

1 Shedding light on the Methylerythritol phosphate (MEP)-pathway: long

- 2 hypocothyl 5 (HY5)/ phytochrome-interacting factors (PIFs) transcription
- 3 factors modulating key limiting steps.
- 4
- 5 Marel Chenge-Espinosa<sup>1</sup>, Elizabeth Cordoba<sup>1</sup>, Cynthia Romero-Guido<sup>1</sup>,
- 6 Gabriela Toledo-Ortiz <sup>2\*</sup> and Patricia León<sup>1\*</sup>
- 7
- 8 <sup>1</sup> Departamento de Biología Molecular de Plantas, Instituto de Biotecnología,
- 9 Universidad Nacional Autónoma de México, Av. Universidad # 2001, Col.
- 10 Chamilpa, Cuernavaca, Morelos, México, C.P. 62210.
- 11 <sup>2</sup> Lancaster University, Lancaster Environment Centre, Lancaster LA1 4YQ, UK
- 12
- 13 \*Corresponding authors:
- 14 Patricia León
- 15 Telephone +52 (55) 5622 7856 Fax: +52 (777) 313 9988
- 16 email: patricia@ibt.unam.mx
- 17 Gabriela Toledo-Ortiz
- 18 Telephone: +44 (0) 1524 595053
- 19 Email: g.toledo-ortiz@lancaster.ac.uk
- 20

Total word count	6998
Summary:	219
Introduction:	949
Results:	2673
Discussion:	1338
Materials and Methods:	723
Acknowledgements:	57
Figure Legends	1036
Figures:	7 (Figs. 1 and 7 in color)
Supporting Information	7 (Fig. S1-S6; Table S1)

21

Formatted: Spanish (Spain, Traditional Sort)

Formatted: Spanish (Spain, Traditional Sort)

#### Significance statement

1 2

- 3
- 4 Light imposes a direct, rapid and potentially multi-faceted effect that leads to
- 5 unique protein dynamics to the main flux-limiting steps of the MEP pathway, a key
- 6 route essential for plants. Through differential direct transcriptional interaction, the
- 7 key-master integrators of light signals HY5 and PIFs, target the main flux-limiting
- 8 steps of the pathway. Our work illustrates how light signals can impose contrasting
- 9 dynamics over a key pathway whose products multi-branch downstream to all
- 10 chloroplastic isoprenoids.

#### 1 Summary

2 The plastidial methylerythritol phosphate (MEP) pathway is an essential route for 3 plants as the source of precursors for all plastidial isoprenoids, many of which are 4 of medical and biotechnological importance. The MEP-pathway is highly sensitive 5 to environmental cues as many of these compounds are linked to photosynthesis and growth and light is one of the main regulatory factors. However, the 6 7 mechanisms coordinating the MEP-pathway with light cues are not fully 8 understood. 9 Here we demonstrate that by a differential direct transcriptional modulation via the 10 key master integrators of light signal transduction HY5 and PIFs that target the genes that encode the rate-controlling DXS1, DXR and HDR enzymes, light 11 12 imposes a direct, rapid and potentially multi-faceted response that leads to unique protein dynamics of this pathway resulting in up to 10-fold difference in the protein 13 14 levels. For DXS1, PIF1/HY5 act as a direct activation/suppression module. In 15 contrast, DXR accumulation in response to light results from HY5 induction with minor contribution of de-repression by PIF1. Finally, HDR transcription increases in 16 17 the light exclusively by suppression of the PIFs repression. This is an example of 18 how light signaling components can differentially multi-target the initial steps of a 19 pathway whose products branch downstream to all chloroplastic isoprenoids. These findings demonstrate the diversity and flexibility of light signaling 20 components that optimize key biochemical pathways essential for plant growth. 21 22 23 Keywords: 24 25 Isoprenoids, MEP-pathway, Light responses, DXS1, DXR and HDR 26 enzymes, Phytochrome interacting factors, Long Hypocotyl 5, Rate-limiting

- 27 enzymes, Arabidopsis thaliana.
- 28

#### 1 Introduction

- 2 3 Isoprenoids constitute a family of natural products synthesized in all 4 organisms with diverse function (Chappell, 1995). Isoprenoids are essential for 5 plant development, participating in several key processes such as photosynthesis, respiration and general plant growth (Bouvier et al., 2005). All isoprenoids are 6 7 produced from the condensation of two universal five-carbon precursors, 8 isopentenyl diphosphate (IPP) and its isomer dimethylallyl diphosphate (DMAPP). 9 Condensation of these basic units gives rise to isoprenoid diversity. 10 Plants synthesize IPP and DMAPP by two non-related pathways present in 11 12 different compartments. The cytoplasmic mevalonic pathway uses acetyl-CoA via mevalonic acid for the synthesis of IPP and DMAPP, and the methyl-D-erythritol 4-13 14 phosphate (MEP) pathway takes place in plastids (Croteau et al., 2000, Hemmerlin 15 et al., 2012, Rodriguez-Concepcion and Boronat, 2015). The MEP pathway uses pyruvate and D-glyceraldehyde 3-phosphate (GAP) for IPP and DMAPP synthesis 16 17 through the activity of seven consecutive enzymes (Phillips et al., 2008). The first 18 step of the MEP pathway is catalyzed by 1-deoxy-D-xylulose 5-phosphate 19 synthase (DXS) that produces 1-deoxy-D-xylulose 5-phosphate (DXP). DXP is 20 rearranged into MEP by the action of the 1-deoxy-D-xylulose 5-phosphate reductoisomerase (DXR), this is the first committed step of the pathway. 21 22 Subsequently MEP is converted to 1-hydroxy-2-methyl-2-(E)-butenyl 4-23 diphosphate (HMBPP) by four additional enzymatic steps and in the final step 24 HMBPP is converted into a mixture of IPP and DMAPP via the HMBPP reductase 25 (HDR) enzyme (Eisenreich et al., 2004). 26 27 The MEP pathway is present in eubacteria, plastids and the apicoplast of

- 28 apicomplexan but is absent in other eukaryotes, including humans (Lange et al.,
- 29 2000). Thus, the MEP pathway is considered an attractive target for development
- 30 of new antibacterial and antiparasitic drugs, and herbicides (Rodríguez-
- 31 Concepción, 2004, Rohdich et al., 2005). In plants the MEP pathway is responsible

1 for the production of essential compounds, such as the precursors of 2 photosynthetic pigments (carotenoids and the side chain of chlorophyll), 3 tocopherols and plastoquinones, hormones including gibberellins, abscisic acid 4 and strigolactone and a variety of monoterpenes, diterpenes and some 5 sesquiterpenes (Bouvier et al., 2005, Umehara et al., 2008). Recent studies have reported that the MEP pathway is also essential for the production of stress-6 7 specific retrograde signaling molecules (de Souza et al., 2017). Thus, in plants the 8 MEP pathway is essential for plant development and also is an important target for 9 biotechnological manipulation. 10 The enzymes in the MEP pathway are subject to modulation at different 11 12 levels, according to the developmental status of the plant and the fluctuating environmental conditions (Guevara-García et al., 2005, Rodriguez-Concepcion, 13 14 2006, Cordoba et al., 2009, Banerjee and Sharkey, 2014). This multi-level dynamic is critical to ensure the supply of IPP and DMAPP precursors with the demand of 15 downstream pathways, many of which are dependent of diurnal and light 16 17 conditions, such as the biosynthesis of chlorophylls and carotenoids (Ruiz-Sola 18 and Rodriguez-Concepcion, 2012). In plants, the plastid-localized enzymes of the 19 MEP pathway come from nuclear-encoded genes. Experimental evidence has 20 demonstrated that the transcript levels of all genes in the pathway are induced during plant development (Guevara-García et al., 2005, Meier et al., 2011), during 21 22 tomato fruit ripening (Lois et al., 2000) and in response to hormones (Oudin et al., 23 2007). Also, biotic interactions (Walter et al., 2000), circadian clock (Cordoba et al., 24 2009, Vranova et al., 2013) and light modulate the MEP transcripts levels in 25 several plant species (Hsieh and Goodman, 2005, Cordoba et al., 2009). Furthermore, post-transcriptional regulation of DXS, one of the rate-limiting steps 26 27 of the pathway, leads to changes in its protein accumulation and enzymatic activity (Guevara-García et al., 2005, Wright et al., 2014). 28 29 30 Light constitutes one of the most critical environmental signals for plant

31 development. From the emergence of the germinating seedling, light changes the

developmental program from skotomorphogenesis to photomorphogenesis (Chen 1 2 et al., 2004). Upon light exposure etioplasts rapidly differentiate into chloroplasts 3 upregulating many genes involved in photosynthesis and other plastid biosynthetic 4 pathways (Jiao et al., 2007). Complex photoreceptor systems allow plants to adjust 5 diverse processes in response to variable light conditions (Franklin and Quail, 2010). The phytochrome photoreceptors fine-tune plant photomorphogenesis in 6 7 response to Red and Far Red light. Mechanistically, light-activated nuclear-8 phytochromes bind directly to members of the bHLH family of phytochrome-9 interacting factors (PIFs), promoting their degradation. PIFs act as important 10 repressors of photomorphogenic development in the dark and key signal integrators (Leivar and Quail, 2011, Leivar and Monte, 2014). In addition, active 11 12 phytochromes prevent the degradation of activators of photomorphogenesis such as ELONGATED HYPOCOTHYL 5 (HY5) and its close homolog HYH. These bZIP 13 14 transcription factors participate in the up-regulation of a variety of genes in response to light and display antagonistic functions to the PIFs in the expression of 15 diverse genes (Tepperman et al., 2001). 16 17 18 Given the strategic role that light has over isoprenoid production in the 19 present study we investigate the mechanisms coordinating the MEP-pathway with light cues. We examined the role of light signaling components PIFs, HY5 and 20 HYH in modulating the expression and imposing a differential and specific dynamic 21 22 on DXS1, DXR and HDR genes encoding for flux-controlling enzymes of the MEP 23 pathway (Estévez et al., 2001, Mahmoud and Croteau, 2001, Botella-Pavía et al., 24 2004, Carretero-Paulet et al., 2006). Transcript analyses and ChIP assays 25 confirmed that HY5 and PIFs are direct regulators of the light-modulated expression of these genes. We show that the function of these factors in tuning the 26 27 MEP-pathway gene expression extends to different developmental stages 28 potentially diversifying the synthesis of multiple compounds of isoprenoid origin, 29 essential for plant growth and development at different stages of the plant life

30 cycle. Furthermore, we demonstrate using protein accumulation studies that the

31 transcriptional regulation mediated by PIF and HY5 impacts the accumulation of

1 DXS1 and HDR enzymes but not DXR. This analysis exemplifies the diverse

2 mechanistic dynamics that the same master light regulators can impose to tune up

3 essential metabolic pathways in response to light.

4

## 5 Results

6

*Cis*-acting elements in the DXS promoter are responsible for the regulation of
 the DXS1 gene.

9 Previous studies have shown that in developing seedlings, transcripts of the 10 MEP pathway genes accumulate upon light exposure (Botella-Pavía et al., 2004, Hsieh and Goodman, 2005, Cordoba et al., 2009). To analyze if the light induction 11 12 response is mediated at the transcriptional level, 3 day-old etiolated transgenic lines containing 1510 bp upstream from the DXS1 gene ATG fused with the GUS 13 14 gene (Estévez et al., 2000) were exposed to light. GUS activity is detected in the cotyledons of dark grown seedlings (Figure 1a). However, after 6 h of light 15 exposure this staining expands to the hypocotyl (Figure 1d). Quantitative 16 17 determination of GUS activity confirmed an approximately 2-fold increase after light 18 exposure in comparison to dark control seedlings (Figure 1g). This result confirms 19 that in response to light, DXS1 transcript levels are transcriptionally up-regulated and also demonstrates that the *cis*-acting elements important for this response are 20 present within the 1.5 Kb upstream regulatory region of this gene. 21 22 23 To further delimit the region involved in the light response two additional 24 transgenic lines containing 750 (Figure b and e) and 670 bp (Figure c and f) 25 upstream from the DXS1 ATG of were generated, and their expression in response 26 to light was analyzed. We observed that the GUS expression in these lines is 27 induced upon light exposure at similar levels to the 1.5 Kb original fragment (Figure 28 1g). These results support that the elements responsible for light response in the 29 DXS1 gene localize within the 670 bp region upstream from the ATG (360 from the 30 transcription initiation site).

#### HY5 and HYH positively regulate transcription of MEP-pathway limiting DXS1 1 2 and DXR genes, but not of HDR. 3 The transcription factor HY5 plays a pivotal role as a positive regulator of 4 photomorphogenesis and greening responses through direct binding to the 5 promoters of diverse light-activated genes (Lee et al., 2007, Zhang et al., 2011). To determine whether HY5 contributes to the light induction of the DXS1 gene, its 6 7 transcript level was analyzed after illumination of dark-adapted wild-type and hy5 8 mutant seedlings; since a clear accumulation by light for MEP genes was observed 9 under these conditions (Cordoba et al., 2009). Seedlings were grown for 3 days 10 under 16 h light: 8h dark photoperiod and then transferred to darkness for 3 additional days (dark-adapted treatment) prior to illumination for 6 h. Northern blot 11 12 analysis shows that the transcript of DXS1 is significantly lower in the light hy5 mutant compared to the Col-0 wild-type counterpart (Figure 2a). In the hy5hyh 13 14 double mutant we observed further reduction of the DXS1 transcript supporting the notion that while minor, HYH also plays a positive role in the expression of DXS1 in 15 response to light. 16 17 18 In addition to DXS, two other enzymes, DXR and HDR, limit the flux through 19 the MEP-pathway (Mahmoud and Croteau, 2001, Botella-Pavía et al., 2004, Carretero-Paulet et al., 2006, Kim et al., 2009). We also analyzed the light 20 responsiveness of DXR and HDR transcripts in the hy5 and hy5hyh mutants. As 21 22 shown in Figure 2b the transcript of DXR accumulates upon light illumination in the 23 Col-0 wild-type plants. This induction is substantially lower in the hy5 mutant, 24 demonstrating that HY5 also acts positively on the expression of the DXR gene in 25 the light. Relative to hy5, the hy5hyh double mutant has no additive effect on the level of DXR transcript abundance, supporting the lack of involvement of HYH in 26 27 the light induced up-regulation of this gene. In contrast, the transcript accumulation 28 in response to light for HDR is marginal in the wild-type (Figure 2a, b) with no clear

- 29 role of HY5 and HYH (Figure 2c).
- 30

To further substantiate these results quantitative RT-PCR (gRT-PCR) on the 1 2 same RNAs samples was performed, confirming the positive role of HY5 on the 3 induction by light of DXS1 and DXR transcripts (Figure 2d, e) but not on HDR. 4 qRT-PCR data further corroborated that HYH does not appear to play a major role 5 in the modulation by light of these genes. As monochromatic red light maximizes the light response we analyzed the expression of these genes in the dark-adapted 6 7 seedlings illuminated with red light (Figure S1). This analysis corroborates an 8 increase of the HDR transcript by light and that HY5 does not play a major role in 9 this response. Red-light experiments also demonstrated that in the absence of 10 these bZIP transcription factors there is still some light responsiveness, supporting 11 the notion that additional unknown factors participate in this response. 12 13 PIFs are negative regulators of DXS1, DXR and HDR genes 14 For diverse light responses, PIF transcription factors act antagonistically to HY5 and HYH (Chen et al., 2013). To test whether PIFs play an opposing role to 15 HY5 in the light-mediated accumulation of the MEP pathway genes, the transcript 16 level of the DXS1, DXR and HDR were analyzed by northern blots in pif1, pif3, pif4 17 18 and pif5 mutants compared to Col-0 wild-type seedlings, using dark-adapted

19 treated seedlings (Leivar *et al.*, 2008). We observed that the transcript levels of the

20 DXS1 (Figure 3a), DXR (Figure 3b) and HDR (Figure 3c) genes are elevated in

21 several of the *pif* single mutants in the dark and in the light, supporting a function of

22 these transcription factors as transcriptional repressors of the expression of these

- 23 rate-limiting MEP pathway genes.
- 24

To further analyze the contribution of the different PIFs we performed qRT-PCR on the same samples illuminated with white light (Figure S2) or with monochromatic red light to maximize response (Figure 3). In the case of *DXS1*, increased transcript accumulation is observed in the *pif1* and *pif3* mutants with a more moderate increment in *pif5*, supporting the repressing function of these factors (Figure 3d and S2a). For *DXR*, transcript levels are higher in the dark and in light-exposed *pif1*, *pif3* and *pif5* mutants demonstrating that these three PIFs

contribute to the low accumulation level of this gene in wild-type plants (Figure 3e). 1 2 Finally, PIF1 appears to have the most significant effect on repressing HDR 3 transcript levels, with some contribution of PIF3 (Figure 3f). Collectively our data 4 demonstrate that the PIF factors in a partially redundant form are required to 5 maintain low expression levels of three key genes of the MEP pathway. 6 7 The functional redundancy of PIFs in regulating DXS1, DXR and HDR 8 genes is further illustrated with the quadruple pifQ mutant lacking PIF1, PIF3, PIF4 9 and PIF5 (Leivar et al., 2009). Because pifQ could suffer from photoxidative 10 damage caused by the overaccumulation of photochlorophyllide in the dark (Chen et al., 2013), pifQ analysis was done in 3 day-old seedlings grown in(the reported?) 11 12 photoperiod and dark-adapted for 36 h before exposure to 6 h of red light. We corroborated that in the pifQ mutant the expression level of DXS1 (Figure 3g), DXR 13 14 (Figure 3h) and HDR (Figure 3i) genes are significantly elevated in the dark and after light exposure compared to wild-type seedlings. Over all, our data 15 demonstrates that PIFs are important negative regulators of the three rate-limiting 16 17 MEP pathway genes. 18 19 HY5 and PIFs regulators modulate the expression of DXS, DXR and HDR 20 genes during de-etiolation It is known that PIFs display differential expression during distinct 21 22 developmental responses (Jeong and Choi, 2013). One of the physiologically 23 relevant responses to light is de-etiolation. Previous data demonstrated that during 24 de-etiolation PIF1 represses PSY gene expression that plays a limiting role in 25 carotenoid biosynthesis, but under the conditions used (1 h of induction) no regulation by PIF1 was detected for DXS1 (Toledo-Ortiz et al., 2010). Based on our 26 27 observations that light induces DXS, DXR and HDR expression at later time points, 28 we re-evaluated the contribution of PIFs and analyzed the role of HY5 by gRT-PCR 29 in 3 day-old etiolated seedlings exposed to red light for 6 h. Similar to the dark-30 adapted seedlings, the transcription factor HY5 was shown to act as an activator of 31 the DXS1 and DXR genes expression (Figure 4a and b). After light exposure the

level of the *DXS1* and *DXR* transcripts is lower in *hy5* compared to the wild-type.
 There was no significant change in the *HDR* expression level in *hy5* (Figure 4c),
 indicating that, at this stage,HY5 does not play a major role in the light response of
 this gene .

5

6 On the other hand, removal of the PIFs results in a significant increase in 7 the transcript levels of the DXS1, DXR and HDR genes at this developmental stage 8 compared to the wild-type seedlings (Figure 4). This result supports and extends 9 the partially redundant, repressive role of PIFs in down-regulating the expression of 10 the DXS1, DXR and HDR genes during de-etiolation. Similar to the dark-adapted seedlings PIF1, PIF3 and PIF5 contribute most significantly to the regulation of 11 12 these genes and we did not observe any major changes in the relative role of the individual PIFs at this developmental stage. In accordance the pifQ mutant 13 14 accumulates at higher transcript levels than the three genes in the dark and in the 15 light. 16 17 PIF1 and HY5 interact with the promoters of the MEP pathway genes in vivo 18 To investigate if the changes in the gene expression result from a direct 19 interaction of the PIFs and HY5 transcription factors with the promoters of the 20 DXS1, DXR and HDR genes, we conducted chromatin immunoprecipitation (ChIP) experiments. We selected as a model PIF1, considering that this factor plays a 21 22 major role in the regulation of the three MEP genes. ChIPs were carried out using

- 23 seedlings that express PIF1 fused to a myc-tag in a *pif1* background (TAP-PIF1)
- 24 (Moon *et al.*, 2008) or HY5 with an HA-tag in *hy5* background (HA-HY5) (Lee *et al.*,
- 25  $\,$  2007). Lines with a TAP-GFP or HA-GUS in a wild-type background were included
- 26 as negative controls for nonspecific binding of DNA to the tags used. Transgenic
- 27 HA-HY5 and TAP-PIF1 are mild-overexpressors, with similar protein expression
- 28 levels to endogenous HY5 and PIF1 and complement the mutant phenotypes in a
- 29 wide range of tested-light responses (Lee et al., 2007, Moon et al., 2008).
- 30

**Commented [HG1]:** Wasn't sure about this when looking at the plots. Certainly for DXS1 and DXR in the light this is true and also for DXS1 in the dark but pif1 looks to be greater than pifQ in DXR and for HDR pif3 in the dark and pif1 in the light. I could be missing something though.

To address if there is any difference in the interaction of these factors 1 2 depending on the developmental stage, the ChIP was conducted in dark-adapted 3 seedlings and etiolated seedlings, both exposed to 6 h of red light using myc or HA 4 antibodies. qRT-PCRs were done using specific primers for selected promoter 5 regions of each gene (Table S1, Figures S3, S5 and S6). 6 7 PIFs transcription factors bind to variants of the E boxes (CANNTG), 8 including the G-box (CACGTG) and the PBE-box (CACATG/CATGTG), or to 9 hexameric sequences G-box coupling elements (GCEs) containing the core 10 "ACGT" elements (Toledo-Ortiz et al., 2003, Zhang et al., 2013, Kim et al., 2016). The G-box, together with diverse GCE related motifs, such as the Z box 11 12 (ATACTGTGT), CA (GACGTA) and CG hybrids (GACGTG), have also been identified as the interaction site of HY5 (Lee et al., 2007, Toledo-Ortiz et al., 2010, 13 14 Zhang et al., 2011). The analysis of the 1.5 kb DXS1 promoter sequence using SOGO New PLACE software, showed no-presence of canonical G boxes. 15 However, several E-box related elements, including a PBE-box (CACATG), a CG 16 17 hybrid box (GACGTG), and a GCE element were found (Figure S3). Four specific 18 oligonucleotide pairs were used to amplify the DNA enriched from the precipitated 19 Protein-DNA complexes from the TAP-PIF1 and HA-HY5 transgenic lines 20 maintained in the dark or exposed to 6 h red light. The gPCR using the oligonucleotide pairs 1 and 2 (Figure S3) showed no significant recovery in 21 22 comparison to the negative TAP-GFP and HA-GUS controls, indicating no binding 23 of PIF1 or HY5 to these promoter fragments. In contrast, enriched DNA sequences 24 were amplified from the TAP-PIF1 and HA-HY5 immunoprecipitated fractions with 25 the oligonucleotide pair P3 (F3/R3) (Figure S3). PIF1 binding was significant in the extracts from dark grown seedlings and a minor enrichment was seen in the light 26 27 extracts (Figure 5a). For the HA-HY5 immunoprecipitate we only observed specific 28 amplification in light-exposed seedlings (Figure 5b). For the P4 (F4/R4) primers 29 pair (Figure S3) a minor amplification was detected from both the TAP-PIF1 and 30 HA-HY5 extracts in comparison to the controls, indicating a very weak, and 31 probably not-significant (just slightly above the in-specific GUS-HA background)

binding of both factors to this fragment (Figure 5c, d). The fragment that showed 1 2 preferential binding to PIF1 and HY5 contains a PBE box, a GCE element and CG 3 hybrid box in close proximity to each other (Figure S3). For PIF1, the PBE box 4 (CACATG) is the most likely candidate binding-element and the GCE element (ACGT) for HY5. Very similar results were obtained from the ChIP experiments in 5 de-etiolated seedlings demonstrating that the same sites are involved in the PIF1 6 7 and HY5 binding in these two light developmental stages (Figure S4). 8 9 We also analyzed the promoter sequences of DXR and HDR genes for 10 potential PIF1 and HY5 binding sites. Within 1300 pb of the upstream sequence of DXR several G box-related sequences were found (Figure S5). Three pairs of 11 12 primers covering the different elements from the DXR promoter were used in ChIP experiments (Table S1 and Figure S5). As shown in Fig. 5f the only significant 13 14 enrichment detected was for HA-HY5 in the light with the primer pair 1 (F1 and R1) containing two GCE box-related sequences. These data demonstrate that DXR is 15 directly upregulated by HY5. In contrast, binding of TAP-PIF1 in the dark was 16 17 minor, although within statistical significance with the two primer pairs and close 18 with the primer pair 3 (p 0.055) (Figure 5e, g and i). In these fragments, several 19 putative G-box and E-box related sequences are present (Fig. S5). However, 20 considering that the binding of PIF1 to the three sites is so close to the negative

21 control in both photoperiodic and de-etiolation experiments (Figure S4e, g, i), this

22 result probably indicates a non-preferential interaction of PIF1 with the promoter of

23 DXR.

24

Finally, the *HDR* gene promoter includes 902 bp with only one PBE box and a GCE/ACE motif (Figure S6). Two pairs of primers were designed to cover the potential PIF1/HY5 binding sites and used to amplify the immunoprecipitates from the ChIPs (Table S1). In contrast to the *DXS1* and *DXR* genes, no enrichment was detected with HA-HY5 (Figure 5I), demonstrating that HY5 does not bind to the upstream sequences of the *HDR* gene. For TAP-PIF1 significant binding was observed in dark samples with the primer pair P2 that includes a PBE-box (Figure

5k). ChIPs results in de-etiolated seedlings (Figure S4k and I) tightly correlate with 1 2 those of the photoperiodic-dark-adapted seedlings (Figure 5k, I). In conclusion, 3 these data support the notion that modulation by light of HDR transcript levels 4 results from its de-repression from the dark activity of PIF1/PIF3 and not from 5 activation in light by HY5. Whether there is another light regulated activation factor, remains to be investigated. Over all, our ChIP studies likely reflect light imposed 6 7 changes in the PIFs/HY5 regulators dynamic behavior and differential promoter-8 binding capacity in response to the light environment. 9 10 Impact of transcriptional imposed regulation over light modulated levels of **MEP** proteins 11 12 The MEP-pathway provides with the intermediaries for the production of multiple compounds essential for photomorphogenic development, including 13 14 multiple hormones (GA, Cytokinin, ABA, Strigolactones) and photopigments 15 (carotenoids and chlorophylls). A previous study demonstrated that carotenoids and chlorophylls 16 17 accumulate upon light exposure and this accumulation is affected in the *pifQ* and 18 hy5 mutants (Toledo-Ortiz et al., 2014). Since the synthesis of these metabolites 19 depends directly on the MEP pathway, the transcriptional regulation of the MEP-20 transcripts by light can impact the accumulation of the final pathway products. To 21 evaluate the importance that light imposed transcriptional regulation of the DXS1, 22 DXR and HDR transcripts has over the pathway, the levels of the corresponding 23 proteins were analyzed in wild-type, pifs and hy5 mutants. Total protein extracts 24 from dark-adapted seedlings during 3 d (Wt, pif1, pif3, pif5 and hy5) or 36 h (pifQ) 25 were obtained and the level of the DXS1, DXR and HDR proteins compared to 26 dark controls. 27 28 We found that the transcriptional regulation mediated by the HY5/PIF1

29 module results in significant changes in the DXS1 enzyme level. Compared to the

30 wild-type, the hy5 mutant has lower accumulation of DXS1 protein whereas pif

31 single mutants and *pifQ* contain up to >10 times higher protein content than wild-

type in the dark (Figure 6B). It is worth noticing that although the DXS1 level in the 1 2 light is higher in the *pifQ* mutant, it does not maintain the same difference observed 3 in the dark (Figure 6b). We hypothesize that this might be the result of post-4 transciptional regulation that keeps this protein within certain levels (Flores-Perez 5 et al., 2008). This response correlates well with the transcript trends observed in pifs. The accumulation of DXS resembles the model of de-repression by PIFs in 6 7 the dark /activation by HY5 in the light. 8 9 We also observed accumulation of HDR protein in *pif3* and *pifQ* with a 10 contributed additive effect of primarily PIF3 and PIF1. We did not detect protein differences in *hy5* vs wild-type. The role of PIFs in de-repressing from the dark 11 12 leads to high protein levels in the light, but no further light-induced upregulation was detected at the time point assessed. 13 14 15 No major differences were detected in the level of DXR protein in the different mutants analyzed (Figure 6), in agreement with the more moderate 16 17 transcript differences observed for the pifs and hy5. 18 19 Discussion 20 21 The MEP pathway is a key biosynthetic route responsible for the synthesis 22 of essential compounds and signals that modulate developmental and stress 23 responses (Bouvier et al., 2005, Umehara et al., 2008, Hemmerlin et al., 2012, 24 Walley et al., 2015, Benn et al., 2016). Also, several of the MEP pathway products 25 have importance for human health and nutrition. Addressing how such a central metabolic pathway is modulated by external and internal cues has big implications 26 27 for future efforts to regulate its outputs including many compounds of 28 biotechnological or pharmaceutical interest 29 30 Light is one of the most relevant signals that affect plant metabolism,

31 including the production of essential photopigments, growth regulators and stress

hormones derived from the MEP-pathway (von Lintig et al., 1997, Rodríguez-1 2 Concepción et al., 2004). Previous work on photopigment production demonstrated 3 that light-responsive transcriptional factors HY5 and PIF1 control central carotenoid 4 and chlorophyll biosynthetic genes (Hug et al., 2004, Toledo-Ortiz et al., 2010). In 5 the case of phytoene synthase PIF1 and HY5 act as a module that antagonistically 6 balance the expression of this key carotenogenic gene (Toledo-Ortiz et al., 2014). 7 Our data demonstrate that via the same master-modulators, an additional higher-8 order layer arises via the light-induced tune up of the MEP-pathway for the 9 coordinated production of the precursors used for multiple compounds involved in 10 photomorphogenesis. Such is the case of carotenoids and the phytol side chain of 11 the chlorophylls. 12 13 The gene products of the MEP pathway accumulate upon light-exposure 14 (Botella-Pavía et al., 2004, Hsieh and Goodman, 2005, Cordoba et al., 2009), 15 however the molecular mechanisms for this upregulation and its impact on the pathway are still not fully understood. The coordinated tuning by light of the MEP 16 17 pathway genes leads to the possibility that common mechanisms regulate the 18 expression of these genes in response to light. Here we show that the master 19 regulators of light signals transduction HY5 and PIFs directly interact with the up-20 steam elements of the flux-controlling DXS1, DXR and HDR genes (Estévez et al., 2001, Botella-Pavía et al., 2004, Carretero-Paulet et al., 2006) and fine-tune their 21 22 expression levels in response to light. This regulation involves the interplay and 23 differential contribution of each factor for each gene, leading to unique

transcriptional dynamics (Figure 7) that lead to changes in protein accumulation.

25 Considering the flux-controlling capacity of DXS, DXR and HDR (Estévez et al.,

26 2001, Botella-Pavía et al., 2004, Enfissi et al., 2005, Carretero-Paulet et al., 2006,

27 Banerjee et al., 2013, Ghirardo et al., 2014), the dynamics observed could

importantly modify the flux through the pathway in response to the prevailing lightenvironment.

30 Our findings on the role of PIFs contrast to previous reports that concluded 31 that *DXS1* expression was not regulated by PIF1 (Toledo-Ortiz *et al.*, 2010). This

discrepancy probably results from differences in the quantification of transcript 1 2 levels through a lower-sensitivity microarray analysis compared to gRT-PCR and 3 the kinetics of light induction at earlier or later time points, as all these genes 4 present light/dark oscillatory patterns (Cordoba et al., 2009). In the case of PIF5, 5 using overexpressing PIF5 cell lines it was concluded that this factor was the main positive regulator of all the MEP pathway genes without major participation for 6 7 PIF1 or PIF3 (Mannen et al., 2014). However, in our study we did not observe 8 major differences of PIF1, PIF3 and PIF5 contribution as negative regulators of the 9 MEP pathway genes during de-etiolation and dark-adaptation. It is possible that 10 these differences result from a dominant negative effect of PIF5-over-expression. Also, we cannot exclude that PIF5 activity could change in other developmental or 11 12 environmental conditions as a result of interaction with other elements (Mannen et al., 2014). 13 14 Our studies demonstrate that the same key master integrators of light signal signals have the capacity to coordinate the core of the MEP pathway and its 15 16 multiple outputs through particular mechanisms for each gene (Figure 7). DXS1 17 expression is repressed in the dark by the direct binding of PIF1, probably through 18 the PBE box (CACATG), whereas HY5 induces the expression of this gene in the 19 light through direct binding to the nearby GCE element (ACGT). Despite the use of mild-over-expressors of PIF1 or HY5 for ChIP assays, the plant material likely 20 resembles the behavior of the endogenous proteins, as these transgenic lines 21 22 express comparable protein levels to the endogenous proteins, complement the 23 pif1 and hy5 mutant phenotypes and maintain the dark/light dynamics imposed on 24 these proteins by phytochromes and COP1, among others. Since PIFs/HY5 25 targeted *cis*-acting elements are located very close to one another, it is likely that the binding of one regulator results in allosteric interference of binding the second 26 27 regulator, resulting in an antagonistic mechanism. This regulation resembles the one described for the phytoene synthase (PSY) gene (Toledo-Ortiz et al., 2010, 28 29 Toledo-Ortiz et al., 2014).

In contrast, the accumulation of *HDR* in the light results exclusively from the
 degradation of the PIF repressors upon light exposure, without contribution from
 HY5. PIF1 directly interacts with *cis*-elements in the regulatory region of *HDR* (Figure 7). The strongest candidate as binding-element is a PBE motif present 156
 bp upstream of the ATG.

6

7 Finally, the expression of *DXR* accumulates in the light as a consequence of the

- 8 direct interaction of HY5 to elements located around 1.2 Kb upstream of the ATG
- 9 of this gene (Figure 7). In contrast, PIF1 has a weak interaction in more than one

10 region of the HDR promoter (Figure 7). The weak interaction observed with PIF1

11 could reflect that its binding capacity might depend on other PIFs. In agreement

12  $\,$  with this possibility our analyses demonstrate that in addition to PIF1, PIF3 and

13 PIF5 also affect the expression level of the DXS1, DXR and HDR genes.

14 Differential affinity of various PIFs for the promoters of MEP-pathway genes could

15 result in modified kinetics of light-responsiveness and modulation by other

- 16 environmental cues such as photoperiodism and the circadian clock.
- 17

18 Other studies have demonstrated high affinity of PIF1 and HY5 for G-box elements

19 in vitro and in vivo (Toledo-Ortiz et al., 2003, Hug et al., 2004, Oh et al., 2009,

20 Zhang et al., 2013). None of the MEP pathway genes analyzed contain canonical

21 G-boxes and our data support that for MEP-genes, the PBE box (CACATG) and

22 the GCE element (ACGT) are the strongest candidates to bind PIF1 for HY5

23 respectively. This differs and extends from the signal integration module

24 established by G-boxes in the case of photopigment related genes. Our analysis

- 25 from the DXS1::GUS transgenic lines carrying different promoter regions of the
- 26 DXS1 also supports our ChIP data, positioning the important cis-acting elements

27 within 300 bp from the transcription initiation site. It is worth to mention that with the

28 DXS1::GUS transgenic constructs although we observed the same response trend,

29 only a 2-fold increase between dark and light conditions was detected with high

30 GUS activity in the dark. This is in contrast to the low DXS1 transcript levels found

31 in the dark and its increase in response to light detected in our quantification

analyzes. This apparent discrepancy probably results from the accumulation of the
 GUS protein in the dark in the transgenic lines, as the stability of this protein is
 known to extend for more than 3 days (Kavita and Burma, 2008). Thus, in the case
 of *DXS1*, GUS is a good marker to identify regulatory *cis* acting elements but not
 for quantitative analyzes of its expression.

7 Hence together, our results indicate that the transcriptional regulation of 8 MEP-pathway genes by PIFs-HY5 results in a unique dynamic behavior for each 9 gene, providing additional flexibility to integrate inputs perceived by these master 10 regulators, such as time keeping or temperature signals. Interestingly, the HY5 and PIF binding motives in DXS1 gene localize proximal to a potential CCA1 binding 11 12 site (AAAATCT). CCA1 encodes a MYB-related protein that binds to Lhcb1\*3 and that participates in the phytochrome regulation of this gene (Wang et al., 1997). 13 14 This factor is also an important component of circadian regulation in coordination of HY5 (Nagel et al., 2015). Since DXS1, as well as other genes of the MEP 15 pathway are regulated by the circadian clock (Cordoba et al., 2009), it is possible 16 17 that the enhanced regulatory dynamic imposed by a PIF/HY5 co-acting module 18 would also bring unique capacity for the light and circadian regulation of DXS1 via 19 interaction with circadian components such as CCA1. 20 Importantly, we provide evidence that the differences in the transcript de-21 22 repression/activation by light mediated by PIFs/HY5 reflects on changes at the 23 protein level leading to unique protein dynamics. DXS1 follows a good 24 correspondence with the transcriptional fluctuations including low protein level in 25 the dark and accumulation in the light in a HY5/PIFs dependent manner, resulting in a significant difference in protein levels. These rapid changes in the protein 26 27 accumulation in response to light supports the possibility of a rapid turnover for this 28 protein and are consistent to its central role as a major flux controller of the 29 pathway in diverse environmental and developmental conditions (Estévez et al., 30 2001, Enfissi et al., 2005, Banerjee et al., 2013, Ghirardo et al., 2014, Wright et al.,

31 2014). However, we observed some discrepancies between the levels of the

2 1.5-fold increase after light exposure is detected. This result probably reflects post-3 translational regulatory events over the DXS1 that adjust the level of this protein in 4 response to the product demand, as has been previously reported (Pulido et al., 5 2013, Pokhilko et al., 2015, Pulido et al., 2016). Thus, while transcription regulation 6 of DXS1 plays an important role in control the levels of this enzyme post-7 translational regulatory events also act as an additional layer of regulation that 8 feedbacks metabolic requirements and impact upon the overall accumulation of 9 this protein. 10 On the other hand, HDR protein levels also reflect on the transcriptional regulation mediated by light, following a different dynamic that results from the de-repression 11 12 of transcript levels in the dark with no further changes associated with the light. The accumulation of HDR in response to light might be important to fulfill the 13 14 synthesis requirement of photopigments in coordination to DXS1. This result is consistent with the co-limiting role previously observed for HDR during carotenoid 15 synthesis in dark/light transition and during fruit ripening (Botella-Pavía et al., 2004, 16 17 Kim et al., 2009). This type or regulation could be particularly important in 18 conditions where different levels of PIFs accumulate, such as photoperiodic 19 conditions, and at the same time could limit the acute responses to the light signal. 20 Finally, changes in the DXR protein accumulation in response to light was reflected in constant protein levels in the light within the time frame analyzed. This result is 21 22 intriguing and it may reflect on particularities of the half-life of this protein or post-23 transcriptional events that control protein abundance. Future analyses in this 24 respect are important to address these possibilities. 25 26 Overall, our data support a model where a differential contribution of the 27 master light regulators PIFs and HY5 to light-modulated transcriptional effects 28 reflects in protein changes in MEP-pathway flux-controlling enzymes DXS1 and

transcript and their corresponding protein such as in the *pifQ* mutant, where only a

1

29 HDR. The significance of this differential regulation may impact on multiple

30 downstream pathways such as chlorophyll and carotenoid biosynthesis as well as

31 hormone and secondary metabolite synthesis, maintaining a very sensitive

1 responsiveness to the prevailing external conditions. Our studies exemplify how

2 differential multi-targeting of the initial steps of a pathway whose products multi-

3 branch downstream could impose a fine and unique modulation of all chloroplastic

- 4 isoprenoids.
- 5

## 6 Methods

7

#### 8 Plant Material and Growth Conditions.

9 The Arabidopsis thaliana lines used in this work are in Columbia (Col-0) background. Seeds from pif1-2, pif3-3, pif4-2, pif5-2, and pifQ were kindly provided 10 by P. Quail (University of California Berkeley). Seeds were grown on 1X Murashige 11 and Skoog (MS) media with Gamborg vitamins (Phytotechology Laboratories, 12 Shawnee Mission, KS) supplemented with 1% (w/v) sucrose and 0.8% (w/v) 13 phytoagar and stratified at 4°C for 4 days. For the light gene expression analysis 14 15 two treatments were used. For the dark-adapted treatment, seedlings were grown for 3 days in a 16 h light: 8h dark photoperiod at 120 µmol m<sup>2</sup> sec<sup>-1</sup>, followed by 3 16 day dark adaptation for the hy5, hy5hy5 and pif single mutants or 36 h for the pifQ. 17 18 Light treatment was done using 6 h with (100 µmol m<sup>2</sup> sec<sup>-1</sup>) cool white (Philips 19 F25T8/TL841) or (40 µmol m<sup>2</sup> sec<sup>-1</sup>) of red (Phillips LED module HF Deep Red 177354) lights. For de-etiolation experiments seedlings were exposed to 3 h of 20 21 white light, transferred to darkness for 3 days and exposed to 6 h of red light (40 22 µmol m<sup>2</sup> sec<sup>-1</sup>). Control seedlings were maintained in darkness. Growth 23 temperature was maintained between 21- 22°C in all cases.

24

#### 25 Expression analysis

- 26 Total RNA was isolated using TRIzol (Invitrogen, Carlsbad, CA, USA)
- 27 according to the protocol provided by the manufacturer. For northern-blot analysis,
- $28-5\,\mu g$  of total RNA was fractionated and transferred onto a Hybond-N+ nylon
- 29 membrane (GE, Buckinghamshire, UK). Hybridizations and washes were
- 30 performed under stringent conditions. Probes were <sup>32</sup>P-radiolabeled using the
- 31 Megaprime DNA labeling system (GE, Buckinghamshire, UK). All probes were

1 obtained by PCR amplification as previously reported (Guevara-García et al.,

2 2005). For qRT-PCR experiments seedlings were harvested in the dark for the

3 dark samples and RNA extraction conducted using the RNeasy Plant Mini Kit

4 (Qiagen) or TRIzol. Complementary DNA (cDNA) was obtained from DNase-

5 treated RNA with M-MLV Reverse Transcriptase kit (Invitrogen, Carlsbad, CA). The

6 qRT-PCR experiments were performed using FastStart DNA Master<sup>PLUS</sup> SYBR

7 Green I (Roche) on an Agilent Technologies Stratagene MX3005P or a Light

8 Cycler 480 Roche. Analyses were done with three independent experiments and

9 technical duplicates were included in each case (n=2). The reference gene used in

10 the qPCR analyses is ACT7 since the expression of this reference gene has been

11 shown to not have major fluctuations in the conditions analyzed.

12

#### 13 **DXS1 promoter analysis.**

14 From the transcriptional DXS1::GUS clone (Estévez et al., 2000), two additional clones were generated containing 750 bp and 660 bp deletions from the 15 ATG of DXS1. The fragments were subcloned into the pBin19 binary vector and 16 17 used to generate transgenic lines through Agrobacterium tumefaciens-mediated 18 transformation into the Col-0 ecotype (Clough and Bent, 1998). At least three 19 independent homozygous lines were selected for each construct and analyzed. 20 21 **GUS** histochemical and fluorimetric analyses 22 Three day-old etiolated seedlings exposed to light or dark for 6 h were 23 stained using the GUS histochemical assay (Jefferson et al., 1987). Plants were 24 clarified as reported (Malamy and Benfey, 1997) and visualized using a

25 stereoscopic microscope (Nikon SMZ1500). For the fluorometric analysis the

26 seedlings were homogenized in GUS extraction buffer (50 mM NaHPO<sub>4</sub>, pH 7.0; 10

27 mM Na<sub>2</sub>EDTA; 1% Triton X-100; 0.1% N-lauroyl sarcosine and 10 mM β-

28 mercaptoetanol). The enzymatic reaction was done using 5 µl of the extracts.

29 Fluorometric quantification was done with TKO 100 fluorimeter (Hoeffer). Specific

30 activity was determined as nmol of methyl-umbelliferone per  $\mu$ g protein<sup>-1</sup> per min<sup>-1</sup>.

#### 1 Protein gel blot analysis

2 Total protein was obtained from seedlings and 20 µg of the samples was 3 separated by SDS-polyacrylamide gel electrophoresis (SDS-PAGE). The proteins 4 were transferred onto nitrocellulose membrane (Amersham Protan Premium 0.45 5 µm NC GE Healthcare Life Science). To verify equal loading, a parallel gel was run and stained with Coomasie blue. Immunodetection was performed as previously 6 7 reported (Guevara-García et al., 2005). Detection was done using the Luminata 8 Crescendo Western HRP Substrate (Millipore, USA). Bands from three independent experiments were quantified by densitometric analysis using ImageJ 9 10 software (Schneider et al., 2012). 11 12 **Chromatin Immunoprecipitation assays** 13 ChIP assays were conducted following the protocol reported previously 14 (Moon et al., 2008) except that in our assays, 2 week old- seedlings were used. Plants were grown as described in (Toledo-Ortiz et al., 2014), and dark adapted for 15 72h before light treatments and sample collection. Samples were collected for dark 16 17 time points (0h, before lights on at the end of dark adaptation) or after 6 h 18 illumination with red light (40 µmol m<sup>-2</sup> s<sup>-1</sup>). Plant material used (35S::HA-HY5 in 19 hy5-215 and 35S::TAP-PIF1 in pif1-2) was previously described (Lee et al., 2007, Moon et al., 2008) and kindly provided by the Deng Lab (Yale) and Hug Lab (UT 20 Austin). Both lines are mild-over expressors that complement the mutant 21 22 phenotypes and show comparative levels and light responses to native PIF1 and 23 HY5. . gRT-PCR was conducted on a Roche 480 Light cyler according to standard 24 protocol by the manufacturer. The oligonucleotides sequences used to amplify 25 upstream promoter regions of individual genes are shown in Table S1. Upstream sequences of the DXS1, DXR and HDR genes were analyzed for possible light 26 27 responsive elements using SOGO New PLACE software (https://sogo.dna.affrc.go.jp/cgibin/sogo.cgi?sid=&lang=en&pj=640&action=page&p 28 age=newplace) and are shown in Figures S2, S3 and S4 (Higo et al., 1999). 29 30

#### 1 Acknowledgements

- 2
- 3 We thank Dr. Nidia Sánchez and Dr. Gabriela Perez for technical assistance and
- 4 Geoff Holrooyd and Phoebe Sutton for proof-reading the manuscript. This research
- 5 was supported by grants from CONACYT (CB 220534 and FC 2016-96) and
- 6 DGAPA-UNAM (IN204617) for PL and by Royal Society Grant (RG150711) to
- 7 GTO. MC and CR were supported by PhD fellowship from CONACYT and DGAPA.
- 8 The authors declare no conflict of interest.
- 9

#### 10 **Bibliography**

12	Ban	lerjee	e, A.	and	Sha	arke	y, T.I	<b>).</b> (2	01	4) N	ſetl	hylery	thrit	ol 4	l-ph	ospha	te (N	ИЕР)	pa	thw	ay
13		me	etab	olic	regi	ulatio	on. No	at P	rod	Re	p, <b>3</b>	<b>1</b> , 104	3-10	)55							
	-			* * *		-		-								-	(0.0	4.00	-		

- Banerjee, A., Wu, Y., Banerjee, R., Li, Y., Yan, H. and Sharkey, T.D. (2013) Feedback
  inhibition of deoxy-D-xylulose-5-phosphate synthase regulates the
  methylerythritol 4-phosphate pathway. *J Biol Chem*, 288, 16926-16936.
  Benn, G., Bjornson, M., Ke, H., De Souza, A., Balmond, E.I., Shaw, J.T. and Dehesh,
- Benn, G., Bjornson, M., Ke, H., De Souza, A., Bannond, E.I., Snaw, J. I. and Denesn,
   K. (2016) Plastidial metabolite MEcPP induces a transcriptionally centered
   stress-response hub via the transcription factor CAMTA3. *Proc Natl Acad Sci U* S A, 113, 8855-8860.
- Botella-Pavía, P., Besumbes, O., Phillips, M.A., Carretero-Paulet, L., Boronat, A.
   and Rodríguez-Concepción, M. (2004) Regulation of carotenoid biosynthesis
   in plants: evidence for a key role of hydroxymethylbutenyl diphosphate
   reductase in controlling the supply of plastidial isoprenoid precursors. *Plant J*,
   40, 188-199.
- Bouvier, F., Rahier, A. and Camara, B. (2005) Biogenesis, molecular regulation and
   function of plant isoprenoids. *Progress in lipid research*, 44, 357-429.
- Carretero-Paulet, L., Cairó, A., Botella-Pavía, P., Besumbes, O., Campos, N.,
   Boronat, A. and Rodríguez-Concepción, M. (2006) Enhanced flux through
   the methylerythritol 4-phosphate pathway in *Arabidopsis* plants
   overexpressing deoxyxylulose 5-phosphate reductoisomerase. *Plant Mol Biol*,
   62, 683-695.
- Chappell, J. (1995) Biochemistry and molecular biology of the isoprenoid pathway in
   plants. Ann. Rev. Plant Mol. Biol., 46, 521-547.
- Chen, D., Xu, G., Tang, W., Jing, Y., Ji, Q., Fei, Z. and Lin, R. (2013) Antagonistic basic
   helix-loop-helix/bZIP transcription factors form transcriptional modules that
   integrate light and reactive oxygen species signaling in Arabidopsis. *Plant Cell*,
   25, 1657-1673.
- Chen, M., Chory, J. and Fankhauser, C. (2004) Light signal transduction in higher
   plants. *Annual review of genetics*, 38, 87-117.

**Clough**, **S.I. and Bent**, **A.F.** (1998) Floral dip: a simplified method for Agrobacterium-1 2 mediated transformation of Arabidopsis thaliana. Plant J, 16, 735-743. 3 Cordoba, E., Salmi, M. and León, P. (2009) Unravelling the regulatory mechanisms 4 that modulate the MEP pathway in higher plants. *J Exp Bot*, **60**, 2933-2943. 5 Croteau, R., Kutchan, T.M. and Lewis, N.G. (2000) Natural products (secondary metabolites). In Biochemistry and Molecular Biology of Plants (Buchanan, B.B., 6 7 Gruissem, W. and Jones, R.L. eds). Rockville, MD: American Society of Plant 8 Biology, pp. 1250-1268. 9 de Souza, A., Wang, J.Z. and Dehesh, K. (2017) Retrograde Signals: Integrators of 10 Interorganellar Communication and Orchestrators of Plant Development. Annu 11 Rev Plant Biol, 68, 85-108. 12 Eisenreich, W., Bacher, A., Arigoni, D. and Rohdich, F. (2004) Biosynthesis of 13 isoprenoids via the non-mevalonate pathway. Cell Mol Life Sci, 61, 1401-1426. 14 Enfissi, E.M.A., Fraser, P.D., Lois, L.M., Boronat, A., Schuch, W. and Bramley, P.M. 15 (2005) Metabolic engineering of the mevalonate and non-mevalonate 16 isopentenyl diphosphate-forming pathways for the production of healthpromoting isoprenoids in tomato . Plant Biotech J, 3, 17-27. 17 18 Estévez, J.M., Cantero, A., Reindl, A., Reichler, S. and León, P. (2001) 1-Deoxy-D-19 xylulose-5-phosphate synthase, a limiting enzyme for plastidic isoprenoid 20 biosynthesis in plants. J Biol Chem, 276, 22901-22909. 21 Estévez, J.M., Cantero, A., Romero, C., Kawaide, H., Jiménez, L.F., Kuzuyama, T., 22 Seto, H., Kamiya, Y. and León, P. (2000) Analysis of the expression of CLA1, a 23 gene that encodes the 1-deoxyxylulose 5-phosphate synthase of the 2-C-24 methyl-D-erythritol-4-phosphate pathway in Arabidopsis. Plant Physiol, 124, 95-103. 25 26 Flores-Perez, U., Sauret-Güeto, S., Gas, E., Jarvis, P. and Rodríguez-Concepción, M. 27 (2008) A mutant impaired in the production of plastome-encoded proteins 28 uncovers a mechanism for the homeostasis of isoprenoid biosynthetic enzymes 29 in Arabidopsis plastids. Plant Cell, 20, 1303-1315. 30 Franklin, K.A. and Quail, P.H. (2010) Phytochrome functions in Arabidopsis development. J Exp Bot, 61, 11-24. 31 32 Ghirardo, A., Wright, L.P., Bi, Z., Rosenkranz, M., Pulido, P., Rodriguez-33 Concepcion, M., Niinemets, U., Bruggemann, N., Gershenzon, J. and 34 Schnitzler, J.P. (2014) Metabolic flux analysis of plastidic isoprenoid 35 biosynthesis in poplar leaves emitting and nonemitting isoprene. Plant Physiol, 36 165.37-51. 37 Guevara-García, A.A., San Roman, C., Arroyo, A., Cortés, M.E., Gutiérrez-Nava, M.L. and León, P. (2005) The characterization of the Arabidopsis *clb6* mutant 38 39 illustrates the importance of post-transcriptional regulation of the Methyl-D-40 Erythritol 4-Phosphate Pathway. Plant Cell, 17, 628-643. 41 Hemmerlin, A., Harwood, J.L. and Bach, T.J. (2012) A raison d'etre for two distinct 42 pathways in the early steps of plant isoprenoid biosynthesis? Progress in lipid research, **51**, 95-148. 43 Higo, K., Ugawa, Y., Iwamoto, M. and Korenaga, T. (1999) Plant cis-acting 44 45 regulatory DNA elements (PLACE) database: 1999. Nucleic Acids Res, 27, 297-300. 46

- Hsieh, M.H. and Goodman, H.M. (2005) The Arabidopsis *IspH* homolog is involved in 1 2 the plastid nonmevalonate pathway of isoprenoid biosynthesis. *Plant Physiol*, 3 138, 641-653. 4 Hug, E., Al-Sady, B., Hudson, M., Kim, C., Apel, K. and Quail, P.H. (2004) 5 Phytochrome-interacting factor 1 is a critical bHLH regulator of chlorophyll biosynthesis. Science, 305, 1937-1941. 6 7 Jefferson, R.A., Kavanagh, T.A. and Bevan, M.W. (1987) GUS fusions: beta-8 glucuronidase as a sensitive and versatile gene fusion marker in higher plants. 9 Embo J, 6, 3901-3907. 10 Jeong, J. and Choi, G. (2013) Phytochrome-interacting factors have both shared and distinct biological roles. Mol Cells, 35, 371-380. 11 12 Jiao, Y., Lau, O.S. and Deng, X.W. (2007) Light-regulated transcriptional networks in 13 higher plants. Nature reviews. Genetics, 8, 217-230. 14 Kavita, P. and Burma, P.K. (2008) A comparative analysis of green fluorescent 15 protein and beta-glucuronidase protein-encoding genes as a reporter system for studying the temporal expression profiles of promoters. J Biosci, 33, 337-16 343. 17 18 Kim, J., Kang, H., Park, J., Kim, W., Yoo, J., Lee, N., Kim, J., Yoon, T.Y. and Choi, G. 19 (2016) PIF1-Interacting Transcription Factors and Their Binding Sequence 20 Elements Determine the in Vivo Targeting Sites of PIF1. Plant Cell, 28, 1388-21 1405. 22 Kim, Y.B., Kim, S.M., Kang, M.K., Kuzuyama, T., Lee, J.K., Park, S.C., Shin, S.C. and 23 Kim, S.U. (2009) Regulation of resin acid synthesis in Pinus densiflora by 24 differential transcription of genes encoding multiple 1-deoxy-D-xylulose 5-25 phosphate synthase and 1-hydroxy-2-methyl-2-(*E*)-butenyl 4-diphosphate 26 reductase genes. Tree Physiol, 29, 737-749. 27 Lange, B.M., Rujan, T., Martin, W. and Croteau, R. (2000) Isoprenoid biosynthesis: 28 the evolution of two ancient and distinct pathways across genomes. Proc Natl 29 Acad Sci USA, 97, 13172-13177. 30 Lee, J., He, K., Stolc, V., Lee, H., Figueroa, P., Gao, Y., Tongprasit, W., Zhao, H., Lee, I. and Deng, X.W. (2007) Analysis of transcription factor HY5 genomic binding 31 32 sites revealed its hierarchical role in light regulation of development. Plant Cell, 33 19,731-749. 34 Leivar, P. and Monte, E. (2014) PIFs: systems integrators in plant development. Plant 35 *Cell*, **26**, 56-78. 36 Leivar, P., Monte, E., Oka, Y., Liu, T., Carle, C., Castillon, A., Hug, E. and Ouail, P.H. 37 (2008) Multiple phytochrome-interacting bHLH transcription factors repress premature seedling photomorphogenesis in darkness. *Curr Biol*, **18**, 1815-38 39 1823. 40 Leivar, P. and Quail, P.H. (2011) PIFs: pivotal components in a cellular signaling hub. Trends Plant Sci, 16, 19-28. 41 42 Leivar, P., Tepperman, J.M., Monte, E., Calderon, R.H., Liu, T.L. and Quail, P.H. 43 (2009) Definition of early transcriptional circuitry involved in light-induced 44 reversal of PIF-imposed repression of photomorphogenesis in young
- 45 Arabidopsis seedlings. *Plant Cell*, **21**, 3535-3553.

1	Lois, L.M., Rodríguez-Concepción, M., Gallego, F., Campos, N. and Boronat, A.	
2	(2000) Carotenoid biosynthesis during tomato fruit development: regulatory	
3	role of 1-deoxy-D-xylulose 5-phosphate synthase. <i>Plant J</i> , <b>22</b> , 503-513.	
4	Mahmoud, S.S. and Croteau, R.B. (2001) Metabolic engineering of essential oil yield	
5	and composition in mint by altering expression of deoxyxylulose phosphate	
6	reductoisomerase and menthofuran synthase. Proc Natl Acad Sci U S A, <b>98</b> ,	
7	8915-8920.	
8	Malamy, J.E. and Benfey, P.N. (1997) Organization and cell differentiation in lateral	
9	roots of Arabidopsis thaliana. Development, <b>124</b> , 33-44.	
10	Mannen, K., Matsumoto, T., Takahashi, S., Yamaguchi, Y., Tsukagoshi, M., Sano,	
11	R., Suzuki, H., Sakurai, N., Shibata, D., Koyama, T. and Nakayama, T. (2014)	
12	Coordinated transcriptional regulation of isopentenyl diphosphate	
13	biosynthetic pathway enzymes in plastids by phytochrome-interacting factor 5.	
14	Biochem Biophys Res Commun, <b>443</b> , 768-774.	
15	Meier, S., Tzfadia, O., Vallabhaneni, R., Gehring, C. and Wurtzel, E.T. (2011) A	
16	transcriptional analysis of carotenoid, chlorophyll and plastidial isoprenoid	
17	biosynthesis genes during development and osmotic stress responses in	
18	Arabidopsis thaliana. BMC Syst Biol, 5, 77.	
19	Moon, J., Zhu, L., Shen, H. and Huq, E. (2008) PIF1 directly and indirectly regulates	
20	chlorophyll biosynthesis to optimize the greening process in Arabidopsis. Proc	
21	Natl Acad Sci U S A, <b>105</b> , 9433-9438.	
22	Nagel, D.H., Doherty, C.J., Pruneda-Paz, J.L., Schmitz, R.J., Ecker, J.R. and Kay, S.A.	
23	(2015) Genome-wide identification of CCA1 targets uncovers an expanded	
24	clock network in Arabidopsis. Proc Natl Acad Sci USA, 112, E4802-4810.	
25	Oh, E., Kang, H., Yamaguchi, S., Park, J., Lee, D., Kamiya, Y. and Choi, G. (2009)	
26	Genome-wide analysis of genes targeted by PHYTOCHROME INTERACTING	
27	FACTOR 3-LIKE5 during seed germination in Arabidopsis. Plant Cell, 21, 403-	
28	419.	
29	Oudin, A., Mahroug, S., Courdavault, V., Hervouet, N., Zelwer, C., Rodriguez-	
30	Concepcion, M., St-Pierre, B. and Burlat, V. (2007) Spatial distribution and	
31	hormonal regulation of gene products from methyl erythritol phosphate and	
32	monoterpene-secoiridoid pathways in Catharanthus roseus. <i>Plant Mol Biol</i> , <b>65</b> ,	
33	13-30.	
34	Phillips, M.A., León, P., Boronat, A. and Rodríguez-Concepción, M. (2008) The	
35	plastidial MEP pathway: unified nomenclature and resources. Trends Plant Sci,	
36	<b>13</b> , 619-623.	
37	Pokhilko, A., Bou-Torrent, J., Pulido, P., Rodriguez-Concepcion, M. and Ebenhoh,	
38	<b>0.</b> (2015) Mathematical modelling of the diurnal regulation of the MEP	
39	pathway in Arabidopsis. <i>The New phytologist</i> , <b>206</b> , 1075-1085.	
40	Pulido, P., Llamas, E., Llorente, B., Ventura, S., Wright, L.P. and Rodriguez-	
41	Concepcion, M. (2016) Specific Hsp100 Chaperones Determine the Fate of the	
42	First Enzyme of the Plastidial Isoprenoid Pathway for Either Refolding or	
43	Degradation by the Stromal Clp Protease in Arabidopsis. <i>PLoS genetics</i> , <b>12</b> ,	
44	e1005824.	
45	Pulido, P., Toledo-Ortiz, G., Phillips, M.A., Wright, L.P. and Rodriguez-	
46	Concepcion, M. (2013) Arabidopsis J-protein J20 delivers the first enzyme of	

**Concepcion, M.** (2013) Arabidopsis J-protein J20 delivers the first enzyme of

1	the plastidial isoprenoid pathway to protein quality control. <i>Plant Cell</i> , <b>25</b> ,
2	4183-4194.
3	Rodriguez-Concepcion, M. (2006) Early steps in isoprenoid biosynthesis: multilevel
4	regulation of the supply of common precursors in plant cells. <i>Phytochem Rev</i> 5,
5	1-15.
6	Rodríguez-Concepción, M. (2004) The MEP pathway: a new target for the
7	development of herbicides, antibiotics and antimalarial drugs. Curr Pharm Des,
8	<b>10</b> , 2391-2400.
9	Rodriguez-Concepcion, M. and Boronat, A. (2015) Breaking new ground in the
10	regulation of the early steps of plant isoprenoid biosynthesis. <i>Curr Opin Plant</i>
11	Biol, <b>25</b> , 17-22.
12	Rodríguez-Concepción, M., Forés, O., Martínez-García, J.F., González, V., Phillips,
13	M.A., Ferrer, A. and Boronat, A. (2004) Distinct light-mediated pathways
14	regulate the biosynthesis and exchange of isoprenoid precursors during
15	Arabidopsis seedling development. <i>Plant Cell</i> , <b>16</b> , 144-156.
16	Rohdich, F., Bacher, A. and Eisenreich, W. (2005) Isoprenoid biosynthetic pathways
17	as anti-infective drug targets. <i>Biochem Soc Trans</i> , <b>33</b> , 785-791.
18	Ruiz-Sola, M.A. and Rodriguez-Concepcion, M. (2012) Carotenoid biosynthesis in
19	Arabidopsis: a colorful pathway. The Arabidopsis book / American Society of
20	Plant Biologists, <b>10</b> , e0158.
21	Schneider, C.A., Rasband, W.S. and Eliceiri, K.W. (2012) NIH Image to Image]: 25
22	years of image analysis. <i>Nat Methods</i> , <b>9</b> , 671-675.
23	Tepperman, J.M., Zhu, T., Chang, H.S., Wang, X. and Quail, P.H. (2001) Multiple
24	transcription-factor genes are early targets of phytochrome A signaling. Proc
25	Natl Acad Sci U S A, 98, 9437-9442.
26	Toledo-Ortiz, G., Huq, E. and Quail, P.H. (2003) The Arabidopsis basic/helix-loop-
27	helix transcription factor family. <i>Plant Cell</i> , <b>15</b> , 1749-1770.
28	Toledo-Ortiz, G., Huq, E. and Rodriguez-Concepcion, M. (2010) Direct regulation of
29	phytoene synthase gene expression and carotenoid biosynthesis by
30	phytochrome-interacting factors. Proc Natl Acad Sci USA, 107, 11626-11631.
31	Toledo-Ortiz, G., Johansson, H., Lee, K.P., Bou-Torrent, J., Stewart, K., Steel, G.,
32	Rodriguez-Concepcion, M. and Halliday, K.J. (2014) The HY5-PIF regulatory
33	module coordinates light and temperature control of photosynthetic gene
34	transcription. <i>PLoS genetics</i> , <b>10</b> , e1004416.
35	Umehara, M., Hanada, A., Yoshida, S., Akiyama, K., Arite, T., Takeda-Kamiya, N.,
36	Magome, H., Kamiya, Y., Shirasu, K., Yoneyama, K., Kyozuka, J. and
37	Yamaguchi, S. (2008) Inhibition of shoot branching by new terpenoid plant
38	hormones. <i>Nature</i> , <b>455</b> , 195-201.
39	von Lintig, J., Welsch, R., Bonk, M., Giuliano, G., Batschauer, A. and Kleinig, H.
40	(1997) Light-dependent regulation of carotenoid biosynthesis occurs at the
41	level of phytoene synthase expression and is mediated by phytochrome in
42	Sinapis alba and Arabidopsis thaliana seedlings. <i>Plant J</i> , <b>12</b> , 625-634.
43	Vranova, E., Coman, D. and Gruissem, W. (2013) Network analysis of the MVA and
44	MEP pathways for isoprenoid synthesis. Annu Rev Plant Biol, 64, 665-700.
45	Walley, J., Xiao, Y., Wang, J.Z., Baidoo, E.E., Keasling, J.D., Shen, Z., Briggs, S.P. and
46	Dehesh, K. (2015) Plastid-produced interorgannellar stress signal MEcPP

1 2	potentiates induction of the unfolded protein response in endoplasmic reticulum. <i>Proc Natl Acad Sci U S A</i> , <b>112</b> , 6212-6217.
3	Walter, M., Fester, T. and Strack, D. (2000) Arbuscular mycorrhizal fungi induce the
4	non-mevalonate methylerythritol phosphate pathway of isoprenoid
5 6	biosynthesis correlated with accumulation of the yellow pigment and the other apocarotenoids. <i>Plant J</i> , <b>21</b> , 571-578.
7	Wang, Z.Y., Kenigsbuch, D., Sun, L., Harel, E., Ong, M.S. and Tobin, E.M. (1997) A
8	Myb-related transcription factor is involved in the phytochrome regulation of
9	an Arabidopsis Lhcb gene. <i>Plant Cell</i> , <b>9</b> , 491-507.
10	Wright, L.P., Rohwer, J.M., Ghirardo, A., Hammerbacher, A., Ortiz-Alcaide, M.,
11	Raguschke, B., Schnitzler, J.P., Gershenzon, J. and Phillips, M.A. (2014)
12	Deoxyxylulose 5-Phosphate Synthase Controls Flux through the
13	Methylerythritol 4-Phosphate Pathway in Arabidopsis. Plant Physiol, 165,
14	1488-1504.
15	Zhang, H., He, H., Wang, X., Wang, X., Yang, X., Li, L. and Deng, X.W. (2011)
16	Genome-wide mapping of the HY5-mediated gene networks in Arabidopsis that
17	involve both transcriptional and post-transcriptional regulation. Plant J, 65,
18	346-358.
19	Zhang, Y., Mayba, O., Pfeiffer, A., Shi, H., Tepperman, J.M., Speed, T.P. and Quail,
20	P.H. (2013) A quartet of PIF bHLH factors provides a transcriptionally centered
21	signaling hub that regulates seedling morphogenesis through differential
22	expression-patterning of shared target genes in Arabidopsis. <i>PLoS genetics</i> , <b>9</b> ,
23	e1003244.
24	
25	Supporting information
26	Complemental Figure 4. Europeanies and size of the DVC4. DVD and UDD server
27	Supplemental Figure 1. Expression analysis of the DXS1, DXR and HDR genes
28 29	in <i>hy5</i> mutants in Red light.
29 30	Supplemental Figure 2. Expression analysis of the DXS1, DXR and HDR genes
31	in the <i>pif</i> single mutants in Red light.
32	
33	Supplemental Figure 3. DXS1 upstream regulatory region.
34	
35	Supplemental Figure 4. Chromatin immunoprecitation assays for 35S::TAP-PIF1
36	and 35S::HA-HY5 in de-etiolated seedlings.
37	3
38	Supplemental Figure 5. DXR upstream regulatory region.
39	
40	Supplemental Figure 6. HDR upstream regulatory region.
41	
42	Supplemental Table I. Sequence of the oligonucleotides used in the work.

#### 1 Figure Legends

2

3 Figure 1. Transcriptional regulation of DXS1 by light. Histochemical expression 4 of 3 day-old transgenic representative lines grown in the dark (a-c) or exposed to 6 5 h of light (d-f) expressing the GUS marker from 1510 bp (a and d), 753 bp (b and e) and 660 bp (c and f) upstream sequences from the ATG of DXS1. (g) GUS specific 6 7 activities from the dark (dark gray) or exposed to light (gray) transgenic lines. The 8 induction level is reported relative to the mean of the specific activity (nmol of 9 methyl-umbelliferone per µg of protein<sup>-1</sup> min<sup>-1</sup>) of the corresponding dark sample. 10 Each bar is the mean of three independent experiments and error bars represent ±SD. The numbers above the bars indicate the P values according to a Student's t 11 12 test. 13 14 Figure 2. Expression analysis of the DXS1, DXR and HDR genes in hy5 and 15 hy5hyh mutants. Representative northern blots of DXS1 (a), DXR (b) and HDR (c), each lane contains 5 µg of RNA from 3 day-old dark-adapted seedlings 16 17 maintained in the dark (D) or exposed to light for 6 h (L) from wild-type (Wt), hy5 18 and *hy5hyh* mutants and hybridized with probes for the DXS1 (a), DXR (b) and 19 HDR (c) genes. The 28S rRNA is shown as loading control. The membrane is 20 representative of three independent biological experiments. gRT-PCR analysis of DXS1 (d), DXR (e) and HDR (f) transcript levels from Col-0, hy5 and hy5hyh dark-21 22 adapted seedlings maintain in the dark (gray column) or exposed to 6 h of light 23 (white column). Expression is normalized to the Col-0 dark samples and adjusted 24 to Actin 7 (ACT7). Bars are means ±SE of triplicate biological experiments (each 25 with n=2 technical replicas). The numbers bars indicate P values p<0.05 and the \* marks statistical difference (p<0.05) between the light samples from the hy5 and 26 27 hy5hyh mutants compared to the induction in the Wt according to a Student's t 28 test. 29

## Figure 3. Expression analysis of the *DXS1*, *DXR* and *HDR* genes in the *pif* single mutants. Representative RNA blots of *DXS1* (a), *DXR* (b) and *HDR* (c)

1 from 3 day-old dark-adapted wild-type Col0 (Wt), pif1, pif3, pif4 and pif5 maintained

2 in darkness (D) or exposed to 6 h light (L). The 28S rRNA is shown as a loading

3 control (28S). Membranes are representative of three independent biological

4 experiments. Analysis by qRT-PCR of DXS1 (d,g), DXR (e,h) and HDR (f,i)

5 transcripts from Col-0, pif1, pif3, pif4 and pif5 (d-f) dark-adapted for 3 days (gray

6 column) or 36 h for *pifQ* (g-i) and exposed 6 h of red light (white column).

7 Expression is reported relative to the dark Col-0 sample and adjusted relative to

8 Actin 7 (ACT7). Bars are means ±SE of triplicate biological experiments (each with

9 n=2 technical replicas). The letter above the bars indicate P values p<0.05

10 between dark (a) or light (b) of Wt compared to the mutants (Student's t test).

11

12 Figure 4. Expression analysis of the DXS1, DXR and HDR genes in the hy5,

13 *pif* single mutants and *pifQ* during de-etiolation. qRT-PCR analysis of *DXS1* 

14 (a), DXR (b) and HDR (c) genes from etiolated seedlings of Col-0, hy5, pif1, pif3,

15 *pif4, pif5* and *pifQ* maintained in the dark (gray columns) or exposed to 6 h of red

16 light (white columns) relative to the Col-0 dark samples and adjusted relative to

17 Actin 7 (ACT7). Bars are means ±SE of triplicate biological experiments (each with

18 n=2 technical replicas). The letter above the bars indicate values (p<0.05)

19 decrease (a and b) or increase (c and d) between the dark (a and c) or light (b and

20 d) values between the Col*0* and the mutants (Student's t test).

21

22 Figure 5. Chromatin immunoprecitation assays for 35S::TAP-PIF1 and

23 35S::HA-HY5 in dark-adapted seedlings. Diagrams of the upstream regions of

24 DXS1, DXR and HDR genes. Primers used for the analyses (arrows) and the

25 potential PIFs (black) and HY5 (gray) binding elements (rectangles). ChIP of three

26 days dark-adapted TAP-PIF and HA-HY5 transgenic seedlings (Lee et al., 2007,

27 Moon *et al.*, 2008) maintained in the dark (grey zone) or exposed to 6h red-light

28 (clear zone). ChIP was conducted using specific antibodies against MYC for PIF1

the HA for HY5. 35S::GFP-TAP or 35S::GUS-HA lines were used as controls for

30 unespecific binding. The ChIP/qPCR was done using specific primer pairs (F,

31 forward primer and R, reverse primer) covering the regions containing putative

- 1 binding elements. ChIP-enriched DNA regions of the TAP-PIF *DXS1* (a and c),
- 2 DXR (e,g and i) and HDR (k) or for HA-HY5 DXS1 (b and d) DXR (f,h and j) and
- 3 HDR (I) samples. The bars are the mean ± SE of triplicate independent
- 4 experiments (each with technical duplicates n=2). The asterisk indicate the values
- 5 with significance (p<0.05) between the negative control according to a Student's t
- 6

test.

7

### 8 Figure 6. Protein accumulation of the DXS1, DXR and HDR in mutants of the

- 9 HY5 and PIFs. (a) Immunoblots with 20 µg of protein extracts from seedlings dark-
- 10 adapted for 3 days (Col-0, hy5, pif1, pif3, pif4 and pif5) or 36 h (pifQ) and
- 11 maintained in the dark (D) or exposed to 6 h light (L) using specific antibodies for
- 12 DXS1, DXR or HDR proteins. A Coomassie blue-stained gel (Coo) is shown as a
- 13 loading control. A representative gel from three independent biological experiments
- 14 is shown. (b) Densitometric analyses of the DXS1, DXR and HDR protein levels
- 15 from Col-0, *hy5*, *pif1*, *pif3*, *pif5* and *pifQ* immunoblots from the dark (grey bars) or
- 16 exposed to 6 h light (white bars) samples. The expression level is reported relative
- 17 to the Col-0 light samples and adjusted to the corresponding loading control. The
- 18 bars correspond to the average of three independent biological experiments ± SD
- 19 of biological triplicates.
- 20

# Figure 7. Model of the differential light regulation of *DXS1*, *DXR* and *HDR*

- 22 gene expression to modulate the MEP pathway. Light via phytochrome (Pfr)
- 23 results in the degradation of PIFs and in the accumulation of the HY5. For the
- 24 DXS1, Pfr accumulation inhibits the direct repression of PIF1 (solid arrow), PIF3
- 25 and PIF5 (dashed arrows) and the activation of HY5 (solid arrow). For DXR, Pfr
- 26 impairs the weak repression of PIF1 in multiple sites (solid arrows), PIF3 and PIF5
- 27 (dashed arrows). Light via HY5 activate DXR (solid line). Finally, Pfr accumulation
- 28 induces HDR expression through the degradation of the PIFs. The dashed arrows
- 29 mean that a direct interaction was not experimentally demonstrated. The thickness
- 30 of the arrows reflects the RNA levels and the enrichment detected in the ChIP

**Commented [HG2]:** In the figure 'f' is incorrectly annotated 'g' so you have two labeled 'g'

1 analyses. The orange arrow reflects the dynamic of the light activation observed for

- 2 each gene.
- 3
- 4