to effector proteins, which in turn would activate specific downstream events. Such a linker function would allow the generation of a greater functional diversity for the Rop GTPase switch and would present a novel mechanism for the activation of G protein effectors (Wu et al., 2001; Yang, 2002).

Arf GTPases

ADP-ribosylation factors (Arfs) were initially identified due to their ability to stimulate the ADP-ribosyltransferase activity of cholera toxin A (Moss and Vaughan, 1998). GTPases that shared significant similarity (40%–60% identity), but that could not activate cholera toxin A or could not rescue *S. cerevisiae* mutants were termed Arf-like GTPases (Arl GTPases; Clark et al., 1993). Today these two subfamilies are classified together as Arf GTPases (Moss and Vaughan, 1998). The Arabidopsis genome (Arabidopsis Genome Initiative, 2000) contains 21 Arf GTPase family members, with isoforms present in both Arf and Arl GTPase subfamilies (Table III; Fig. 4).

Arf GTPases Recruit Coat Proteins to Transport Vesicles

Arf GTPases play important roles during membrane trafficking steps in eukaryotic cells (for review, see Chavrier and Goud, 1999). In all cases, Arf GTPases act to recruit cytosolic coat proteins to sites of vesicle budding. Three types of protein coats have been described for transport vesicles, COP-I, COP-II,

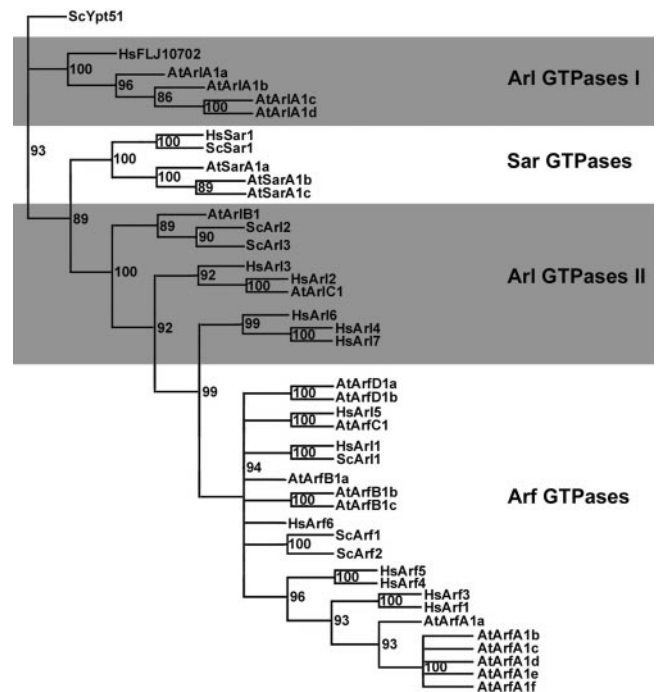


Figure 4. The Arf GTPase family of Arabidopsis. A neighbor-joining tree of *A. thaliana* Arf and Arl GTPases including representative Arf and Arl GTPase sequences of *S. cerevisiae* and *H. sapiens* was generated using ClustalW and scoring for amino acid differences (Thompson et al., 1994). The tree was rooted with the *S. cerevisiae* Rab GTPase, Ypt51, and branches with percentage bootstrap values of less than 80% (of 1,000) were collapsed to simplify the tree.

and clathrin coats (Kirchhausen, 2000), and formation of these coats are attributed to two distinct subsets of the Arf GTPase family. Arf GTPases recruit COPI and clathrin protein coats to transport vesicles, whereas a specific subset of the Arf GTPase family, the Sar1p GTPases, recruit COP-II coats.

The best understood mechanism by which Arf GTPases recruit coat proteins to transport vesicles is in *S. cerevisiae*, where recruitment of COP-II protein coat by Sar1p has been extensively studied (for review, see Kirchhausen, 2000). Sar1p GTPases regulate formation of COP-II coated vesicles carrying cargo from the ER to the cis-Golgi compartment (Wieland and Harter, 1999). In Arabidopsis, three Sar1p homologs have been identified (Fig. 4). Because these GTPases cosegregated with animal and yeast Sar1 GTPases, we have named them AtSARA1a, AtSARA1b, and AtSARA1c, respectively (Fig. 4; Table III). *AtSARA1a*, previously identified as *AtSAR1*, was shown to functionally complement mutants in *S. cerevisiae* (d'Enfert et al., 1992), and *AtSARA1a* expression was correlated to levels of secretion activity from ER membranes in plant cells, because blockage of transport from the ER resulted in up-regulation of mRNA levels for *AtSARA1a* (Bar-Peled et al., 1995).

In *S. cerevisiae*, two Arf GTPases, Arf1p and Arf2p, are functionally interchangeable and act to recruit COP-I coat proteins to transport vesicles during

transport of cargo from cis-Golgi compartments back to the ER (Cosson and Letourneur, 1997; Gaynor and Emr, 1997). The yeast Arf1p and Arf2p GTPase functions are distinct from that of Sar1p, which acts in ER-to-Golgi trafficking (Barlowe et al., 1994). In Arabidopsis, 12 Arf GTPase isoforms that cosegregated with members of the Arf GTPase subfamily were identified (Table III; Fig. 4). It should be noted, however, that we also observed mammalian and yeast Arl GTPase sequences within this grouping. Six AtARF GTPases cosegregated with human Arf1 and Arf3 sequences (AtARFA subfamily). This included *AtARFA1a*, previously identified as *AtArf1* (Regad et al., 1993), which has been localized to peripheral Golgi stacks along with At γ -COP, an Arabidopsis homolog of the COP-I coat protein complex (Ritzenhaller et al., 2002). Six additional Arabidopsis sequences cosegregated with the Arf GTPase subfamily (AtARFB-AtARFD), however, no significance could be ascribed to these groupings outside that of the AtARFA subfamily, which cosegregated with a distinct subset of mammalian Arf GTPases.

Arl GTPases: Regulatory Proteins with Mystery Functions

Six of the 21 Arabidopsis Arf GTPases segregate in groups that contain neither Sar1p nor Arf GTPase subfamily members (Fig. 4). As such, they have been classified as Arl GTPases. Little is known of the function of these GTPases, although at least one member provides essential functions as demonstrated in fruitfly (Tamkun et al., 1991). In plants, mutation of the *TITAN5* gene, which corresponds to *AtARLC1*, results in dramatic alterations of mitosis and cell cycle control during seed development (McElver et al., 2000). The authors speculated that this gene might play a role in membrane trafficking steps necessary for proper cell plate deposition during cytokinesis in developing embryos (McElver et al., 2000). However, much remains to be determined regarding the role(s) of these proteins in any eukaryotic organism.

Arf GTPase-Interacting Proteins

Like other small GTPases, Arf GTPases cycle between an active, membrane-bound form when associated with GTP, and an inactive, predominantly cytosolic form when bound to GDP. Similarly, this GTPase cycle is regulated by GEFs (ArfGEFs) and GAPs (ArfGAPs; Donaldson and Jackson, 2000). However, unlike Rho and Rab GTPases, Arf GTPases do not require GDIs to catalyze their delivery to membranes. Whereas Rab and Rho GTPases are lipid modified at their carboxy terminus, Arf GTPases are myristoylated at their amino terminus. Due to this myristoylation, GDP-bound Arf GTPases display a weak interaction with membranes (Franco et al., 1995). Activation of the membrane-associated Arf GTPase through GEF-catalyzed exchange of GDP for

GTP causes conformational changes resulting in stabilization of the Arf GTPase-membrane interaction (Antonny et al., 1997).

Much of the research into Arf GTPase function has focused upon their roles in coat protein recruitment during vesicle formation (see above), and several proteins that interact with Arf GTPases during these processes such as Sec12p and Sec23p, display Sar1p-specific GEF and GAP activities, respectively (Springer et al., 1999). However, Arf GTPases also have other roles such as alteration of membrane lipid compositions, and actin remodeling upon various membrane compartments of the cell (Donaldson, 2000). Distinct ArfGEF or ArfGAP proteins regulate these additional activities.

To date, all ArfGEFs possess a Sec7 domain (Donaldson and Jackson, 2000). This 200-amino acid domain is sufficient to catalyze GDP for GTP exchange on Arf GTPases in vitro (Chardin et al., 1996). Arabidopsis encodes eight proteins containing Sec7 domains. One of these ArfGEFs, *GNOM*, was identified in mutant screen for Arabidopsis with defective body organization in embryos (Mayer et al., 1991, 1993). Polarized distribution of the auxin-efflux carrier, *PIN1*, was disrupted in *gnom* mutants, indicating that this ArfGEF is involved in trafficking of *PIN1* proteins during development (Steinmann et al., 1999; Geldner et al., 2001). Little is known of the functions of the other seven Arabidopsis ArfGEF proteins. However, given the central role of the ArfGEF *GNOM* during organization of the early embryo, this class of proteins is likely to fulfill important roles in growth and development in plants.

In animals, ArfGAPs are a diverse family of multidomain proteins (Donaldson, 2000) that contain a zinc-finger motif and a conserved Arg residue within the ArfGAP catalytic domain (Cukierman et al., 1995; Randazzo et al., 2000). Arabidopsis contains 15 proteins with ArfGAP domains, and we have termed these Arf GAP domain (AGD) proteins. We grouped the AtAGD proteins into four distinct classes based on phylogenetic analysis (data not shown) and overall domain organization (Fig. 5). The first class of AtAGD proteins, consisting of AtAGD1-AtAGD4, is a novel, plant-specific family of putative Arf-GAP proteins. These proteins all contain pleckstrin homology (PH) domains, and two or three ankyrin repeat domains in addition to the AGD (Fig. 5). Presence of PH domains in these AtAGD proteins raises the possibility that they may coordinate their ArfGAP activity with phospholipid signaling events as PH domains bind to phosphoinositides (for review, see Lemmon et al., 2002). In addition, AtAGD1, AtAGD2, and AtAGD3 contain amino-terminal Bin1-amphiphysin-Rvs167p/Rvs161p (BAR) domains. BAR domains are found in adaptor proteins implicated in numerous biological functions including actin regulation and synaptic vesicle endocytosis (Wigge and McMahon, 1998; Balguerie et al., 1999). However,

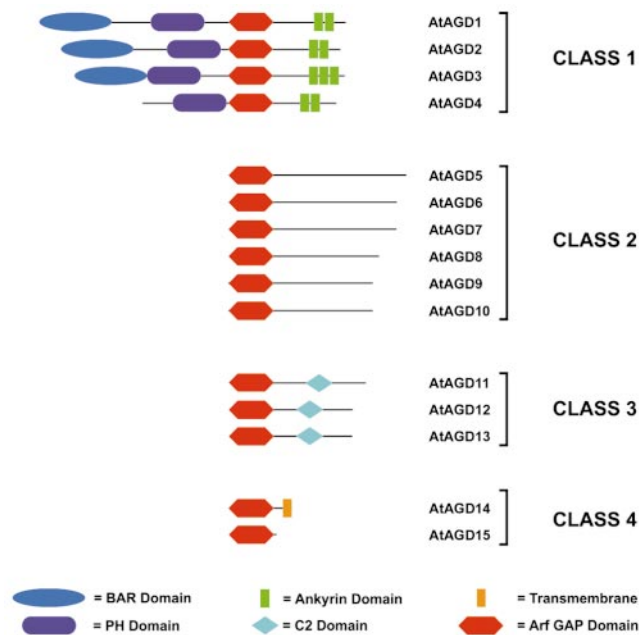


Figure 5. ArfGAP proteins of Arabidopsis. AtAGD proteins were identified using BLASTP (Altschul et al., 1997) with the AGD of ASAP1 (GenBank accession no. NP_060952). Domains within the AtAGD sequences were detected using SMART (Letunic et al., 2002). AGI numbers of *AtAGD1* to *AtAGD15* are as follows: *At5g61980* (*AtAGD1*), *At1g60680* (*AtAGD2*), *At4g13300* (*AtAGD3*), *At1g10870* (*AtAGD4*), *At5g54310* (*AtAGD5*), *At3g53710* (*AtAGD6*), *At2g37550* (*AtAGD7*), *At4g17890* (*AtAGD8*), *At5g46750* (*AtAGD9*), *At2g35210* (*AtAGD10*), *At3g07490* (*AtAGD11*), *At4g21160* (*AtAGD12*), *At4g05330* (*AtAGD13*), *At1g08680* (*AtAGD14*), and *At3g17660* (*AtAGD15*).

presence of BAR domains in ArfGAP proteins appears to be plant specific. Class 2 AtAGD proteins (*AtAGD5*–*AtAGD10*; Fig. 5) contain AGDs at the amino terminus and no other discernible protein domains. These proteins show similarity both at sequence level and in overall domain organization to Golgi-localized ArfGAPs in mammals and in yeast (Cukierman et al., 1995; Poon et al., 1999). Class 3 AtAGD proteins (*AtAGD11*–*AtAGD13*; Fig. 5) are distinguished by the presence of a centrally located C2 domain. C2 domains bind a range of ligands, such as phospholipids, phosphoinositides, and other proteins, in calcium-dependent fashion (Sutton et al., 1995; Essen et al., 1996; Shao et al., 1997). In Class 4 AtAGD proteins (*AtAGD14* and *AtAGD15*; Fig. 5), the AGD comprises almost the entire open reading frames of these proteins. In *AtAGD14*, the AGD is followed by an apparent membrane-spanning α -helix, suggesting this ArfGAP could be an integral membrane protein. As in mammals, plant ArfGAPs have diversified into a large family of proteins. Several classes of the AtAGD proteins contain additional domains, and these may act to coordinate the timing of ArfGAP activity within the cell or may serve functions beyond simply acting as negative regulators of Arf GTPases. Understanding the mechanisms by

which temporal and spatial activities of plant Arf-GAPs are regulated will be an interesting area for future investigation.

In yeast and animals, increasing attention is being paid to the roles of Arf GTPases in lipid modification. Mammalian and yeast Arf GTPases activate phosphatidylinositol 4-phosphate, 5-kinases (Honda et al., 1999; Walch-Solimena and Novick, 1999). Arabidopsis also contains phosphatidylinositol 4-phosphate, 5-kinase proteins, but whether AtARF GTPases interact with these remains unknown. A new class of Arf GTPase effector proteins that facilitate membrane trafficking at the TGN has recently been described. These Golgi-localizing, γ -adaptin ear homology, Arf-binding (GGA) proteins contain multiple domains: an amino-terminal VHS (Vps27, Hrs, and STAM) domain, followed by the GAT (GGA1 and TOM proteins) region and the γ -adaptin homology domain at the carboxy terminus. The VHS domain is thought to mediate interactions with cargo molecules, whereas the GAT domain is responsible for association with Arf GTPases, and the γ -adaptin homology domain interacts with clathrin (for review, see Boman, 2001). Five GGA-like proteins are present in the Arabidopsis genome. Intriguingly, although these proteins contain the amino-terminal VHS and GAT domains, they appear to lack the carboxy-terminal γ -adaptin homology domain. Given the apparent central role of GGA proteins during sorting of cargo proteins and recruitment of vesicle coat proteins, it will be interesting to determine the role of this class of proteins in plant Arf GTPase functions.

Ran GTPases

In animals, Ran GTPases together with their regulatory factors, the Ran-binding proteins (RanBPs), RCC1 (a GEF, RanGEF) and RanGAP (a GAP), play key roles in controlling nuclear processes throughout the cell mitotic cycle (for review, see Clarke and Zhang, 2001). Ran GTPases, like other small GTPases, cycle between GDP- and GTP-bound states. However, for Ran GTPases, GTP binding and hydrolysis is linked to transport into or out of the nucleus (Moore, 1998). Also, unlike other small GTPases, Ran GTPases are not posttranslationally lipid modified and do not associate with cellular membranes (Rush et al., 1996). Four Ran GTPases are present in Arabidopsis (Table IV). *AtRAN1*, *AtRAN2*, and *AtRAN3* were identified by sequence similarity (Haizel et al., 1997; see below). The fourth gene, *AtRAN4*, is annotated as “salt stress-inducible small GTP-binding protein Ran1-like protein,” but so far no data has been published.

At the protein level, *AtRAN1*, *AtRAN2*, and *AtRAN3* are nearly identical (95%–96% of identity) differing only in their C-terminal regions, whereas *AtRAN4* is more divergent with only 65% identity to the other *AtRAN* sequences. All *AtRan* GTPases con-

tain sequence motifs involved in GTP binding/hydrolysis and an effector-binding domain for interaction with RanGAPs. This effector-binding motif is 100% identical in AtRAN1 to AtRAN3 and in tomato and tobacco Ran GTPases (KKYEPTIGVEV) but diverges strikingly in AtRAN4, with only five residues of 11 conserved. Moreover, although other plant Ran GTPases possess conserved C-terminal acidic domains (DDDDD/E), this sequence is absent in AtRAN4. In animals, this acidic domain is necessary for interaction with Ran-binding proteins (Haizel et al., 1997; see below). These observations suggest that AtRAN1 to AtRAN3 are likely Ran orthologs involved in nucleocytoplasmic transport (see below), whereas AtRAN4 may have distinct functions in Arabidopsis.

Ran GTPases Establish Nuclear Compartment Identity

In interphase cells, Ran GTPases direct nucleocytoplasmic transport. RanGAP and RanBP2 (which stimulate the intrinsic GTPase activity of the Ran GTPase) are cytoplasmic proteins, whereas RCC1 (which generates Ran:GTP) is confined to the nucleus. These distributions ensure that Ran GTPases are bound with GTP in the nucleus and GDP in the cytoplasm. The restriction of Ran:GTP to the nuclear interior has been proposed to be crucial for establishing directional transport of proteins and RNA through the nuclear pore (Moore and Blobel, 1993; Izaurralde et al., 1997). Ran GTPases also play important roles during cellular mitosis (Kalab et al., 1999; Wilde and Zheng, 1999). Association of RCC1 with chromatin results in localized generation of Ran:GTP in the vicinity of chromosomes, which in turn promotes microtubule spindle assembly (Carazo-Salas et al., 1999; Ohba et al., 1999; Wiese et al., 2001). At the end of mitosis, the cycling of Ran:GDP and Ran:GTP induces nuclear envelope reassembly, probably by controlling vesicle binding and fusion (Hetzer et al., 2000; Zhang and Clarke, 2000).

Functions of Arabidopsis Ran GTPases and Associated Proteins

AtRAN1 to AtRAN3 are ubiquitously expressed during development, with higher levels in meristematic tissues and developing embryos (Haizel et al., 1997). Overexpression of plant Ran GTPases suppressed cell cycle defects in mutant fission yeast, indicating that they functioned similarly to their yeast homologs (Ach and Gruissem, 1994; Merkle et al., 1994; Haizel et al., 1997). In animals, Exportin-1 is the nuclear export receptor for proteins carrying nuclear export signals (NES). Exportin-1 binds NES-containing proteins and Ran GTPases cooperatively, and this triple complex undergoes nuclear export (Fornerod et al., 1997). In Arabidopsis, AtRAN1 interacts with AtXPO1, an Arabidopsis Exportin-1 ho-

molog, and a NES-containing protein, AtRanBP1a (Haizel et al., 1997; Haasen et al., 1999). This suggests the nuclear export machinery may be functionally conserved in plants (Haasen et al., 1999). Antisense expression of an additional AtRanBP isoform in Arabidopsis, AtRanBP1c, enhanced primary root growth, suppressed lateral root growth and rendered transgenic roots hypersensitive to auxin (Kim et al., 2001). The authors proposed a model where AtRanBP1c plays key roles in delivery of nuclear proteins responsible for suppression of auxin action and regulation mitosis in root tips. However, direct evidence is still required to clarify the role(s) of Ran GTPases and RanBPs in plant growth and development.

The mammalian RanGEF, RCC1, contains seven tandem repeats of a 50- to 60-amino acid domain, and these repeats constitute the majority of the protein. Arabidopsis contains 18 proteins containing one to five RCC1 domains (see supplemental data). One of these, *uvr8*, is a mutant hypersensitive to UV-B (Kliebenstein et al., 2002). The UVR8 protein contains five RCC1 repeats but interestingly does not possess nuclear localization sequences conserved in animal and yeast RCC1 homologs (Kliebenstein et al., 2002). Further work is necessary to determine whether UVR8, or other Arabidopsis proteins with RCC1 domains act as RanGEFs.

Two RanGAP sequences have been identified in Arabidopsis: AtRanGAP1 (accession no. AF2214559) and AtRanGAP2 (accession no. AF214560). Both AtRanGAPs complemented yeast RanGAP mutants (Pay et al., 2002), suggesting that these proteins are functional orthologs of the yeast RanGAP. AtRanGAP-GFP fusions associate with the nuclear envelope, and this localization is dependent upon a unique N-terminal domain (Rose and Meier, 2001). AtRanGAP1 localization is consistent with a role for AtRAN GTPases in nucleocytoplasmic transport. However, whether plant Ran GTPases function to regulate these transport steps or different plant Ran GTPases, especially the divergent AtRAN4, have distinct functions remains undetermined.

CONCLUSIONS

From these studies and observations, it is clear that small GTPases represent an important and diverse set of regulatory molecules in plants, with Arabidopsis containing members of the Rab, Rho, Arf, and Ran classes of small GTPases. In many cases, plant small GTPases appear to maintain similar functions as their animal and yeast counterparts. The identification of additional families of putative GAP and GEF proteins for most of these families further highlights the highly conserved nature of these regulatory molecules throughout evolution of eukaryotes. However, plants contain some unique variations on this conserved regulatory mechanism.

Although small GTPases related to Rab, Rho, Arf and Ran GTPases were identified in Arabidopsis, Ras

GTPases were not detected. This, in principal, concurs with the evolution of eukaryotic regulatory systems. The notable absence of plant Ras GTPases correlates with lack of Tyr kinase receptors, which act upstream of Ras signaling in animals. Plant Rop GTPases segregate as a distinct subfamily within the Rho GTPase family. While maintaining conserved functions in regulation of actin cytoskeleton and cellular signaling pathways, newly established roles in ABA-signaling events and interactions with plant RLKs highlight potential plant-specific variations of the functions of these Rop GTPases. Plant Rab GTPases segregate into distinct subfamilies that appear to be organized around the types of compartments upon which they are localized. Increased numbers of Arabidopsis Rab GTPases in subfamilies with predicted functions in vacuolar and post-Golgi trafficking may indicate plant-specific elaborations of these pathways. Plant Arf GTPases and Ran GTPases functionally complement mutations of their respective counterparts in yeast. However, plants contain multiple, novel ArfGEFs and ArfGAPs, probably with novel functions. Also, AtRAN4 is significantly different in sequences conserved throughout Ran GTPases from plants, yeast, and animals, perhaps indicating novel roles during plant growth and development. A wealth of knowledge of the mechanisms of action of small GTPases and of some of their important activator proteins (GEFs) and inhibitor proteins (GAPs) will clearly serve as a strong basis for understanding how these conserved regulatory mechanisms have been modified to carry out plant-specific processes.

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