1	Mutations in HUA2 restore flowering in the Arabidopsis trehalose
2	6-phosphate synthase1 (tps1) mutant
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28 Abstract

Plant growth and development are regulated by many factors, including carbohydrate 29 availability and signaling. Trehalose 6-phosphate (T6P), which is synthesized by 30 TREHALOSE-6-PHOSPHATE SYNTHASE 1 (TPS1), is positively correlated with and 31 32 functions as a signal that informs the cell about the carbohydrate status. Mutations in TPS1 negatively affect the growth and development of Arabidopsis thaliana and complete 33 loss-of-function alleles are embryo lethal, which can be overcome using inducible expression 34 of TPS1 (GVG::TPS1) during embryogenesis. Using EMS mutagenesis in combination with 35 genome re-sequencing we have identified several alleles in the floral regulator HUA2 that 36 restore flowering and embryogenesis in *tps1-2 GVG::TPS1*. Genetic analyses using a *HUA2* 37 T-DNA insertion allele, *hua2-4*, confirmed this finding. RNA-seq analyses demonstrated that 38 hua2-4 has widespread effects on the tps1-2 GVG::TPS1 transcriptome, including key genes 39 40 and pathways involved in regulating flowering. Higher order mutants combining tps1-2 GVG::TPS1 and hua2-4 with alleles in the key flowering time regulators FLOWERING 41 LOCUS T (FT), SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1 (SOC1), and 42 FLOWERING LOCUS C (FLC) were constructed to analyze the role of HUA2 during floral 43 44 transition in tps1-2 in more detail. Taken together, our findings demonstrate that loss of HUA2 can restore flowering and embryogenesis in tps1-2 GVG::TPS1 in part through 45 activation of FT, with contributions of the upstream regulators SOC1 and FLC. Interestingly, 46 we found that mutation of FLC is sufficient to induce flowering in tps1-2 GVG::TPS1. 47 48 Furthermore, we observed that mutations in HUA2 modulate carbohydrate signaling and that this regulation might contribute to flowering in *hua2-4 tps1-2 GVG::TPS1*. 49

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51 **Keywords:** carbohydrate signaling, Trehalose 6-phosphate (T6P), *TREHALOSE*

52 PHOSPHATE SYNTHASE1 (TPS1), HUA2, Flowering time, Arabidopsis thaliana

53 Introduction

Plants have evolved intricate signaling mechanisms that enable them to monitor a wide range of environmental and endogenous cues and adjust their physiology, growth, and development accordingly. Adjustments occur more or less constantly, but developmental phase transitions such as germination, the switch from juvenile to adult growth, or the induction of flowering and reproductive development are under particularly stringent control.

In Arabidopsis thaliana, the floral transition is controlled by environmental factors including 59 exposure to prolonged periods of cold (vernalization), ambient temperature, day length 60 (photoperiod), light quality, and endogenous signals such as plant age, diverse hormones 61 including gibberellic acid (GA), and carbohydrate signaling (Srikanth and Schmid, 2011; 62 Romera-Branchat et al., 2014; Cho et al., 2017). Eventually, these signaling pathways 63 converge on and regulate the expression of key floral integrator genes such as FLOWERING 64 65 LOCUS T (FT) and SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1 (SOC1) (Kardailsky et al., 1999; Moon et al., 2005; Kobayashi and Weigel, 2007; Turck et al., 2008; 66 Lee and Lee, 2010; Jung et al., 2012). FT is induced in response to permissive photoperiod in 67 the leaf vasculature where it is also translated. The FT protein is then transported via the 68 69 phloem to the shoot apical meristem (SAM) where it interacts with the bZIP transcription factor FD and 14-3-3 proteins to form the florigen activation complex (FAC) (Abe et al., 70 2005; Wigge et al., 2005; Mathieu et al., 2007; Taoka et al., 2011; Collani et al., 2019). In 71 contrast, SOC1 is induced and acts largely at the SAM, both downstream and in parallel to FT 72 (Yoo et al., 2005; Lee and Lee, 2010). Eventually, these factors induce flower meristem 73 identity genes such as LEAFY (LFY) and APETALA1 (AP1) at the SAM, thus completing the 74 floral transition (Weigel and Nilsson, 1995; Liljegren et al., 1999; Blázquez and Weigel, 75 2000). 76

Apart from photoperiod, carbohydrate signaling has been shown to be necessary for *FT* expression (Wahl et al., 2013). Sucrose is the major product of photosynthesis and most common transport-sugar. However, rather than measuring sucrose concentration directly, plants employ trehalose 6-phosphate (T6P) as a readout and signal of sucrose availability (Goddijn and van Dun, 1999; Lunn et al., 2006; Martins et al., 2013; Yadav et al., 2014;

Figueroa and Lunn, 2016). T6P is the intermediate of trehalose synthesis. It is synthesized from glucose 6-phosphate and uridine diphosphate glucose by TREHALOSE 6-PHOSPHATE SYNTHASE (TPS) and subsequently dephosphorylated by TREHALOSE 6-PHOSPHATE PHOSPHATASE (TPP) (Cabib and Leloir, 1958).

In Arabidopsis thaliana, there are 11 TPS genes (AtTPS1-AtTPS11), which can be divided 86 into two subclades, class I and class II, and 10 TPP genes (TPPA-TPPJ) (Leyman et al., 2001; 87 Lunn, 2007; Vandesteene et al., 2012). Of the class I TPS genes (AtTPS1-AtTPS4), only 88 AtTPS1, AtTPS2, and AtTPS4 have demonstrable catalytic activity, whereas AtTPS3 harbors a 89 90 premature translational stop codon and is likely a pseudogene (Blázquez et al., 1998; Van Dijck et al., 2002; Lunn, 2007; Delorge et al., 2015). Class II TPS genes (AtTPS5-AtTPS11), 91 for which no TPS activity, has been detected, which have been reported to participate in cell 92 size regulation, thermotolerance, and cold and salt resistance, but the underlying molecular 93 94 mechanisms remain largely unclear (Chary et al., 2008; Ramon et al., 2009; Singh et al., 2011; Tian et al., 2019; Van Leene et al., 2022). The main T6P synthase in Arabidopsis thaliana is 95 TPS1. TPS1 loss-of-function mutants are embryonic lethal (Eastmond et al., 2002), but 96 homozygous tps1-2 mutants could be established by dexamethasone-inducible expression of 97 TPS1 (GVG::TPS1) during embryogenesis (van Dijken et al., 2004). Interestingly, the 98 resulting homozygous tps1-2 GVG:TPS1 plants flower extremely late compared to wild type 99 100 under both short- and long-day conditions. At the molecular level, late flowering of tps1-2 GVG::TPS1 has been attributed to the combined misregulation of key flowering time genes. 101 In particular, tps1-2 GVG::TPS1 mutant plants fail to induce FT in leaves even under 102 permissive photoperiod. In addition, MIR156 and its targets, the SOUAMOSA PROMOTER 103 BINDING PROTEIN LIKE (SPL) genes, which together constitute the age pathway, are also 104 misregulated in tps1-2 GVG::TPS1 (Wahl et al., 2013). Nevertheless, many questions 105 regarding the regulation of plant growth and development by the T6P pathway remain open. 106

In an EMS suppressor screen, we have recently reported dozens of mutations that partially 107 108 restored flowering and seed set in tps1-2 GVG::TPS1, including several alleles in SNF1 109 KINASE HOMOLOG 10 (*KIN10*) and HOMOLOG OFYEAST **SUCROSE** NONFERMENTING 4 (SNF4), two subunits of Arabidopsis thaliana SNF1-Related Kinase 1 110

(*SnRK1*) (Jung et al., 2012; Zacharaki et al., 2022), an evolutionary conserved regulator of
cellular energy homeostasis.

Here, we identified several new alleles in HUA2 (At5g23150) that partially rescue the tps1-2113 GVG::TPS1 phenotype. Mutations in HUA2 were originally identified in a genetic screen as 114 enhancers of the AGAMOUS (AG) allele ag-4 (Chen and Meyerowitz, 1999). In addition, 115 HUA2 has also been reported to affect shoot morphology and function as a repressor of 116 flowering (Doyle et al., 2005; Wang et al., 2007). At the molecular level, HUA2 has been 117 suggested to function as a putative transcription factor but has also been implicated in RNA 118 119 processing (Cheng et al., 2003). We show that three different EMS-induced point mutations in HUA2 restore flowering in tps1-2 GVG::TPS1 and verify this finding using a previously 120 described T-DNA insertion allele, hua2-4. RNA-seq analyses revealed widespread effects of 121 hua2-4 on the tps1 GVG::TPS1 transcriptome, including activation of flower integrator genes 122 123 such as SOC1 and AGAMOUS-LIKE 24 (AGL24). Genetic analyses demonstrated that induction of flowering in tps1-2 GVG::TPS1 required functional FT. Furthermore, we 124 observed that loss of FLOWERING LOCUS C (FLC) is sufficient to induce flowering in 125 tps1-2 GVG::TPS1. Interestingly, hua2-4 also attenuated the induction of known SnRK1 126 target genes in response to carbon starvation. Taken together, our results identify mutations in 127 HUA2 as suppressors of the non-flowering phenotype of tps1-2 GVG::TPS1 and provide 128 insights into the underlying genetic and molecular pathways. 129

130

131 **Results**

132 Mutations in *hua2* restore flowering in *tps1-2 GVG::TPS1*

To identify novel components of the T6P pathway, we recently conducted a suppressor screen in which the non-flowering *tps1-2 GVG::TPS1* mutant was subjected to ethyl methane sulfonate (EMS) mutagenesis. In total, 106 M2 mutant plants in which flowering and seed set was at least partially restored were isolated, and EMS-induced SNPs were identified by whole genome sequencing in a subset of 65 mutants (Zacharaki et al., 2022). To identify

additional candidate suppressor genes in which SNPs were overrepresented, we expanded thislist to 92 by sequencing the genomes of another 27 mutants (Table S1).

Analysis of these 92 genome sequences for genes with multiple independent EMS-induced 140 mutations identified three SNPs in the coding sequence of HUA2 (AT5G23150) (Table S2, 141 142 S3). The three alleles result in non-synonymous amino acid substitutions, namely A983T, P455S, and R902C. We refer to these new EMS-induced suppressor lines as hua2-11 (line 143 #8-1-1), hua2-12 (line #233-14-1), and hua2-13 (line #164-9-1), respectively (Fig. 1A). The 144 polymorphism R902C resides at the C-terminal end of the HUA2 CID motif (RNA Pol-II 145 146 C-terminal domain (CTD) interaction domain). The hua2-11 (line #8-1-1) allele was also detected in two additional suppressor lines, #57-2-1 and #30-34 (Table S2, S3). As these 147 three lines share most EMS-induced SNPs genome-wide, we assume these lines originate 148 from the same parental plant. 149

Importantly, flowering was restored in all three *hua2* alleles, even though all three mutant 150 lines produced substantially more leaves before making the transition to flowering than Col-0 151 152 control plants (Fig. 1B, C). The flowering time of hua2-11 was 32.15 days, whereas hua2-12 and hua2-13 flowered after 46.5 and 50.9 days, respectively, compared to Col-0, which 153 flowered after 25.2 days. Thus, the three mutants form an allelic series with *hua2-11* being 154 the strongest and hua2-13 being the weakest allele. As HUA2 has previously been implicated 155 in flowering time regulation and has been shown to regulate the expression of a group of 156 MADS-box transcription factors known to form a floral repressive complex in Arabidopsis 157 thaliana (Doyle et al., 2005; Wang et al., 2007; Lee et al., 2013; Posé et al., 2013; Jali et al., 158 2014; Yan et al., 2016), we considered mutations in this gene as likely to be causal for the 159 160 restoration of flowering in the tps1-2 GVG::TPS1 suppressor lines.

Since the three *hua2* alleles described above were generated through EMS mutagenesis, it is possible that other independent mutations not linked to *HUA2* could be involved in partially rescuing the *tps1-2 GVG::TPS1* phenotype. To confirm that mutations in *HUA2* are causal for the suppression of the *tps1-2* non-flowering phenotype, we crossed *tps1-2 GVG::TPS1* with *hua2-4*, a previously described *hua2* loss-of-function mutant that carries a T-DNA insertion in the 2^{nd} intron (Fig. 2A) (Doyle et al., 2005). Of the F2 plants homozygous for the *tps1-2* mutations, only those approx. 25% that were homozygous for the *hua2-4* T-DNA insertion flowered without application of dexamethasone. Similar to *hua2-11 tps1-2 GVG::TPS1* (Fig. 1B,C), *hua2-4 tps1-2 GVG::TPS1* double mutants displayed a bushy shoot phenotype and were moderately late flowering (Fig. 2B,C). Taken together, our findings confirm that recessive mutations in *HUA2* are responsible for the induction of flowering in *tps1-2 GVG::TPS1*. Our findings also suggest that *HUA2* normally functions by repressing flowering either directly or indirectly through the promotion of floral repressors.

hua2-4 has widespread effects on the *tps1-2 GVG::TPS1* transcriptome

To identify possible downstream targets of *HUA2* whose misexpression might explain the induction of flowering in the suppressor mutant, we performed RNA-seq analysis in leaves of 21-d-old *tps1-2 GVG::TPS1* plants, *tps1-2 GVG::TPS1* plants treated with dexamethasone, and the *hua2-4 tps1-2 GVG::TPS1* double mutant. Plants were grown under long days (16 h light, 8 h dark) in the presence or absence of dexamethasone and samples were collected at ZT4 (Zeitgeber time 4, means 4 h after lights on). Genes that were differentially expressed in three independent replicates per genotype and treatment were identified using Cuffdiff.

We observed that dexamethasone treatment significantly affected the expression of 9600 182 genes in tps1-2 GVG::TPS1. Of these, 4830 and 4770 genes were upregulated and 183 downregulated, respectively (Fig. 3A). In contrast, mutation of hua2 affected the expression 184 185 of only 2066 genes, of which 988 and 1078 genes were upregulated and downregulated in hua2-4 tps1-2 GVG::TPS1, respectively (Fig. 3A). In total our RNA-seq analysis identified 186 1437 genes that are differentially expressed in tps1-2 GVG::TPS1 in response to 187 dexamethasone application and the hua2-4 mutation. Importantly, HUA2 expression is not 188 189 changed in tps1-2 GVG::TPS1 in response to dexamethasone application, suggesting that *hua2* might induce flowering largely by activating a pathway not normally regulated by the 190 T6P pathway (Fig. S1). 191

Since both, dexamethasone application and mutations in *hua2* can induce flowering in *tps1-2* GVG::TPS1, we next searched for genes that were repressed or induced in response to either treatment. We identified 412 genes that were downregulated in *tps1-2* GVG::TPS1 in response to dexamethasone application and mutations in *hua2* (Fig. 3A). Gene ontology (GO) analysis revealed that among others, processes such as flavonoid metabolism (GO:0009812), carbohydrate transport (GO:0008643), and starvation response (GO:0009267) were significantly enriched, which is in line with the well-established role of *TPS1* in remodeling carbohydrate metabolism (Fig. 3B; Table S4).

In addition, we identified 243 genes that were induced in response to dexamethasone and in 200 hua2-4 tps1-2 GVG::TPS1. Among these genes, GO categories related to the response to 201 gibberellin (GO:0009739) and the regulation of timing of meristematic phase transition 202 203 (GO:0048506) are of particular interest as they are directly linked to the transition to flowering (Fig. 3C; Table S5). Importantly, among the genes induced in tps1-2 GVG::TPS1 204 by dexamethasone and hua2 were SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1 205 (SOC1) and AGAMOUS-LIKE 24 (AGL24), two MADS-domain transcription factors known 206 207 to promote the transition to flowering (Fig. 3D; Table S6). In contrast, other known flowering time regulators such as CONSTANS (CO), FT, and TWIN SISTER OF FT (TSF) are either 208 hardly detectable (Fig. S2A), possibly because of the collection time of the RNA-seq samples 209 at ZT 4 or did not change significantly in *hua2* and in response to dexamethasone treatment 210 (Fig. S2B). In summary, our transcriptome analysis identified several downstream genes and 211 pathways whose misregulation could contribute to the induction of flowering in tps1-2 212 GVG::TPS1 in response to dexamethasone application or loss of hua2 (Fig. S2; Table S6). 213

Induction of flowering of *tps1-2 GVG::TPS1* by *hua2-4* requires *FT*

To test whether SOC1, which we found to be differentially expressed in response to 215 dexamethasone application or in hua2-4 tps1-2 GVG::TPS1, is a major target of HUA2 in the 216 regulation of flowering time in tps1-2 GVG::TPS1 we constructed the soc1-2 hua2-4 tps1-2 217 GVG::TPS1 triple mutant. We observed that the triple mutant flowered only moderately later 218 than the hua2-4 tps1-2 GVG::TPS1 double mutant (Fig. 4A,B). This indicates that even 219 though SOC1 is significantly induced in our RNA-seq experiment in hua2-4 tps1-2 220 GVG::TPS1 (Fig. 3D; Table S6) and in RT-qPCR experiments (Fig. 4C), SOC1 is largely 221 dispensable for the induction of flowering in tps1-2 GVG::TPS1 by loss of hua2. 222

SOC1 is known to act partially upstream of the flowering time integrator gene and florigen 223 FT. We, therefore, decided to test if induction of flowering in tps1-2 GVG::TPS1 by hua2-4 224 required functional FT. Interestingly, mutation of FT completely abolished the effect of 225 hua2-4 on flowering of tps1-2 GVG::TPS1 and the ft-10 hua2-4 tps1-2 GVG::TPS1 triple 226 mutant failed to flower even after four months of growth in inductive long-day conditions 227 (Fig. 4D,E). In line with this observation, we detected increased expression of FT at the end 228 of the long day (ZT 16) in the hua2-4 tps1-2 GVG::TPS1 double mutant when compared to 229 *tps1-2 GVG::TPS1* (Fig. 4F). It is interesting to note that *FT* expression was barely detectable 230 at ZT 4 according to our RNA-seq analysis (Fig. S2A), which is in agreement with the 231 diurnal expression pattern reported for FT (Kobayashi et al., 1999). Taken together, our 232 genetic and molecular analyses indicate that hua2-4 induces flowering of tps1-2 GVG::TPS1 233 in part through activation of FT, with minor contributions of the upstream regulators SOC1. 234

Loss of *FLC* induces flowering in *tps1-2 GVG::TPS1*

HUA2 has previously been reported to regulate flowering at least in part by regulating the 236 expression of floral repressors of the MADS-domain transcription factor family, including 237 FLOWERING LOCUS C (FLC) and FLOWERING LOCUS M (FLM) (Doyle et al., 2005). To 238 239 test if hua2-4 induces flowering in tps-2 GVG::TPS1 through these repressors we constructed the flc-3 hua2-4 tps1-2 GVG::TPS1 triple mutant. We found that this triple mutant flowered 240 moderately earlier than hua2-4 tps1-2 GVG::TPS1 (Fig. 4G,H). In agreement with these 241 findings, RT-qPCR analysis failed to detect FLC expression in the hua2-4 tps1-2 GVG::TPS1 242 mutant, whereas FLC expression was readily detectable by RT-qPCR in tps1-2 GVG::TPS1 243 (Fig. 4I). 244

Furthermore, we found that the expression of *FLC* was significantly upregulated in 18-day-old *tps1-2 GVG::TPS1* seedlings when compared to Col-0 in publicly available RNA-seq data (Zacharaki et al., 2022) (Fig. 5A). This prompted us to test loss off *FLC* on its own might be sufficient to suppress the non-flowering phenotype of *tps1-2 GVG::TPS1*. Indeed, we observed that *flc-3* alone is capable of inducing flowering in the otherwise non-flowering *tps1-2 GVG::TPS1* mutant background, even though the *flc-3 tps1-2 GVG::TPS1* double mutant flowered significantly later than wild-type and *flc-3* (Fig. 5B,C). These findings suggest that the failure of *tps1-2 GVG::TPS1* to flower could in part be due to *FLC*, possibly in conjunction with other MADS-box repressors such as *MADS AFFECTING FLOWERING 5 (MAF5)*, the expression of which was also elevated in *tps1-2 GVG::TPS1* (Fig. 5A). In contrast, expression of *HUA2* was not changed in *tps1-2 GVG::TPS1* when compared to Col-0 according to publicly available RNA-seq data (Fig. S3).

257 *hua2-4* attenuates carbon starvation responses

The above data indicate that mutations in HUA2 bypass the requirement for TPS1 to induce 258 flowering by reducing expression of MADS-box floral repressors and ultimately inducing 259 floral integrator genes such as FT and SOC1. However, carbohydrate signaling has been 260 shown to also indirectly regulate phase transitions, including flowering, in A. thaliana 261 (Corbesier et al., 1998; Gibson, 2005; Xing et al., 2015; Wang et al., 2020). In part, this 262 response is mediated by SnRK1, which in response to stress conditions such as extended 263 264 darkness phosphorylates a range of proteins, including several C- and S1-class bZIP transcription factors. Activation of these transcription factors by SnRK1 induces expression 265 of including SENESCENCE5 (SEN5) DARK 266 stress response genes, and INDUCED6/ASPARAGINE SYNTHASE1 (DIN6/ASN1), which can be used as a proxy for 267 268 SnRK1 activity (Delatte et al., 2011; Dietrich et al., 2011; Mair et al., 2015). To test if loss of HUA2 might affect flowering also more indirectly by modulating cellular energy responses, 269 we analyzed the expression of SEN5 and DIN6. Interestingly, we found that induction of 270 SEN5 and DIN6 in response to extended night was strongly attenuated in hua2-4 (Fig. 6A, B) 271 272 similar to what we had previously observed in mutants affected in SnRK1 subunits (Zacharaki et al., 2022). This finding indicates that mutations in HUA2 might modulate 273 carbohydrate signaling more directly and that this regulation might contribute to the induction 274 of flowering in hua2-4 tps1-2 GVG::TPS1. In agreement with this hypothesis, we found that 275 276 expression of SEN5 and DIN6 was even further attenuated in three independent hua2-4 tps1-2 277 GVG::TPS1 lines (Fig. 6A,B).

278

279 **Discussion**

Arabidopsis thaliana HUA2 has been reported to play a crucial role in various aspects of 280 plant growth and development. HUA2 was initially identified as an enhancer of the 281 AGAMOUS (AG) allele ag-4 (Chen and Meyerowitz, 1999). Later, HUA2 was found to also 282 play a role as a repressor of flowering (Doyle et al., 2005; Wang et al., 2007). At the 283 molecular level, HUA2 has been suggested to function as a putative transcription factor but 284 has also been implicated in RNA processing (Cheng et al., 2003). HUA2 is expressed 285 throughout the whole plant growth period (Chen and Meyerowitz, 1999), indicating the 286 importance and widespread effects on plant growth. Here, our study showed that loss of 287 HUA2 can partially restore flowering and embryogenesis in tps1-2 GVG::TPS1. 288

It is interesting to note that in our EMS suppressor screen, we did not identify mutations in 289 any of the HUA2-like genes, HULK1, HULK2, and HUL3, present in A. thaliana (Jali et al., 290 2014). One possible explanation is that our genetic screen might not have been saturated or 291 292 that HUA2-like genes were missed due to the relatively low sequencing coverage. However, we believe this to be rather unlikely given that our approach has recovered multiple alleles in 293 HUA2 (this study) as well as two SnRK1 subunits (Zacharaki et al., 2022). Furthermore, 294 flowering time is unaffected in the *hua2-like* single mutants and *hulk2 hulk3* double mutants 295 have been shown to be late flowering (Jali et al., 2014). Thus, it seems unlikely that mutation 296 in any of the HUA2-like genes would suppress the non-flowering phenotype of tps1-2 297 GVG::TPS1. 298

HUA2 has been reported to exert its function in part by regulating the expression of 299 MADS-box transcription factors (Doyle et al., 2005), named after MINICHROMOSOME 300 MAINTENANCE 1 (MCM1) in yeast, AGAMOUS (AG) in Arabidopsis, DEFICIENS (DEF) 301 in Antirrhinum, and serum response factor (SRF) in humans. MADS-BOX domain 302 transcription factors contribute to all major aspects of the life of land plants, such as female 303 gametophyte, floral organ identity, seed development, and flowering time control (Portereiko 304 et al., 2006; Colombo et al., 2008; Koo et al., 2010; Lee et al., 2013; Posé et al., 2013). In this 305 context, it is interesting to note that our transcriptome and genetic analysis identified several 306 307 MADS-box transcription factors to be misregulated in tps1-2 GVG::TPS1. In particular, the well-known floral repressors FLOWERING LOCUS C (FLC) and MADS AFFECTING 308

FLOWERING5 (*MAF5*) were found to be induced in *tps1-2 GVG::TPS1* compared to Col-0
(Fig. 5A). Moreover, loss of *FLC* was sufficient to induce flowering in *tps1-2 GVG::TPS1*(Fig. 5B,C), suggesting that these floral repressors are partially responsible for the non-flowering phenotype of *tps1-2 GVG::TPS1*. Our transcriptome analyses further identified two MADS-box transcription factors, *SUPPRESSOR OF OVEREXPRESSION OF CONSTANS 1* (*SOC1*) and *AGAMOUS-LIKE 24* (*AGL24*), both known to promote flowering in Arabidopsis, to be upregulated in *hua2-4 tps1-2 GVG::TPS1*.

The molecular mechanism by which HUA2 regulates the expression of these MADS-box 316 317 flowering time regulators is currently unclear. However, since HUA2 localizes to the nucleus, it seems possible that HUA2 is directly involved in regulating the expression of these genes. 318 For example, HUA2 could (directly) promote the expression of FLC, which has previously 319 been shown to directly bind to and repress the expression of FT and SOC1 (Chen and 320 321 Meyerowitz, 1999; Doyle et al., 2005; Deng et al., 2011). In such a scenario, the increased expression of FT, SOC1, and AGL24 in hua2-4 tps1-2 GVG::TPS1 would be the result of 322 reduced expression of floral repressors such as FLC and MAF5. However, the regulation of 323 flowering is a very complex process full of intricate feedback loops and HUA2 might regulate 324 SOC1 and AGL24 directly, rather than indirectly. In this context, it is interesting to note that a 325 nonfunctional *hua2* allele may compensate for the loss of *FLC* in Ler accession (Lemus et al., 326 2023). Alternatively, HUA2 might affect the expression of these important flowering time 327 genes through interaction with RNA Pol-II via its CID domain, which is affected by the 328 hua2-13 alleles (R902C). Interestingly, polymorphisms resulting in amino acid substitutions 329 in natural accessions of A. thaliana have been reported for R902 and A983, but not for P455 330 (The 1001 Genomes Consortium, 2016). Even though the molecular mechanisms underlying 331 HUA2 function remain elusive, our results confirm HUA2 as a central regulator of flowering 332 time in Arabidopsis thaliana. 333

We have previously identified mutations in two subunits of *SNF1-Related Kinase 1* (*SnRK1*), *KIN10* and *SNF4*, that partially restore flowering and seed set in *tps1-2 GVG::TPS1* (Zacharaki et al., 2022). Identification of these suppressor mutations was in line with the role of SnRK1 as a downstream regulator of the T6P pathway and other stresses. Antagonizing

SnRK1 in the regulation of energy homeostasis in plants is target of rapamycin (TOR), the 338 activity of which is inhibited under energy-limiting conditions (Baena-González and Hanson, 339 340 2017; Belda-Palazón et al., 2022). In contrast to mutations in KIN10 and SNF4, mutations in HUA2 appear, at first glance, to be bypass mutations that induce flowering independently of 341 T6P signaling. However, and rather unexpectedly, we did observe that mutation of HUA2 342 attenuated the induction of the carbon starvation markers SEN5 and DIN6 in response to 343 extended night treatments (Fig. 6A, B), indicating that mutations in HUA2 might modulate 344 345 carbohydrate signaling more directly than anticipated. How exactly HUA2 modulates carbon responses in Arabidopsis remains to be established. It is well-known that T6P signaling 346 through SnRK1 affects processes such as carbon starvation response, germination, flowering, 347 and senescence in opposition to the TOR (target of rapamycin) pathway (Figueroa and Lunn, 348 2016; Baena-González and Lunn, 2020). The regulatory network controlling this central 349 metabolic hub is still not fully understood and new players are constantly added. For example, 350 it has recently been shown that class II TPS proteins are important negative regulators of 351 *SnRK1* (Van Leene et al., 2022). 352

Regarding a possible role of HUA2 in integrating carbon responses, it is worth noting that 353 flavonoid-related genes (GO:0009812) were downregulated in tps1-2 GVG::TPS1 in 354 response to dexamethasone application and the *hua2* mutant (Fig. 3B). This is interesting as 355 HUA2 is known to promote anthocyanin accumulation (Ilk et al., 2015), whereas SnRK1 has 356 been shown to repress sucrose-induced anthocyanin production (Li et al., 2014; Meng et al., 357 2018; Brouke et al., 2023). Thus, HUA2 might constitute an important hub in coordinating 358 metabolic responses. However, as expression of SnRK1 subunits is not affected in hua2-4 359 tps1-2 GVG::TPS1 when compared to tps1-2 GVG::TPS1 (Fig. S4), such a role would likely 360 be indirect. 361

Understanding the interplay between energy metabolism, in particular SnRK1, TOR, and T6P signaling, and plant growth and development is of utmost importance for developing plants capable of withstanding future challenges. The suppressor mutants generated in the *tps1-2 GVG::TPS1* background comprise an important resource in our hunt for additional factors that, like *HUA2*, link energy metabolism to plant development.

367

368 Material and methods

369 Plant materials and growth conditions

All T-DNA insertion mutants and transgenic lines used in this work are in the Col-0 370 background. The tps1-2 GVG::TPS1 line used in this work is referred to as ind-TPS1 #201 in 371 the original publication (Dijken et al., 2004). The hua2-4 (SALK_032281C) was obtained 372 from NASC and the presence of the T-DNA insertion was confirmed by PCR (Table S5). 373 374 ft-10 (GABI-Kat: 290E08) was provided by Dr. Yi Zhang, Southern University of Science and Technology, *flc-3* (Kim et al., 2006) by Dr. Liangyu Liu, Capital Normal University, and 375 soc1-2 (Lee et al., 2000) by Dr. Jie Luo, Chinese Academy of Sciences. tps1-2 GVG::TPS1 376 hua2-4 plants were generated by crossing and double homozygous mutants were identified by 377 378 phenotyping and genotyping of F2 individuals. Higher order mutants were obtained by crossing soc1-2, flc-3, and ft-10 mutants with the tps1-2 GVG::TPS1 hua2-4 double mutant 379 and homozygous triple mutants were identified in the F2 and F3 generation. All mutant 380 genotypes were confirmed by PCR, see Table S7 for details. Plants were planted onto nutrient 381 soil with normal water supply and grown under either long days (LD) with a photoperiod of 382 16 hours light at 22°C and 8 hours darkness at 20°C or in short days (SD) with a photoperiod 383 of 8 hours light at 22°C and 16 hours darkness at 20°C. Flowering time are presented as 384 average rosette leaf number, cauline leaf number, and total leaf number. 385

Genome sequencing and analysis

Young leaves were used for DNA extraction for sequencing using the NovaSeq 6000 387 Sequencing platform (Novogene). Adapters and low-quality sequences of raw reads were 388 trimmed using Trimmomatic (Bolger et al., 2014), and the clean reads were mapped to the 389 reference genome of Col-0 using BWA-MEM (v0.7.15) (Cingolani et al., 2012). SNP calling 390 performed using Genome Analysis Toolkit 4 (GATK4; 391 was https://gatk.broadinstitute.org/hc/en-us) with default parameters. Variants were annotated 392 using snpEff 4.3 (Li and Durbin, 2009) based on TAIR 10 annotation. Next, we identified the 393 protein-coding genes with multiple non-redundant mutations and found three mutant lines 394

harboring unique non-synonymous mutations in the *HUA2* gene. The method was inspired by
our previous study that multiple EMS-induced mutants with unique mutation sites in the
coding regions of *SnRK1* alpha subunit rescued the non-flowering phenotype of *tps1*(Zacharaki et al., 2022).

399 Gene expression analysis by RNA-seq

For RNA-seq analyses, plants were grown on soil for 3 weeks in LD conditions. Leaves from 400 21-day-old Arabidopsis thaliana were collected, immediately snap-frozen and stored at 401 -80 °C. Total RNA was extracted using RNAprep Pure Plant Plus Kit (Tiangen, China, 402 DP441). RNA integrity was assessed using the RNA Nano 6000 Assay Kit on the 403 Bioanalyzer 2100 system (Agilent Technologies, CA, USA). RNA-seq libraries were 404 generated with three independent biological replicates and sequenced on the Illumina 405 NovaSeq platform by Annoroad Gene Technology. The raw RNA-seq reads were quality 406 407 trimmed by Trimmomatic (v 0.11.9) (Bolger et al., 2014). The qualified reads were mapped to TAIR10 version genome guided by gene annotation model using HISAT2 (v2.1.0) (Kim et 408 al., 2015). The expression level for each gene was determined by StringTie (v1.3.4) (Pertea et 409 al., 2016). The differential expressed genes were identified by DESeq2 (Love et al., 2014). 410

411 **RNA isolation and RT-qPCR data analysis**

Total RNA was extracted from Arabidopsis seedlings using the RNA Isolation Kit (Tiangen, 412 China, DP441) according to the manufacturer's instructions. cDNA was synthesized from 3 413 µg total RNA in a 10 µl reaction volume using the RevertAid Premium First Strand cDNA 414 Synthesis Kit (Fermentas, Thermo Fisher Scientific, Rochester, NY). Quantitative real-time 415 PCR (qRT-PCR) was performed using TB Green[™] Premix Ex Taq[™] II (Takara, Dalian, 416 China). Relative gene expression was calculated using the $2-\Delta\Delta Ct$ method (Rao et al., 2013). 417 All analyses were repeated three times. The primer used for qRT-PCR are listed in 418 Supplemental Tables S5. 419

420 Accession numbers

- 421 Identifiers of key genes used in this study: TPS1 (At1g78580), HUA2 (AT5G23150), SOC1
- 422 (AT2G45660), FLC (AT5G10140), FT (AT1G65480). RNA-seq data generated in this study
- have been deposited with NCBI under the BioProject PRJNA1005425.

424 **Data availability**

425 The data and material that support the findings of this study are available from the 426 corresponding author upon reasonable request.

427 Author contributions

LZ and MS designed the experiments. LZ carried out the SNP detection and genetic analyses
with input from VZ and MS. LZ carried out the gene expression analyses. LP and MS wrote
the manuscript with contributions from all authors.

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654 **Figure legends**

Figure 1. EMS-induced mutations in HUA2 induce flowering in tps1-2 GVG::TPS1 655 background. A) Schematic drawing of HUA2 indicating the position and the amino acid 656 changes caused by the EMS-induced mutations hua2-11 (P455S), hua2-12 (R902C), and 657 hua2-13 (A983T). B) Phenotype of 9-week-old tps1-2 GVG::TPS1, hua2-11 tps1-2 658 GVG::TPS1, hua2-12 tps1-2 GVG::TPS1, hua2-13 tps1-2 GVG::TPS1 and wild-type Col-0 659 *plants* grown in LD at 23°C. C) Flowering time of genotypes is given as total leaf number 660 (rosette (grey); cauline leaves (white)) determined after bolting. Error bars represent the 661 standard deviation. ANOVA Tukey's multiple comparisons test was applied, and letters 662 represent the statistical differences among genotypes (P < 0.001). 663

664

Figure 2. A T-DNA insertion in *HUA2* partially rescues the flowering time phenotype of

tps1-2 GVG::TPS1. A) Schematic drawing of the *HUA2* locus indicating the position of the T-DNA insertion (SALK_032281C) in the 2nd intron in *hua2-4*. **B-C**) Phenotypic analysis (**B**) and flowering time(**C**) of 9-week-old wild-type Col-0, *tps1-2 GVG::TPS1*, *hua2-4 tps1-2 GVG::TPS1* and *hua2-4* plants grown in LD at 23°C. Flowering time was scored as total leaf number (rosette (grey) and cauline leaves (white)) after bolting. Error bars represent the standard deviation. ANOVA Tukey's multiple comparisons test was applied, and letters represent the statistical differences among genotypes (P < 0.001).

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Figure 3. Characterization of hua2-4 tps1-2 GVG::TPS1 transcriptome. A) 4-way Venn 674 diagram of genes that are differentially expressed in tps1-2 GVG::TPS1 in response to 675 dexamethasone treatment and/or differentially expressed in hua2-4 tps1-2 GVG::TPS1 when 676 compared to tps1-2 GVG::TPS1. B) GO analysis of 412 genes downregulated in tps1-2 677 GVG::TPS1 in response to dexamethasone treatment and in hua2-4 tps1-2 GVG::TPS1. C) 678 GO analysis of 243 genes upregulated in tps1-2 GVG::TPS1 in response to dexamethasone 679 treatment and in hua2-4 tps1-2 GVG::TPS1. D) Relative expression of AGL24 and SOC1 in 680 tps1-2 GVG::TPS1 (white), tps1-2 GVG::TPS1 treated with dexamethasone (black), and 681 hua2-4 tps1-2 GVG::TPS1 (grey). AGL24 and SOC1 are significantly differentially expressed. 682

Error bars indicate the standard deviation. ANOVA Tukey's multiple comparisons test was applied, and letters represent the statistical differences among genotypes (P < 0.001).

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Figure 4. Genetic interactions between tps1-2, hua2-4, and floral regulators SOC1, FT, 686 and FLC. A-B) Phenotypes (A) and flowering time (B) of Col-0, hua2-4, tps-2 GVG::TPS1, 687 and soc1-2 mutant combinations. D-E) Phenotypes (D) and flowering time (E) of Col-0, 688 hua2-4, tps-2 GVG::TPS1, and ft-10 mutant combinations. G-H) Phenotypes (G) and 689 flowering time (H) of Col-0, hua2-4, tps-2 GVG::TPS1, and flc-3 mutant combinations. 690 Flowering time (**B**, **E**, **H**) was scored as total leaf number (rosette (grey) and cauline leaves 691 (white)) after bolting. C, F, I) Relative expression of SOC1 (C), FT (F), and FLC (I) in 692 tps1-2 GVG::TPS1 and hua2-4 tps-2 GVG::TPS1. Gene expression was determined by 693 RT-qPCR at the end of the long day (ZT 16). Error bars represent the standard deviation. 694 ANOVA Tukey's multiple comparisons test was applied, and letters represent the statistical 695 differences among genotypes (P < 0.001). 696

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698 Figure 5. Loss of *FLC* rescues the non-flowering phenotype of *tps1-2 GVG::TPS1*. A) VST expression estimates for MADS-box floral repressors in 18-day-old plants. RNA-seq 699 expression data retrieved from Zacharaki et al., 2022. Columns indicate mean VST 700 expression estimates as implemented in DEseq2 calculated from three individual biological 701 replicates per genotype. Circles indicate expression estimates for individual biological 702 replicates. Asterisks indicate differential gene expression with a statistical significance of 703 Padj < 0.01. B-C) Phenotypes (B) and total leaf number (C) of Col-0, tps1-2 GVG::TPS1, 704 flc-3, and flc-3 tps1-2 GVG::TPS1 double mutant. Flowering time was scored as total leaf 705 number (rosette (grey) and cauline leaves (white)) after bolting. Error bars represent the 706 standard deviation. ANOVA Tukey's multiple comparisons test was applied, and letters 707 represent the statistical differences among genotypes (P < 0.001). 708

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Figure 6. Expression of SnRK1 target genes SEN5 and DIN6 in hua2-4 and hua2-4
tps1-2 GVG::TPS1 double mutant. A-B) Induction of SEN5 (A) and DIN6 (B) in response
to extended night is attenuated in 14-day-old of hua2-4 single mutant and three independent

- 713 lines of the *hua2-4 tps1-2 GVG::TPS1* double mutant. Plants were grown for 14 days in LD
- (grey) before being exposed to a single extended night (12h additional darkness; black). LD,
- ⁷¹⁵ long days. Error bars represent the standard deviation. ANOVA Tukey's multiple comparisons
- test was applied, and letters represent the statistical differences among genotypes (P < 0.001).

717 Supplemental Material

718	Supplemental Figure S1	Relative expression of HUA2 in tps1-2 GVG::TPS1 treated
719		with dexamethasone or untreated.
720	Supplemental Figure S2	Relative expression of important floral regulators in tps1-2
721		GVG::TPS1, tps1-2 GVG::TPS1 treated with dexamethasone,
722		and hua2-4 tps1-2 GVG::TPS1.
723	Supplemental Figure S3	VST expression estimates for HUA2 in 18-day-old plants.
724	Supplemental Figure S4	Relative expression of SnRK1 subunits in tps1-2
725		GVG::TPS1, tps1-2 GVG::TPS1 treated with dexamethasone,
726		and hua2-4 tps1-2 GVG::TPS1.
727	Supplemental Table S1	Number of SNPs identified in individual suppressor mutants.
728	Supplemental Table S2	Number of SNPs identified in EMS suppressor lines carrying
729		mutations in HUA2.
730	Supplemental Table S3	EMS suppressor lines bearing non-synonymous mutations in
731		HUA2.
732	Supplemental Table S4	GO analysis of 412 genes downregulated in tps1-2
733		GVG::TPS1 in response to dexamethasone application and in
734		hua2-4.
735	Supplemental Table S5	GO analysis of 243 genes induced in tps1-2 GVG::TPS1 in
736		response to dexamethasone application and in hua2-4.
737	Supplemental Table S6	Expression of flowering time genes in hua2-4 tps1-2
738		GVG::TPS1 and tps1-2 GVG::TPS1.
739	Supplemental Table S7	List of oligonucleotides used in this study.

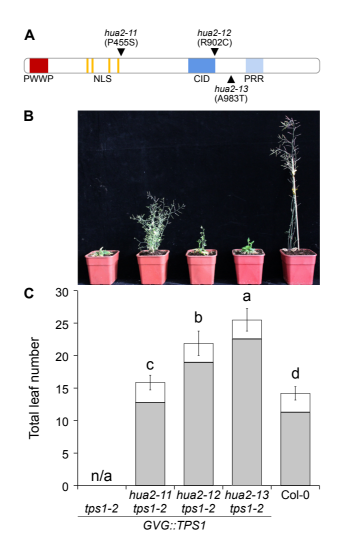


Figure 1. EMS-induced mutations in *HUA2* induce flowering in *tps1-2 GVG::TPS1* background. A) Schematic drawing of HUA2 indicating the position and the amino acid changes caused by the EMS-induced mutations *hua2-11* (P455S), *hua2-12* (R902C), and *hua2-13* (A983T). B) Phenotype of 9-week-old *tps1-2 GVG::TPS1*, *hua2-11 tps1-2 GVG::TPS1*, *hua2-12 tps1-2 GVG::TPS1*, *hua2-13 tps1-2 GVG::TPS1* and wild-type Col-0 *plants* grown in LD at 23°C. C) Flowering time of genotypes is given as total leaf number (rosette (grey); cauline leaves (white)) determined after bolting. Error bars represent the standard deviation. ANOVA Tukey's multiple comparisons test was applied, and letters represent the statistical differences among genotypes (P < 0.001).

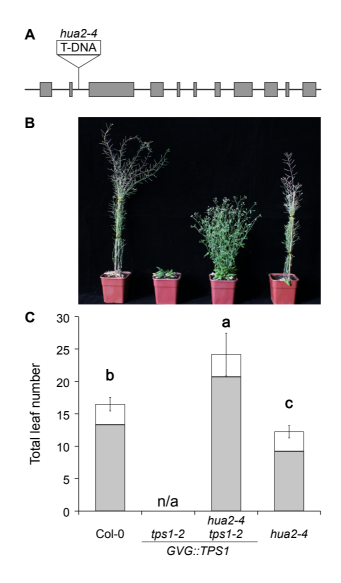


Figure 2. A T-DNA insertion in *HUA2* partially rescues the flowering time phenotype of *tps1-2 GVG::TPS1*. A) Schematic drawing of the *HUA2* locus indicating the position of the T-DNA insertion (SALK_032281C) in the 2nd intron in *hua2-4*. **B-C**) Phenotypic analysis (**B**) and flowering time(**C**) of 9-week-old wild-type Col-0, *tps1-2 GVG::TPS1*, *hua2-4 tps1-2 GVG::TPS1* and *hua2-4* plants grown in LD at 23°C. Flowering time was scored as total leaf number (rosette (grey) and cauline leaves (white)) after bolting. Error bars represent the standard deviation. ANOVA Tukey's multiple comparisons test was applied, and letters represent the statistical differences among genotypes (P < 0.001).

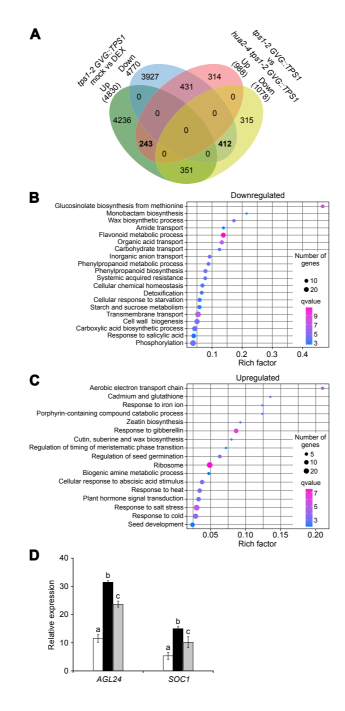


Figure 3. Characterization of *hua2-4 tps1-2 GVG::TPS1* transcriptome. A) 4-way Venn diagram of genes that are differentially expressed in *tps1-2 GVG::TPS1* in response to dexamethasone treatment and/or differentially expressed in *hua2-4 tps1-2 GVG::TPS1* when compared to *tps1-2 GVG::TPS1*. B) GO analysis of 412 genes downregulated in *tps1-2 GVG::TPS1* in response to dexamethasone treatment and in *hua2-4 tps1-2 GVG::TPS1*. C) GO analysis of 243 genes upregulated in *tps1-2 GVG::TPS1* in response to dexamethasone treatment and in *hua2-4 tps1-2 GVG::TPS1*. C) GO analysis of 243 genes upregulated in *tps1-2 GVG::TPS1* in response to dexamethasone treatment and in *hua2-4 tps1-2 GVG::TPS1*. D) Relative expression of *AGL24* and *SOC1* in *tps1-2 GVG::TPS1* (white), *tps1-2 GVG::TPS1* treated with dexamethasone (black), and *hua2-4 tps1-2 GVG::TPS1* (grey). *AGL24* and *SOC1* are significantly differentially expressed. Error bars indicate the standard deviation. ANOVA Tukey's multiple comparisons test was applied, and letters represent the statistical differences among genotypes (P < 0.001).

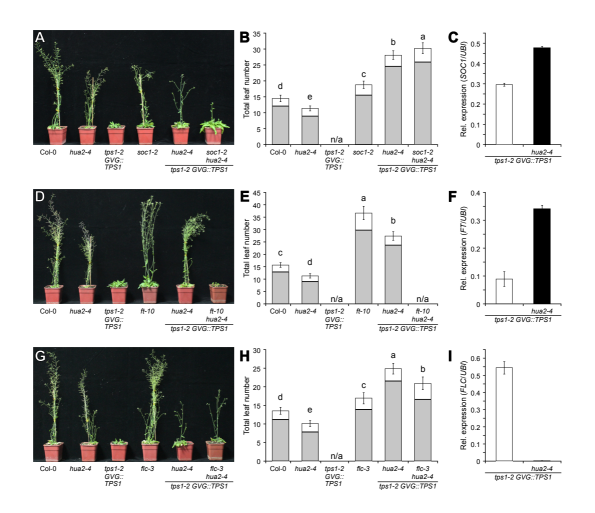


Figure 4. Genetic interactions between *tps1-2*, *hua2-4*, and floral regulators *SOC1*, *FT*, and *FLC*. A-B) Phenotypes (A) and flowering time (B) of Col-0, *hua2-4*, *tps-2 GVG::TPS1*, and *soc1-2* mutant combinations. D-E) Phenotypes (D) and flowering time (E) of Col-0, *hua2-4*, *tps-2 GVG::TPS1*, and *ft-10* mutant combinations. G-H) Phenotypes (G) and flowering time (H) of Col-0, *hua2-4*, *tps-2 GVG::TPS1*, and *flc-3* mutant combinations. Flowering time (B, E, H) was scored as total leaf number (rosette (grey) and cauline leaves (white)) after bolting. C, F, I) Relative expression of *SOC1* (C), *FT* (F), and *FLC* (I) in *tps1-2 GVG::TPS1* and *hua2-4 tps-2 GVG::TPS1*. Gene expression was determined by RT-qPCR at the end of the long day (ZT 16). Error bars represent the standard deviation. ANOVA Tukey's multiple comparisons test was applied, and letters represent the statistical differences among genotypes (P < 0.001).

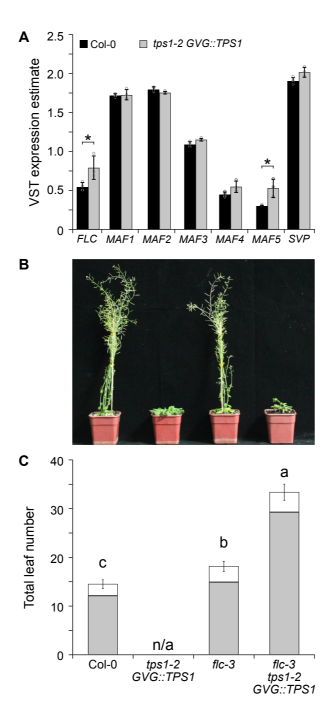


Figure 5. Loss of *FLC* rescues the non-flowering phenotype of *tps1-2 GVG::TPS1*. A) VST expression estimates for MADS-box floral repressors in 18-day-old plants. RNA-seq expression data retrieved from Zacharaki et al., 2022. Columns indicate mean VST expression estimates as implemented in DEseq2 calculated from three individual biological replicates per genotype. Circles indicate expression estimates for individual biological replicates. Asterisks indicate differential gene expression with a statistical significance of *Padj* < 0.01. **B-C**) Phenotypes (**B**) and total leaf number (**C**) of Col-0, *tps1-2 GVG::TPS1, flc-3*, and *flc-3 tps1-2 GVG::TPS1 double mutant*. Flowering time was scored as total leaf number (rosette (grey) and cauline leaves (white)) after bolting. Error bars represent the standard deviation. ANOVA Tukey's multiple comparisons test was applied, and letters represent the statistical differences among genotypes (P < 0.001).

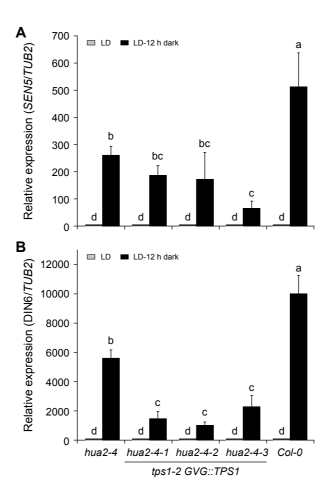


Figure 6. Expression of SnRK1 target genes SEN5 and DIN6 in hua2-4 and hua2-4 tps1-2 GVG::TPS1 double mutant. A-B) Induction of SEN5 (A) and DIN6 (B) in response to extended night is attenuated in 14-day-old of hua2-4 single mutant and three independent lines of the hua2-4 tps1-2 GVG::TPS1 double mutant. Plants were grown for 14 days in LD (grey) before being exposed to a single extended night (12h additional darkness; black). LD, long days. Error bars represent the standard deviation. ANOVA Tukey's multiple comparisons test was applied, and letters represent the statistical differences among genotypes (P < 0.001).

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