- 1 Short title: PHYs temperature perception and plastid metabolism
- 2 PHYTOCHROME-DEPENDENT TEMPERATURE PERCEPTION MODULATES
- 3 ISOPRENOID METABOLISM
- 4 Ricardo Bianchetti^{1‡}, Belen De Luca^{2‡}, Luis A de Haro^{2†}, Daniele Rosado^{1††}, Diego
- 5 Demarco¹, Mariana Conte³, Luisa Bermudez^{3,4}, Luciano Freschi¹, Alisdair R. Fernie⁵,
- 6 Louise V Michaelson⁶, Richard P Haslam⁶, Magdalena Rossi^{1‡}, Fernando Carrari^{2,4‡*}
- 7 Departamento de Botânica, Instituto de Biociências, Universidade de São Paulo, São
- 8 Paulo, Brasil.
- ²Instituto de Fisiología, Biología Molecular y Neurociencias (IFIBYNE-UBA-CONICET)
- 10 Ciudad Universitaria, C1428EHA Buenos Aires, Argentina.
- ³Instituto de Agrobiotecnología y Biología Molecular (IABIMO), CICVyA INTA;
- 12 CONICET, Argentina.
- ⁴Cátedra de Genética, Facultad de Agronomía, Universidad de Buenos Aires, Buenos Aires,
- 14 Argentina.
- ⁵Max Planck Institute of Molecular Plant Physiology, Wissenschaftspark Golm, Am
- 16 Mühlenberg 1, Potsdam-Golm, D-14476, Germany.
- ⁶Department of Plant Sciences, Rothamsted Research, Harpenden, Herts, AL5 2JQ, UK.
- †Current address: Department of Plant and Environmental Sciences, Weizmann Institute of
- 19 Science, Rehovot 76100, Israel
- 20 ††Current address: Cold Spring Harbor Laboratory, 1 Bungtown Road, Cold Spring Harbor,
- 21 New York 11724, USA
- * Corresponding Author.
- 23 Fernando Carrari
- 24 Phone: +54 11 4576-3386 / 3368 ext 238
- email: <u>fcarrari@fbmc.fcen.uba.ar</u>
- 26 ‡ These authors contributed equally to this work.
- 28 One-sentence summary: Phytochrome-mediated temperature perception compromises
- 29 plastidial development and function, impairing isoprenoid metabolism in tomato leaves and
- 30 fruits.

31

Author contributions

- RB performed most of the experiments and analyzed the data; BDL and LAH performed
- experiments and analyzed data; DR, DD and LB performed experiments; MC, LF, ARF,
- 35 LVM and RPH contributed to experimental design and provided technical assistance; RB,
- 36 MR, ARF and FC conceived the project, designed experiments and wrote the paper which
- was revised and approved by all authors. FC agrees to serve as the author responsible for
- 38 contact and ensures communication.

FUNDING

32

39

- 40 This work was supported by FAPESP (Fundação de Amparo à Pesquisa do Estado de São
- 41 Paulo, Grant Number #2016/01128-9); CNPq (Conselho Nacional de Desenvolvimento
- 42 Científico e Tecnológico, Grant Number 440976/2016-2); European Union Horizon 2020
- 43 Research and Innovation Programme (Grant Agreement Number 679796), and ANPCyT
- 44 Agencia Nacional de Promoción Científica y Tecnológica, Argentina, Grant Number 2014-
- 45 0984 to F.C.). R.P.H and L.V.M. were supported by the BBSRC (Biotechnology and
- 46 Biological Sciences Research Council, UK) through the Tailoring Plant Metabolism
- 47 Institute Strategic Grant (BBS/E/C/000I0420) supporting R.P.H and the UK-Brazil
- 48 Alliance for Sustainable Agriculture scheme (BBS/OS/NW/000001). R.B. and D.R. were
- 49 recipient of FAPESP fellowships (#2017/24354-7 and #2015/14658-3). B.D.L and L.A.de
- 50 H. are CONICET fellows. L.B. and F.C are members of CONICET. M.R. was a recipient
- of CNPq fellowship.

Abstract

53

54

55

56

59

63

65

66

67

68 69

73

Changes in environmental temperature influence many aspects of plant metabolism; however, the underlying regulatory mechanisms remain poorly understood. In addition to their role in light perception, phytochromes (PHYs) have been recently recognized as temperature sensors affecting plant growth. In particular, in Arabidopsis thaliana, high 57 temperature reversibly inactivates PHYB, reducing photomorphogenesis-dependent 58 responses. Here, we show the role of phytochrome-dependent temperature perception in modulating the accumulation of isoprenoid-derived compounds in tomato (Solanum 60 61 lycopersicum) leaves and fruits. The growth of tomato plants under contrasting temperature regimes revealed that high temperatures resulted in co-ordinated up-regulation of 62 chlorophyll catabolic genes, impairment of chloroplast biogenesis, and reduction of carotenoid synthesis in leaves in a PHYB1B2-dependent manner. Furthermore, by 64 assessing a triple phyAB1B2 mutant and fruit-specific PHYA- or PHYB2-silenced plants, we demonstrated that biosynthesis of the major tomato fruit carotenoid, lycopene, is sensitive to fruit-localized PHY-dependent temperature perception. The collected data provide compelling evidence concerning the impact of PHY-mediated temperature perception on plastid metabolism in both leaves and fruit, specifically on the accumulation of isoprenoidderived compounds. 70

- Keywords: carotenoid, chlorophyll, chloroplast, fleshy fruit, isoprenoids, phytochrome, 71
- temperature, tomato. 72

INTRODUCTION

74

Temperature cues regulate plant primary and secondary metabolism, affecting several 75 76 agronomically important traits in crop species (Suwa et al., 2010; Bita and Gerats, 2013; Zhao et al., 2017). It has been established that heat disrupts chloroplast integrity leading to 77 further deficiencies of plastid-associated metabolites and a subsequent decline in plant 78 performance (Yamori and von Caemmerer, 2009; Spicher et al., 2017). The ability to 79 perceive stress conditions allows plants to adapt their metabolism in order to minimize any 80 harmful effects on fitness (Saidi et al., 2011). 81 82 PHYTOCHROMES (PHYs) have been extensively described as light receptors. The biologically inactive PHY form (Pr) remains in the cytosol; however, once activated by red 83 light, the active form (Pfr) is translocated towards the nucleus where it assembles into 84 photobodies and triggers photomorphogenesis-associated responses (Rockwell et al., 2006). 85 The conversion of Pr to Pfr can be reversed by far-red light or darkness (Burgie and 86 Vierstra, 2014). PHYTOCHROME B (PHYB) has been associated with a quantitative trait 87 88 loci interval for thermoresponsive growth in Arabidopsis thaliana (Box et al., 2015). High temperature reduces the abundance of Pfr by a quick and spontaneous reversion to Pr in a 89 90 light-independent manner, consequently decreasing the size of nuclear bodies in a process termed thermoreversion (Legris et al., 2016). By contrast, lack of thermoreversion was 91 detected in the hyperactive phyB mutant resulting in the constitutive presence of 92 photobodies regardless of temperature condition (Huang et al., 2019). As such, the A. 93 thaliana phyB null mutant mimics the transcriptional profile and physiological parameters 94 95 of the wild-type counterpart grown under high temperature (Jung et al., 2016). In tomato (Solanum lycopersicum) PHYs belong to a multigenic family encompassing five 96 members: PHYA, PHYB1, PHYB2, PHYE, and PHYF (Alba et al., 2000b). We have 97 previously demonstrated that PHYA, PHYB1, and PHYB2 positively control the 98 biosynthesis of isoprenoid-derived compounds in tomato fruits in response to light 99 (Bianchetti et al., 2018; Gramegna et al., 2018). PHYs post-translationally downregulate a 100 group of helix-loop-helix proteins named PHYTOCHROME INTERACTING FACTORS 101 (PIFs) (Park et al., 2018). PIFs derive from a multigenic family and have undergone sub-102 and neofunctionalization at the mRNA level (Rosado et al., 2016). It has been shown that 103 SIPIF1a, SIPIF3, and SIPIF4 regulate carotenogenesis (Llorente et al., 2016), tocopherol 104

biosynthesis (Gramegna et al., 2018), and sugar metabolism (Rosado et al., 2019), 105 respectively, by (a) light-dependant mechanism(s). 106 Here, by analyzing the metabolic and transcriptional profile of wild-type and phy-mutant 107 tomato plants grown under contrasting temperature conditions, we showed that high 108 109 temperature results in the reduction of leaf chlorophyll (Chl) and carotenoid levels in a PHYB1/B2-mediated manner through the regulation of Chl degradation and carotenoid 110 111 biosynthetic genes, respectively. Furthermore, our data also demonstrate that high temperature or PHYAB1B2 impairment leads to the transcriptional downregulation of 112 carotenoid biosynthetic genes in fruits, resulting in reduced levels of lycopene. Data 113 114 obtained from fruit-specific PHYA- and PHYB2-silenced plants corroborated the role of fruit-localized PHYs in carotenoid accumulation through an intricate network in which 115 master ripening transcription factors participate as mediators of temperature perception in 116 tomato fruits. 117 118

RESULTS

120 121

122

123

124

125

126

127

128 129

130

131

132

133

134

135

136

137

High temperature affects plants growth phenotype in tomato

To investigate the role played by PHYs in response to high temperature, 20-day-old plants from wild type cv. MoneyMaker (MM), phyB1 and phyB2 single mutants, and the phyB1B2 double mutant were transferred to ambient-temperature (AT, 24°C/18°C) and hightemperature (HT, 30°C/24°C) growing conditions (Supplemental Figure S1). Thirty days post-transfer, phy mutants displayed more elongated internodes than MM adult plants, as previously described (Kerckhoffs et al., 1997; 1997) under both temperature regimes, indicating that this is a genotype-dependent phenotype. On the other hand, for all the genotypes, HT promoted a narrower stem diameter, reduced leaf area, and less branching (Figure 1A), exposing a HT-dependent phenotype. Thus, in these plants, no PHYdependent HT phenotype was observed, which contrasted with that described in A. thaliana seedlings, where HT increases elongation in a PHY-mediated manner (Jung et al., 2016). Moreover, no differences in relative water contents in leaves and fruits were observed between either genotypes or treatments (Supplemental Figure S2). These results indicate that the applied temperature treatments affected plant growth without changing the water status of the plants, rendering the experimental setup suitable to study the impact of temperature on tomato metabolism in a PHY-dependent manner.

138139

140141

142

143144

145

146147

148

149

150

High temperature alters chlorophyll metabolism and fluorescence parameters in a PHYB1/B2-dependent manner

Evidences about the role of PHYB on Chl biosynthesis (Inagaki et al., 2015) and its function as a thermosensor in *A. thaliana* (Jung et al., 2016; Legris et al., 2017) led us to investigate the effects of PHYB1/B2-dependent temperature perception on tomato Chl metabolism. To address this question, Chl levels and fluorescence parameters were analyzed in leaves from 85-day-old plants of *phyB1*, *phyB2*, and *phyB1B2* mutants alongside the corresponding wild-type genotype. Notably, HT resulted in a significant reduction of total leaf Chl content in MM genotype. However, the mutants were virtually insensitive to this temperature effect. Interestingly, regardless of the temperature treatment, leaf Chl content in the *phyB1B2* double mutant was the same as that observed in MM under

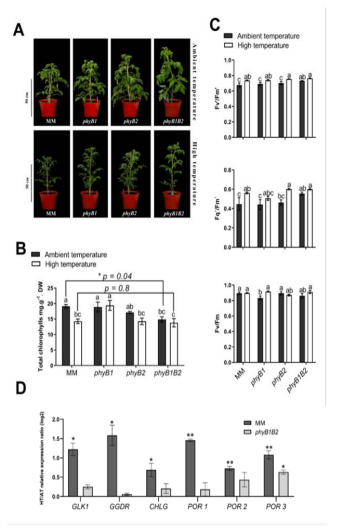


Fig. 1. PHYB1/B2 are involved in temperature perception impacting leaf chlorophyll metabolism and fluorescence parameters in tomato. (A) Side view of 50-day-old *S. lycopersicum* cv. Moneymaker (MM) plants and phyB1, phyB2 and phyB1B2 knockout mutants grown under ambient (AT, day/night 24 °C/18 °C) and high-temperature (HT, day/night 30 °C/24 °C). (B) Quantification of total chlorophylls in the seventh fully expanded leaf from 85-days-old plants. Each bar represents mean \pm SE (C) PSII maximum efficiency (Fv'/Fm'), PSII operating efficiency (Fq'/Fm') and maximum quantum efficiency of PSII (Fv/Fm) measured in the sixth fully expanded leaf from 85-days-old plants. n = at least five biological replicates. Each bar represents mean \pm SE. Different letters indicate statistically significant differences according to Fisher's multiple comparison test (p < 0.05). Asterisks (* p < 0.05, ** p < 0.01) indicate statistically significant differences by two-tailed Student t test between MM and phyB1B2 in the same environmental condition. (D) HT/AT relative expression ratio of GLK1, GGDR, CHLG, POR1, POR2 and POR3 mRNA abundance in MM and phyB1B2 mutant leaf samples from 85-days-old plants. n = at least three biological replicates. Each bar represents mean \pm SE. Asterisks (* p < 0.05, ** p < 0.01) indicate statistically significant differences by two-tailed Student t test between AT and HT within the same genotype. Genes are denoted according to the abbreviations: GLK1, GOLDEN2-LIKE1; GGDR, GERANYLGERANYL DIPHOSPHATE REDUCTASE; CHLG, CHLOROPHYLL SYNTHASE; POR, PROTOCHLOROPHYLLIDE OXIDOREDUCTASE.

152	mutant, but not the phyB1 or phyB2 single mutants, exhibited values of light-adapted PSII
153	maximum efficiency (Fv'/Fm') and PSII operating levels (Fq'/Fm') similar to those
154	observed in MM under HT conditions and higher than MM under AT conditions. No
155	impact of HT on dark-adapted PSII maximum quantum efficiency (Fv/Fm) was observed in
156	any genotype (Figure 1C).
157	In order to gain insights into the regulatory role of PHYB1/B2 as temperature mediators in
158	Chl metabolism, we further investigated the mRNA levels of genes encoding key proteins
159	involved in Chl biosynthesis that previously showed PHY-dependent transcriptional
160	regulation (Gramegna et al., 2018; Alves et al., 2020). These genes included
161	GERANYLGERANYL DIPHOSPHATE REDUCTASE (GGDR), the last enzyme for phytyl
162	diphosphate (PDP) chain synthesis (Almeida et al., 2015); the tetrapyrrole biosynthetic
163	gene PROTOCHLOROPHYLLIDE OXIDOREDUCTASE (POR; three loci exist in the
164	tomato genome named POR1 (Solyc12g013710), POR2 (Solyc10g006900) and POR3
165	(Solyc07g054210), as revealed by the phylogenetic analyses presented in Supplemental
166	Figure S3); and CHLOROPHYLL SYNTHASE (CHLG), which condensates the tetrapyrrole
167	ring with the PDP (Almeida et al., 2015). Additionally, as a marker of chloroplast
168	biogenesis and activity, we also included in the analysis the master transcriptional factor
169	GOLDEN2-LIKE1 (GLK1) (Nguyen et al., 2014), which regulates the expression of Chl
170	biosynthetic genes (Nakamura et al., 2009). HT resulted in significant increases in GLK1,
171	GGDR, CHLG, POR1, POR2, and POR3 mRNA levels in the leaves of MM genotype,
172	whereas phyB1B2 double mutant displayed a constitutive HT phenotype regarding
173	expression of Chl biosynthetic genes (except for the case of POR3, Figure 1D,
174	Supplemental Table S1), exposing a PHYB1/B2-dependent induction of Chl biosynthesis
175	under HT conditions.
176	Taken together, these results show that Chl accumulation is regulated by temperature in a
177	PHYB1/B2-dependent manner synergistically, and reveal that temperature-induced Chl
178	reduction is not the consequence of PHY-mediated impairment in Chl biosynthesis.

High temperature and PHYB1/B2 deficiency impact chloroplast biogenesis and differentiation

The effect of PHYs (Oh and Montgomery, 2014; Martin et al., 2016) and temperature 182 (Takahashi and Murata, 2008; Van Eerden, et al., 2015) on chloroplast biogenesis and 183 structure have already been studied separately in several plant species. However, these two 184 variables have not been previously assayed in an integrated manner in tomato, nor have 185 they been assessed in any other crop species where chloroplasts determine nutritional 186 quality. Here, we evaluated whether the HT-induced impact on chloroplast development is 187 mediated by PHYB1/B2 in tomato. Similarly to that observed for 85-day-old plants (Figure 188 1B), 21-day-old plants of MM, grown under HT conditions, and the phyB1B2 mutant, 189 190 regardless of the temperature condition, showed chlorotic symptoms (red arrows Figure 191 2A). This visual phenotypic difference agreed with lower measurements of leaf Chl content (Figure 2B), strengthening the idea of PHYB1/B2 as a mediator of temperature perception 192 modulating leaf Chl metabolism. Thus, to address whether differences in Chl levels were 193 associated with changes in chloroplast biology, plastid quantification and ultrastructure 194 analyses were performed in mesophyll cells. Microscopy data revealed a reduction in the 195 number of chloroplasts per mesophyll cell in MM plants grown under HT and in the 196 phyB1B2 double mutant cells under AT and HT compared to MM genotype under AT 197 (Figure 2C). Interestingly, the phyB1B2 double mutant still showed a slight temperature 198 199 response, indicating the existence of PHY-independent temperature sensing, as reported in A. thaliana (Fujii et al., 2017; Ma et al., 2016). Additionally, leaves developed under HT 200 exhibited remarkable changes in chloroplasts ultrastructure (Figure 2D). Chloroplasts of 201 MM developed under HT and those of the phyB1B2 double mutant, under both temperature 202 203 regimes, showed reduced grana stacking and dilated thylakoid lumen in comparison to MM 204 under AT. PIF transcription factors have shown to mediate PHY-dependent temperature perception 205 206 and, in particular PIF1 and PIF3, have been shown to regulate chloroplast development in 207 A. thaliana (Kim et al., 2016, Stephenson et al., 2009). In tomato PIF4 has been shown to participate in temperature-dependent seedling elongation (Rosado et al., 2019). Although 208 209 PIFs accumulation is mostly regulated post-translationally, we found that PIF1b mRNA levels increase under HT in the MM genotype but the phyB1B2 double mutant was 210 insensitive (Supplemental Figure S4A). We also identified significant upregulation of PIF3 211 mRNA in response to HT in MM and in the phyB1B2 mutant regardless of temperature. 212

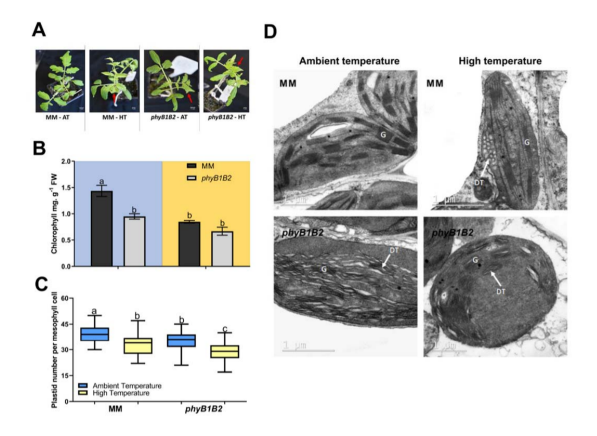


Figure 2. High temperature affects plastid biogenesis and development in leaves in a PHYB1/B2-dependent manner. (A) Visualization of a representative leaf of 21-day-old *S. lycopersicum* cv. Moneymaker (MM) and phyB1B2 knockout mutants after two weeks under ambient (AT, day/night 24 °C/18 °C) and high-temperature (HT, 30 °C/24 °C). Red arrows indicate chlorotic leaves (MM at HT, phyB1B2 at AT and phyB1B2 at HT). (B) Quantification of total chlorophylls in leaves cultivated under AT (blue background) and HT (yellow background). n = at least three biologically replicates. Each bar represents mean \pm SE. (C) Plastid density per mesophyll cell. Values represent chloroplast quantification of \pm 70 cells. Different letters indicate statistically significant differences according to Fisher's multiple comparison test (p< 0.05). Each bar represents mean \pm SE. (D) Representative TEM images of chloroplast from MM and phyB1B2 leaves grown under AT and HT. G indicates grana and DT indicated dilated thylakoids.

These alterations in *PIF1b* and *PIF3* mRNA levels may explain, at least in part, the reduced number of plastids and their altered ultrastructure.

213

214

215

216

217

In conclusion, our data suggest that temperature perception mediated by PHYB1/2 has an impact on chloroplast development that ultimately determines their structure and function.

PHYB1/B2 inactivation mediated by high temperature enhances chlorophyll 218 degradation pathway in leaves 219 In A. thaliana, PHYB regulates leaf senescence in response to light conditions through 220 PIF4/PIF5, which in turn induce Chl breakdown (Sakuraba et al., 2014). However, it is 221 unknown how PHYB-mediated temperature perception operates on Chl catabolism. The 222 Chl degradation pathway operates in tomato leaves during senescence (Lira et al., 2014; 223 Guyer et al., 2014), leading to phytol chain removal followed by the linearization of the 224 tetrapyrrole ring involving several enzymatic steps (Hörtensteiner, 2013; summarized in 225 Figure 3A). In brief, Chl a is converted to an intermediate phytol-free chlorophyllide a 226 227 (Chlide a) or a magnesium-free pheophytin a (Pheo a) form by CHLOROPHYLLASE (CLH) and STAY GREEN (SGR), respectively. Chlide a and Pheo a are subsequently 228 229 converted into pheophorbide a (Pheide a) through dechelation, mediated by STAY GREEN-LIKE (SGR-like), and dephytylation, mediated by PHEOPHYTINASE (PPH), 230 respectively. Finally, Pheide a is linearized by PHEOPHORBIDE A OXYGENASE (PAO) 231 232 to yield a red chlorophyll catabolite (RCC). To understand Chl reductions observed under HT and in the phyB1B2 mutant, we therefore measured the mRNA levels of CLH1, CLH2, 233 CLH4, SGR, PPH, SGR-like, and PAO. 234 mRNA levels of CLH2, CLH4, PPH, SGR-like, and PAO were at least two-fold up-235 regulated in MM plants cultivated under HT (Figure 3B). Consistent with our proposed role 236 of PHYB1/B2 in temperature perception, regardless of the temperature condition, the 237 phyB1B2 double mutant showed mRNA levels similar to those observed in MM under HT 238 for the CLH4, PPH, SGR-like, and PAO genes (Figure 3A and Figure 3B). It has been 239 reported that Chl degradation is triggered by dark-induced senescence and mediated by 240 PHY perception and PIF signaling (Sakuraba et al., 2014). In particular, SGR and PAO 241 242 enzyme-encoding genes are directly upregulated by PIF4 and PIF5 in A. thaliana (Song et 243 al., 2014; Zhang et al., 2015). Even though the PIF4 mRNA profile did not respond to temperature in our experiment (Supplemental Figure S4A), we cannot formally rule out its 244 245 involvement in the regulation of this process, considering that we were able to identify PIFbinding motifs in the four genes that showed PHYB1/B2-mediated temperature modulation, 246 specifically CLH4, PPH, SGR-like, and PAO (Supplemental Figure S4B). This observation 247 suggests that the PHY-PIF module acts similarly in tomato as in A. thaliana. 248

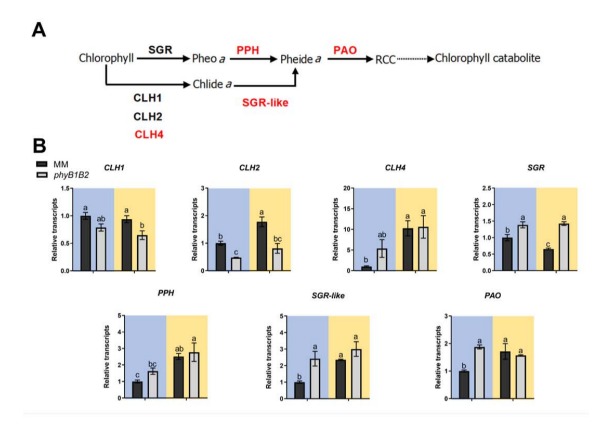


Figure 3. High temperature enhances chlorophyll degradation in leaves in a PHYB1/B2-dependent manner.

(A) Schematic model of chlorophyll degradation pathway. Enzymes and metabolites are denoted according to the following abbreviations: Pheo a, pheophytin a; Chlide a, chlorophyllide a; Pheide a, pheophorbide a; RCC, red chlorophyll catabolite; CLH, CHLOROPHYLLASE; SGR, STAY GREEN; SGR-like, PPH, PHEOPHYTINASE; STAY GREEN-LIKE; PAO, PHEOPHORBIDE a OXYGENASE. The enzymes highlighted in red are those that showed to be regulated by temperature in a PHYB1/B2-dependent manner according to Figure 3B. (B) Relative mRNA levels of chlorophyll degrading enzymes encoding genes in Moneymaker (MM) and phyB1B2 mutant leaf samples from 85-day-old plants grown under ambient temperature (AT, 24 °C/18 °C - blue background) and high-temperature (HT, 30 °C/24 °C - yellow background). Expression levels are relative to MM – AT condition. n = at least three biological replicates. Each bar represents mean \pm SE. Different letters indicate statistically significant differences according to Fisher's multiple comparison test (p< 0.05).

Collectively, these results demonstrated that PHYB1/B2 represses Chl catabolism genes in a temperature-dependent manner in tomato leaves.

249

250

251

252

253

PHYB1/B2-dependent temperature perception regulates leaf carotenoid biosynthesis at the transcriptional level

Alongside the induction of the Chl degradation pathway, the HT treatment and the 254 phyB1B2 double mutation resulted in over 30% reduction of total leaf carotenoids (Figure 255 4A). To understand the molecular basis of this reduction, we analyzed the transcriptional 256 profile of carotenoid biosynthetic enzyme-encoding genes. GERANYLGERANYL 257 DIPHOSPHATE SYNTHASE 1 (GGPS1), which produces the carotenoids precursor 258 geranylgeranyl diphosphate (Quadrana et al., 2013), was shown to be downregulated both 259 260 in response to HT and in the phyB1B2 double mutant (Figure 4B). By contrast, genes encoding carotenoid downstream enzymes, such as PHYTOENE SYNTHASE 2 (PSY2), 261 PHYTOENE DESATURASE (PDS), CHROMOPLAST-SPECIFIC LYCOPENE β 262 263 CYCLASE (CYCβ), and CHLOROPLAST-SPECIFIC LYCOPENE β CYCLASE (LYCβ), did not show PHYB1/B2-mediated temperature modulation (Supplemental Figure S5). 264 Thus, the results presented above indicate that, in tomato leaves, the accumulation of 265 266 chloroplast photosynthetic pigments is controlled through transcriptional adjustments of Chl degradation and carotenoid biosynthesis genes by the ambient temperature in a 267 268 PHYB1/B2-dependent manner.

269

- Fruit carotenoid contents are modulated by temperature through PHYA and
- 271 PHYB1/B2
- A role for PHYs in tomato fruit carotenoid accumulation has long been proposed (Alba et
- 273 al., 2000a; Gupta et al., 2014), and we recently demonstrated that fruit-localized PHYA and
- 274 PHYB2 positively influence fruit carotenoid accumulation (Bianchetti et al., 2018). Aiming
- 275 to evaluate the temperature effects on carotenogenesis, we followed two complementary
- approaches: i) investigate the single phyA, phyB1, and phyB2 mutants and the triple
- 277 phyAB1B2 mutant and ii) analyze fruit-specific RNAi PHYA- and PHYB-silenced lines in
- the Micro-Tom (MT) background (PHYA^{RNAi} and PHYB2^{RNAi}). This would allow us to
- 279 unravel whether fruit-localized PHYs regulate carotenogenesis in a temperature-dependent
- 280 manner in this organ and if this mechanism is genotype-independent.
- 281 Except for the *phyAB1B2* triple mutant, ripe fruits harvested from HT-grown plants
- exhibited reductions in total carotenoid content compared to that in AT counterparts (Figure
- 283 5A). Ripe fruits collected from single phyA-, phyB1-, and phyB-mutant plants or double
- 284 phyB1B2-mutant plants grown under AT displayed lower total carotenoid levels those that

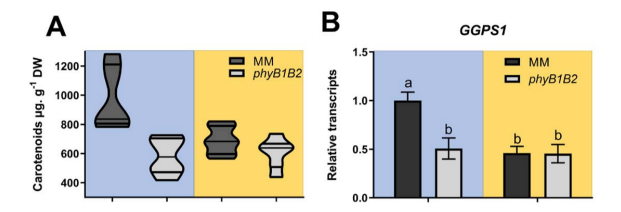


Figure 4. PHYB1/B2-dependent temperature perception transcriptionally regulates leaf carotenogenesis.

(A) Total carotenoid levels expressed in μ g per g of dry weight. n = at least five biological replicates. (B) Relative mRNA levels of *GGPS1* gene in Moneymaker (MM) and *phyB1B2* mutant leaf samples from 85-day-old plants grown under ambient temperature (AT, 24 °C/18 °C - blue background) and high-temperature (HT, 30 °C/24 °C - yellow background). Expression levels are relative to MM – AT condition. n = at least three biological replicates. Each bar represents mean \pm SE. Different letters indicate statistically significant differences according to Fisher's multiple comparison test (p< 0.05). Gene is denoted according to the abbreviation: *GGPS1*, *GERANYLGERANYL PHOSPHATE SYNTHASE 1*.

in MM counterparts. Interestingly, an enhanced reduction was observed in the *phyAB1B2* triple mutant, which displayed similar total fruit carotenoid levels under both temperature regimes (Figure 5A). High-performance liquid chromatography (HPLC) carotenoid profiling revealed that accumulation of lycopene, alongside its precursors phytoene and phytofluene, was reduced by HT compared to AT in all genotypes aside from in the triple *phyAB1B2* mutant, which showed similar profiles of phytoene, phytofluene, and lycopene in fruits developed under both temperature regimes (Supplemental Figure S6 and Figure 5B).

To investigate whether the temperature-mediated reduction in carotenoid content is a consequence of the transcriptional regulation of carotenoid biosynthetic genes, we further profiled *GGPS2*, *PSY1*, and *PDS* mRNA abundances in MM and the *phyAB1B2* triple mutant. In agreement with enhanced carotenoid biosynthesis during ripening, the three analyzed genes displayed a clear induction in MM under AT at the onset of ripening (from mature green to breaker stages). At the mature green stage, no significant difference in expression was observed attributable to genotype or temperature treatment. However, at

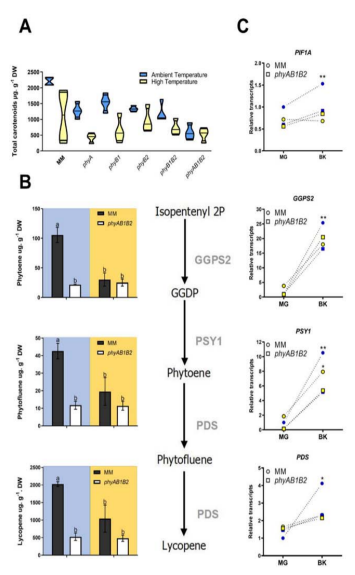


Figure 5. PHYA/B1/B2-dependent temperature perception transcriptionally regulates fruit carotenogenesis. (A) Total carotenoid (phytoene, phytofluene, lycopene, lutein and β-carotene) levels quantified from ripe fruits of Moneymaker (MM) and phyA, phyB1, phyB2, phyB1B2 and phyAB1B2 mutant plants grown under ambient (AT, blue, day/night 24 °C/18 °C) and high-temperature (HT, yellow, day/night 30 °C/24 °C). (B) Center: schematic model of lycopene biosynthetic pathway, the dotted line represents more than one enzymatic step. Left: levels of lycopene, phytoene and phytofluene in ripe fruits. AT: blue background; HT: yellow background. Each bar represents mean ± SE. Different letters indicate statistically significant differences according to Fisher's multiple comparison test (p< 0.05). Right: Relative mRNA levels from carotenoids biosynthetic genes GGPS2, PSY1 and PDS in fruits at mature green (MG) and breaker (BK) stages harvested from plants grown at AT (blue) and HT (yellow). Transcripts levels are expressed relative to MM MG – AT condition. Asterisks (* p<0.05, ** p<0.01) indicate differences in the analysis of variance in a multiple comparison test within the same fruit stage. Metabolites and genes are denoted according to the following abbreviations: GGDP, geranylgeranyl diphosphate; GGPS2, GERANYLGERANYL PHOSPHATE SYNTHASE 2; PSY1, PHYTOENE SYNTHASE 1; PDS, PHYTOENE DESATURASE. (C) Relative mRNA levels of PIF1a carotenogenesis regulator gene in fruits at mature green (MG) and breaker (BK) stages harvested from plants grown at AT (blue) and HT (yellow). Transcripts levels are expressed relative to MM MG – AT condition. Asterisks (** p<0.01) indicate differences in the analysis of variance in a multiple comparison test within the same fruit stage.

both temperature and the triple phyAB1B2 mutation (Figure 5B). The role of PIF1a in 301 tomato fruit carotenogenesis has been previously reported. Although, upon induction of 302 ripening, PIF1a transcript levels increased; concomitantly, Chl degradation alters the 303 quality of the light filtered through the fruit pericarp, increasing the relative red/far-red 304 ratio. As a consequence, PIF degradation increases, enhancing PSYI expression and 305 carotenoid accumulation (Llorente et al., 2016). Indeed, our results revealed PIF1a 306 307 upregulation from MG to BK stage in MM grown under AT, but its levels remained invariable in both genotypes under HT (Figure 5C). 308

In summary, the temperature-insensitive phenotype observed in the phyAB1B2 triple-

mutant fruits revealed that the inductive effect of PHYA/B1/B2 over the transcription of

311 carotenogenesis-associated genes is abolished by HT conditions.

In order to mitigate the influence of any possible pleiotropic effect of *phy* mutations at the

313 whole-plant level, we further analyzed fruit carotenoid accumulation in fruit-specific

314 PHYA- and PHYB2-silenced lines, namely PHYARNAi and PHYB2RNAi. Regardless of

315 temperature, the lycopene content in *PHYA*^{RNAi} or *PHYB*2^{RNAi} fruit was reduced by half in

316 comparison to control fruits developed under AT, the latter of which were affected by HT

317 conditions (Figure 6A). It is worth noting that this effect was observed in two independent

318 transgenic lines for each genotype (Supplemental Figure S7). In agreement, *PSY1* transcript

accumulation accompanied the variations observed in lycopene content (Figure 6B),

320 confirming that fruit-localized PHY-mediated temperature perception controls carotenoids

321 accumulation through transcriptional regulation of the biosynthetic enzyme-encoding

322 genes.

323

324

309

310

PHYA- and PHYB2-mediated temperature perception controls carotenoids

325 biosynthesis through master ripening transcription factors

Besides PIF1a (Llorente et al., 2016), PSYI expression has been demonstrated to be

regulated by ripening-associated transcription factors in tomato (Liu et al., 2015). For this

328 reason, we profiled the mRNA levels of candidate genes encoding master ripening

regulators, namely APETALA2a (AP2a), NON-RIPENING (NOR), RIPENING INHIBITOR

330 (RIN), TOMATO AGAMOUS-LIKE (TAGL1), and FRUITFULL1 and 2 (FUL1/2) (Klee and

Giovannoni, 2011), in *PHY* RNAi lines subjected to both temperature regimes.

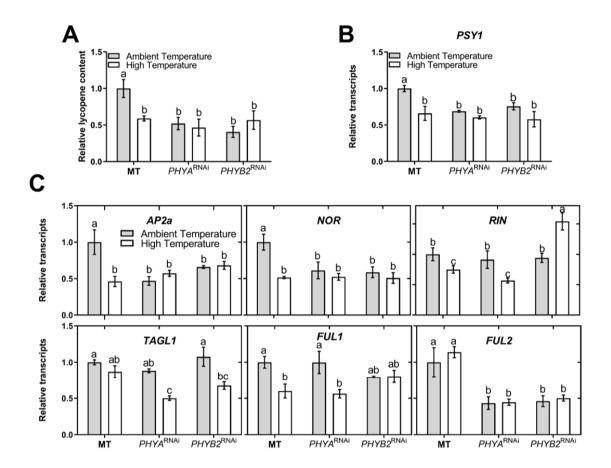


Figure 6. Fruit-localized PHYA and PHYB2 are involved in temperature perception impacting lycopene synthesis and master fruit ripening regulators. (A) Lycopene levels quantified in ripe fruits from Micro-Tom (MT) control genotype and fruit-specific *PHYA-* (*PHYA*^{RNAi}) and *PHYB2-* (*PHYB2*^{RNAi}) knockdown transgenic lines grown under ambient (24 °C/18 °C) and high temperature (30 °C/24 °C). Lycopene levels were quantified and expressed relative to MT fruits at ambient temperature and values are means of at least three biological replicates from two independent lines for each genotype. Each bar represents mean ± SE. (B,C) Relative mRNA levels of (B) *PSY1* (C) and master fruit ripening regulator genes in MT, *PHYA*^{RNAi} and *PHYB2*^{RNAi} breaker fruit samples harvested under AT and HT. Expression levels are relative to MT – AT condition. n = at least three biological replicates. Each bar represents mean ± SE. Different letters indicate statistically significant differences according to Fisher's multiple comparison test (*p*< 0.05). *RIN*; *RIPENING INHIBITOR*, *NOR*; *NON-RIPENING*, *FUL1*; *FRUITFULL1*, *FUL2*; *FRUITFULL2*, *AP2a*; *APETALA2a*, *TAGL1*; *TOMATO AGAMOUS-LIKE1*, *PSY1*; *PHYTOENE SYNTHASE 1*.

Three expression patterns were observed; i) *RIN*, *TAGL1*, and *FUL1* mRNA levels responded to temperature treatment and in certain cases these responses varied in the absence of *PHYA* or *PHYB2*, ii) *FUL2* was downregulated only in the lack of functional PHYs (*PHYA*^{RNAi} or *PHYB2*^{RNAi} lines), and iii) *AP2a* and *NOR* mRNAs displayed a clear

332

333334

response to PHY-mediated temperature perception (Figure 6C). These results suggest that fruit-localized PHY-mediated temperature perception controls carotenoid accumulation via transcriptional regulation of ripening master controller genes.

Combined, these results decrypt the role of fruit-localized PHYA and PHYB1/B2 as temperature sensors in tomato fruits, which regulate carotenoid accumulation through the transcriptional control of genes involved in their biosynthesis by alternative and converging molecular pathways.

DISCUSSION

346

345

Studies performed on plants under warm treatment or employing PHY-signalling defective 347 mutants have demonstrated the synergistic influence of light and temperature on the 348 regulation of chloroplast metabolism (Stephenson et al., 2009; Zhao et al., 2016; Spicher et 349 al., 2016; Dubreuil, et al., 2017; Spicher et al., 2017). Despite this fact, the two factors have 350 351 been studied independently, limiting our comprehensive understanding of light- and temperature-perception mechanisms at the molecular level. Recently, the role of PHYs as 352 353 thermosensors by the gradual inactivation of PHYB by increasing temperature was reported 354 in A. thaliana (Legris et al., 2017). However, whether this mechanism operates in crop species evolved in different environments, and how this PHY-dependent temperature 355 signaling cascade impacts major metabolic pathways, remains poorly understood. It is 356 worth mentioning that PHY-independent temperature responses have also been described 357 associated with other photoreceptors (Fujii et al., 2017; Ma et al., 2016) and also in 358 photoreceptor-impaired conditions (Legris et al., 2016). In accordance, we also observed 359 PHY-independent temperature responses here (Figure 2C). In the current study, we present 360 experimental evidence that central enzyme-encoding genes of Chl and carotenoid 361 metabolism are regulated at the transcriptional level by PHYB1/B2- and PHYA/B1/B2-362 mediated temperature perception in tomato leaf chloroplasts and fruit chromoplasts, 363 364 respectively. Constitutive HT-induced features were previously observed in the A. thaliana phyB loss-of-365 function mutant (Jung et al., 2016; Huang et al., 2019). In contrast to the single-copy gene 366 PHYB found in A. thaliana (Sharrock and Quail, 1989), the tomato genome harbors two 367 PHYB paralogs, PHYB1 and PHYB2, which originated during a genome triplication event 368 369 in the Solanaceae common ancestor (Tomato Genome Consortium, 2012). Indeed, results 370 from our current work indicate that loss-of-function in the double phyB1B2 mutant, rather than single phyB1 or phyB2 mutations, is necessary to generate a thermoinsensitive 371 phenotype in tomato leaves leading to a reduction in Chl content, indicating that PHYB1 372 and PHYB2 play additive functions as temperature sensors (Figure 1B). The phyB1B2 373 double mutant did not show changes in the mRNA abundances of Chl biosynthetic genes in 374 response to HT (Figure 1D). However, these genes were upregulated by HT in MM leaves, 375

which does not explain the low-Chl phenotype shown by these plants (Figure 1B) and 376 suggests a highly complex regulatory mechanism. 377 Interestingly, together with the reduction in Chl content, increased Chl fluorescence 378 parameters were registered (Figure 1C). Our results are consistent with the finding in A. 379 thaliana that faster electron transport occurred under HT conditions acting as an electron 380 sink for the increment in photorespiration (Zhang and Sharkey, 2009). 381 382 The observable decrease in total Chl has been associated with a decrease in the LHCII, serving as a protective mechanism in plants undergoing abiotic stress (Ishida et al., 2000). 383 384 In agreement, we observed a reduction in chloroplast number and alterations in grana 385 stacking (Figure 2), probably mediated by higher PIF3 mRNA level in response to PHYB1B2-mediated temperature inactivation (Supplemental Figure S4A). In line with this, 386 387 PIF3 protein accumulation led to impaired chloroplast development in A. thaliana (Stephenson et al., 2009). Moreover, our results provide genetic evidence that HT and the 388 phyB1B2 mutation trigger Chl catabolism via the upregulation of genes associated with Chl 389 degradation, i.e., CLH4, PPH, SGR-like, and PAO (Figure 3). It was recently demonstrated 390 that PIF4 regulates senescence in tomato (Rosado et al., 2019). Additionally, PHYs trigger 391 PIF4 degradation (Lorrain et al., 2008), avoiding its inductive role on Chl breakdown 392 (Sakuraba et al., 2014; Song et al., 2014; Zhang et al., 2015). In this sense, several 393 canonical PIF binding motifs, i.e., G- and PBE-box (Martínez-Garcia et al., 2000; Zhang et 394 al., 2013), were found in the promoter of the Chl degrading-associated genes mentioned 395 above (Supplemental Figure S4B). Further reports extensively associate the inhibition of 396 PIF4 degradation in response to HT impacting several events throughout the plant cycle 397 (Koini et al., 2009; Qiu et al., 2019; Zhou et al., 2019). Together, these data suggest that 398 inactivation of PHYB1/B2 under HT affects Chl accumulation through altered chloroplast 399 400 biogenesis (Figure 2) and degradation (Figure 3), likely via PIF1b/PIF3- and PIF4-401 dependent mechanism(s), respectively. Our findings additionally indicate a down-regulation of GGPS1 mRNA levels by 402 403 temperature and the phyB1B2 mutation (Figure 4B). Since GGPS is the last shared step 404 between the Chl and carotenoid biosynthetic pathways (Cordoba et al., 2009), this result is

in accordance with the impairment of Chl and carotenoid synthesis in the leaves under these

405

406

conditions (Figure 2B,4A).

Besides the impact on vegetative organs, our data also indicated that HT negatively 407 influences carotenoid accumulation in tomato fruits (Figure 5A). Alterations of isoprenoid-408 derived compounds in response to thermal stress have been increasingly demonstrated over 409 the last years (Velikova et al., 2011; Spicher et al., 2016). Exposure of single, double, and 410 triple PHY mutants as well as the MM control plants to two independent temperature 411 regimes revealed a combinatory effect of temperature and the action of PHYs on carotenoid 412 content of ripe fruits (Figure 5A). In contrast to the other genotypes analyzed, the reduced 413 levels of fruit lycopene and their precursors observed in phyAB1B2 might be due to 414 415 impaired temperature perception in this genotype (Figure 5B). Consistent with this view, 416 our findings indicate that the phyAB1B2 mutations, as well as HT conditions, downregulate the expression of the major carotenoid biosynthetic genes, e.g., GGPS2, PSY1, and PDS 417 (Figure 5B). This is in agreement with our previous report showing that these genes are 418 transcriptionally regulated by PHY-mediated light perception in tomato fruit (Bianchetti et 419 al., 2018). 420 421 To further examine whether the observed impact on fruit carotenoids was the effect of whole-plant PHY deficiency or those localized in the fruits, we analyzed fruit-specific 422 RNAi PHYA- and PHYB-silenced lines in MT genetic background (Bianchetti et al., 2018). 423 Results showed equivalent reductions of lycopene accumulation and PSY1 transcript levels 424 in both transgenic lines, showing temperature insensitivity (Figure 6A, 6B), demonstrating 425 that fruit-localized PHYs act as thermosensors and that this effect is genotype independent. 426 Indeed, the central role played by temperature and PHYs controlling PSYI expression 427 (Figure 6B) is supported by previous studies demonstrating that PIF1a downregulates 428 carotenoid biosynthesis via *PSY1* transcriptional repression in tomato fruits (Llorente et al., 429 2016), similarly as its ortholog in A. thaliana in response to temperature (Toledo-Ortiz et 430 al., 2014). In addition to PIF1a, carotenoid biosynthesis in tomato fruit is markedly 431 regulated at the transcriptional level by the master ripening transcription factors RIN, NOR, 432 FUL1, FUL2, TAGL1, and AP2a (Klee and Giovanonni, 2011). Whereas a non-combined 433 effect of fruit-specific PHY expression and temperature on RIN, TAGL1, FUL1, or FUL2 434 transcript abundance was observed, AP2a and NOR mRNA levels responded to temperature 435 in a PHY-dependent manner (Figure 6C), thus contributing to a reduction in lycopene 436 content (Figure 6A), most likely by the transcriptional regulation of *PSYI*, as previously 437

described (Chung et al., 2010; Karlova et al., 2011; Yuan et al., 2016; Cruz et al., 2018). 438 These results exposed an interesting network that regulates ripening in response to light and 439 temperature in an independent or integrated way, warranting this key evolutionarily 440 selected process. 441 442 Overall, our results support a model (Figure 7) where increases in temperature induce the inactivation of PHYs, probably through the conformational change from the biologically 443 444 active Pfr to the inactive Pr form. The thermosensing role of PHYB1/B2 impacts leaf Chl and carotenoid levels through combined control of chloroplast biogenesis and the transcript 445 levels of carotenoid biosynthetic and Chl degrading enzyme genes. Moreover, our data 446 447 demonstrate that PHYA/PHYB1/PHYB2-mediated temperature perception modulates carotenoid metabolism in fruit. The data also showed the involvement of master ripening 448 449 regulator genes as mediators of PHY-dependent temperature regulation of the carotenoid biosynthetic pathway. In conclusion, this study demonstrates the effect of PHY-mediated 450 temperature perception on both photosynthetic and heterotrophic tomato plastid 451 metabolism. Moreover, the results presented here identify the PHYs as critical hubs that 452 can be manipulated to maintain and/or improve the nutritional quality of edible fruits in the 453 context of global increasing temperatures. 454 455

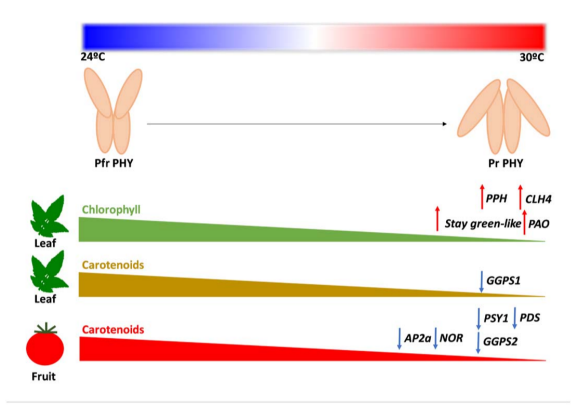


Figure 7. The effect of PHY-mediated temperature perception on tomato metabolism regulation. The rise of ambient temperature shifts balance to the inactive phytochrome (Pr) form, which promotes chlorophyll degradation pathway in source leaves through the transcriptional up-regulation of chlorophyll catabolic enzyme-associated genes. Additionally, reduced levels of Pfr impair carotenoid accumulation in both leaves and ripe fruits, through the transcriptional downregulation of carotenoid biosynthetic and master ripening regulator genes.

METHODS

Plant material, growth conditions, and sampling

Solanum lycopersicum plants (cv. MoneyMaker) harboring loss-of-function mutations in phyA, phyB1, phyB2, phyB1B2, and phyAB1B2 were previously characterized (Kerckhoffs et al., 1996; Kerckhoffs et al., 1997; Lazarova et al., 1998a; Lazarova et al., 1998b; Kerckhoffs et al., 1999; Weller et al., 2000). Micro-Tom fruit-specific PHYA- and PHYB2-silenced lines (PHYA^{RNAi} and PHYB2^{RNAi}) were previously obtained and characterized by Bianchetti et al. (2018). Although the Micro-Tom cultivar is deficient in brassinosteroid biosynthesis due to the weak mutation d, it has been extensively demonstrated that it represents a convenient and adequate model system to study fruit biology (Campos et al.,

2010). In this work, we used Micro-Tom PHYA^{RNAi} and PHYB2^{RNAi} lines as a proof of 468 concept that fruit-localized PHYs regulate carotenogenesis in a temperature-dependent 469 manner in this organ and that this mechanism is genotype independent. 470 Tomato seeds (cv. MoneyMaker and Micro-Tom) were sown under standard greenhouse 471 conditions (day/night 24°C/18°C, 16/8 h light/dark, and 60% air relative humidity) in 472 Floricultura Z substrate. Twenty-day-old plants were transplanted to 9-L pots and cultivated 473 474 for 120 days (16/8 h light/dark and 60% air relative humidity) in greenhouses under two distinct temperature regimens: ambient (24°C/18°C) and high (30°C/24°C) temperature 475 with a mean daily difference of 5°C (Supplemental Figure S1A, B). These temperatures 476 477 have been previously described as optimal and suboptimal high for tomato (Ayenan et al., 2019). Plants were cultivated under 250 umol m⁻² s⁻¹ light intensity, which is lower than the 478 tomato saturation point (approximately 800 µmol m⁻² s⁻¹). To avoid unwanted effects on 479 plant water status (Fahad et al., 2017), the plants were watered twice a day and the relative 480 ambient humidity was monitored (Supplemental Figure S1C). Leaves (seventh fully 481 expanded leaf from bottom to top) and fruits [at mature green, breaker, and ripe (7 days 482 after breaker) stages] from MoneyMaker background plants were harvested 65 and 80-110 483 days after the beginning of the temperature treatment. Fruits from Micro-Tom plants were 484 harvested 75-90 days after the start of temperature treatment. Samples were ground in 485 liquid N₂ and freeze-dried prior to subsequent analysis. 486

487

488

Pigment quantification

Chlorophyll (Chl) extraction was carried out from 5 mg of freeze-dried leaf tissue in 1 mL 489 of 80:20 acetone:Tris-HCl 100 mM pH 7.5. Samples were sonicated for 5 min at 42 kHz 490 and further centrifuged at 16000 × g for 2 min. The supernatant was collected and the 491 procedure was repeated until the green color was totally removed from the tissue. 492 493 Spectrophotometer measurements were performed at 537, 647, 663, and 750 nm. Total Chl content was estimated by the equation: 0.01373 X (A663-A750) - 0.000897 X (A537-494 A750) - 0.003046 X (A647-A750) (Sims and Gamon, 2003). For leaf carotenoid 495 extraction, 5 mg of freeze-dried samples were immersed in 1 ml of N,N-496 dimethylformamide, sonicated for 5 min at 42 kHz, and centrifuged at 16000 × g for 5 min. 497 Supernatant absorbance was recorded at 480, 647, and 664 nm, then carotenoid contents 498

were determined by the equation: {1000 X A480 – [1.12 X (12 X A664) – (3.11 X A647)] + [34.07 X (20.78 X A647) – (4.88 X A664)]} / 245 (Wellburn, 1994). Fruit carotenoid extraction was carried out from 20 mg of freeze-dried fruit tissue according to Bianchetti et al., 2018. Phytoene, phytofluene, lycopene, β-carotene, and lutein were determined by high-performance liquid chromatography (HPLC). Eluted compounds were detected at 450 nm (lycopene, β-carotene, and lutein), 286 nm (Phytoene), and 347 nm (Phytofluene) as described in Fraser et al. (2000).

506 507

508

509

510

511

512

513

514

515

516

Chlorophyll fluorescence measurements

Chl fluorescence parameters were determined according to Lira et al. (2017) using a portable open gas-exchange system (LI-6400XT system; LI-COR) equipped with an integrated modulated Chl fluorometer (LI-6400-40; LI-COR). Briefly, the second leaflet of the sixth fully expanded leaf from 85-day-old plants was kept under dark adaptation for 60 min, then weak and saturating white light pulses were applied to determine, respectively, initial fluorescence and maximum fluorescence emission. Further, the same procedure was applied on light-adapted leaves to determining the light-adapted initial fluorescence and maximum fluorescence emission. The values were used to calculate maximum quantum efficiency of PSII, PSII operating efficiency, and PSII maximum efficiency.

517518

Plastid abundance and ultrastructure

- Two-week-old MoneyMaker and phyB1B2 plants grown under standard conditions were
- 520 transferred to the distinct temperature conditions described above. After one week, the
- 521 fourth fully expanded leaf was used for determining plastid abundance and ultrastructure
- 522 analysis.
- Plastid abundance was determined in the leaf mesophyll as described in Bianchetti et al.
- 524 (2017). In brief, leaf samples were incubated in 3.5% glutaraldehyde for 60 min and then in
- 525 0.1 M Na-EDTA (pH 9.5) at 60°C for 180 min. Isolated cells were visualized through
- optical microscopy and plastid number per cell was estimated using the ImageJ program
- 527 (https://imagej.nih.gov/ij/).
- For ultrastructure analysis, leaf segments were fixed at 4°C in Karnovsky's solution (2.5%
- 529 [v/v] glutaraldehyde and 2% [v/v] paraformaldehyde in 0.1 M sodium phosphate buffer, pH

- 530 7.2) for 24 h. After washing in phosphate buffer, the samples were post-fixed in buffered
- 531 1% (w/v) osmium tetroxide, washed, dehydrated in a graded series of acetone, and
- embedded in Spurr's resin. The resin was polymerized at 60°C. Ultrathin sections were
- stained with saturated uranyl acetate (Watson, 1958) and lead citrate (Reynolds, 1963) and
- observed using a JEM 1011 transmission electron microscope.

535

536

RNA extraction and reverse transcription quantitative PCR (RT-qPCR)

- 537 RNA extraction, cDNA synthesis, primer design, and qPCR assays were performed as
- described in Quadrana et al. (2013). The primers used for RT-qPCR analyses are listed in
- 539 Supplementary Table S2. qPCR reactions were performed in a QStudio6 A1769 PCR
- Real-Time thermocycler using 2X Power SYBR Green Master Mix in a final volume of 10
- 541 μL. Absolute fluorescence data were analyzed using LinRegPCR software to obtain Ct and
- primer efficiency values. Relative mRNA abundance was calculated and normalized with
- 543 the $\Delta\Delta$ Ct method using two reference genes (Expósito-Rodríguez et al., 2008):
- 544 EXPRESSED and TIP4.1 for leaves and EXPRESSED and CAC for fruits (Quadrana et al.,
- 545 2013).

546 547

Phylogenetic analysis

- 548 Amino acid sequences of A. thaliana PROTOCHLOROPHYLLIDE
- OXIDOREDUCTASES (PORA, PORB, and PORC) were BLAST against the tomato
- 550 genome in Sol Genomics network database (http://solgenomics.net). Homologous
- 551 sequences from Solanum tuberosum, A. thaliana, Arapidopsis lyrata, Brasica oleracea,
- 552 Brasica rapa, Sorghum bicolor, Zea mays, and Setaria viridis were retrieved by BLASTp
- 553 against Viridiplantae in Phytozome (http://phytozome.jgi.doe.gov/pz/portal.html).
- 554 MUSCLE package available in MEGA software 10.0.3 was used to perform multiple
- sequences alignments. Phylogenetic reconstruction was performed with the maximum-
- 556 likelihood method with 5,000 bootstrap replications.

557 558

Data analysis

- The values in the figures represent the mean of at least three biological replicates \pm
- standard error. Statistical differences in parameters were analyzed with InfoStat/F software

- 561 (http://www.infostat.com.ar). Two-way analysis of variance (ANOVA) was performed to
- determine genotype (G), environmental (E) or GxE interaction. A Fisher test (p < 0.05) was
- performed to compare GxE interaction and a t-test (p < 0.05) was applied to discriminate
- means of the sample within genotypes.

565

- **Accession numbers**
- Sequence data from this article can be found in the GenBank/EMBL data libraries under
- accession numbers: AJ001913 (phyA); AJ002281 (phyB1) and AF122901 (phyB2).

569

570 Supplemental Data

571

- 572 **Supplemental Figure S1.** Time course of temperature and relative humidity measurements
- 573 registered along the plant growth cycle.
- 574 Supplemental Figure S2. Hydric status of Moneymaker and knockout-phytochrome
- 575 mutant plants at two temperatures regimes.
- 576 Supplemental Figure S3. Phylogenetic construction of the PROTOCHLOROPHYLLIDE
- 577 OXIDOREDUCTASE protein family.
- 578 Supplemental Figure S4. PHYTOCHROME INTERACTION FACTORS involvement in
- 579 the regulation of temperature-induced chlorophyll reduction.
- Supplemental Figure S5. Expression profile of carotenoid biosynthetic genes in leaves.
- 581 Supplemental Figure S6. Carotenoid profile in ripe fruits from Moneymaker and
- 582 knockout- phytochrome mutants.
- 583 Supplemental Figure S7. Lycopene content in ripe fruits from fruit-specific PHYA- and
- 584 PHYB2-knockdown transgenic lines.
- 585 **Supplemental Table S1.** Relative expression of chlorophyll biosynthetic genes.
- 586 **Supplemental Table S2.** Primer sequences used in this study.

- 588 Figure Legends
- 589 Figure 1. PHYB1/B2 are involved in temperature perception impacting leaf
- 590 chlorophyll metabolism and fluorescence parameters in tomato. (A) Side view of 50-
- day-old S. lycopersicum ev. Moneymaker (MM) plants and phyB1-, phyB2-, and phyB1B2-

knockout mutants grown under ambient- (AT, day/night 24°C/18°C) and high-temperature 592 (HT, day/night 30°C/24°C) conditions. (B) Quantification of total chlorophyll (Chl) in the 593 seventh fully expanded leaf from 85-day-old plants. Each bar represents mean \pm SE. (C) 594 PSII maximum efficiency (Fv'/Fm'), PSII operating efficiency (Fq'/Fm'), and maximum 595 quantum efficiency of PSII (Fv/Fm) measured in the sixth fully expanded leaf from 85-day-596 old plants. n = at least five biological replicates. Each bar represents mean \pm SE. Different 597 598 letters indicate statistically significant differences according to Fisher's multiple comparison test (p < 0.05). Asterisks (* p < 0.05, ** p < 0.01) indicate statistically significant 599 differences by two-tailed Student's t-test between MM and phyB1B2 under the same 600 601 environmental conditions. (D) HT/AT relative expression ratio of GLK1, GGDR, CHLG, POR1, POR2, and POR3 mRNA abundance in MM and phyB1B2-mutant leaf samples 602 603 from 85-day-old plants. n = at least three biological replicates. Each bar represents mean \pm SE. Asterisks (* p < 0.05, ** p < 0.01) indicate statistically significant differences by two-604 tailed Student's t-test between AT and HT within the same genotype. Genes are denoted 605 according to the abbreviations: GLK1, GOLDEN2-LIKE1; GGDR, GERANYLGERANYL 606 *DIPHOSPHATE* CHLG, 607 *REDUCTASE*; CHLOROPHYLL SYNTHASE; POR. PROTOCHLOROPHYLLIDE OXIDOREDUCTASE. 608

609

610

611

612613

614

615

616

617

618

619

620

621

Figure 2. High temperature affects plastid biogenesis and development in leaves in a PHYB1/B2-dependent manner. (A) Visualization of a representative leaf from 21-day-old *S. lycopersicum* cv. Moneymaker (MM) and *phyB1B2*-knockout mutants after two weeks under ambient- (AT, day/night 24°C/18°C) and high-temperature (HT, 30°C/24°C) conditions. Red arrows indicate chlorotic leaves (MM at HT, *phyB1B2* at AT, and *phyB1B2* at HT). (B) Quantification of total Chl in leaves cultivated under AT (blue background) and HT (yellow background) conditions. n = at least three biological replicates. Each bar represents mean \pm SE. (C) Plastid density per mesophyll cell. Values represent chloroplast quantification of \pm 70 cells. Different letters indicate statistically significant differences according to Fisher's multiple comparison test (p< 0.05). Each bar represents mean \pm SE. (D) Representative TEM images of chloroplasts from MM and *phyB1B2* leaves grown under AT and HT conditions. G indicates grana and DT indicates dilated thylakoids.

Figure 3. High temperature enhances chlorophyll degradation in leaves in a 623 PHYB1/B2-dependent manner. (A) Schematic model of chlorophyll (Chl) degradation 624 pathway. Enzymes and metabolites are denoted according to the following abbreviations: 625 Pheo a, pheophytin a; Chlide a, chlorophyllide a; Pheide a, pheophorbide a; RCC, red 626 chlorophyll catabolite; CLH, CHLOROPHYLLASE; SGR, STAY GREEN; SGR-like, 627 PPH, PHEOPHYTINASE; STAY GREEN-LIKE; PAO, PHEOPHORBIDE 628 OXYGENASE. The enzymes highlighted in red are those that showed to be regulated by 629 630 temperature in a PHYB1/B2-dependent manner according to Figure 3B. (B) Relative 631 mRNA levels of Chl degrading enzyme-encoding genes in Moneymaker (MM) and phyB1B2 mutant leaf samples from 85-day-old plants grown under ambient- (AT. 632 24°C/18°C; blue background) and high-temperature (HT, 30°C/24°C; yellow background) 633 conditions. Expression levels are relative to MM - AT conditions. n = at least three 634 biological replicates. Each bar represents mean \pm SE. Different letters indicate statistically 635 significant differences according to Fisher's multiple comparison test (p < 0.05). 636

637

638

639

640

641

642

643

644

645

646

647

Figure 4. PHYB1/B2-dependent temperature perception transcriptionally regulates leaf carotenogenesis. (A) Total carotenoid levels expressed in μg per g of dry weight. n = at least five biological replicates. (B) Relative mRNA levels of *GGPS1* gene in Moneymaker (MM) and *phyB1B2*-mutant leaf samples from 85-day-old plants grown under ambient- (AT, 24°C/18°C; blue background) and high-temperature (HT, 30°C/24°C; yellow background) conditions. Expression levels are relative to MM – AT conditions. n = at least three biological replicates. Each bar represents mean \pm SE. Different letters indicate statistically significant differences according to Fisher's multiple comparison test (p< 0.05). Gene is denoted according to the abbreviation: *GGPS1*, *GERANYLGERANYL PHOSPHATE SYNTHASE 1*.

648

Figure 5. PHYA/B1/B2-dependent temperature perception transcriptionally regulates fruit carotenogenesis. (A) Total carotenoid (phytoene, phytofluene, lycopene, lutein, and β-carotene) levels quantified from ripe fruits of Moneymaker (MM) and *phyA*-, *phyB1*-, *phyB2*-, *phyB1B2*-, and *phyAB1B2*-mutant plants grown under ambient- (AT, day/night 24°C/18°C; blue fill) and high-temperature (HT, day/night 30°C/24°C; yellow fill) conditions. (B) Center: schematic model of the lycopene biosynthetic pathway, the dotted line represents more than one enzymatic step. Left: levels of lycopene, phytoene, and phytofluene in ripe fruits. AT: blue background; HT: yellow background. Each bar represents mean ± SE. Different letters indicate statistically significant differences according to Fisher's multiple comparison test (p < 0.05). Right: Relative mRNA levels of carotenoid biosynthetic genes GGPS2, PSYI, and PDS in fruits at mature green (MG) and breaker (BK) stages harvested from plants grown under AT (blue) and HT (vellow) conditions. Transcripts levels are expressed relative to MM MG – AT condition. Asterisks (* p < 0.05, ** p < 0.01) indicate differences in the analysis of variance in a multiple comparison test within the same fruit stage. Metabolites and genes are denoted according to the following abbreviations: GGDP, geranylgeranyl GGPS2, diphosphate; GERANYLGERANYL PHOSPHATE SYNTHASE 2; PSY1, PHYTOENE SYNTHASE 1; PDS, PHYTOENE DESATURASE. (C) Relative mRNA levels of PIF1a carotenogenesis regulator gene in fruits at mature green (MG) and breaker (BK) stages harvested from plants grown under AT (blue) and HT (yellow) conditions. Transcripts levels are expressed relative to MM MG - AT conditions. Asterisks (** p<0.01) indicate differences in the analysis of variance in a multiple comparison test within the same fruit stage.

671672

673

674

675

676

677

678

679

680

681

682

683

653

654

655

656

657

658

659

660

661

662

663 664

665

666

667

668

669

670

Figure 6. Fruit-localized PHYA and PHYB2 are involved in temperature perception impacting lycopene synthesis and master fruit ripening regulators. (A) Lycopene levels quantified in ripe fruits from Micro-Tom (MT) control genotype and fruit-specific *PHYA-(PHYA*^{RNAi)} and *PHYB2-(PHYB2*^{RNAi)} knockdown transgenic lines grown under ambient-(24°C/18°C) and high-temperature (30°C/24°C) conditions. Lycopene levels were quantified and expressed relative to MT fruits under AT conditions and values are means of at least three biological replicates from two independent lines for each genotype. Each bar represents mean \pm SE. (B,C) Relative mRNA levels of (B) *PSYI* (C) and master fruit ripening regulator genes in MT, *PHYA*^{RNAi}, and *PHYB2*^{RNAi} breaker fruit samples harvested under AT and HT conditions. Expression levels are relative to MT – AT conditions. n = at least three biological replicates. Each bar represents mean \pm SE. Different letters indicate statistically significant differences according to Fisher's multiple comparison test (p<

684	0.05). RIN; RIPENING INHIBITOR, NOR; NON-RIPENING, FUL1; FRUITFULL1, FUL2;
685	FRUITFULL2, AP2a; APETALA2a, TAGL1; TOMATO AGAMOUS-LIKE1, PSY1;
686	PHYTOENE SYNTHASE 1.
687	
688	Figure 7. The effect of PHY-mediated temperature perception on tomato metabolism
689	regulation. The rise of ambient temperature shifts balance to the inactive phytochrome (Pr)
690	form, which promotes the Chl degradation pathway in source leaves through the
691	transcriptional up-regulation of Chl catabolic enzyme-associated genes. Additionally,
692	reduced levels of Pfr impair carotenoid accumulation in both leaves and ripe fruits, through
693	the transcriptional downregulation of carotenoid biosynthetic and master ripening regulator
694	genes.
695	
696	

Parsed Citations

Alba R, Cordonnier-Pratt MM, Pratt LH. (2000a). Fruit-localized phytochromes regulate lycopene accumulation independently of ethylene production in tomato. Plant Physiology 123: 363–370.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Alba R, Kelmenson PM, Cordonnier-Pratt MM, Pratt LH. (2000b). The phytochrome gene family in tomato and the rapid differential evolution of this family in angiosperms. Molecular Biology and Evolution 17: 362–373.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Almeida J, Asís R, Molineri VN, Sestari I, Lira BS, Carrari F, Peres LEP, Rossi M. (2015). Fruits from ripening impaired, chlorophyll degraded and jasmonate insensitive tomato mutants have altered tocopherol content and composition. Phytochemistry 111: 72–83.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Alves FRR, Lira BS, Pikart FC, Monteiro SS, Furlan CM, Purgatto E, Pascoal GB, Andrade SCS, Demarco D, Rossi M, Freschi L. (2020). Beyond the limits of photoperception: constitutively active PHYTOCHROME B2 overexpression as a means of improving fruit nutritional quality in tomato. Plant Biotechnology Journal https://doi.org/10.1111/pbi.13362.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Ayenan MAT, Danquah A, Hanson P, Ampomah-Dwamena C, Sodedji FAK, Asante IK, Danquah EY. (2019) Accelerating Breeding for Heat Tolerance in Tomato (Solanum lycopersicum L.: An Integrated Approach. Agronomy 9: 720.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Bianchetti RE, Cruz AB, Oliveira BS, Demarco D, Purgatto E, Peres LEP, Rossi M, Freschi L. (2017). Phytochromobilin deficiency impairs sugar metabolism through the regulation of cytokinin and auxin signaling in tomato fruits. Scientific Reports 7: 7822.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Bianchetti RE, Lira BS, Monteiro SS, Demarco D, Purgatto E, Rothan C, Rossi M, Freschi L. (2018). Fruit-localized phytochromes regulate plastid biogenesis, starch synthesis, and carotenoid metabolism in tomato. Journal of Experimental Botany 69: 3573–3586.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Bita CE, Gerats T. (2013). Plant tolerance to high temperature in a changing environment: scientific fundamentals and production of heat stress-tolerant crops. Frontiers in Plant Science 4: 273.

Pubmed: Author and Title

Google Scholar: <u>Author Only Title Only Author and Title</u>

Box MS, Huang BE, Domijan M, Jaeger KE, Khattak AK, Yoo SJ, Sedivey EL, Jones DM, Hearn TJ, Webb ARR, Grant A, Locke JCW, Wigge PA (2015). ELF3 controls thermoresponsive growth in Arabidopsis. Current Biology 25: 194–199.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Burgie ES, Vierstra RD. (2014). Phytochromes: An Atomic Perspective on Photoactivation and Signaling. 26: 4568–4583.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Campos ML, Carvalho RF, Benedito VA, Peres LEP. (2010). The Micro-Tom model system as a tool to discover novel hormonal functions and interactions. Plant Signaling & Behavior 5: 267-270.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Chung MY, Vrebalov J, Alba R, Lee J, McQuinn R, Chung JD, Klein P, Giovannoni J. (2010). Atomato (Solanum lycopersicum) APETALA2/ERF gene, SIAP2a, is a negative regulator of fruit ripening. Plant Journal 64: 936–947.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Cordoba E, Salmi M, León P. (2009). Unravelling the regulatory mechanisms that modulate the MEP pathway in higher plants. Journal of Experimental Botany 60: 2933–2943.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Cruz AB, Bianchetti RE, Alves FRR, Purgatto E, Peres LEP, Rossi M, Freschi L. 2018. Light, Ethylene and Auxin Signaling Interaction Regulates Carotenoid Biosynthesis During Tomato Fruit Ripening. Frontiers in Plant Science 9: 1370.

Pubmed: Author and Title

Google Scholar: <u>Author Only Title Only Author and Title</u>

Dubreuil C, Ji Y, Strand Å, Grönlund A (2017). A quantitative model of the phytochrome-PIF light signalling initiating chloroplast development. Scientific Reports 7: 13884\to aded from on May 14, 2020 - Published by www.plantphysiol.org
Copyright © 2020 American Society of Plant Biologists. All rights reserved.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Expósito-Rodríguez M, Borges AA, Borges-Pérez A, Pérez JA (2008). Selection of internal control genes for quantitative real-time RT-PCR studies during tomato development process. BMC Plant Biology 8: 131.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Fahad S, Bajwa AA, Nazir U, Anjum SA, Farooq A, Zohaib A, Sadia S, Nasim W, Adkins S, Saud S, Ihsan MZ, Alharby H, Wu C, Wang D, Huang J. (2017). Crop Production under Drought and Heat Stress: Plant Responses and Management Options. Frontiers in Plant Science. 8: 1147.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Fraser PD, Pinto M, Holloway DE, Bramley P. (2000). Application of high-performance liquid chromatography with photodiode array detection to the metabolic profiling of plant isoprenoids. The Plant Journal 24: 551-558.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Fujii Y, Tanaka H, Konno N, Ogasawara Y, Hamashima N, Tamura S, Hasegawa S, Hayasaki Y, Okajima K, Kodama T. (2017). Phototropin perceives temperature based on the lifetime of its photoactivated state. Proceedings of the National Academy of Sciences 114: 9206–9211.

Pubmed: <u>Author and Title</u>

Google Scholar: Author Only Title Only Author and Title

Gramegna G, Rosado D, Carranza APS, Cruz AB, Simon-Moya M, Llorente B, Rodríguez-Concepción, M, Freschi L, Rossi M. (2018). PHYTOCHROME - INTERACTING FACTOR 3 mediates light - dependent induction of tocopherol biosynthesis during tomato fruit ripening. Plant Cell & Environment 42: 1328–1399.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Gupta SK, Sharma S, Santisree P, Kilambi HV, Appenroth K, Sreelakshmi Y, Sharma R. (2014). Complex and shifting interactions of phytochromes regulate fruit development in tomato. Plant, Cell and Environment 37: 1688–1702.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Guyer L, Hofstetter SS, Christ B, Lira BS, Rossi M, Hörtensteiner S. (2014). Different Mechanisms Are Responsible for Chlorophyll Dephytylation during Fruit Ripening and Leaf Senescence in Tomato. Plant Physiology 166: 44–56.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Hörtensteiner S. (2013). Update on the biochemistry of chlorophyll breakdown. Plant Molecular Biology 82: 505-517.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Huang H, McLoughlin KE, Sorkin ML, Burgie ES, Bindbeutel RK, Vierstra RD, Nusinow DA (2019). PCH1 regulates light, temperature, and circadian signaling as a structural component of phytochrome B-photobodies in Arabidopsis. Proceedings of the National Academy of Sciences 116: 8603–8608.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Inagaki N, Kinoshita K, Kagawa T, Tanaka A, Ueno O, Shimada H, Takano M. (2015). Phytochrome B Mediates the Regulation of Chlorophyll Biosynthesis through Transcriptional Regulation of ChlH and GUN4 in Rice Seedlings. PLoS ONE 10: 10.1371/journal.pone.0135408.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Ishida A, Toma T, Marjenah M. (2000) Leaf gas exchange and canopy structure under wet and drought years in Macaranga conifera, a tropical pioneer tree. Rainforest Ecosystems of East Kalimantan 140:129-42.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Jung JH, Domijan M, Klose C, Biswas S, Ezer D, Gao M, Khattak AK, Box MS, Charoensawan V, Cortijo S, Kumar M, Grant A, Locke JC, Schäfer E, Jaeger KE, Wigge PA (2016). Phytochromes function as thermosensors in Arabidopsis. Science 354: 886–889.

Pubmed: Author and Title

Google Scholar: <u>Author Only Title Only Author and Title</u>

Karlova R, Rosin FM, Busscher-Lange J, Parapunova V, Do PT, Fernie AR, Fraser PD, Baxter C, Angenent GC, de Maagd RA (2011). Transcriptome and Metabolite Profiling Show That APETALA2a Is a Major Regulator of Tomato Fruit Ripening. The Plant Cell 23: 923–941.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Kerckhoffs LHJ, Kelmenson PM, Schreuder MEL, Kendrick CI, Kendrick RE, Hanhart CJ, Koornneef M, Pratt LH, Cordonnier-Pratt MM.

Downloaded from on May 14, 2020 - Published by www.plantphysiol.org

Copyright © 2020 American Society of Plant Biologists. All rights reserved.

(1999). Characterization of the gene encoding the apoprotein of phytochrome B2 in tomato, and identification of molecular lesions in two mutant alleles. Molecular and General Genetics 261: 901–907.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Kerckhoffs LHJ, Schreuder MEL, VanTuinen A, Koornneef M, Kendrick RE. (1997). Phytochrome control of anthocyanin biosynthesis in tomato seedlings: Analysis using photomorphogenic mutants. Photochemistry and Photobiology 65: 374–381.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Kerckhoffs LHJ, Van Tuinen A, Hauser BA, Cordonnier Pratt MM, Nagatani A, Koornneef M, Pratt LH, Kendrick RE. (1996). Molecular analysis of tri-mutant alleles in tomato indicates the Tri locus is the gene encoding the apoprotein of phytochrome B1. Planta 199: 152–157.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Kim K, Jeong J, Kim J, Lee N, Kim ME, Lee S, Kim SC, Choi G. (2016). PIF1 regulates plastid development by repressing photosynthetic genes in the endodermis. Mol. Plant 9: 1415–1427

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Klee HJ, Giovannoni JJ. (2011). Genetics and Control of Tomato Fruit Ripening and Quality Attributes. Annual Review of Genetics 45: 41–59.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Koini MA, Alvey L, Allen T, Tilley CA, Harberd NP, Whitelam GC, Franklin KA (2009). Report High Temperature-Mediated Adaptations in Plant Architecture Require the bHLH Transcription Factor PIF4. Current Biology 19: 408–413.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Lazarova GI, Kerckhoffs LHJ, Brandstädter J, Matsui M, Kendrick RE, Cordonnier-Pratt MM, Pratt LH. (1998a). Molecular analysis of PHYA in wild-type and phytochrome A-deficient mutants of tomato. Plant Journal 14: 653–662.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Lazarova GI, Kubota T, Frances S, Peters JL, Hughes MJ, Brandstadter J, Széll M, Matsui M, Kendrick RE, Cordonnier-Pratt MM, Pratt LH. (1998b). Characterization of tomato PHYB1 and identification of molecular defects in four mutant alleles. Plant Molecular Biology 38: 1137–1146.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Legris M, Klose C, Burgie ES, Rojas CCR, Neme M, Hiltbrunner A, Wigge PA, Schäfer E, Vierstra RD, Casal JJ. (2016). Phytochrome B integrates light and temperature signals in Arabidopsis. Science 354: 897–900.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Legris M, Nieto C, Sellaro R, Prat S, Casal JJ. (2017). Perception and signalling of light and temperature cues in plants. Plant Journal 90: 683–697.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Lira BS, Gramegna G, Trench B, Alves FRR, Silva EM, Silva GFF, Thirulmalaikumar VP, Lupi ACD, Demarco D, Purgatto E, Nogueira FTS, Balazadeh S, Freschi L, Rossi M. (2017). Manipulation of a senescence-associated gene improves fleshy fruit yield. Plant Physiology 175: 77-91.

Pubmed: Author and Title

Google Scholar: <u>Author Only Title Only Author and Title</u>

Lira BS, de Setta N, Rosado D, Almeida J, Freschi L, Rossi M. (2014). Plant degreening: Evolution and expression of tomato (Solanum lycopersicum) dephytylation enzymes. Gene 546: 359–366.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Liu L, Shao Z, Zhang M, Wang Q. (2015). Regulation of carotenoid metabolism in tomato. Molecular Plant 8: 28-39.

Pubmed: Author and Title

Google Scholar: <u>Author Only Title Only Author and Title</u>

Llorente B, D'Andrea L, Ruiz-Sola MA, Botterweg E, Pulido P, Andilla J, Loza-Alvarez P, Rodriguez-Concepcion M. (2016). Tomato fruit carotenoid biosynthesis is adjusted to actual ripening progression by a light-dependent mechanism. Plant Journal 85: 107–119.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Lorrain S, Allen T, Duek PD, Whitelam GC, Fankhauser C. (2008). Phytochrome-mediated inhibition of shade avoidance involves degradation of growth-promoting bHLH transcription factors. Plant Journal 53: 312–323.

Downloaded from on May 14, 2020 - Published by www.plantphysiol.org

Downloaded from on May 14, 2020 - Published by www.plantphysiol.org Copyright © 2020 American Society of Plant Biologists. All rights reserved. Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Ma D, Li X, Guo Y, Chu J, Fang S, Yan C, Noel JP, Liu H. (2016). Cryptochrome 1 interacts with PIF4 to regulate high temperature-mediated hypocotyl elongation in response to blue light. Proceedings of the National Academy of Sciences 113: 224–229.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Martin G, Leivar P, Ludevid D, Tepperman JM, Quail PH, Monte E. (2016). Phytochrome and retrograde signalling pathways converge to antagonistically regulate a light-induced transcriptional network. Nature Communications 7: 11431.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Martínez-García JF, Huq E, Quail PH. (2000). Direct Targeting of Light Signals to a Promoter. Science 288: 859-859.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Nakamura H, Muramatsu M, Hakata M, Ueno O, Nagamura Y, Hirochika H, Takano M, Ichikawa H, (2009). Ectopic overexpression of the transcription fator OsGLK1 induces chloroplast development in non-green rice cells. Plant Cell Physiology 50: 1933-1949.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Nguyen C V, Vrebalov JT, Gapper NE, Zheng Y, Zhong S, Fei Z, Giovannoni JJ. (2014). Tomato GOLDEN2-LIKE Transcription Factors Reveal Molecular Gradients That Function during Fruit Development and Ripening. Plant Cell 26: 585–601.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Oh S, Montgomery BL. (2014). Phytochrome-dependent coordinate control of distinct aspects of nuclear and plastid gene expression during anterograde signaling and photomorphogenesis. Frontiers in Plant Science 5: 171.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Park E, Kim Y, Choi G. (2018). Phytochrome B Requires PIF Degradation and Sequestration to Induce Light Responses across a Wide Range of Light Conditions. Plant Cell 30: 1277–1292.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Qiu Y, Li M, Kim RJ, Moore CM, Chen M. (2019). Daytime temperature is sensed by phytochrome. Nature Communications 10: 140.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Quadrana L, Almeida J, Otaiza SN, Duffy T, Corrêa da Silva JV, de Godoy F, Asís R, Bermúdez L, Fernie AR, Carrari F, Rossi M. (2013). Transcriptional regulation of tocopherol biosynthesis in tomato. Plant Molecular Biology 81: 309–325.

Pubmed: <u>Author and Title</u>

Google Scholar: Author Only Title Only Author and Title

Reynolds ES. (1963). The use of lead citrate at high pH as an electron-opaque stain in electron microscopy. The Journal of Cell Biology 17: 208–212.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Rockwell NC, Su Y-S, Lagarias JC. (2006). Phytochrome Structure and Signaling Mechanisms. Annual Review of Plant Biology 57: 837–858.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Rosado D, Gramegna G, Cruz A, Lira BS, Freschi L, De Setta N, Rossi M. (2016). Phytochrome Interacting Factors (PIFs) in Solanum lycopersicum: Diversity, evolutionary history and expression profiling during different developmental processes. PLoS ONE 11: 10.1371/journal.pone.0165929.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Rosado D, Trench B, Bianchetti R, Zuccarelli R, Alves FRR, Purgatto E, Floh ElS, Nogueira FTS, Freschi L, Rossi M. (2019). Downregulation of PHYTOCHROME-INTERACTING FACTOR 4 influences plant development and fruit production. Plant Physiology 181: 1360-1370.

Pubmed: Author and Title

Google Scholar: <u>Author Only Title Only Author and Title</u>

Saidi Y, Finka, A, Goloubinoff, P. (2017). Heat perception and signalling in plants: a tortuous path to thermotolerance. New Phytologist 190: 556-565.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Sakuraba Y, Jeong J, Kang MY, Kim J, Paek NC, Choi G. (2014). Phytochrome-interacting transcription factors PIF4 and PIF5 induce leaf senescence in Arabidopsis. Nature Control of Copyright 2020 American Society of Plant Biologists. All rights reserved.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Sharrock RA, Quail PH. (1989). Novel phytochrome sequences in Arabidopsis thaliana: structure, evolution, and differential expression of a plant regulatory photoreceptor family. Genes & Development 3: 1745–1757.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Sims DA, Gamon JA (2003). Estimation of vegetation water content and photosynthetic tissue area from spectral reflectance: a comparison of indices based on liquid water and chlorophyll absorption features. Remote Sensing of Environment 84: 526–537.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Song Y, Yang C, Gao S, Zhang W, Li L, Kuai B. (2014). Age-triggered and dark-induced leaf senescence requires the bHLH transcription factors PIF3,4, and 5. Molecular Plant 7: 1776-1787.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Spicher L, Almeida J, Gutbrod K, Pipitone R, Dörmann P, Glauser G, Rossi M, Kessler F. (2017). Essential role for phytol kinase and tocopherol in tolerance to combined light and temperature stress in tomato. Journal of Experimental Botany 68: 5845–5856.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Spicher L, Glauser G, Kessler F. (2016). Lipid Antioxidant and Galactolipid Remodeling under Temperature Stress in Tomato Plants. Frontiers in Plant Science 7: 167.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Stephenson PG, Fankhauser C, Terry MJ. (2009). PIF3 is a repressor of chloroplast development. Proceedings of the National Academy of Sciences 106: 7654–7659.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Suwa R, Hakata H, Hara H, El-Shemy HA, Adu-Gyamfi JJ, Nguyen NT, Kanai S, Lightfoot DA, Mohapatra PK, Fujita K. (2010). High temperature effects on photosynthate partitioning and sugar metabolism during ear expansion in maize (Zea mays L.) genotypes. Plant Physiology and Biochemistry 48: 124–130.

Pubmed: <u>Author and Title</u>

Google Scholar: Author Only Title Only Author and Title

Takahashi S, Murata N. (2008). How do environmental stresses accelerate photoinhibition? Trends in Plant Science 13: 178-182.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Toledo-Ortiz G, Johansson H, Lee KP, Bou-torrent J, Stewart K, Steel G, Rodríguez-Concepción M, Halliday KJ. (2014). The HY5-PIF Regulatory Module Coordinates Light and Temperature Control of Photosynthetic Gene Transcription. PLoS Genetics 10: 10.1371/journal.pgen.1004416.

Pubmed: Author and Title

Google Scholar: <u>Author Only Title Only Author and Title</u>

Tomato Genome Consortium. (2012). The tomato genome sequence provides insights into fleshy fruit evolution. Nature 485: 635-641.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Van Eerden FJ, De Jong DH, De Vries AH, Wassenaar TA, Marrink SJ. (2015). Characterization of thylakoid lipid membranes from cyanobacteria and higher plants by molecular dynamics simulations. Biochimica et Biophysica Acta - Biomembranes 1848: 1319–1330.

Pubmed: Author and Title

Google Scholar: <u>Author Only Title Only Author and Title</u>

Velikova V, Varkonyi Z, Szabo M, Maslenkova L, Nogues I, Kóvacs L, Peeva V, Busheva M, Garab G, Sharkey TD, Loreto F. (2011). Increased Thermostability of Thylakoid Membranes in Isoprene-Emitting Leaves Probed with Three Biophysical Techniques. Plant Physiology 157: 905–916.

Pubmed: <u>Author and Title</u>

Google Scholar: Author Only Title Only Author and Title

Watson ML. (1958). Staining of Tissue Sections for Electron Microscopy with Heavy Metals: II. Application of Solutions Containing Lead and Barium. The Journal of Cell Biology 25: 727–730.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Wellburn A (1994). The spectral determination of chlorophylls a and b, as well as total carotenoids, using various solvents with spectrophotometers of different resolution. Journal of Plant Physiology 144: 307-313.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Weller JL, Schreuder MEL, Smith H, Moormagef MnKendrick ₹50(2000) bRhysiological interportions of phytochromes A, B1 and B2 in the Copyright © 2020 American Society of Plant Biologists. All rights reserved.

control of development in tomato. Plant Journal 24: 345-356.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Yamori W, von Caemmerer S. (2009). Effect of Rubisco Activase Deficiency on the Temperature Response of CO2 Assimilation Rate and Rubisco Activation State: Insights from Transgenic Tobacco with Reduced Amounts of Rubisco Activase. Plant Physiology 151: 2073–2082.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Yuan XY, Wang RH, Zhao XD, Luo YB, Fu DQ. (2016). Role of the tomato Non-Ripening mutation in regulating fruit quality elucidated using iTRAQ protein profile analysis. PLoS ONE 11: 10.1371/journal.pone.0164335.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Zhang Y, Liu Z, Chen Y, He, J, Bi Y. (2015). PHYTOCHROME-INTERACTING FACTOR 5 (PIF5) positively regulates dark-induced senescence and chlorophyll degradation in Arabidopsis. Plant Science 237: 57-68.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Zhang Y, Mayba O, Pfeiffer A, Shi H, Tepperman JM, Speed TP, Quail PH. (2013). A Quartet of PIF bHLH Factors Provides a Transcriptionally Centered Signaling Hub That Regulates Seedling Morphogenesis through Differential Expression-Patterning of Shared Target Genes in Arabidopsis. PLoS Genetics 9: 10.1371/journal.pgen.1003244.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Zhang R, Sharkey TD. (2009). Photosynthetic electron transport and proton flux under moderate heat stress. Photosynthesis Research 100: 29-43.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Zhao C, Liu B, Piao S, Wang X, Lobell DB, Huang Y, Huang M, Yao Y, Bassu S, Ciais P, Durand JL, Elliott J, Ewert F, Janssens IA, Li T, Lin E, Liu Q, Martre P, Müller C, Peng S, Peñuelas J, Ruane AC, Wallach D, Wang T, Wu D, Liu Z, Zhu Y, Zhu Z, Asseng S. (2017). Temperature increase reduces global yields of major crops in four independent estimates. Proceedings of the National Academy of Sciences 114: 9326–9331.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Zhao F, Zhang D, Zhao Y, Wang W, Yang H, Tai F, Li C, Hu X. (2016). The Difference of Physiological and Proteomic Changes in Maize Leaves Adaptation to Drought, Heat, and Combined Both Stresses. Frontiers in Plant Science 7: 1471.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Zhou Y, Xun Q, Zhang D, Lv M, Ou Y, Li J. (2019). TCP Transcription Factors Associate with PHYTOCHROME INTERACTING FACTOR 4 and CRYPTOCHROME 1 to Regulate Thermomorphogenesis in Arabidopsis thaliana. iScience 15: 600-610.

Pubmed: Author and Title

Google Scholar: Author Only Title Only Author and Title

Competing interests

The authors declare no competing interests.