- 1 REVEILLE 7 inhibits the expression of the circadian clock gene EARLY
- 2 FLOWERING 4 to fine-tune hypocotyl growth in response to warm

# 3 temperatures

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- 15 **Summary for the icon:** Warm temperature-induced RVE7 fine-tunes 16 thermoresponsive hypocotyl growth by inhibiting the expression of *ELF4* in 17 *Arabidopsis*, indicating that ELF4 is important for thermomorphogenesis in 18 plants.
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27 Abstract

28 The circadian clock maintains the daily rhythms of plant growth and anticipates predictable ambient temperature cycles. The evening complex (EC), 29 30 comprising EARLY FLOWERING 3 (ELF3), ELF4, and LUX ARRHYTHMO, plays an essential role in suppressing thermoresponsive hypocotyl growth by 31 32 negatively regulating PHYTOCHROME INTERACTING FACTOR 4 (PIF4) activity and its downstream targets in Arabidopsis thaliana. However, how EC 33 activity is attenuated by warm temperatures remains unclear. Here, we 34 demonstrate that warm temperature-induced REVEILLE 7 (RVE7) fine-tunes 35 thermoresponsive growth in Arabidopsis by repressing *ELF4* expression. *RVE7* 36 transcript and RVE7 protein levels increased in response to warm temperatures. 37 Under warm temperature conditions, an rve Closs-of-function mutant had 38 shorter hypocotyls, while overexpressing RVE7 promoted hypocotyl elongation. 39 PIF4 accumulation and downstream transcriptional effects were reduced in the 40 rve7 mutant but enhanced in RvE7 overexpression plants under warm 41 conditions. RVE7 associates with the evening element in the ELF4 promoter 42 43 and directly represses its transcription. ELF4 is epistatic to RVE7, and 44 overexpressing *ELF4* suppressed the phenotype of the *RVE7* overexpression line under warm temperature conditions. Together, our results identify RVE7 as 45 an important regulator of thermoresponsive growth that functions (in part) by 46 47 controlling *ELF4* transcription, highlighting the importance of ELF4 for 48 thermomorphogenesis in plants.

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50 **Keywords:** *Arabidopsis thaliana*, ELF4, Hypocotyl growth, RVE7, Warm 51 temperatures

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#### 54 Introduction

55 Plant growth and development are widely influenced by environmental conditions, including ambient temperatures (Vu et al., 2019). Plants sense 56 57 elevated ambient temperatures and transduce the warm temperature signal to downstream transcription factors to regulate gene expression and trigger 58 59 various These physiological responses. responses include rapid 60 hypocotyl/petiole growth, increased leaf hyponasty, and accelerated flowering 61 via a process known as thermomorphogenesis (Casal and Balasubramanian, 2019; Sun et al., 2020; Wang et al., 2021). 62

In Arabidopsis (Arabidopsis thaliana), warm temperatures are sensed by at 63 least three thermosensors: phytochrome B (phyB) EARLY FLOWERING 3 64 (ELF3), and PHYTOCHROME INTERACTING FACTOR 7 (PIF7) (Lin et al., 65 2020), phyB and ELF3 are repressors of the transcription factor PIF4, which 66 plays a key role in thermomorphogenesis (Jung et al., 2016; Legris et al., 2016). 67 phyB, a well-known photoreceptor, rapidly reverts from its active form Pfr to its 68 inactive form Pr under warm temperature conditions (Klose et al., 2020), which 69 reduces its inhibition of PIF4 Jung et al., 2016; Legris et al., 2016). ELF3 70 undergoes liquid-liquid phase separation (deactivation) and ubiquitin-mediated 71 degradation under warm temperature conditions (Jung et al., 2020; Zhang et 72 al., 2021b; Zhang et al., 2021c), both of which release the inhibitory effects of 73 74 ELF3 on PIF4 (Nomoto et al., 2012; Box et al., 2015; Nieto et al., 2015; Jung et al., 2020; Silva et al., 2020). PIF7 was recently reported as a new type of 75 76 thermosensor in Arabidopsis (Chung et al., 2020). Indeed, the secondary 77 structure of PIF7 RNA in the 5' untranslated region (5' UTR) undergoes a 78 conformational change under warm temperature conditions, which leads to enhanced translation of PIF7 (Chung et al., 2020). Both PIF7 and PIF4 are 79 80 basic helix-loop-helix (bHLH) transcription factors that recognize G-box (CACGTG)-containing cis-elements and regulate the transcription of 81

downstream genes involved in auxin biosynthesis and signaling to promote
thermoresponsive hypocotyl growth (Gray et al., 1998; Franklin et al., 2011;
Sun et al., 2012).

85 ELF3, together with ELF4 and LUX ARRHYTHMO (LUX), assemble into the evening complex (EC) to regulate the circadian clock (Thines and Harmon, 86 87 2010; Huang and Nusinow, 2016). LUX is a MYB domain transcription factor that binds to DNA with high affinity. However, the LUX-ELF3 complex has 88 relatively poor DNA binding activity, but adding ELF4 to this complex restores 89 its DNA-binding activity in in vitro DNA binding assays (Silva et al., 2020). 90 Similarly, the complete EC strongly binds to DNA at 4°C and weakly binds to 91 DNA at 27°C in vitro. Adding an excess of ELF4 restores strong DNA binding 92 for the EC, even at 27°C (Silva et al., 2020), augesting that ELF4 is a key 93 modulator of thermosensitive EC activity. However, how ELF4 functions in 94 thermomorphogenesis in plants has not yet been reported. 95

The circadian clock consists of a series of repressors and activators that 96 form interconnected feedback loops (Zhang et al., 2021a). Besides the EC 97 repressor, other transcription factors from the MYB family are also key 98 99 components of the circadian clock. In particular, CIRCADIAN CLOCK ASSOCIATED 1 (CCA1) and LATE ELONGATED HYPOCOTYL (LHY) are 100 transcriptional repressors belonging to a small MYB subfamily, which also 101 102 includes eight REVEILLE (RVE) transcription factors (Rawat et al., 2009). 103 Among the RVEs, RVE4/6/8 were shown to be activators of gene expression 104 that act antagonistically with CCA1/LHY within the plant oscillator network to provide rhythmic robustness across environmental conditions (Xie et al., 2014; 105 106 Shalit-Kaneh et al., 2018). RVE7, also known as EARLY-PHYTOCHROME-RESPONSIVE 1 (EPR1), was previously reported to primarily function as a 107 circadian output rather than a circadian regulator at ambient temperature 108 conditions (23°C) (Kuno et al., 2003). 109

In the current study, we uncovered the essential role of warm-induced RVE7 in thermomorphogenesis. We demonstrate that RVE7 represses the expression of the circadian clock gene *ELF4* and fine-tunes hypocotyl growth under warm temperature conditions. Thus, RVE7 is not only an output factor, as previously described, but it is also an important modulator of the circadian clock under specific thermal conditions.

116

117 **Results** 

### 118 **RVE7** promotes thermoresponsive hypocotyl elongation in Arabidopsis

Similar to CCA1/LHY, the expression of RVE1/2/3/4/8, but not RVE5/6, is 119 regulated by the circadian clock in seedlings, with an expression peak occurring 120 near subjective dawn at ambient temperature (Rawat et al., 2011). We 121 examined the expression levels of these genes under ambient (22°C) and warm 122 (29°C) temperature conditions. At ZT24 (zeitgeber time: 24 h, or dawn), the 123 expression of CCA1/LHY/RVE1/3/4/8 decreased while that of RVE2 increased 124 at 29°C (Figure S1). The expression of RVE5/6 decreased slightly at ZT16 and 125 ZT24 but increased slightly at 2732 at 29°C compared to at 22°C (Figure S1). 126 By contrast, the expression of RVE7 increased at ZT16, ZT24, and ZT32 under 127 warm temperature conditions (Figure 1A). These differences in RVE7 transcript 128 levels under warm conditions prompted us to focus on this gene. 129

130 Since heat stress elements (HSEs; 5'-AGAAnnTTCT-3') are present in the 131 upstream sequences of RVE7, we measured the expression of RVE7 at ZT24 132 in a guadruple knockout (gk) mutant of Arabidopsis HEAT SHOCK FACTOR A1 (HSFA1) genes (HSFA1A, HSFA1B, HSFA1C, and HSFA1D). RVE7 133 134 expression did not increase in *hsfa1qk* seedlings at 29°C as it did in the wild type (WT; Figure 1B). Therefore, the induction of *RVE7* expression by warm 135 temperatures is dependent on these HSFA1s. We then examined RVE7 protein 136 accumulation under warm temperature conditions in seedlings overexpressing 137

RVE7-MYC driven by the constitutive cauliflower mosaic virus (CaMV) 35S promoter (Figure S2A) via immunoblot analysis. RVE7-MYC protein levels in these seedlings were higher at 29°C than at 22°C (Figure 1C), suggesting that RVE7-MYC might be degraded at 22°C. Indeed, RVE7-MYC was stabilized at 22°C when MG132, a potent 26S proteasome inhibitor, was added to the assays (Figure S3). We concluded that both *RVE7* transcript levels and RVE7 protein stability are regulated by warm temperatures.

To investigate the role of RVE7 in warm temperature-mediated growth, we 145 generated two independent alleles (rve7-11 and rve7-12) via clustered regularly 146 interspaced short palindromic repeats (CRISPR)/CRISPR-associated nuclease 147 9 (Cas9)-mediated gene editing (Figure S4). The hypocotyl lengths of both the 148 rve7-11 and rve7-12 mutants were similar to that of WT seedlings at 22°C. 149 However, the hypocotyls of *rve7-11* and *rve7-12* seedlings were significantly 150 shorter than those in WT at 29°C (R < 0.05, Figure 1D and F). We also 151 generated RVE7 overexpression (ines (Figure S2A) and measured their 152 hypocotyls. Consistent with the notion that RVE7 promotes hypocotyl growth at 153 warm temperatures, the RVET overexpression seedlings (RVE7ox-1 and 154 *RVE7ox-2* lines) were about 1.5-fold taller than WT seedlings at 29°C, but not 155 at 22°C (Figure 1E and G). Taken together, these results demonstrate that 156 RVE7 is a positive regulator of thermomorphogenesis that is important for 157 158 hypocotyl growth under warm conditions.

### 159 **RVE7 functions upstream of PIF4 during thermomorphogenesis**

The bHLH transcription factor PIF4 is a central regulator of seedling and plant morphogenesis (Koini et al., 2009; Quint et al., 2016). To analyze the genetic relationship between *RVE7* and *PIF4*, we generated the *rve7-11 pif4-101* double mutant and performed an epistatic analysis. Similar to the *pif4-101* single mutant, the hypocotyls of *rve7-11 pif4-101* seedlings did not elongate at 29°C relative to seedlings grown at 22°C (Figure 2A and C). Thus, *PIF4* is 166 epistatic to RVE7 during thermomorphogenesis. To examine whether the effect 167 of RVE7 in promoting thermomorphogenesis depends on PIF4, we overexpressed RVE7 in both the WT (RVE7ox) and PIF4 mutant backgrounds 168 169 (*pif4-101 RVE7ox*) (Figure S2B) and measured hypocotyl length. Unlike the *RVE7* overexpression seedlings in the WT background, the hypocotyl length of 170 171 pif4-101 RVE7ox seedlings was similar to that in WT at 29°C (Figure 2B and D). Thus, the function of RVE7 in controlling thermoresponsive hypocotyl 172 growth is largely dependent on PIF4. 173

To investigate how RVE7 affects PIF4 activity under warm temperature 174 conditions, we performed reverse transcription quantitative PCR (RT-qPCR) 175 and immunoblot analysis of WT, rve7-11, and RVE70x-1 seedlings. Compared 176 to the WT, the expression of *PIF4* was higher in *RVE7ox-1* seedlings but lower 177 in rve7-11 seedlings at 29°C, whereas such differences were modest at 22°C 178 (Figure 3A-B). In agreement with these results, the accumulation of 179 endogenous PIF4 protein decreased in rve7-11 seedlings and increased in 180 RVE7ox-1 seedlings only at 29°G (Figure 2E-F). 181

We also measured the expression levels of genes that function 182 downstream of PIF4 (Wang et al., 2018). In agreement with the accumulation 183 of PIF4 at 29°C, the transcript levels of At1g73120 (encoding an F-box protein), 184 XYLOGLUCAN ENDOTRANSGLYCOSYLASE 7 (XTR7, At4g14130), (IAA19, 185 186 At3g15540), and YUCCA 8 (YUC8, At4g28720) were higher in RVE7ox-1 seedlings at 29°C compared to WT seedlings (Figure 3C, E, G, I). By contrast, 187 188 At1g73120 and XTR7 transcript levels were lower in rve7-11 seedlings at this temperature (Figure 3D, F). We observed little effect on IAA19 or YUC8 189 190 transcript levels in *rve7-11* seedlings (Figure 3H, J), likely due to the functional redundancy of PIF4 with other regulators such as PIF5/7 (Koini et al., 2009; 191 192 Fiorucci et al., 2020). The differences in the expression levels of the abovementioned genes among WT, rve7-11, and RVE7ox-1 seedlings were 193

modest at 22°C (Figure 3C-J). Taken together, these results support the notion
that RVE7 functions upstream of PIF4 in thermomorphogenesis to control PIF4
accumulation and the expression of its downstream target genes under warm
temperature conditions.

# 198 RVE7 regulates the expression of the circadian clock gene *ELF4* under 199 warm conditions

ELF3 inhibits PIF4 by both suppressing its accumulation via the EC (Nomoto et 200 al., 2012) and preventing PIF4 from activating its transcriptional targets 201 independently of the complete EC (Nieto et al., 2015). Therefore, we measured 202 the expression levels of clock genes, including the three EC genes, in WT, rve7-203 11, and RVE7ox-1 seedlings at both 22°C and 29°C At 29°C, the expression 204 of ELF4 decreased at ZT16, ZT20, and ZT24 in RVE7ox-1 seedlings but 205 increased at ZT16 and ZT24 in rve7-11 seedlings, relative to the WT (Figure 206 4A-B). However, the expression levels of *ELF4* in WT and *rve7-11* seedlings 207 were similar at 22°C, while the expression levels of ELF4 were lower in 208 RVE7ox-1 compared to WT seedings at 22°C (Figure 4A-B). By contrast, the 209 expression levels of both ELF3 and LUX were similar between WT, rve7-11, 210 and RVE7ox-1 seedlings at both 22°C and 29°C (Figure 4C-F). The expression 211 levels of ELF4 were lower in RVE7ox-1 seedlings at ZT16 and ZT20 at 22°C, 212 likely because RVE7 was constitutively overexpressed in these lines. Finally, 213 214 the expression of *ELF4* was anti-phase to the expression of *RVE7* and *PIF4* in WT seedlings, both at 22°C and 29°C (Figure S5). Together, these results 215 216 indicate that RVE7 regulates ELF4 expression under warm temperature 217 conditions.

# 218 **RVE7 directly binds to the evening element (EE) in the ELF4 promoter and**

# 219 inhibits transcription

To explore how RVE7 regulates gene expression, we performed an effectorreporter assay with the *ELF4* promoter region (Figure 5A). RVE7 exhibited a

similar repressor activity as CCA1 in this assay (Figure 5B). We then performed 222 223 electrophoretic mobility shift assays (EMSAs) using recombinant purified 224 maltose binding protein (MBP)-RVE7 and the biotin-labeled clock gene-225 associated *cis*-element Evening Element (EE) derived from the *ELF4* promoter. When MBP-RVE7 was incubated with biotin-labeled EE (5'-AAATATCT-3'), we 226 227 observed a shift in mobility for the labeled probe (Figure 5C). Adding nonlabeled cold probes competed with this binding, while adding the mutated form 228 (5'-AAATCGAG-3') as a cold probe did not (Figure 5C), indicating that the 229 binding of MBP-RVE7 to the EE is sequence specific. These results 230 demonstrate that RVE7 specifically binds to the ELF4 promoter via the EE. 231

To examine the in vivo binding of RVE7 to the ELF4 promoter, we 232 performed chromatin immunoprecipitation qPCR (ChIP-qPCR) using RVE7-233 MYC overexpression lines grown at both 22°C and 29°C. After the RVE7-MYC 234 fusion protein was precipitated, we successfully amplified the ELF4 genomic 235 sequence (-364 bp to -170 bp relative to the TSS [transcription start site]) by 236 PCR (Figure 5D). Thus, RVE7 binds to the ELF4 promoter in planta. 237 238 Furthermore, warm temperatures enhanced the occupancy of RVE7 at ELF4 (Figure 5D). Since CCA1 and LHY were previously shown to associate with the 239 PIF4 promoter (Sun et al., 2019), we also examined the possible in vivo binding 240 of RVE7-MYC to the PIF4 promoter. We detected a slight enrichment of RVE7-241 242 MYC at the PIF4 promoter region (-577 to -415 bp relative to the TSS) at 29°C (Figure S6). Therefore, RVE7 directly inhibits the expression of *ELF4* by binding 243 244 to the EE *cis*-element in its promoter at warm temperatures.

# **Overexpressing ELF4 alleviates the inhibitory effect of RVE7 on hypocotyl**

246 growth under warm temperature conditions

To explore the genetic relationship between *RVE7* and *ELF4*, we generated the *rve7-11 elf4-209* double mutant and performed a phenotypic analysis under warm temperature conditions. *elf4-209* seedlings (Kolmos et al., 2009) had long

hypocotyls at both 22°C and 29°C, while rve7-11 seedlings had short 250 251 hypocotyls at 29°C (Figure 6A-B). By contrast, the hypocotyl length of rve7-11 252 elf4-209 seedlings was similar to that of elf4-209 seedlings at both 22°C and 253 29°C (Figure 6A-B). Thus, *ELF4* is epistatic to *RVE7*. We also generated lines overexpressing both RVE7 and ELF4 (Figure S2C) and determined that 254 overexpressing ELF4 partially suppresses the long hypocotyl phenotype 255 caused by RVE7 overexpression at warm temperatures (Figure 6C-D). 256 Seedlings overexpressing RVE7 had long hypocotyls at 29°C (Figure 1E, G). 257 We crossed the RVE7ox-3 overexpression line with the elf4-209 mutant and 258 performed a phenotypic analysis. The hypocotyl length of elf4-209 RVE7ox-3 259 seedlings was similar to that of elf4-209 seedlings at both 22°C and 29°C 260 (Figure 6E-F). Finally, we measured PIF4 abundance in RVE7 and ELF4 261 under warm temperature overexpression lines conditions. 262 double Overexpressing ELF4 prevented PIF4 accumulation, while overexpressing 263 RVE7 had the opposite effect. However, overexpressing ELF4 suppressed 264 PIF4 accumulation in RVE7 overexpression lines at 29°C (Figure 6G-H). These 265 266 results confirm the notion that RVE7 regulates hypocotyl growth by inhibiting the expression of clock genes such as ELF4 under warm temperature 267 conditions. 268

# 269 RVE7 functions redundantly with CCA1/LHY in controlling hypocotyl 270 growth under warm temperature conditions

*CCA1* and *LHY* play partially redundant functions in maintaining circadian rhythms and controlling temperature compensation in Arabidopsis (Mizoguchi et al., 2002; Salome et al., 2010). CCA1 and LHY play negative roles in lightinduced *ELF4* expression (Kikis et al., 2005), and CCA1 represses *ELF3* expression by associating with its promoter. ELF3 acts downstream of CCA1 to mediate the repression of *PIF4* and *PIF5* to control hypocotyl elongation under ambient temperature conditions (Lu et al., 2012). Therefore, we

278 investigated the possible functional redundancy between RVE7 and CCA1/LHY. 279 The hypocotyls of the cca1 lhy double mutant are shorter than those of WT 280 seedlings when grown at 22°C under red-light conditions (Yamashino et al., 281 2008). However, under white light conditions at 20°C, the difference in hypocotyl length between WT and cca1 lhy plants is marginal (Sun et al., 2019). 282 283 We crossed rve7-11 to the cca1-1 lhy-20 double mutant (Marshall et al., 2016) and generated the rve7-11 cca1-1 and rve7-11 lhy-20 double mutants, as well 284 as the rve7-11 cca1-1 lhy-20 triple mutant, and measured their hypocotyl 285 lengths at both 22°C and 29°C. The hypocotyl lengths of the cca1-1 and lhy-20 286 single mutants were similar to that of rve7-11 seedlings, and the rve7-11 cca1-287 1 and rve7-11 lhy-20 double mutants were indistinguishable from their 288 constituent single mutants (Figure 7A-D). However, the hypocotyls of the rve7-289 11 cca1-1 lhy-20 triple mutant were shorter than those of the rve7-11 single 290 mutant and the cca1-1 lhy-20 double mutant (Figure 7E-F). These results are 291 consistent with the notion that RVE7 plays redundant roles with CCA1/LHY in 292 controlling hypocotyl elongation under warm temperature conditions. 293

294

#### 295 Discussion

Circadian rhythms are generated in plants via the input of light and temperature 296 297 signals and are sustained by interconnected feedback loops (Creux and 298 Harmer, 2019). One output pathway of the circadian clock controls diurnal hypocotyl growth (Farre, 2012). Accumulating evidence indicates that the 299 300 circadian clock is tightly associated with the adaptive growth of hypocotyls in plants (Gil and Park, 2019). CCA1 and LHY are core components of the 301 302 circadian clock. The loss of CCA1 and LHY function confers early flowering at ambient temperatures (Mizoguchi et al., 2002) and reduces hypocotyl growth 303 at warm temperatures (Figure 7). The genetic inactivation of RVE1, a paralog 304 of CCA1/LHY, did not affect circadian rhythms, but did lead to a short-hypocotyl 305

306 phenotype at normal ambient growth temperature (Rawat et al., 2009). The 307 constitutive overexpression of RVE2 (also named CIRCADIAN 1 [CIR1]) leads to a shorter circadian period, delayed flowering, and long hypocotyls at ambient 308 309 temperature (Zhang et al., 2007). By contrast, the *rve4 rve6 rve8* triple mutant has a longer circadian period, delayed flowering, and a long hypocotyl 310 phenotype at normal growth temperature (Gray et al., 2017), suggesting that 311 RVE4/6/8 play a role opposite from that of CCA1/LHY/RVE1/RVE2 under 312 313 ambient temperature conditions.

In the current study, we demonstrated that RVE7 is functionally redundant with CCA1/LHY under warm temperatures and is involved in thermoresponsive hypocotyl growth (Figure 8). These findings expand our understanding of the functions of the RVE protein family and highlight the connection between circadian clock control with thermomorphogenesis.

Previous studies have indicated that RVE1, RVE2, and RVE7 are not 319 closely associated with the circadian oscillator, but these experiments have 320 been carried out under normal growth conditions (Kuno et al., 2003; Rawat et 321 al., 2009). In the current study, RVE7 showed transcriptional repression activity 322 and directly inhibited the expression of circadian clock genes, including ELF4, 323 at warm temperatures (Figure 4 and Figure 5). ELF4 is one of three 324 components of the EC (Huang and Nusinow, 2016). ELF4 accelerates the 325 326 nuclear localization of ELF3, which functions as a scaffolding protein to bring ELF4 together with LUX, a MYB domain transcription factor that directly binds 327 328 to DNA (Nusinow et al., 2011; Herrero et al., 2012; Silva et al., 2020). The EC inhibits the expression of PIF4 and PIF5, which is suppressed at dawn; 329 330 therefore, elevated levels of PIF4 and/or PIF5 promote gene expression associated with hypocotyl growth (Nomoto et al., 2012). ELF3 also inhibits the 331 activity of PIF4 independently of the EC (Nieto et al., 2015), which is released 332

by warm temperatures (Jung et al., 2020; Zhang et al., 2021b; Zhang et al.,2021c).

The expression of *ELF4* was not highly responsive to 29°C treatment in 335 336 WT seedlings, but it was altered in both *RVE7* overexpression lines and *RVE7* mutant seedlings (Figure 4), suggesting that other unknown regulators function 337 338 in an opposite manner to RVE7 to maintain ELF4 expression at warm temperatures in the WT. When this balance is disrupted due to reduced or 339 enhanced levels of RVE7 transcript levels under warm conditions, the 340 expression levels of *ELF4* and other downstream genes are likewise altered. 341 leading to the phenotypes observed in the current study. 342

Interestingly, LUX transcript accumulation was fully responsive to 29°C 343 conditions in various RVE7 genotypes as in WD seedlings. This observation 344 supports the notion that RVE7 and other factors that regulate the response to 345 warm temperatures are required to counteract the upregulation of LUX under 346 warm conditions. The EC has previously been shown to be crucial for this type 347 of autoregulation, whereby ELF3 is subjected to protein degradation, leading to 348 reduced EC activity under warm conditions (Ding et al., 2018; Zhang et al., 349 2021b; Zhang et al., 2021c). The current findings support the notion that RVE7 350 is essential for regulating ELF4 expression under warm temperature conditions. 351 352 Thus, in addition to ELF3 levels, the regulation of ELF4 levels is also essential 353 for thermomorphogenic growth in plants.

We showed that *RVE7* transcript levels increase as RVE7 protein levels increased at 29°C (Figure 1A-B), supporting the role of RVE7 in plant responses to warm temperature conditions. The expression levels of *ELF4* were reduced in *RVE7ox-1* seedlings, which is consistent with the increased PIF4 accumulation and the increased expression of *PIF4* downstream genes in this line (Figure 2 and Figure 3). In addition, overexpressing *ELF4* substantially suppressed the hypocotyl phenotype of *RVE7ox-1* seedlings (Figure 6).

361 Therefore, the regulation of *ELF4* expression by RVE7 is important for 362 thermoresponsive hypocotyl growth.

RVE7 regulates ELF4 expression at dusk (ZT16) under warm conditions, 363 364 as *ELF4* was expressed at higher levels at this time point in *rve7-11* seedlings than in the WT at 29°C (Figure 4B). The EC has previously been shown to 365 366 inhibit PIF4 activity at both the transcriptional and posttranslational levels (Nomoto et al., 2012; Zhang et al., 2021a). Indeed, the protein abundance of 367 PIF4 also decreased at dusk in rve7-11 seedlings at 29°C relative to the WT 368 (Figure 2E-F). We cannot exclude the possibility that RVE7 inhibits PIF4 369 expression through other mechanisms under warm temperatures, as the 370 expression of PIF4 and its downstream genes was also reduced in rve7-11 371 seedlings at 29°C at ZT24, when ELF4 expression in this mutant showed little 372 change from the WT (Figure 3 and Figure 4). Since RVE7 directly binds to the 373 EE *cis*-elements in its target promoters (Figure 5C), and the EE is present in 374 the promoters of many circadian clock genes (Nagel et al., 2015), besides ELF4, 375 RVE7 might also regulate *PIF4* expression via other clock components. 376

377

#### 378 Conclusion

In summary, we propose a model describing the positive role of RVE7 in thermomorphogenesis (Figure 8). According to this model, RVE7 represses the expression of *ELF4*, encoding an important component of the EC, to negatively regulate PIF4 levels during thermoresponsive hypocotyl growth.

383

#### 384 Materials and Methods

#### 385 Plant materials and hypocotyl length measurements

All Arabidopsis (*Arabidopsis thaliana*) genotypes used in this study were in the Columbia-0 (Col-0) background. The *cca1-1*, *elf4-209*, *hsfa1qk*, *lhy-20*, and *pif4-101* lines were described previously (Kolmos et al., 2009; Zhang et al.,

2013; Ding et al., 2018; Han et al., 2020). The rve7 single mutants were 389 390 generated using the CRISPR/Cas9 system (Yan et al., 2015). Two mutant alleles were selected for analysis: rve7-11, with a 16-bp deletion in the coding 391 392 sequence; and *rve7-12*, with a 1-bp insertion in the coding sequence (Figure S4). Both mutations lead to a frame shift and premature termination of 393 394 translation (Figure S4). To produce the overexpression lines, the coding sequences of RVE7 and ELF4 were amplified and inserted into pSKM36 or 395 396 pCAMBIA1306, respectively. These constructs were subsequently transformed into Agrobacterium (Agrobacterium tumefaciens) strain GV3101 via the freeze-397 thaw method and introduced into plants via the floral-dip method (Clough and 398 Bent, 1998). Higher-order mutants were generated by genetic crossing, as 399 400 mentioned in the text.

Seeds were surface sterilized for 15 min in 0.01% (w/v) sodium 401 hypochlorite and washed four times with sterile water. The seeds were sown 402 on half-strength Murashige and Skoog (MS) medium with vitamins (containing 403 1.2% [w/v] sucrose and 0.8% [w/v] agar, pH 5.7) and stratified at 4°C for 2 days, 404 405 after which they were transferred to a standard plant incubator at 22°C under a 16-h-light/8-h-dark photoperiod (long-day conditions) and 60% relative humidity. 406 For phenotypic assays seedlings were grown at 22°C for 3 days and 407 transferred to 29°C or maintained at 22°C for 4 days. To measure hypocotyl 408 409 length, the seedlings were photographed, and the hypocotyl lengths of the 410 seedlings were measured using ImageJ software (Zhang et al., 2021b; Zhang 411 et al., 2021c). All primers used in this study are listed in Table S1.

412 **RNA extraction and RT-qPCR** 

Five- or six-day-old seedlings grown at 22°C were transferred to 29°C at ZTO, while the control seedlings were maintained at 22°C. The seedlings were harvested at the indicated times and immediately frozen in liquid nitrogen for gene expression analysis. For comparisons between *RVE7ox-1* seedlings and 417 WT seedlings or between *rve7-11* seedlings and WT seedlings, the same batch 418 of WT seedlings was used for the control. Total RNA was extracted from the samples using an RNA Prep Pure Plant kit (Tiangen, Beijing. China). For 419 420 reverse transcription, 2 µg of RNA and oligo (dT) primers were used to 421 synthesize first-strand cDNA in a 20-µL reaction using M-MLV reverse 422 transcriptase (TaKaRa, Dalian, China). The resulting cDNAs were used for PCR or qPCR analysis. qPCR was performed using SuperReal PreMix Color 423 (Tiangen, Beijing, China) with a CFX96 real-time system (Bio-Rad, CA, USA) 424 with the gene-specific primers listed in Table S1. 425

#### 426 ChIP-qPCR

The ChIP assay was performed using an integrated method with a Chelex 427 resin-based ChIP procedure and protein A agarose beads (Millipore, CA, USA) 428 using an anti-myc antibody (Abmart, Shanghai, China). RVE7-MYC 429 overexpression lines were grown for 13 days and transferred to 29°C or 430 maintained at 22°C for the indicated times. The samples were fixed in 1% (w/v) 431 formaldehyde for 2×10 min under a vacuum, and fixation was stopped by 432 433 adding 0.15 M glycine to a final concentration of 0.125 M. The materials were then frozen in liquid nitrogen) After sonication in 0.8% (w/v) SDS buffer, protein 434 A-agarose beads (Millipore, CA, USA) and an anti-MYC antibody were used to 435 precipitate the DNA; IgG served as a serum control. The purified DNA was 436 437 quantified by qPCR. All primers used for qPCR are listed in Table S1.

#### 438 Immunoblot analysis

To analyze protein abundance, total proteins were extracted from the samples in extraction buffer (125 mM Tris-HCI [pH 8.0], 375 mM NaCI, 2.5 mM EDTA, 1% [w/v] SDS and 1% [w/v] beta-mercaptoethanol), and the protein concentrations were determined using a bicinchoninic acid assay (BCA) protein assay kit (Solarbio, Shanghai, China). The proteins were separated by 10% (w/v) SDS-PAGE and analyzed by immunoblotting using anti-MYC, anti-FLAG

(Abmart, Shanghai, China), anti-tubulin (Sigma, CA, USA), or anti-PIF4
(Abiocode, Shanghai, China) antibodies. The blots were scanned, and the
densitometry signal intensity of each band was quantified using ImageJ
software. The results are from the analyses of three immunoblots.

#### 449 Electrophoretic Mobility Shift Assay (EMSA)

450 The coding sequence of RVE7 was subcloned into pETMAL-H to produce and purify the recombinant MBP-RVE7 fusion protein according to standard 451 protocols (Zhang et al., 2021c). The DNA (-264 to -301 bp relative to the TSS) 452 containing the EE (5'-AAATATCT-3') derived from the ELF4 promoter was 453 454 synthesized and biotinylated using a biotin 3'-end DNA Labeling Kit (Thermo Fisher Scientific, CA, USA). A mutated form of the EE (5-AAATCGAG-3') was 455 used for the competition experiment. EMSA was performed using a LightShift 456 Chemiluminescent EMSA Kit (Thermo Fisher Scientific, CA, USA) according to 457 the manufacturer's protocols. Briefly, each binding reaction (20 mM HEPES, 458 pH 7.2, 80 mM KCl, 0.1 mM EDTA, 10% [v/v] glycerol, 2.5 mM DTT, 0.07 mg/ 459 mL BSA, 8 ng/mL poly dl-dC) was incubated for 20 min at room temperature, 460 461 and the reaction mixtures were resolved by electrophoresis through a 5% (w/v)non-denaturing polyacrylamide gel. After transferring to a nylon membrane, the 462 membrane was crosslinked under UV light and examined with a 463 Chemiluminescent Nucleic Acid Detection Module (Thermo Fisher Scientific, 464 465 CA, USA).

#### 466 Effector-reporter assay

The *ELF4* promoter sequence (-1,150 to +3 bp relative to the TSS) was PCR amplified and cloned into pGreen0800-II upstream of the firefly luciferase gene but downstream of the CaMV promoter to generate the reporter vector; the Renilla luciferase gene driven by the 35S promoter served as an internal control. The coding sequence of *RVE7* or *CCA1* was inserted into the pSKM36 vector to generate the respective effector construct. Different combinations of

473 constructs were transiently infiltrated in *Nicotiana benthamiana* leaves via 474 Agrobacterium (strain GV3101)-mediated infiltration. Three days after 475 infiltration, luciferase activity was measured with a Dual-luciferase Reporter 476 Assay kit (Promega, CA, USA). All primers are listed in Table S1.

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484 1 Ihy-20 and hsfa1qk mutant seeds, respectivel

485

# 486 Author contributions

487 J.X.L. and Y.Y.T. designed the experiments; Y.Y.T., W.L., M. J. W, and J.Y. L.

performed the experiments; J.X.L. and Y.Y.T. analyzed the data; J.X.L. and

489 S.J.D wrote the paper.

490

# 491 Declaration of interests

492 The authors declare no competing interests.

493

#### 494 Data availability statement

495 The data that support the findings of this study are available in the

496 supplementary materials of this article.

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#### 681 Supplemental information

Additional Supporting Information may be found online in the SupportingInformation section at the end of the article:

Figure S1. Warm temperature–regulated *CCA1*, *LHY*, and *RVE* gene expression. Six-day-old wild-type (WT) seedlings grown at 22°C were maintained at 22°C or transferred to 29°C and sampled at the indicated time points for gene expression analysis. Relative gene expression is the expression level of the target gene normalized to that of *PP2A*. Data are means  $\pm$  standard error (SE, n = 3).

Figure S2. Validation of transgenic lines. Six-day-old wild-type (WT), *pif4-101*, and various transgenic overexpression lines grown at 22°C were maintained at 22°C or transferred to 29°C and sampled at ZT24 for RT-PCR analysis. The expression of *UBQ5* was used as an internal control.

Figure S3. Protein stability assay. Seven-day-old *RVE7ox-1* seedlings grown at 22°C were maintained at 22°C or transferred to 29°C for 16 h in the presence or absence of the 26S proteasome inhibitor MG132 and sampled for immunoblotting with anti-myc antibody. Tubulin served as a protein loading control.

Figure S4. Characterization of gene-edited *rve7* mutant plants. Alignment
of the partial coding sequences of *RVE7* and their deduced amino acid
sequences in the wild type (WT) and the *rve7* mutants (*rve7-11* and *rve7-12*).
The sgRNA sequences used for vector construction are shown in red. \*, stop
codon.

Figure S5. Expression patterns of *RVE7*, *ELF4*, and *PIF4*. Five-day-old wildtype (WT) seedlings grown at 22°C were maintained at 22°C or transferred to 29°C and sampled at different time points (ZT) for gene expression analysis. The expression level of each gene was normalized to that of *PP2A*. Data are means  $\pm$  SE (n = 3).

Figure S6. Binding of RVE7 to the PIF4 promoter. Thirteen-day-old 709 transgenic seedlings overexpressing RVE7-MYC grown at 22°C were 710 maintained at 22°C or transferred to 29°C for 16 h and sampled for ChIP-qPCR 711 712 using anti-MYC antibody. Relative enrichment of each sample was normalized to that the anti-GST sample (IgG control) at 16 h at 22°C, both of which were 713 714 normalized to the TA3 control. Data are means  $\pm$  SE (n = 3). Different letters above the bars indicate significant differences, as determined by post hoc test 715 (*P* < 0.05). 716

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719 **Table S1. Primers used in this study.** 

s used in this study.

#### 720 FIGURE LEGENDS

Figure 1. RVE7 is responsive to warm temperatures and positively 721 regulates thermomorphogenesis. A-B, Upregulation of RVE7 transcript 722 723 levels by warm temperatures. Six-day-old wild-type (WT) seedlings grown at 724 22°C were maintained at 22°C or transferred to 29°C at ZT0 and sampled at 725 the indicated time for gene expression analysis (A). The hsfa1a hsfa1b hsfa1c hsfa1d quadruple mutant (hsfa1qk) was also treated like the WT and sampled 726 at ZT24 (B). Relative gene expression is the expression level of RVE7 in each 727 sample normalized to that of PP2A. Data are means  $\pm$  SE (n = 3). C, 728 729 Accumulation of RVE7 under warm temperature conditions. Seven-day-old RVE7-MYC overexpression seedlings grown at 22°C were maintained at 22°C 730 or transferred to 29°C and sampled for immunobletting with anti-myc antibody. 731 Tubulin served as a protein loading control. **D-G**. Phenotypic analysis. 732 Seedlings of WT, RVE7 loss-of-function mutants (rve7-11 and rve7-12), and 733 RVE7 overexpression lines (RVE7ox-1 and RVE7ox-2) were grown at 22°C for 734 3 days and kept at 22°C or transferred to 29°C for 4 days, after which 735 representative seedlings were imaged (D-E) and their hypocotyl lengths 736 measured (F-G). *pif4-10* was used as a control. Data are means ± standard 737 deviation (SD, n = 24). Different lowercase letters indicate significant 738 differences, as determined by post hoc test (P < 0.05). Scale bars = 5 mm. 739

740 Figure 2. RVE7 functions upstream of PIF4 in thermomorphogenesis. A-D, Genetic analysis of the roles of RVE7 and PIF4 in thermoresponsive 741 742 hypocotyl growth. Seedlings of the WT, rve7-11, pif4-101, the rve7-11 pif4-101 743 double mutant, RVE7ox, and pif4-101 RVE7ox were grown at 22°C for 3 days 744 and kept at 22°C or transferred to 29°C for 4 days, after which representative 745 seedlings were imaged (A-B) and their hypocotyl lengths measured (C-D). Data are means  $\pm$  SD (n = 24). Scale bars = 5 mm. E-F, Accumulation of PIF4. 746 Seven-day-old WT, rve7-11, and RVE7ox-1 seedlings grown at 22°C were 747

maintained at 22°C or transferred to 29°C for 16 h and sampled for immunoblotting with anti-PIF4 antibody (E). Tubulin served as a protein loading control. The band intensities in three immunoblots were quantified (F). Data are means  $\pm$  SE (n = 3). Different lowercase letters indicate significant differences, as determined by post hoc test (P < 0.05).

753 Figure 3. RVE7 regulates the expression of PIF4 and its downstream genes under warm conditions. A-J, Expression of *PIF4* and its downstream 754 genes under ambient and warm temperature conditions. Five-day-old WT, 755 rve7-11, and RVE7ox-1 seedlings grown at 22°C were maintained at 22°C or 756 transferred to 29°C and sampled at three different time points (ZT) for 757 quantitative gene expression analysis. The expression level of each gene was 758 normalized to that of the WT at ZT16 at 22°C, which was normalized to that of 759 *PP2A*. Data are means  $\pm$  SE (n = 3). 760

Figure 4. RVE7 regulates EC gene expression under warm conditions. A-F, Expression of three EC genes under ambient and warm temperature conditions. Five-day-old WT, *rve7-11*, and *RVE7ox-1* seedlings grown at 22°C were maintained at 22°C or transferred to 29°C and sampled at three different time points (ZT) for quantitative gene expression analysis. The expression level of each gene was normalized to that of the WT at ZT16 at 22°C, which was normalized to that of *PP2A*. Data are means  $\pm$  SE (n = 3).

768 Figure 5. RVE7 directly inhibits the expression of ELF4. A-B, Transcriptional repression activity assay. RVE7-MYC, CCA1-MYC, or MYC 769 770 (vector control) driven by the 35S promoter was used as the effector, and the 771 firefly luciferase driven by the ELF4 promoter (pELF4) linked to the 35S 772 promoter was co-expressed as the reporter in effector-reporter assays. The activity of Renilla luciferase, whose encoding gene was constitutively 773 expressed, was used as an internal control. Relative luciferase activity is firefly 774 luciferase activity normalized to Renilla luciferase activity, which was then 775

normalized to the vector control. Data are means  $\pm$  SE (n = 3). C, Direct binding 776 777 of RVE7 to the EE. Recombinant MBP-RVE7 was incubated with biotin-labeled DNA containing the EE (5'-AAATATCT-3') derived from the ELF4 promoter, and 778 779 electrophoretic mobility shift assays (EMSAs) were performed. Non-labeled native or mutated (5'-AAATCGAG-3') cold probes were added to the reaction 780 781 for competition assays. **D**, Binding of RVE7 to the *ELF4* promoter in seedlings under two temperature conditions. Thirteen-day-old transgenic seedlings 782 overexpressing RVE7-MYC grown at 22°C were maintained at 22°C or 783 transferred to 29°C for 16 h and sampled for ChIP-gPCR using anti-MYC 784 antibody. The relative enrichment of *ELF4* DNA in each sample was normalized 785 to that in the anti-GST sample (IgG control) at 22°C, both of which were 786 normalized to that of the TA3 control. Data are means  $\pm$  SE (n = 3). Different 787 lowercase letters indicate significant differences, as determined by post hoc 788 test (P < 0.05). 789

Figure 6. Overexpressing *ELF4* suppresses the long hypocotyl phenotype 790 caused by RVE7 overexpression under warm temperature conditions. A-791 F, Genetic analysis of the relevant RVE7 and ELF4 in thermomorphogenesis. 792 Seedlings of WT, rve7-11, elf4-209, rve7-11 elf4-209, RVE7ox-1 and ELF4 793 overexpression (ELF40x,1), RVE7 and ELF4 double overexpression (ELF40x-794 1 RVE7ox-1) lines, and elf4-209 RVE7ox-3 grown at 22°C for 3 days were kept 795 796 at 22°C or transferred to 29°C for 4 days, after which representative seedlings were imaged (A, C, E) and their hypocotyl lengths measured (B, D, F). Data are 797 798 means ± SD (n = 24). G-H, Accumulation of PIF4. Seven-day-old WT, ELF4ox-799 21, RVE7ox-1, and RVE7ox-1 ELF4ox-11 seedlings grown at 22°C were 800 maintained at 22°C or transferred to 29°C for 16 h and sampled for immunoblotting with anti-PIF4 antibody (G). Tubulin served as a protein loading 801 802 control. The band intensities in three immunoblots were quantified (H). Data are

803 means  $\pm$  SE (n = 3). Different lowercase letters indicate significant differences, 804 as determined by post hoc test (*P* < 0.05). Scale bars = 5 mm.

RVE7 functions redundantly CCA1/LHY 805 Figure 7. with in 806 thermomorphogenesis. A-F, Genetic analysis of the roles of RVE7 and CCA1/LHY in thermomorphogenesis. WT, rve7-11, cca1-1, lhv-20, rve7-11 807 808 cca1-1, rve7-11 lhy-20, and rve7-11 cca1-1 lhy-20 seedlings grown at 22°C for 3 days were kept at 22°C or transferred to 29°C for 4 days, after which 809 representative seedlings were imaged (A, C, E) and their hypocotyl lengths 810 measured (B, D, F). Data are means  $\pm$  SD (n = 24). Different lowercase letters 811 indicate significant differences, as determined by post hoc test (P < 0.05); scale 812 813 bars = 5 mm.

Figure 8. A simplified working model for the role of RVE7 in 814 thermoresponsive hypocotyl growth. The hypocotyl growth-promoting 815 bHLH transcription factor PIF4 is negatively regulated by the evening complex 816 (EC) consisting of ELF3, ELF4, and LUX. Under warm temperature conditions 817 (29°C), the MYB transcription factor RVE7 accumulates and reduces the 818 expression of *ELF4*, allowing *PH*<sup>2</sup>4 to reach a certain level in wild-type (WT) 819 seedlings. In RVE7 overexpression (RVE7ox-1) seedlings, higher RVE7 protein 820 abundance leads to tower ELF4 transcript levels and higher accumulation of 821 822 PIF4, thereby triggering higher expression of PIF4 downstream genes and 823 faster hypocotyl growth under warm temperature conditions. By contrast, in RVE7 mutant (rve7-11) seedlings, higher ELF4 expression levels lead to 824 825 greater repression of PIF4, resulting in shorter hypocotyls. The positive regulators of *ELF4* and *PIF4* expression are not depicted in the model. Positive 826 827 and negative regulatory activities are indicated by arrows and lines with bars, respectively. The thickness of the lines and the depth of color of the shapes 828 829 reflect the degree of regulation.

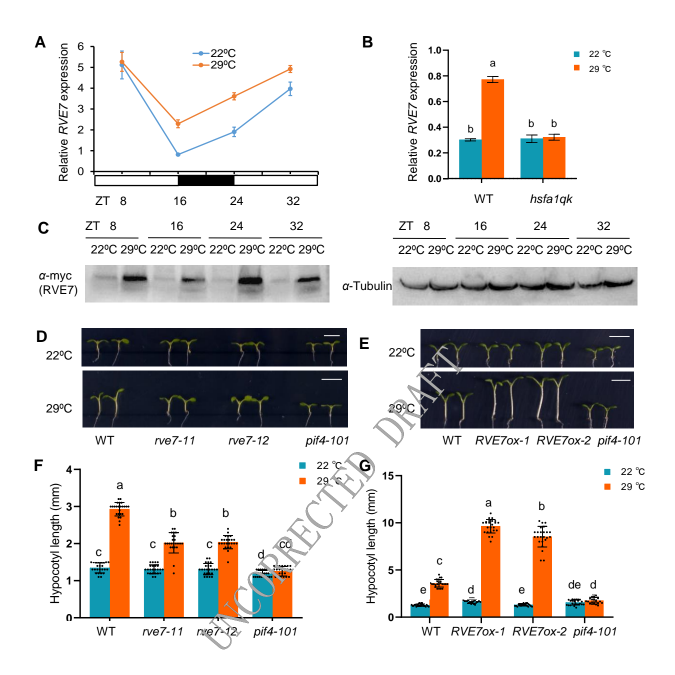
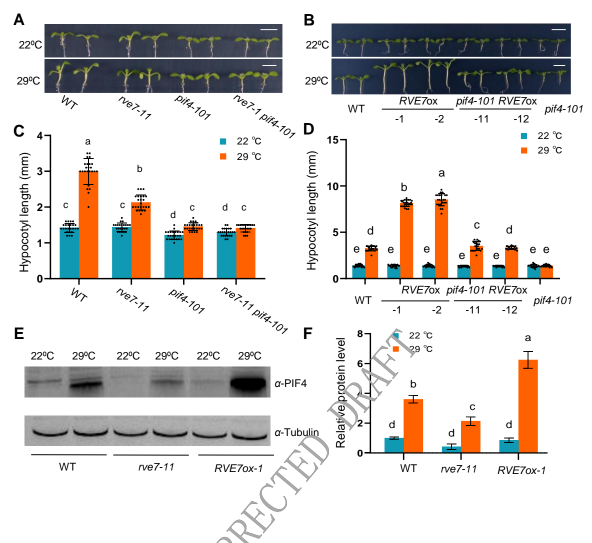


Figure 1. *RVE7* is responsive to warm temperatures and positively regulates thermomorphogenesis. A-B, Upregulation of *RVE7* transcript levels by warm temperatures. Six-day-old wild-type (WT) seedlings grown at 22° C were maintained at 22° C or transferred to 29° C at ZT0 and sampled at the indicated time for gene expression analysis (A). The *hsfa1a hsfa1b hsfa1c hsfa1d* quadruple mutant (*hsfa1qk*) was also treated like the WT and sampled at ZT 24 (B). Relative gene expression is the expression level of *RVE7* in each sample normalized to that of *PP2A*. Error Data are means  $\pm$  SE (n=3). **C**, Accumulation of RVE7 under warm temperature conditions. Seven-day-old *RVE7-MYC* overexpression seedlings grown at 22° C were maintained at 22° C or transferred to 29° C and sampled for immunoblotting with anti-myc antibody. Tubulin served as a protein loading control. **D-G**, Phenotypic analysis. Seedlings of WT, *RVE7* loss-of-function mutants (*rve7-11* and *rve7-12*), and *RVE7* overexpression lines (*RVE7ox-1*) were grown at 22° C for 3 days and kept at 22° C or transferred to 29° C for 4 days, after which representative seedlings were imaged (D-E) and their hypocotyl lengths measured (F-G). *pif4-101* was used as a control. Data are means  $\pm$  standard deviation (SD, n=24). Different lowercase letters indicate significant differences, as determined by post hoc test (*P* < 0.05). Scale bars = 5 mm.



**Figure 2. RVE7 functions upstream of PIF4 in thermomorphogenesis.** A-D, Genetic analysis of the roles of *RVE7* and *PIF4* in thermoresponsive hypocotyl growth. Seedlings of the WT, *rve7-11*, *pif4-101*, the *rve7-11 pif4-101* double mutant, *RVE7ox* and *pif4-101 RVE7ox* were grown at 22° C for 3 days and kept at 22° C or transferred to 29° C for 4 days, after which representative seedlings were imaged (A-B) and their hypocotyl lengths measured (C-D). Data are means  $\pm$  SD (n=24). Scale bars = 5 mm. **E-F**, Accumulation of PIF4. Seven-day-old WT, *rve7-11*, and *RVE7ox-1* seedlings grown at 22° C were maintained at 22° C or transferred to 29° C for 16 h and sampled for immunoblotting with anti-PIF4 antibody (E). Tubulin served as a protein loading control. The band intensities in three immunoblots were quantified (F). Data are means  $\pm$  SE (n = 3). Different lowercase letters indicate significant differences, as determined by post hoc test (P < 0.05).

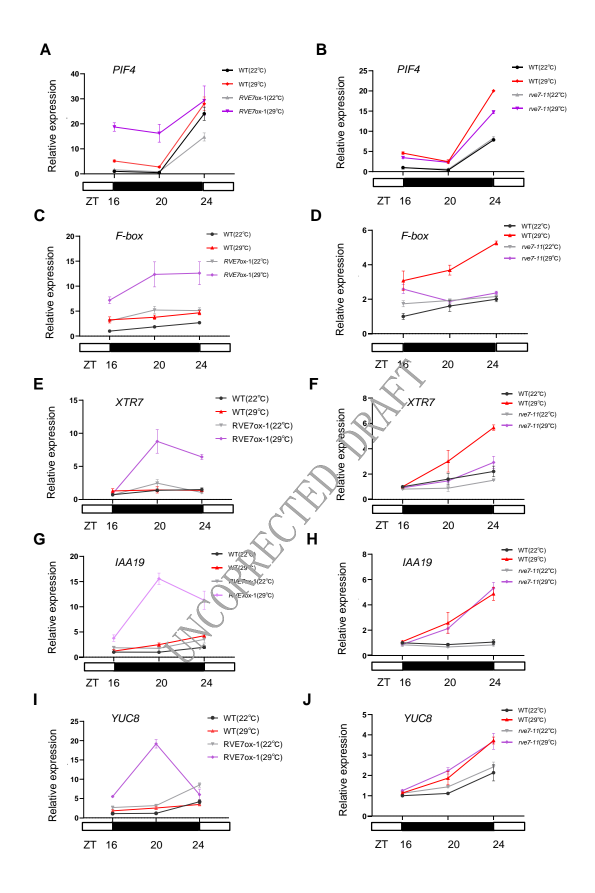
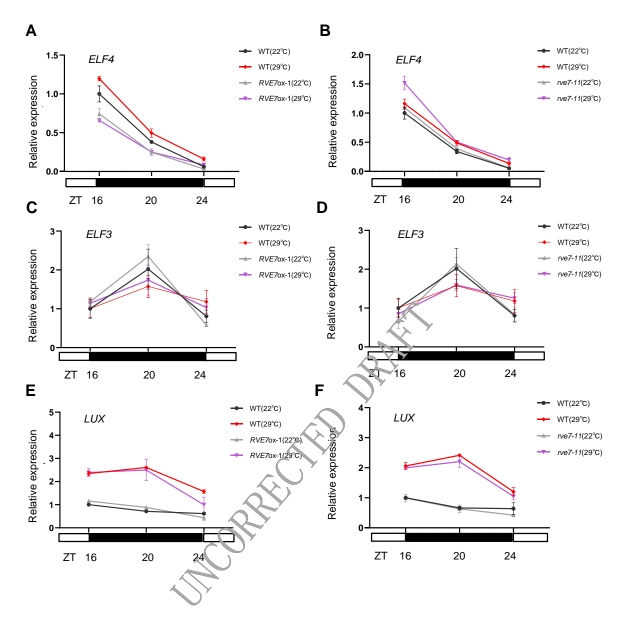
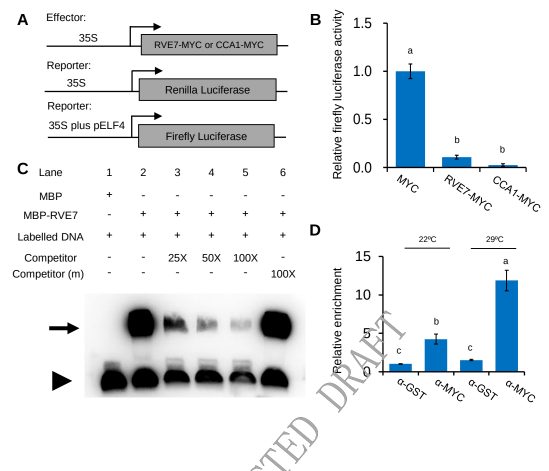


Figure 3. RVE7 regulates the expression of *PIF4* and its downstream genes under warm conditions. A-J, Expression of *PIF4* and its downstream genes under ambient and warm temperature conditions. Five-day-old WT, *rve7-11*, and *RVE7ox-1* seedlings grown at 22° C were maintained at 22° C or transferred to 29° C and sampled at three different time points (ZT) for quantitative gene expression analysis. The expression level of each gene was normalized to that of WT at ZT16 at 22° C, which was normalized to that of *PP2A*. Data are means  $\pm$  SE (n = 3).



**Figure 4. RVE7 regulates EC gene expression under warm conditions. A-F**, Expression of three EC genes under ambient and warm temperature conditions. Five-day-old WT, *rve7-11* and *RVE7ox-1* seedlings grown at  $22^{\circ}$  C were maintained at  $22^{\circ}$  C or transferred to  $29^{\circ}$  C and sampled at three different time points (ZT) for quantitative gene expression analysis. The expression level of each gene was normalized to that of WT at ZT16 at  $22^{\circ}$  C, which was normalized to that of *PP2A*. Data are means  $\pm$  SE (n = 3).



**Figure 5. RVE7 directly inhibits the expression of ELF4. A-B**, Transcriptional repression activity assay. RVE7-MYC, CCA1-MYC, or MYC (vector control) driven by the 35S promoter was used as the reporter in effector, and the firefly luciferase driven by the *ELF4* promoter (pELF4) linked to the 35S promoter was co-expressed as the reporter in effector-reporter assays. The activity of Renilla luciferase, whose encoding gene was constitutively expressed, was used as an internal control. Relative luciferase activity is firefly luciferase activity is firefly luciferase activity is firefly luciferase activity which was then normalized to the vector control. Data are means  $\pm$  SE (n = 3). **C**, Direct binding of RVE7 to the EE. Recombinant MEP-RVE7 was incubated with biotin-labeled DNA containing the EE (5'-AAATATCT-3') derived from the *ELF4* promoter and electrop oreic mobility shift assays (EMSAs) were performed. Non-labeled native or mutated (5'-AAATCGAG-3') cold probes were added to the reaction for competition assays. **D**, Binding of RVE7 to the *ELF4* promoter in seedlings under two temperature conditions. Thisteen-day-old transgenic seedlings overexpressing *RVE7-MYC* grown at 22° C were maintained at 22° C or transferred to 29° C for 16 h and sampled for ChIP-qPCR using anti-MYC antibody. The relative enrichment of *ELF4* DNA in each sample was normalized to that in the anti-GST sample (IgG control) at 22° C, both of which were normalized to that of the *TA3* control. Data are means  $\pm$  SE (n = 3). Different lowercase letters indicate significant differences, as determined by post hoc test (*P* < 0.05).

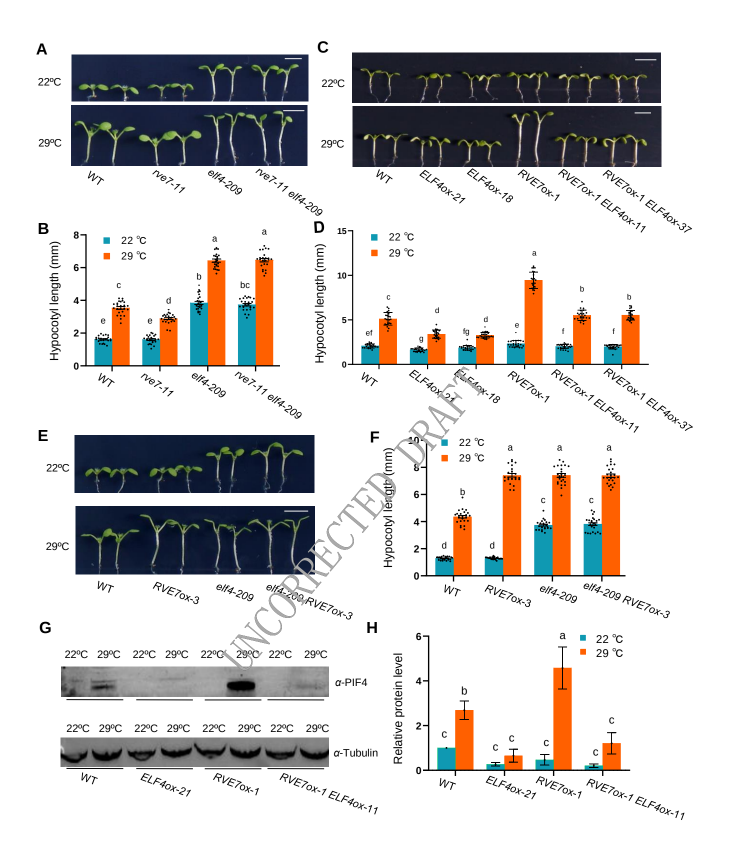
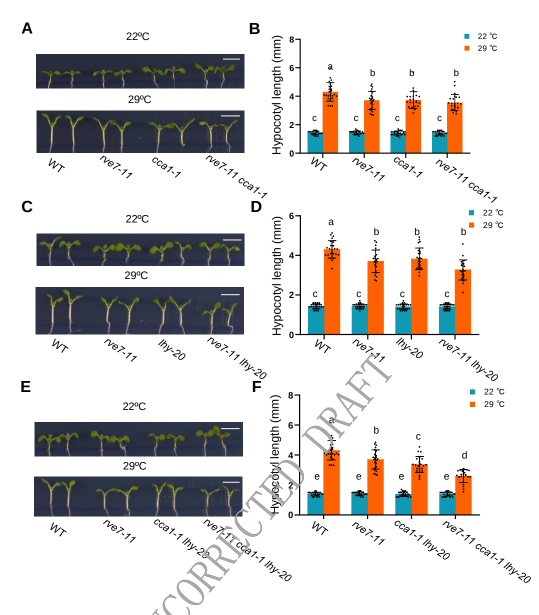
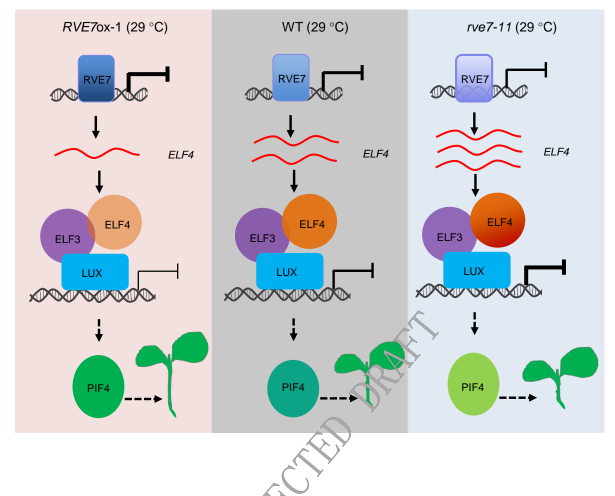


Figure 6. Overexpressing *ELF4* suppresses the long hypocotyl phenotype caused by *RVE7* overexpression under warm temperature conditions. A-F, Genetic analysis of the roles of *RVE7* and *ELF4* in thermomorphogenesis. Seedlings of WT, *rve7-11*, *elf4-209*, *rve7-11* elf4-209, *RVE7*ox-1 and *ELF4* overexpression (*ELF4*ox-1), *RVE7* and *ELF4* double overexpression (*ELF4*ox-1) *RVE7*ox-1) lines, and *elf4-209*, *RVE7*ox-3 grown at 22° C for 3 days were kept at 22° C or transferred to 29° C for 4 days, after which representative seedlings were imaged (A, C, E) and their hypocotyl lengths measured (B, D, F). Data are means  $\pm$  SD (n=24). G-H, Accumulation of PIF4. Seven-day-old WT, *ELF4ox-21*, *RVE7ox-1* and *RVE7ox-1 ELF4ox-11* seedlings grown at 22° C were maintained at 22° C or transferred to 29° C for 16 h and sampled for immunoblotting with anti-PIF4 antibody (G). Tubulin served as a protein loading control. The band intensities in three immunoblots were quantified (H). Data are means  $\pm$  *SE* (n = 3). Different lowercase letters indicate significant differences, as determined by post hoc test (*P* < 0.05). Scale bars = 5 mm.



**Figure 7. RVE7 functions redundantly with CCA1/LHY in thermomorphogenesis. A-F**, Genetic analysis of the roles of *RVE7* and *CCA1/LHY* in thermomorphogenesis. WT, *rve7-11, cca1-1, lhy-20, rve7-11 cca1-1, rve7-11 lhy-20,* and *rve7-11 cca1-1 lhy-20* seedlings grown at 22° C for three days were kept at 22° C or transferred to 29° C for 4 days, after which representative seedlings were imaged (A, C, E), and their hypocotyl lengths measured (B, D, F). Data are means  $\pm$  *SD* (n=24). Different lowercase letters indicate significant differences, as determined by post hoc test (*P* < 0.05); Scale bars = 5 mm.



**Figure 8. A simplified working model for the role of RVE7 in thermoresponsive hypocotyl growth.** The hypocotyl growthpromoting bHLH transcription factor PIF4 is negatively regulated by the evening complex (EC) consisting of ELF3, ELF4, and LUX. Under warm temperature conditions (29° C), the MYb transcription factor RVE7 accumulates and reduces the expression of *ELF4*, allowing PIF4 to reach a certain level in wild-type (W1) seedlings. In *RVE7* overexpression (*RVE70x*-1) seedlings, higher RVE7 protein abundance leads to lower *ELF4* transcript levels and higher accumulation of PIF4, thereby triggering higher expression of PIF4 downstream genes and faster hypocotyl growth under warm temperature conditions. By contrast, in *RVE7* mutant (*rve7-11*) seedlings, higher *ELF4* expression levels lead to greater repression of PIF4, resulting in shorter hypocotyls. The positive regulators of *ELF4* and *PIF4* expression are not depicted in the model. Positive and negative regulatory activities are indicated by arrows and lines with bars, respectively. The thickness of the lines and the depth of color of the shapes reflect the degree of regulation.