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RESEARCH PAPER



Comparative phylogenetic and expression analysis of small GTPases families in legume and non-legume plants

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ABSTRACT

Background: Small monomeric GTPases act as molecular switches in several processes that involve polar cell growth, participating mainly in vesicle trafficking and cytoskeleton rearrangements. This gene superfamily has largely expanded in plants through evolution as compared with other Kingdoms, leading to the suggestion that members of each subfamily might have acquired new functions associated to plant-specific processes. Legume plants engage in a nitrogen-fixing symbiotic interaction with rhizobia in a process that involves polar growth processes associated with the infection throughout the root hair. To get insight into the evolution of small GTPases associated with this process, we use a comparative genomic approach to establish differences in the Ras GTPase superfamily between legume and non-legume plants.

Results: Phylogenetic analyses did not show clear differences in the organization of the different subfamilies of small GTPases between plants that engage or not in nodule symbiosis. Protein alignments revealed a strong conservation at the sequence level of small GTPases previously linked to nodulation by functional genetics. Interestingly, one Rab and three Rop proteins showed conserved amino acid substitutions in legumes, but these changes do not alter the predicted conformational structure of these proteins. Although the steady-state levels of most small GTPases do not change in response to rhizobia, we identified a subset of Rab, Rop and Arf genes whose transcript levels are modulated during the symbiotic interaction, including their spatial distribution along the indeterminate nodule.

Conclusions: This study provides a comprehensive study of the small GTPase superfamily in several plant species. The genetic program associated to root nodule symbiosis includes small GTPases to fulfill specific functions during infection and formation of the symbiosomes. These GTPases seems to have been recruited from members that were already present in common ancestors with plants as distant as monocots since we failed to detect asymmetric evolution in any of the subfamily trees. Expression analyses identified a number of legume members that can have undergone neo- or sub-functionalization associated to the spatio-temporal transcriptional control during the onset of the symbiotic interaction.

ARTICLE HISTORY

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KEYWORDS

Arf; biological nitrogen fixation; comparative genomics; Rab; Rop; symbiosis

Background

Most legume plants have the capacity to establish an intracellular symbiotic interaction with nitrogen-fixing soil bacteria called rhizobia. In this association, bacteria are accommodated inside the cell, surrounded by a host-derived membrane in a subcellular structure specialized in nitrogen fixation. This interaction, known as root nodule symbiosis (RNS), is present only in legumes, with the exception of a small tropical tree of the genus *Parasponia*. Some of the molecular components required to establish this symbiotic interaction apparently evolved by recruitment of pre-existing components from a more ancient interaction between plants and fungi from the phylum Glomeromycota, known as arbuscular mycorrhiza symbiosis (AMS).^{2,3} In this interaction, fungi are accommodated inside the root cortical cells in structures called arbuscules. The genetic program shared

between RNS and AMS is known as the common Sym pathway and its components participate mainly in the molecular changes that occur during infection. On the other hand, RNS involves the formation of a new organ, the nodule, whereas there is not an organogenesis counterpart in AMS. In addition, the recognition of signals derived from the bacteria or the fungi in these two interactions is mediated by different receptors (with the exception of Parasponia), whereas some of the transcription factors required for the establishment of the interaction are also specific for each type of symbiosis. In RNS, the most common infection mechanism begins with the redirectioning of the polar growth of the root hair, which curls to surround the bacterial microcolony, forming an infection focus.⁴ From this point, a tubular structure, known as the infection thread, grows inward

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• Supplemental data for this article can be accessed on the publisher's website.

the root hair, guiding the bacteria through the epidermis towards the dividing cortical cells, where bacteria will be released, surrounded by a host-derived membrane and differentiated to bacteroids to form a structure called the symbiosome. ^{5,6} Several molecular components associated with bacterial infection and release have been identified in the last years, including vesicle trafficking components, 7-12 cytoskeleton related-proteins. 13-16 and cell-wall degrading enzymes. 17 Among proteins involved in vesicle trafficking are the small GTPases, some of which play critical roles in early events of the infection process and in the signaling pathway required for nodulation, 9-11 but also at later stages during symbiosome maturation.^{7,8} In a more general context, small GTPases act as molecular binary switches, fluctuating between an activated state when they are bound to GTP and an inactive state bound to GDP. 18 They participate in multiple cellular processes across all eukaryotic organisms, being one of the most important groups of regulatory proteins. The superfamily of small GTPases has been divided into five families based on sequence identity.¹⁹ Ras GTPases were the first members identified in viruses that can lead to cancer (hence the name Rat sarcoma),²⁰ giving origin to the name that has been adopted for the family and sometimes for the superfamily as well. Ras proteins are part of the signal transduction pathways that regulate cell proliferation in mammals and yeast, but they are absent from plant genomes.²¹ Proteins of the two other families, Rab (Ras-related in brain) and Arf (ADP ribosylation factor), are involved in the four steps of vesicle trafficking in eukaryotic cells: budding of vesicles, transport, tethering and membrane fusion. 22,23 Members of the Rho (Ras homologous) family, known as Rop in plants (Rho of plants), participate in cytoskeleton dynamics and remodeling in response to external stimuli.²⁴ Finally, members of the Ran (Ras-related nuclear protein) family regulate nuclear-cytoplasmic trafficking in both directions across the nuclear pore.25

In the last years, advances in sequencing techniques have facilitated the generation of genomic and transcriptomic datasets, allowing the comparison between species in order to infer the evolutionary history of genomes. The limitation of the RNS to a group of phylogenetically related species, mainly legumes, provides a good opportunity to use comparative genomics and meta-analysis of transcriptomes to understand how different components of the genetic programs have evolved from common ancestors with non-nodulating species, giving origin to new functions in the context of this new biological process. We focused on the small GTPase superfamily, whose members seem to have evolved from functioning in other biological processes to acquire specific functions during the infection process initiated by rhizobia.26 Based on a previous analysis of expressed sequence tags (ESTs), it was suggested that some clades of the Ras superfamily of small GTPases have expanded and undergone neofunctionalization in legumes as a consequence of the new functions required for the establishment of the nitrogen-fixing symbiosis.^{27,28} In order to re-evaluate this hypothesis, we took advantage of the genomic and transcriptomic datasets recently generated in different species to conduct a comprehensive comparison of the small GTPase gene families and expression patterns of their members in nodulating and non-nodulating species. Our results support the notion that sub- or neo-functionalization are not associated with asymmetric evolution of the gene families or subfamilies, but with the spatio-temporal regulation of gene expression of specific members during the establishment of the symbiotic interaction.

Results and discussion

Phylogenetic analysis of small GTPases in legume and non-legume plants

In order to perform a comprehensive comparison of small GTPases subfamilies between legume and non-legume plants, we selected a group of species whose genomes have been sequenced and transcriptomic datasets are publicly available. Among legumes, the two model plants Lotus japonicus and Medicago truncatula (both associated with forage species) and the two grain legumes common bean (Phaseolus vulgaris) and soybean (Glycine max) were selected. As representatives of nonlegumes we included the dicotyledonous Arabidopsis thaliana and tomato (Solanum lycopersicum) and the monocotyledonous rice (Oryza sativa) and maize (Zea mays). Members of the small GTPases superfamily were identified through tBLASTn searches using the amino acid sequence of the monomeric GTPases annotated and classified in Arabidopsis. 21-29 as queries and the genome database of each species. Amino acid sequences of these proteins were used to generate phylogenetic trees, allowing their classification in different families and subfamilies (Fig. 1 and Figs. S1-8). As previously described in Arabidopsis, members of Rab, Arf, Rop and Ran families were identified in all species; however, genes encoding proteins of the Ras family were not identified in any of the plant genomes explored in this work (Figs. S1-8).²¹ No significant differences in the number of members of each family were observed between legume and nonlegume species (Table 1). The higher number of soybean GTPases compared with other species is most likely related to the partially diploidized tetraploid nature of its genome.³⁰ As each family is subdivided into subfamilies in Arabidopsis, we classified small GTPases from the other selected species using phylogenetic trees that included Arabidopsis members that are representative of each subfamily (Figs. S9-24). Again, we did not observe appreciable differences in the number of genes constituting each subfamily of the monomeric GTPases between the two groups of plants, legume and non-legume species (Tables 2 & 3). Taken together, our phylogenetic analysis in different species did not reveal asymmetric evolution of the small GTPase subfamilies in legume plants associated with the evolution of nitrogen-fixing symbiosis.

Small GTPases involved in nodulation are structurally conserved among legume and non-legumes

We next wondered whether differences in the amino acid sequence could be linked to specialized functions of small GTPases in the context of physiological changes that occur during rhizobia infection. As described in the introduction, several small GTPases participate in the establishment of symbiosis between legumes and rhizobia. These proteins were used to identify members from other species that were evolutionary related based on phylogenetic analysis (Figs. S25-S29) and

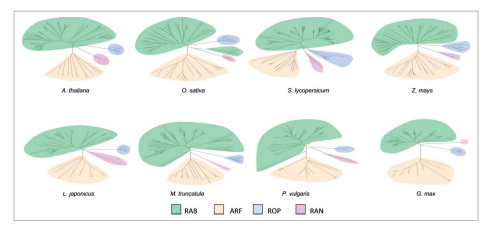


Figure 1. Phylogenetic analysis of the small GTPase superfamily in legume and non-legume plants. Amino acid sequences corresponding to small GTPases from *A thaliana*, *O. sativa*, *S. lycopersicum*, *Z. mays*, *L. japonicus*, *M. truncatula*, *P. vulgaris*, and *G. max* were retrieved from genomic databases. Unrooted neighbor-joining trees were obtained using the Mega 7 software. Subfamilies were identified for each species: Rab (green), Arf (orange), Rop (blue) and Ran (pink).

sequence identity. We selected PvRabA2 from common bean, MtRab7 (corresponding to a Rab of the G3 group according to the nomenclature suggested by Vernoud et al., 2003,²¹ MtRop9 and MtRop10 from M. truncatula and LjRop6 from L. japonicus. ArfA1 was also included considering its function in root growth³¹ and RNS (Dalla Via and Blanco, unpublished results). Multiple sequence alignments were generated for each of these proteins (Figs. 2-4 and Figs. S30-32). All alignments showed strong amino acid sequence conservation across the different species analyzed here, suggesting that Rab, Arf and Rop proteins were subjected to strong selective pressure. No differences in amino acid sequences were observed in ArfA1 (Fig. S30) and Rab7 (Fig. S31) that could be correlated with legume and nonlegume species. PvRabA2 and their putative ortholog genes contain a conserved substitution: whereas a valine residue is in the position 177 in the four legumes, isoleucine is present in that position in the four non-legume species (Fig. 2). Interestingly, three positions show no conservative substitutions in ROP9 proteins (amino acids 53, 129 and 151 are isoleucine, cysteine and asparagine in legumes and threonine, phenylalanine and glycine in non-legumes, respectively). The first substitution is at the end of the effector domain (domain II), whereas the second one affects the RHO insert region (domain V) (Fig. 3). Rop6 is in the same clade that MtRop9 and their amino acid sequences are very similar, but it shows the same single substitution in position 151 that was observed for Rop9 (Fig. S32). Rop10 also contains substitutions in legume species (Fig. 4), but in this case all changes are conservative and only the substitution of lysine in non-legumes by arginine in legumes affects a conserved region.

Since the conservation of amino acids across the eight studied species -including monocots and dicots- is an indication of a strong selective pressure, variation of some of these conserved residues in LjRop6, MtRop10, MtRop9 and PvRabA2 in legumes might reflect specialized functions of these proteins in the symbiotic interaction of legumes with rhizobia. However, the comparison of the predicted three dimensional conformations of RabA2, ROP9 and ROP10 suggests that the structure of the proteins is not strongly modified by these substitutions (Fig. 5). Nevertheless, functional analysis would be required to determine if the observed polymorphisms at the sequence level are associated with neo- or sub-functionalization in legume plants.

Expression of small GTPases in different tissues

Another possible difference between proteins of the small GTPase families among different species is the spatio-temporal control of gene expression of their members. Using publicly available datasets for each species, we retrieved expression data of each member of small GTPase families (Additional files 2–9; Tables S1-8). Values were normalized using the root sample as a reference and compared to other tissues. Considering the diversity of tissues selected in each study and the biological conditions and/or treatments applied, we selected samples that could be systematically compared among the eight species. In all species, legumes and non-legumes, GTPases of the four families were mainly accumulated at higher levels in roots as compared with aerial tissue (stem and leaf were considered in most cases) or had similar levels in different tissues (Additional files 2–9; Tables S1-

Table 1. Number of members of each small GTPases subfamily.

		Non-le	gumes	Legumes						
Sub-family	Arabidopsis thaliana	Oryza sativa	Solanum lycopersicum	Zea mays	Lotus japonicus	Medicago truncatula	Phaseolus vulgaris	Glycine max		
RAB	57	37	56	53	30	64	50	94		
ARF	21	21	23	25	13	19	20	41		
ROP	11	8	10	9	8	7	11	20		
RAN	4	2	4	3	2	4	3	7		
Total	93	68	93	90	53	94	84	162		

Table 2. Number of members of the small GTPases Rab subfamily.

		Non-le	gumes	Legumes					
Group	Arabidopsis thaliana	Oryza sativa	Solanum lycopersicum	Zea mays	Lotus japonicus	Medicago truncatula	Phaseolus vulgaris	Glycine max	
A	26	17	26	23	12	23	23	41	
В	3	3	5	4	2	7	1	4	
C	3	0	4	3	4	6	5	11	
D	4	4	5	6	1	4	4	7	
E	5	3	5	5	3	6	5	8	
F	3	4	4	3	3	5	4	7	
G	8	4	4	5	3	9	4	8	
Н	5	2	3	4	2	4	4	8	
Total	57	37	56	53	30	64	50	94	

8). With the exception of some Rops from common bean and soybean, very few small GTPases showed higher levels of mRNA in aerial tissue as compared with roots (Table 4). The distribution of small GTPases with differential expression in root or aerial tissue was not associated with a particular group of Rabs, Arfs or Rops in Arabidopsis (Fig. S1). The spatial expression pattern of small GTPases points toward predominant specific roles in polar and transport processes specific of the root, but the comparison of legume and non-legume plants suggests that these roles are present in all species and probably involve molecular mechanisms that have evolved before the legume family diverged from other groups of plants.

Next, we analyzed the expression of small GTPases in nodules of the four legume species. A small number of Rab and Arf members accumulated at higher levels in nodules as compared to roots. This number was comparable among the four legumes, but lower than the number of small GTPases that exhibited higher transcript levels in roots than in nodules. A similar pattern was observed in Rop mRNA accumulation (Table 5). The small GTPases that were up- or down-regulated in roots as compared to nodules do not cluster together in any particular branch of the phylogenetic trees of L. japonicus, M. truncatula or P. vulgaris (Figs. S5-7), suggesting that regulation of their expression in response to rhizobia has evolved independently in different members of each gene family. To validate the expression of a group of selected small GTPases we conducted reverse transcription reactions followed by quantitative PCR (RTqPCR) in M. truncatula roots and nodules. We selected one ROP (Medtr4g088055), one Arf (Medtr5g016540) and two Rab proteins belonging to the A and C families (Medtr2g090865 and Medtr1g062760, respectively). Consistent with the microarray data, our RT-qPCR data showed that mRNA levels of these genes were higher in roots than in nodules for Medtr4g088055, Medtr5g016540 and Medtr1g062760, whereas an opposite expression pattern was observed for Medtr2g090865 (Fig. 6). These results showed that the expression data obtain by high throughput analysis could be reproduced using a more reliable technique for all the genes analyzed in M. truncatula.

Considering that infections are initiated in the root, before the cortical divisions give origin to the nodule, we compared inoculated with uninoculated roots at very early stages of the interaction (i.e., short time periods after rhizobia inoculation). Surprisingly, only Phvul.004G011900, one member of the Arf family showed a significant difference in the expression level in common bean, with higher transcript levels at 24 hours post-inoculation with rhizobia (Additional file 8; Table S7). Likewise, none of the small GTPases was significantly regulated in roots of L. japonicus at 24 hours after infection with rhizobia (Additional file 6; Table S5) or in M. truncatula root hairs at early stages of the interaction (1, 3 and 5 dpi) when roots inoculated with a wild type strain were compared with roots inoculated with a non-fixing strain of Sinorrhizobium meliloti³² (Additional file 7; Table S6). These results suggest that, besides the biological important functions played by the small GTPases during early stages of legume-rhizobia symbiosis, these molecular switches are not strongly regulated at transcriptional level and/or at the level of their mRNA stability.

Spatial distribution of small GTPases transcripts in indeterminate nodules

Indeterminate nodules are divided into developmental zones according to the specific functions played by the different cell types found within the organ.⁶ We took advantage of the lasercapture microdissection strategy applied to dissect the transcriptome of different zones of M. truncatula indeterminate nodules to characterize the spatial distribution of transcripts

Table 3. Number of members of the small GTPases Arf subfamily.

		Non-le	gumes		Legumes						
Group	Arabidopsis thaliana	Oryza sativa	Solanum lycopersicum	Zea mays	Lotus japonicus	Medicago truncatula	Phaseolus vulgaris	Glycine max			
A	6	6	5	6	4	5	4	10			
$B+C+D^a$	6	6	8	9	4	7	6	12			
ARLA	4	2	4	4	1	3	4	5			
ARLB	1	2	1	2	1	0	1	2			
ARLC	1	1	1	1	1	1	1	2			
SARA	3	4	4	3	2	3	4	10			
Total	21	21	23	25	13	19	20	41			

^aGroups B, C and D of ARF were associated since it was difficult to identify the clades as described for Arabidopsis.

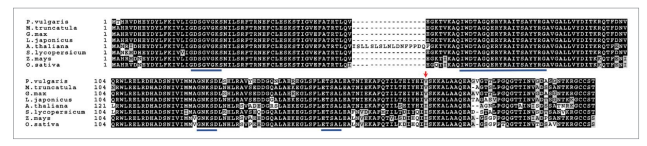


Figure 2. Multiple sequence alignment of P. vulgaris RABA2 (Phvul.011G061100) and proteins with the highest sequence identity from M. truncatula (Medtr4g064897), G. max (Glyma11g14360), L. japonicus (chr3.CM0792.300.r2.d), A. thaliana (At1g07410), S. lycopersicum (Solyc06g076450), Z. mays (GRMZM2G473906) and O. sativa (LOC_Os03g60870). Black boxes indicate identical residues and gray ones indicate conservative substitutions. Alignments were generated with Clustal Omega in MEGA7 and formatted with Boxshade. The red arrow indicates a conservative amino acid substitution in legume versus non-legume sequences. The conserved domains of Rabs are indicated by blue lines.

encoding each member of the small GTPase family.³³ In that study, different zones from the mature nodule were excised using a laser microdissection microscope and RNA from each nodule zone was subjected to transcriptome analysis. Based on clustering analysis, 13 expression patterns were defined according to mRNA abundance in the meristematic (FI), the infection zone (FII), the interzone (IZ) or the fixation zone (FIII). These data allow us to study the transcript levels of individual members of the small GTPase family in the context of the nodule organogenesis (Additional file 10; Table S9). Several members of Rab, Arf, Rop and Ran subfamilies showed differential accumulation of mRNA across the different nodule zones (Table 6). Clusters 1 to 5 showed a high relative transcript abundance in the meristematic and infection zones, whereas clusters 6, 11, 12 exhibited higher expression around the fixation zone. Interestingly, four Rab genes were grouped in cluster 1 and 2, showing a strong expression in the meristem (Medtr8g027220) or the meristem and the pre-infection zone (FIId; Medtr2g075950, Medtr3g086980 and Medtr7g051940). Only one Rop showed enhanced expression in the meristem (Medtr6g087980), whereas no Arf proteins were grouped in these clusters. Expression patterns of these genes suggest that the encoded small GTPase can play roles in nodule meristem activity/maintenance or at very early events of the infection process. On the other hand, several Rabs, Rops and Arf were grouped in the clusters 11 and 12, showing higher accumulation of their mRNAs in the fixation zone, where bacteria are already differentiated inside the symbiosomes (Table 6). Five Rabs and one Arf were included in cluster 13, which includes genes that are expressed in all nodule zones, but preferentially in the fixation zone. GTPases encoded by genes from cluster 11, 12 and 13

could be involved in the maintenance of symbiosome membrane identity or the exchange of metabolites across the symbiosome membrane between the macro- and the microsymbionts. Taken together, these results revealed that the expression of distinct small GTPases is differentially regulated at the spatial level in nodule zones that exert highly specialized functions during root nodule symbiosis and suggest that symbiotic functions of these proteins can be associated with their expression patterns.

Concluding remarks

The development of comparative genomics over the past years has contributed to understand functionalization of genes in an evolutionary context. In this work, we aimed to unveil how members of the small GTPase superfamily have acquired specific functions in the legume family associated with the unique capacity to establish RNS. Our analyses suggest that the number of family members and the primary sequence of small GTPases are well conserved between legume and non-legume plants. More importantly, small GTPases that have been linked to the symbiotic association by genetic approaches have highly conserved homolog genes in non-legume plants. However, we were able to identify specific amino acid residues in some of these GTPases that have changed during the course of evolution, beside the extreme amino acid conservation among genetically distant species. It will be of great interest to explore the functional relevance of these substitutions in the context of the symbiotic interaction established between legumes and rhizobia. Another interesting aspect of our analysis is

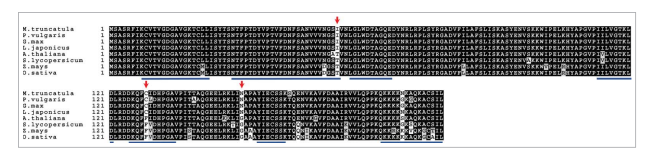


Figure 3. Multiple sequence alignment of M. truncatula ROP9 (Medtr5g022600) and proteins with the highest sequence identity from, P. vulgaris (Phvul.002G106600), G. max (Glyma01g36880), L. japonicus ROP6 (chr2.CM0272.860.r2.m), A. thaliana (At2g17800), S. lycopersicum (Solyc02g083580), Z. mays (GRMZM2G375002) and O. sativa (LOC_Os02g58730). Black boxes indicate identical residues and gray ones indicate conservative substitutions. Alignments were generated with Clustal Omega in MEGA7 and formatted with Boxshade. Red arrows indicate amino acid substitutions in legumes versus non-legumes. The conserved domains of ROPs are indicated by blue lines.



Figure 4. Multiple sequence alignment of *M. truncatula* ROP10 (Medtr3g078260) and proteins with the highest sequence identity from *P. vulgaris* (Phvul.009G180800), *G. max* (Glyma04g35110), *L. japonicus* (chr1.CM0166.830.r2.m), *A. thaliana* (At3g48040), *S. lycopersicum* (Solyc03g114070), *Z. mays* (GRMZM2G415327) and *O. sativa* (LOC_Os02g50860). Black boxes indicate identical residues and gray ones indicate conservative substitutions. Alignments were generated with Clustal Omega in MEGA7 and formatted with Boxshade. Red arrows indicate amino acid substitutions in legumes versus non-legumes. The conserved domains of ROPs are indicated by blue lines. The sequence from *Z. mays* is truncated at its C terminus.

the identification of specific members of each of the Rab, Arf and Rop families that show changes in the steady-state level of their mRNAs during the onset of the symbiotic interaction. Spatial information provided by laser-capture microdissection allowed us to identify several members that are also regulated along the different zones of the indeterminate nodule, suggesting that these small GTPases could have acquired specific functions that operate during infec-

tion, progression of the IT, symbiosome release or allocation and maintenance of nitrogen-fixing symbiosomes. In addition to provide insight into these interesting aspects of the symbiotic interaction, we believe this study presents a comprehensive compendium of a relevant and complex gene family in eight different species, establishing the basis for the study of the diverse biological processes in which small GTPases could be involved.

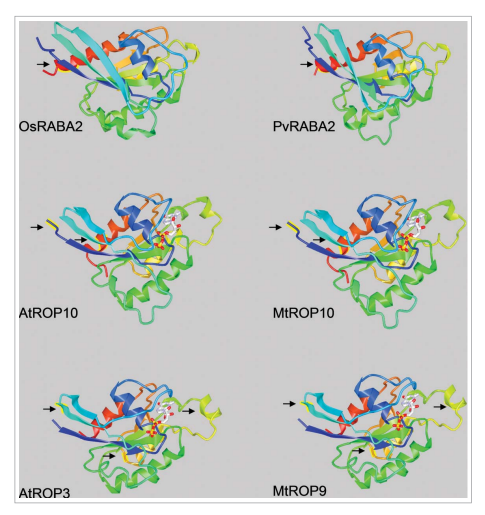


Figure 5. Three dimensional models of RABA2 from *O. sativa* and *P. vulgaris*, ROP10 from *A. thaliana* and *M. truncatula*, ROP3 from *A. thaliana* and ROP9 from *M. truncatula*. Arrows and yellow boxes indicate the position of the substitutions observed in legume versus non-legume species.

Table 4. Number of each subfamily members that shows no changes (0), reduced (-) or increased (+) levels of mRNA in aerial tissue compared with the root.

	Rab				Arf			Rop			Ran		
	_	+	0	_	+	0	_	+	0	_	+	0	
A. thaliana ^a	16	5	36	6	2	13	4	0	7	0	0	4	
O. sativa ^b	14	2	20	4	1	16	2	2	4	0	1	1	
S. lycopersicum ^a	15	1	34	5	2	14	5	0	5	1	0	3	
Z. mays ^c	17	0	34	5	1	18	0	1	8	0	0	3	
L. japonicus ^d	1	0	29	0	0	13	2	0	6	0	1	1	
M. truncatula ^a	17	2	26	5	0	13	2	0	5	2	0	2	
P. vulgaris ^a	14	3	28	5	0	13	1	4	6	1	0	2	
G. max ^a	50	2	42	19	2	20	5	6	9	4	0	3	

^acompared to leaf.

Methods

Identification of small GTPases from different species

Members of the small GTPases superfamily were identified through tBLASTn searches³⁴ using the amino acid sequence of all members of the small GTPase family previously described and classified in Arabidopsis^{21,29} as queries and the genomic dataset of each species (see Availability of data and materials). The list of identified genes were manually curated and classified according to our phylogenetic analyses (see below).

Phylogenetic analysis

Amino acid sequences were aligned using Clustal Omega http://www.ebi.ac.uk/Tools/msa/clustalo³⁵ and the phylogenetic analysis was performed with the MEGA7 package³⁶ http://www.megasoftware.net using the neighbor-joining method³⁷ with 1000 trials to obtain bootstrap values. The evolutionary distances were computed using the number of differences method.³⁸ All positions containing gaps and missing data were eliminated from the dataset. The phylogenetic analysis is available in Figs. S1-29. For Figs. S25-29, the branches with bootstrap values lower than 29% were collapsed to simplify the trees. Alignment files are provided in Additional file 11.

Protein alignments

The sequences of small GTPases involved in the establishment of symbiosis between legumes and rhizobia were used to identify members from other species using BLASTP. We selected PvRabA2 from common bean, MtRab7, MtRop9 and MtRop10 from *M. truncatula* and LjRop6 from *L. japonicus* and PvArfA1 as queries and the genome database of each species. The amino acid sequences with the lowest E value were used to generate a multiple sequence alignment.

Alignments were generated with Clustal Omega (http://www.ebi.ac.uk/Tools/msa/clustalo) in MEGA7 (http://www.megasoftware.net) and formatted with Boxshade (http://embnet.vital-it.ch/software/BOX_form.html).

3D modeling

Proteins were modeled using Swiss Model.³⁹ (https://swissmodel.expasy.org/) and analyzed using the 3D structure viewer iCn3D (https://www.ncbi.nlm.nih.gov/Structure/icn3d/full.html).

Genomic datasets

Sequences were obtained from public datasets: *Arabidopsis thaliana* TAIR10, *Oryza sativa* v7_JGI, *Zea mays* Ensembl-18, *Solanum lycopersicum* iTAG2.3, *Medicago truncatula* Mt4.0v1. *Glycine max* Wm82.a2.v1 and *Phaseolus vulgaris* v1.0 are available at Phytozome v11.0 (https://phytozome.jgi.doe.gov/pz/por tal.html) and *Lotus japonicus* v2.5, is available in Miyakogusa v2.5 (http://www.kazusa.or.jp/lotus).

Transcriptomic datasets

Gene expression of small GTPases was explored using publicly available databases.

Table 5. Number of each subfamily members that shows no changes (0), reduced (-) or increased (+) levels of mRNA in nodule tissue compared with the root.

	Rab				Arf			Rop			Ran		
	_	+	0	_	+	0	_	+	0	_	+	0	
L. japonicus ^a M. truncatula ^b	6 16	2	22 27	0	2	11 14	2	1	5	1	0	1 3	
P. vulgaris ^c G. max ^d	17 52	6 7	25 35	6 27	3 2	10 12	4 8	3 4	2 8	1 3	0 2	2 2	

^a14 day-old nodules vs roots.

^bcompared to shoot 2w.

 $^{^{}c}$ compared to stem + SAM, only expression values of the primary transcript were considered.

dcompared to leaf 6 w 5 mM nitrate.

^b4, 10, 14 and 28 day-old nodules vs roots.

^c5 and 21 day-old fixing nodules vs roots.

^dNodules collected 20–25 days after inoculation vs roots.



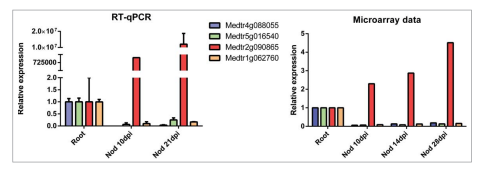


Figure 6. Comparison of expression data obtained by RT-qPCR with microarray data of selected M. truncatula genes encoding small GTPases. Expression levels of the four transcripts in roots and nodules (Nod) at 10 or 21 days post inoculation (dpi) with S. meliloti were measured by RT-qPCR or by microarray analysis. 48 Bars in qPCR graphs represent media and SE of two biological replicates. Expression levels were normalized with HIS3L and presented relative to the values of root tissue, which was set at 1.

- A. thaliana: expression data were obtained from the Transcriptome Variation Analysis database (TraVA, http://travadb. org. 40 That study analyzed RNA-seq data from 79 samples, each with two biological replicates, corresponding to different

Table 6. Expression of M. truncatula GTPases in different regions of the mature

nodule	•							
Туре	Gene ID	Cluster	Reads	FI ^b	FIId	FIIp	ΙZ	ZIII
RAB	Medtr8g027220	1	487.0	93.3°	5.9	0.8	0.0	0.0
RAB	Medtr2g075950	2	4680.8	52.9	25.6	7.1	8.6	5.8
RAB	Medtr3g086980	2	1248.4	66.1	17.0	13.3	2.2	1.6
RAB	Medtr7g051940	2	857.9	50.5	35.4	9.4	2.6	2.0
RAB	Medtr2g005510	4	1068.4	29.6	38.2	11.5	12.4	8.2
RAB	Medtr1g090717	4	311.0	36.0	37.8	5.4	8.0	12.8
RAB	Medtr5g013900	5	1873.2	23.3	24.6	28.0	9.8	14.2
RAB	Medtr4g079350	5	554.1	26.9	28.6	30.1	9.7	4.8
RAB	Medtr4g088090	5	1845.6	33.1	26.1	15.3	17.6	7.8
RAB	Medtr4g063160	5	11458.1	24.6	32.1	17.1	12.6	13.6
RAB	Medtr8g090215	6	2000.7	2.4	3.1	8.3	50.6	35.6
RAB	Medtr2g005500	11	2508.5	10.6	22.0	15.0	31.3	21.2
RAB	Medtr3g064390	11	9915.2	9.0	14.4	26.2	29.2	21.2
RAB	Medtr2g090865	11	4080.8	13.1	25.5	22.6	18.8	19.9
RAB	Medtr4g099470	11	16424.1	8.0	15.3	25.0	27.2	24.5
RAB	Medtr2g101890	11	1673.9	9.3	14.0	18.7	37.8	20.1
RAB	Medtr4g012940	11	4250.8	9.6	12.6	23.9	34.7	19.3
RAB	Medtr8g093740	11	537.4	6.0	8.5	32.2	38.1	15.2
RAB	Medtr8g089985	11	10940.8	4.0	20.4	26.0	29.7	19.9
RAB	Medtr2g027660	12	4490.6	3.7	7.9	21.0	34.5	32.9
RAB	Medtr8g006970	13	20837.8	7.1	14.5	17.5	32.1	28.7
RAB	Medtr7g103070	13	1483.5	12.8	13.0	12.9	23.4	38.0
RAB	Medtr4g069850	13	15666.8	8.8	12.8	18.6	35.2	24.5
RAB	Medtr7g081700	13	25501.2	12.3	14.4	19.2	29.8	24.4
RAB	Medtr5g076590	13	5298.2	6.1	11.1	19.4	32.3	31.2
ARF	Medtr5g007810	5	1566.1	20.7	33.8	17.7	11.1	16.7
ARF	Medtr2g034640	5	15828.0	18.5	23.2	23.0	22.7	12.6
ARF	Medtr4g077640	11	3825.1	10.1	16.5	23.2	28.2	21.9
ARF	Medtr4g121860	11	14082.7	2.9	6.1	30.6	41.2	19.2
ARF	Medtr7g109960	11	11655.0	11.5	16.3	22.0	35.0	15.2
ARF	Medtr6g005820	11	111069.1	10.0	21.2	25.3	22.5	21.2
ARF	Medtr1g076550	13	5677.5	13.2	20.0	15.7	27.5	23.6
ROP	Medtr6g087980	1	1104.4	89.3	8.9	0.4	0.3	1.0
ROP	Medtr4g088055	3	149.1	23.6	52.4	19.7	4.3	0.0
ROP	Medtr4g073250	11	9576.4	14.6	14.3	17.2	30.4	23.5
ROP	Medtr2g090875	12	5019.7	3.9	6.3	9.8	27.6	52.4
ROP	Medtr3g078260	13	1370.5	19.5	30.1	10.2	4.5	35.6
RAN	Medtr3g107707	5	6828.1	27.1	20.1	25.5	16.6	10.7
RAN	Medtr3g107713	5	43402.5	24.3	32.5	23.9	9.3	10.0

^aData were retrieved from the laser-capture microdissection analysis of indeterminate nodules from M. truncatula.33

developmental stages and parts of roots, leaves, flowers, seeds, siliques and stems from A. thaliana ecotype Col-0. Differential gene expression analysis between all possible pairs of samples were performed using DESeq. 41 A false discovery rate (FDR) of 0.05 and a fold change of 2.0 were chosen as the threshold for significantly differential expression.

- S. lycopersicum expression data were obtained from the Tomato Functional Genomics Database (http://ted.bti.cornell. edu).42 That study contains Illumina RNA-seq data from root, stem, leaf, flower and fruit at 3 different maturation stages: mature green, breaker and ripe of tomato cultivar Heinz. Two biological replicates were analyzed. Differentially expressed genes were identified using DESeq and raw p values of multiple tests were corrected using FDR.
- O. sativa: expression data were retrieved from. 43 That study includes gene expression data obtained using Illumina RNAseq technology from rice roots and shoots in different stages of Pi deprivation. Three biological replicates of Oriza sativa cv Nipponbare were used. Identification of differentially expressed genes was performed using Cuffdiff⁴⁴ to analyze the effects of Pi status on the transcriptome of roots and shoots. Additional expression data were obtained from, 45 which contains RNAseq data from nine distinct tissues including immature leaves, pre and post-emergence flowers, anther, pistil, whole seed at two different maturation stages, embryo, endosperm and biological replicates of rice cv. Nipponbare.
- Z. mays: expression data were obtained from RNAseq transcriptomic analyses⁴⁶ that include expression data from 18 tissues representing five organs. Samples were obtained from the reference inbred line B73. Three replicates were used for all the analyses. RNA-Seq expression data were compared with the previously published microarray based gene atlas.⁴⁷
- M. truncatula: data from The Medicago truncatula Gene Expression Atlas MtGEA (http://mtgea.noble.org/v3, 48) was used. The study was performed using the Affymetrix GeneChip Medicago Genome Array. Gene expression values were obtained from three independent biological replicates of each organ: roots, nodules, stems, petioles, leaf blades, vegetative buds, flowers, and seed pods of M. truncatula cv Jemalong A17. In addition, three stages of nodule and six stages of seed development were profiled. Hierarchical clustering analysis was conducted with Spotfire Decision-Site 8.1 (Spotfire Inc., http://spotfire.tibco.com/). For the nodule developmental series, transcript levels were expressed relative to the level in roots just prior to inoculation and clustered using the

^bNodule zones dissected in the study are: the nodule meristematic region, FI; distal and proximal adjacent regions to the meristem, FIID and FIIP, respectively; the interzone II-III, IZ; and the nitrogen-fixation zone, ZIII.

Values displayed on the table correspond to the mean percentage obtained from 3 biological replicates.

Pearson correlation coefficient. Additionally, we explored the transcriptome data of the laser-capture microdissection strategy applied to M. truncatula indeterminate nodules³³ to characterize of gene expression of the small GTPase family in different nodule zones. In addition, whole roots, whole nodules and aerial organs were profiled. Three biological repetitions were analyzed in all cases except for pooled aerial organs (one experiment). Based on clustering analysis, 13 expression patterns were defined according to mRNA abundance in the meristematic (FI), the infection zone (FII), the interzone (IZ) or the fixation zone (FIII). LCM methodology was validated by Roux et al with a set of 37 marker genes previously characterized by other methods.³³ Additional expression data were obtained from root hairs prior to and during the initial stages of infection with S. meliloti.³² Three biological replicates were performed for each treatment. Wild-type (Jemalong A17) or skl-1 mutant plants were used at different days post inoculation with Sinorhizobium meliloti or a S. meliloti non-fixing strain. A17 was also profiled at 1 d post-treatment following addition of Nod factors.

- L. japonicus: data were obtained from The Lotus japonicus Gene Expression Atlas LjGEA (http://ljgea.noble.org/v2, 49). This work presents an integrated genome-wide analysis of transcriptome landscapes of wild-type (ecotype Gifu) and symbiotic mutant plants using the Affymetrix GeneChip Lotus Array. Five different organs, five stages of the sequentially developed determinate root nodules, and eight mutants impaired at different stages of the symbiotic interaction were analyzed in triplicates from three spatially and temporally separated batches of plants. Significant genes were identified using the Limma package. 50 A FDR (corrected p-value < 0.05) was used as the criterion for significance, in combination with a $|M| \ge 1$ filter, where M is the log₂ ratio of average expression values from any two conditions.

- P. vulgaris: expression data were obtained from the Common Bean Gene Expression Atlas (http://plantgrn.noble.org/ PvGEA,⁵¹), which presents gene expression values obtained by RNAseq from 24 unique samples collected from seven distinct tissues of P. vulgaris cv. Negro Jamapa; roots, nodules, leaves, stems, flowers, seeds, and pods. Plants were inoculated with either effective or ineffective strains of rhizobia. Transcripts differentially expressed between libraries were identified using NOIseq.⁵² In that study, differential expression was confirmed by reverse transcription followed by quantitative PCR (RTqPCR) analyses for 92% of the 85 genes selected. Additional expression data were retrieved from our transcriptomic analysis of P. vulgaris cv. NAG12 plants inoculated with different Rhizobium etli strains at 24 hours.⁵³ Two biological replicates were used for RNAseq and differentially expressed genes were identified using Cuffdiff. RT-qPCR analysis of differentially expressed genes confirmed expression data obtained by RNAseq for nine out of 11 genes.⁵³

- Glycine max: data were obtained from the SoyBase Database (https://www.soybase.org/soyseq, 54) which includes RNA-Seq from seven tissues (leaf, flower, pod, two stages of podshell, root, nodule) and seven stages in seed development. General trends in expression profiles for all genes were examined by a comparison of the transcription count for every tissue to every other tissue using a Fisher 's exact test with a FDR correction of 0.05.

Expression data for all small GTPase members from each species are available in Additional files 2-10, Tables S1-9. The Join two Datasets tool available at Galaxy platform⁵⁵ was used to merge expression data from different datasets.

Validation of Expression Data by qRT-PCR

We selected a set of four differentially expressed genes corresponding to small GTPases (Medtr2g090865, Medtr5g016540, Medtr4g088055 and Medtr1g062760) to validate their expression using RT-qPCR and to compare with that previously obtained by microarray by Benedito et al.⁴⁸ RT-qPCR was performed essentially as previously described.⁵⁶ For each pair of primers (Additional file 12, Table S10), the presence of a unique PCR product of the expected size was verified in ethidium bromide-stained agarose gels. Absence of contaminant genomic DNA was confirmed in reactions with DNAse-treated RNA as template. Expression values were normalized to HIS3L, which has been validated by GNORM.⁵⁷ Two biological replicates were performed.

Statistical analysis of expression data

Using publicly available datasets for each species, we retrieved expression data of each member of small GTPases families. We selected roots as the reference organ and tested their expression in other organs for the eight species. The expression values were normalized using the root sample and the log₂ (fold change) was calculated for each tissue. For A. thaliana and G. max, genes were considered as differentially expressed according to the statistical analyses performed by the authors. 40,54 We analyzed the data retrieved from 43 for O. sativa and data from common bean using CuffDiff⁴⁴ to identify genes differentially expressed between root and shoot. Genes whose transcript levels showed at least a 2-fold change between samples (log₂ fold change <-1 or >1), a p value < 0.05 and an expression > 1 fragment per kilobase per million (FPKM) in at least one sample were considered as differentials.

For the others species, genes assigned with reduced (-) or increased (+) transcript level were those that exhibited at least a log₂ fold change <-1 or >1, respectively. Transcripts with -1<log₂ fold change<1 were considered as genes with no changes in gene expression (0). Only values with FPKM >1 or reads per kilobase per million reads (RPKM) >1 in at least one sample were considered.

In order to analyze the expression of small GTPases in the nodules of the four legume species, we compared their expression to root samples. For M. truncatula and G. max, the genes considered as differentially expressed were assigned according to the statistical analyses from the authors. 48,54 For L. japonicus and P. vulgaris, differentially expressed transcripts were those with at least a 2-fold change between root tissue and at least one sample of nodule tissue (\log_2 fold change >1 or <-1).

Abbreviations

3D Three dimensional

AMS arbuscular mycorrhiza symbiosis

ARF ADP ribosylation factor **EST** expressed sequence tag

GDP Guanosine-5'-diphosphate GTP Guanosine-5'-triphosphate NFR5 Nod Factor Receptor 5 RAB Ras-related in brain

RAN Ras-related nuclear protein

RAS Rat sarcoma RHO Ras homologous ROP Rho of plants

RNS root nodule symbiosis

vs versus

Declarations

Ethics approval and consent to participate: Not applicable Consent for publication: Not applicable Availability of data and materials:

Genomic datasets

Sequences were obtained from public datasets: *Arabidopsis thaliana* TAIR10, *Oryza sativa* v7_JGI, *Zea mays* Ensembl-18, *Solanum lycopersicum* iTAG2.3, *Medicago truncatula* Mt4.0v1, *Glycine max* Wm82.a2.v1 and *Phaseolus vulgaris* v1.0 are available on the Phytozome v11.0 (https://phytozome.jgi.doe.gov/pz/portal.html) and *Lotus japonicus* v2.5, is available on Miyakogusa v2.5 (http://www.kazusa.or.jp/lotus).

Transcriptomic datasets

Gene expression atlases were explored in order to characterize the transcriptional patterns of the *A. thaliana* (TraVA, Transcriptome Variation Analysis, http://travadb.org, 40), *S. lycopersicum* (Tomato Functional Genomics Database http://ted.bti.cornell.edu, 42), *M. truncatula* (The *Medicago truncatula* Gene Expression Atlas MtGEA http://mtgea.noble.org/v3, 48), *L. japonicus* (The *Lotus japonicus* Gene Expression Atlas LjGEA http://ljgea.noble.org/v2, 49), *P. vulgaris* (A Common Bean Gene Expression Atlas http://plantgrn.noble.org/PvGEA, 51) and *Glicine max* (SoyBase Database https://www.soybase.org/soyseq. 54) Additional expression data were retrieved for *O. sativa*. 43 *Z. mays*, 46 *P. vulgaris*. 53 and *M. truncatula*. 32,33 The expression data for each species is available in Additional files 2–10; Tables S1-9.

Competing interest

The authors declare that they have no competing interests.

Authors' contributions

FAB design research; AF, VDV, VS and UM collected data and performed the analysis. AF, MEZ and FAB wrote the article.

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