Involvement of SchRabGDI1 from *Solanum chilense* in endocytic trafficking and tolerance to salt stress

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**ABSTRACT**

Physiological responses of plants to salinity stress require the coordinated activation of many genes. A salt-induced gene was isolated from roots of the wild tomato species *Solanum chilense* and named SchRabGDI1 because it encodes a protein with high identity to GDP dissociation inhibitors of plants. These proteins are regulators of the RabGTPase cycle that play key roles in intracellular vesicular trafficking. The expression pattern of SchRabGDI1 showed an early up-regulation in roots and leaves under salt stress. Functional activity of SchRabGDI1 was shown by restoring the defective phenotype of the yeast sec19-1 mutant and the capacity of SchRabGDI1 to interact with RabGTPase was demonstrated through BiFC assays. Expression of SchRabGDI1 in *Arabidopsis thaliana* plants resulted in increased salt tolerance. Also, the root cells of transgenic plants showed an early up-regulation of SchRabGDI1 expression in response to salt. SchRabGDI1 expression correlated with the increased rate of endocytosis under normal growth conditions and higher accumulation of sodium in vacuoles and small vesicular structures under salt stress than wild type. Our results suggest that in salt tolerant species such as *S. chilense*, bulk endocytosis is one of the early mechanisms to avoid salt stress, which requires the concerted expression of regulatory genes involved in vesicular trafficking of the endocytic pathway.

**1. Introduction**

Membrane trafficking in eukaryotes depends on the accurate targeting of transport vesicles to and from defined membrane-bound compartments whereby different proteins participate in distinct steps of the process. Among the proteins involved in this vesicular trafficking is the Rab/Ypt family (RabGTPases), which form the largest branch of the Ras superfamily of the small GTPases that exist in all eukaryotic cells [1]. Different members of the RabGTPase family localized in the cytoplasmic side of organelles have been shown to have specific roles in targeting and/or tethering transport vesicles during exocytosis and endocytosis in eukaryotic cells [1,2]. RabGTPases function depends on their interaction with accessory proteins and their capacity to bind and hydrolyze GTP, which translates in the alternation between “active” and “inactive” states. RabGTPases carrying a geranylgeranyl group require guanine nucleotide exchange factors (GEFs) that facilitate GDP dissociation, GTPase activating proteins (GAP) that stimulate GTP hydrolysis, and guanine dissociation inhibitors (GDI) that form soluble complexes with small GTPases by shielding their lipid group. GDI proteins play a critical role in regulating the recycling of RabGTPases, allowing their rapid recycling [3] and maintaining a cytosolic pool of available RabGTPases to be delivered to vesicle membranes [4].

Although specific GDIs for Rab GTPases (RabGDIs) have been well characterized in yeast and animals, in plants, only few genes that share homology with members of the GDI family have been reported. Whereas 57 RabGTPases members have been identified in the *Arabidopsis thaliana* genome [5], only three RabGDIs homologues have been identified, AtRabGDI1, AtRabGDI2 and AtRabGDI3 [6–8]. Two cDNAs encoding RabGDI have been isolated from rice, OsGDI1 and OsGDI2 [9], one GDI has been cloned from tobacco [10] and chickpea [11], and three RabGDIs have been identified in grapevine [12]. Among all characterized RabGDI family members, five structurally and functionally sequence-conserved regions (SCRs) have been identified [4]. Mutations of residues within SCRs interrupt the binding of RabGDIs to RabGTPase proteins and may lead to a decreased RabGTPase recycling [13,14]. The ability of RabGDI to interact with distinct RabGTPases has...
been demonstrated, however different RabGTPase members present distinct binding affinity [15].

Plants display different mechanisms to tolerate salt stress including ion extrusion from the cell and ion sequestration in the vacuole to prevent excess accumulation in the cytosol [16,17]. The buildup of sodium into the vacuole is mediated by the action of Na⁺/H⁺ antiporters located in the tonoplast [18,19]. In addition, a rapid increase in the root vacuolar volume leading to increased vacuolar salt content has been shown under salt stress [20]. During this process, endosomes are fused to the main vacuole, suggesting that vesicle trafficking might play an important role in the response to salt stress [21]. Overexpression of the AtRabG3e gene (encodes a RabGTPase) in Arabidopsis triggered accelerated endocytosis in roots, leaves, and protoplasts and resulted in accumulation of sodium in vacuoles and increased tolerance to salt and osmotic stresses [22]. Similar results have been observed with the overexpression of an AtRabG3e homologous gene, PgRab7 from Pennisetum glaucum in tobacco [23]. Loss of function of RabA1 members involved in vesicle transport between the trans-Golgi network and the plasma membrane caused hypersensitivity to salt stress, most likely due to the participation of these proteins in the localization of cell-surface proteins, such as ion channels and pumps [24]. The involvement of RabGTPases in the response to salinity stress suggests that other interactive members controlling their function could also play a role in this response. Whereas one study has implicated an Arabidopsis RabGEF in mediating an endocytic pathway affecting stress tolerance [25], the role of RabGDIs in plant tolerance to salt stress has not been yet reported.

Cultivated species of the Solanaceae family are susceptible to a wide range of environmental stresses. For example, salinity is known to negatively affect seed germination, inhibit growth and decrease fruit productivity [26]. The Solanum section Lycopersicon includes S. chilense, a wild tomato species with a notable capacity to withstand salinity and drought [27]. When subjected to salt stress conditions, S. chilense activates the expression of a set of genes that may be associated with its capacity to adapt to its natural habitat [28–30]. Among the genes that are differentially expressed in salt treated S. chilense roots, we identified one encoding a protein with high homology to AtRABGD1, hence named SchRABGD1. Here, we analyzed the molecular function of SchRabGD1 and its ability to bind RabGTPases in vivo. Expression of SchRabGD1 in Arabidopsis thaliana resulted enhanced tolerance to salt stress and in an increase of both the endocytosis rate in root cells and vacuolar Na⁺ content upon high salt exposure. Our results suggest that salt-induced expression of SchRabGD1 contributes to endocytic trafficking in S. chilense and to its natural salt tolerance.

2. Material and methods

2.1. Plant materials and growth conditions

Solanum chilense (Dunal) seeds were obtained from plants collected in Northern Chile at a 2500 m.s.n.m. 18°26' lat. S 69° 45' long. Plants were clonally propagated in pots containing a mixture of perlite, vermiculite and peat moss (1:1:1 v/v) and grown under greenhouse conditions at 23–25 °C and a 16 h:8 h light:dark photoperiod. Plants were fertilized with commercial Hoagland’s solution (1/4 strength) every 10 days. For gene expression analyses under non-stressed conditions organ samples including root (R), young and mature leaves (YL, ML), stem (S), flower bud (FB) and flower (F) were taken at flowering time. Salt stress in S. chilense was applied to 7-week-old plants grown in 21 pots containing a mixture of perlite:vermiculite:peat moss (1:1:1 v/v) and fertilized with Hoagland solution by irrigating once with 400 ml of 300 mM NaCl. Leaves and roots samples were collected at 0, 3, 6, 12, 24, 48 and 72 h after salt treatment and immediately frozen in liquid nitrogen and stored at −80 °C.

Wild-type (Col-0) and transgenic Arabidopsis thaliana plants were grown in a chamber at 21 °C and a 16 h light/8 h dark photoperiod. For saline stress, 5-days-old Arabidopsis seedlings grown in solid half-strength MS (Murashige and Skoog, basal salt mixture) were transferred to half-strength MS medium containing 0 or 75 mM NaCl and maintained for 15 days. Then, biomass production (fresh weight) and leaf oxidative damage were evaluated. To assess germination rate under saline stress, seeds were surface sterilized with a 2% sodium hypochlorite solution. One-hundred seeds from wild-type and transgenic lines were sown in half-strength MS plates containing 0, 50 or 75 mM NaCl and kept at 4 °C in darkness for 3 days for seed stratification. Radicle emergence was examined every 24 h during 96 h. All treatments were done in three independent experiments.

2.2. RNA isolation

Total RNA was extracted from different organs of S. chilense and Arabidopsis leaves using the SV Total RNA Isolation System kit (Promega). All RNA extractions for gene expression assay were done in triplicate for each organ and condition. RNA integrity was visualized by 2% agarose gel electrophoresis and RNA concentration and purity (OD260/OD280 ratio > 1.95) were determined with a NanoDrop ND-1000 spectrophotometer (NanoDrop Technologies). RNA samples were treated with RNase free DNase I (Ambion) to remove contaminant DNA traces.

2.3. Subtractive library construction

The Clontech PCR-Select cDNA Subtraction kit was used for the preparation of a subtractive cDNA library to identify differentially expressed genes in roots of S. chilense under salt stress (half-strength MS supplemented with 400 mM NaCl) in hydropnic cultivation. Double-stranded cDNA synthesis was carried out on total mRNA derived from roots of stressed (tester) and normal (driver) plants. The tester and driver cDNAs were then digested with RsaI yielding blunt end fragments of approximately 400 bp length on average and processed following the manufacturer’s instructions with some modifications. The tester cDNA was aliquoted into two halves, and each half was ligated with different cDNA adaptors. Adapter ligation was followed by two rounds of hybridization with an excess of driver cDNA as per manufacturer’s protocol. The resultant products were subjected to two cycles of PCR with adaptor targeting primers to amplify the differentially expressed sequences. Amplifications were performed on a Stratagene Mx3000P (Agilent Technologies). First PCR master mix contained 10x PCR reaction buffer, 0.2 mM dNTPs, 0.4 μM PCR primer 1 and Advantage cDNA polymerase (Clontech). PCR was performed under the following conditions: 94 °C (25 s) followed by 30 cycles each consisting of a denaturation step at 94 °C (10 s), an annealing step at 66 °C (30 s) and an elongation step at 72 °C (1.5 min). Before using the primary PCR products as templates for secondary PCR, these were diluted 10-fold with sterile water. The second PCR master mix contained 10x PCR re- action buffer, 0.2 mM dNTPs, 0.4 μM nested PCR primer 1, 0.4 μM nested PCR primer 2 and Advantage cDNA polymerase. PCR was run through 20 cycles each consisting of 94 °C for 15 s, 66 °C for 30 s and 72 °C for 1.5 min. cDNA molecules were cloned non-directionally into the pGEM-T-Easy Vector (Promega). The ligation products were used to transform the E. coli DH5α strain via electroporation. Positive clones were collected and used for plasmid isolation and sequencing.

2.4. Analysis of gene expression

Gene transcript levels were analyzed by quantitative PCR (qRT-PCR) using a Stratagene Mx3000P (Agilent Technologies) system and the Brilliant SYBR Green Master Mix (Stratagene). To prepare first-strand cDNA, 2 μg of total RNA were reverse transcribed in a 20 μl reaction using the oligo d(T) and AffinityScript QPCR cDNA Synthesis Kit (Stratagene) following manufacturer’s instructions. For each sample (three biological replicates), qPCR was carried out in triplicate...
(technical repeats) using 10 μl Master Mix, 0.5 μl of 250 nM primers, 1 μl of diluted cDNA and nuclease-free water to a final volume of 20 μl. Amplification was followed by a melting curve analysis with continuous fluorescence acquisition during the 55–95 °C melt. The 2−ΔΔCt method was applied to calculate the fold change of gene transcript levels [30]. The tomato transcripts were normalized against tomato AtBox (accession X58253) and GAPDH (accession U97257) genes [29,32] and Arabidopsis transcripts were normalized against AtP-box (accession AT5G15710) [33]. The primers used for qPCR analysis are shown in supplementary Table S1. All qPCR products obtained from S. chilense were previously sequenced in order to corroborate primer specificity and gene identity.

2.5. Yeast complementation assay

For yeast complementation experiments using SchRabGDI1, the sec19-1 yeast mutant strain was employed. The Saccharomyces cerevisiae strains RSY274 (sec19-1) and its parental wild-type strain RSY249 were provided by R. Schekman (University of California, Berkeley, CA). The cDNA of SchRabGDI1 was PCR-amplified (primers described in Table S2) and cloned into the entry vector pENTR™/SD/D-TOPO™ (Invitrogen) to generate pENTR-SchRabGDI1 and verified by sequencing. pENTR-SchRabGDI1 was recombined by LR Clonase™ reaction (Invitrogen) into the destination vector pYES-dest52, under control of the GAL1 promoter. The resulting construct pYES-dest52-SchRabGDI1 and the empty vector pYES-dest52 were transformed into the sec19 mutant, whereas the wild-type strain was transformed with the empty vector. Yeast transformation was performed using the S. cerevisiae EasyComp™ Transformation kit (Thermo Fisher Scientific) and selection was done on Synthetic complete (SC) medium lacking uracil and supplemented with glucose to repress the GAL1 promoter. To evaluate the sec19 mutant phenotype and complementation capacity of SchRabGDI1, transformed yeast cells were grown in SC medium supplemented with galactose at 28 °C or 37 °C for 3 days.

2.6. Genetic construct and plant transformation

The coding sequence of SchRabGDI1 was amplified by PCR using the Platinum® Taq DNA Polymerase (Invitrogen) and cloned into pGEM-T vector (Promega) and sequenced (primers listed in Table S2). The PCR product was inserted into the XbaI-Sacl sites of the pBI121 binary vector to replace the β-glucuronidase (GUS) gene, resulting in the SchrabGDI1 gene being under the control of the CaMV 35S promoter. For yeast transformation, the SchRabGDI1 construct was recombined into pYES-dest52, under control of the GAL1 promoter. The resulting construct pYES-dest52-SchRabGDI1 and the empty vector pYES-dest52 were transformed into the sec19 mutant, whereas the wild-type strain was transformed with the empty vector. Yeast transformation was performed using the S. cerevisiae EasyComp™ Transformation kit (Thermo Fisher Scientific) and selection was done on Synthetic complete (SC) medium lacking uracil and supplemented with glucose to repress the GAL1 promoter. To evaluate the sec19 mutant phenotype and complementation capacity of SchRabGDI1, transformed yeast cells were grown in SC medium supplemented with galactose at 28 °C or 37 °C for 3 days.

2.7. Structure prediction by homology modeling

The structural models of SchRabGDI1 and SchRabG3e proteins were built by homology modeling based on crystal structures of homologous proteins. SWISS-MODEL [35] was used to select 3D models crystallized with the closest sequence homology and also to construct comparative model structures. The crystalline structure of a GDI from Saccharomyces cerevisiae (PDB code 1UKV; [36]) and Ypt7, a Rab7 class from S. cerevisiae (PDB code 4PHH; [37]) were selected based in the best homology to SchRabGDI1 and SchRabG3e, respectively. The best models for both proteins obtained by SWISS-MODEL were improved by molecular dynamic simulation and equilibration methods using Nano Molecular Dynamics (NAMDv.2.10; [38]), the Chemistry of Harvard Molecular Modeling (CHARMM27) force field [39] and the TIP3P model for water [40]. A short initial minimization of 15,000 steps was used to remove wrong contacts and for energy optimization. The molecular dynamics were done using the following conditions: 12 ns of molecular dynamics, a periodic bordering condition box (80 Å, 100 Å, 80 Å), 150 mM NaCl and 300 °K with default parameters [38]. The final 3D model of each structure was checked for its stereochemical quality and atomic coordinates with a Ramachandran map using PROCHECK [41].

2.8. Protein-protein docking

Protein-protein docking between SchRabGDI1 and SchRabG3e with GLUpro software [42] and structural alignment with 1UKV Rab-GDI tridimensional model of S. cerevisiae were used to determine the binding platform. Then, a molecular dynamic of 12 ns with NAMD was made to evaluate the interaction among amino acids of the binding platform and determine the free end of both proteins. This information was used to predict the location of the SCYCE and VYNE fluorescent proteins. The Python Molecular Viewer 1.4.5 [43] and the visual molecular dynamics (VMD; [44]) softwares were used for final visualizations.

2.9. In vivo interaction of SchRabGDI1 and SchRabG3e by BIFC

Full-length cDNA encoding SchRabGDI1 lacking the stop codon and SchRabG3e were amplified and cloned into the entry vector pDONR207 (Invitrogen, USA) to generate pENTR-GDI1 and pENTR-G3e, respectively (primers shown in Table S2). pENTR-GDI1 was recombined into pDEST-GWYNE [45], generating GDI-YWNE and pENTR-SchRabG3e into pDEST-SCYCE(R)GW [45] to generate SCYCE-SchRabG3e. The NIX5 construct used in validation experiments of the specific interaction of Rab with GDI was obtained used the same entry vector as Bassil et al. [46]. The resulting constructs were transformed into A. thaliana Col-0 and transformed bacteria were infiltrated into four-week-old leaves of tobacco (Nicotiana benthamiana). Infiltrated plants were grown in a growth chamber for 3 days at 23 °C with 16 h/8 h light/dark photoperiod. Fluorescence was evaluated in the abaxial face of infiltrated leaves and images were acquired using Zeiss LSM 710 confocal microscope. Green fluorescence signal (500–550 nm) and mRFP (585–620 nm) were collected following excitation with 488- and 561-nm lasers as described before [47].

2.10. Endocytosis of FM4-64 and sodium green staining

Plasma membrane internalization was evaluated with the endocytic tracer FM4-64 (Thermo Fisher Scientific #T-13320) in root cells of seven-day-old wild-type and transgenic Arabidopsis seedlings grown in solid MS (half-strength). For each genotype, 4 seedlings were analyzed. Seedlings were treated with 4 μM FM4-64 in half-strength liquid MS for 10 min at 4 °C. and then washed and incubated in half-strength liquid MS at 25 °C for 5 and 30 min to allow FM4-64 internalization. Images of root cells from the transition zone were captured with a LSM 710 Zeiss confocal microscope. FM4-64 was excited using a 543 nm laser and the fluorescence collected in the range of 560–650 nm. Image analyses and fluorescence quantification were performed using the FIJ-Image J software [48]. The rate of FM4-64 internalization was calculated by the ratio between the mean of the intracellular fluorescence and the mean of the whole cell fluorescence (including the plasma membrane) from 20 to 50 cells. Three biological replicates were performed.

To monitor the intracellular localization of Na+ in root cells of wild-type and SchRabGDI1-expressing Arabidopsis lines, 7-day-old seedlings grown in half-strength MS plates were incubated in liquid half-strength MS supplemented with 100 mM NaCl and 4 μM FM4-64 for 12 h [21].
After treatment, seedlings were incubated with 5 μM of the sodium tracer Sodium Green™ tetraacetate (ThermoFisher Scientific #S-6901) for 30 min. The Na⁺-dependent fluorescence of Sodium Green and the FM-64 fluorescence from cells of the transition zone of the primary root were visualized under a confocal microscope. Confocal images were captured with a LSCM 710 Zeiss confocal microscope using a predetermined setting for FM-64 and Sodium Green fluorescence obtained from the software ZEN 2012 edition. Three biological replicates were performed.

2.11. NBT staining

For the histochemical detection of superoxide radical (O₂⁻) in Arabidopsis plants subjected to salt stress (described above), the nitroblue tetrazolium (NBT) method was employed [49]. After 15 d of growth in the presence of 75 mM NaCl, plants were imersed in a solution of NBT (0.5 mg/ml dissolved in sodium phosphate buffer pH 8.0) and incubated in the dark for 12 h. Chlorophyll was removed immersing the plants in an ethanol:acetic acid:glycerol (3:1:1) solution and then heating at 90 °C for 15 min to complete discoloration.

2.12. Chemical analysis

Arabidopsis Col-0 seedlings and 35S:SchRabGDI1 lines were grown in MS (half-strength) media for seven days and then transferred to MS (half-strength) supplemented with 75 mM NaCl for 15 days. The seedlings were collected and roots and leaves were separated to quantify sodium content. The leaves and root samples were washed with bi-distilled water and dried up in a heater at 70 °C. For digestion, 0.5 g of sample was weighed out and 5 ml of Suprapur nitric acid was added. The samples were then dried almost completely under an extractor fan, with constant stirring, using a heating plate set to 90 °C. Finally, the solutions were filtered using 0.45 μm filters. The filtering process was performed to a final 50 ml volume with bi-distilled water. Sodium measurements were done by flame atomic absorption spectroscopy (air/acetylene) using a Unicam spectrophotometer model 969. The analysis methodology was validated using certified reference material (BIMEP-432), supplied by the Wageningen Evaluating Programs for Analytical Laboratories (WEPAL). The reagents used were of high purity (Suprapur, Merck, Darmstadt, Germany) and the standard solutions for the various metals were prepared from concentrated solutions (Fisher Scientific International Company).

2.13. Statistical analysis

All data was subjected to different types of statistical analyses by using the software R and Remdr package (http://knuth.uca.es/R/doku.php?id=instalacion_de_r_y_remdr-r-uca). They included t-student (endocytosis rate, plant biomass and ROS quantification), one way ANOVA (signal fluorescence of sodium green quantification and sodium contents in roots and shoots). Statistical differences are referred to as significant when P ≤ 0.05.

3. Results

3.1. Identification and isolation of GDP dissociation inhibitor gene SchRabGDI1 from Solanum chilense

Subtractive cDNA hybridization of stressed (salt-treated) and unstressed plants (control treatment) of S. chilense was carried out to isolate differentially up-regulated genes in response to salt stress. This strategy identified several genes and their sequences were compared with the NCBI database since no annotation is available for the S. chilense genome. One of these genes, named SchRabGDI1 (Accession number AY778206), showed high similarity to a GDP dissociation inhibitor (GDI) gene from Solanum lycopersicum. The full-length clone of SchRabGDI1 (1335 nucleotides) encodes a protein of 444 amino acid residues. A search conducted in the Solgenomic database (https://solgenomics.net/) predicted three putative RabGDI genes in the genome of the cultivated tomato (S. lycopersicum), which were named SIGDI1, SIGDI8 and SIGDI12 according to their chromosome location.

The deduced protein sequence of SchRabGDI1 was 99% identical to SIGDI12 (Solycol2g017570.1) from S. lycopersicum and StGDI1 (NP_001274860.1) from S. tuberosum. The amino acid sequence similarity to other GDIs were: 93.5% with SIGDI8 (Solycol0g015860.2), 84.41% with AtGDI1, 84.23% with AtGDI2 and 82.21% with SIGDI1 (Solycol1g0105810.2). At the structural level, SchRabGDI1 presented five classical domains of the GDIs called SCR (Sequence Conserved Region), composed of invariant sequences of tri or tetra peptides (Fig. S1A). The phylogenetic analysis revealed that SchRabGDI1 is closely associated with GDI proteins from the Solanacea family, such as those of S. tuberosum RabGDI, S. lycopersicum RabGDI12 and RabGDI8 and Nicotiana tabacum RabGDI1 (AAAB80717; [50]), whereas RabGDI1 from S. lycopersicum grouped separately from the rest of the plant RabGDI proteins (Fig. S1B). These results strongly suggested that SchRabGDI1 encodes a RabGDI in wild tomato S. chilense.

3.2. SchRabGDI1 is induced under salt stress in S. chilense

Due to the genetic redundancy of RabGDIs in plants, we studied the organ-specific transcriptional profile of SchRabGDI1. When S. chilense plants were grown under control conditions, SchRabGDI1 expression was detected in all the analyzed tissues (Fig. 1), being its transcript levels slightly higher in roots than in flowers and other vegetative tissues (Fig. 1). To assess the inducibility of SchRabGDI1 by salt, S. chilense plants were exposed to a severe salt stress treatment (400 mM NaCl) and the expression patterns of SchRabGDI1 and two stress-responsive genes TSW12 [32,51] and AREB1 [32,52] were analyzed. An increase in the transcript levels of SchRabGDI1 were observed in roots 3 h after treatment (Fig. 2A), reaching a maximum level (4-fold) at 6 h and then decreased gradually to control levels after 72 h. In leaves, up-regulation of SchRabGDI1 was detected 3 h after treatment, with the highest expression levels (4-fold) attained 12 h after treatment, declining to basal levels after 24 h and remained constant thereafter (Fig. 2B). SchRabGDI1 expression pattern under salt stress was similar to the profile displayed by the stress-induced genes TSW12 and AREB1 (Fig. 2C and D), suggesting that they may respond to a similar signaling pathway and a possible participation of this RabGDI protein in mechanisms associated with stress response.

![Fig. 1. Gene expression profile of SchRabGDI1 in vegetative and reproductive tissues of S. chilense. The relative abundance of the transcripts was determined by qPCR from total RNA of the indicated organs: R root, T stem, YL Young leaf, ML mature leaf, FB floral buttons and F flowers. Values represent mean ± SE (n = 3). The constitutive expression of the gene SchUBI3 and the 2-ΔΔCt method described by [31] were used for normalization. The transcript levels obtained for YL were taken to assign the value one.](https://www.plantphysiol.org/doi/fig/10.1104/pp.17.00646.s1.00646.F1-00646.F1)
3.3. SchRabGDI1 encodes a RabGDI protein

Functional characterization of SchRabGDI1 was carried out by functional complementation of a yeast mutant with a defect in membrane trafficking. S. cerevisiae contains a single copy gene encoding a Rab-GDP dissociation inhibitor (GDI1) which is allelic to the SEC19 gene [53,54]. Mutations in SEC19 (i.e. the sec19-1 mutation) cause a thermosensitive growth defect, thus allowing the sec19 mutant strain to grow at a permissive temperature (25 °C) but not at the restrictive temperature of 37 °C. The sec19-1 mutant was transformed with the construct GAL1:SchRabGDI1, which allows to switch on and off the expression of the SchRabGDI1 by adding galactose or glucose in the media, respectively. SchRabGDI1 expression rescued the defective mutant phenotype at 37 °C in a medium containing galactose (Fig. 3). However the mutant transformed with GAL1:SchRabGDI1 still displayed a growth defect at 37 °C in glucose-containing media which repressed the GAL1 promoter. Therefore these results demonstrated that SchRabGDI1 encoded a functional RabGDI protein able to restore the function of a defective SEC19 gene in S. cerevisiae.

3.4. SchRabGDI1 interacts with SchRabG3e in vivo

RabGDI1s are regulatory proteins that interact with RabGTPases and mediate their recycling [4]. In addition to SchRabGDI1, we identified and isolated a SchRabG3e which is identical to AtRabG3e from Arabidopsis thaliana. In silico analysis predicted the interaction between SchRabGDI1 and SchRabG3e (Fig. 4A). Protein-protein docking between the tri-dimensional structures of SchRabGDI1 and SchRabG3e and the structural alignment with the RabGTPase-RabGDI crystalized structure PDB code 1UKV, showed the binding platform between both proteins (Figs. S1A and S2). Indeed, the molecular dynamics analysis revealed the essential residues that participate in the interaction interface (Figs. S1A and S2). To confirm that these two proteins can interact in vivo, we followed a bimolecular fluorescence complementation (BiFC) assay which is based on the restoration of fluorescence after two fluorescent protein halves are brought together due to the protein-protein interaction of two proteins of interest fused to the fluorescent halves [55]. To develop the assay, we first evaluated (in silico) the effect that fusion proteins could have on RabGTPase-RabGDI binding. Visual evaluation of the structures formed by the half of the fluorescent

Fig. 2. Expression analysis of SchRabGDI1 in S. chilense exposed to salt stress. SchRabGDI1 (A, B), SchAREB1 (C, D) and SchTSW12 (E, F), were evaluated by qRT-PCR using total RNA extracted from leaves and roots of 12 weeks old plants after salt stress treatment (300 mM NaCl) for a period of 0, 3, 6, 12, 24, 48 and 72 h. Values are the mean ± SE (n = 3). Expression of the constitutive gene SchUBI3 and the 2-ΔΔCt method described by [31] were used for normalization. The transcript levels obtained in roots at time 0 were assign the value of one for each gene analyzed.
protein Venus fused to the C-terminal end of SchRabGDI1 and half of the super cyan fluorescent protein (SCFP) bound to the free N-terminal end of SchRabG3e, together with a molecular dynamics analysis of each system, indicated that there was no obstruction of the RabGTPase-RabGDI binding platform (Fig. 4B). Subsequently, the constructs GDI-VYNE (SchRabGDI1 fused to the N-terminal half of Venus) and SCYCE-RabG3e (C-terminal half of SCFP fused to N-terminal of SchRabG3e) were transiently expressed in epidermal cells of tobacco (Nicotiana benthamiana) leaves. As a positive control of the transformation, a construct that allows the accumulation of the plasma membrane intrinsic protein PIP1;4 fused to red fluorescent protein (PIP1:4-RFP) was also used. The green fluorescence signal showed the interaction

Fig. 3. Complementation assay of the sec19 yeast strain with SchRabGDI1. RSY249 (WT) and RSY274 (sec19-1) yeast cells were transformed with pYES-DEST52 and sec19-1 (gdi1) cells transformed with GAL1-SchRabGDI1 were streaked on glucose (Glu; SC-URA) and galactose (Gal; SC-URA) plates and incubated at 37 °C (restrictive) and 28 °C (permissive) for 3 days.

Fig. 4. Structural docking between SchRabGDI1, SchRabG3e and fusion proteins. (A) The energy of protein-protein docking SchRabGDI1-SchRabG3e calculated by the CLUSPRO software was −693 Kcal/mol. (B) Complex of Rab-GDI-YC155-YN155 proteins. It is presented to VYNE (yellow) fused to the C-terminus end of SchRabGDI1 (orange) and SCYCE (cyan) fused to the N-terminus of SchRabG3e (purple). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
between GDI-VYNE and SCYCE-RabG3e fusion proteins confirming the in vivo interaction of SchRabGDI1 and SchRabG3e (Fig. 5). In both constructs, the SchRabGDI1 and SchRabG3e maintained their own targeting sequences for their localization. The endosome-like and membrane-like pattern of green fluorescent SchRabGDI1-SchRabG3e complex was consistent with the molecular function of the complex in membrane recruiting, supporting the role of SchRabGDI1 in endomembrane trafficking (Fig. 5). To verify that the interaction of SchRabGDI1 and SchRabG3e was specific, AtNHX5 which is known to localize in endosomal compartments [46] was used as negative control for the interaction with SchRabG3e. As shown in Fig. S3B, BiFC signal of AtNHX5 and SchRabG3e was not observed, while the plasma membrane expression control showed fluorescence signal of RFP. Similarly, Fig. S3C showed no BiFC signal of SchRabGDI1 and AtRem1.2 (encoding Remorin, a membrane raft-bound protein) interaction. BiFC signal of AtNHX5 and AtRem1.2 in Fig. S3D showed that the NHX5-VYNE and SCYCE-Rem1.2 genes were properly expressed and interacted. These results indicated that the BiFC interaction between SchRabGDI1 and SchRabG3e was specific.

3.5. Expression of SchRabGDI1 in Arabidopsis enhances tolerance to salt stress

Given the salt-induced SchRabGDI1 expression in S. chilense (Fig. 2), we assessed whether the heterologous expression of SchRabGDI1 could contribute to enhance salt tolerance in Arabidopsis thaliana. Homozygous T3 transgenic Col-0 Arabidopsis overexpressing SchRabGDI1 were generated (Fig. S4) and their growth performance under 75 mM NaCl was tested (Fig. 6). No phenotypical differences were observed when the plants were grown under control conditions and their biomass was not affected (Fig. 6A). On the other hand, when plants were subjected to 75 mM NaCl, growth of transgenic plants expressing higher levels of SchRabGDI1 (L15 and L32) (Fig. S5) was less affected (Fig. 6B). Salt stress induces the accumulation of toxic reactive oxygen species (ROS) (Miller et al., 2010). Histochemical staining with nitroblue tetrazolium (NBT) showed that transgenic lines L15 and L32 accumulated less superoxide radical (O$_2^-$) than wild type plants after salt treatment (Fig. 7). The results showed that the salt tolerance displayed by the transgenic plants correlated with SchRabGDI1 expression levels (Figs. 7B and S4), strongly suggesting an active role of SchRabGDI1 in stress tolerance. Besides, the total Na$^+$ content was determined before and after salt treatment. The content of Na$^+$ in shoot and roots from transgenic and WT plants was similar before the salt treatment however after the treatment, the shoot Na$^+$ content in the transgenics was lower than in WT, while the root Na$^+$ content was higher than in WT (Fig. 8A and B).

3.6. Endocytosis is induced by expressing SchRabGDI1 in Arabidopsis

To analyze the involvement of SchRabGDI1 in the endocytic pathway, Arabidopsis transgenic plants overexpressing SchRabGDI1 were used to monitor endocytosis by using the endocytic tracer dye FM4-64 [56] which initially binds the plasma membrane and then is internalized to endosomes. Fig. 9 shows FM4-64 internalization at 5 and 30 min in root cells of wild-type (control) and SchRabGDI1 transgenic lines. An enhanced internalization of FM4-64 was already observed at 5 min after the endocytosis time course in SchRabGDI1 transgenic lines compared to the wild-type (Fig. 9A). The increased internalization of the tracer dye was evidenced by the higher abundance of small punctate FM4-64-endosomes at both 5 and 30 min of endocytosis dynamics. Quantification of the FM4-64 fluorescence signal showed significant differences in the endocytic rates between wild-type and overexpressing SchRabGDI1 lines (~15%) (Fig. 9B), suggesting that an increment in the cellular RabGTPases pool brought about by
SchRabGDI1 increased the rate of endocytic trafficking, evidencing the cellular functionality of SchRabGDI1.

In addition, the intracellular Na⁺ distribution in root cells of wild type and SchRabGDI1 transgenic lines subjected to 100 mM NaCl was monitored using the fluorescent Na⁺ indicator Sodium Green and the endocytic tracer FM4-64. Fig. S5 shows root cells from the transition zone in which Na⁺ ions are detected in the lumen of the main vacuole and smaller vesicular structures (Fig. S5A). Quantification of the fluorescence intensity of Sodium Green revealed a higher accumulation of Na⁺ in the transgenic lines compared to wild type (Fig. S5B). These results indicate that the increase in salt tolerance conferred by over-expression of SchRabGDI1 may be consequence of an intensification of endocytosis and sodium accumulation in root cells.

4. Discussion

The physiological response of plants to salt stress is a complex process that requires the coordinated function of many genes. Although members of the Rab family genes has been associated to endosomal trafficking [57], little is known about the participation of those associated with the control of the molecular switch of RabGTPases in the response to saline stress. In this work, we report that SchRabGDI1, one of the genes involved in the control of cycling between active and inactive state of Rab proteins, plays a role in the tolerance of plants to salt stress.

4.1. SchRabGDI1 encodes a functional GDP dissociation inhibitor and is induced by salt stress

SchRabGDI1 was identified among salt stress-induced transcripts of Solanum chilense roots. SchRabGDI1 expression was induced early by salinity in both leaves and roots as two stress-responsive genes TSW12 [32,51] and AREB1 [32,52] (Fig. 2). SchRabGDI1 expression patterns was similar to that of AtRabGDI1 (AT2G44100) in Arabidopsis (http://bar.utoronto.ca/efp/cgi-bin/efpWeb.cgi), but differed to the expression of RabGDI from Mangifera indica L, which is down-regulated in leaves.

![Fig. 6. Analysis of Arabidopsis plants expressing SchRabGDI1 subjected to salt stress. Phenotypes and fresh weight of transgenic lines expressing SchRabGDI1 under normal growth condition (MS) (A) and 75 mM NaCl (B). Three independent experiments with five plants of each genotype were used in the analysis. Data are the mean +/- S.D. (n = 3). Asterisks represent significant difference from the wild type (P < 0.05).](image)

![Fig. 7. ROS accumulation in wild-type and Arabidopsis plants expressing SchRabGDI1 under salt stress. (A) ROS accumulation was evaluated on seedlings of wild-type and three transgenic lines (4 weeks-old) exposed under normal (MS) or saline media (MS + 100 mM NaCl) during 15 days. Fifteen seedlings were used for each line, and three representative pictures of each line are shown. (B) Quantification was performed using the software image J. from fifteen plants per line. Values are mean +/- S.E. (n = 15). Asterisks represent significant difference from the wild type (P < 0.05).](image)
under similar salt stress conditions [58]. However, the timing of the induction of SchRabGDI1 in response to such stimulus coincides with that described for genes that belong to the RabGTPase family such as McRab5-b from *Mesembryanthemum crystallinum* [56], AtRabG3e (AT1G49500.1) and AtRabA1(At1g06400) from *Arabidopsis thaliana* [22,24], (http://bar.utoronto.ca/efp/cgi-bin/efpWeb.cgi) and SchRabG3e from *S. chilense* (data not shown). These results would suggest that SchRabGDI1 expression is required during the vesicular trafficking events occurring during salt stress.

The analysis of the deduced amino acid sequence showed that...
SchRabGDI1 possesses high sequence similarity with other RabGDIs from plants and its evolutionary proximity to RabGDIs of *S. tuberosus*, *S. lyco-
copercum* and *Nicotiana tabacum* (Fig. S1B). *SchRabGDI1* contains the five domains that structurally and functionally define sequence conserved residues (SCRs) which are characteristic of all RabGDIs (Fig. S1A). Functionally, *SchRabGDI1* was able to complement the yeast secretory mutant sec19 RSY273, restoring the lethal temperature-sensitive phenotype [53].

Protein-protein interaction between Rab and GDI proteins is critical for maintaining an efficient vesicular trafficking, since GDI retrieves the GDP-bound form of Rab from the membrane to form a heterodimeric complex that is used as a cytosolic pool for the reuse of inactive Rab during a series of vesicle budding and fusion [4,59]. We used bioin-
formatics tools to determine the essential residues for the interaction of *SchRabGDI1* and SchRabG3e, which revealed a high interaction affinity with very low ΔG (Figs. 4 A and S2). The results obtained using bi-
molecular fluorescence complementation (BIFC) assays ratified the in silico predictions given that the in vivo interaction between *SchRabGDI1* and SchRabG3e was observed when both genes were transiently co-
expressed in epidermal cells of tobacco leaves (Fig. 5). This demonstr-
ated that *SchRabGDI1* was able to form a heterodimer with SchRabG3e, suggesting its functional participation in the cycle of RabGTPases and their likely involvement in intracellular vesicular trafficking.

4.2. *SchRabGDI1* confers tolerance to saline stress in Arabidopsis

In recent years, evidences indicate that intracellular vesicular traffic plays an important role in the adaptation of plants to salt stress [21,60]. Ectopic expression and silencing have been used to show that members of Rab GTPases and SNARES (soluble N-ethylmaleimide sensitive factor attachment protein receptor) families, two major regulators of vesicular trafficking, participates in the tolerance of plants to salt stress [22,24,61–64]. Under standard growth conditions wild and transgenic plants do not exhibit a differential phenotype (Fig. 6A), probably be-
cause the protein encoded by the transgene requires the expression of the other genes that constitute the molecular switch of RabGTPases, which are not expressed under this conditions.

To demonstrate its in-
volvement in the response to salt stress, we analyzed the effect of ec-
topic expression of *SchRabGDI1* in Arabidopsis plants. Transgenic plants showed increased tolerance to salt (75 mM NaCl), displaying higher fresh weight and reduced accumulation of reactive oxygen species compared with wild type plants (Figs. 6 and 7). The improved salinity tolerance of Arabidopsis plants harboring *SchRabGDI1* is comparable to results described in Arabidopsis, tobacco and rice overexpressing Rab7 (Rab3g3e) when exposed to salt stress [22,23,65]. In addition to accu-
mulate Na⁺ into their vacuoles and to extrude Na⁺ to the apoplast, plants also restrict the movement of Na⁺ ion from the roots to the shoots [19]. The enhanced salt tolerance displayed by the transgenic plants overexpressing *SchRabGDI1* correlated with the increased Na⁺ accumulation in the root vacuoles and the differential Na⁺ content between roots and shoots.

4.3. *SchRabGDI1* regulates endocytotic pathway and Na⁺ intracellular distribution

Leschem et al. [66] reported that salt stress induced bulk-flow endocytosis in Arabidopsis roots. The increased endocytosis induced by salt stress has been confirmed subsequently in other studies [60,67,68]. The bulk endocytosis induced by salt has been shown to promote a rapid increase in vacuolar volume and accumulation of sodium in the vacuole of roots cells [20]. In this work, the overexpression of *SchRabGDI1* in Arabidopsis roots led to an increase in endocytosis as determined by the tracer dye FM 4–64 (Fig. 8). Moreover, when ex-
posed to 100 mM NaCl, the transgenic plants showed higher accumu-
lution of sodium in their root vacuoles (as measured by the fluorescence of the dye sodium-green) (Fig. S4). These results suggest the partici-
pation of *SchRabGDI1* in the regulation of endocytosis and postulates a possible contribution of salt-induced endocytosis to sodium accumula-
tion in the vacuole, in addition to the action of vacuolar Na⁺/H⁺ anti-
porters [19,69]. In conclusion, our findings suggest that salt tolerant species such as *S. chilense* use bulk endocytosis as one of the early mechanisms to avoid the saline stress, for which they require the con-
certed expression of regulatory genes of the vesicular trafficking of the endocytic pathway.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.plantsci.2017.06.007.

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