1	Transcriptome profiles of eggplant (Solanum melongena) and its wild relative S.
2	dasyphyllum under different levels of osmotic stress provide insights into
3	response mechanisms to drought
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23	Abstract
24	Defence mechanisms to abiotic stresses, like drought, are very broad and RNA
25	sequencing (RNA-Seq) can help in understanding the complex responses triggered. In
26	this study, we performed RNA-Seq of the transcriptomes of eggplant (Solanum
27	melongena) and its related wild species (S. dasyphyllum) under two PEG concentrations

29 plant phenological stages (three and five true fully developed leaves). Solanum

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(20% and 30%), two different times (after 0.5 h and 2 h of osmotic stress) and at two

30 *dasyphyllum* was more tolerant to osmotic stress, and a differential expression pattern of 31 drought-related genes was identified between the two species. Plants subjected to a 32 higher osmotic potential, at a more adult stage and at a higher stress exposure time 33 displayed a higher number of DEGs (differential expressed genes). Gene ontology (GO) 34 enrichment analysis revealed that, compared to S. melongena, S. dasyphyllum triggered 35 the regulation of a wide range of transcription factors (AP2/ERF, DREB, bZIP, WRKY 36 and bHLH). In both species, the abscisic acid (ABA) signaling response pathway played 37 a crucial role leading to stomatal closure. Other important pathways involved in abiotic 38 stresses tolerance including flavonoid, carotenoid and phenylpropanoid biosynthesis, 39 chlorophyll metabolism and photosynthesis pathway among others were found to have a 40 relevant role under both moderate and severe osmotic stresses. Our results reveal that S. 41 *dasyphyllum* is a potential source of genes for breeding resilient eggplant varieties.

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Keywords: transcriptome, RNA Seq, osmotic stress, drought, *Solanum melongena*, *S. dasyphyllum*.

46

47 **1. Introduction**

48 Drought spells occur naturally in many areas of the world, but climate change has 49 accelerated and intensified them, with dramatic consequences on agriculture [1]. 50 Projections indicate that the risk and severity of drought episodes will increase across 51 the subtropics and mid-latitudes in both hemispheres as a consequence of global 52 warming and decreased regional precipitation [2,3]. Drought stress triggers 53 morphological, physiological, biochemical, cellular and molecular response mechanisms in plants with a potentially severe reduction in plant growth and crop 54 55 production as a major consequence. Therefore, determining plant response and tolerance 56 mechanisms against drought stress is fundamental to mitigating its effects [4,5].

The development of new molecular and bioinformatics tools has allowed the expansion of applied knowledge in breeding programs. In this way, transcriptomics has provided new potential resources for studying the molecular response of abiotic stress in crops [6], being RNA sequencing (RNA-Seq) the general method of choice. This method allows a broad coverage of the transcriptome, providing a significant characterization of

62 mRNA transcripts of specific tissue and time and, in addition, is a quantitative method

that yields a digital gene expression atlas at a genomic scale [7].

64 Drought tolerance is a complex trait involving different components at the physiological, biochemical and genetic levels [8]. Osmotic stress, resulting in an 65 increased difficulty for water uptake by the roots, is one of the most important factors in 66 67 drought [9]. To unravel the effects of water deficit in genetic networks, the use of a 68 solution containing polyethylene glycol (PEG) in hydroponic culture is a common practice to induce osmotic stress and reduce the water potential of tissues in plants 69 70 [10,11]. In this way, the transcriptome of PEG-treated plants provides information regarding drought-related genes, which can be primarily classified in protective and 71 72 regulatory genes [7]. Regarding the former, these are genes that encode LEA proteins, 73 chaperones, osmoprotectants, water channels, ion exchangers, and enzymes involved in 74 the osmolyte biosynthesis and the reactive oxygen species (ROS), among others 75 [12,13]. On the other hand, genes encoding regulatory proteins act on the expression of 76 stress-responsive, including transcription factors, protein kinases and phosphatases, 77 enzymes involved in phospholipid metabolism and abscisic acid (ABA) biosynthesis 78 and epigenetic-related genes [14,15].

79 Crop wild relatives (CWRs) are an increasingly fundamental resource for plant breeding 80 to improve the adaptative capacity of agricultural systems to climate change-related 81 stresses [16]. Among vegetable crops, eggplant (Solanum melongena L.) can be highly 82 benefited by introgression breeding, as many eggplant CWRs thrive in areas affected by 83 moderate to severe drought [17]. Eggplant is an important crop, being the eighth 84 vegetable crop in terms of cultivated area in the world, being widely grown in Asia, 85 Africa and Europe [18]. It has been described as a relatively drought-tolerant crop and 86 different degrees of drought tolerance have been observed in cultivated accessions and 87 CRWs [19–21]. Among these CWRs, S. dasyphyllum Schumach. and Thonn. grows 88 naturally in areas where drought spells are frequent and it has been reported to exhibit 89 significant drought tolerance both under field and experimental conditions [19,22]. It is 90 considered the wild ancestor of the gboma eggplant (S. macrocarpon L.) [23,24] and is classified in the Anguivi clade, which includes several African and Southeast Asian 91 92 "prickly" species [17,25]. Solanum dasyphyllum is a member of the secondary genepool 93 of eggplant [26], and interspecific hybrids and advanced backcross materials of *S*.
94 *dasyphyllum* with *S. melongena* have been obtained [27,28].

95 In the present study, we analyzed the transcriptomes of a cultivated S. melongena and a drought-tolerant S. dasyphyllum accessions under PEG-induced osmotic stress in two 96 97 different plant phenological stages and at two times for each phenological stage. By evaluating its physiological responses in conjunction with the analysis of the gene 98 99 expression we aimed at a better comprehensive understanding of the different response 100 mechanisms against osmotic stress in these materials. The results are of great interest 101 for a better understanding of drought tolerance and to foster introgression breeding of drought-tolerant resilient cultivars in eggplant. 102

103

104 2. Material and Methods

105 *2.1. Plant materials and growth conditions*

Solanum melongena MEL1 and S. dasyphyllum DAS1 accessions were used for the present study. Seeds were germinated according to Ranil et al. [29] protocol for uniform eggplant CWRs germination and plants were grown in hydroponic culture according to Renau-Morata et al. [30] with Hoagland solution [31] in a growth chamber with a 16/8 h light/dark photoperiod, 25°C temperature and 60-65% of humidity. The nutrient solution was resupplied every four days and an air compressor was used to supply aeration.

113 2.2. PEG-induced osmotic stress

To evaluate the effect of the plant phenological stage and the stress response, two 114 115 osmotic stress experiments were conducted using PEG 6000 (Bio Basic Inc., Ontario, 116 Canada). One experiment was performed with 20% PEG at a phenological stage of three fully developed true leaves (Ex 1), while the other with 30% PEG at the five fully 117 developed true leaves stage (Ex 2). In each experiment, leaves of three biological 118 replicates (i.e., three different plants uniformly developed, each one constituting a 119 120 replicate and for each one a library was developed) were taken for each species at three 121 times: 0 h (control; T0), 0.5 h (T0.5) and 2 h (T2) after initiation of the stress treatment. 122 Immediately, leaf samples were frozen with liquid nitrogen and stored at -80°C for RNA 123 extraction. Plant symptoms were registered at different times of the treatments.

124 2.3. RNA extraction, sequencing and data processing

Total RNA was extracted from leaves samples of each biological replicate using 125 126 TRIzolTM Reagent (Invitrogen, Carlsbad, CA, USA). For each of the 36 replicates, the RNA library was performed by Novogene Co., LTD (Beijing, China) and sequenced on 127 an Illumina NovaSeq 6000 (paired-end 150 bp). Raw data in FASTQ format were 128 filtered by removing reads with adaptor contamination, reads containing N > 10% and 129 130 low-quality reads (Qscore of over 50% bases below 5). Error rate (%), Q20 (%), Q30 131 (%) and GC content (%) were calculated for data quality control of clean data. Gene expression levels were estimated by calculating fragments per kilobase of transcript 132 sequence per millions of base pairs sequenced (FPKM). 133

134 *2.4. Transcriptomic analysis*

Differentially expressed genes (DEGs) analysis was performed using the DEseq2 R package [32], and the resulting *p*-values were adjusted using Benjamini and Hochberg's correction for controlling the false discovery rate (FDR) [33]. Genes with adjusted *p*value $\square < \square 0.05$ and $||og_2(fold change)|\square > \square 1$ were considered as differentially expressed. DEGs were annotated based on the functional annotation information of genes of the eggplant reference genome "67/3" V3 [34]. Venn diagrams of DEGs were displayed using jvenn, a plug-in for the jQuery JavaScript library [35].

Hierarchical clustering analysis was carried out of log₂ (FPKM+1) of union differential
expression genes, within all comparison groups. Heatmaps were performed selecting
drought-related DEGs, based on the scientific literature, which were classified
according to their function into four groups: osmoprotectants, phytohormones, protein
kinases and transcription factors using the web tool ClustVis [36].

Gene ontology (GO, http://www.geneontology.org/) and Kyoto Encyclopedia of Genes and Genomes (KEGG, http://www.genome.jp/kegg/) enrichment analyses of the DEGs were performed. The tomato (*S. lycopersicum* L.) KEGG pathways annotated database was used for the analysis, being the closest species with more comprehensive and reliable information. GO and KEGG terms with an adjusted *p*-value $\square < \square 0.05$ were considered significantly enriched for the DEGs.

153

154 **3. Results**

155 *3.1. Physiological responses to osmotic stress*

As a general trend, in both experiments, *S. dasyphyllum* (DAS) displayed a better water stress tolerance than *S. melongena* (MEL). In Ex_1, DAS presented visual symptoms only at T2 while MEL started to show symptoms of stress at T0.5 (Figure 1). In Ex_2, manifestations of water stress in plants were observed at T0.5 and T2 in both species in a faster way with more severe symptoms compared with Ex_1, although DAS, again, exhibit more tolerance, with fewer symptoms of wilting (Figure 1).

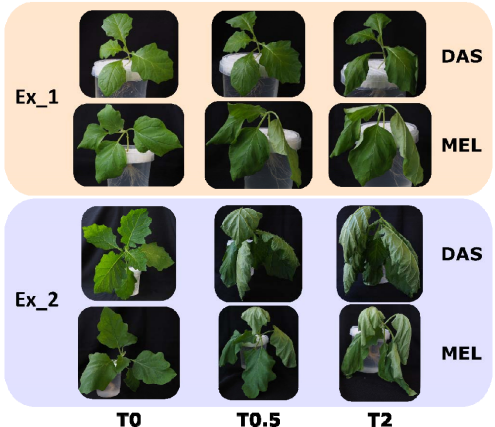


Figure 1. Representative phenotypes of *S. melongena* (MEL) and *S. dasyphyllum* (DAS) after 0, 0.5 and 2h of PEG stress in hydroponic culture in both experiments (Ex_1 and Ex_2).

165

166 *3.2. Differential gene expression over time in response to PEG treatment*

After filtering raw sequencing data, clean reads showed an error rate between 0.02 and 0.03%, an average Q30 of 93.85% and GC content of 43.17% (Table S1). For each

169 experiment, DEGs with an adjusted *p*-value < 0.05 and a $|\log_2 (\text{fold change})| > 1$ were

selected by performing pairwise comparisons at each time of PEG treatment (T0.5 and

171 T2) with the non-stressed control (T0).

172 In Ex_1 (20% PEG and three fully developed true leaves stage), a total of 894 and 433 DEGs were detected for DAS and MEL, respectively. For DAS a total of 114 (74 up-173 regulated [UR], 40 down-regulated [DR] and 33 related to drought stress) and 840 174 DEGs (475 UR, 365 DR and 171 related to drought stress) were detected at T0.5 and 175 176 T2, respectively (Table 1). For MEL, a total of 327 (273 UR, 54 DR and 89 related to 177 drought stress) and 117 DEGs (76 UR, 41 DR and 24 related to drought stress) were detected at T0.5 and T2, respectively (Table 1). Venn diagram analysis showed that in 178 DAS 52 DEGs were commonly regulated at T0.5 and T2 while 49 and 707 DEGs were 179 specific at 0.5 and T2, respectively (Figure 2A). In MEL, seven DEGs were commonly 180 regulated after both times of treatment, 273 and 67 DEGs at T0.5 and T2 respectively 181 182 (Figure 2A).

In Ex_2 (30% PEG and five fully developed true leaves) a total of 2,037 and 4,375183 184 DEGs were detected for DAS and MEL, respectively. For DAS, a total of 147 (109 UR 185 38 DR and 53 related to drought stress) and 1,999 DEGs (1,040 UR, 959 DR and 363 related to drought stress) were detected at T0.5 and T2, respectively (Table 1). For 186 187 MEL, a total of 198 (134 UR, 64 DR and 62 related to drought stress) and 4,360 DEGs 188 (2,252 UR, 2,108 DR and 774 related to drought stress) were detected at T0.5 and T2, respectively (Table 1). Venn diagram analysis showed that 31 and 80 DEGs were 189 commonly regulated at T0.5 and T2 exclusively in DAS and MEL respectively (Figure 190 191 2B). A total of 20 and 1,065 DEGs were detected only in DAS at T0.5 and T2 192 respectively. In MEL, 11 and 3,363 DEGs were detected exclusively at T0.5 and T2 193 respectively. A total of 38 common DEGs were detected for both times and both 194 accessions. (Figure 2B).

Table 1. Differentially expressed genes that were up-regulated or down-regulated after
0.5 h (T0.5) and 2 h (T2) of PEG stress in *S. dasyphyllum* (DAS) and *S. melongena*

197 (MEL) in experiments 1 and 2 (Ex_1 and Ex_2).

Experiment	Time	Number of DEGs		Up-Regulated DEGs		Down-Regulated DEGs		Number of DEGs related to drought	
	comparison	DAS	MEL	DAS	MEL	DAS	MEL	DAS	MEL
Ex_1	T0 vs T0.5	114	327	74	273	40	54	33	89

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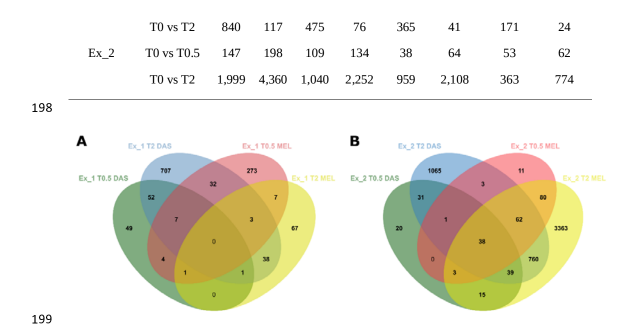
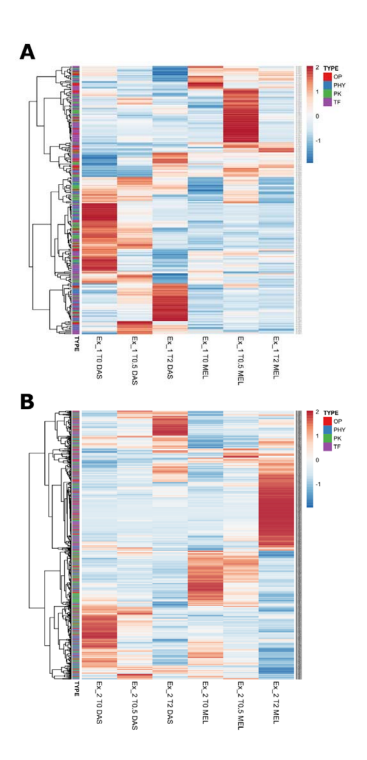


Figure 2. Venn diagram of DEGs under 0.5 and 2h of PEG stress of *S. dasyphyllum*(DAS) and *S. melongena* (MEL) in experiment 1 (Ex_1; A) and experiment 2 (Ex_2;

202 **B**).

203 Drought-responsive DEGs were classified according to their function into four groups: osmoprotectants, phytohormones, protein kinases and transcription factors related to the 204 205 drought stress response. A total of 264 DEGs related to drought were observed in Ex 1, 206 of which 38 of them were genes related to osmoprotectants, 46 were related to the synthesis of phytohormones, 67 were protein kinases genes and 113 were transcription 207 factors. In Ex_2 a total of 953 DEGs were detected, of which 150 were genes that 208 209 encode for proteins related to osmoprotectants, 180 were related to phytohormones, 296 210 for protein kinases and 327 transcription factors genes (Figure 3).

In both experiments, in general, the expression pattern of drought-responsive genes changed over time for both accessions, allowing clear differentiation between accessions and time of exposure to stress. In all cases, up-regulated and down-regulated genes from the different groups of the classification were observed (Figure 3).



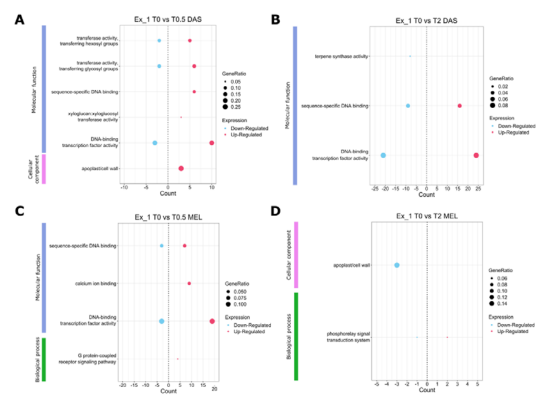
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Figure 3. Heatmap of DEGs related to drought stress, osmoprotectants (OP), phytohormones (PHY), protein kinases (PK) and transcription factor (TF) after 0, 0.5 and 2 h of PEG stress of *S. dasyphyllum* (DAS) and *S. melongena* (MEL) in experiments 1 (Ex_1; A) and 2 (Ex_2; B).

3.3. GO and KEGG enrichment in DEGs according to phenological stage and stress conditions

222 A gene ontology (GO) analysis was performed with DEGs being annotated as a 223 biological process (BP), cellular components (CC) and molecular function (MF). In 224 Ex 1, for DAS at T0.5, 37 DEGs were annotated as MF, 13 of them as DNA-binding transcription factor activity (10 UR and three DR), three as xyloglucan:xyloglucosyl 225 transferase activity (UR), six as sequence-specific DNA binding (UR), eight as 226 227 transferase activity (transferring glycosyl groups; six UR and two DR), seven as 228 transferase activity (transferring hexosyl groups; five UR and two DR). Regarding CC, 229 three were annotated as apoplast and cell wall (Figure 4A). After 2 h of osmotic stress 230 (T2), in DAS, all significant DEGs were annotated as MF, 45 as DNA-binding transcription factor activity (24 UR and 21 DR), 25 as sequence-specific DNA binding 231 232 (16 UR and nine DR) and eight as terpene synthase activity (DR; Figure 4B). For MEL, 233 at T0.5, a total of 45 DEGs were annotated as MF, 22 of them as DNA-binding 234 transcription factor activity (19 UR and three DR), nine as calcium ion binding (UR) 235 and 10 as sequence-specific DNA binding (seven UR and three DR) and four as G protein-coupled receptor signaling pathway as biological process (UR; Figure 4C). In 236 237 MEL at T2, three CC DEGs were annotated as apoplast and cell wall (DR) and three BP DEGs as phosphorelay signal transduction system (two UR and one DR; Figure 4D). 238

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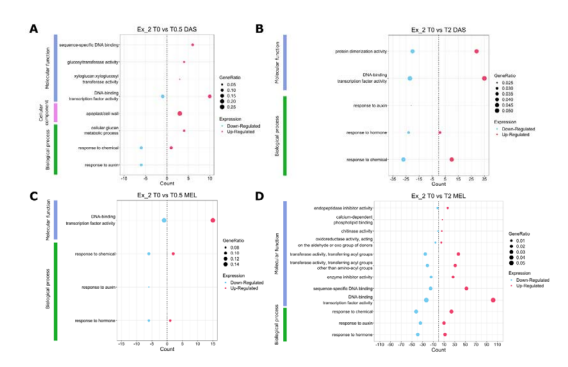
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Figure 4. Gene ontology (GO) terms enrichment scatter plot in DEGs of *S. dasyphyllum*(DAS) after 0.5 (T0.5) (A) and 2 h (T2) (B) versus 0 h of PEG stress and *S. melongena*(MEL) after 0.5 (T0.5) (C) and 2 h (T2) (D) compared with 0 h of PEG stress in
experiment 1 (Ex_1).

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In Ex 2 for DAS at T0.5, a total of 44 DEGs were annotated, 17 of them as BP, three as 246 CC and 24 as MF. Within the biological process category, seven DEGs were annotated 247 as a response to chemical (one UR and six DR), six as response to auxin (DR) and four 248 249 as cellular glucan metabolic process (UR), while as CC, three as apoplast and cell wall 250 (UR). As molecular function, 11 were annotated as DNA-binding transcription factor activity (10 UR and one DR), three as xyloglucan:xyloglucosyl transferase activity 251 252 (UR), four as glucosyltransferase activity (UR) and six as sequence-specific DNA 253 binding (UR; Figure 5A). At T2, in DAS, a total of 82 DEGs were annotated as BP, 37 as response to chemical (10 UR and 27 DR), 24 as response to hormone (one UR and 23 254 DR) and 21 as response to auxin (DR). As MF, 49 DEGs as protein dimerization 255 activity (29 UR and 20 DR) and 57 as DNA-binding transcription factor activity (35 UR 256 257 and 22 DR) (Figure 5B). For MEL, significant GO terms annotated for BP at T0.5 were seven to response to hormone (one UR and six DR), six to response to auxin (DR) and 258

eight to response to chemical (two UR and six DR). Also, 16 DEGs were annotated as 259 DNA-binding transcription factor activity (15 UR and one DR) in MF classification 260 (Figure 5C). At T2, 161 DEGs were classified as BP, 51 as response to hormone (12 UR 261 and 39 DR), 44 as response to auxin (10 UR and 34 DR) and 66 as response to chemical 262 (24 UR and 42 DR). As molecular function, 125 were annotated as DNA-binding 263 transcription factor activity (102 UR and 23 DR), 72 as sequence-specific DNA binding 264 (52 UR and 15 DR), 41 as enzyme inhibitor activity (27 UR and 14 DR), 52 as 265 266 transferase activity (transferring acyl groups other than amino-acyl groups; 31 UR and 21 DR), 62 as transferase activity (transferring acyl groups; 37 UR and 25 DR), 11 267 oxidoreductase activity (acting on the aldehyde or oxo group of donors as; five UR and 268 six DR), seven as chitinase activity (six UR and one DR), seven as calcium-dependent 269 phospholipid binding (UR) and 19 as endopeptidase inhibitor activity (17 UR and two 270 271 DR) (Figure 5D). Enriched genes annotated in each GO term classification were included in Table S2. 272



273

Figure 5. Gene ontology (GO) terms enrichment scatter plot in DEGs of *S. dasyphyllum*(D) after 0.5 h (T0.5) (A) and 2 h (T2) (B) versus 0 h of PEG stress and *S. melongena*(M) after 0.5 h (T0.5) (C) and 2 h (T2) (D) compared with 0h of PEG stress experiment
277 2 (Ex_2).

279 A pathway enrichment analysis using the Kyoto Encyclopedia of Genes and Genomes (KEGG) was performed to identify significant (padj < 0.05) enriched metabolic or 280 signal transduction pathways associated with differentially expressed genes (DEGs) 281 282 comparing the whole genome background. In Ex_1 more DEGs were assigned to KEGG pathways in DAS than in MEL. For DAS, at T0.5 and T2, plant hormone signal 283 284 transduction and MAPK (mitogen activated protein kinase) signaling pathway were 285 identified as enriched pathway (five UR and five DR DEGs). For T2 were also 286 determined circadian rhythm (seven UR and two DR DEGs), sesquiterpenoid and 287 triterpenoid biosynthesis (seven DR DEGs), galactose metabolism (seven UR DEGs) and zeatin biosynthesis (one UR and eight DR) as enriched pathways. For MEL at T0.5, 288 289 plant-pathogen interaction (10 UR DEGs) and, at T2, circadian rhythm (eight UR and 290 two DR DEGs) were enriched pathways detected (Table 2). In Ex_2, more expressed 291 genes were assigned to metabolic pathways in MEL than for DAS. For DAS at T0.5, 292 DEGs were assigned to plant hormone signal transduction (six UR and eight DR) and 293 also to MAPK signaling pathway (three UR and four DR). At T2, plant hormone signal 294 transduction (16 UR and 29 DR DEGs) and the phenylpropanoid biosynthesis (17 UR 295 and 11 DR) were determined as enriched pathways. For MEL at T0.5, plant hormone 296 signal transduction (nine UR and six DR DEGs), MAPK signaling pathway (three UR 297 and four DR), fatty acid elongation (three DR), and carotenoid biosynthesis (three UR) were found to be enriched pathways. At T2, DEGs were linked to porphyrin and 298 chlorophyll metabolism (seven UR and 21 DR), plant hormone signal transduction (58 299 UR and 37 DR), photosynthesis and antenna proteins (10 DR), α -linolenic acid 300 metabolism (24 UR and two DR MAPK signaling pathway (53 UR and 16 DR), 301 302 flavonoid biosynthesis (nine UR and nine DR and glutathione metabolism (21 UR and 303 seven DR) (Table 2). Enriched genes annotated in each KEGG pathway classification 304 were included in Table S3.

305

Table 2. Significant Kyoto Encyclopedia of Genes and Genomes (KEGG) enriched
pathways and its ID of tomato database in *S. dasyphyllum* (DAS) and *S. melongena*(MEL) after 0.5 h (T0.5) and 2 h (T2) of PEG stress in experiments 1 (Ex_1) and 2
(Ex_2).

Experiment	Time	Accession	Pathway Terms	ID	Count	Up-	Down-
Experiment	comparison	Accession	Tatilway Terms	ID	Count	Regulated	Regulated

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	T0 vs T0.5	DAS	Plant hormone signal transduction	sly04075	10	5	5
			MAPK signaling pathway - plant	sly04016	8	3	5
		MEL	Plant-pathogen interaction	sly04626	10	10	0
	T0 vs T2	DAS	Plant hormone signal transduction	sly04075	32	12	20
Ex_1			MAPK signaling pathway - plant	sly04016	27	10	17
			Circadian rhythm - plant	sly04712	9	7	2
			Sesquiterpenoid and triterpenoid biosynthesis	sly00909	7	0	7
			Galactose metabolism	sly00052	7	7	0
			Zeatin biosynthesis	s1y00908	8	1	7
	-	MEL	Circadian rhythm - plant	sly04712	10	8	2
		DAS	Plant hormone signal transduction	sly04075	14	6	8
	- T0 vs T0.5		MAPK signaling pathway - plant	sly04016	7	3	4
		MEL	Plant hormone signal transduction	sly04075	15	9	6
			MAPK signaling pathway - plant	sly04016	10	7	3
			Fatty acid elongation	sly00062	3	0	3
			Carotenoid biosynthesis	sly00906	3	3	0
	T0 vs T2	DAS	Plant hormone signal transduction	sly04075	45	16	29
Ex_2			Phenylpropanoid biosynthesis	sly00940	28	17	11
		MEL	Porphyrin and chlorophyll metabolism	s1y00860	28	7	21
			Plant hormone signal transduction	sly04075	95	58	37
			Photosynthesis - antenna proteins	sly00196	10	0	10
			α-Linolenic acid metabolism	sly00592	26	24	2
			MAPK signaling pathway - plant	sly04016	69	53	16
			Flavonoid biosynthesis	sly00941	18	9	9
			Glutathione metabolism	sly00480	28	21	7

311

312 4. Discussion

313 Eggplant has been considered a relatively drought-tolerant crop since a long time ago 314 [37] and several studies to evaluate the physiological and biochemical responses to 315 water stress of different eggplant cultivars and wild relatives have been performed [21,38,39]. However, detailed molecular mechanisms in response to drought stress in 316 eggplant are not well known and, to our knowledge, transcriptional analysis by RNA-317 Seq method has not been reported so far. In the current study, we evaluated plants of the 318 cultivated eggplant S. melongena and its wild relative S. dasyphyllum under two 319 320 concentrations of PEG (20% and 30%) at two different phenological stages (three and five fully developed true leaves) in hydroponic conditions in order to obtain a general 321 322 overview of their response to osmotic stress and get insight in the gene expression 323 involved in response and tolerance to drought. Solanum dasyphyllum displayed a better 324 water deficit tolerance than S. melongena, confirming its already recently reported 325 drought tolerance in field and experimental conditions [19,22]. PEG concentration had a 326 visually significant effect in physiological response, with more symptoms in Ex_2, in 327 which plants were subjected to a higher PEG concentration, resulting in a higher 328 osmotic potential [41].

RNA sequencing is a tool for transcriptome analysis that has allowed a better 329 330 understanding of the functions of the genome [42]. In this research, the analysis of 331 differential gene expression has enabled the study of the response to osmotic stress in both species at the genomic level. One of the most important components of drought 332 333 stress is osmotic stress and it has been widely used to study drought tolerance in many 334 species [9]. In our study, in general, osmotic stress treatments mainly triggered an 335 activation response, as more significantly up-regulated than down-regulated DEGs were 336 observed. The number of DEGs increased as PEG concentration was higher and longer 337 in time, as was previously reported in potato (Solanum tuberosum L.) [43,44]. The 338 expression pattern of drought-responsive genes displayed large differences between S. dasyphyllum and S. melongena, revealing very divergent response mechanisms under an 339 osmotic stress according to plant physiological observations. 340

This study has disclosed the main functions and pathways expressed of two related species with large differences in osmotic stress response. GO enrichment of the

identified DEGs has allowed establishing the biological functions associated to those 343 genes. Solanum dasyphyllum expressed genes were involved in diverse functions related 344 to osmotic stress response. On one side, genes involved in the modification of cell wall 345 346 and apoplast structure, such as xyloglucan:xyloglucosyl transferases [45,46], were enriched in the wild species. Other up-regulated genes in S. dasyphyllum in response to 347 osmotic stress were classified in the DNA-binding transcription factor activity and 348 349 sequence-specific DNA binding GO terms, including a wide range of transcription 350 factors (TFs), which exert crucial roles in diverse signaling pathways in different abiotic 351 stress response as AP2/ERF (APETALA2/Ethylene Response Factor) family [47] and 352 two of its major subfamilies such as dehydration-responsive element binding proteins 353 (DREBs) and ethylene-responsive element (ERE) binding factors [48,49]. The same occurs with TFs, from homeobox-leucine zipper family [50–52], basic leucine zipper 354 355 (bZIP) [53] and WRKY family [54]. Meanwhile, in the case of S. melongena, the expression of AP2/ERF, WRKY and bZIP TFs was also observed, however, in general, 356 357 the number of differential genes expressed under the stress treatments was fewer. When 358 plants were subjected to the higher osmotic potential, the overall gene expression was 359 also higher and included DEGs classified in response to chemical and hormones and 360 also down-regulated genes related to auxin response. Auxins are involved in the regulation of plant growth and development and auxin response factors (ARFs) gene 361 family play an essential role in the regulation of auxin-relative genes in abiotic stress 362 responses in tomato (S. lycopersicum) [55]. Basic helix-loop-helix (bHLH) transcription 363 factors were overexpressed in S. dasyphyllum, which they have been reported to be 364 365 involved in the response to abiotic stresses in potato (S. tuberosum) [56] and pepper 366 (Capsicum annuum L.) [57]. For S. melongena, exposure to a higher osmotic stress 367 resulted in the differential expression of genes related to enzyme inhibitor activity, transferases, chitinase and oxidoreductase activities, among others. The overall response 368 observed was very broad, with the wild species (S. dasyphyllum) showing a greater and 369 370 more diverse expression of genes involved in drought response, which could be related 371 to its increased tolerance.

KEGG analysis revealed significant enriched pathways related to osmotic stress such as
plant hormone signal transduction and MAPK signaling. In these pathways, genes
encoding for the three main components of the core Abscisic Acid (ABA) signaling
response were up-regulated, a pathway that has been widely reported as a key drought

stress response [58]. Among those genes, protein phosphatases type-2C (PP2Cs), ABA 376 receptors PYRPYR/PYL/RCAR (PYRABACTIN-RESISTANCE 1/PYRABACTIN 377 RESISTANCE LIKE/REGULATORY COMPONENT OF ABA RECEPTOR) and 378 379 SNF1-Related Protein Kinases type 2 (SnRK2s) were identified as DEGs [59]. Although PP2Cs are negative regulators of ABA signalling, an increased relative 380 expression under drought stress conditions has been reported in other similar studies 381 382 [60–62], suggesting that these apparent contrasting effects need to be further 383 investigated. AREB/ABF transcription factors and MAPKKs (mitogen activated protein 384 kinase kinase) were also activated as a response to ABA signaling, which leads to 385 stomatal closure, one of the most important drought responses [59,63]. Solanum 386 dasyphyllum displayed a wide variety of response mechanisms along with the ABA pathway. These included galactinol synthase and transferases related genes, which have 387 388 been reported to improve drought tolerance [64]. Also, zeatin biosynthesis was downregulated, in particular the cytokinin signaling repressors A-type ARABIDOPSIS 389 390 RESPONSE REGULATORS (ARRs), which have been reported to negatively regulate 391 by drought stress, promoting cell division in meristems [65-67]. In addition, 392 GIGANTEA (GI) protein synthesis was activated, which is a regulator in the circadian 393 rhythm plant pathway and improves drought tolerance [68]. Finally, phenylpropanoid 394 biosynthesis pathway was detected, which exhibits different important roles in the regulation under abiotic stress conditions [69]. On the other hand, S. melongena showed 395 different drought response pathways, including the carotenoid biosynthesis, which has 396 been reported to have a similar regulation in S. tuberosum [60], the inactivation of 397 398 porphyrin, chlorophyll metabolism and photosynthesis pathways as a consequence of 399 the osmotic stress [70]. Furthermore, the regulation of flavonoid biosynthesis, which 400 has an important role in coping with environmental stress [69], the expression of plant 401 glutathione transferases (GSTs), which has been reported to be involved in responses to biotic and abiotic stress [71], and the synthesis of the stress signaling molecule, such as 402 jasmonic acid (JA) by the metabolism of α -Linolenic acid [72] were linked to osmotic 403 404 stress. When the plants are more adult and under a more intense osmotic stress, ABA signaling response leads to stomatal closure and to the down regulation of small auxin 405 406 up-regulated RNA (SAUR) genes, which induce plant growth [73]. In our study, a 407 common response as stress adaptation has been observed, including ABA signaling 408 response and inhibition of plant growth.

409

410 **5.** Conclusions

411 The present work provides an overview of the osmotic stress response at the 412 transcriptomic level of cultivated eggplant (S. melongena) and its drought-tolerant wild 413 relative S. dasyphyllum. We have found that osmotic potential and plant phenological stage play a crucial role in the response, which is increased when the exposure time was 414 415 longer and osmotic stress was more intense. Our data showed that response 416 mechanisms at the gene expression level were very wide-ranging, including transcription factors, phytohormones, osmoprotectants and protein kinases, being ABA 417 response signaling an important pathway. Clear differences observed between the two 418 species in the response to osmotic stress and overall gene expression pattern confirmed 419 420 that S. dasyphyllum is a potential source for breeding to drought tolerance in eggplant. 421 Overall, our work provided insights into the gene expression mechanisms of tolerance 422 to osmotic stress in eggplant and its wild relative S. dasyphyllum, which is of great 423 relevance in the improvement of drought tolerance of cultivated eggplant.

424

425 Supplementary information

426 Supplementary Table S1. Data quality summary of samples of experiment 1 and 2,
427 after 0, 0.5 and 2 h of PEG stress of *S. dasyphyllum* (DAS) and *S. melongena* (MEL).

428 Supplementary Table S2. Gene ontology (GO) terms enrichment and regulation, ID,

description and transcription factor family of DEGs in *S. dasyphyllum* after 0.5 (T0.5)

(A) and 2 h (T2) versus 0h of PEG stress and *S. melongena* (M) after 0.5 (T0.5) and 2 h

431 (T2) compared with 0h of PEG stress experiment 2 (Ex_2).

Supplementary Table S3. Significant Kyoto Encyclopedia of Genes and Genomes
(KEGG) enriched pathways, its ID of tomato database, and regulation ID, description
and transcription factor family of DEGs in *S. dasyphyllum* (D) and *S. melongena* (M)
after 0.5 h (T0.5) and 2 h (T2) of PEG stress in experiments 1 (Ex_1) and 2 (Ex_2).

436

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447

448 **Declaration of Competing Interest**

449 The authors declare that they have no known competing financial interests or personal

450 relationships that could have appeared to influence the work reported in this paper.

451

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