Mechanism of Dual Targeting of the Phytochrome Signaling Component HEMERA/pTAC12 to Plastids and the Nucleus[OPEN]

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HEMERA (HMR) is a nuclear and plastidial dual-targeted protein. While it functions in the nucleus as a transcriptional coactivator in phytochrome signaling to regulate a distinct set of light-responsive, growth-relevant genes, in plastids it is known as pTAC12, which associates with the plastid-encoded RNA polymerase, and is essential for inducing the plastomic transcriptome in response to R light (Tepperman et al., 2010), including the red (R) and far-red light-sensing phytochromes (phys) that are essential for establishing photomorphogenesis (Franklin and Quail, 2010). Phys are bilin-containing proteins that can be photoconverted between two relatively stable forms: an R light-absorbing inactive Pr form and a far-red light-absorbing active Pfr form (Rockwell et al., 2006; Nagatani, 2010; Burgie and Vierstra, 2014). Phys mediate almost all the reprogramming of the nuclear transcriptome in response to R light (Tepperman et al., 2006; Leivar et al., 2009; Hu et al., 2013). One of the first light responses at the cellular level is the translocation of photoactivated phy from the cytoplasm to the nucleus, where phy are further compartmentalized to subnuclear photosensory domains named photobodies (Sakamoto and Nagatani, 1996; Kircher et al., 1999; Yamaguchi et al., 1999; Chen et al., 2003, 2005, 2010; Van Buskirk et al., 2012, 2014). A central mechanism by which phy control gene expression, possibly at photobodies, is by regulating the stability and activity of a massive transcriptional reprogramming in both the nuclear and plastidial genomes and sophisticated communications between these two genome-bearing subcellular compartments.

Deetiolation is a pivotal developmental transition when a seedling emerges from the ground and establishes a light-dependent photoautotrophic lifestyle or photomorphogenesis. In Arabidopsis (Arabidopsis thaliana), photomorphogenesis entails restriction of hypocotyl growth and promotion of leaf development and chloroplast biogenesis. These diverse morphological changes reflect massive transcriptional reprogramming in both the nuclear and plastidial genomes and sophisticated communications between these two genome-bearing subcellular compartments.

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group of nodal basic helix-loop-helix transcriptional regulators, the Phytochrome-Interacting Factors (PIFs; Shen et al., 2005; Al-Sady et al., 2006; Oh et al., 2006; Lorrain et al., 2008; Leivar and Quail, 2011; Park et al., 2012; Kaiserli et al., 2015; Qiu et al., 2015). Eight PIFs have been reported, including PIF1, PIF3-8, and PIL1 (PIF3-Like1), which perform overlapping and distinct antagonistic roles in photomorphogenesis (Ni et al., 1998; Huq and Quail, 2002; Huq et al., 2004; Khanna et al., 2004; Leivar et al., 2008a; Li et al., 2012; Luo et al., 2014). PIF1, PIF3, PIF4, PIF5, and PIF7 promote hypocotyl growth by activating growth-relevant genes, including genes in the biosynthesis and signaling of the plant growth hormone auxin (Huq and Quail, 2002; Huq et al., 2004; Al-Sady et al., 2008; Leivar et al., 2008a, 2008b; Lorrain et al., 2009; Shin et al., 2009; Li et al., 2012; Pfeiffer et al., 2014; Procko et al., 2016). PIF1, PIF3, and PIF5 inhibit chloroplast development by repressing nuclear-encoded photosynthetic genes (Huq et al., 2004; Moon et al., 2008; Shin et al., 2009; Stephenson et al., 2009; Kim et al., 2011). Most PIFs accumulate to high levels in dark-grown seedlings, and photoactivated phy mediate photomorphogenetic reprogramming by interacting directly with PIFs and repressing their stability and activity in the light (Shen et al., 2005; Al-Sady et al., 2006; Oh et al., 2006; Lorrain et al., 2008; Leivar and Quail, 2011; Park et al., 2012; Qiu et al., 2015).

Light also regulates the activity of plastid-encoded genes and, in turn, gene expression in plastids can influence nuclear genes through plastid-to-nucleus or retrograde signaling (de Souza et al., 2016; Woodson and Chory, 2008). The plastome in land plants carries over 120 genes encoding key components of the plastidial transcriptional, translational, and photosynthetic apparatuses (Sugiura, 1992; Wakasugi et al., 2001). Plastomic genes are transcribed by two types of plastid RNA polymerases: a phage-type nuclear-encoded plastid RNA polymerase (NEP) and a eubacterial-type plastid-encoded plastid RNA polymerase (PEP; Lerbs-Mache, 1993; Allison et al., 1996; Hricová et al., 2006; Liere et al., 2011). Although most plastid genes are transcribed by both the NEP and PEP (Legen et al., 2002; Demarsy et al., 2006, 2012), housekeeping genes are preferentially transcribed by the NEP, whereas photosynthetic genes are transcribed mainly by the PEP (Allison et al., 1996; Hajdukiewicz et al., 1997; Hübschmann and Börner, 1998). During deetiolation, phy signaling in the nucleus predominantly induces the plastid-encoded photosynthetic genes by the PEP (DuBell and Mullet, 1995; Chun et al., 2001; Thum et al., 2001). The PEP is comprised of eubacteria-like $\alpha$, $\beta$, $\beta'$, and $\beta''$ core subunits, a sigma factor, and a battery of plant-specific PEP-associated proteins (Pfalz and Pfannschmidt, 2013). Whereas the $\alpha$, $\beta$, $\beta'$, and $\beta''$ core subunits are encoded by the plastome, the sigma factors and PEP-associated proteins are encoded by nuclear genes (Hu and Bogorad, 1990; Pfannschmidt et al., 2000; Suzuki et al., 2004; Pfalz et al., 2006; Steiner et al., 2011; Melonek et al., 2012). Recent proteomic studies of purified soluble PEP-complex and membrane-bound transcriptionally active chromosome (TAC) from Arabidopsis and mustard (Sinapis alba) identified 12 PEP-associated proteins named PAPs and/or pTACs, which are essential for the activity of the PEP (Pfalz et al., 2006; Steiner et al., 2011; Pfalz and Pfannschmidt, 2013). The activity of the PEP can also influence phy-regulated nuclear gene expression through retrograde signaling (Woodson et al., 2013). Defects in chloroplast biogenesis, including deficiency in PEP activity, are able to repress the expression of the nuclear-encoded photosynthetic genes (Koussevitzky et al., 2007; Woodson et al., 2013; Martin et al., 2016). Despite the essential roles of PAPs/pTACs for the function of the PEP, their precise functions in PEP activity and roles in nuclear-plastidial communication are still not well understood.

Recent studies have uncovered that pTAC12, an essential PEP-associated protein (Pfalz et al., 2006; Steiner et al., 2011), is a key phy signaling molecule named HEMERA (HMR; Chen et al., 2010). Although the biochemical function of pTAC12/HMR in the PEP is still unclear, HMR in maize (Zea mays) was shown to bind single-stranded DNA and RNA and plays a role in the accumulation of the PEP (Pfalz et al., 2015). In addition, pTAC12/HMR interacts directly with two other PEP-associated proteins, pTAC14 and pTAC7 (Gao et al., 2011; Yu et al., 2013). Interestingly, both Arabidopsis and maize pTAC12/HMR are dual localized to the nucleus and plastids (Chen et al., 2010; Pfalz et al., 2015). The nuclear function of pTAC12/HMR was revealed through a forward genetic screen for mutants defective in the subnuclear localization of Arabidopsis phytochrome B (phyB) to photobodies (Chen et al., 2010). The hmr mutant represents the founding member of a new class of photomorphogenetic mutants that show a combination of long hypocotyl and albino phenotypes (Chen et al., 2010; Chen and Chory, 2011). HMR transduces light signals by interacting directly with photoactivated phy and all PIFs, and it acts as a transcriptional coactivator that regulates the degradation and activity of PIF1 and PIF3 (Galvão et al., 2012; Qiu et al., 2015). Knocking out PIF1, PIF3, PIF4, and PIF5 rescues hmr's long hypocotyl phenotype but not its albino phenotype, indicating that HMR plays two separable roles: a PIF-dependent role in regulating hypocotyl elongation and a PIF-independent role in chloroplast biogenesis (Chen et al., 2010; Qiu et al., 2015). These two roles likely reflect the functions of HMR in the nucleus and plastids, respectively.

The discovery of HMR raises a new hypothesis that nuclear-plastidial communication could be coordinated by nuclear/plastidial dual-targeted proteins such as HMR. As an initial step to test this hypothesis, we characterized the subcellular localization signals of HMR and investigated the mechanisms that target HMR to the nucleus and plastids. We show that HMR can be directly imported into chloroplasts through a transit peptide, which is cleaved upon chloroplast import. However, the predicted nuclear localization signals of HMR are unexpectedly nonfunctional. We
determined the mature form of HMR and showed that mature HMR retains the biochemical functions required for phytochrome signaling. Surprisingly, characterization of subcellular localization and function of mature HMR revealed an unexpected relationship between HMR’s plastidial localization and nuclear accumulation.

RESULTS
HMR Can Be Imported Directly into Chloroplasts by a Transit Peptide Residing in the N-Terminal 50 Amino Acids

HMR isolated from the nucleus and chloroplasts show the same protein size (Fig. 1A; Chen et al., 2010; Pfalz et al., 2015). It was speculated that this mature form of HMR is smaller than the full-length HMR due to the loss of its transit peptide upon chloroplast import. However, it has not been demonstrated experimentally that HMR can be directly imported into chloroplasts. In fact, the ChloroP algorithm predicts that Arabidopsis HMR lacks a chloroplast-targeting sequence (Emanuelsson et al., 1999; Chen et al., 2010). Thus, we sought to determine whether HMR has an N-terminal transit peptide and, if so, whether this signal could be cleaved after chloroplast import. To address these questions, we compared the protein size of HA-tagged HMR (HMR-HA) transiently expressed in tobacco (Nicotiana benthamiana) with that of the full-length HMR-HA made by in vitro translation. The tobacco-expressed HMR-HA was clearly smaller than the full-length form, indicating that the mature HMR in vivo is a cleaved version (Fig. 1B). We then performed chloroplast import assays to confirm that HMR can be directly imported into chloroplasts. These experiments show that in vitro-translated HMR fused with FLAG and His tags (HMR-FLAG-His) could be...
imported into isolated Arabidopsis chloroplasts and resulted in smaller mature HMR-FLAG-His (Fig. 1C). To define HMR’s transit peptide, we fused various N-terminal fragments of HMR, ranging from 20 to 60 amino acids (N20-N60), to YFP (Fig. 1D) and generated transgenic lines in the Col-0 background expressing these recombinant YFP proteins. The ability of these HMR peptides in targeting YFP to the chloroplasts in the transgenic lines was examined by confocal microscopy. While N20-YFP and N31-YFP were localized in the nucleus and cytoplasm and failed to localize to the chloroplasts, both N50-YFP and N60-YFP were localized in the chloroplasts (Fig. 1E). In addition, N20-YFP and N31-YFP from the transgenic lines were the same sizes as their corresponding in vitro translated forms, whereas N50-YFP and N60-YFP from the transgenic lines were smaller than their respective in vitro translated forms (Fig. 1F). These results indicate that the N-terminal 50 amino acids of HMR are sufficient for chloroplast import. However, surprisingly, although N60-YFP is larger than N50-YFP, the mature forms of N50-YFP and N60-YFP had similar molecular mass (Fig. 1F), suggesting that N60-YFP was further proteolytically processed after the removal of the transit peptide. Together, these results demonstrate that Arabidopsis HMR can be directly targeted into the chloroplasts by a transit peptide, which resides in its N-terminal 50 amino acids. After cleavage of the transit peptide, the N terminus of HMR may be further proteolytically processed to yield mature HMR.

Mature HMR Lacks the N-Terminal 57 Amino Acids

We then determined the mature form of HMR by mass spectrometry. We isolated HMR-HA from 4-d-old, red light-grown HMR-HA seedlings (Galvão et al., 2012; Qiu et al., 2015) by affinity purification with anti-HA antibody-conjugated agarose beads. HMR-HA proteins were separated by SDS-PAGE and stained with SimplyBlue SafeStain. The HMR band was excised from the gel and subjected to in-gel digestion with endoproteinase GluC, which cleaves peptide bonds C terminal to either Glu or Asp (Drapeau et al., 1972). The peptides digested from the excised band were analyzed by liquid chromatography-tandem mass spectrometry (LC-MS/MS). The LC-MS/MS analysis identified 94 unique HMR peptides (Supplemental Table S1), which spanned 76% of the full-length HMR sequence (Fig. 2A). The most N-terminal peptide identified is between amino acids 58 and 78, KKEGDGDVSEGKKSEGEEGF (Fig. 2, A and B). Given the number of Glu and Asp residues within its sequence, this peptide likely resulted from incomplete digestion. Because the N-terminal residue preceding the peptide is a Cys, which cannot be cleaved by GluC, the cleavage between Cys-57 and Lys-58 was most likely catalyzed by an endogenous peptidase, suggesting that the mature form of HMR (HMRm) begins from Lys-58.

HMRm Retains Binding Activities with Phytochromes and PIFs

We showed previously that HMR acts as a phy signaling component in the nucleus via direct interactions with phys and PIFs. HMR interacts preferentially with the Pfr form of phyA and phyB (Galvão et al., 2012), and it binds to all seven PIFs and PIL1 (Qiu et al., 2015). Both the HMR-phy and HMR-PIF interactions are mediated by the N-terminal half of HMR, in particular the phyA-interacting region 1 (PIR1) within the N-terminal 115 amino acids. Therefore, it raised the question of whether HMRm retains the ability to interact directly with phys and PIFs. To address this question, we...

Figure 2. Identification of HMRm by LC-MS/MS. A, LC-MS/MS analysis of HMR-HA. HMR-HA was immunoprecipitated from 4-d-old HMR-HA seedlings, digested with endoproteinase GluC, and analyzed by LC-MS/MS. Residues identified by MS analysis are highlighted in yellow. The red box indicates the N-terminal-most identified peptide. B, Annotated MS/MS spectrum (mascot ion score 26.7) of the peptide KKEGDGDVSEGKKSEGEEGF on Q Exactive high-resolution mass spectrometer illustrating product ion coverage throughout sequence.
performed GST pull-down assays using *Escherichia coli* produced GST-HMRm as the bait to pull down in vitro translated HA-tagged phyA, phyB, PIFs, and PIL1. As shown in Figure 3A, GST-HMRm was able to pull down the apoprotein, Pr, and Pfr forms of phyA and phyB, and its interactions with the Pfr forms of phyA and phyB were stronger than those with the Pr forms. GST-HMRm could also pull down all PIFs and PIL1 but failed to pull-down SPATULA and ALCATRAZ, which lack PIFs’ conserved APB motif required for HMR interaction (Fig. 3B; Khanna et al., 2004; Qiu et al., 2015). These results indicate that HMRm retained the previously characterized biochemical properties in phytochrome signaling.

**HMR’s Predicted Nuclear Localization Signals Are Nonfunctional**

The observation that nuclear and plastidial HMR share the same *M* raises the possibility that HMR is targeted to the plastids first before translocating to the nucleus. If this hypothesis is correct, one might expect that deleting the transit peptide would block not only HMR’s chloroplast localization but also its nuclear accumulation. However, HMR does contain two predicted bipartite nuclear localization signals, NLS1 and NLS2 (Fig. 4A). We therefore tested whether these two nuclear localization signals are functional. To make a combined *M* greater than the nuclear pore’s passive diffusion limit of 60 kD (Wang and Brattain, 2007), we fused the two nuclear localization signals, either individually or combined, to YFP and GUS (YG; Fig. 4A). These recombinant proteins were transiently expressed in tobacco and their localization patterns were characterized by confocal microscopy. The results showed that neither NLS1, NLS2, nor combined NLS1/NLS2 (2NLS) was able to effectively import YG into the nucleus (Fig. 4, B and C). Although a small fraction of cells showed nuclear YFP signal, the majority of YG-NLS1, YG-NLS2, and YG-2NLS were localized to the cytoplasm, similar to the YG control (Fig. 4, B and C). These results indicate that the two predicted nuclear localization signals of HMR are nonfunctional.

![Figure 3](https://example.com/figure3.png)

**Figure 3.** HMRm interacts with phyA, phyB, and all PIFs in vitro. A, GST pull-down assays showing that HMRm interacts with phyA and phyB in vitro. The pull-down assays were performed with recombinant GST-HMRm or GST produced in *E. coli* to pull down in vitro-translated HA-tagged phyA and phyB (phyA-HA and phyB-HA) that were either apoproteins (apo) or holoproteins in the Pfr or Pr form. Input and bound phy fractions were detected by immunoblot using anti-phyA or -phyB antibodies (top). Immobilized GST-HMRm and GST are shown in the Coomassie blue-stained SDS-PAGE gels at the bottom. The relative amounts of the bound phy proteins are normalized to the amount of the corresponding GST-HMRm bait and are shown below the top. B, GST pull-down assays showing that HMRm interacts with all PIFs in vitro. *E. coli*-expressed recombinant GST-HMRm or GST were used to pull down in vitro-translated HA-tagged PIF1, PIF3−8, PIL1, SPATULA, and ALCATRAZ. Input and bound fractions of the prey proteins were detected by immunoblot using anti-HA antibodies (top). Immobilized GST-HMRm and GST are shown in the Coomassie blue-stained SDS-PAGE gels at the bottom. I, 10% input of the prey proteins; G, GST; H, GST-HMRm.
HMRm Could Rescue Neither the Albino nor Long Hypocotyl Phenotypes of hmr

To examine the function of HMRm in vivo, we generated transgenic lines in the hmr-5 background expressing HA-tagged HMRm (HMRm-HA) under the 35S promoter, the same as the previously described HMR-HA line (Galvão et al., 2012). We analyzed two independent HMRm-HA transgenic lines. As expected, without the transit peptide, HMRm-HA could not rescue hmr-5’s albino phenotype (Fig. 5A) or its defects in the expression of PEP-dependent genes (Fig. 5B). However, HMRm also failed to rescue hmr-5’s long hypocotyl phenotype (Fig. 5, A and C), which presumably depends on the nuclear function of HMR (Qiu et al., 2015). These results suggest that HMRm alone is able to rescue neither the nuclear nor plastidial functions of HMR.

HMRm Failed to Rescue the Defects of HMR-Regulated Nuclear Genes in hmr

We showed previously that nuclear HMR is required for the regulation of a subset of growth-promoting PIF-induced direct target-genes: HMR-repressed Class A genes and HMR-induced Class B genes (Qiu et al., 2015). To demonstrate that HMRm-HA does not show nuclear function, we asked whether the HMRm-HA lines could rescue expression of the HMR-dependent nuclear genes. However, because the expression of light responsive genes in the nucleus is also controlled by retrograde signaling from plastids, deficiencies in chloroplast biogenesis could dramatically alter nuclear gene expression (Woodson and Chory, 2008). Therefore, one major challenge in characterizing the nuclear function of HMRm-HA in the albino HMRm-HA/hmr-5 lines was to distinguish whether alterations in nuclear gene expression were due to nuclear HMRm-HA or the

Figure 4. HMR’s predicted nuclear localization signals are nonfunctional. A, Schematic illustration of recombinant proteins of YG fused with either one or both of HMR’s predicted NLSs. YG with the SV40 NLS (PKKKRKV), YG-NLS, was used as a control for nuclear targeting. B, Confoical images of representative tobacco cells expressing the indicated recombinant proteins. Tissue was collected 3 d after agroinfiltration, stained with DAPI, and imaged by confocal microscopy. White arrowheads indicate nuclei. Scale bars = 50 μm. C, Quantitation of subcellular localization data in B. The percentage of cells with YFP signal in the nucleus or cytoplasm for each of the recombinant proteins was determined by examining at least 60 DAPI- and YFP-positive cells. Error bars represent the sd of three independent experiments.

Figure 5. HMRm requires a transit peptide to rescue both the tall and albino phenotypes of the hmr mutant. A, Images of 4-d-old seedlings grown under 10 μmol m⁻² s⁻¹ red light from Col-0, hmr-5, HMRm-HA/hmr-5, and rbcStp-HMRm-HA/hmr-5 lines. B, qRT-PCR data showing the transcript levels of representative PEP-dependent genes relative to PP2A in 4-d-old red light grown Col-0, hmr-5, HMRm-HA/hmr-5, and rbcStp-HMRm-HA/hmr-5 lines. Error bars represent the se of three replicates. C, Hypocotyl length measurements of seedlings in A. Error bars represent the sd of at least 20 seedlings.
concomitant defect in plastid biogenesis. To this end, we first tested whether the expression levels of the HMR-regulated genes were influenced by defects in plastid biogenesis through treatment with lincomycin, a potent inhibitor of plastid translation (Nott et al., 2006; Koussevitzky et al., 2007). Lincomycin inhibits the 70s ribosome, leading to the loss of the plastid-encoded core subunits of the PEP RNA polymerase. The resulting defects in expression of plastid-encoded photosynthetic genes subsequently lead to repression of nuclear-encoded photosynthesis-associated nuclear genes through retrograde signaling (Koussevitzky et al., 2007; Woodson et al., 2013). Interestingly, although the Class A and B genes are misregulated in the albino hmr-5 mutant, lincomycin treatment of Col-0 did not have a significant effect or the same effect on these HMR-regulated PIF-induced genes (Fig. 6). Therefore, these two groups of HMR-regulated PIF-induced genes can be used as markers for assessing the nuclear function of HMR. These results are consistent with a recent study by Martín et al., who showed that lincomycin, through retrograde signaling, controls PIF-repressed but not PIF-induced genes (Martín et al., 2016). We then examined the expression of the Class A and B genes in the HMRm-HA/ hmr-5 lines. Compared with hmr-5, HMRm-HA/hmr-5 showed similar expression levels in three of the four Class A genes, At5g14180, At2g45210, and At1g19530, and in three of the four Class B genes, At2g46870, At4g16780, and At4g14130 (Fig. 6, A and B). These results support the conclusion that HMRm fails to rescue the nuclear defects of hmr-5.

HMRm Failed to Accumulate in the Nucleus

We then compared the relative amount of the nuclear fraction of HMRm-HA with that of HMR-HA. Strikingly, in the two HMRm-HA lines that we examined, the nuclear fractions of HMRm were more than 5-fold less...
than those of the HMR-HA lines (Fig. 7A). Consistent with the notion that HMR’s predicted nuclear localization signals are nonfunctional, these results demonstrate that HMRm-HA is unable to efficiently accumulate in the nucleus.

We had previously shown that when an YFP is fused to the N terminus of the full-length HMR to block the function of its transit peptide, YFP-HMR was localized to the nucleus and cytoplasm, and it fails to rescue the hmr mutant (Chen et al., 2010). To test whether the defect of YFP-HMR might also be due to the failure in nuclear accumulation, we reexamined YFP-HMR localization in the YFP-HMR lines. These results show that despite observable nuclear localization in 25% of the nuclei, YFP-HMR was mainly localized to the cytoplasm (Supplemental Fig. S1, A and B). As shown previously, YFP-HMR did not rescue the long hypocotyl phenotype of hmr-5 (Supplemental Fig. S1C); also, it did not rescue the defect in the expression of the HMR- and PIF-induced Class B genes (Supplemental Fig. S1D). These results support the notion that HMR does not contain a functional NLS and suggest that the reason why YFP-HMR did not rescue the nuclear defect of the hmr mutant is likely due to the lack of nuclear accumulation.

Fusing the Transit Peptide of rbcS to HMRm Rescues Its Defects in Nuclear Accumulation and Function

To further test the hypothesis that HMR’s nuclear accumulation is dependent on its plastidial localization, we asked whether fusing a heterologous transit peptide to HMRm would rescue its nuclear accumulation and function. To this end, we fused the transit peptide of the small subunit of Rubisco, rbcStp (Roesler et al., 1997), to HMRm-HA and generated transgenic lines in the hmr-5 background. We picked two independent rbcStp-HMRm-HA/hmr-5 lines for further characterization. Both transgenic lines rescued hmr-5’s albino phenotype (Fig. 5A) as well as its defects in the expression of PEP-dependent genes (Fig. 5B), indicating that HMRm is fully functional in plastids. Interestingly, rbcStp-HMRm-HA also rescued the long hypocotyl phenotype of hmr-5 (Fig. 5C). In addition, the expression of the Class A genes, AT5G14180, AT2G45210, and AT1G19530, as well as all of the Class B genes were rescued in the rbcStp-HMRm-HA lines (Fig. 6).

Therefore, rbcStp-HMRm-HA was able to rescue both the plastidial and nuclear defects of hmr-5. In line with these observations, the levels of nuclear HMR-HA in the rbcStp-HMRm-HA lines were greatly enhanced compared with those in the HMRm-HA/hmr-5 direct connections between chloroplasts and the nucleus, such as stromules. In the first two scenarios, HMRm could be posttranslationally modified (represented as the red circle) to facilitate its nuclear localization.

Figure 7. HMR’s nuclear accumulation requires its localization to chloroplasts. A, Semiquantitative measurements of the nuclear fractions of HMR-HA, HMRm-HA, and rbcStp-HMRm-HA from transgenic lines. Total and nuclear HMR-HA protein fractions were isolated and analyzed with western blots using anti-HA antibodies. Pol II and SiR were used as nuclear and plastidial controls, respectively. Band intensities from the HA blots were normalized to their respective Pol II controls, and the nuclear to total ratio (N/T) was calculated for each line and is listed below the HA western blots. B, Immunoblots showing the endogenous HMR-HA proteins are the same size as or similar to the size of their corresponding in vitro translated mature forms. Left, The endogenous total and nuclear HMR-HA proteins are the same size as the in vitro translated (IVT) HMRm-HA and are clearly smaller than the in vitro translated full-length HMR-HA. Right, The endogenous rbcStp-HMRm-HA proteins are similar in size to the in vitro translated HMRm-HA and are smaller than the in vitro translated full-length rbcStp-HMRm-HA. Red arrowheads indicate the bands of full-length in vitro translated HMR proteins. C, Model for the mechanism of nuclear and plastidial targeting of HMR. Upon translation in the cytoplasm, the HMR preprotein (pre-HMR) is imported into the chloroplasts by its transit peptide (TP, labeled in green). The TP and additional N-terminal residues are cleaved in the chloroplast to yield HMRm, which participates in the PEP-mediated transcriptional activation of plastid-encoded photosynthetic genes. HMRm can be then translocated to the nucleus by at least three possible ways: (1) with an unknown retrograde translocon, (2) due to leakage from damaged chloroplasts, and (3) through posttranslational modifications.
lines and were comparable to those in the HMR-HA/hmr-5 lines (Fig. 7A). Also, similar to HMR-HA, the size of endogenous rbcStp-HMRm-HA was similar to, if not the same as, that of HMRm-HA (Fig. 7B), suggesting that the rbcS transit peptide was removed from rbcStp-HMRm-HA and that it was the mature form of rbcStp-HMRm-HA that had accumulated in the nuclei of the rbcStp-HMRm-HA/hmr-5 lines. Together, these results support the hypothesis that HMR’s nuclear accumulation depends on localization to the plastids.

DISCUSSION

HMR is the first identified phy signaling component dual targeted to the nucleus and plastids (Chen et al., 2010), which draws a direct link between the activity of the PEP RNA polymerase in the plastids and phy signaling in the nucleus (Pfalz et al., 2006, 2015; Chen et al., 2010; Galvão et al., 2012; Qiu et al., 2015). Molecular understanding of the mechanisms targeting HMR to the nucleus and plastids holds promise for gaining novel insight into the mechanisms of nuclear-plastidial communication (Krause and Krupinska, 2009; Krause et al., 2012). In this study, we have shown that HMR can be directly imported into chloroplasts, where it is processed to the mature form starting from Lys-58 (Figs. 1 and 2). HMRm retains the abilities to interact with phy and PIFs in phy signaling (Fig. 3; Galvão et al., 2012; Qiu et al., 2015). Unexpectedly, the predicted nuclear localization signals in HMR are non-functional (Fig. 4). We showed that HMR’s nuclear accumulation and function rely on its transit peptide (Figs. 5–7). Fusing the transit peptide of rbcS to HMRm rescues its defects in nuclear accumulation and function (Figs. 5–7). Together, these results, combined with the observation that nuclear HMR has the same molecular mass as plastidial HMR (Fig. 1; Chen et al., 2010; Pfalz et al., 2015), provide biochemical, cellular, and genetic evidence supporting a plastid-to-nucleus retrograde protein translocation mechanism, in which nuclear accumulation of HMR depends on localization to the plastids (Fig. 7C).

HMRm

HMR/pTAC12 was shown to be dual targeted to the nucleus and plastids in Arabidopsis and maize (Chen et al., 2010; Pfalz et al., 2015). It is intriguing that the nuclear and plastidial HMR proteins showed the same size (Fig. 1; Chen et al., 2010; Pfalz et al., 2015), which raised the possibility that HMRm is derived from the removal of its plastidial targeting transit peptide. However, it had not been experimentally shown that HMR could be directly imported into plastids, and HMRm had not been determined. In this study, we show that HMR can be directly imported into chloroplasts (Fig. 1C). Chloroplast localization leads to reduction in the size of HMR as well as recombinant proteins with N-terminal fragments of HMR in both tobacco and Arabidopsis (Fig. 1), demonstrating that full-length pre-HMR is proteolytically processed upon chloroplast localization. We determined that mature HMR begins from Lys-58. Several lines of evidence support this conclusion. First, when purified HMR-HA was subjected to Glu-C digestion and LC-MS/MS analysis, the most N-terminal fragment identified begins at Lys-58 (Fig. 2). Because Glu-C is not expected to cleave between Cys-57 and Lys-58, this peptide likely defines the N terminus of HMRm. Second, although the N-terminal 50 amino acids of HMR are sufficient to mediate chloroplast import, when the N-terminal 60 amino acids of HMR were fused to YFP, the mature protein was further processed to a residue between amino acids 50 and 60 (Fig. 1), which is consistent with the conclusion that HMRm begins at residue 58. Third, HMRm-HA from the HMR-HA/hmr-5 lines showed the same size as HMRm-HA from the HMRm-HA/hmr-5 lines (Fig. 7A). Fourth, HMRm retains the binding activities with phyA, phyB, PIFs, and PIF1 (Fig. 3). Lastly, fusing the transit peptide of rbcS to HMRm rescues both the nuclear and plastidial phenotypes of the hmr mutant, indicating that HMRm is functional in both the nucleus and plastids in vivo (Figs. 5–7). Based on these results, we conclude that HMRm in Arabidopsis begins from Lys-58.

Mechanism of Dual-Targeting HMR to Plastids and the Nucleus

It is well documented in yeast (Saccharomyces cerevisiae), animals, and plants that a single gene can produce proteins targeted to dual destinations (Karnieli and Pines, 2005; Carrie et al., 2009; Krause et al., 2012). Dual distribution between subcellular compartments can be achieved by two general strategies. One strategy is that a single gene produces two protein variants that differ in subcellular targeting signals due to alternative transcription or translation initiation sites. For example, the Arabidopsis GSTF8 (glutathione S-transferase F8) gene utilizes multiple transcription start sites that produce two distinct protein forms: a large protein with an N-terminal transit peptide that localizes to the plastids and a small protein without the transit peptide that localizes to the cytoplasm (Thatcher et al., 2007). The alternative strategy is that a single protein can be targeted to two subcellular compartments. The dual-localization of HMR is likely due to the latter strategy for several reasons. First, HMR has not been found to produce alternative transcripts from distinct transcription start sites. In addition, because both the nuclear and plastidial HMR show the same size (Fig. 1; Chen et al., 2010; Pfalz et al., 2015), the two forms are unlikely to be produced by alternative transcripts like GSTF8. Second, although HMRm is smaller than full-length HMR, this is unlikely the result of alternative translation start sites. The second Met in HMR is at position 91, so if this Met were used as an alternative translation start site, it would produce a protein much smaller than the one observed. Furthermore,
this second AUG is unlikely to be an initiator codon, because it does not contain the caA(A/C)aAUGGCg consensus sequence of plant translation initiator codons (Joshi et al., 1997). Taken together, the current evidence supports the idea that a single translational product of HMR is dual-targeted to plastids and the nucleus.

The best-recognized mechanism targeting a single protein to the nucleus and an organelle is that a single polypeptide harbors two targeting signals, a nuclear localization signal and an organelar targeting signal. In this case, the partitioning of the protein between these two compartments is determined by competition and accessibility of the two targeting signals (Karnieli and Pines, 2005; Krause and Krupinska, 2009; Krause et al., 2012). For instance, yeast Pir1p contains both a nuclear localization signal and a mitochondrial targeting signal, and its partitioning to these two subcellular compartments is regulated by masking and unmasking of the nuclear localization signal by its binding partner Apn1p (Vongsamphanh et al., 2001). It was hypothesized that the dual targeting of HMR was mediated by its predicted nuclear localization signals and transit peptide (Pfalz et al., 2006; Chen et al., 2010); however, the functionality of these targeting signals had never been experimentally validated. In this study, we demonstrate that HMR possesses a functional transit peptide within its N-terminal 50 amino acids, which is required and sufficient for delivering a YFP reporter into the chloroplasts in Arabidopsis (Fig. 1). Surprisingly, neither of these two predicted nuclear localization signals in HMR is highly functional (Fig. 4). Consistent with this notion, HMRm fails to accumulate and exert functions in the nucleus (Figs 5–7). Together, these results argue against the original hypothesis of dual-targeting signals. On the contrary, these results, combined with the fact that both nuclear and plastidial HMR show the same size (Fig. 1; Chen et al., 2010; Pfalz et al., 2015), support the alternative model that HMR is first localized to and proteolytically processed in the plastids before being relocated to the nucleus, a mechanism that requires plastid-to-nucleus retrograde protein translocation (Fig. 7C). This plastid-to-nucleus retrograde protein translocation model has been proposed for the subcellular localization of another nuclear and plastidial localized protein, Whirly1 (Why1; Grabowski et al., 2008). Why1 is a single-stranded DNA and RNA binding protein (Desveaux et al., 2000, 2002; Prikryl et al., 2008). In the nucleus, Why1 binds to single-stranded telomeric DNA to modulate telomere length homeostasis (Yoo et al., 2007) and acts as a transcriptional regulator for pathogen responsive genes (Desveaux et al., 2000; Xiong et al., 2009). Plastidial Why1 is required for chloroplast biogenesis in maize (Prikryl et al., 2008) and plays a possible role in plastidial genome repair (Cappadocia et al., 2010, 2012). Interestingly, a fraction of Why1 is also associated with the pTAC complex (Pfalz et al., 2006; Melonek et al., 2010). Compared with the full-length protein, nuclear Why1 exhibits the same reduced molecular mass as the plastidial form (Grabowski et al., 2008). More importantly, plastomically expressed Why1 in tobacco could translocate to the nucleus, providing solid evidence that Why1, at least in tobacco, could move from plastids to the nucleus (Isemer et al., 2012). Here, we showed that nuclear accumulation of HMR is dependent on its localization to the plastids (Figs. 5–7). Although the localization data of plastomically expressed HMR are still missing, the results shown in this study strongly support the idea that HMR’s nuclear localization in Arabidopsis is mediated by a retrograde protein translocation pathway (Fig. 7C).

The mechanism of retrograde protein translocation from plastids to the nucleus is still largely unknown. There are at least three possible pathways (Fig. 7C). First, HMR can be retrieved from the plastids by retrograde movement through an unidentified translocon or secretory pathway. Second, HMR can be redistributed as a result of leakage from chloroplasts, either due to organelar damage or during autophagy (Woodson et al., 2015). Because HMRm cannot effectively accumulate in the nucleus, in these two scenarios, HMRm might be modified in plastids to enhance the activity of its own nuclear localization signals or to promote interactions with other proteins that facilitate HMR’s nuclear accumulation (Fig. 7C). Third, HMR can be transported from plastids to the nucleus through physical connections between these two compartments such as stromules (Kwok and Hanson, 2004). With the increasing number of described nuclear and plastidial dual-targeted proteins, it has become essential to understand the mechanisms of dual targeting to the nucleus and plastids and the significance of these proteins in both antero- and retrograde signaling between the two gene-bearing compartments (Krause and Krupinska, 2009; Krause et al., 2012). This study represents one of the initial attempts toward these goals.

MATERIALS AND METHODS

Plant Growth Conditions and Hypocotyl Measurements

Seeds were surface sterilized in 50% bleach with 0.01% Triton X-100 for 10 min and then washed four times with sterile double distilled water. Seeds were then plated on half-strength Murashige and Skoog with Gamborg’s vitamins (MSP0506; Caisson Laboratories), 0.5 mM MES (pH 5.7), and 0.8% (w/v) agar (A038; Caisson Laboratories). For lincomycin treatments, plates were supplemented with 0.5 mM lincomycin hydrochloride (L2774; Sigma-Aldrich). Seeds were then plated on half-strength Murashige and Skoog supplemented with 0.5 mM MES (pH 5.7), and 0.8% (w/v) agar (A038; Caisson Laboratories). For lincomycin treatments, plates were supplemented with 0.5 mM lincomycin hydrochloride (L2774; Sigma-Aldrich). Seeds were stratified in the dark at 4°C for 4 d. For microscopy, hypocotyl measurements, nuclear fractionation, and RNA extraction, seedlings were grown at 21°C in an LED chamber (Percival Scientific), and light intensity was measured using an Apogee PS200 spectroradiometer (Apogee Instruments). For hypocotyl length measurements, an Epson Perfection V700 photo scanner was used to capture images of seedlings. At least 20 hypocotyls from each genotype and treatment were measured with NIH ImageJ software.

Plasmid Construction and Generation of Transgenic Lines

For producing HMR-HA by in vitro translation, HMR-HA was subcloned from pCHF3-HMR-HA (Qu et al., 2015) and ligated into the XmaIII and PstI sites of the pCMX-PL1 vector (Umesono et al., 1991). The constructs for identifying HMR’s transit peptide were generated from pCHF3-YFP, which was generated by ligating YFP flanked by Sall and XhoI sites into the pSall site after the 3SS promoter in pCHF3 (Fankhauser et al., 1999). The

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segments of HMR were amplified by PCR using the primers in Supplemental Table S2 and ligated into the BamHI and SalI sites of pCHF3-YFP. These constructs were then transfected into Col-0 plants with Agrobacterium tumefaciens strain GV3101 carrying the appropriate plasmids. Transformants were identified by screening on kanamycin plates.

Constructs producing recombinant HA-tagged prey proteins in vitro for GST pulldowns were previously described (Qiu et al., 2015). The bait construct encoding GST-tagged HMRn was cloned by amplifying HMRn by PCR with the primers in Supplemental Table S2 and ligating into the EcoRI and SalI sites of pET24b (Novagen).

For constructs testing the predicted NLS of HMR, YFP without a stop codon was ligated into the KpnI and BamHI sites of pCHF3 to give pCHF3-YFP (N). Then Gus, lacking a stop codon and flanked by BglII and BamHI sites, was amplified by PCR with primers in Supplemental Table S2 and ligated into the BamHI site of pCHF3-YFP/N to give pCHF3-YFP-GUS. The control constructs, YG-NLS2, YG-NLS1, and YG-NLS2 were generated by amending segments of HMR by PCR with the primers described in Supplemental Table S2 and then ligating them into the BamHI and SalI sites of pCHF3-YFP-GUS. HMRm-HA was made by amending HMRm-HA by PCR with the primers described in Supplemental Table S2 using pCHF3-HMR-HA as a template. The PCR product was then inserted into the SalI site after the 35S promoter in pCHF3 by Gibson assembly (NEB). Similarly, rcsHm-HMRm-HA was cloned by amending the rcsHm from cDNA, and HMRm-HA from pCHF3-HMR-HA, by PCR with primers listed in Supplemental Table S2. Both fragments were then inserted simultaneously into the SalI site of pCHF3 by Gibson assembly (NEB). Transgenic lines were generated by transforming hmr-5 heterozygous plants with A. tumefaciens strain GV3101 carrying either pCHF3-HMRm-HA or pCHF3-rcsHm-HMRm-HA plasmids, and transgenic seedlings were identified by screening on kanamycin plates.

Chloroplast Import Assays

Import assays were performed as previously described (Li and Schnell, 2006). Substrate proteins for import were produced and labeled with [35S] Met using 10 mM [35S] Met or [35S] Cys (PerkinElmer Life and Analytical Sciences) in the presence of 10 mM EDTA for 10 min. Then 20 to 30 g of chloroplasts were recovered by isolation over a 35% Percoll cushion and washed for 20 min at 26°C in import buffer (330 mM sorbitol, 50 mM HEPES-KOH, pH 7.5, 25 mM KOAc, and 5 mM MgOAc). Thermolysin treatment was then performed by diluting the import reactions with 3-fold excess ice-cold import buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM NaCl, 1 mM EDTA, 0.5% Triton X-100, and 1% EDTA-free protease inhibitor cocktail (Roche). The crude extract was centrifuged at 10,000g for 15 min, and the supernatant was then further cleared through a 0.45-mm filter. The clarified lysate was mixed with 50 mM of anti-HA conjugated agarose beads (ThermoFisher Scientific) and incubated at 4°C for 4 h. Washing and elution with 2× LDS loading buffer were done using an anti-HA spin column kit (ThermoFisher Scientific) according to the manufacturer’s directions.

The Duke Proteomics Core Facility received samples in LDS Loading buffer. Samples were reduced with 10 mM DTT at 70°C for 10 min prior to SDS-PAGE separation on a 4 to 12% Bis-Tris acrylamide gel (NuPAGE; ThermoFisher Scientific) with colloidal Coomassie blue staining. Bands were excised and subjected to standardized in-gel trypsin or protease V-8 (Gluc) digestion. Extracted peptides were lyophilized to dryness and resuspended in 20 mM of 0.2% formic acid/2% acetonitrile.

Each sample was subjected to chromatographic separation on a Waters NanoAcquity UPLC equipped with a 1.7-μm BEH130 C18 75-μm × 20-mm reversed-phase column. The mobile phase consisted of (A) 0.1% formic acid in water and (B) 0.1% formic acid in acetonitrile. Following a 2-min injection, peptides were trapped for 3 min on a 5-μm Symmetry PuriPro X2 130 A C18 precolumn (Waters) and then eluted with a linear gradient of 5% B to 100% B over 60 min at 400 nL/min. The analytical column was connected to a fused silica PicoTip emitter (New Objective) with a 10-μm tip orifice and coupled to a Thermo Q Exactive Plus mass spectrometer through an electrospray interface operating in a data-dependent mode of acquisition. The instrument was set to acquire a precursor MS scan from m/z 375 to 1,600 with MS/MS spectra acquired for the 10 most abundant precursor ions. For all experiments, charge-dependent CID energy settings were employed and a 60± dynamic exclusion was employed for previously fragmented precursor ions.

Raw LC-MS/MS data files were processed in Mascot distiller (Matrix Science) and then submitted to independent Mascot searches (Matrix Science) against a TAIR10 database containing both forward and reverse entries of each protein (32,790 forward entries). Search tolerances were 5 ppm for precursor ions and 0.02 D for product ions using semi-GluC specificity with up to seven missed cleavages. Carbamidomethylation (+57.0214 D on C) was set as a fixed modification, whereas oxidation (+15.9949 D on M) was considered a dynamic mass modification. All searched spectra were imported into Scaffold (v4.3; Proteome Software) and scoring thresholds were set to achieve a protein false discovery rate of 0.7% using the PeptideProphet algorithm.

Affinity Purification and MS

To purify HMR-HA for transit peptide identification, HMR-HA lines were grown in a continuous white light incubator (Percival Scientific) for 96 h. Five grams of seedlings were collected and ground in liquid nitrogen with a mortar and pestle. The powder was then resuspended in comminutionprecipitation buffer containing 50 mM Tris-HCl (pH 7.5), 100 mM NaCl, 1 mM EDTA, 0.5% Triton X-100, and 1% EDTA-free protease inhibitor cocktail (Roche). The crude extract was centrifuged at 10,000g for 15 min, and the supernatant was then further cleared through a 0.45-μm filter. The clarified lysate was mixed with 50 μL of anti-HA conjugated agarose beads (ThermoFisher Scientific) and incubated at 4°C for 4 h. Washing and elution with 2× LDS loading buffer were done using an anti-HA spin column kit (ThermoFisher Scientific) according to the manufacturer’s directions.

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Confocal Microscopy

Confocal microscopy was performed using a Zeiss LSM510 inverted confocal microscope with either 10×/0.3, 20×/0.8, or 40×/0.95 Plan-Neofluar air objective. EVFP fluorescence was monitored using a 565±50-nm band-pass emission filter and 514-nm excitation from an argon laser. DAPI was monitored with a 420- to 480-nm band-pass emission filter and 405-nm diode laser excitation. Chlorophyll was observed with a long-pass 575-nm filter and 405-nm diode laser excitation.
Quantification of Nuclear Localization

For quantifying nuclear localization, YFP-CUS-tagged constructs were transiently expressed in tobacco, and confocal microscopy was used to take a Z-stack of images through the epidermal layer of each sample at 10X magnification. For each image, all DAPI-positive cells with YFP signal were marked in NIH ImageJ (https://imagej.nih.gov/ij/; NIH image) and 20 cells were randomly chosen for analysis. All optical sections for each cell were examined for localization of DAPI and YFP. Localization was scored as “not nuclear” if there was no detectable YFP signal inside a given nucleus, or if in any optical section the YFP signal appeared as a ring around, but not within, a nucleus. Conversely, localization was scored as “nuclear” if YFP was clearly visible within the center of a nucleus in the optical section with maximum DAPI signal. Each cell was marked as having cytosolic localization if there was YFP signal visible around the cell’s border.

RNA Extraction and qRT-PCR

Seedlings for RNA extraction were collected, frozen in liquid nitrogen, and stored at −80°C before processing. Samples were ground in liquid nitrogen and RNA was extracted using the Quick-RNA MiniPrep kit with on-column DNase I digestion (Zymo Research). cDNA synthesis was performed with the SuperScript II first-strand cDNA synthesis kit (Invitrogen), using oligo(dT) along with gene-specific primers for plastidial genes listed in Supplemental Table S3. For qRT-PCR, cDNA was reverse transcribed with First Strand Universal SYBR Green Master Mix (Roche) and primers. qRT-PCR reactions were performed in triplicate with a Roche LightCycler 96 thermal cycler. Primers for qRT-PCR of PEP-dependent gene sequences are listed in Supplemental Table S3, while nuclear gene primers are listed in Supplemental Table S4.

GST Pull-Down Assays

Escherichia coli BL21(DE3) cells, expressing either GST or GST-HMRm from pET42b vectors (Novagen), were harvested and centrifuged at 5,000g for 10 min and then resuspended in E buffer containing 50 mM Tris-Cl (pH 7.5), 100 mM NaCl, 1 mM EDTA, 1 mM EGTA, 1% DMSO, 2 mM DTT, and bacterial protease inhibitor cocktail (Sigma-Aldrich). All subsequent purification and binding steps were carried out at 4°C. Cells were lysed by French press, and the lysate was centrifuged at 10,000g for 20 min. Proteins were then precipitated with 3.3 M ammonium sulfate and incubated for 4 h at 4°C. After centrifugation at 10,000g for 30 min, protein pellets were resuspended in E buffer. Insoluble protein was removed by centrifugation at 10,000g for 1 h, and the supernatant was dialyzed against E buffer overnight at 4°C.

To immobilize bait proteins, protein extracts were incubated with glutathione Sepharose beads (GE Healthcare) equilibrated in E buffer for 2 h, and then beads were washed four times in E buffer with 0.1% Nonidet P-40. HA-tagged prey proteins were produced in vitro using the TNT coupled reticulocyte lysate system (Promega) according to the manufacturer’s protocol and with previously described plasmids (Qu et al., 2015). TNT products were then diluted in E buffer with 0.1% Nonidet P-40 and incubated with bead-immobilized bait proteins at 4°C for 2 h. After binding, beads were washed four times in E buffer with 0.1% Nonidet P-40, and then protein elution was boiled in 1× Laemmli sample buffer. Bound proteins were separated by SDS-PAGE, and prey proteins were detected with immunoblots using either polyclonal goat anti-HA antibodies (Genescript) or monoclonal anti-phospho antibodies. Bait proteins were visualized by staining SDS-PAGE gels with Coomassie Brilliant Blue.

Nuclear Fractionation

Nuclei were isolated from Arabidopsis seedlings with some modifications to our previously described protocol (Galvão et al., 2012). Seedlings were ground in liquid nitrogen and homogenized with volumes (mg μL) of nuclear extraction buffer containing 20 mM PIPES-KOH (pH 7.0), 10 mM MgCl2, 12% hexylene glycol, 0.25% Triton X-100, 5 mM β-mercaptoethanol, and 1× EDTA-free protease inhibitor cocktail (Roche) added to the ground tissue. The crude lysate was filtered through a 40-μm cell strainer and incubated on ice for 10 min. The filtrate was then loaded on top of 2 mL of 30% Percoll (Sigma-Aldrich) and centrifuged at 700g for 5 min at 4°C. Filtration through Percoll gradients was repeated five times for each sample to ensure removal of all chloroplast debris. Nuclear pellets were resuspended in extraction buffer containing 100 mM Tris-HCl (pH 7.5), 100 mM NaCl, 1% SDS, 5 mM EDTA, 5 mM DTT, 10 mM β-mercaptoethanol, and 1× EDTA-free protease inhibitor cocktail (Roche). Protein fractions were analyzed with western blots.

Accession Numbers

Sequence data from this article can be found in The Arabidopsis Information Resource or GenBank/EMBL data libraries under the following accession numbers: HEMERA/pTAC12 (At1g34640), PIF1 (At1g09530), PIF4 (At1g43010), PIF5 (At3g59060), PIF6 (At3g56290), PIF7 (At5g61270), PIF8 (At1g00050), PIF11 (At1g46970), ALCATRAZ (At1f57710), and SPATULA (At4g56930).

Supplemental Data

The following supplemental materials are available.

Supplemental Figure S1. YFP-HMR is localized mainly to the cytoplasm and fails to rescue the nuclear defect of the hmr mutant.

Supplemental Table S1. HMR peptides identified by LC-MS/MS analysis.

Supplemental Table S2. Oligonucleotides used for generating constructs used in this study.

Supplemental Table S3. Primers used for cDNA synthesis and qRT-PCR analysis of PEP-dependent plastidial genes.

Supplemental Table S4. Primers used for qRT-PCR analysis of nuclear-encoded genes.

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