HEMERA Couples the Proteolysis and Transcriptional Activity of PHYTOCHROME INTERACTING FACTORs in Arabidopsis Photomorphogenesis

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Phytochromes (phys) are red and far-red photoreceptors that control plant development and growth by promoting the proteolysis of a family of antagonistically acting basic helix-loop-helix transcription factors, the PHYTOCHROME-INTERACTING FACTORs (PIFs). We have previously shown that the degradation of PIF1 and PIF3 requires HEMERA (HMR). However, the biochemical function of HMR and the mechanism by which it mediates PIF degradation remain unclear. Here, we provide genetic evidence that HMR acts upstream of PIFs in regulating hypocotyl growth. Surprisingly, genome-wide analysis of HMR- and PIF-dependent genes reveals that HMR is also required for the transactivation of a subset of PIF direct-target genes. We show that HMR interacts with all PIFs. The HMR-PIF interaction is mediated mainly by HMR's N-terminal half and PIFs' conserved active-phymochrome B binding motif. In addition, HMR possesses an acidic nine-amino-acid transcriptional activation domain (9aaTAD) and a loss-of-function mutation in this 9aaTAD impairs the expression of PIF target genes and the destruction of PIF1 and PIF3. Together, these in vivo results support a regulatory mechanism for PIFs in which HMR is a transcriptional coactivator binding directly to PIFs and the 9aaTAD of HMR couples the degradation of PIF1 and PIF3 with the transactivation of PIF target genes.

INTRODUCTION

Light is one of the most influential environmental cues for plants, not only because it is the ultimate energy source for photosynthesis, but also because it reflects the local growth conditions as well as diurnal and seasonal time (Franklin and Quail, 2010; Kami et al., 2010). Therefore, plants have evolved a high degree of phenotypic plasticity to fine-tune their developmental programs in response to changes in environmental light cues. During seedling development, the absence or presence of light leads to morphologically distinct developmental programs. Arabidopsis thaliana seedlings that germinate under the ground or in the dark adopt a dark-grown developmental program called skotomorphogenesis, which promotes the elongation of the embryonic stem, or hypocotyl, and represses leaf expansion and chloroplast development. In contrast, when emerging from the ground or exposed to light, seedlings switch to a light-grown developmental program called photomorphogenesis, which restricts hypocotyl growth and promotes leaf expansion and chloroplast biogenesis (Chen and Chory, 2011). The switch to the photomorphogenetic program is driven by massive reprogramming of the transcriptome (Leivar et al., 2009). Up to one-third of Arabidopsis nuclear-encoded genes are differentially expressed between dark- and light-grown wild-type seedlings (Ma et al., 2001).

Photomorphogenesis is initiated by a suite of photoreceptors, which can collectively sense the entire light spectrum ranging from UV-B to far-red light (Kami et al., 2010; Rizzini et al., 2011; Christie et al., 2012; Wu et al., 2012). Among these photoreceptors, the red (R) and far-red (FR) light-sensing phytochromes (phys) are essential for establishing photomorphogenesis (Franklin and Quail, 2010). Phys are bilin-containing proteins that can be photoconverted between two relatively stable forms: a R light-absorbing inactive Pr form and a FR light-absorbing active Pfr form (Rockwell et al., 2006; Nagatani, 2010). In Arabidopsis, phyA and phyB are the most prominent phys, and they monitor continuous FR and R light, respectively.

Phys are responsible for almost the entire reprogramming of the transcriptome in response to R light (Tepperman et al., 2006; Leivar et al., 2009; Hu et al., 2013). A central mechanism by which phys initiate photomorphogenesis is by repressing the steady state levels of a family of antagonizing transcription factors called PHYTOCHROME INTERACTING FACTORs (PIFs) (Leivar and Quail, 2011; Park et al., 2012). The PIFs belong to subfamily 15 of the basic helix-loop-helix (bHLH) super protein family in Arabidopsis, which includes seven members: PIF1 and PIF3-8 (Bailey et al., 2003; Heim et al., 2003; Toledo-Ortiz et al., 2003; Leivar and Quail, 2011). All PIFs contain a C-terminal bHLH domain for DNA binding and dimerization as well as an N-terminal Active Phytochrome B binding (APB) motif, which preferentially binds to the Pfr form of phyB (Khanna et al., 2004; Leivar and Quail, 2011). PIF1 and PIF3 contain an additional Active Phytochrome A binding (APA) motif in their N termini for interacting with activated phyA (Khanna et al., 2004; Leivar and Quail, 2011). In addition to
PIFs, PHYTOCHROME INTERACTING FACTOR3-LIKE1 (PIL1), another member of subfamily 15, also contains an APB motif and can interact with phyB (Khanna et al., 2004; Luo et al., 2014). PIFs act as either transcriptional activators or repressors (Huq et al., 2004; Leivar et al., 2009, 2012; Shin et al., 2009; Toledo-Ortiz et al., 2010; Leivar and Quail, 2011; Hornitschek et al., 2012). A number of PIFs, including PIF1, PIF3, PIF4, PIF5, and PIF7, promote hypocotyl growth (Huq and Quail, 2002; Fujimori et al., 2004; Huq et al., 2004; Khanna et al., 2004; Oh et al., 2004; Al-Sady et al., 2008; Lorrain et al., 2009; Li et al., 2012). Direct-target genes induced by PIF1, 3, 4, and 5, such as PIL1, ARABIDOPSIS THALIANA HOMEOBOX PROTEIN2 (ATHB-2), INDOLEACETIC ACID-INDUCED PROTEIN29 (IHA29), and XYLOGLUCAN ENDOTRANSGLYCOSYLASE7 (XTR7), encode transcription factors or enzymes involved in plant growth (Leivar et al., 2009, 2012; Oh et al., 2009, 2012; Hornitschek et al., 2012; Li et al., 2012; Zhang et al., 2013; Bernardo-García et al., 2014). PIF1, PIF3, and PIF5 also regulate nuclear-encoded genes required for chloroplast development (Huq et al., 2004; Moon et al., 2008; Leivar et al., 2009; Shin et al., 2009; Stephenson et al., 2009). Most PIFs accumulate in dark-grown seedlings, and photoactivated phyb bind to PIFs and trigger their phosphorylation and subsequent degradation by ubiquitin proteasome-dependent proteolysis to initiate photomorphogenetic responses (Al-Sady et al., 2006; Lorrain et al., 2008; Shen et al., 2008; Ni et al., 2013, 2014). Consistent with this model, a quadruple pif1 pif3 pif4 pif5 (pifq) mutant shows constitutive photomorphogenetic phenotypes in the dark, including a short hypocotyl phenotype as hmr-5 pifq seedlings showed the same defects in the expression of plastid-encoded marker genes transcribed by the plastid-encoded plastid RNA polymerase (PEP) but an enhanced expression of those transcribed by the nuclear-encoded plastid RNA polymerase (Figure 1C) (Pfalz et al., 2006; Chen et al., 2010b), indicating that HMR plays a separate, PIF-independent role in PEP-mediated plastidial gene expression. This conclusion is consistent with the nuclear and plastidial dual localization of HMR and the proposed role of plastid-localized HMR, also called pTAC12, as an essential component of PEP (Pfalz et al., 2006; Chen et al., 2010b), indicating that HMR plays a separate, PIF-independent role in PEP-mediated plastidial gene expression. This conclusion is consistent with the nuclear and plastidial dual localization of HMR and the proposed role of plastid-localized HMR, also called pTAC12, as an essential component of PEP (Pfalz et al., 2006; Chen et al., 2010b; Williams-Carrier et al., 2014). Together, these genetic data demonstrate that HMR regulates downstream photomorphogenetic responses by playing separate roles: a PIF-dependent role in regulating hypocotyl growth and a PIF-independent role in promoting chloroplast development.

RESULTS

HMR Acts Genetically Upstream of PIFs in Regulating Hypocotyl Growth

The hmr mutant is impaired in both the light-mediated inhibition of hypocotyl growth and promotion of chloroplast development (Chen et al., 2010b; Chen and Chory, 2011). Because PIF1 and PIF3 fail to be degraded in the hmr mutant in the light (Chen et al., 2010b; Galvão et al., 2012), we asked whether the phenotypes of hmr depend on the accumulation of PIFs. To that end, we crossed a null hmr allele, hmr-5, with the pifq quadruple mutant (Leivar et al., 2008b) to generate a hmr-Spifq quintuple mutant. Interestingly, the long hypocotyl phenotype of hmr-5 was largely rescued in hmr-Spifq seedlings (Figures 1A and 1B), indicating that the long hypocotyl phenotype of hmr-5 is due to the accumulation of PIFs. The fact that hmr-5 pifq seedlings are still slightly taller than pifq seedlings implies that HMR might regulate other PIFs besides the four tested. Similar to the phenotypes in the light, hmr-5 pifq seedlings showed the same hypocotyl phenotype as pifq in the dark (Figures 1A and 1B), indicating that pifq is epistatic to hmr in regulating hypocotyl elongation. Together with our previous results that hmr partially represses the short hypocotyl phenotype of the constitutively active phyB allele, YHB, in the dark (Chen et al., 2010b; Galvão et al., 2012), these results indicate that HMR acts genetically between phy and PIFs in regulating hypocotyl growth.

In contrast to the hypocotyl phenotype, the albino phenotype of hmr-5 was not rescued in hmr-5 pifq seedlings (Figure 1A). Similar to hmr-5, the hmr-5 pifq seedlings showed the same defects in the expression of plastid-encoded marker genes transcribed by the plastid-encoded plastid RNA polymerase (PEP) but an enhanced expression of those transcribed by the nuclear-encoded plastid RNA polymerase (Figure 1C) (Pfalz et al., 2006; Chen et al., 2010b), indicating that HMR plays a separate, PIF-independent role in PEP-mediated plastidial gene expression. This conclusion is consistent with the nuclear and plastidial dual localization of HMR and the proposed role of plastid-localized HMR, also called pTAC12, as an essential component of PEP (Pfalz et al., 2006; Chen et al., 2010b; Williams-Carrier et al., 2014). Together, these genetic data demonstrate that HMR regulates downstream photomorphogenetic responses by playing separate roles: a PIF-dependent role in regulating hypocotyl growth and a PIF-independent role in promoting chloroplast development.

HMR Is Required for the Transactivation of a Set of PIF Target Genes

PIFs act as transcriptional activators in the dark or under shade conditions to promote hypocotyl growth (Huq et al., 2004; Al-Sady et al., 2008; de Lucas et al., 2008; Shen et al., 2008; Hornitschek et al., 2009). To investigate how HMR regulates the functions of PIFs, we performed microarray analysis to examine how PIF-regulated genes are altered genome-wide in the hmr mutant. Our microarray analysis identified 1348 genes that were changed statistically significantly and by 2-fold between 4-d-old red-light-grown hmr-5 and wild-type Columbia-0 (Col-0) (Supplemental Data Set 1). To determine how many
HMR-dependent genes are PIF regulated, we compared the 1348 HMR-dependent genes with a previously defined set of 1028 PIF-regulated genes (Leivar et al., 2009), and we found that 203 PIF-regulated genes were significantly changed by at least twofold in hmr-5 (Figure 2A; Supplemental Data Set 2). The 203 HMR-dependent, PIF-regulated genes include 62 PIF-induced genes and 141 PIF-repressed genes (Supplemental Data Set 2) (Leivar et al., 2009). Because the steady state levels of PIF1 and PIF3 are elevated in hmr mutants (Chen et al., 2010b), it was expected that PIF-induced genes would be upregulated and PIF-repressed genes downregulated in hmr-5. However, this was not entirely the case. While the majority of the 141 PIF-repressed genes were downregulated in hmr-5 (Supplemental Data Set 2), surprisingly, among the 62 HMR-dependent PIF-induced genes, only 27 were upregulated in hmr-5; we named this group the Class A genes (Figure 2B). In contrast, more than half of the 62 HMR-dependent PIF-induced genes were downregulated (Figure 2B; Supplemental Data Set 2), indicating that the transactivation of this set of PIF-induced genes is impaired in hmr-5. Because the 1028 PIF-regulated genes were defined based on their expression in pifq mutants and Col-0, some of them might not be direct-target genes of PIFs (Leivar et al., 2009; Pfeiffer et al., 2014). Recent ChIP-seq and RNA-seq analyses have identified genes that are bound and regulated by individual PIFs or a combination of PIF1, 3, 4, and 5 (Hornitschek et al., 2012; Oh et al., 2012; Zhang et al.,

**Figure 1.** HMR Regulates Hypocotyl Growth and Chloroplast Biogenesis through Separate PIF-Dependent and PIF-Independent Pathways.  
(A) Images of 4-d-old Col-0, pifq, hmr-5, and hmr-5 pifq mutants grown in 10 μmol m⁻² s⁻¹ R light (upper panel) and in the dark (lower panel).  
(B) Hypocotyl measurements of the seedlings shown in (A). Error bars represent standard errors.  
(C) qRT-PCR analyses of the relative expression levels of representative PEP- and nuclear-encoded plastid RNA polymerase (NEP)-dependent genes in 4-d-old Col-0, pifq, hmr-5, and hmr-5 pifq seedlings grown in 10 μmol m⁻² s⁻¹ R light. Transcript levels were calculated relative to those of PP2A. Error bars represent the sd of three replicates.
A total of 338 genes have been recognized as PIF direct-target genes (Pfeiffer et al., 2014), among which 301 genes are represented in the Affymetrix Arabidopsis ATH1 array (Supplemental Data Set 3). To further determine whether HMR affects the transactivation of PIF direct-target genes, we analyzed the expression of these 301 PIF direct-target genes in hmr-5 mutants. We found that 43 of them were changed statistically significantly and by 2-fold between 4-d-old red-light-grown hmr-5 and wild-type Col-0 (Figure 2A) and that 106 genes, or 31%, of the 301 PIF direct-target genes were changed statistically significantly by 1.5-fold in hmr-5 (Supplemental Data Set 4). Strikingly, 41 genes, or 71%, of the 58 HMR-dependent PIF-induced direct-target genes were downregulated in hmr-5 (Figure 2B), and we named these 41 genes the Class B genes. Interestingly, the Class B genes include some of the best characterized PIF direct-target genes.

Figure 2. HMR Is Required for the Activation of a Distinct Set of PIF Target Genes.

(A) Venn diagram showing that 203 of the 1348 HMR-dependent genes overlap with the previously defined 1028 PIF-regulated genes (Leivar et al., 2009) and that 43 HMR-dependent genes belong to the 301 PIF direct-target genes (Pfeiffer et al., 2014).

(B) Pie charts showing that the majority of HMR-dependent, PIF-induced genes are downregulated in hmr-5. Among the 62 HMR-dependent PIF-induced genes (Supplemental Data Set 2), 27 genes (the Class A genes), or 43%, were induced in hmr-5; surprisingly, 57% of these genes were downregulated in hmr-5. Among the 58 PIF-induced direct targets that were changed statistically significantly by 1.5-fold in hmr-5 (Supplemental Data Set 4), 41 genes (the Class B genes), or 71%, were downregulated.

(C) qRT-PCR analyses of selected PIF-induced genes in 4-d-old Col-0, pifq, hmr-5, and hmr-5 pifq seedlings grown in 10 μmol m⁻² s⁻¹ R light. Transcript levels were calculated relative to those of PP2A. Error bars represent the SD of three replicates. Red and blue arrows indicate increases and decreases in gene expression, respectively. The expression of the Class A genes in hmr-5 was compared with that in Col-0 using Welch’s two-sample t test. **P < 0.01 and ****P < 0.0001.

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involved in plant growth, such as ATHB-2, IAA29, and XTR7 (Hornitschek et al., 2012; Oh et al., 2012; Zhang et al., 2013). To confirm the microarray data, we examined the expression of eight representative PIF-regulated genes in Col-0, hmr-5, pifq, and hmr-5 pifq seedlings using quantitative RT-PCR (qRT-PCR). The representative genes include four Class A genes, three Class B genes, and PIL1, a well-characterized PIF-induced direct-target gene that is not included in the Affymetrix ATH1 microarray (Hornitschek et al., 2012; Oh et al., 2012; Zhang et al., 2013). The qRT-PCR results for the expression of the Class A and B genes in hmr-5 and Col-0 were consistent with those from the microarray experiments (Figure 2C). In addition, the expression of PIL1 was also downregulated in hmr-5 mutants, indicating that PIL1 also belongs to the Class B genes (Figure 2C). Moreover, the data show that the elevated expression of three of the four Class A genes in hmr-5 was reversed in hmr-5 pifq, indicating that the upregulation of the Class A genes in hmr-5 mutants is largely due to the accumulation of PIFs. Therefore, the long hypocotyl phenotype of hmr-5, indicating that the upregulation of the Class A genes in hmr-5 is partially due to the upregulation of some of the Class A genes. Strikingly, the expression of the Class B genes shows least partial downregulation in hmr-5, indicating that despite the enhanced steady state levels of PIF1 and PIF3 in hmr, they are surprisingly not transcriptionally active to induce expression of these target genes in the absence of HMR. Together, these results unexpectedly reveal that HMR plays dual and seemingly opposing roles in promoting the degradation of PIF1 and PIF3 and facilitating the transcriptional activity of PIFs to the Class B genes.

**HMR’s N-Terminal Half Interacts Directly with PIF1 and PIF3**

To elucidate the mechanisms by which HMR regulates the stability and transcriptional activity of PIF1 and PIF3, we asked whether HMR can regulate PIFs through direct interaction. We used a previously described transgenic line expressing HA-tagged HMR (HMR-HA) (Galvão et al., 2012) to examine whether PIF1 and PIF3 could be coimmunoprecipitated with HMR-HA using anti-HA antibodies. The data show that both PIF1 and PIF3 can be coimmunoprecipitated with HMR-HA (Figure 3A), indicating that HMR is associated with PIF1 and PIF3 in vivo. To demonstrate that HMR interacts directly with PIF1 and PIF3, we performed glutathione S-transferase (GST) pull-down assays using recombinant GST-HMR to pull down in vitro-translated HA-PIF1 and HA-PIF3. These experiments show that GST-HMR, but not GST alone, can pull down HA-PIF1 and HA-PIF3 (Figure 3B). The in vitro pull-down data also show that GST-HMR interacts with HA-PIF1 more strongly than HA-PIF3, although this difference was not noticeable in the in vivo coimmunoprecipitation experiments (Figures 3A and 3B). Based on these results, we conclude that HMR interacts directly with both PIF1 and PIF3.

To map the subdomains of HMR required for PIF1 binding, we utilized HMR truncation fragments fused with GST to pull down HA-PIF1 in vitro. These experiments showed that HMR’s N-terminal half (GST-HMR-N, amino acids 1 to 352), but not its C-terminal half (GST-HMR-C, amino acids 352 to 527), could pull down HA-PIF1 (Figure 3C), indicating that the N-terminal half of HMR is both required and sufficient for PIF1 binding.

The HMR-PIF1 Interaction Is Mediated by a Dimer or Oligomer Form of PIF1’s APB Motif

To determine PIF1 and PIF3’s subdomains that are required for their interaction with HMR, we tested the interaction between GST-HMR and a series of N-terminal deletion fragments of PIF1. These experiments indicate that the strong interaction between PIF1 and HMR requires the APB motif of PIF1 (Figure 4A). HMR also interacts weakly with PIF1’s C-terminal fragment (amino acids 209 to 478), including the bHLH domain. To test whether the N-terminal halves of PIF1 and PIF3 are sufficient for binding with HMR, we used GST-HMR to pull down either PIF1 or PIF3’s N termini, which contain the APA and APB motifs (Khanne et al., 2004), or their C termini, which contain the bHLH domain (Figure 4B). Surprisingly, although GST-HMR interacts strongly with full-length PIF1, its interaction with the N- and C-terminal fragments of PIF1 was substantially reduced (Figure 4B). Additionally, GST-HMR could pull down neither the N- nor the C-terminal fragments of PIF3 (Figure 4B). These data indicate that both the N- and C-terminal halves of PIF1 and PIF3 are required for the interaction with HMR. We reasoned that one possibility could be that the C-terminal fragments of PIF1 and PIF3 are required structurally for the interaction with HMR, perhaps through dimerization or oligomerization of the bHLH domain (Murre et al., 1989). Consistent with this hypothesis, when PIF1’s N-terminal fragment was fused to the dimeric Gal4 DNA binding domain (DBD) (Carey et al., 1989), it regained the strong interaction with HMR, whereas fusing it to the monomeric Gal4 activation domain (GAD) did not affect HMR binding (Figure 4B). These results indicate that HMR mainly interacts with the N-terminal half of PIF1 and that this interaction requires the dimerization or oligomerization domain of PIF1’s C-terminal half, likely through the bHLH domain.

To further test that the N terminus of PIF1 is responsible for its strong interaction with HMR, we took advantage of the difference in the HMR binding affinities between PIF1 and PIF3 and swapped the N- and C-terminal fragments of PIF1 and PIF3 to generate HA-tagged chimeric proteins with either PIF1’s N-terminal domain fused to PIF3’s C-terminal domain (HA-1N3C) or PIF3’s N-terminal domain fused to PIF1’s C-terminal domain (HA-3N1C) (Figure 4C). GST-HMR interacted strongly with HA-1N3C but not HA-3N1C (Figure 4C), confirming that the strong HMR-PIF1 interaction is determined by PIF1’s N-terminal fragment.

We then asked which region within the PIF1 N terminus is responsible for the HMR interaction. Because the C-terminal fragment of PIF3 does not interact with HMR (Figure 4B), we fused two truncation fragments of the N-terminal half of PIF1 to PIF3’s C-terminal domain, HA-1N1-3C and HA-1N2-3C, and tested their interaction with GST-HMR (Figure 4C). The data demonstrate that HA-1N2-3C, which contains the PIF1 fragment spanning amino acids 1 through 163, confers strong interaction with HMR; this region of PIF1 contains both the APA and APB motifs (Figure 4C). When the APB of PIF1 alone was fused with the C-terminal fragment of PIF3, the recombinant protein, HA-1APB-3C, also interacted strongly with GST-HMR (Figure 4C), indicating that the APB motif of PIF1 is sufficient for the interaction with HMR. The APB and APA motifs of PIFs are involved in the interaction with active forms of phyB and phyA, respectively (Khanne et al., 2004; Al-Sady et al., 2008; Shen et al., 2008). It has been shown that the
bHLH domain is either not required for the interaction between phyB and PIFs or it attenuates this interaction (Khanna et al., 2004). Therefore, although both HMR and phyB interact with the APB motif, HMR and phyB might bind to different interfaces or distinct residues in APB. The conserved residues Glu-41 and Leu-42 in the APB of PIF1 are essential for the interaction with phyB, and mutations N144A and L95A in the APA of PIF1 abolish its interaction with phyA (Shen et al., 2008). However, mutations in these residues did not affect PIF1’s interaction with HMR (Supplemental Figure 1), confirming that the HMR binding sites differ from those for phy. Taken together, these results indicate that the APB motif of PIF1 is both required and sufficient for its strong interaction with HMR. The HMR-PIF1 interaction appears to involve residues different from the phy-PIF1 interaction and require the dimerization or oligomerization function of the bHLH domain from the C-terminal half of PIF1.

HMR Interacts with All PIFs and PIL1 through the Conserved APB Motif

PIFs belong to subfamily 15 of the Arabidopsis bHLH superfamily (Toledo-Ortiz et al., 2003). Among the 15 members of subfamily 15, 12 contain an APB motif (Khanna et al., 2004). These include the seven PIFs (PIF1 and PIF3-8), PIL1, and four less-characterized bHLH proteins (bHLH127, bHLH023, bHLH119, and bHLH056) (Khanna et al., 2004). We fused the respective APB motifs from PIF3-PIF8 and PIL1 with the C-terminal fragment of PIF3 and examined their ability to bind GST-HMR using pull-down assays. The data show that GST-HMR interacts with the APB motifs from all PIFs and from PIL1 (Figure 5A). Consistent with this, GST-HMR could also pull down all full-length PIFs and PIL1 (Figure 5B). In contrast, GST-HMR failed to pull down ALCATRAZ (ALC), another subfamily 15 bHLH protein that does not contain an APB motif (Figure 5B) (Khanna et al., 2004). Together, these results demonstrate that HMR interacts directly with all PIFs and PIL1 through their conserved APB domains.

HMR Possesses an Acidic 9-Amino-Acid Transcription Activation Domain

The direct interaction between HMR and PIFs raises a hypothesis that HMR binds to PIFs to modulate their stability and activity. Because HMR is required for the activation of the Class B genes (Figure 2), one possibility is that HMR might act as...
Figure 4. The HMR-PIF1 Interaction Is Mediated Mainly by PIF1’s APB Motif and Requires the bHLH Dimerization Domain.

(A) The APB motif of PIF1 is required for the strong interaction with HMR. GST pull-down assays were performed using E. coli-expressed GST-HMR or GST to pull down in vitro-translated HA-tagged full-length PIF1 or PIF1 truncation fragments. Input and pull-down fractions of HA-tagged PIF1 fragments were detected by immunoblots using anti-HA antibodies. The corresponding SDS-PAGE gels show the amount of GST or GST-HMR immobilized in each assay. I, 10% input of the indicated PIF fragment; G, GST; H, GST-HMR; PIR1, phytochrome interacting region 1; GLU, glutamate-rich region; NLS, nuclear localization signal; PIR2, phytochrome interacting region 2; APB, active phytochrome B binding motif.

(B) HMR interacts more strongly with a dimer or oligomer form of PIF1’s N-terminal half. GST pull-down assays were performed using E. coli-expressed GST-HMR or GST to pull down in vitro-translated HA-tagged full-length PIF1 and PIF3, the N- or C-terminal fragments of PIF1 and PIF3, and the N-terminal fragment of PIF1 fused with either Gal4 DBD or Gal4 activation domain. Input and pull-down fractions of the PIF fragments were detected by immunoblots using anti-HA antibodies. The corresponding SDS-PAGE gels show the amount of GST or GST-HMR immobilized in each assay.

(C) The APB motif of PIF1 confers the strong interaction with HMR in the presence of PIF’s bHLH domain. GST pull-down assays were performed using E. coli-expressed GST-HMR or GST to pull down in vitro-translated HA-tagged PIF1/PIF3 chimeric proteins. Input and pull-down fractions of the PIF fragments were detected by immunoblots using anti-HA antibodies. The corresponding SDS-PAGE gels show the amount of GST or GST-HMR immobilized in each assay.
a transcriptional coactivator. Supporting this hypothesis, full-length HMR and its C-terminal half between amino acids 254 and 527, when fused to the Gal4 DBD, were able to activate transcription in yeast (Supplemental Figure 2), suggesting that the C-terminal half of HMR contains a transcriptional activation domain. Transcriptional activation domains have been classified based on their amino acid composition into acidic, glutamine-rich, proline-rich, and serine/threonine-rich types (Mitchell and Tjian, 1989). Studies of yeast and mammalian acidic transcriptional activators have identified a nine-amino-acid transcriptional activation domain.

Figure 5. HMR Interacts with All PIFs and PIL1 through the Conserved APB Motif.

(A) HMR interacts with the APB motifs from all PIFs and PIL1. GST pull-down assays were performed using E. coli-expressed GST-HMR or GST to pull down in vitro-translated HA-tagged APB motif from indicated PIFs or PIL1 fused with the C-terminal fragment of PIF3. Input and pull-down fractions of the prey proteins were detected by immunoblots using anti-HA antibodies. The corresponding SDS-PAGE gels show the amount of GST or GST-HMR immobilized in each assay. I, 10% input of the indicated prey protein; G, GST; H, GST-HMR.

(B) HMR interacts with all PIFs and PIL1 in vitro. GST pull-down assays were performed using E. coli-expressed GST-HMR or GST to pull down in vitro-translated HA-tagged PIF4-8, PIL1, and ALC. Input and pull-down fractions of HA-tagged PIFs, PIL1, and ALC were detected by immunoblots using anti-HA antibodies. The corresponding SDS-PAGE gels show the amount of GST or GST-HMR immobilized in each assay.
activation domain (9aaTAD) conserved among well-characterized transcriptional activators, such as Gal4, Gcn4, Myc, and VP16 (Piskacek et al., 2007). Using the 9aaTAD prediction utility (Piskacek et al., 2007), we identified six putative 9aaTADs in the C-terminal half of HMR with match scores greater than 80% (Figure 6A). To test whether these predicted 9aaTADs function in yeast, we examined the transactivation activity of a series of truncation fragments of the C-terminal half of HMR. These experiments showed that the 9aaTAD from amino acids 512 to 520 is both required and sufficient for HMR’s transactivation activity in yeast (Figure 6A). This particular 9aaTAD is highly conserved among HMR orthologs from moss to higher plants (Figure 6B).

Within the 9aaTAD sequence of Arabidopsis HMR, ENLTDFLMD, the three acidic residues at the first, fifth, and ninth positions (italicized) are conserved among all HMR orthologs (Figure 6B). Substitutions of the conserved acidic residues individually to alanine greatly reduced the transactivation activity of the 9aaTAD (Figure 6C). Together, these data indicate that HMR is an acidic transcriptional activator.

The Transcriptional Activity of HMR’s 9aaTAD Is Required for the Activation of the Class B PIF Targets and Degradation of PIF1 and PIF3

To demonstrate the function of HMR’s 9aaTAD in vivo, we employed the Seattle Arabidopsis TILLING service (http://tilling.fhcrc.org/) to search for missense mutations in the 9aaTAD (Till et al., 2003). We had previously screened for mutations in the N-terminal half of HMR and reported 11 hmr alleles named hmr-4 to hmr-14 (Supplemental Figure 3) (Galvão et al., 2012). The second round of TILLING identified eight additional missense alleles named hmr-15 to hmr-22 (Supplemental Figure 3). One of these alleles, hmr-22, carries a single amino acid substitution of the conserved acidic Asp-516 in the 9aaTAD to a noncharged Asn residue (Figure 6B). The D516N mutation reduced the transactivation activity of the 9aaTAD in yeast by 46% (Figure 6C).

To determine whether the D516N mutation affects HMR’s function in phy signaling, we measured the hypocotyl growth in wild-type Arabidopsis seedlings under a series of intensities of either R or FR light. Similar to the null hmr-2 allele, hmr-22 was hypersensitive to both continuous red and far-red light (Figures 7A and 7B), and it had normal cryptochrome-mediated responses in blue light (Supplemental Figure 4), indicating that hmr-22 is impaired specifically in both phyA and phyB signaling. Seedlings of hmr-22 are also impaired in chloroplast biogenesis (Supplemental Figure 5A). Despite the obvious defect in chloroplast development at the seedling stage, both cotyledons and leaves of hmr-22 are able to turn green when they become fully expanded (Supplemental Figure 5B).

However, the rosette of the hmr-22 mutant is substantially smaller than that of Col-0 (Supplemental Figure 5B). Taken together, these results indicate that hmr-22 is a weak, viable, loss-of-function hmr allele and that the D516N mutation attenuates HMR’s functions in phy signaling and chloroplast biogenesis in vivo.

To examine the effect of the D516N mutation on the expression of the HMR-dependent PIF direct-target genes, we performed genome-wide transcriptome analysis on 4-d-old red-light-grown Col-0 and hmr-22 seedlings using the Affymetrix ATH1 microarray. The microarray analysis identified 385 genes changed significantly by twofold between Col-0 and hmr-22 (Supplemental Data Set 5) and 48 PIF direct-target genes were changed statistically significantly by 1.5-fold (Supplemental Data Set 6). The set of genes changed in hmr-22 largely overlaps with those changed in hmr-5. Among the 385 genes changed in hmr-22 mutants, 352 genes, or 91%, were also changed in hmr-5 mutants (Supplemental Figure 6). Similarly, 42, or 88%, of the 48 PIF direct-target genes changed in hmr-22 were also changed in hmr-5 (Supplemental Figure 6). Strikingly, 21 of the 25 PIF-induced direct-target genes were downregulated in hmr-22 mutants, including the four Class B marker genes (Figure 7C).

These data indicate that the expression of the Class B PIF target genes is dependent on the transcriptional activity of HMR’s 9aaTAD in vivo. Interestingly, similar to hmr-5, the expression of the four Class A genes was upregulated in hmr-22 (Supplemental Figure 7), suggesting that PIF levels might also be enhanced in hmr-22. Indeed, PIF1 and PIF3 failed to be degraded in hmr-22 in both R and FR light (Figure 7D). Because the D516N mutation does not alter the interaction between HMR and PIF1 in vitro (Supplemental Figure 8A), and because the level of HMRD516N (HMR22) in hmr-22 is similar to that of the wild-type HMR in Col-0 in both dark and light conditions (Supplemental Figure 8B), the defect in the degradation of PIF1 and PIF3 in hmr-22 is most likely dependent on the activity of HMR’s 9aaTAD. Together, these data indicate that HMR’s 9aaTAD mediates both the degradation of PIF1 and PIF3 as well as the activation of the Class B PIF targets.

HMR Plays Dual Opposing Roles in Regulating Hypocotyl Growth

The degradation of PIF1 and PIF3 inhibits hypocotyl growth (Leivar et al., 2008b; Shin et al., 2009); in contrast, the expression of the Class B genes, including PIL1, IAA29, ATHB2, and XTR7, contributes to hypocotyl growth (Leivar et al., 2009; Hornitschek et al., 2012; Zhang et al., 2013). Therefore, HMR appears to play dual opposing roles in regulating hypocotyl growth. We hypothesize that the phenotype of hmr mutants should reflect a balance between these two opposing roles of HMR. We reasoned that the long hypocotyl phenotypes of hmr mutants in continuous R and FR light is mainly due to the accumulation of PIF1 and PIF3 because the Class B genes are expressed at low basal levels under continuous light (Leivar et al., 2009; Hornitschek et al., 2012; Zhang et al., 2013) and a further decrease in the expression of the Class B genes might not have a major impact on hypocotyl growth. Therefore, a better condition to demonstrate the physiological significance of HMR’s transactivation activity is where PIF degradation is attenuated and the Class B genes are expressed at higher levels. One such condition is in the phyB-9 background, where PIF3 degradation is impaired in the light and the four Class B genes are induced (Figures 7E and 7F) (Hornitschek et al., 2012). We then generated hmr-22 phyB-9 double mutant. Indeed, the hmr-22phyB-9 mutant had a similar level of PIF3 as phyB-9 in the light (Figure 7E), confirming that the HMR-mediated PIF3 degradation is also phyB dependent.

As predicted, three of the four Class B genes were downregulated in hmr-22 phyB-9 compared with phyB-9 (Figure 7F). More interestingly, in contrast to the long hypocotyl phenotype of hmr-22 seedlings, the hmr-22 phyB-9 seedlings were shorter compared with phyB-9 seedlings (Figures 7G and 7H), indicating that the
activity of HMR’s 9aaTAD is required for promoting hypocotyl growth. Together with the tall hypocotyl phenotype of hmr mutants, these results demonstrate that HMR plays opposing roles in regulating hypocotyl growth.

**Fusion of VP16 to HMR22 Rescues Its Defects in the Transactivation of the Class B Genes but Not PIF3 Degradation**

To further confirm the role of HMR in the activation of the Class B genes and to dissect the relationship between transactivation of the Class B genes and PIF degradation, we asked whether the defects of HMR22 could be rescued by fusing the trans-activation domain of the Herpes simplex virus activator VP16. To this end, we generated transgenic lines expressing either HMR-HA or HMR22-HA-VP16 under the 35S promoter in hmr-22. We picked two independent transgenic lines for each construct for further analysis. As expected, the two HMR-HA/hmr-22 lines (#3 and #9) rescued the long hypocotyl phenotype of hmr-22 (Figure 8A). In contrast, the two HMR22-HA-VP16/hmr-22 lines had even longer hypocotyls than that of hmr-22 (Figure 8A). Both HMR-HA and HMR22-HA-VP16 were able to rescue the defects

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**Figure 6. HMR Possesses a Conserved Acidic Nine-Amino-Acid Transactivation Domain.**

(A) Dissection of the transcriptional activation domain in the C-terminal half of HMR using yeast. A series of truncation fragments of HMR’s C-terminal half were fused to the GAL4 DBD in pGBKT7. The schematics of the constructs are shown in the left panel. The six predicted 9aaTADs are labeled with either pink or red blocks. TAD1, amino acids 255 to 263; TAD2, amino acids 307 to 315, TAD3, amino acids 339 to 347; TAD4, amino acids 449 to 457; TAD5, amino acids 502 to 510; and TAD6, amino acids 512 to 520. Murine p53 (53) and lamin (Lam) were used as positive and negative controls, respectively. Serial dilutions of Y2HGold (Clontech) yeast strains containing the indicated vectors were grown on either SD/-Trp or SD/-Trp/+Aur-eobasidin A media (right panels). Strains showing positive transactivation results are outlined with a red frame. (B) Amino acid sequence alignment of TAD6 among HMR orthologs. The blue bar labels the 9aaTAD, and the red stars indicate the three acidic residues that are conserved among all HMR orthologs. (C) The acidic residues of HMR’s TAD are required for its full transcriptional activity in yeast. Yeast β-galactosidase liquid assays showing the transactivation activity of the wild type and mutated TADs with the indicated amino acid substitution. The D516N mutation in hmr-22 reduces the activity of HMR’s 9aaTAD by 46%. Error bars represent the standard deviations of three replicates. ****P < 0.0001.
Figure 7. HMR’s 9aaTAD Mediates Both the Expression of the Class B Genes and the phyB-Dependent Degradation of PIF1 and PIF3 in Vivo.

(A) Fluence response curves for R light. Relative hypocotyl length of 4-d-old Col-0, phyB-9, hmr-2, and hmr-22 seedlings grown in various intensities of R light or in the dark.

(B) Fluence response curves for FR light. Relative hypocotyl length of 4-d-old Col-0, phyA-211, hmr-2, and hmr-22 seedlings grown in various intensities of FR light or in the dark. Hypocotyl lengths in (A) and (B) are given relative to the average hypocotyl length of each genotype in the dark, and error bars represent standard errors from three replicates.

(C) Microarray and qRT-PCR data for the mRNA levels of the indicated PIF-induced genes in 4-d-old Col-0 and hmr-22 seedlings grown in 10 μmol m⁻² s⁻¹ R light. In each panel, the filled columns represent data from the microarray analysis and the open columns represent confirmation data by qRT-PCR. Transcript levels from the qRT-PCR experiments were calculated relative to those of PP2A. Blue arrows indicate decrease in gene expression. Error bars represent the SD of three replicates.

(D) Immunoblot analysis of PIF1 and PIF3 protein levels in 4-d-old Col-0 and hmr-22 seedlings grown in the dark, 10 μmol m⁻² s⁻¹ R light, or 10 μmol m⁻² s⁻¹ FR light. RPN6 was used as a loading control.
of hmr-22 in the expression of the Class B genes (Figure 8B), further confirming that the activation of these PIF targets is dependent on the transactivation activity of HMR. Interestingly, whereas the HMR-HA/hmr-22 lines rescued the defect of hmr-22 in PIF3 degradation, the HMR22-HA-VP16/hmr-22 lines accumulated the same level of PIF3 as hmr-22 (Figure 8C), indicating that VP16 is not able to mediate PIF3 degradation. The combination of the elevated levels of PIF3 and the enhanced expression of the Class B genes in the HMR22-HA-VP16/hmr-22 lines provide an explanation for their long hypocotyl phenotype (Figure 8A). Therefore, these results indicate that the degradation of PIF3 is specifically dependent on HMR’s 9aaTAD. Mutations in the 9aaTAD could lead to a separation of the dual functions of HMR.

DISCUSSION

We have previously shown that the degradation of PIF1 and PIF3 in the light is dependent on HMR (Chen et al., 2010b; Galvão et al., 2012). However, the biochemical function of HMR and the mechanism by which HMR regulates PIF degradation were unclear. This study establishes the genetic and molecular functions of HMR in phy signaling. We show that hmr-5 pifq seedlings exhibit a similar hypocotyl phenotype as pifq seedlings in both light and dark conditions (Figures 1A and 1B). Combined with our previously published data that hmr-1 YHB partially rescues the short hypocotyl phenotype of YHB in the dark (Galvão et al., 2012), these results indicate that HMR acts genetically between phs and PIFs in regulating hypocotyl growth. This PIF-dependent role of HMR in regulating hypocotyl growth is distinct from a PIF-independent role of HMR in plastidial gene expression and chloroplast development (Figure 1C). This study elucidates the molecular mechanism by which HMR regulates PIFs. We demonstrate that HMR is a transcriptional coactivator possessing an acidic 9aaTAD and interacting directly with all PIFs and PIL1. Surprisingly, the transactivation activity of the 9aaTAD is required for both the activation of a distinct set of growth-relevant PIF target genes, the Class B genes, including PIL1, ATHB-2, IAA29, and XTR7, as well as the degradation of PIF1 and PIF3. These in vivo data support a mechanism in which the 9aaTAD of HMR couples the degradation of PIF1 and PIF3 with the transactivation of PIF target genes (Figure 8D). We propose that HMR imposes a regulatory module to remove “spent” PIF1 and PIF3 during the transactivation of a distinct set of growth-relevant PIF target genes, and this function of HMR enables a tightly controlled mode of hypocotyl growth in the light.

HMR Is a Transcriptional Coactivator Required for the Expression of a Distinct Set of PIF Targets

The function of HMR as a transcriptional coactivator for PIF targets is first supported by the genetic evidence that despite the elevated levels of PIF1 and PIF3 in hmr-5 mutants, the Class B PIF target genes fail to be activated (Figure 2C). Our data demonstrate that HMR is an acidic transcriptional coactivator (Figure 6) interacting directly with all PIFs and PIL1 (Figures 4 and 5). The 9aaTAD of HMR is highly conserved among HMR orthologs from various land plants, and substitutions of the conserved acidic residues in the 9aaTAD individually to alanine greatly reduced its transactivation activity in yeast (Figures 6B and 6C), indicating that the function of HMR as a transcriptional coactivator is evolutionarily conserved from moss to higher plants. The characterization of the weak allele hmr-22, which carries a single amino acid substitution of the conserved acidic Asp-516 in the 9aaTAD to an uncharged Aan residue, provides in vivo evidence that the transactivation activity of HMR is required for the function of HMR in phy signaling, in particular the activation of some of the Class B genes (Figures 7A to 7D). Fusion of VP16 to HMR22 rescues its defects in the transactivation of the Class B genes (Figure 8B), further confirming the conclusion that the expression of the Class B PIF targets is dependent on the transactivation activity of HMR. Consistent with these results, our previous studies have shown that although the hmr-1 YHB double mutant had a similar level of PIF3 as the control PBG line in the dark, the expression of the Class B genes in hmr-1 YHB was much lower compared with that in PBG (Galvão et al., 2012), indicating that HMR is also required for the activation of these PIF targets in YHB.

Because expression of the Class B genes, including PIL1, ATHB-2, IAA29, and XTR7, correlates with hypocotyl growth (Leivar et al., 2009; Hornitschek et al., 2012; Zhang et al., 2013), the function of HMR in the transactivation of these genes is expected to promote hypocotyl growth. To demonstrate this function of HMR, we examined the effect of the hmr-22 mutation in the phyB-9 mutant, where PIF3 degradation is impaired and the expression of the Class B genes is activated (Figures 7E and 7F). The hmr-22 phyB-9 double mutant had reduced expression of the Class B genes and hypocotyl growth compared with phyB-9 (Figures 7E to 7H), demonstrating that the transactivation activity of HMR plays a role in promoting hypocotyl growth. This conclusion is further supported by the long hypocotyl phenotype of the HMR22-HA-VP16/hmr-22 lines, in which the defects in the transactivation activity of HMR22 were

Figure 7. (continued).

(E) Immunoblot analysis of PIF3 protein levels in 4-d-old phyB-9 and hmr22 phyB9 seedlings grown in continuous 10 μmol m⁻² s⁻¹ R light. Dark-grown Col-0 and pifq were used as controls for the PIF3 band. RPN6 was used as a loading control.
(F) qRT-PCR data for the mRNA levels of the indicated PIF-induced genes in 4-d-old Col-0, phyB-9, and hmr22 phyB-9 seedlings grown in 10 μmol m⁻² s⁻¹ R light. Transcript levels from the qRT-PCR experiments were calculated relative to those of PP2A. Blue arrows indicate decrease in gene expression. Error bars represent the mean of the three replicates. The expression of the Class B genes in hmr-22 phyB-9 was compared with that in hmr-22 using Welch’s two-sample t test. ***P < 0.001 and **P < 0.01.
(G) Representative images of 4-d-old Col-0, phyB-9, and hmr22 phyB9 seedlings grown in continuous 10 μmol m⁻² s⁻¹ R light.
(H) Hypocotyl measurements of seedlings shown in (G). Error bars represent standard errors.
rescued by VP16 but the PIF3 level remained high (Figures 8A to 8C). Together, these results show that HMR is a transcriptional coactivator binding directly to PIFs to activate the expression of the Class B PIF targets; this function of HMR promotes hypocotyl growth, which opposes the other role of HMR in mediating the degradation of PIF1 and PIF3 (Figure 8D).

How the 9aaTAD of HMR activates PIF targets is still unknown. Studies on prototypic acidic transcription activators, such as VP16, have shown that transcriptional activation domains interact directly with subunits of basal transcription factors, including TFIIA, TFIIB, TFIID, TFIILH, and subunits of the mediator complex to facilitate the assembly and/or the stability of the RNA polymerase II preinitiation complex at the transcriptional initiation site (Kobayashi et al., 1995; Uesugi et al., 1997; Hall and Struhl, 2002; Langlois et al., 2008; Borggreve and Yue, 2011; Vojnic et al., 2011). These basic transcriptional activation mechanisms are likely conserved among eukaryotes because prototypic acidic transcriptional activation domains, such as the ones from Gal4 and VP16, are potent activators in yeast, animal, and plant cells (Ptashne, 1988; Sadowski et al., 1988; Schwechheimer et al., 1998). Consistent with this notion, this study shows that HMR’s 9aaTAD can also recruit protein degradation machineries for PIF1 and PIF3, such as E3 ubiquitin ligases or the proteasome. As such, HMR mediates the turnover of “spent” PIF1 and PIF3 during the transaction of the Class B genes. HMR enables a mechanism to tightly control hypocotyl growth by light.
to PIFs and the 9aaTAD at its C terminus interacts with subunits of the general transcriptional machinery (Figure 8D). Because only the expression of the Class B genes, but not the Class A genes, depends on the transcriptional activity of HMR (Figure 2C; Supplemental Figure 7), these data suggest that HMR might be associated only with the promoter regions of the Class B genes (Figure 8D). To test this model, we used the HMR-HA lines and examined whether the promoter regions of PIL1, ATHB-2, IAA29, and XTR7 can be pulled down by HMR-HA using chromatin immunoprecipitation (ChiP). However, these ChiP experiments yielded negative results, which might be due to low efficiency of the ChiP experiments because HMR does not directly bind to DNA; alternatively, it is also possible that the PIF-HMR complex is rather unstable at the promoter regions. Further investigation is needed to distinguish these possibilities. This new model in Figure 8D also suggests that HMR might interact directly with subunits of the general transcriptional machinery. One possible candidate is the mediator subunit MED25. MED25 interacts directly with the transcriptional activation domain of VP16 (Vojnic et al., 2011). The ortholog of MED25 in Arabidopsis, known as PFT1 (Phytochrome and Flowering Time1) (Cerdán and Chory, 2003; Bäckström et al., 2007), was identified as a factor involved in phy signaling and flowering time (Cerdán and Chory, 2003; Iñigo et al., 2012a, 2012b; Klose et al., 2012). Our future investigation will test the hypothesis that HMR’s 9aaTAD interacts directly with PFT1 to promote gene activation.

The 9aaTAD of HMR Mediates Both Degradation of PIF1 and PIF3 and Transactivation of PIF Targets

The current model for phy signaling suggests that phys promote the degradation of PIFs to repress the expression of PIF target genes, including the growth-relevant PIL1, ATHB-2, IAA29, and XTR7 (Al-Sady et al., 2006; Ni et al., 2014). In this model, the phy-dependent degradation of PIFs serves as a restraint for the function of PIFs in promoting hypocotyl growth and degradation of PIFs is a separate process from PIF’s transcriptional activity. Our results reveal an alternative mechanism, in which degradation of PIF1 and PIF3 is linked to the transactivation of the Class B PIF target genes by HMR’s 9aaTAD. This counterintuitive relationship between the degradation of PIF1 and PIF3 and transactivation of PIF targets was unexpected revealed by our analysis of HMR-PIF-dependent genes. We found that both the degradation of PIF1 and PIF3 as well as the transactivation of the Class B PIF target genes are impaired in hmr (Figure 2C) (Chen et al., 2010b; Enjalón et al., 2012). The identification of hmr-22, which carries a single loss-of-function mutation in HMR’s 9aaTAD (Figures 6C and 7C), allowed us to specifically test whether the transactivation activity of the 9aaTAD is required for the degradation of PIF1 and PIF3. The accumulation of PIF1 and PIF3 in hmr-22 provides strong in vivo evidence supporting the notion that degradation of PIF1 and PIF3 is dependent on the transactivation activity of the 9aaTAD of HMR (Figure 7D). Because the level of PIF3 remained the same in hmr-22 phyB-9 and phyB-9 (Figure 7E), the HMR-dependent PIF3 degradation is part of the phyB-mediated mechanism of PIF3 degradation. It is intriguing that fusion of VP16 to HMR-22 only rescues the defects of HMR-22 in the activation of the Class B genes but not in PIF3 degradation (Figures 8B and 8C), indicating that not all transcriptional activation domains can mediate PIF degradation, there is something unique to the 9aaTAD of HMR that gives it the ability to mediate the degradation of PIF1 and PIF3.

Coupled degradation and activity has been shown for a number of prototypic unstable transcriptional activators in yeast and metazoan models, including Myc (Kim et al., 2003b; von der Lehr et al., 2003), VP16 (Salghetti et al., 2001), Gcn4 (Chi et al., 2001; Lipford et al., 2005), and the estrogen receptor (Reid et al., 2003). In fact, the transcriptional activation domains of most unstable transcriptional activators overlap with their degrons, the sequences for proteolysis (Muratani and Tansey, 2003). In particular, it is an acidic type, but not proline-rich or glutamine-rich, transcriptional activation domain that can mediate activator degradation (Salghetti et al., 2001). The molecular basis linking proteolysis and transcriptional activation remains elusive. However, accumulating evidence suggests that ubiquitination and subsequent proteasome-mediated degradation of activators are an integral part of transcriptional activation (Lipford and Deshaies, 2003; Muratani and Tansey, 2003; Geng and Tansey, 2012). For example, the transcriptional activity of VP16 in yeast requires its ubiquitin ligase Met30 (Salghetti et al., 2001). In the absence of Met30, VP16 is stabilized but not transcriptionally active; fusion of ubiquitin to VP16 can bypass the requirement for Met30 for its transcriptional activity (Salghetti et al., 2001). Similarly, the transcriptional activity of the protooncogene Myc in human cells is dependent on Skp2, the substrate recognition subunit of a Cullin-based E3 ubiquitin ligase for Myc degradation (Kim et al., 2003b). Therefore, one possible mechanism is that the transcriptional activation domains of VP16 and Myc are required for recruiting the E3 ubiquitin ligases for their degradation. Alternatively, the ubiquitylation of activators could be required to recruit the proteasome, which has been suggested to play an important role in transcription activation besides its conventional role in protein degradation (Lipford and Deshaies, 2003; Muratani and Tansey, 2003; Geng and Tansey, 2012). Little is known about the relationship between degradation and activity of transcriptional activators in plants. However, a few lines of evidence have begun to reveal the biological importance of protein degradation in transcriptional activation in plants. First, proteasome-dependent degradation of a transcription coactivator NPR1 (Nonexpressor of Pathogenesis-Related PR genes) has been shown to stimulate the expression of pathogen responsive genes in plant immunity (Spoel et al., 2009). Second, turnover of the mediator subunit MED25 is coupled to the activation of FLOWERING LOCUS T in floral initiation (Iñigo et al., 2012a). Lastly, proteolysis of the transcription factor MYC2 is required for its transcriptional activity in plant immune responses (Zhai et al., 2013). Interestingly, the transcriptional activation domain of MYC2 is also an acidic type (Zhai et al., 2013). In this study, we identified another acidic transcriptional activation domain that is capable of coupling protein degradation and transcriptional activation. We propose that the mechanism of coupled degradation and activation for acidic activators is an ancient regulatory mechanism evolved prior to the divergence of the plant and animal/fungal lineages.

The 9aaTAD of HMR could work in a similar mechanism as VP16 and Myc, in which it is required to recruit either E3
ubiquitin ligases or the proteasome for the degradation of PIF1 and PIF3 during the transactivation of the Class B PIF targets (Figure 8D). Components for PIF degradation could be recruited directly by HMR or through subunits of the general transcriptional machinery (Figure 8D). In either case, PIF degradation is dependent on a unique function of HMR’s 9aaTAD that is missing in VP16. It has been reported recently that the degradation of PIF3 is mediated by Cullin3-based E3 ubiquitin ligases with LRBs (Light-Response Broad-Complex/Tramtrack/Bric-a-brac) as the substrate recognition subunits (Ni et al., 2014). Interestingly, despite enhanced abundance of PIF3 in the light, the brb123 mutant shows a short hypocotyl phenotype (Ni et al., 2014), which resembles a pif3 mutant as opposed to a PIF3 overexpression line (Kim et al., 2003a; Al-Sady et al., 2008). Although it was suggested that the phenotype of brb123 is due to an enhanced level of phyB, an alternative explanation is that the LRB-containing E3 ubiquitin ligases are required for the function of PIF3 in vivo. These observations are consistent with our model that the ubiquitin proteasome-mediated degradation of PIF3 is coupled to the expression of PIF targets that promote hypocotyl growth. It would be interesting to investigate if HMR is involved in the ubiquitylation of PIF3 by LRBs. Another well-characterized E3 ubiquitin ligase in light signaling is COP1 (CONSTITUTIVE PHOTOMORPHOGENETIC1) (Chen et al., 2010a). Recently, COP1 has been shown to interact directly with PIF1 (Xu et al., 2014). However, because COP1 is not required for PIF3 degradation (Bauer et al., 2004), it is unlikely that COP1 is an E3 ubiquitin ligase directly regulating the activity of PIF1 and PIF3. However, further investigations are needed to examine these possibilities.

**Biological Significance of the HMR-Dependent Regulation of PIF Stability and Activity**

It has been proposed that linking stability and activity of transcription activators is required to remove “spent” activators and thus to tightly control transcription and downstream activator-mediated responses (Salgheetti et al., 2001; Lipford and Deshaies, 2003). We suggest that the HMR-dependent PIF degradation mechanism removes “spent” PIF1 and PIF3 and enables phy to tightly control hypocotyl growth in the light (Figure 8D). It is intriguing that all prototypic unstable transcriptional activators are regulated by their own transcriptional activation domains (Muratan and Tansey, 2003), whereas PIF1 and PIF3 are regulated by the 9aaTAD of the cofactor HMR. We propose that this unique configuration of the HMR-PIF system accommodates the need to switch between two dramatically different modes of hypocotyl growth: rapid hypocotyl growth in the dark and quantitatively controlled hypocotyl growth in the light. In the dark, PIFs accumulate to high levels and the expression of the Class B genes is highly activated (Leivar et al., 2009), indicating that the degradation and activity of PIFs are uncoupled in dark-grown seedlings to maximize the speed of hypocotyl growth. Because hmr mutants show no obvious hypocotyl phenotype in the dark (Chen et al., 2010b) and because the level of HMR remains low in the absence of light (Galvão et al., 2012), HMR does not play a major role in hypocotyl growth the dark and HMR’s 9aaTAD is likely not responsible for the expression of the Class B genes in the dark. Although the transcriptional activation domains of PIFs have not been precisely identified, PIFs have been shown to contain intrinsic transactivation activity (Hu et al., 2004; Al-Sady et al., 2008; de Lucas et al., 2008; Leivar et al., 2008a; Shen et al., 2008; Hornitschek et al., 2009). It is conceivable that the Class B genes in the dark are activated by the intrinsic transcriptional activation domains of PIFs. If this is the case, our model predicts that the intrinsic transcriptional activation domains in PIFs are not capable of coupling their degradation with their transcriptional activity. The onset of light activates the phy-mediated PIF degradation and enhances the accumulation of HMR (Al-Sady et al., 2006; Galvão et al., 2012). Binding of HMR to PIFs enables the coupling of the phyB-mediated degradation of PIF1 and PIF3 with the transactivation of the Class B PIF targets (Figure 8D). Therefore, phyB and HMR together serve as the switch to couple the stability and activity of PIF1 and PIF3 in the light, this mechanism allows hypocotyl growth to be tightly controlled in the light.

**METHODS**

**Plant Materials, Growth Conditions, and Hypocotyl Measurement**

The hmr-1 through hmr-14 alleles have been previously characterized (Chen et al., 2010b; Galvão et al., 2012). The ABRC accession number for the newly identified hmr-15 to hmr-22 alleles as well as their genotyping cleaved amplified polymorphic sequences (Konieczny and Ausubel, 1993) markers, are listed in Supplemental Table 1. All hmr-TILLING alleles were backcrossed to Col-0 at least three times before being used for experiments. Wild-type Col-0, phyB-9 (Col-0), and phyA-217 (Col-0) mutants were used as controls for physiological studies. The pifq mutant was previously described (Leivar et al., 2008b). Seeds were surface-sterilized and plated on half-strength Murashige and Skoog growth medium without sucrose as described previously (Chen et al., 2010b). Seeds were stratified in the dark at 4°C for 5 d. Seedlings were grown at 21°C in an LED chamber (Percival Scientific) under the indicated light conditions. Fluence rates of light were measured using an Apogee PS200 spectroradiometer (Apogee Instruments).

For the measurement of hypocotyl length, seedlings were scanned using an Epson Perfection V700 photo scanner, and hypocotyls were measured using NIH ImageJ software (http://rsb.info.nih.gov/ij-image). Data were collected from at least 30 seedlings per genotype per treatment.

**Protein Extraction and Immunoblot**

Protein was extracted as previously described (Shen et al., 2008) with some changes. The extraction buffer consisted of 100 mM Tris-HCl, pH 7.5, 100 mM NaCl, 5 mM EDTA, pH 8.0; 5% SDS, 20% glycerol, 20 mM DTT, 40 mM β-mercaptoethanol, 2 mM PMSF, 1× EDTA-free protease inhibitor cocktail (Roche), 80 μM MG132 (Sigma-Aldrich), 80 μM MG115 (Sigma-Aldrich), 1% phosphatase inhibitor cocktail 3 (Sigma-Aldrich), and 10 mM N-ethylmaleimide. Seedlings were ground directly in extraction buffer in a 1:2 (mg/mg) ratio in dim green light, boiled for 10 min, and then centrifuged at 15,000g for 10 min at room temperature. The supernatant was then saved for further analysis.

For immunoblots, proteins were separated on a SDS-PAGE mini-gel, transferred onto a nitrocellulose membrane, probed with the indicated primary antibodies, and then incubated with secondary goat anti-rabbit or anti-mouse antibodies (Bio-Rad) as the substrate recognition subunits (Ni et al., 2014). In experiments involving PIFs, we used commercially available antibodies (Enzo Life Sciences) and polyclonal anti-phyB antibodies were used at 1:500 dilution. Polyclonal anti-RPN6 antibodies (Enzo Life Sciences) were used at 1:1000 dilution. Monoclonal anti-phyB antibodies were used at

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1:1000 dilution. Polyclonal anti-PIF1 and anti-PIF3 antibodies were used at 1:500 dilution.

RNA Extraction and qRT-PCR
Total RNA from seedlings of the indicated genotypes and growth conditions was isolated using the Spectrum Plant Total RNA kit (Sigma-Aldrich) with on-column DNase I (Sigma-Aldrich) treatment. cDNA was synthesized using an Invitrogen SuperScript II First-Strand cDNA synthesis kit according to the manufacturer’s recommendations. Oligo(dT) primers were used for the analysis of nuclear gene expression, and a mixture of oligo(dT) and gene-specific primers was used for the analysis of plastidial genes. qRT-PCR was performed with FastStart Universal SYBR Green Master Mix (Roche). All primers used are listed in Supplemental Tables 2 and 3.

Microarray Analysis
Col-0, hmr-5, and hmr-22 mutant seedlings were grown in continuous red light (10 μmol m⁻² s⁻¹) for 4 d before harvesting. Total RNA isolation was performed as described above. Three different biological replicates of each genotype were grown and separated, and extracted and processed independently. Total RNA was assessed for quality with a NanoDrop 8000 spectrophotometer (Thermo Scientific). cDNA was synthesized using an Invitrogen Superscript II First-Strand cDNA synthesis kit. Synthesis conditions was isolated using the Spectrum Plant Total RNA kit (Sigma-Aldrich) with on-column DNase I (Sigma-Aldrich) treatment. cDNA was incubated with glutathione Sepharose beads (GE Healthcare) equilibrated in E buffer at 4°C for 2 h. The beads with the immobilized GST fusion proteins were washed four times with E buffer supplemented with 0.1% Nonidet P-40. Prey proteins with a single N-terminal HA tag were synthesized using plasmid pCMX-PL2 and the TNT T7 Coupled Reticulocyte Lysate System (Promega) according to the manufacturer’s protocol. The in vitro-translated prey proteins were diluted with E buffer supplemented with 0.1% Nonidet P-40 and incubated with the affinity-purified GST fusion protein immobilized on the beads at 4°C for 2 h. Then, the beads were washed four times with E buffer supplemented with 0.1% Nonidet P-40. Bound proteins were eluted by boiling in 1× Laemmli protein sample buffer and subjected to 8% SDS-PAGE. Input and immunoprecipitated prey proteins were detected by immunoblots using goat anti-HA polyclonal antibodies (GenScript). The amount of GST fusion proteins used in each pull-down assay was visualized by staining the SDS-PAGE with Coomassie Brilliant Blue. All primers used for making the constructs used in the GST pull-down experiments are listed in Supplemental Table 4.

GST Pull-Down
GST pull-down assays were performed as described previously (Chen et al., 2010b; Galvão et al., 2012). Briefly, GST fusion proteins were expressed in *Escherichia coli* strain BL21(DE3) host strains (Agilent Technologies) carrying pET28b vectors (Novagen). After harvesting by centrifugation at 5000g for 10 min at 4°C, the cells were lysed by French press in E buffer containing 50 mM Tris-HCl, pH 7.5, 100 mM NaCl, 1 mM EDTA, 1 mM EGTA, 1% DMSO, 2 mM DTT, and protease inhibitor cocktail (Sigma-Aldrich). The cell extract was prepared by centrifugation at 10,000g for 20 min at 4°C, followed by ultracentrifugation at 60,000g for 15 min at 4°C. The cleared cell extract was incubated with glutathione Sepharose beads (GE Healthcare) equilibrated in E buffer at 4°C for 2 h. The beads with the immobilized GST fusion proteins were washed four times with E buffer supplemented with 0.1% Nonidet P-40.

Prey proteins with a single N-terminal HA tag were synthesized using plasmid pCMX-PL2 and the TNT T7 Coupled Reticulocyte Lysate System (Promega) according to the manufacturer’s protocol. The in vitro-translated prey proteins were diluted with E buffer supplemented with 0.1% Nonidet P-40 and incubated with the affinity-purified GST fusion protein immobilized on the beads at 4°C for 2 h. Then, the beads were washed four times with E buffer supplemented with 0.1% Nonidet P-40. Bound proteins were eluted by boiling in 1× Laemmli protein sample buffer and subjected to 8% SDS-PAGE. Input and immunoprecipitated prey proteins were detected by immunoblots using goat anti-HA polyclonal antibodies (GenScript). The amount of GST fusion proteins used in each pull-down assay was visualized by staining the SDS-PAGE with Coomassie Brilliant Blue. All primers used for making the constructs used in the GST pull-down experiments are listed in Supplemental Table 4.

Yeast Transactivation Assay
Yeast cultures were incubated at 20°C, and pictures were taken on the third day. For liquid culture assay, healthy colonies were selected and mated with yeast strain Y187 containing pGADT7 vector (Clontech). The mated diploids were cultured in SD/-Leu/-Trp media, and the transactivation assay was performed using o-nitrophenyl-β-D-galactoside as a substrate according to the Yeast Protocols Handbook (Clontech). The activity of β-D-galactosidase was calculated in Miller units from three replicates. Primers used to make the constructs for the yeast transactivation assays are listed in Supplemental Table 5. The wild-type and mutant versions of *HMR*-bAtAD were cloned into pGBKTK7 vector and subsequently transformed into Y2HGold yeast (Clontech). Overnight yeast cultures were first diluted to an OD₆₀₀ of 0.2, from which 5-fold serial dilutions were spotted on SD/-Trp and SD/-Trp/+Aureobasidin A. The plates were incubated at 30°C, and pictures were taken on the third day. Yeast transactivation activity in yeast.

Accession Numbers
Sequence data from this article can be found in the Arabidopsis Genome Initiative or GenBank/EMBL databases under the accession numbers listed in Supplemental Tables 2 and 3.

Supplemental Data
Supplemental Figure 1. Residues in the APB and APA essential for phyB and phyA interaction are not required for the interaction with HMR.

Supplemental Figure 2. The C-terminal half of HMR possesses transactivation activity in yeast.

Supplemental Figure 3. The hmr allele collection.

Supplemental Figure 4. *hmr-22* has a normal hypocotyl growth response in blue light.

Supplemental Figure 5. *hmr-22* is a weak, viable allele of *hmr*.
Supplemental Figure 6. A similar set of genes was misregulated in hmr-22 and hmr-5.

Supplemental Figure 7. The expression of the Class A HMR-PIF-dependent genes is upregulated in hmr-22.

Supplemental Figure 8. A D516N mutation in HMR22 affects neither the affinity of HMR to PIF1 in vitro nor HMR’s accumulation in vivo.

Supplemental Table 1. CAPS markers for genotyping hmr-14 to -22 mutants.

Supplemental Table 2. qRT-PCR primers for the nuclear-encoded genes examined in this study.

Supplemental Table 3. Primers used to measure the transcript levels of plastidial-encoded genes.

Supplemental Table 4. Primers used for making the bait and prey constructs for the GST pull-down assays.

Supplemental Table 5. Primers used to amplify HMR fragments for the constructs used in the yeast transactivation assays.

Supplemental Table 6. Primers used to introduce single amino acid substitutions in HMR’s 9aaTAD.

Supplemental Data Set 1. HMR-dependent genes in continuous R light.

Supplemental Data Set 2. HMR-PIF-dependent genes.

Supplemental Data Set 3. A list of 301 PIF direct-target genes represented in the Affymetrix ATH1 array.

Supplemental Data Set 4. HMR-dependent PIF direct-target genes.

Supplemental Data Set 5. Genes changed statistically significantly and by twofold in hmr-22.

Supplemental Data Set 6. PIF direct-target genes changed in hmr-22.

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AUTHOR CONTRIBUTIONS


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