

Phospholipases $D\zeta 1$ and $D\zeta 2$ have distinct roles in growth and antioxidant systems in Arabidopsis thaliana responding to salt stress

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1	Phopholipases D(1 and D(2 have distinct roles in growth and antioxidant systems in Arabidopsis
2	thaliana responding to salt stress
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26	Short running title
27	Roles of phopholipases Dζ during salt stress in Arabidopsis
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35 Main conclusion

36 Phospholipases Dζ play different roles in Arabidopsis salt tolerance affecting the regulation of ion
37 transport and antioxidant responses.

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42 Abstract

Lipid signalling mediated by phospholipase D (PLD) plays essential roles in plant growth including 43 stress and hormonal responses. Here we show that $PLD\zeta 1$ and $PLD\zeta 2$ have distinct effects on 44 45 Arabidopsis responses to salinity. A transcriptome analysis of a double $pld\zeta lpld\zeta 2$ mutant revealed 46 a cluster of genes involved in abiotic and biotic stresses, such as the high salt stress responsive 47 genes DDF1 and RD29A. Another cluster of genes with a common expression pattern included 48 ROS detoxification genes involved in electron transport and biotic and abiotic stress responses. 49 Total SOD activity was induced early in the shoots and roots of all $pld\zeta$ mutants exposed to mild or 50 severe salinity with the highest SOD activity measured in $pld\zeta^2$ at 14 days. Lipid peroxidation in 51 shoots and roots was higher in the *pld* $\zeta 1$ mutant upon salt treatment and *pld* $\zeta 1$ accumulated H₂O₂ 52 earlier than other genotypes in response to salt. Salinity caused less deleterious effects on K⁺ 53 accumulation in shoots and roots of the $pld\zeta^2$ mutant than of wild type, causing only a slight 54 variation in Na⁺/K⁺ ratio. Relative growth rates of wild-type plants, $pld\zeta 1$, $pld\zeta 2$ and $pld\zeta 1 pld\zeta 2$ 55 mutants were similar in control conditions but strongly affected by salt in WT and $pld\zeta l$. The 56 efficiency of photosystem II, estimated by measuring the ratio of chlorophyll fluorescence (Fv/Fm 57 ratio), was strongly decreased in *pld* $\zeta 1$ under salt stress. In conclusion, *PLD* $\zeta 2$ plays a key role in 58 determining Arabidopsis sensitivity to salt stress allowing ion transport and antioxidant responses to 59 be finely regulated.

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⁶⁵ Keywords: Ion relations; Phospholipase D; PLDζ; Reactive oxygen species; Salt stress;
66 Transcriptome.

69 Introduction

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Salinity is a major environmental constraint on the growth, productivity and diversity of plants. The effect of salt stress on plants depends on the salt concentration, the duration of exposure and the plant genotype (Ellouzi et al. 2011). The presence of salt in the environment induces water deficit in plants because the external water potential is lowered, while ion toxicity and nutritional alterations disturb ion transport systems (Munns and Tester 2008; Julkowska and Testerink 2015). Salt stress also causes membrane damage, alters levels of growth regulators, inhibits some enzymes, and disrupts photosynthesis, and may thus lead to plant death.

78 One of the known plant responses to salt stress is ROS production (for review see Miller et al. 2010; 79 Ben Rejeb et al. 2014). Plant cells need to regulate ROS production as excess ROS is potentially 80 harmful to nucleic acids, proteins and lipids, and may therefore lead to cell injury and death (Gill 81 and Tuteja 2010). ROS produced through NAPDH oxidase activity was shown to be mediated by 82 phospholipid signalling (Zhang et al. 2009). The second messenger phosphatidic acid (PA) is a 83 phospholipid which targets specific proteins to bring about cellular and physiological changes that 84 allow plants to adapt to abiotic stresses (for review see Hong et al. 2010; Hou et al. 2016). PA is 85 formed when phospholipase D (PLD) hydrolyses structural phospholipids at the terminal 86 phosphoesteric bond with release of the hydrophilic head group. In plants, phospholipase D (PLD) 87 is predominant among the phospholipase families (for review see Li et al. 2009; McLoughlin and 88 Testerink, 2013). PLD activity increases rapidly in response to various environmental stresses such 89 as cold, drought, and salinity (Vergnolle et al. 2005; Bargmann et al. 2009; Hong et al. 2010). 90 Proline accumulation, another common physiological response to stress, was shown to be 91 negatively regulated by PLD activity in Arabidopsis thaliana (Thiery et al. 2004).

92 In A. thaliana, the PLD family includes 12 members that are classified into six types, PLDa (3 93 isoforms), $-\beta$ (2 isoforms), $-\gamma$ (3 isoforms), $-\delta$, $-\varepsilon$, and $-\zeta$ (2 isoforms), according to their sequences 94 and enzymatic properties (Bargmann and Munnik 2006). Biochemical studies have revealed that the 95 phospholipid-hydrolysing activities of PLD are either calcium-dependent through a C2 domain (C2-PLD) or calcium-independent having pleckstrin homology (PH) and phox homology (PX) domains 96 97 (PXPH-PLD) (Meijer and Munnik 2003). Several PLD isoforms have been functionally 98 characterized. The two PLD(s (PLD(1 and PLD(2))) are structurally different from other PLDs (for 99 review see Li et al. 2009). PLD ζ 1 and PLD ζ 2 do not have C2 domains and do not require Ca²⁺ for 100 enzymatic activity (Qin and Wang 2002), but they do have PH and PX domains (Qin and Wang 101 2002). *PLD*\[2] gene expression is five-fold greater in roots than in leaves and the gene function is 102 required for root hair morphogenesis (Li et al. 2006). The $PLD\zeta 2$ gene is mainly expressed in roots 103 (Li et al. 2006). *PLD* ζ ² gene expression is triggered by exogenous auxin (Li and Xue 2007) and 104 phosphate (Li et al. 2006). PLD ζ 1 and PLD ζ 2 were shown to be involved in the process by which 105 root architecture adapts to the lack of phosphate (Li et al. 2006). PLD ζ 2 is also involved in vesicle 106 trafficking and auxin transport (Li and Xue 2007). The *pld* ζ 2 mutant has a reduced halotropic 107 response, i.e. the capacity to change the direction of root growth to avoid salt, due to impaired

- 108 vesicle trafficking (Galvan-Ampudia et al. 2013).
- 109 In this study, the roles of $PLD\zeta 1$ and $PLD\zeta 2$ were investigated in Arabidopsis responding to salt
- 110 stress. The global transcriptome of the $pld\zeta lpld\zeta 2$ double mutant treated with salt was analysed to
- 111 identify candidate genes regulated by these PLDs. Changes in growth, ion balance, ROS content,
- 112 antioxidants, and lipid peroxidation were compared in wild type, $pld\zeta 1$, $pld\zeta 2$, and $pld\zeta 1pld\zeta 2$ after
- short and long periods of salt treatment. $pld\zeta 1$ and $pld\zeta 2$ mutants responded differently to salt stress
- 114 suggesting they have distinct physiological roles in Arabidopsis.
- 115

117 Materials and methods

118 Plant materials and growth conditions

119 Arabidopsis thaliana (L.) Heynh ecotype Columbia-0 (Col-0) was used as the wild type in this 120 study and was obtained from Nottingham Arabidopsis Stock Centre, Loughborough, United 121 Kingdom. Homozygous T-DNA insertion mutant lines *pldZ1* (SALK 083090) and pldZA 122 (SALK_094369) were obtained from the Salk Institute, La Jolla, USA (Alonso and Stepanova 123 2003) and have been described by Li et al. (2006). Double homozygous mutant $pld\zeta 1pld\zeta 2$ plants 124 were isolated from the F2 progeny of crosses between $pld\zeta l$ and $pld\zeta 2$. The presence and 125 homozygosity of the T-DNA alleles were checked by triplex PCR in all experimental lines using a 126 primer specific to the left border of the T-DNA and two gene-specific primers flanking the insertion 127 site (Suppl. Fig. S1 and Suppl. Table S1).

128 Surface sterilized seeds of wild-type and mutant lines were sown onto grids placed on Petri dishes

129 containing $0.5 \times$ Murashige and Skoog (MS; Murashige and Skoog 1962) solid medium (0.8 %

130 agar) according to Parre et al. (2007). Seeds were placed in a cold room at 4 $^{\circ}$ C for 1 day to break

- 131 dormancy and then transferred to a growth chamber at 22 °C with continuous light (90 μ mol 132 photons m⁻² s⁻¹). Twelve-day-old Arabidopsis seedlings were exposed to 200 mM NaCl for up to 24 133 h.
- 134 For physiological experiments, wild type, $pld\zeta 1$, $pld\zeta 2$ and $pld\zeta 1pld\zeta 2$ seeds were germinated in 135 plastic pots (70 mL) filled with inert sand and watered daily with distilled water for one week. 136 Seedlings of each genotype were then irrigated with Hewitt nutrient solution (Hewitt 1966) for 3 137 weeks. The experiments were performed in a glasshouse under controlled conditions, photoperiod 138 of 16h/8h (day/night), at 20-25 °C and 65-75 % relative humidity. Four-week-old plants were 139 treated with 75 mM or 150 mM NaCl and collected after 3 h, 24 h and 72 h for short exposure to 140 salt stress and after 7 and 14 days for long exposure to salinity. Non-stressed plants grown without 141 added salt were collected and used as controls.
- 142
- 143 Transcriptome analysis of the $pld\zeta 1pld\zeta 2$ double mutant

144 Analysis of the *pld\zeta1pld\zeta2* transcriptome was conducted at the Unité de Recherche en Génomique 145 Végétale (Evry, France) using the Complete Arabidopsis Transcriptome MicroArray (CATMA) 146 containing 24576 gene-specific tags from Arabidopsis (Crowe et al. 2003). Six biological replicates 147 and 14 dye swaps (technical replicates) were made. Twelve-day-old seedlings treated for 3 h with 148 either 200 mM NaCl or 400 mM mannitol for stress conditions or with 0.5 × MS for control 149 conditions were rapidly washed with water and immediately frozen in liquid nitrogen and stored at

150 -80 °C. Total RNA was isolated from seedlings by the guanidinium thiocyanate-CsCl purification 151 method (Sambrook et al. 1989). Labelling of cRNAs with Cy3-dUTP or Cy5-dUTP (Perkin-Elmer-152 NEN Life Science Products, Courtaboeuf, France), hybridization to slides, and scanning were 153 performed as described in Lurin et al. (2004). Methods and data were deposited in CATMA 154 database RS06-01_PLD (http://urgv.evry.inra.fr/cgi-bin/projects/CATdb/consult_expce.pl? 155 experiment id=113) and GEO (https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE9459). 156 Statistical analysis of transcriptome data was done as in Planchais et al. (2014). 157

158 Reverse transcription PCR and quantitative PCR analysis of gene expression

159 Total RNA was extracted from 100 mg of homogenized tissue from different genotypes using the 160 RNeasy Plant Mini Kit (Qiagen, Hilden, Germany) following the manufacturer's instructions. RNA 161 was quantified by measuring the absorbance at 260 nm using a Nanovue spectrophotometer 162 (Nanodrop Spectrophotometer ND 1000, Wilmington, DE, USA). RNA was incubated with DNase 163 I (Sigma-Aldrich, Saint-Louis, MO, USA) to eliminate genomic DNA. For reverse transcription, 164 first-strand cDNA was synthesized from 1.5 µg of total RNA using the RevertAid reverse transcriptase kit (Thermo Fisher Scientific Inc., Waltham, MA, USA). Complementary DNAs were 165 166 amplified using DreamTaq Green polymerase (Thermo Fisher Scientific Inc.) and gene-specific primers (Supplementary Table S1). The APT1 gene (At1g27450) was used as a positive control for 167 168 quantifying relative amounts of cDNA. Amplified PCR fragments were separated on 2 % (w/v) 169 agarose gels and visualized by staining with ethidium bromide and analysing the image with 170 GelDoc (Bio-Rad, Hercules, USA). For real-time PCR, 5 µL of diluted cDNA was used with 10 µL 171 of SYBR[®]GreenqPCR Master Mix (Thermo Fisher Scientific Inc.) and gene-specific primers in an 172 Eppendorf Master cycler[®] (Eppendorf France SAS, Montesson, France). Critical thresholds (Ct) were determined by using the Eppendorf Master cycler[®] realplex software and ratios were 173 174 calculated by using the method described by Pfaffl (2001). For each gene, a standard curve made 175 with dilutions of cDNA pools was used to calculate reaction efficiencies, and the levels of gene 176 transcript were normalized and expressed relative to the amounts observed under control conditions 177 with APT1 (At1g27450) as housekeeping gene. All semi-quantitative RT-PCR and real-time PCR 178 experiments were carried out with three biological replicates.

179

180 Measurements of plant biomass, maximum efficiency of PSII photochemistry, and leaf water status

Fresh weight (FW) and dry weight (DW) of rosettes and roots were measured at each experimental time point to evaluate leaf and root growth. The accumulation of biomass was estimated by determining the relative growth rate (RGR), which is a measure of biomass production relative to

- 184 treatment duration and/or initial plant size. RGR was calculated as $\Delta \ln(DW)/\Delta t$, where DW is the dry 185 weight, ln is the natural logarithm and Δ represents the difference between final and initial values for a given 186 time interval Δt (Hunt, 1990). The maximum efficiency of PSII photochemistry (*Fv/Fm*) was 187 monitored throughout the experiment using a portable mini-PAM fluorometer (ADC BioScientific
- 188 LC pro System Serial, Hoddesdon, UK).

189 The relative leaf water content (RWC) was calculated as 100×(FW-DW)/(TW-DW), where TW was

190 the turgid weight of leaves saturated for 24 h in deionized water at 4 °C in darkness and DW was

- 191 obtained after oven-drying the leaves at 60 °C for 72 h according to Ellouzi et al. (2013).
- 192
- 193 Measurement of ion content

Desiccated leaf and root tissues were ground to a fine powder and then broken down with concentrated 0.5 % HNO₃ according to Deal (1954). Na⁺ and K⁺ contents were determined using flame emission photometry (Corning, Tewksbury, MA, USA). In emission photometry the soluble mineral component is injected into an air-propane flame. Following thermal excitation a characteristic spectrum of lines is emitted that are selected by monochromator filters. Behind each filter a photoreceptor cell detects the intensity of the emitted light which is proportional to the amount of the emitting element contained in the vaporized solution.

- 201
- 202 Histochemical detection of hydrogen peroxide

The localization of hydrogen peroxide was determined according to Ben Rejeb et al. (2015). Fresh leaf and root samples were collected from each mutant and infiltrated with a solution containing 0.5

205 mg ml⁻¹ diaminobenzidine (DAB) and 10 mM Mes buffer pH 5.8. Samples were then kept at room

- 206 temperature until the brown residue generated by H₂O₂-DAB polymerization developed. Samples
- 207 were incubated for 60 min in boiling ethanol before being photographed.
- 208
- 209 Measurement of hydrogen peroxide concentration

210 Fresh leaf and root samples were ground in 0.2 % trichloroacetic acid (TCA) in a mortar and pestle

chilled on ice. The homogenate was centrifuged at 15000 g for 20 min at 4 °C. The supernatant was

212 mixed with 10 mM sodium phosphate buffer pH 7 and 1 M KI. The H₂O₂ content was determined

213 by comparing the absorbance of the sample at 390 nm with a standard calibration curve, expressing

214 values in μ mol H₂O₂ g⁻¹ DW.

- 215
- 216 Membrane lipid peroxidation assays

- Levels of lipid peroxidation were assessed by measuring the amount of malondialdehyde (MDA) in tissue. Fresh leaf and root samples were homogenized in 10 % TCA. The homogenate was centrifuged at 15000 *g* for 20 min at 4 °C. The supernatant was collected and mixed with 0.5 % thiobarbituric acid in 20 % TCA. Samples were heated at 95 °C for 25 min in a water bath, and then cooled on ice. The samples were centrifuged at 10000 *g* for 10 min and the absorbance of solutions at 532 and 600 nm was recorded. The MDA level was calculated using the extinction coefficient for MDA [$\varepsilon = 155 \mu$ M cm⁻¹] expressed in nmol MDA g⁻¹ DW.
- 224
- 225 Protein extraction and antioxidant enzyme assays

226 Leaf and root samples of each genotype were homogenized in a chilled mortar containing 10 % 227 (w/w) poly-vinyl-polypyrrolidone in 50 mM potassium phosphate buffer pH 7, 0.1 mM 228 ethylenediaminetetraacetic acid (EDTA) and 1 mM dithiothreitol. Homogenates were then 229 centrifuged at 10000 g for 15 min at 4 °C. The supernatants were collected and stored at -20 °C for 230 protein and enzyme assays. The soluble protein content of leaves and roots was estimated according 231 to Bradford (1976) with bovine serum albumin as the standard. Superoxide dismutase (SOD, 232 EC.1.15.1.1) activity was assayed based on the inhibition of nitro blue tetrazolium (NBT) reduction 233 (Beyer and Fridovich 1987). The assay was conducted in 50 mM potassium phosphate pH 7.8 234 containing 50 µL of enzyme extract, 2.25 mM NBT, 13 mM methionine, 2 mM riboflavin and 1 235 mM EDTA. The reaction was started by illuminating the sample for 15 min and stopped by 236 switching the light off. The amount of blue formazan formed was determined by measuring the 237 absorbance at 560 nm. One unit of SOD activity (U) was defined as the amount of enzyme that 238 caused 50 % inhibition of NBT reduction.

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240 Statistical analysis

241 Differences between measurements and between genotypes at different times were evaluated by 242 analysis of variance (one-way ANOVA) using SPSS (Chicago, IL, USA). Differences were 243 considered as statistically significant when $P \le 0.05$.

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251 Results

252 Analysis of the $pld\zeta 1pld\zeta 2$ double mutant transcriptome

253 If Arabidopsis PLD ζ act in lipid signalling in response to salt stress then a *pld\zeta1pld\zeta2* double 254 mutant, lacking both PLD⁽², would be expected to differ from WT in the way it responds 255 to salt stress. We first investigated gene expression in WT after 3 h exposure to salt or mannitol. 256 This was to test whether the ionic stress component of the response could be distinguished from the 257 osmotic stress component. A large common set of genes was induced (560 genes) or repressed (221 258 genes) by the two individual treatments, which was attributed to the similar effects of osmotic stress 259 on the plants (Supplementary Fig. S2). However a smaller subset of genes was specifically induced 260 (170 genes) or repressed (82 genes) only by salt. For example, the known salt-stress responsive 261 gene DDF1 was specifically upregulated in response to NaCl in WT but was not affected by 262 hyperosmotic stress caused by mannitol. Global transcriptome analysis of the $pld\zeta lpld\zeta 2$ double 263 mutant also revealed common sets of genes that were triggered or repressed by both NaCl and 264 mannitol, but also differential expression of subsets of genes in a salt-specific and mannitol-specific 265 manner (data not shown). In further experiments we were thus able to distinguish between ionic and 266 osmotic stress effects, focusing on the former. As some genes were differentially and specifically 267 regulated by NaCl, the perception and transduction of the ionic component of the salt stress signal 268 must have occurred within the 3 h of the stress treatment.

269 We then compared differences in the NaCl stress transcriptomes of $pld\zeta lpld\zeta 2$ and WT with the 270 WT NaCl stress response. In WT genes were either induced (642 genes) or repressed (241 genes) 271 after 3 h salt stress (Fig. 1). Twenty of the salt-induced genes in WT were found to be expressed at lower levels in $pld\zeta lpld\zeta 2$ after 3 h salt treatment than in WT in the same conditions (intersection in 272 273 Fig. 1a). This cluster of genes induced in WT, but less strongly induced in $pld\zeta 1pld\zeta 2$ was defined 274 as cluster 1 (Fig. 1 and Suppl. Fig. S3). Analysis of gene ontology using BAR Classification 275 SuperViewer (http://bar.utoronto.ca) showed an over-representation of genes involved in abiotic 276 and biotic stresses in cluster 1 (Suppl. Fig. S4). It was noted that the salt-stress responsive genes 277 DDF1 and RD29A (Magome et al. 2008) were present in cluster 1. We investigated the expression 278 of these two genes in WT, the pld $\zeta 1$ and pld $\zeta 2$ single mutants and the pld $\zeta 1$ pld $\zeta 2$ double mutant 279 using RT-PCR and quantitative PCR (Fig. 2a, c). The RT-PCR analysis confirmed that DDF1 280 expression was induced by salinity in WT plants (Fig. 2a). DDF1 gene expression was induced in 281 $pld\zeta lpld\zeta 2$ but not as strongly (4.9-fold) as in WT (6-fold) confirming the transcriptome data (Fig. 282 2c). In comparison, DDF1 expression was induced 9.7-fold in $pld\zeta_2$ and 7.8-fold in $pld\zeta_1$ (Fig. 2a, 283 c). Similarly, RD29A expression was strongly induced by salt treatment in WT (Fig. 2a, c) but less

so in *pld\zeta1pld\zeta2* as expected from the transcriptomic data. Of the four genotypes tested *pld\zeta2* again showed the highest level of salt-induction with a 10.4-fold increase in *RD29A* expression.

Another cluster of genes, cluster 2, was identified by comparing the differences in the $pld\zeta 1pld\zeta 2$ and WT transcriptomes between control and salt stress conditions. In this cluster 23 genes were deregulated in the double mutant in both conditions, 8 upregulated and 15 downregulated (Fig. 1 b, Suppl. Fig. S5). Cluster 2 gene functions mainly related to electron transport and biotic and abiotic stresses (Suppl. Fig. S4). The ROS detoxification genes *SOD1* (At1g08830) and *CCS1* (*Copper Chaperone for SOD1*, At1g12520) were overexpressed in $pld\zeta 1pld\zeta 2$ compared to WT after salt treatment and in control conditions (Suppl. Fig. S5).

- 293
- 294 ROS detoxification was impaired in $pld\zeta$ mutants

We verified whether *SOD1* expression was altered in *pld* ζ single and double mutants by RT-PCR, and compared it to expression of two other ROS detoxifying genes, ascorbate peroxidase 1 (*APX1*) and catalase (*CAT2*), which are commonly used as oxidative burst markers (Fig. 2b). *APX1* expression was slightly higher in *pld* ζ 2 than in WT and *pld* ζ 1, but was lower in *pld* ζ 1*pld* ζ 2. *SOD1* gene expression was upregulated by salt treatment in both *pld* ζ single mutants but not in WT. Expression of *CAT2* was highest in *pld* ζ 2 compared to the other genotypes even in control conditions (Fig. 2).

302 To test whether the lack of both or either PLD^{\zet} affected the ROS detoxification activity of SOD, 303 SOD activity was monitored for 14 days in the shoots and roots of WT and *pld* ζ mutants exposed to 304 moderate (75 mM NaCl) or severe salinity (150 mM NaCl) (Fig. 3). In leaves treated with 75 mM 305 NaCl SOD activity increased transiently in the first 72 h up to 316 % in WT, 169 % in $pld\zeta l$ and 306 108 % in *pld\zeta1pld\zeta2*, declining less rapidly then remaining constant until the end of the time course. 307 In roots, the kinetics of SOD activity were markedly different in $pld\zeta 1pld\zeta 2$ in comparison to other 308 genotypes as the sharp transient increase occurred within the first three days of mild salt treatment 309 (Fig. 3). Under the same salt conditions, no significant variations were detected in stressed roots of 310 WT and *pld* $\zeta 1$ mutants. By contrast SOD activity in both the shoots and roots of *pld* $\zeta 2$ treated with 311 75 mM NaCl increased linearly to reach a plateau that remained higher than levels in the other 312 genotypes at the end of the 14-day experiment. Upon severe NaCl stress, SOD activity in leaves of 313 $pld\zeta^2$ and $pld\zeta^2 pld\zeta^2$ mutants showed a distinct profile with a transient maximum activity at 3 h, 314 remaining higher than that of WT and *pld\zeta1* mutants until the end of the salt treatment (Fig. 3). No 315 substantial changes in SOD activity were observed in shoots of WT and $pld\zeta l$ mutants during stress 316 from 150 mM NaCl. SOD activities increased distinctly in the roots of all genotypes except $pld\zeta l$ 317 under severe salt stress. Although there was a transient maximal peak in SOD activity in WT during

318 the first 72 h of exposure to 150 mM NaCl, the highest SOD activities were measured in $pld\zeta^2$ and

319 $pld\zeta 1pld\zeta 2$ mutants after 14 days.

- 320
- 321 *pld* ζ mutants differentially accumulate ROS in response to salt stress
- The presence of H_2O_2 is an indicator of oxidative stress. The differential accumulation of H_2O_2 in leaves and roots over the duration of the stress was evaluated in the four genotypes using DAB, which stains tissues brown in the presence of H_2O_2 (Fig. 4). Leaves and roots grown in the absence of salt did not stain brown with DAB. After 3 h salt treatment, whole leaves and roots of WT and *pld* ζ *1pld* ζ *2* and *pld* ζ *1* were stained brown with DAB (Fig. 4). Although the assay is qualitative, the intensity of the staining consistently increased over the duration of the experiment. Roots and shoots of *pld* ζ *2* mutant were the most faintly stained, indicating these tissues had the lowest
- $329 \quad \text{ amounts of } H_2O_2.$
- The differences observed in DAB staining in *pld* ζ mutants led us to quantify H₂O₂ content. H₂O₂ measurements corresponded with the DAB staining. In the absence of salt stress, H₂O₂ did not accumulate above basal levels in any of the four genotypes. Under mild salt stress, H₂O₂ content increased steadily in shoots and roots in all four genotypes, accumulating faster during the first 24 h (Fig. 5). H₂O₂ accumulated at a similar rate in WT and *pld* ζ 1, but *pld* ζ 2 and *pld* ζ 1*pld* ζ 2 accumulated less than half of the amount measured in WT (Fig. 5). Under severe salt stress, *pld* ζ 2 leaves and roots had the lowest levels of H₂O₂ compared to the other *pld* ζ genotypes.
- Oxidative stress can lead to damage of structural macromolecules including lipids. Lipid peroxidation in leaves and roots was estimated by quantifying malondialdehyde (MDA), which forms when ROS attack lipids. The accumulation of MDA greatly increased in both leaves and roots growing under salt stress (Fig. 6). However the genotypes accumulated MDA differently. The highest amounts of MDA were found in leaves and roots of salt-stressed *pld* $\zeta 1$, while *pld* $\zeta 2$ accumulated the least MDA in leaves and roots in response to both moderate and severe salt stresses (Fig. 6).
- 344
- 345 Ion distribution was altered in $pld\zeta$ mutants responding to salt stress

The distribution of ions between leaves and roots of WT and the *pld* ζ mutants exposed to 14 days of either moderate or severe salinity is shown in Table 1. Changes in Na⁺ and Cl⁻ content followed the same profile over the 14 days in all genotypes and under both salt treatments, increasing linearly (data not shown) and reaching high levels by the end of the experiment. These final levels were higher in roots than in leaves (Table 1). There were some differences between genotypes. *pld* ζl mutants stressed by 150 mM NaCl had the most Na⁺ in leaves (7.3 mmol g⁻¹ DW) and roots (8.4 mmol g⁻¹ DW) (Table 1). This increase was concomitant with the highest Cl⁻ accumulation, which was greater in roots (5.8 mmol g⁻¹DW) than in leaves (3.9 mmol g⁻¹ DW) (Table 1). However, Na⁺ content increased more than Cl⁻ content within leaves and roots upon salt stress. When grown in 150 mM NaCl, *pldζ2* mutants contained the least Na⁺ (3 mmol g⁻¹ DW in leaves and 3.9 mmol g⁻¹ DW in roots) (Table 1) and the least Cl⁻ (2.1 mmol g⁻¹ DW in leaves and 3.1 mmol g⁻¹ DW in roots). Na⁺ accumulation concomitant with K⁺ loss was higher in roots than in leaves in all genotypes under salt stress (Table 1). Salt-stressed *pldζ1* mutants showed the most drastic decrease in K⁺

content with 6-fold and 22-fold decreases in leaves and roots, respectively, when treated with 150

- 360 mM NaCl (Table 1). By contrast, only a slight variation in Na⁺/K⁺ ratio was observed in $pld\zeta 2$
- 361 mutant. Salinity effects on K^+ accumulation were less severe in shoots (2-fold decrease) and roots
- 362 (7-fold decrease) in $pld\zeta^2$ than in other genotypes.
- 363

Relative growth rate, relative water content and efficiency of chlorophyll fluorescence in $pld\zeta$ mutants during salt stress

- 366 Relative growth rate (RGR) of WT, $pld\zeta 1$, $pld\zeta 2$ and $pld\zeta 1pld\zeta 2$ had decreased after 14 days of exposure to NaCl (Fig. 7). RGRs of *pld\zeta1pld\zeta2* and *pld\zeta2* mutants were less affected by salt than 367 368 those of $pld\zeta l$ and WT, which were severely inhibited by even mild salt stress. By measuring the 369 change in relative water content (RWC) over time, we found that salt stress induced dehydration in 370 leaves of all genotypes (Fig. 8). *pld* $\zeta 1$ mutants became the most dehydrated with RWC decreases of 371 50 % in 75 mM NaCl and 66 % in 150 mM NaCl compared to the same genotype at the same 372 developmental stage in control conditions (Fig. 8). The $pld\zeta lpld\zeta 2$ leaves only lost 40 % of RWC 373 compared to plants grown in control conditions.
- 374 The Fv/Fm ratio is a convenient measure of photosynthesis efficiency at the photosystem II level
- 375 (PSII). The Fv/Fm ratios for all genotypes were constant in control conditions but decreased under
- 376 salt stress (Fig. 8). After 14 days of salt treatment, the Fv/Fm ratios of $pld\zeta l$ plants had decreased
- 377 by as much as 64 % in 150 mM NaCl. By contrast $pld\zeta 1pld\zeta 2$ plants maintained higher Fv/Fm ratio
- 378 values, which were 1.16 and 1.32-fold higher than those measured in WT, respectively under mild
- and severe stress, suggesting PSII functions better under stress in the absence of PLD ζ (Fig. 8).
- 380
- 381

382 **Discussion**

Recent studies have provided valuable insights into the molecular and cellular mechanisms by which plants respond to and tolerate salinity stress (Deinlein et al. 2014; Julkowska and Testerink 2015; Slama et al. 2015). Although *Arabidopsis thaliana* is considered to be a glycophyte, it has 386 been widely used as a genetic model to investigate salt signalling mechanisms. Salt treatment 387 significantly inhibits growth of A. thaliana with intracellular Na⁺ concentration increasing at the expense of K⁺ (Ghars et al. 2008; Ellouzi et al. 2011). The replacement of K⁺ by Na⁺ affects the 388 389 cell's Na⁺-sensitive enzymes, including components of the photosynthetic machinery, that 390 determine plant growth and yield. Tolerance to salt stress depends on complex signalling networks 391 enabling plants to respond rapidly and efficiently to this constraint (Zhu 2002). Many signal 392 transduction pathways have been shown to be stimulated in response to high salinity (Bragmann et 393 al. 2009; Julkowska and Testerink 2015). Lipid mediators are key components in the signalling 394 network of plant stress adaptation, including adaptation to salinity (Julkowska and Testerink 2015). 395 As an enzyme responsible for producing PA, we focused on the roles of the two A. thaliana PLDC

396 in salt tolerance.

397 To identify changes in gene expression affected by PLD ζ activity, the transcriptome of a double 398 $pld\zeta 1pld\zeta 2$ mutant was compared to the WT transcriptome. In gene cluster 1, the stress-responsive 399 genes DDF1 and RD29A (Thiery et al. 2004; Magome et al. 2004; Magome et al. 2008) were 400 strongly induced by severe salt stress (200 mM NaCl) within 3 h in the WT but were less strongly 401 induced in $pld\zeta lpld\zeta 2$ under the same conditions. Transgenic Arabidopsis lines overexpressing 402 DDF1 were shown to have increased tolerance to high salt stress (170 mM NaCl) by repressing 403 plant growth through the induction of genes coding gibberellin (GA) deactivating enzymes 404 (Magome et al. 2004; Magome et al. 2008). We found that the $pld\zeta^2$ single mutant showed higher 405 expression of *DDF1* and *RD29A* under salt stress compared with the WT and other *pld* ζ mutants 406 (Fig. 2). The expression of auxin-responsive genes such as *IAA5*, *IAA19*, and *GH3-3* was previously 407 found to be reduced in *pldZ* mutants of Arabidopsis in response to external IAA (Li and Xue 408 2007). Therefore, *PLD* ζ ² may be specifically involved in the regulation of gene expression in stress 409 responses and hormonal signalling for growth. Controlled growth reduction may be an effective 410 strategy to save energy and minimize ROS accumulation while facing the deleterious impact of salt 411 stress (Rangani et al. 2016).

412 Transcriptomic analysis revealed a second gene cluster that included ROS detoxification genes such 413 as SOD1 and CCS1, which were overexpressed in $pld\zeta 1pld\zeta 2$ mutants (Suppl. Fig. S5). It is known 414 that ROS act as second messengers in intracellular signalling cascades to trigger plant tolerance to 415 various abiotic and biotic stresses (Ben Rejeb et al. 2014; Mittler, 2016). When Cakile maritima, a 416 salt-tolerant plant, is treated with salt, there is a transient and rapid increase in SOD activity and 417 H₂O₂ content within 4 h. The SOD activity measured in *C. maritima* was eight-fold higher than was 418 found in Arabidopsis (Ellouzi et al. 2013). In the present study, the highest SOD1, CAT2 and APX 419 gene expression was observed in $pld\zeta^2$ mutant under salt stress (Fig. 2). High expression of SOD 420 under salt stress corresponded with high SOD activity and low H_2O_2 levels in leaves and roots of 421 *pldζ2* mutants, indicating that *PLDζ2* participates in the regulation of ROS generation upon salt 422 stress in Arabidopsis. Several reports show that different PLD family enzymes function in concert 423 with ROS to mediate tolerance responses to various abiotic stresses (Hong et al. 2010, Singh et al. 424 2012). For example, mutation in *PLDα1* leads to PA deficiency, reduced plasma membrane

- 425 NADPH oxidase activity, and less ROS in stomata guard cells in response to abscisic acid (Zhang et
- 426 al. 2009). As a consequence, $pld\alpha l$ mutants can not fully close stomata leading to increased water 427 loss (Zhang et al. 2009). We found that $pld\zeta l$ mutants accumulated H₂O₂ faster in leaves and roots
- 428 than *pld* ζ^2 mutants do in response to salt stress. Moreover, *pld* ζ^1 plants displayed the lowest SOD
- 429 activity which, associated with the highest levels of H_2O_2 , implies that *pld* ζl is more sensitive to
- 430 salt stress than WT and the other *pld* ζ mutants. Interestingly other *pld* mutants such as *plda1*, *plda3*
- 431 and $pld\delta$ have also been shown to be hypersensitive to salt treatment (Hong et al. 2008; Bargmann
- 432 et al. 2009).
- 433 Salinity had several striking effects on physiological indexes with distinctions becoming clear 434 between WT and *pld* $\zeta 1$ versus *pld* $\zeta 2$ and *pld* $\zeta 1$ *pld* $\zeta 2$. The *pld* $\zeta 1$ mutant showed the highest Na⁺ 435 content in leaves and roots compared with the other genotypes upon salt stress, leading to a drastic 436 decrease in K^+ content in roots and shoots (Table 1). The loss of K^+ could be explained by 437 downregulation of the expression of genes involved in K⁺ transport like the HAK5 transporter gene. 438 HAK5 is expressed under the control of DDF2 gene, a transcription factor which is homologous to 439 DDF1 (Hong et al. 2013). The massive K efflux could also be mediated by the opening of outward-440 rectifying depolarisation-activated (GORK) channels, an outward K⁺ channel (for review see 441 Anschütz et al., 2014). Phospholipase D ζ could be involved in the regulation of ion channels by
- 442 modulating their lipid environment. For example in yeast cells, many ion channels have been shown
- to be located in plasma membrane microdomains called lipid rafts with channel activities dependent
- 444 on the lipid raft composition (for review see Mollinedo 2012).
- 445 RGR measurements showed that WT and $pld\zeta l$ have only a limited capacity to withstand the
- 446 presence of salt. Sensitivity of WT and $pld\zeta l$ to salt stress was associated with a decline in the leaf
- 447 water content, increase in Na⁺ and Cl⁻ preferentially accumulated in roots, early induction of
- 448 oxidative stress (excess H_2O_2 and MDA), and inhibition of photosynthesis from the toxicity of the
- salt ions (low Fv/Fm ratio). For most markers $pld\zeta l$ was more sensitive than WT, indicating that
- 450 PLD(1 is required in the regulation of growth in normal conditions. Interestingly and as described
- 451 by Ohashi et al. (2003) and Chen et al. (2013), this PLD isoform is greatly involved in the root
- 452 development and growth and the lack of this PLD may lead to enhanced salt stress sensitivity
- 453 (Wang, 2005).

454 Conversely, *pld* ζ^2 mutants tolerated salt stress better than WT. This response may be related to its 455 ability to minimize redistribution of Na^+ and Cl^- to the roots, resulting in the lowest Na^+/K^+ ratio 456 and consequently better retention of K^+ in these organs. pld $\zeta 2$ grew less than pld $\zeta 1$ pld $\zeta 2$ double 457 mutant, which produced more biomass in both roots and leaves. We can consider growth reduction 458 in *pld* $\zeta 2$ to be a strategy to maintain performance under salinity by first controlling ion homeostasis. 459 The activation of the Salt Overly Sensitive (SOS) signalling pathway is a key mechanism for Na⁺ 460 exclusion in roots (Zhu, 2000). PLDs were shown to interact strongly with SOS and plasma 461 membrane transporters under salt stress (Yu et al. 2010). Notably, the activation of PLD in salt 462 stressed tobacco elevated the PA level which is a direct stimulator of SOS1, a central regulator of 463 ion homeostasis (Gardiner et al. 2001). Other studies suggested that there is a functional connection 464 between the activation of *PLD* ζ ² and SOS3, which is more prominent in the roots and has a crucial 465 role in Na^+/K^+ dynamics (Muzi et al. 2016). Much less is known about the cross-talk between PLD 466 and SOS signalling pathways under salt stress. However, the above findings will guide future 467 research to elucidate the mechanism of the salt tolerance in $pld\zeta 2$. 468 It was clear from our results that the *pldZ* mutant was the most tolerant of salt stress as evidenced

by the lowest Na^+/K^+ ratio. It has long been known that salinity stress triggers a dramatic increase in

470 ROS accumulation in plant tissues. Interestingly, our results showed that salt-stressed $pld\zeta^2$ plants

471 also displayed the lowest levels of H_2O_2 and MDA detected in both leaves and roots. In addition,

472 mild and severe salt stresses triggered higher SOD activity in $pld\zeta^2$ than in the other genotypes

473 which could explain why H_2O_2 and MDA levels were lower in both leaves and roots. The oxidative

474 burst indicators SOD, H₂O₂ and MDA indicate the presence of an efficient antioxidant defence in

475 $pld\zeta^2$ mutant. The lack of PLD ζ^2 may therefore allow a better performance of the plants facing

476 moderate and severe salinity due to a primed antioxidant defence system.

In conclusion, our study suggests that in Arabidopsis *PLD* ζ genes encode isoforms that have distinct roles in plant growth particularly under salt stress. The *pld* ζ 2 KO mutant was more salt-tolerant than the WT as growth, water status, ion homeostasis and antioxidant defence systems were all better adjusted to withstanding salinity. Possibly *pld* ζ are differently regulated in halophytes enabling them to tolerate salinity. Looking for natural variants or mutations in *PLD* ζ 2 gene might be a way to improve salt tolerance in glycophytic crops.

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484

485 Author contribution statement

486 AS and CA conceived and designed the research. Data collection, analysis and interpretation were 487 performed by AO, HE, SP and BF. SP conducted the transcriptomic analysis. DDV was involved in 488 the molecular analysis of pld ζ mutant and in obtaining the double pld ζ lpld ζ mutant. PC 489 contributed to study conception and design. AO, HE, SP wrote the manuscript. AS and CA 490 supervised manuscript preparation and correction. All authors read and approved the manuscript.

491

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647Table 1. Ion content (mmol g-1 DW) of leaves and roots of WT and *pld\zeta1*, *pld\zeta2* and *pld\zeta1pld\zeta2*648mutants after 14 days of mild or severe salt stress. Data are means of three replicates ± S.E. Means649indicated by the same superscript letters are not significantly different at P ≤ 0.05 in one-way650ANOVA.

652 Leaves

	Genotyp	e WT	pldζ1	pldζ2	pld[]pld[2
NaCl (mM)	Ion		1 5	1 5	1 5 1 5
	content				
0	Na ⁺	0.14 ± 0.01 ^a	0.14 ± 0.00 ^{ab}	0.14 ±0.01 ^{ab}	0.15 ± 0.00 ^b
75	Na^+	2.36 ± 1.09 ^b	3.30 ± 0.91 ^d	1.33 ± 0.19 a	2.89 ± 0.87 $^{\rm c}$
150	Na^+	$4.50\pm1.06~^{c}$	$7.31 \pm 1.06^{\ d}$	$3.00\pm1.10\ ^a$	$3.92\pm1.08~^{b}$
0	\mathbf{K}^+	2.79 ± 0.24 ^b	1.86 ± 0.02 ^a	2.32 ± 0.21 ab	2.07 ± 0.17 $^{\rm a}$
75	\mathbf{K}^+	1.33 ± 0.15 ^b	0.76 ± 0.03 $^{\rm a}$	1.30 ± 0.27 ^b	0.65 ± 0.01 a
150	\mathbf{K}^+	0.77 ± 0.05 $^{\rm b}$	0.31 ± 0.00 a	1.12 ± 0.22 b	0.28 ± 0.02 a
0	Cl-	0.03 ± 0.00^{a}	0.03 ± 0.00^{a}	0.09 ± 0.01 b	0.02 ± 0.00^{a}
75	Cl-	2.37 ± 0.55 ^c	2.14 ± 0.32 ^a	1.66 ± 0.32 ^b	1.70 ± 0.09 ^a
150	Cl	$3.08\pm0.85~^{\text{b}}$	$3.88\pm0.66^{\ d}$	2.09 ± 0.53 a	3.23 ± 1.021 ^c
0	Na ⁺ /K ⁺	0.04 ± 0.00 ^a	0.07 ± 0.00 ^c	0.06 ± 0.00^{b}	0.07 ± 0.00 ^c
75	Na^+/K^+	1.80 ± 0.06 ^b	4.38 ± 0.19 ^c	1.05 ± 0.11 a	4.56 ± 0.30 ^c
150	Na^+/K^+	$5.88\pm0.44~^{b}$	$23.88\pm0.22~^{d}$	2.88 ± 0.51 a	14.30 ± 1.40 ^c
Roots					
0	Na ⁺	$0.12\pm0.00~^a$	0.11 ± 0.01 ^a	0.22 ± 0.01 ^b	0.13 ± 0.00 ^a
75	Na^+	3.73 ± 1.11 ^b	$3.72 \pm 1.19^{\text{ b}}$	$2.91\pm0.66~^a$	2.89 ± 0.97 $^{\rm a}$
150	Na ⁺	6.43 ± 1.31 ^b	$8.42\pm1.16\ ^{c}$	$3.92\pm0.91~^a$	5.88 ± 1.08 ^b
0	\mathbf{K}^+	1.14 ± 0.21 ^b	0.65 ± 0.24 $^{\rm a}$	0.98 ± 0.07 ab	0.94 ± 0.02 ^{ab}
75	\mathbf{K}^+	0.40 ± 0.01 ^c	0.10 ± 0.00 ^a	0.52 ± 0.02 d	$0.28\pm0.00~^{b}$
150	\mathbf{K}^+	$0.09\pm0.01~^{\text{b}}$	0.04 ± 0.00 a	0.15 ± 0.00 c	$0.08\pm0.00~^{\text{b}}$
0	Cl-	0.04 ± 0.00 ^a	0.03 ± 0.00 ^a	0.06 ± 0.01 ^b	$0.03 \pm 0.00^{\text{ a}}$
75	Cl-	4.10 ± 0.91 ^c	4.08 ± 0.90 ^c	1.91 ± 0.36 ^a	3.50 ± 0.33 ^b
150	Cl	5.08 ± 1.05 $^{\rm c}$	$5.72\pm0.28~^{d}$	3.00 ± 0.83 ^a	$4.00\pm0.83~^{b}$
0	Na ⁺ /K ⁺	$0.11 \pm 0.00^{\text{ a}}$	0.21 ± 0.06 ^{ab}	0.23 ± 0.01 ^b	0.13 ± 0.01 ab
75	Na^+/K^+	9.61 ± 0.52 ^b	35.14 ± 2.75 ^c	5.54 ± 0.13^{a}	10.22 ± 1.05 ^b
150	Na^+/K^+	70.67 ± 2.81^{b}	$246.95 \pm 40.1^{\text{d}}$	55.41 ± 20.0^{a}	$72.91 \pm 1.30^{\circ}$



Figure 1. Numbers of differentially expressed genes in WT and $pld\zeta 1pld\zeta 2$ transcriptome comparisons. Venn diagrams show how gene clusters were defined. (**a**) Cluster 1 (intersection in bold) includes genes which are induced by salt stress in WT and are expressed at lower levels in the double mutant under salt stress than in WT. (**b**) Cluster 2 (intersection in bold) includes genes which are up or down regulated in $pld\zeta 1pld\zeta 2$ compared to WT in both control and salt stress conditions.

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Figure 2. Gene expression in wild-type (WT), $pld\zeta 1$, $pld\zeta 2$ and $pld\zeta 1\zeta 2$ plants treated (+) or not treated (-) with 200 mM NaCl for 3 h. *DDF1* and *RD29A* (**a**), and *APX1*, *SOD1* and *CAT2* (**b**) expression was estimated by RT-PCR with *APT1* as a loading control. (**c**) Quantitative RT-PCR of *DDF1* and *RD29A* expression in WT, $pld\zeta 1$, $pld\zeta 2$ and $pld\zeta 1\zeta 2$ seedlings. *DDF1* transcript abundance was expressed as a ratio of the value for WT in control conditions. *APT1* gene expression was used as a standard. Mean and standard errors are based on three technical repeats





Figure 3. Changes in superoxide dismutase (SOD) activity in leaves and roots of wild type and *pld* ζ 1, *pld* ζ 2 and *pld* ζ 1*pld* ζ 2 mutants of *A. thaliana* plants grown for 14 days under moderate (75 mM NaCl) or severe salinity (150 mM NaCl). Data are means ± SE of 3 replicates. Results of statistical analysis are given in each panel (ANOVA, $P \le 0.05$). NS, not significant.



Figure 4 Histochemical detection of hydrogen peroxide in leaves (a) and roots (b) of wild type and *pld\zeta 1, pld\zeta 2* and *pld\zeta 1pld\zeta 2* mutants of *A. thaliana* plants grown for 14 days under moderate (75 mM NaCl) or severe salinity (150 mM NaCl). Brown residue from diaminobenzidine staining indicates sites of H₂O₂ accumulation.



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Figure 5. Changes in hydrogen peroxide levels in leaves and roots of wild type and *pld* $\zeta 1$, *pld* $\zeta 2$ and *pld* $\zeta 1$ *pld* $\zeta 2$ mutants of *A. thaliana* plants grown for 14 days with moderate (75 mM NaCl) or severe salinity (150 mM NaCl). Data are means \pm SE of 3 replicates. Results of statistical analysis are given in each panel (ANOVA, $P \le 0.05$). NS, not significant.



Figure 6. Changes in malondialdehyde (MDA) levels in leaves and roots of wild type and *pld* $\zeta 1$, *pld* $\zeta 2$ and *pld* $\zeta 1$ *pld* $\zeta 2$ mutants of *A. thaliana* plants grown for 14 days under moderate (75 mM NaCl) or severe salinity (150 mM NaCl). Data are means \pm SE of 3 replicates. Results of statistical analysis are given in each panel (ANOVA, $P \le 0.05$). NS, not significant.



Figure 7. Relative growth rate (RGR) in *A. thaliana* single and double $pld\zeta$ mutants grown for 14 days under control conditions or moderate (75 mM NaCl) or severe salinity (150 mM NaCl). Means indicated by different letters are significantly different (P \leq 0.05) as determined by one-way ANOVA.



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Figure 8. Changes in the relative water content (RWC) and in *Fv/Fm* ratios of leaves and roots of wild type, and *pldζ1*, *pldζ2* and *pldζ1pldζ2* mutants of *A. thaliana* plants grown for 14 days under moderate (75 mM NaCl) or severe salinity (150 mM NaCl). Data are means \pm SE of 3 replicates. Results of statistical analysis (ANOVA, $P \le 0.05$) are given in each panel. NS, not significant

b





Supplementary Figure S1 Map and genotyping of $pld\zeta 1$, $pld\zeta 2$ and $pld\zeta 1pld\zeta 2$ mutants. (a) Map of T-DNA insertion site in $pld\zeta 1$ and $pld\zeta 2$ genes. Primers used for PCR are represented by arrows. (b) 1 % agarose gels of PCR products for $PLD\zeta 1$ WT allele (A+B), $pld\zeta 1$ T-DNA flanking sequence (A+C), or both (A+B+C) in hemizygous plants (top gel). PCR products for WT PLDZ WT allele (D+E), $pld\zeta 2$ T-DNA flanking sequence (D+C), or both (D+E+C) in hemizygous plants (middle gel). WT allele amplification (A+B), $pld\zeta 1$ T-DNA flanking sequence (A+C) and $pld\zeta 2$ T-DNA flanking sequence (D+C) in $pld\zeta 1$ T-DNA flanking sequence (A+C) and $pld\zeta 2$ T-DNA flanking sequence (D+C).



Supplementary Figure S2 Venn diagram of WT transcriptomes under salt and mannitol stresses in comparison to control growth conditions. Numbers of differentially expressed genes for each comparison are shown.

		WT+NaCl / WI	` control	pldζ1pldζ2+Na	Cl / WT+NaCl
AGI	Name	Log2 Ratio	Pvalue	Log2 Ratio	Pvalue
AT1G12610	DDF1 transcription factor	1.96	0.00E+0	-1.89	0.00E+0
AT1G73480	alpha/beta hydrolase family	3.40	0.00E+0	-1.04	0.00E+0
AT5G52310	COR78 (COLD REGULATED 78) (RD29A)	4.04	0.00E+0	-1.03	5.60E-12
AT2G16500	ADC1_ arginine decarboxylase 1	0.89	1.17E-4	-0.98	1.12E-10
AT1G58360	AAP1; amino acid permease 1	1.95	0.00E+0	-0.86	1.77E-7
AT2G38240	oxidoreductase. 2OG-Fe(II) oxygenase	1.23	2.24E-11	-0.81	2.50E-6
AT5G13750	ZIFL1 zinc induced facilitator-like 1	1.81	0.00E+0	-0.78	9.68E-6
AT2G46370	JAR1 (JASMONATE RESISTANT 1)	0.83	1.15E-3	-0.76	2.66E-5
AT4G26080	ABI1 (ABA INSENSITIVE 1) PP2C family	1.48	0.00E+0	-0.75	4.51E-5
AT1G51090	metal ion binding	2.99	0.00E+0	-0.74	1.05E-4
AT1G72770	HAB1; PP2C homologue to ABI1	2.53	0.00E+0	-0.73	1.22E-4
AT3G07700	Protein kinase superfamily protein	0.90	8.80E-5	-0.73	1.57E-4
AT1G01650	SPPL4(SIGNAL PEPTIDE PEPTIDASE-LIKE 4)	0.73	4.31E-2	-0.71	3.41E-4
AT3G48990	AMP-dependent synthetase and ligase	0.75	2.14E-2	-0.71	3.61E-4
AT1G29400	AML5 (ARABIDOPSIS MEI2-LIKE PROTEIN 5)	0.79	5.23E-3	-0.70	5.50E-4
AT5G17760	P-loop nucleoside triphosphate hydrolase	1.43	0.00E+0	-0.70	6.32E-4
AT1G76960	unknown protein	0.85	7.24E-4	-0.67	2.25E-3
AT2G35940	EDA29 (BEL1-like homeodomain 1)	0.80	3.45E-3	-0.65	5.59E-3
AT4G15530	PPDK pyruvate orthophosphate dikinase	0.90	1.04E-4	-0.64	8.57E-3
AT5G55970	RING/U-box superfamily protein	1.25	5.59E-12	-0.61	2.51E-2

Supplementary Figure S3 List of genes in Cluster 1. Genes are involved in abiotic or biotic stimulus and response to stress. Expression data is given as log_2 ratios.

	Normed to Freq. In Arabidopsis set (± bootstrap		
Cluster 1 (Figure 1)	StdDev, p-value)		
Biological process	Frequence in Cluster 1	Std dev	P value
response to abiotic or biotic stimulus (Input set freq.: 0.65; 0.13)	4.96	0.795	0.0000009492
other biological processes (Input set freq.: 0.45; 0.12)	3.65	0.874	0.0002548
response to stress (Input set freq.: 0.5; 0.14)	3.45	0.681	0.0001537
developmental processes (Input set freq.: 0.4; 0.13)	3.02	0.91	0.002149
transport (Input set freq.: 0.35; 0.12)	2.82	0.814	0.006256
signal transduction (Input set freq.: 0.15; 0.07)	2.12	1.185	0.115

	Normed to Freq. In Arab	oidopsis s	et (± bootstrap
Cluster 2 (Figure 4)	StdDev, p-value)		
Biological process	Frequence in Cluster 2	Std dev	P value
electron transport or energy pathways (Input set freq.: 0.28; 0.02)	13.07	4.609	4.157e-06
response to stress (Input set freq.: 0.76; 0.13)	5.47	0.807	1.850e-10
response to abiotic or biotic stimulus (Input set freq.: 0.61; 0.13)	4.62	0.885	2.780e-07
other biological processes (Input set freq.: 0.57; 0.13)	4.28	0.815	2.565e-06
transport (Input set freq.: 0.38; 0.1)	3.57	1.227	7.785e-04
cell organization and biogenesis (Input set freq.: 0.33; 0.11)	2.81	0.94	6.493e-03
developmental processes (Input set freq.: 0.33; 0.12)	2.62	0.781	9.223e-03
signal transduction (Input set freq.: 0.14; 0.05)	2.53	1.084	0.083

http://bar.utoronto.ca/ntools/cgi-bin/ntools_classification_superviewer.cgi#annotation_list

Supplementary Figure S4 SuperViewer classification for genes listed in clusters 1 and 2.

		n14/21/22 / X	VT control	n11/21/2+NaC	u / WT±N₀Cl
AGI	Name	$I \circ \sigma^2 Ratio$	Pyalua	Log2 Ratio	Pyalua
AT5G35935	unknown protein copia-like	1.64	0.00E+0	1.73	0.00E+0
AT1G23390	unknown protein	1.28	0.00E+0	0.78	1 40E-5
AT1G08830	CSD1: copper, zinc superoxide dismutase	1.27	0.00E+0	0.78	1.10E 5
AT1G66100	toxin receptor binding thionin putative	1.23	0.00E+0	0.65	4 41E-3
AT1G12520	CCS1: superoxide dismutase copper chaperone	1.04	5 30E-9	0.81	2.04E-6
AT3G02380	COL2 transcription factor	0.81	3.05E-4	0.64	7.26E-3
AT5G48485	DIR1 lipid transfer protein (LTP)	0.74	3 99E-3	0.60	4 57E-2
AT4G09320	NDPK1 nucleoside diphosphate kinase type 1	0.72	7 93E-3	0.73	1.57E 2
AT3G48990	AMP-dependent synthetase and ligase	-0.84	7 33E-5	-1.06	0.00E+0
AT5G08530	CI5. NADH-ubiquinone oxidoreductase 51 kDa	-0.85	5 72E-5	-0.95	9 52E-10
AT3G23820	GAE6	-0.87	2.15E-5	-1.09	0.00E+0
AT5G05170	CESA3 (CELLULASE SYNTHASE 3)	-0.92	2.63E-6	-0.98	1 12E-10
AT4G32410	CESA1 (CELLULASE SYNTHASE 1)	-0.94	9.08E-7	-1.00	3.92E-11
AT3G46970	ATPHS2	-1.01	3.21E-8	-1.09	0.00E+0
AT1G59870	ABC transporter family protein	-1.08	8.23E-10	-1.04	0.00E+0
AT2G25490	EBF1 (EIN3-BINDING F BOX PROTEIN 1)	-1.15	1.12E-11	-1.31	0.00E+0
AT4G38770	PRP4 (PROLINE-RICH PROTEIN 4)	-1.29	0.00E+0	-0.94	1.15E-9
AT3G60750	transketolase transketolase. putative	-1.31	0.00E+0	-1.01	1.12E-11
AT3G33002	ribosomal protein S2p family	-1.33	0.00E+0	-1.11	0.00E+0
AT5G42020	ATP binding luminal binding protein 2 (BiP-2)	-1.38	0.00E+0	-1.53	0.00E+0
AT3G09440	HSP70-3	-1.61	0.00E+0	-1.38	0.00E+0
AT4G30650	low temperature and salt responsive protein	-1.82	0.00E+0	-1.11	0.00E+0
AT5G02500	HSC70-1	-1.83	0.00E+0	-1.39	0.00E+0

Supplementary Figure S5 List of genes in cluster 2. Genes are involved in electron transport, biotic and abiotic stress and stress responses. Expression data are given as log_2 ratios.

Supplementary Table S1 List of primer sequences used for RT-PCR analysis and genotyping.

For RT-PCR

Gene name	Accession number	5'-3' Sequence
DDF1-F	AT1G12610	GGGACTTATCCCACAGCAGA
DDF1-R	AT1G12610	ATCATTGGATTCCGGCACC
RD29AF	AT5G52310	CAAAACAGAGCACTTACACAGAGAA
RD29AR	AT5G52310	CATAATCTCTACCCGACACACTTTT
APT1F	AT1G27450	GAGACATTTTGCGTGGGATT
APT1R	AT1G27450	CGGGGATTTTAAGTGGAACA
APX1F	X59600.1	CTGACATTCCTTTCCACCCT
APX1R	X59600.1	CAGACCTATCCTTGTGGCAT
CSD1F	AT1G08830	GGTTTCCATGTCCATGCTCT
CSD1R	AT1G08830	ATTGTGAAGGTGGCAGTTCC
CAT2F	AT4G35090.1	AACTCTGGTGCTCCTGTATGG
CAT2R	AT4G35090.1	CTCCAGTTCTCTTGGATGTG

For genotyping

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А	PLDzeta1F	At3g16785	TCAGAATCACTTAAGAGGAGATGGG
В	PLDzeta1R	At3g16785	TTTTCGCATAGTCACTTGCTGT
С	LBb1	T-DNA left border	GCGTGGACCGCTTGCTGCAACT
D	PLDzeta2F	At3g05630	TCTCTGTTTTGGGCGGTACGA
Е	PLDzeta2R	At3g05630	AAAATGTTCAGCGTTCTGGAT