

1 **Short Title:** Monocot ELF3 functions in dicots

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17 **Cross-species complementation reveals conserved functions for EARLY FLOWERING 3**
18 **between monocots and dicots**

19

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28

29 **One Sentence Summary:**

30 Orthologs of a key circadian clock component ELF3 from grasses functionally complement the
31 Arabidopsis counterpart at the molecular and physiological level, in spite of high sequence
32 divergence.

33

34 **Footnotes:**

35

36 H.H., M.A.G., T.C.M., and D.A.N. conceived the project; T.C.M., B.S.E. and D.A.N. supervised
37 the experiments; H.H., M.A.G., S.E.H., S.A., C.L., E.L.G., J.G., M.J.N., R.K.B. and D.A.N.
38 performed the experiments; H.H., M.A.G., S.E.H., S.A., C.L., M.J.N. and D.A.N. analyzed the
39 data; H.H., M.A.G., and D.A.N. wrote the manuscript. All authors edited the manuscript.

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48

49 **ABSTRACT**

50

51 Plant responses to the environment are shaped by external stimuli and internal signaling
52 pathways. In both the model plant *Arabidopsis thaliana* and crop species, circadian clock factors
53 have been identified as critical for growth, flowering and circadian rhythms. Outside of *A.*
54 *thaliana*, however, little is known about the molecular function of clock genes. Therefore, we
55 sought to compare the function of *Brachypodium distachyon* and *Setaria viridis* orthologs of
56 *EARLY FLOWERING3*, a key clock gene in *A. thaliana*. To identify both cycling genes and
57 putative ELF3 functional orthologs in *S. viridis*, a circadian RNA-seq dataset and online query
58 tool (Diel Explorer) was generated as a community resource to explore expression profiles of
59 *Setaria* genes under constant conditions after photo- or thermo-entrainment. The function of
60 *ELF3* orthologs from *A. thaliana*, *B. distachyon*, and *S. viridis* were tested for complementation
61 of an *elf3* mutation in *A. thaliana*. Despite comparably low sequence identity versus AtELF3
62 (less than 37%), both monocot orthologs were capable of rescuing hypocotyl elongation,
63 flowering time and arrhythmic clock phenotypes. Molecular analysis using affinity purification
64 and mass spectrometry to compare physical interactions also found that BdELF3 and SvELF3
65 could be integrated into similar complexes and networks as AtELF3, including forming a
66 composite evening complex. Thus, we find that, despite 180 million years of separation, BdELF3
67 and SvELF3 can functionally complement loss of *ELF3* at the molecular and physiological level.

68

69 INTRODUCTION

70

71 Plants have developed sophisticated signaling networks to survive and thrive in diverse
72 environments. Many plant responses are shaped, in part, by an internal timing mechanism known
73 as the circadian clock, which allows for the coordination and anticipation of daily and seasonal
74 variation in the environment (Greenham and McClung, 2015). Circadian clocks, which are
75 endogenous oscillators with a period of approximately 24 hours, are critical for regulating the
76 timing of physiology, development, and metabolism in all domains of life (Bell-Pedersen et al.,
77 2005; Doherty and Kay, 2010; Edgar et al., 2012; Harmer, 2009; Wijnen and Young, 2006). In
78 plants and blue-green algae, circadian clocks provide an experimentally observable adaptive
79 advantage by synchronizing internal physiology with external environmental cues (Dodd et al.,
80 2005; Ouyang et al., 1998; Woelfle et al., 2004). Currently, circadian oscillators are best
81 understood in the reference plant *Arabidopsis thaliana*, in which dozens of clock or clock-
82 associated components have been identified using genetic screens and non-invasive, luciferase-
83 based oscillating reporters (Hsu and Harmer, 2014; Nagel and Kay, 2012). These morning-,
84 afternoon-, and evening-phased clock oscillators form multiple interconnected transcription-
85 translation feedback loops and compose a complex network (Hsu and Harmer, 2014; Pokhilko et
86 al., 2012). The *A. thaliana* circadian clock regulates a significant portion of physiology,
87 including photosynthesis, growth, disease resistance, starch metabolism, and phytohormone
88 pathways (Covington et al., 2008; Graf et al., 2010; Harmer et al., 2000; Michael et al., 2008;
89 Wang et al., 2011b), with up to 30% of gene expression under circadian control (Covington et
90 al., 2008; Michael et al., 2008).

91 Within the *Arabidopsis* clock network, a tripartite protein complex called the evening complex
92 (EC) is an essential component of the evening transcription loop (Huang and Nusinow, 2016b).
93 The EC consists of three distinct proteins, EARLY FLOWERING 3 (ELF3), EARLY
94 FLOWERING 4 (ELF4) and LUX ARRHYTHMO (LUX, also known as PHYTOCLOCK1),
95 with transcript and protein levels peaking in the evening (Doyle et al., 2002; Hazen et al., 2005;
96 Hicks et al., 2001; Nusinow et al., 2011; Onai and Ishiura, 2005). The EC plays a critical role in
97 maintaining circadian rhythms, by repressing expression of key clock genes (Dixon et al., 2011;
98 Helfer et al., 2011; Herrero et al., 2012; Kolmos et al., 2011; Mizuno et al., 2014). Loss-of-

99 function mutation of any EC component in *A. thaliana* results in arrhythmicity of the circadian
100 clock and causes excessive cellular elongation and early flowering regardless of environmental
101 photoperiod (Doyle et al., 2002; Hazen et al., 2005; Hicks et al., 2001; Khanna et al., 2003; Kim
102 et al., 2005; Nozue et al., 2007; Nusinow et al., 2011; Onai and Ishiura, 2005).

103
104 In *A. thaliana*, ELF3 directly interacts with ELF4 and LUX, functioning as a scaffold to bring
105 ELF4 and LUX together (Herrero et al., 2012; Nusinow et al., 2011). Additional protein-protein
106 interaction studies and tandem affinity purification coupled with mass spectrometry (AP-MS)
107 have identified many ELF3-associating proteins and established ELF3 as a hub of a complex
108 protein-protein interaction network, which consists of key components from the circadian clock
109 pathway and light signaling pathways (Huang et al., 2016a; Huang and Nusinow, 2016b; Liu et
110 al., 2001; Yu et al., 2008). In this network, ELF3 directly interacts with the major red light
111 photoreceptor phytochrome B (phyB), and CONSTITUTIVE PHOTOMORPHOGENIC 1
112 (COP1), which is an E3 ubiquitin ligase required for proper regulation of photomorphogenesis
113 and also interacts with phyB (Liu et al., 2001; Yu et al., 2008). The physical interaction among
114 ELF3, phyB, and COP1, together with recruitment of direct interacting proteins to the network,
115 provides biochemical evidence for cross-talk between circadian clock and light signaling
116 pathways (Huang and Nusinow, 2016b). Although much work does translate from *A. thaliana* to
117 other plant species, interaction between ELF3 and other proteins have yet to be tested in species
118 outside *A. thaliana*. Whether evening complex-like protein assemblages or a similar ELF3-
119 containing protein-protein interaction network exists in species outside *A. thaliana* is an
120 interesting question to ask.

121
122 Identification and characterization of clock genes in diverse plant species has revealed that many
123 clock components are broadly conserved (Filichkin et al., 2011; Khan et al., 2010; Lou et al.,
124 2012; Song et al., 2010). Furthermore, comparative genomics analysis has found that circadian
125 clock components are selectively retained after genome duplication events, suggestive of the
126 importance of their role in maintaining fitness (Lou et al., 2012). Recently, mutant alleles of
127 *ELF3* were identified associated with the selection of favorable photoperiodism phenotypes in
128 several crops, such as pea, rice, soybean and barley (Faure et al., 2012; Lu et al., 2017;

129 Matsubara et al., 2012; Saito et al., 2012; Weller et al., 2012; Zakhrebekova et al., 2012). These
130 findings are consistent with the reported functions of *A. thaliana* ELF3 in regulating the
131 photoperiodic control of growth and flowering (Hicks et al., 2001; Huang and Nusinow, 2016b;
132 Nozue et al., 2007; Nusinow et al., 2011). However, opposed to the early flowering phenotype
133 caused by *elf3* mutants in *A. thaliana*, pea, and barley (Faure et al., 2012; Hicks et al., 2001;
134 Weller et al., 2012), loss of function mutation of the rice or soybean *ELF3* ortholog results in
135 delayed flowering (Lu et al., 2017; Saito et al., 2012), suggesting ELF3-mediated regulation of
136 flowering varies in different plant species. The molecular mechanisms underlying this difference
137 have not been thoroughly elucidated.

138
139 *Brachypodium distachyon* is a C3 model grass closely related to wheat, barley, oats, and rice.
140 *Setaria viridis*, is a C4 model grass closely related to maize, sorghum, sugarcane, and other
141 bioenergy grasses. Both grasses are small, transformable, rapid-cycling plants with recently
142 sequenced genomes, making them ideal model monocots for comparative analysis with
143 *Arabidopsis* (Bennetzen et al., 2012; Brutnell et al., 2010). Computational analysis of *B.*
144 *distachyon* has identified putative circadian clock orthologs (Higgins et al., 2010), including
145 *BdELF3*. However, no such comparative analysis has been done systematically in *S. viridis* to
146 identify putative orthologs of circadian clock genes. Therefore, we generated a RNA-seq time-
147 course dataset to analyze the circadian transcriptome of *S. viridis* after either photo- or thermo-
148 entrainment and developed an online gene-expression query tool (Diel Explorer) for the
149 community. We found that the magnitude of circadian regulated genes in *S. viridis* is similar to
150 other monocots after photo-entrainment, but much less after thermal entrainment. We further
151 analyzed the functional conservation of SvELF3, together with previously reported BdELF3, by
152 introducing both ELF3 orthologs into *A. thaliana elf3* mutant for physiological and biochemical
153 characterization. We found that *B. distachyon* and *S. viridis* ELF3 can complement the hypocotyl
154 elongation, flowering time and circadian arrhythmia phenotypes caused by the *elf3* mutation in
155 *A. thaliana*. Furthermore, AP-MS analyses found that *B. distachyon* and *S. viridis* ELF3 were
156 integrated into a similar protein-protein interaction network *in vivo* as their *A. thaliana*
157 counterpart. Our data collectively demonstrated the functional conservation of ELF3 among *A.*

158 *thaliana*, *B. distachyon* and *S. Viridis* is likely due to the association with same protein partners,
159 providing insights of how ELF3 orthologs potentially function in grasses.

160

161 **RESULTS**

162

163 **Identifying and cloning ELF3 orthologs from *B. distachyon* and *S. viridis***

164

165 ELF3 is a plant-specific nuclear protein with conserved roles in flowering and the circadian
166 clock in multiple plant species (Faure et al., 2012; Herrero et al., 2012; Liu et al., 2001; Lu et al.,
167 2017; Matsubara et al., 2012; Saito et al., 2012; Weller et al., 2012; Zakhrabekova et al., 2012).
168 To identify ELF3 orthologs in monocots, we used the protein sequence of *Arabidopsis thaliana*
169 ELF3 (AtELF3) to search the proteomes of two model monocots *Brachypodium distachyon* and
170 *Setaria viridis* using BLAST (Altschul et al., 1990). Among the top hits, we identified a
171 previously reported ELF3 homolog in *B. distachyon* (*Bradi2g14290.1*, *BdELF3*) (Calixto et al.,
172 2015; Higgins et al., 2010) and two putative *ELF3* homologous genes *Sevir.5G206400.1*
173 (referred as *SvELF3a*) and *Sevir.3G123200.1* (referred as *SvELF3b*) in *S. viridis*. We used
174 Clustal Omega (<http://www.ebi.ac.uk/Tools/msa/clustalo/>) to conduct multiple sequence
175 alignments of comparing protein sequences of ELF3 orthologs with that of AtELF3 (Sievers et
176 al., 2011). *BdELF3*, *SvELF3a* and *SvELF3b* encode proteins with similar identity compared to
177 AtELF3 (34.7–36.8%) (**Supplemental Figure S1**). When compared to *BdELF3*, *SvELF3b* was
178 74.3% identical while *SvELF3a* was 57.4% identical (**Supplemental Figure S1**). Therefore, to
179 maximize the diversity of ELF3 sequences used in this study, we cloned full length cDNAs
180 encoding *BdELF3* and *SvELF3a*.

181

182 **Diel Explorer of *S. viridis* circadian data**

183

184 In *Arabidopsis*, ELF3 cycles under diel and circadian conditions (constant condition after
185 entrainment) with a peak phase in the evening (Covington et al., 2001; Hicks et al., 2001;
186 Nusinow et al., 2011). We queried an available diurnal time course expression dataset for *B.*
187 *distachyon* from the DIURNAL website, and found that *BdELF3* expression cycles under diel

188 conditions (LDHH, 12 h light / 12 h dark cycles with constant temperature), but not under
189 circadian conditions in available data (LDHC-F or LDHH-F, **Supplemental Figure S2**)
190 (Filichkin et al., 2011; Mockler et al., 2007). Also different from *AtELF3*, transcript levels of
191 *BdELF3* accumulate at dawn rather than peak in the evening (**Supplemental Figure S2**) when
192 grown under diel conditions, suggesting different regulations on *ELF3* expression between
193 monocot and dicot plants. Neither diel nor circadian expression data for *S. viridis* was available.
194 Therefore, we generated RNA-seq time-course data to examine *SvELF3* expression as well as the
195 circadian expression of other clock orthologs after both photocycle and thermocycle entrainment.
196 In addition, we developed the Diel Explorer tool
197 (<http://shiny.bioinformatics.danforthcenter.org/diel-explorer/>) to query and visualize *S. viridis*
198 circadian-regulated gene expression (**Supplemental Figure S3**). 48,594 *S. viridis* transcripts are
199 represented in the two datasets entrained under either photocycles (LDHH-F) or thermocycles
200 (LLHC-F). With Diel Explorer users can manually enter a list of transcript identifiers, gene
201 ontology (GO) terms, or gene orthologs, plot gene expression, and download data. Alternatively,
202 users can upload files of transcript identifiers or gene orthologs, and/or filter the datasets by
203 entrainment, phase, or significance cut-offs. Data and graphs can be downloaded directly using
204 Diel Explorer. The tool serves as a community resource that can be expanded to include other
205 circadian or diurnal data in the future. The underlying code is available on Github
206 (<https://github.com/danforthcenter/diel-explorer>).

207
208 Under photoperiod entrainment (LDHH-F), 5,585 of the 48,594 *S. viridis* transcripts are
209 circadian regulated (Bonferroni-adjusted P-Value < 0.001). This proportion of photoperiod-
210 entrained circadian genes (~11.5%) is similar to maize (10.8%), rice (12.6%), and poplar
211 (11.2%) data sets, but much smaller than the approximately 30% reported for *A. thaliana*
212 (Covington et al., 2008; Filichkin et al., 2011; Khan et al., 2010). Under thermocycle
213 entrainment (LLHC-F), 582 of the 48,594 *S. viridis* transcripts are circadian regulated.
214 Therefore, only ~1.2% of *S. viridis* transcripts are circadian cycling under thermocycle
215 entrainment. The ~10-fold reduction in circadian cycling genes between photocycle and
216 thermocycle entrainment (**Supplemental Figure S4**) is interesting considering that there was
217 less than 1% difference in the number of genes with a circadian period between photocycle and

218 thermocycle entrainment in C3 monocot rice (*Oryza japonica*) (Filichkin et al., 2011). The
219 reduction in cycling genes between the two entrainment conditions in *S. viridis* compared to *O.*
220 *japonica* is an indication that circadian regulation could vary greatly among monocots. Also, the
221 difference in number of cycling genes between monocots and dicots may represent a significant
222 reduction of the role of the circadian clock between these lineages.

223
224 In addition to the overall reduction in circadian genes, the phase with the most number of cycling
225 genes was ZT18 after light entrainment (LDHH-F; **Supplemental Figure S4**), but ZT12 with
226 temperature entrainment (LLHC-F; **Supplemental Figure S4**), which is consistent with previous
227 studies that have found significant differences in temperature and light entrainment of the
228 circadian clock (Boikoglou et al., 2011; Michael et al., 2008; Michael et al., 2003). There are 269
229 genes that are considered circadian-regulated and are cycling under both LDHH-F and LLHC-F
230 conditions (Bonferroni Adjusted P-Value < 0.001). The list of 269 genes that overlap between
231 photocycle and thermocycle entrainment includes best matches for Arabidopsis core clock
232 components *TIMING OF CAB EXPRESSION 1* (*TOC1*, *AT5G61380.1*; *Sevir.1G241000.1*),
233 *LATE ELONGATED HYPOCOTYL* (*LHY*, *AT1G01060*; *Sevir.6G053100.1*), and *CCA1*-like gene
234 *REVEILLE1* (*RVE1*, *AT5G17300.1*; *Sevir.1G280700.1*). However, putative *S. viridis* orthologs of
235 *TOC1*, *LHY*, and *RVE1* all have different circadian phases under LDHH entrainment compared
236 to LLHC entrainment (**Figure 1**). In fact, the majority (233/269) of overlapping circadian genes
237 in *S. viridis* have a distinct circadian phase under thermocycle compared to photocycle entrainment
238 (**Supplemental Figure S4**). We also found that putative orthologs of *PSEUDO-RESPONSE*
239 *REGULATOR 7* (*PRR7*, *AT5G02810*; *Sevir.2G456400.1*; related to *OsPRR73* (Murakami et
240 al., 2003)) and *LUX ARRHYTHMO* (*LUX*, *AT3G46640*; *Sevir.5G474200.1*) cycle significantly
241 under LDHH-F but not LLHC-F conditions (**Figure 1**). Neither *SvELF3a* nor *SvELF3b* cycle
242 under circadian conditions after photo- or thermo-entrainment (**Figure 1**), similar to *ELF3*
243 orthologs in *B. distachyon* (**Supplemental Figure S2**) (Mockler et al., 2007) and *O. sativa*
244 (Filichkin et al., 2011). This is different from *AtELF3*, which continues to cycle under constant
245 condition after either photo- or thermos-entrainment (**Supplemental Figure S5**) (Mockler et al.,
246 2007). The difference in expression of these putative orthologs between *A. thaliana* and

247 monocots *S. viridis*, *B. distachyon*, and *O. sativa*, suggest that the architecture of the circadian
248 clock may have significant differences in response to environmental cues in these two species.

249

250 **BdELF3 and SvELF3 rescue growth and flowering defects in *Arabidopsis elf3* mutant**

251

252 Although the circadian expression pattern of *B. distachyon ELF3* and *S. viridis ELF3* is different
253 from that of *A. thaliana ELF3*, it is still possible that the ELF3 orthologs have conserved
254 biological functions. To test this, we sought to determine if BdELF3 or SvELF3a could
255 complement the major phenotypic defects of the *elf3* mutant in *A. thaliana*, namely hypocotyl
256 elongation, time to flowering, or circadian rhythmicity. To this end, we constitutively expressed
257 *BdELF3*, *SvELF3a* (hereafter referred as *SvELF3*) and *AtELF3* cDNAs by the *35S Cauliflower*
258 *mosaic virus* promoter in the *A. thaliana elf3-2* mutant expressing a *LUCIFERASE* reporter
259 driven by the promoter of *CIRCADIAN CLOCK ASSOCIATED 1 (CCA1) (elf3-2 [CCA1:LUC])*
260 (Pruneda-Paz et al., 2009). All three ELF3 coding sequences were fused to a C-terminal His₆-
261 3xFlag affinity tag (HFC), which enables detection by western blotting and identification of
262 protein-protein interaction by affinity purification and mass spectrometry (AP-MS) (Huang et al.,
263 2016a). After transforming these constructs, we identified and selected two biologically
264 independent transgenic lines with a single insertion of each At/Bd/SvELF3-HFC construct.
265 Western blot analysis using FLAG antibodies detected the expression of all ELF3-HFC fusion
266 proteins (**Supplemental Figure S6**).

267

268 Next, we asked if expressing At/Bd/SvELF3-HFC fusion proteins could rescue the mutation
269 defects caused by *elf3-2*. When plants are grown under light/dark cycles (12 hour light: 12 hour
270 dark), *elf3-2* mutant plants elongate their hypocotyls much more than wild type plants (4.75±0.48
271 mm vs. 1.95±0.27 mm, respectively. ± = standard deviation) (**Figure 2**). The long hypocotyl
272 defect in *elf3-2* was effectively suppressed by expressing either *AtELF3*, or *ELF3* orthologs
273 (*BdELF3* or *SvELF3a*) (**Figure 2**). These data show that the monocot *ELF3* orthologs function
274 similarly to *A. thaliana ELF3* in the regulation of hypocotyl elongation in seedlings.

275

276 In addition to regulating phenotypes in seedlings, ELF3 also functions in adult plants to suppress
277 the floral transition. Loss-of-function in Arabidopsis *ELF3* results in an early flowering
278 phenotype regardless of day-length (Hicks et al., 2001; Liu et al., 2001; Zagotta et al., 1992). To
279 determine how monocot ELF3 orthologs compared to *A. thaliana* ELF3 in flowering time
280 regulation, we compared flowering responses under long day conditions among wild type, *elf3-2*,
281 and *elf3-2* transgenic lines expressing *AtELF3*, *BdELF3*, or *SvELF3* (At/Bd/SvELF3-HFC).
282 Constitutive over-expression of *AtELF3* led to a delay in flowering in long days (**Figure 3**) as
283 previously observed (Liu et al., 2001). Similarly, constitutive expression of *BdELF3* or *SvELF3*
284 caused plants to flower significantly later than the *elf3* mutants. These data show that all *ELF3*
285 orthologs can function to repress the rapid transition to flowering of the *elf3* mutation when
286 constitutively expressed in adult plants.

287

288 **BdELF3 and SvELF3 restore the circadian rhythmicity in Arabidopsis *elf3* mutant**

289

290 ELF3 is a key component of the *A. thaliana* circadian clock and is critical for maintaining the
291 periodicity and amplitude of rhythms as shown using the *CCA1* promoter driven luciferase
292 reporter (*CCA:LUC*) (Covington et al., 2001; Hicks et al., 1996; Nusinow et al., 2011). To
293 determine if *BdELF3* or *SvELF3a* could rescue the arrhythmic phenotype of the *elf3* mutation,
294 we analyzed the rhythms of the *CCA1::LUC* reporter under constant light conditions after diel
295 entrainment (12 hours light: 12 hours dark at constant 22 °C). Relative amplitude error (RAE)
296 analysis found that 100% of wild type and all three *elf3-2* transgenic lines expressing *AtELF3*,
297 *BdELF3*, and *SvELF3* were rhythmic, while only 50% of the *elf3-2* lines had rhythms (RAE <
298 0.5) (**Supplemental Figure S7**). Comparison of average period length found that the *AtELF3*
299 and *SvELF3* expressing lines rescued the period and amplitude defects in the *elf3* mutant (**Figure**
300 **4A, 4C and 4D**). While the *BdELF3* lines rescued the amplitude defect (**Figure 4B**), their period
301 was significantly divergent from wild type (compare 23.21 ± 0.59 hours for wild type to *BdELF3*
302 #2= 26.03 ± 1.33 hours, *BdELF3* #3= 26.86 ± 0.81 hours, *elf3-2*= 26.54 ± 5.02 hours, \pm =
303 standard deviation, **Figure 4D**). In summary, these data show that expression of any of the ELF3
304 orthologs is sufficient to recover the amplitude and restore circadian rhythms of the *CCA1::LUC*
305 reporter.

306

307 **BdELF3 and SvELF3 are integrated into a similar protein-protein interaction network in**

308 *A. thaliana*

309

310 Despite relatively low sequence conservation at the protein level, the *ELF3* orthologs can
311 complement a wide array of *elf3* phenotypes (**Figures 2 to 4**). As ELF3 functions within the
312 evening complex (EC) in Arabidopsis, which also contains the transcription factor LUX and the
313 DUF-1313 domain containing protein ELF4 (Herrero et al., 2012; Nusinow et al., 2011), we
314 reasoned that the monocot *ELF3* orthologs may also be able to bind to these proteins when
315 expressed in *A. thaliana*. To determine if a composite EC could be formed, we tested if BdELF3
316 or SvELF3a could directly interact with AtLUX or AtELF4 in a yeast two-hybrid assay. Similar
317 to AtELF3 (Nusinow et al., 2011), both BdELF3 and SvELF3a directly interact with both
318 AtELF4 and the C-terminal portion of AtLUX (**Figure 5**). We cannot conclude that whether
319 monocot ELF3 orthologs are also able to interact with the N-terminal AtLUX, since this
320 fragment auto-activated the reporter gene in the yeast two-hybrid assay (**Figure 5**).

321

322 ELF3 functions not only as the scaffold of the EC, but also as a hub protein in a protein-protein
323 interaction network containing multiple key regulators in both the circadian clock and light
324 signaling pathways (Huang et al., 2016a; Huang and Nusinow, 2016b). We hypothesize that
325 BdELF3 and SvELF3 could rescue many of the defects of the *elf3* mutant because both monocot
326 versions were integrated into the same protein-protein interaction network. To test this
327 hypothesis, we used affinity purification and mass spectrometry (AP-MS) to identify the proteins
328 that co-precipitate with monocot ELF3s when expressed in *A. thaliana*. AP-MS on two
329 biological replicates for each sample with the above-mentioned independent insertion lines were
330 included for each ELF3 ortholog. For comparison, the same AP-MS experiment was done with
331 one of the 35S promoter-driven AtELF3-HFC transgenic lines (AtELF3-2). To detect specific
332 co-precipitating proteins, we manually removed commonly identified contaminant proteins from
333 plant affinity purifications and mass spectrometry experiments (Van Leene et al., 2015), and
334 proteins identified from a control transgenic line expressing GFP-His₆-3xFlag described

335 previously (Huang et al., 2016a) (**Table 1**, the full list of identified proteins can be found in
336 **Supplemental Table S2**).

337
338 We have previously reported proteins that co-precipitated with ELF3 driven from its native
339 promoter using a similar AP-MS methodology (Huang et al., 2016a). When using the 35S
340 promoter driven AtELF3 transgenic line, we were able to generate a curated list of 22 proteins
341 that specifically co-precipitate with AtELF3, including all previously identified proteins, such as
342 all five phytochromes, PHOTOPERIODIC CONTROL OF HYPOCOTYL1 (PCH1) (Huang et
343 al., 2016c), and COP1 (**Table 1**). In addition, we also identified LIGHT-REGULATED WD 2
344 (LWD2) and SPA1-RELATED 4 (SPA4) as now co-precipitating with AtELF3. These additional
345 interactions may be a result of a combination of altered seedling age, expression level of the
346 ELF3 bait, or tissue-specificity of expression due to these purifications are from tissues where
347 the epitope-tagged transgene is constitutively over-expressed. However, since LWD2 is a known
348 component of the circadian clock (Wu et al., 2008) and SPA4 is a known component of the
349 COP1-SPA complex (Zhu et al., 2008), these interactions are likely to be relevant.

350
351 In comparing the list of BdELF3 and SvELF3 co-precipitated proteins with that of AtELF3, we
352 found that neither SvELF3 nor BdELF3 co-precipitated SPA2 and SPA4, components of the
353 COP1-SPA complex. In addition, SvELF3 did not co-precipitate MUT9-LIKE KINASE1, a
354 kinase with roles in chromatin modification and circadian rhythms as AtELF3 did (Huang et al.,
355 2016a; Wang et al., 2015). However, BdELF3 and SvELF3 associated with most of the proteins
356 found in AtELF3 AP-MS (20 out of 22 for BdELF3, 19 out of 22 for SvELF3), in at least one of
357 the replicate purifications from each monocot ortholog AP-MS. Therefore our data suggest that
358 BdELF3 and SvELF3 are integrated into a similar protein-protein interaction network as
359 AtELF3, which likely underlies their ability of broadly complementing *elf3* mutants.

360 361 **DISCUSSION**

362
363 Recent work in diverse plant species has found that the circadian clock plays critical roles in
364 regulating metabolism, growth, photoperiodism, and other agriculturally important traits (Bendix

365 et al., 2015; McClung, 2013; Shor and Green, 2016). While the relevance of the circadian clock
366 to plant fitness is unquestioned, it is unclear if the circadian clock components have conserved
367 function among different plant species. This is particularly true for the majority of clock proteins,
368 whose biological functions are currently poorly understood at the molecular level (Hsu and
369 Harmer, 2014). Also, the divergent modes of growth regulation and photoperiodism between
370 monocots and dicots suggest that the clock evolved to have altered roles in regulating these
371 physiological responses between lineages (Matos et al., 2014; Poire et al., 2010; Song et al.,
372 2014). Here we asked if orthologs of ELF3 from two monocots could complement any of the
373 loss-of-function phenotypes in the model dicot plant *A. thaliana*. In this study we found that
374 ELF3 from either *B. distachyon* or *S. viridis* could complement the hypocotyl elongation, early
375 flowering, and arrhythmic clock phenotype of the *elf3* mutant in *A. thaliana*, despite the
376 variations in protein sequences and evolutionary divergence between monocot and dicot plants.
377 These data suggest that monocot ELF3s can functionally substitute for *A. thaliana* ELF3, albeit
378 with varying efficacy. Since monocot and dicot ELF3 are largely different in the protein
379 sequences, functional conservation of ELF3 orthologs also leads to the next open question of
380 identifying the functional domains within ELF3.

381
382 Previously, comparison of ELF3 homologs has identified at least five conserved regions that may
383 be important for function (**Supplemental Figure S1**) (Liu et al., 2001; Saito et al., 2012; Weller
384 et al., 2012). Our multiple sequence alignments also show that at least two regions of AtELF3,
385 namely the N-terminus (AA 1~49) and one middle region (AA 317~389) share many conserved
386 residues with ELF3 orthologs in grasses (**Supplemental Figure S1**). These regions fall within
387 known fragments that are sufficient for binding to phyB (Liu et al., 2001), COP1 (Yu et al.,
388 2008), or ELF4 (Herrero et al., 2012). Consistent with the hypothesis that these conserved
389 regions are critical for proper ELF3 function, a single amino acid substitution (A362V) within
390 this middle region results in defects of ELF3 nuclear localization and changes in the circadian
391 clock period (Anwer et al., 2014). In addition, our protein-protein interaction study and AP-MS
392 analysis show that both monocot ELF3 can form composite ECs (**Figure 5**) and that all three
393 ELF3 homologs interact with an almost identical set of proteins *in vivo* (**Table 1**), further
394 suggesting that one or more of the conserved regions may mediate the binding between ELF3

395 and its known interacting proteins. Furthermore, the similar pool of ELF3 interacting proteins
396 identified by Bd/SvELF3 AP-MS suggests that the overall conformation of ELF3 ortholog
397 proteins is conserved and that similar complexes and interactions with ELF3 orthologs may form
398 in monocot species. However, whether these interactions form *in planta* and have the same effect
399 on physiology is unclear. For example, *S. viridis* data generated here and public data for *B.*
400 *distachyon* and *O. sativa*, showed that ELF3 does not cycle under circadian conditions, which
401 differs from Arabidopsis. Further, different from the fact that the clock plays a key role in
402 regulating elongation in *A. thaliana* (Nozue et al., 2007), the circadian clock has no influence on
403 growth in C3 model grass *B. distachyon*, despite robust oscillating expression of putative clock
404 components (Matos et al., 2014). Similarly, ELF3 from rice (*Oryza sativa*) and soybean
405 promotes flowering and senescence (Lu et al., 2017; Saito et al., 2012; Sakuraba et al., 2016;
406 Yang et al., 2013; Zhao et al., 2012), while in *A. thaliana*, ELF3 represses these responses (Liu et
407 al., 2001; Sakuraba et al., 2014; Zagotta et al., 1992), which suggests significant rewiring of
408 ELF3 regulated photoperiodic responses of flowering between short-day (rice/soybean) and
409 long-day (*A. thaliana*) plants. Alternatively, ELF3 may form distinct interactions and complexes
410 in monocot species that were not identified in our trans-species complementation analysis.
411 Clearly, further work is required to understand ELF3 function in monocots beyond the studies
412 presented here.

413
414 In addition to the molecular characterization of ELF3, our analysis of circadian-regulated genes
415 in *S. viridis* after photo- and thermo-entrainment found significant differences in the behavior of
416 the clock when compared to other monocots. Although the number of circadian regulated genes
417 is comparable to studies done in corn and rice after photo-entrainment (between 10-12%)
418 (Filichkin et al., 2011; Khan et al., 2010), we found that very few genes (~1%) continue to cycle
419 after release from temperature entrainment in *S. viridis* (**Supplemental Figure S4**) when
420 compared to rice (~11%) (Filichkin et al., 2011). This may reflect a fundamental difference in
421 how the clock interfaces with temperature between these monocot species. Furthermore,
422 proportions of circadian regulated genes upon photo-entrainment in all three monocot plants
423 (**Supplement Figure S4**) (Filichkin et al., 2011; Khan et al., 2010) are much smaller than the
424 approximately 30% reported for *A. thaliana* (Covington et al., 2008), suggesting the divergence

425 of clock functions through evolution or domestication. Further comparisons of circadian
426 responses among monocots or between monocots and dicots will help to determine the molecular
427 underpinning of these differences.

428
429 In summary, we find that BdELF3 and SvELF3 form similar protein complexes *in vivo* as
430 AtELF3, which likely allows for functional complementation of loss-of-function of *elf3* despite
431 relatively low sequence conservation. We also present an online query tool, Diel Explorer that
432 allows for exploration of circadian gene expression in *S. viridis*, which illustrate fundamental
433 differences in clock function among monocots and between monocots and dicots. Collectively,
434 this work is a first step toward functional understanding of the circadian clock in two model
435 monocots, *S. viridis* and *B. distachyon*.

436

437 **MATERIALS AND METHODS**

438

439 **Plant materials and growth conditions**

440

441 For *A. thaliana*, wild type (Columbia-0) and *elf3-2* plants carrying the CCA1::LUC reporter were
442 described previously (Nusinow et al., 2011; Pruneda-Paz et al., 2009). Seeds were surface
443 sterilized and plated on 1/2x Murashige and Skoog (MS) basal salt medium with 0.8% agar + 1%
444 (w/v) sucrose. After 3 days of stratification, plates were placed horizontally in a Percival
445 incubator (Percival Scientific, Perry, IA), supplied with $80 \mu\text{mol m}^{-2} \text{sec}^{-1}$ white light and set to a
446 constant temperature of 22°C. Plants were grown under 12 h light / 12 h dark cycles (12L:12D)
447 for 4 days (for physiological experiments) or for 10 days (for AP-MS) before assays.

448

449 For *S. viridis* circadian expression profiling by RNA-seq, seeds were stratified for 5 days at 4°C
450 before being moved to entrainment conditions. Plants were grown under either LDHH or LLHC
451 (L: light, D: dark, H: hot, C: cold) entrainment condition, and then sampled for RNA-seq in
452 constant light and constant temperature (32°C) conditions (F, for free-running) every 2 hours for
453 48 hours. Light intensity was set to $400 \mu\text{mol m}^{-2} \text{sec}^{-1}$ white light. In LDHH-F, stratified *S.*
454 *viridis* seeds were grown for 10 days under 12L:12D and constant temperature (32°C) before

455 sampling in constant light and constant temperature. In LLHC-F, stratified *S. viridis* seeds were
456 grown for 10 days under constant light conditions and cycling temperature conditions 12 h at
457 32°C (subjective day) / 12 h at 22°C (subjective night) before sampling in constant conditions.
458 Two experimental replicates were collected for each entrainment condition.

459

460 **Setaria circadian RNA-seq**

461

462 The second leaf from the top of seventeen *S. viridis* plants was selected for RNA-seq sampling at
463 each time point for each sampling condition. Five replicate samples were pooled after being
464 ground in liquid nitrogen and resuspended in lithium chloride lysis binding buffer (Wang et al.,
465 2011a). RNA-seq libraries from leaf samples were constructed according to the previous
466 literature (Wang et al., 2011a) with one major modification. Rather than extracting RNA then
467 mRNA from ground leaf samples (Wang et al., 2011a), mRNA was extracted directly from
468 frozen ground leaf samples similar to the method described in (Kumar et al., 2012), except that
469 two additional rounds of wash, binding, and elution steps after treatment with EDTA were
470 necessary to remove rRNA from samples. mRNA quantity was assessed using a Qubit with a
471 Qubit RNA HS Kit and mRNA quality was assessed using a Bioanalyzer and Plant RNA PiCO
472 chip. 96 library samples were multiplexed 12 per lane, for a total of 8 lanes of Illumina HiSeq
473 2000 sequencing. Paired end 101 bp Sequencing was done at MOgene (St. Louis, MO). Raw
474 data and processed data can be found on NCBI's Gene Expression Omnibus (GEO; (Barrett et
475 al., 2013; Edgar et al., 2002)) and are accessible with identification number GSE97739
476 (<https://www.ncbi.nlm.nih.gov/geo/query/acc.cgi?acc=GSE97739>).

477

478 RNA-seq data was trimmed with BBTools (v36.20) using parameters: ktrim=r k=23 mink=11
479 hdist=1 tpe tbo ktrim=l k=23 mink=11 hdist=1 tpe tbo qtrim=rl trimq=20 minlen=20 (Bushnell,
480 2016). Any parameters not specified were run as default. Before trimming we had 1,814,939,650
481 reads with a mean of 18,905,621 reads per sample and a standard deviation of 2,875,187. After
482 trimming, we have 1,646,019,593 reads with a mean of 17,146,037 reads per sample and a
483 standard deviation of 2,411,061. Kallisto (v 0.42.4; (Bray et al., 2016)) was used to index the
484 transcripts with the default parameters and the *S. viridis* transcripts fasta file (*Sviridis_311_v1.1*)

485 from Phytozome (Goodstein et al., 2012). The reads were quantified with parameters:-t 40 -b
486 100. Any parameters not specified were run as default. Kallisto output was formatted for
487 compatibility with JTKCycle (v3.1; (Hughes et al., 2010)) and circadian cycles were detected.
488 To query the *S. viridis* expression data we developed Diel Explorer. The tool can be found at
489 <http://shiny.bioinformatics.danforthcenter.org/diel-explorer/>. Underlying code for Diel Explorer
490 is available on Github (<https://github.com/danforthcenter/diel-explorer>).

491

492 **Plasmid constructs and generation of transgenic plants**

493

494 Coding sequences (without the stop codon) of *AtELF3* (*AT2G25930*) and *SvELF3a*
495 (*Sevir.5G206400.1*) were cloned into the pENTR/D-TOPO vector (ThermoFisher Scientific,
496 Waltham, MA), verified by sequencing and were recombined into the pB7HFC vector (Huang et
497 al., 2016a) using LR Clonease (ThermoFisher Scientific, Waltham, MA). Coding sequence of
498 *BdELF3* (*Bradi2g14290.1*) was submitted to the U.S. Department of Energy Joint Genome
499 Institute (DOE-JGI), synthesized by the DNA Synthesis Science group, and cloned into the
500 pENTR/D-TOPO vector. Sequence validated clones were then recombined into pB7HFC as
501 described above. The pB7HFC-At/Bd/SvELF3 constructs were then transformed into *elf3-2*
502 [*CCA1::LUC*] plants by the floral dip method (Zhang et al., 2006). Homozygous transgenic
503 plants were validated by testing luciferase bioluminescence, drug resistance, and by PCR-based
504 genotyping. All primers used in this paper were listed in **Supplemental Table S1**.

505

506 **Hypocotyl and flowering time measurement**

507

508 20 seedlings of each genotype were arrayed and photographed with a ruler for measuring
509 hypocotyl length using the ImageJ software (NIH) (Schneider et al., 2012). The procedure was
510 repeated three times. For measuring flowering time, 12 plants of each genotype were placed in a
511 random order and were grown under the long day condition (light : dark = 16 : 8 hours). The
512 seedlings were then observed every day at 12:00 PM; the date on which each seedling began
513 flowering, indicated by the growth of a ~1 cm inflorescence stem, was recorded along with the
514 number of rosette leaves produced up to that date.

515

516 **Circadian assays in *A. thaliana***

517

518 Seedlings were transferred to fresh 1/2x MS plates after 5 days of entrainment under the
519 12L:12D condition and sprayed with sterile 5 mM luciferin (Gold Biotechnology, St. Louis, MO)
520 prepared in 0.1% (v/v) Triton X-100 solution. Sprayed seedlings were then imaged in constant
521 light ($70 \mu\text{mol m}^{-2} \text{sec}^{-1}$, wavelengths 400, 430, 450, 530, 630, and 660 set at intensity 350
522 (Heliospectra LED lights, Göteborg, Sweden)). Bioluminescence was recorded after a 120-180s
523 delay to diminish delayed fluorescence (Gould et al., 2009) over 5 days using an ultra-cooled
524 CCD camera (Pixis 1024B, Princeton Instruments) driven by Micro-Manager software (Edelstein
525 et al., 2010; Edelstein et al., 2014). The images were processed in stacks by Metamorph software
526 (Molecular Devices, Sunnyvale, CA), and rhythms determined by fast Fourier transformed non-
527 linear least squares (FFT-NLLS) (Plautz et al., 1997) using the interface provided by the
528 Biological Rhythms Analysis Software System 3.0 (BRASS) available at <http://www.amillar.org>.

529

530 **Yeast two-hybrid analysis**

531

532 Yeast two-hybrid assays were carried out as previously described (Huang et al., 2016a). In brief,
533 the DNA binding domain (DBD) or activating domain (AD)-fused constructs were transformed
534 using the Li-Ac transformation protocol (Clontech) into *Saccharomyces cerevisiae* strain Y187
535 (MAT α) and the AH109 (MATa), respectively. Two strains of yeast were then mated to generate
536 diploid with both DBD and AD constructs. Protein-protein interaction was tested in diploid yeast
537 by replica plating on CSM –Leu –Trp –His media supplemented with extra Adenine (30mg/L
538 final concentration) and 2mM 3-Amino-1,2,4-triazole (3AT). Pictures were taken after 4-day
539 incubation at 30 °C. All primers used for cloning plasmid constructs were listed in

540 **Supplemental Table S1.**

541

542 **Protein extraction and western blotting**

543

544 Protein extracts were made from 10-day-old seedlings as previously described (Huang et al.,
545 2016a) and loaded 50 µg to run 10% SDS-PAGE. For western blots, all of the following primary
546 and secondary antibodies were diluted into PBS + 0.1% Tween and incubated at room
547 temperature for 1 hour: anti-FLAG®M2-HRP (Sigma, A8592, diluted at 1:10,000) and anti-
548 Rpt5-rabbit (ENZO Life Science, BML-PW8245-0025, diluted at 1:5000) and anti-Rabbit-HRP
549 secondary antibodies (Sigma, A0545, diluted at 1:10,000).

550

551 **Affinity purification and mass spectrometry**

552

553 Protein extraction methods and protocols for AP-MS were described previously (Huang et al.,
554 2016a; Huang et al., 2016b; Huang and Nusinow, 2016a; Huang et al., 2016c)}. In brief,
555 transgenic seedlings carrying the At/Bd/SvELF3-HFC constructs were grown under 12L:12D
556 conditions for 10 days and were harvested at dusk (ZT12). 5 grams of seedlings were needed per
557 replicate to make protein extracts, which underwent tandem affinity purification utilizing the
558 FLAG and His epitopes of the fusion protein. Purified samples were reduced, alkylated and
559 digested by trypsin. The tryptic peptides were then injected to an LTQ-Orbitrap Velos Pro
560 (ThermoFisher Scientific, Waltham, MA) coupled with a U3000 RSLCnano HPLC
561 (ThermoFisher Scientific, Waltham, MA) with settings described previously (Huang et al.,
562 2016a).

563

564 **AP-MS data analysis**

565

566 Data analysis was done as previously described (Huang et al., 2016a). The databases searched
567 were TAIR10 database (20101214, 35,386 entries) and the cRAP database
568 (<http://www.thegpm.org/cRAP/>). Peptide identifications were accepted if they could be
569 established at greater than 95.0% probability and the Scaffold Local FDR was <1%. Protein
570 identifications were accepted if they could be established at greater than 99.0% probability as
571 assigned by the Protein Prophet algorithm (Keller et al., 2002; Nesvizhskii et al., 2003). A full
572 list of all identified proteins (reporting total/exclusive unique peptide count and percent
573 coverage) can be found in **Supplemental Table S2**. The mass spectrometry proteomics data

574 have been deposited to the ProteomeXchange Consortium (Vizcaino et al., 2014) via the PRIDE
575 partner repository with the dataset identifier PXD006352 and 10.6019/PXD006352.

576

577

578 **Supplemental Data**

579 **Supplemental Table S1.** List of all primers used.

580 **Supplemental Table S2.** A full list of At/Bd/SvELF3 associated proteins identified from AP-
581 MS.

582 **Supplemental Figure S1.** Multiple sequence alignments of ELF3 orthologs **Supplemental**

583 **Figure S2.** Diel and circadian expression of *BdELF3* from the DIURNAL database.

584 **Supplemental Figure S3.** Example of the Diel Explorer interface.

585 **Supplemental Figure S4.** Summary of circadian regulated genes in *S. viridis*.

586 **Supplemental Figure S5.** Circadian expression of selected *A. thaliana* clock genes from the
587 DIURNAL database.

588 **Supplemental Figure S6.** Anti-FLAG western of ELF3 transgenic lines used for
589 complementation analysis.

590 **Supplemental Figure S7.** Relative Amplitude Error vs period plots of transgenic *elf3-2* plants
591 expressing At/Bd/SvELF3.

592

Table 1. Proteins co-purified with ELF3 orthologs from AP-MS

AGI number	Protein Name	Molecular Weight	Exclusive Unique Peptide Count/Percent Coverage									
			AtELF3		SvELF3 #2		SvELF3 #3		BdELF3 #2		BdELF3 #3	
			rep1	rep2	rep1	rep2	rep1	rep2	rep1	rep2	rep1	rep2
n/a	AtELF3-HFC	84 kDa	20/36%	19/31%	—	—	—	—	—	—	—	—
n/a	SvELF3-HFC	87 kDa	—	—	19/40%	29/48%	22/42%	33/54%	—	—	—	—
n/a	BdELF3-HFC	86 kDa	—	—	—	—	—	—	25/42%	26/43%	21/35%	19/33%
AT2G18790	phyB	129 kDa	23/37%	24/37%	31/54%	28/42%	32/50%	30/45%	22/34%	22/33%	24/40%	24/37%
AT5G35840	phyC	124 kDa	22/29%	23/29%	27/37%	32/39%	27/37%	33/42%	20/24%	18/22%	19/23%	22/27%
AT5G43630	TZP	91 kDa	14/21%	12/17%	13/23%	20/33%	13/22%	24/36%	12/18%	13/19%	14/20%	14/22%
AT4G18130	phyE	123 kDa	11/16%	19/27%	12/18%	18/23%	11/19%	17/24%	6/7%	10/14%	14/19%	13/18%
AT2G16365.2	PCH1b	51 kDa	9/25%	9/25%	9/30%	14/43%	11/36%	16/49%	9/26%	9/28%	11/32%	11/32%
AT2G46340	SPA1	115 kDa	8/14%	7/9%	2/2%	4/8%	2/3%	4/9%	—	—	6/8%	5/6%
AT3G42170c	DAYSLEEPER	79 kDa	8/19%	5/11%	—	7/16%	—	12/27%	3/7%	4/8%	2/4%	—
AT2G32950	COP1	76 kDa	8/16%	9/17%	6/17%	4/8%	3/6%	5/12%	1/2%	2/4%	6/11%	4/8%
AT3G22380	TIC	165 kDa	5/5%	5/5%	4/4%	12/12%	3/3%	15/15%	4/4%	3/3%	1/1%	1/1%
AT4G11110	SPA2	115 kDa	5/11%	6/12%	—	—	—	—	—	—	—	—
AT2G40080	ELF4	12 kDa	4/50%	4/50%	3/42%	5/68%	4/60%	4/60%	4/50%	4/50%	5/68%	5/68%
AT1G09340c	CRB	43 kDa	4/16%	3/12%	1/4%	1/4%	1/4%	4/18%	1/4%	1/4%	—	—
AT3G13670	MLK4	79 kDa	3/10%	3/13%	—	—	—	1/2%	6/21%	3/10%	3/11%	5/13%
AT5G61380	TOC1	69 kDa	2/4%	2/4%	2/5%	—	3/7%	—	—	—	1/2%	1/2%
AT1G53090	SPA4	89 kDa	2/6%	5/14%	—	—	—	—	—	—	—	—
AT5G18190	MLK1	77 kDa	2/12%	2/11%	—	—	—	—	2/14%	2/14%	2/11%	2/11%
AT3G26640	LWD2	39 kDa	2/21%	1/21%	—	1/16%	—	2/22%	3/27%	1/16%	—	—
AT1G12910	LWD1	39 kDa	2/21%	3/30%	2/21%	4/31%	1/17%	2/21%	2/22%	2/19%	0/11%	0/11%
AT4G16250	phyD	129 kDa	1/10%	1/11%	2/12%	1/12%	3/15%	2/12%	2/11%	1/10%	2/14%	2/12%
AT1G09570	phyA	125 kDa	1/1%	1/1%	7/11%	5/5%	7/11%	2/2%	—	2/2%	6/7%	3/4%

AT2G25760	MLK3	76 kDa	1/5%	1/6%	—	1/4%	—	1/4%	3/13%	2/9%	2/9%	2/9%
AT3G03940	MLK2	78 kDa	1/8%	1/8%	—	1/7%	—	0/2%	3/20%	2/13%	3/13%	1/9%
AT3G46640	LUX	35 kDa	1/3%	1/3%	0/10%	3/19%	0/10%	3/23%	1/3%	1/3%	1/3%	2/11%
AT1G17455	ELF4-L4	13 kDa	—	—	—	—	—	—	1/23%	1/23%	—	—
AT1G72630	ELF4-L2	13 kDa	—	1/11%	—	1/13%	—	—	1/22%	1/22%	1/22%	1/11%

Proteins co-purified with ELF3 orthologs (AtELF3, SvELF3 and BdELF3, with C-terminal His6-3xFLAG tag) were identified from affinity purification coupled with mass spectrometry (AP-MS) analyses using 12L:12D grown, 10-day-old transgenic plants (in *elf3-2 null* mutant backgrounds) harvested at ZT12.

^a All listed proteins match 99% protein threshold, minimum number peptides of 2 and peptide threshold as 95%. Proteins not matching the criteria were marked with "—".

^b Percent coverage for PCH1 is calculated using protein encoded by *At2g16365.2*

^c These proteins have been noted as frequently identified proteins in AP-MS experiments (see Van Leene, 2015).

593 **FIGURE LEGENDS**

594

595 **Figure 1.** Circadian expression profiles of putative *S. viridis* clock components from Diel
596 Explorer using time-course RNA-seq data. *S. viridis* plants were entrained by either photocycle
597 (LDHH) or thermocycle (LLHC), followed by being sampled every 2 hours for 48 hours under
598 constant temperature and light conditions (Free-Running; F) to generate time-course RNA-seq
599 data. Mean values of Transcripts per Kilobase Million (TPM) from two experimental replicates
600 for each timepoints per gene were plotted.

601

602 **Figure 2.** ELF3 orthologs suppress hypocotyl elongation defects in *elf3-2*. The hypocotyls of 20
603 seedlings of wild type, *elf3-2* mutant, AtELF3 *elf3-2*, BdELF3 *elf3-2*, and SvELF3 *elf3-2* (two
604 independent transgenic lines for each ELF3 ortholog) were measured at 4 days after germination
605 under 12-hour light :12-hour dark growth conditions at 22 °C. Upper panel shows representative
606 seedlings of each genotype, with scale bar equal to 5 mm. Mean and 95% confidence intervals
607 are plotted as crosshairs. This experiment was repeated three times with similar results. ANOVA
608 analysis with Bonferroni correction was used to generate adjusted P values, * < 0.05, ** < 0.01,
609 **** < 0.0001.

610

611 **Figure 3.** ELF3 orthologs suppress time to flowering of *elf3-2*. 12 wild type, *elf3-2* mutant,
612 AtELF3 *elf3-2*, BdELF3 *elf3-2*, and SvELF3 *elf3-2* seedlings from two independent
613 transformations were measured for days (A) and number of rosette leaves (B) at flowering (1 cm
614 inflorescence). Mean and 95% confidence intervals are plotted as crosshairs. This experiment
615 was repeated twice with similar results. ANOVA analysis with Bonferroni correction was used to
616 generate adjusted P values, ** < 0.01, *** < 0.001, **** < 0.0001, of measurements when
617 compared to the *elf3-2* mutant line.

618

619 **Figure 4.** ELF3 orthologs can recover *CCA1::LUC* rhythms and amplitude in *elf3-2* mutants. 8
620 seedlings of wild type, *elf3-2* mutant, AtELF3 *elf3-2* (A), BdELF3 *elf3-2* (B), and SvELF3 *elf3-2*
621 (C) from two independent transformations were imaged for bioluminescence under constant light
622 after entrainment in 12-hour light :12-hour dark growth conditions at 22 °C. Each plot shows

623 average bioluminescence of all seedlings along with 95% confidence interval (error bars). This
624 experiment was repeated twice with similar results. Note that wild type and *elf3-2* mutant data
625 was plotted on all graphs for comparison. **(D)** Periods of seedlings. Only periods with a Relative
626 Amplitude Error below 0.5 (see **Supplemental Figure S7**) were plotted. Mean and 95%
627 confidence intervals are plotted as crosshairs. ANOVA analysis with Bonferroni correction was
628 used to generate adjusted P values, ** < 0.01, *** < 0.001, **** < 0.0001, of measurements
629 when compared to the wild type.

630

631 **Figure 5.** Both BdELF3 and SvELF3 can directly bind to AtELF4 and AtLUX. Yeast two-hybrid
632 analysis of testing if either BdELF3 **(A)** or SvELF3 **(B)** can directly interact with either AtELF4,
633 the N-terminal half of AtLUX (AtLUX-N, a.a. 1-143) or the C-terminal half of AtLUX (AtLUX-
634 C, a.a. 144-324). –LW tests for the presence of both bait (DBD) and pray (AD) vectors, while the
635 –LWH + 3AT tests for interaction. Vector alone serves as interaction control. This experiment
636 was repeated twice with similar results.

637

638

639

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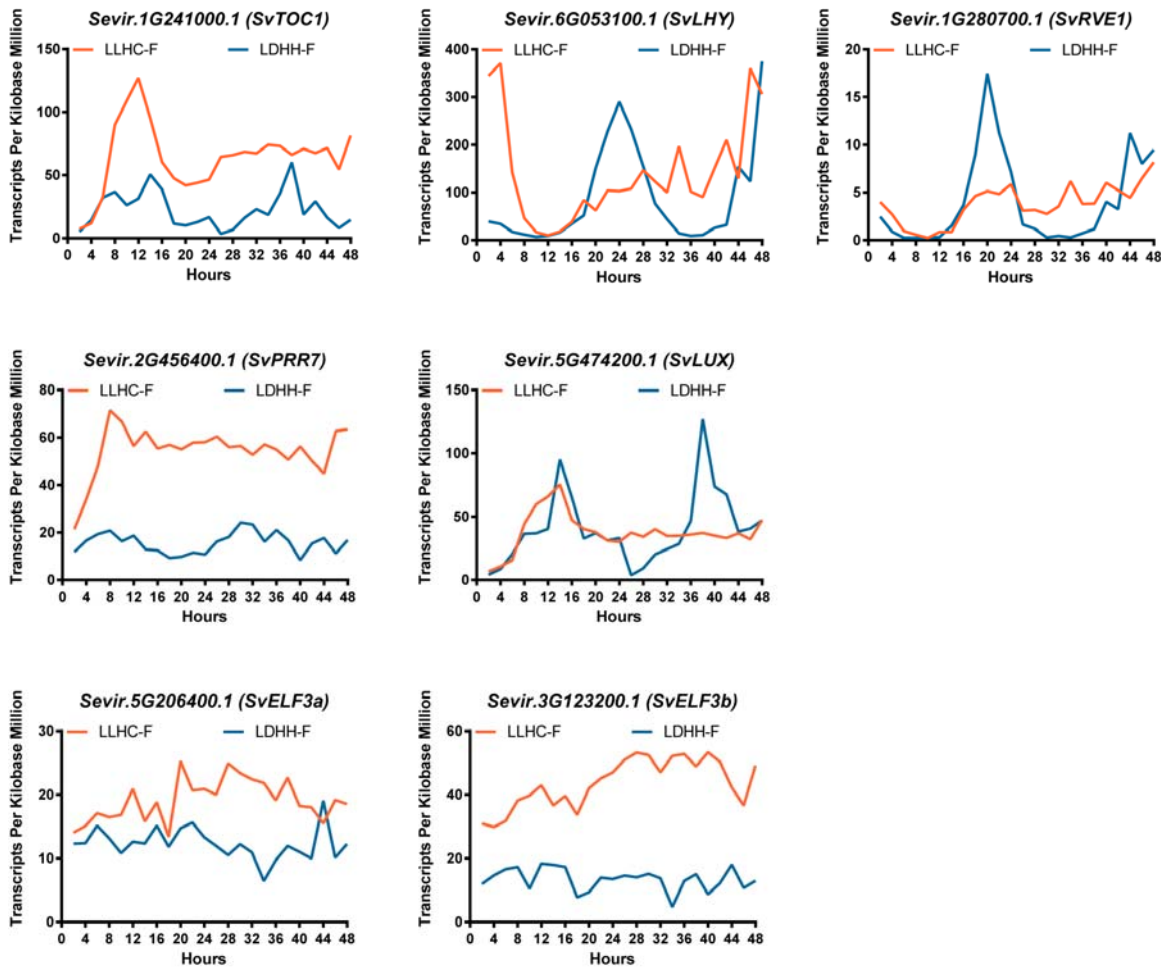


Figure 1. Circadian expression profiles of putative *S. viridis* clock components from Diel Explorer using time-course RNA-seq data. *S. viridis* plants were entrained by either photocycle (LDHH) or thermocycle (LLHC), followed by being sampled every 2 hours for 48 hours under constant temperature and light conditions (Free-Running; F) to generate time-course RNA-seq data. Mean values of Transcripts per Kilobase Million (TPM) from two experimental replicates for each timepoints per gene were plotted.

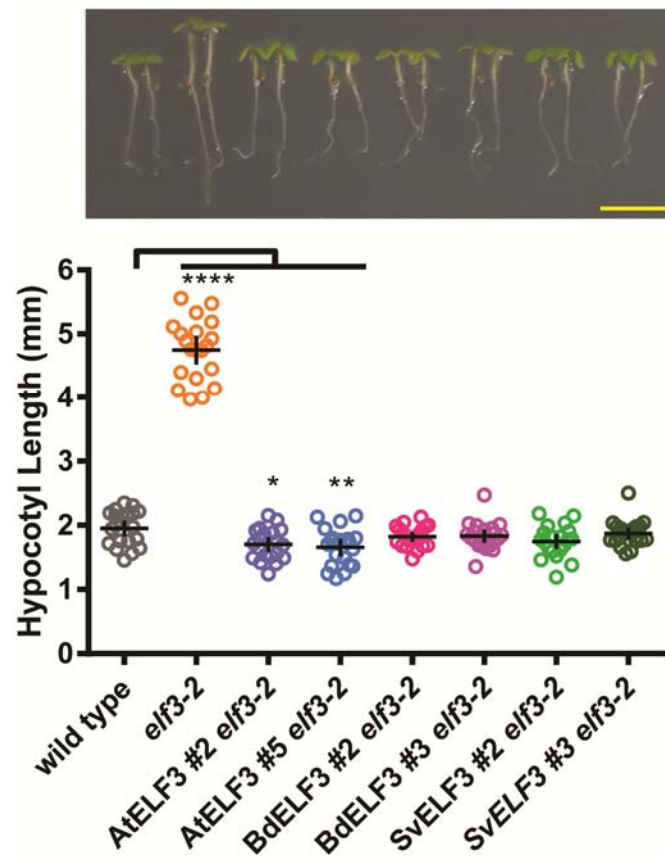


Figure 2. ELF3 orthologs suppress hypocotyl elongation defects in *elf3-2*. The hypocotyls of 20 seedlings of wild type, *elf3-2* mutant, AtELF3 *elf3-2*, BdELF3 *elf3-2*, and SvELF3 *elf3-2* (two independent transgenic lines for each ELF3 ortholog) were measured at 4 days after germination under 12-hour light :12-hour dark growth conditions at 22 °C. Upper panel shows representative seedlings of each genotype, with scale bar equal to 5 mm. Mean and 95% confidence intervals are plotted as crosshairs. This experiment was repeated three times with similar results. ANOVA analysis with Bonferroni correction was used to generate adjusted P values, * < 0.05, ** < 0.01, **** < 0.0001.

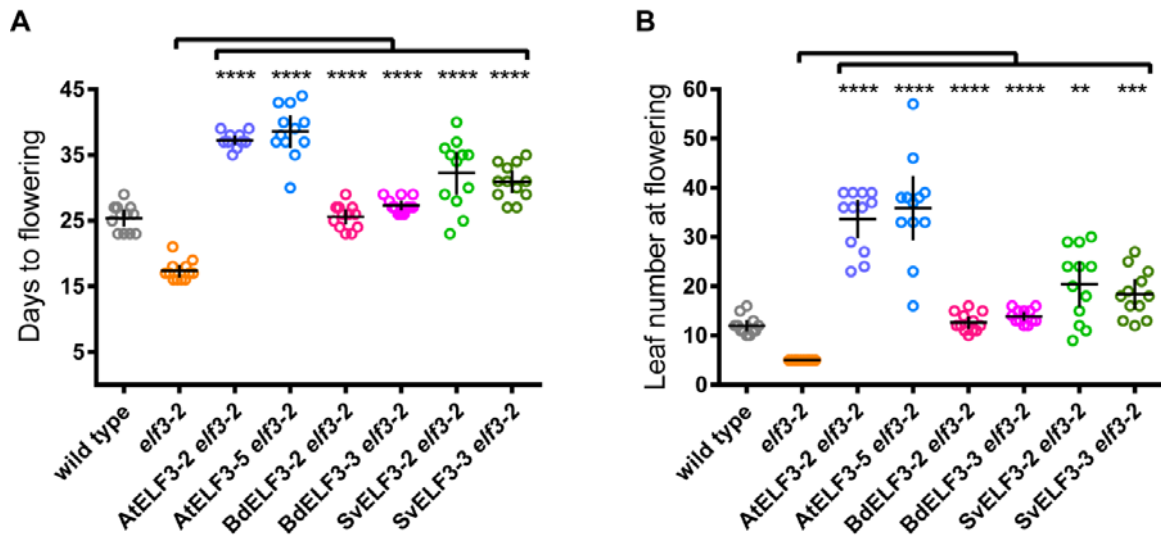


Figure 3. ELF3 orthologs suppress time to flowering of *elf3-2*. 12 wild type, *elf3-2* mutant, AtELF3 *elf3-2*, BdELF3 *elf3-2*, and SvELF3 *elf3-2* seedlings from two independent transformations were measured for days (A) and number of rosette leaves (B) at flowering (1 cm inflorescence). Mean and 95% confidence intervals are plotted as crosshairs. This experiment was repeated twice with similar results. ANOVA analysis with Bonferroni correction was used to generate adjusted P values, ** < 0.01, *** < 0.001, **** < 0.0001, of measurements when compared to the *elf3-2* mutant line.

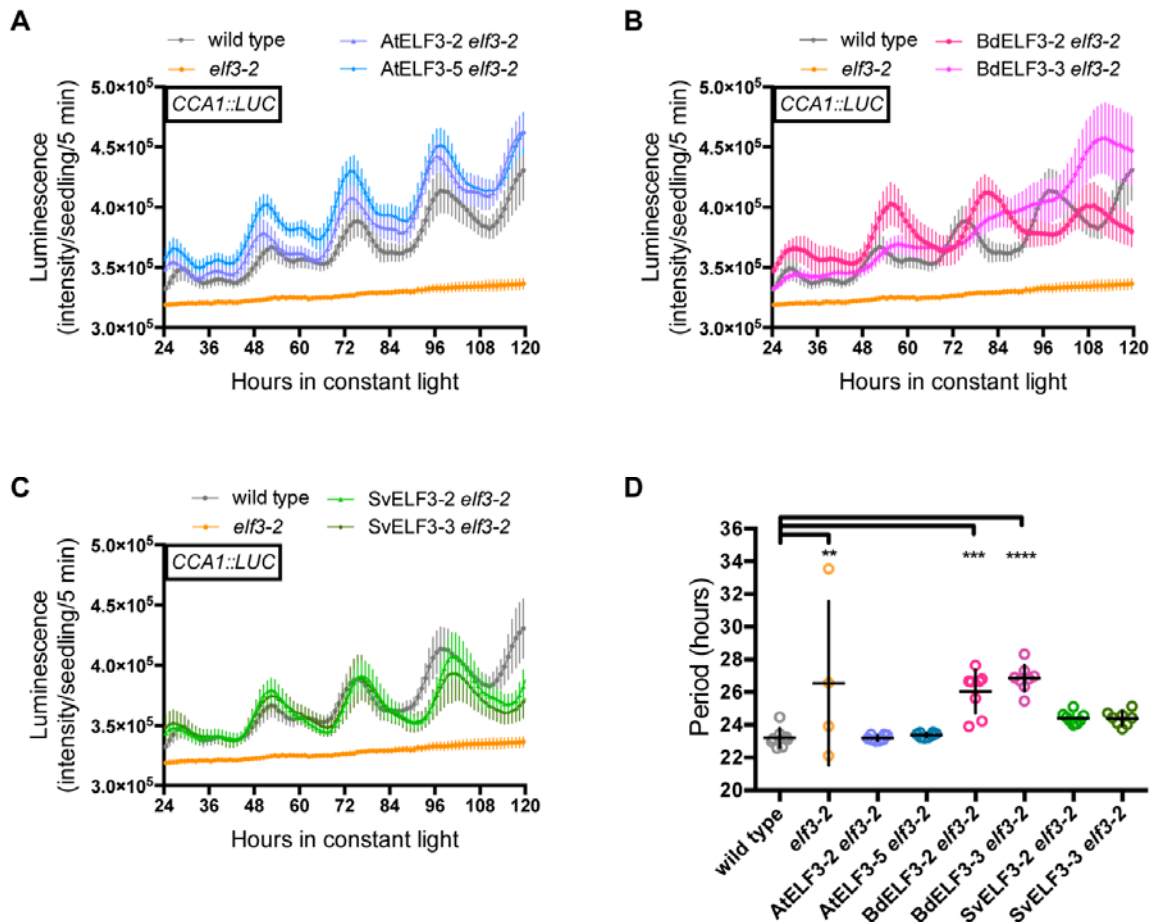


Figure 4. ELF3 orthologs can recover *CCA1::LUC* rhythms and amplitude in *elf3-2* mutants. 8 seedlings of wild type, *elf3-2* mutant, AtELF3 *elf3-2* (A), BdELF3 *elf3-2* (B), and SvELF3 *elf3-2* (C) from two independent transformations were imaged for bioluminescence under constant light after entrainment in 12-hour light :12-hour dark growth conditions at 22 °C. Each plot shows average bioluminescence of all seedlings along with 95% confidence interval (error bars). This experiment was repeated twice with similar results. Note that wild type and *elf3-2* mutant data was plotted on all graphs for comparison. (D) Periods of seedlings. Only periods with a Relative Amplitude Error below 0.5 (see Supplemental Figure 7) were plotted. Mean and 95% confidence intervals are plotted as crosshairs. ANOVA analysis with Bonferroni correction was used to generate adjusted P values, ** < 0.01, *** < 0.001, **** < 0.0001, of measurements when compared to the wild type.

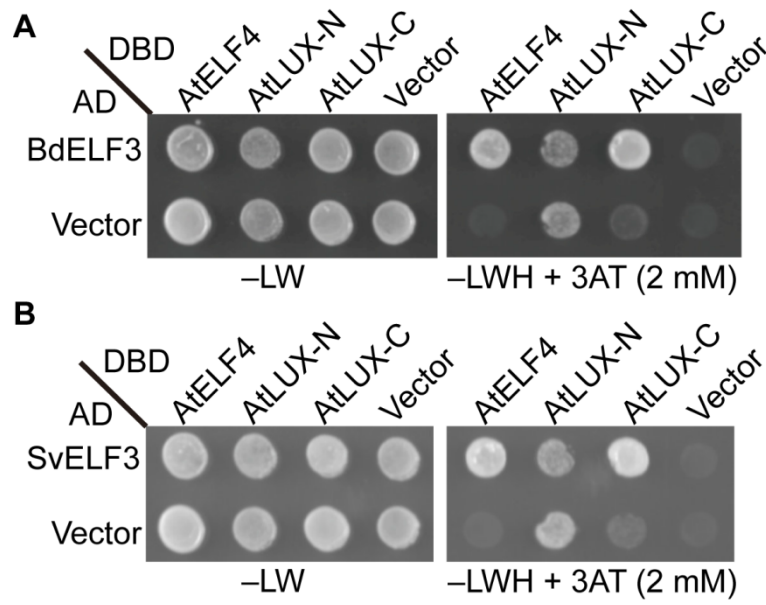


Figure 5. Both BdELF3 and SvELF3 can directly bind to AtELF4 and AtLUX. Yeast two-hybrid analysis of testing if either BdELF3 (**A**) or SvELF3 (**B**) can directly interact with either AtELF4, the N-terminal half of AtLUX (AtLUX-N, a.a. 1-143) or the C-terminal half of AtLUX (AtLUX-C, a.a. 144-324). -LW tests for the presence of both bait (DBD) and prey (AD) vectors, while the -LWH + 3AT tests for interaction. Vector alone serves as interaction control. This experiment was repeated twice with similar results.