Comparative functional analysis of full-length and N-terminal fragments of phytochrome C, D and E in red light-induced signaling

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Summary

- Phytochromes (phy) C, D and E are involved in the regulation of red/far-red light-induced photomorphogenesis of Arabidopsis thaliana, but only limited data are available on the mode of action and biological function of these lesser studied phytochrome species.
- We fused N-terminal fragments and/or full-length phyC, D and E to YELLOW FLUORESCENT PROTEIN (YFP), and analyzed the function, stability and intracellular distribution of these fusion proteins in planta.
- The activity of the constitutively nuclear-localized homodimers of N-terminal fragments was comparable with that of full-length PHYC, D, E-YFP, and resulted in the regulation of various red light-induced photomorphogenic responses in the studied genetic backgrounds. PHYE-YFP was active in the absence of phyB and phyD, and PHYE-YFP controlled responses, as well as accumulation, of the fusion protein in the nuclei, was saturated at low fluence rates of red light and did not require functional FAR-RED ELONGATED HYPOCOTYL1 (FHY-1) and FHY-1-like proteins.
- Our data suggest that PHYC-YFP, PHYD-YFP and PHYE-YFP fusion proteins, as well as their truncated N-terminal derivatives, are biologically active in the modulation of red-light-regulated photomorphogenesis. We propose that PHYE-YFP can function as a homodimer and that low-fluence red light-induced translocation of phyE and phyA into the nuclei is mediated by different molecular mechanisms.

These photoreceptors form two functionally distinct groups, designated as type I and type II phytochromes. Type I phyA plays an important role in the transition from heterotrophic to phototrophic growth, that is, in seedling establishment, which is one of the most critical stages of the plant life cycle. Accordingly, it has been shown that phyA is the primary photoreceptor for very low-fluence responses (VLFR) in a broad spectrum of wavelengths and for high-irradiance responses in continuous FR light (Neff & Chory, 1998). In contrast with other phys, phyA exists only as a homodimer (Sharrock & Clack, 2004). Type II phytochromes regulate R/FR photoreversible low-fluence responses (Rockwell & Lagarias, 2006); phyB is the most prominent type II phytochrome. phyB null mutants display very characteristic deficiencies in R/FR reversible responses and in shade avoidance responses (Reed et al., 1993; Lorrain et al., 2008), whereas phenotypes of phyC, phyD and phyE null mutants are more subtle (Aukerman et al., 1997; Devlin et al., 1998; Monte et al., 2003).

Introduction

Plants are sessile organisms and must adapt to changes in their environment. Light is an important environmental factor, which not only provides energy for photosynthesis, but also regulates the growth and development of plants throughout their entire life cycle. To monitor changes in their ambient light environment, plants have evolved a number of photoreceptors, including the UVB-sensing photoreceptor UVB-RESISTANCE 8 (Rizzini et al., 2011), the blue/UV-monitoring cryptochromes, phototropins, the family of Zeitlupe/FKF1/LKP2 proteins (Demarsy & Funkhauser, 2009; Yu et al., 2010) and the red (R)/far-red (FR) light-absorbing phytochromes (Nagy & Schäfer, 2002). Plants contain multiple forms of phytochromes; there are five genes in Arabidopsis thaliana encoding phytochromes (PHYA–PHYE).

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Homodimers of phyB (Wagner et al., 1996) and phyD (Clack et al., 2009) have been observed in wild-type (WT) and over-expressing lines, and it was assumed until recently that type II phytochromes exist only as homodimers, similar to phyA. Two recent papers (Sharrock & Clack, 2004; Clack et al., 2009), however, have provided compelling evidence that phyB and phyD heterodimerize with each other, phyC and phyE form obligate heterodimers with phyB and phyD in planta, and these heterodimers can bind PHYTOCHROME INTERACTING FACTOR 3 (PIF3), similar to homodimers of phyB. These data suggest a possible novel mechanism by which phyC, phyD and phyE contribute to the fine tuning of phyB-mediated physiological responses. Molecular analysis of PHYA-GREEN FLUORESCENT PROTEIN (GFP)- and PHYB-GFP-controlled signaling cascades has demonstrated that light in a quality- and quantity-dependent fashion regulates nucleo/cytoplasmic partitioning of phytochromes (Kircher et al., 1999, 2002; Yamaguchi et al., 1999). The same authors noticed that phyA and phyB-GFP fusion proteins localized in the nucleus are not distributed evenly in the nucleoplasm, but are associated with specific subnuclear structures, termed speckles, nuclear bodies (NBs) or photobodies. As for phyB, it has been shown that short exposure to R light induces the formation of transiently appearing phyB NBs (termed early NBs), whereas extended irradiation promotes the formation of more stable phyB-associated NBs, also called late NBs (for a recent review, see Van Buskirk et al., 2012). Subsequent work has demonstrated that phyA does not contain an authentic nuclear localization signal (NLS), and it was shown that the nuclear import of phyA is mediated by the FAR-RED ELONGATED HYPOCHOTYL 1/FHY-1 LIKE (FHY1/FHL) proteins (Hiltbrunner et al., 2005, 2006; Fankhauser & Chen, 2008; Pfeiffer et al., 2009). The molecular machinery mediating the nuclear import of phyB is less well understood. It has been reported (Chen et al., 2005) that a short domain within the PAS-PAS region of the C-terminal part of phyB contains an intrinsic NLS that mediates nuclear import, and it has been postulated that R light-induced conformational change of the C-terminal domain facilitates the interaction of phyB with the import machinery. However, more recently, it has been demonstrated that phyB does not possess a functional NLS motif, and that PIF3 promotes light-regulated nuclear import of the photoreceptor in vitro and is also required for translocation of PHYB-GFP into the nucleus during the early phase of the dark-to-light transition in planta (Pfeiffer et al., 2012). In contrast with phyB, the mechanism and identity of the factors involved in translocating cytosolic phyC, phyD and phyE into the nucleus remain to be elucidated.

It has been shown that nuclear localized dimers of short N-terminal PHYB fragments can complement phyB-deficient mutants (Matsushita et al., 2003; Oka et al., 2004; Palagi et al., 2010). These data indicated that the N-terminal fragment of phyB, consisting of the GAF and PHY domains, is sufficient to control and launch signaling cascades that underlie many aspects of R/FR-regulated photomorphogenesis. In contrast with phyB, homodimers of the N-terminal fragment of phyA localized in the nucleus are inactive in launching phyA-controlled signaling (Wolf et al., 2011; Viczian et al., 2012), whereas no data are available to prove whether N-terminal fragments of phyC, phyD and phyE are functional or can substitute to any extent the function of the native photoreceptors. In this work, we have investigated the biological activity of full-length and N-terminal fragments of PHYC, PHYD and PHYE fused to the YELLOW FLUORESCENT PROTEIN (YFP). Our data show that homodimers of the N-terminal fragments of these phytochromes are biologically active, they complement specific mutants lacking phyC, phyD and phyE, and the nuclear localization of these fusion proteins is necessary to launch efficient signaling. Moreover, we demonstrate that homodimers of full-length PHYE-YFP are imported into the nucleus in an R light-induced fashion, this process is induced and saturated at low fluence rates of R light, and translocation of phyE-YFP into the nucleus does not require heterodimerization with phyB and/or phyD.

Materials and Methods

Cloning of the constructs

Full-length PHYC, D and E cDNA fragments were subcloned from the 35S:PHYC, D, E-GFP pPCV plasmids (Kircher et al., 2002) into the 35S:YFP-NOS3’ pPCV812 (PHYC) or 35S:YFP-NOS3’ pPCVB812 (PHYD, PHYE) vectors as BamH-I-EcoRI vectors as BamH-I-EcoRI. The cDNA fragments encoding the N-terminal domain of Arabidopsis PHYC, PHYD and PHYE fused to the YELLOW FLUORESCENT PROTEIN (YFP). Our data show that homodimers of the N-terminal fragments of these phytochromes are biologically active, they complement specific mutants lacking phyC, phyD and phyE, and the nuclear localization of these fusion proteins is necessary to launch efficient signaling. Moreover, we demonstrate that homodimers of full-length PHYE-YFP are imported into the nucleus in an R light-induced fashion, this process is induced and saturated at low fluence rates of R light, and translocation of phyE-YFP into the nucleus does not require heterodimerization with phyB and/or phyD.

Plants transformation and regeneration of transgenic lines

Arabidopsis thaliana (L.) plants were transformed by the Agrobacterium-mediated floral dip method (Clough & Bent, 1998). Details of the raised transgenic lines and the mutant backgrounds used are given in Table S2. From each of these transformations, transgenic seedlings expressing the fusion proteins were selected by their resistance to hygromycin or Basta, and grown to maturity in the glasshouse (for details, see Bauer et al., 2004). Independent homozygous lines expressing one copy of the transgene were selected for further analysis.

Seedling and plant growth conditions and growth measurements

For hypocotyl length measurements, seeds were sown on four layers of filter paper and imbibed in water for 48 h at 4°C. For cotyledon area measurements, seeds were placed on Murashige and Skoog (MS) medium without sucrose. Cold-treated seeds were
then irradiated with white light for 3 h at 22°C to induce seed germination, and transferred to dark for an additional 18 h at 22°C. The plates were then placed under various light conditions for 4 d or otherwise, specified in the figure legends. Seedlings were placed horizontally on the surface of agar medium and scanned (n = 50). Images of scanned seedlings were analyzed using MetaMorph Software (Universal Imaging, Downingtown, PA, USA). Hypocotyl length values, measured at different fluences of light, were normalized to the corresponding dark-grown hypocotyl length to reflect solely the light-dependent regulation.

Flowering time measurement

Seeds were sown in water and incubated for 2 d in the dark at 4°C. They were then placed on the surface of soil and transferred to short-day conditions (SD, 8 h white light : 16 h dark at 22°C). Light sources were fluorescent (cool-white) tubes producing a fluence rate of 60 μmol m⁻² s⁻¹. The flowering time was recorded as the number of rosette leaves at the time at which inflorescences reached a height of 1 cm (n = 40).

Epifluorescence microscopy

Seeds were sown on a four-layer filter paper and imbibed in water in the dark for 48 h at 4°C. Cold-treated seeds were then transferred to 25°C and irradiated with 18 h of white light to induce homogeneous germination, and grown for additional days in the dark. Six-day-old dark-grown seedlings were then subjected to various light treatments, as described in the text. The standard epifluorescence microscopy set-up and observation techniques have been described previously (Bauer et al., 2004; Viczian & Kircher, 2010; Sokolova et al., 2012). For semiquantitative epifluorescence microscopy, etiolated seedlings were irradiated for 6 h with R light at the fluence rates given in the figure legends. Using an Axioplan microscope (Zeiss, Germany) equipped with a CoolSnap HQ camera (Photometrics, USA), 12-bit TIFF images, not containing saturated pixels, were taken of nuclei in epidermal cells of hypocotyls. In order to minimize the effect of the microscopic light, images were taken within the first 120 s after the onset of excitation light. Adjusted and identical exposure times and excitation light intensity settings were applied throughout the analysis of each genotype. The average intensity of nuclear pixels was calculated using ImageJ software (National Institutes of Health, USA), including the subtraction of background signals in each image. The mean value of the data obtained from at least 25 independent nuclei was normalized to the corresponding dark control. For each genotype, three independent biological replica experiments were performed.

Plant protein extraction and western blot hybridization

Seedling protein extracts were prepared in extraction buffer (50 mM Tris, pH 8.0, 150 mM NaCl, 0.1% NP-40), as described previously (Sharrock & Clack, 2004). Proteins were fractionated on 6% sodium dodecylsulfate-polyacrylamide gel electrophoresis (SDS/PAGE) gels and transferred to Hybond-ECL membranes (GE Healthcare). Membranes were blocked overnight at 4°C with blocking buffer (5% non-fat dry milk, 0.2% Tween 20 in TBS-T buffer, pH 7.6). Membranes were probed in blocking buffer containing the following primary monoclonal antibodies: anti-phyC C11 and C13, anti-phyD 2C1, and anti-phyE 7B3 (Hirschfeld et al., 1998). After three washes with TBS-T buffer, chemiluminescent detection of primary antibodies was performed with horseradish peroxidase-conjugated secondary antibody and Supersignal West Pico reagents (Thermo Fisher Scientific). Total protein was analyzed by the Bio-Rad Protein Assay. For native gel electrophoresis, 7-d-old seedlings grown under the described light conditions were ground at 0°C under dim green safe light at a 1 : 1 weight : volume ratio in non-denaturing extraction buffer (25 mM Tris-HCl, pH 7.5, 10 mM NaCl, 5 mM EDTA) containing Complete EDTA-free Protease Inhibitor Cocktail (Roche Diagnostics, Indianapolis, IN, USA), and the extracts were centrifuged for 3 min at 4°C. Proteins were separated on 4–20% gradient PAGE gels in Tris/borate/EDTA buffer for 40 h at 4°C. Gel blotting was the same as for SDS-PAGE gels, and blots were probed with anti-GFP antibody GF28R (Thermo Fisher Scientific). Secondary antibodies coupled to horseradish peroxidase (Bio-Rad) were used according to the manufacturer’s recommendations.

Analysis of transcript levels

Seeds were surface sterilized and plated onto MS medium, stratified at 4°C in the dark for 3 d, and exposed to white light for 3 h to induce germination. Subsequent growth conditions and light inductions were carried out as described in the corresponding figure legends. RNA samples were prepared from whole seedlings using RNeasy Miniprep Kits (Qiagen) according to the manufacturer’s instructions, and DNA was removed by DNase I treatment. cDNA was synthesized from 1 μg of total RNA using the Revert Aid First Strand cDNA Synthesis Kit (Fermentas). Real-time reverse transcription-polymerase chain reaction (RT-PCR) analysis was carried out with a 7500 Real-Time PCR System with SYBR Green JumpStart TaqReadyMix (Sigma). The expression levels were normalized to the expression of TUBULIN2/3. The experiments were performed at least three times, and a representative dataset is presented. The primers used in qRT-PCRs are listed in Table S1.

Accession numbers

PHYB, AT2G18790; PHYC, AT5G35840; PHYD, AT4G16250; PHYE, AT4G18130; FHY1, AT2G37678; FHL, AT5G02200; PRR9, At2g46790; CAB2, At1g29920; ELIP1, AT3G22840; EXPANSINS5, AT3G29030; EXPANSIN9, AT5G02260; BBX23, AT4G10240; HB4, AT2G44910; TUBULIN2/3, AT5G62690.

Results

To test whether homodimers of N-terminal fragments of phyC, phyD and phyE, similar to those of phyB, can launch R
light-induced signaling, we constructed chimeric genes consisting of the N-terminal domains of PHYC, PHYD and PHYE fused to the YFP reporter under the control of the viral 35S promoter. These domains were selected on homology to the N-terminal 1–651-amino-acid region of PHYB (Fig. S1). Appropriate dimerization and targeted, constitutive, subcellular localization were achieved by the addition of the leucine-zipper domain of the CPRF transcription factor (DD) and the SV 40 NLS peptide or nuclear exclusion signal (NES), as described by Palágyi et al. (2010) and Wolf et al. (2011). The domain structures of the various fusion proteins are shown in Fig. S1. The synergistic and, in some cases, antagonistic action of phytochrome species makes the elucidation of the roles of minor family members in single mutants difficult. Therefore, we used double-, triple- or quadruple-mutant combinations showing characteristic deficient phenotypes to assess the functions of these fusion proteins. We regenerated 15 independent transgenic lines for each construct and used quantitative western blot hybridization to select lines which expressed the individual full-length PHYC-YFP, PHYD-YFP and PHYE-YFP, and their N-terminal derivative fusion proteins, approximately at the same or, at least, comparable levels. The genetic backgrounds of the transgenic lines and the expression levels of the various fusion proteins relative to their endogenous phyC, phyD and phyE counterparts are shown in Tables S2 and S3, respectively.

Homodimers of phyC N-terminal fragments localized in the nucleus are functional in regulating R light-induced signaling

The selected transgenic phyC/phyD mutant lines over-expressed the PHYC-YFP c. two-fold, whereas the expression level of the PHYC602-YFP-DD-NLS and PHYC602-YFP-DD-NES fusion proteins was c. 40% or 70% of the endogenous phyC, respectively (Table S3). PHYC-YFP fully, and the PHYC602-YFP-DD-NLS fusion protein partially, restored R light-induced hypocotyl growth inhibition of the phyC/phyD mutant (Fig. 1a). R light-induced expansion of the cotyledon area of the mutant was also restored (Fig. 1b) and, in this case, PHYC-YFP-expressing seedlings exhibited a weak over-expression phenotype, whereas the PHYC602-YFP-DD-NLS seedlings displayed a nearly fully complemented phenotype. These fusion proteins also restored phyC function in regulating the flowering time under SD conditions (Fig. 1c). By contrast, the PHYC602-YFP-DD-NES photoreceptor was biologically inactive in all responses tested, as the PHYC602-YFP-DD-NES-expressing seedlings invariably
displayed the original phyC/phyD mutant phenotype in all experiments performed (Fig. 1a–c). The abundance of the native phyC and the various PHYC-YFP fusion proteins was down-regulated by constant R light (cR) irradiation. Irradiation with 120 h of cR light (25 μmol m⁻² s⁻¹) reduced the level of the endogenous phyC below the detection level (Fig. S2a), and decreased the levels of all other phyC fusion proteins c. four-fold (Fig. S2b).

Microscopic analysis of the nucleo/cytoplasmic distribution of the various PHYC-YFP fusion proteins demonstrated that, in etiolated as well as R light-irradiated seedlings, PHYC602-YFP-DD-NLS was detectable only in the nucleus (Fig. S3a–c), PHYC602-YFP-DD-NES was localized exclusively in the cytoplasm (Fig. S3d–f) and PHYC-YFP was observed in both the nucleus and cytoplasm in etiolated as well in R light-irradiated seedlings (Fig. S3g–i). A 5-min R light treatment induced the formation of early PHYC-YFP NBs, whereas a prolonged, 24-h R light treatment promoted the formation of late, stable PHYC-YFP NBs. Under these conditions, we could not detect the appearance of either early or late NBs associated with the truncated forms of PHYC. Quantitative analysis of R light-induced accumulation of the fusion protein indicated that a 6-h cR light treatment did not modify significantly the level of PHYC-YFP in the nucleus (Fig. 1d).

Homodimers of phyD N-terminal fragments localized in the nucleus restore phyD signaling

The selected transgenic phyA/phyB/phyD plants over-expressed PHYD-YFP two-fold and the PHYD-N654-YFP-DD-NLS and PHYD-N654-YFP-DD-NES fusion proteins c. 25 and 20-fold, respectively, when compared with endogenous phyD (Table S3). Analysis of R light-induced hypocotyl growth inhibition (Fig. 2a), cotyledon expansion (Fig. 2b) and flowering time (Fig. 2c) under SD conditions demonstrated that the PHYD654-YFP-DD-NLS and PHYD654-YFP-DYNES fusion proteins associated with the nucleus restore phyD signaling.

PHYD654-DD-YFP-NES was detected exclusively in the cytoplasm, and these truncated proteins never associated with NBs (Fig. S5a–c and d–f, respectively). PHYD-YFP was clearly visible in the nucleus in etiolated seedlings, and R light treatments did not affect the amount of photoreceptor localized in the nucleus or induce the formation of early/late PHYD-YFP-containing NBs (Fig. S5h–i). Quantitative analysis of the accumulation of PHYD-YFP in the nucleus corroborated these observations (Fig. 2d).

Homodimers of phyE N-terminal fragments localized in the nucleus are capable of signaling in red light

The selected transgenic phyA/phyB/phyE plants expressed the PHYE-YFP and PHYE593-YFP-DD-NES fusion proteins eight- to ten-fold and the PHYE593-YFP-DD-NLS protein c. three- to four-fold more strongly than the native phyE (Table S3). Analysis of R light-induced inhibition of hypocotyl growth and cotyledon expansion demonstrated that, with the exception of PHYE593-YFP-DD-NES, these fusion proteins efficiently complemented the phyA/phyB/phyE mutant, and even displayed characteristic over-expression phenotypes when compared with phyA/phyB mutants (Fig. 3a,b). These responses were induced at very low fluence rates (<0.001 μmol m⁻² s⁻¹) and saturated at low fluence rates (0.01 μmol m⁻² s⁻¹). Transgenic plants expressing nuclear localized homodimers of N-terminal fragments of PHYE, but not the full-length PHYB, were similarly hypersensitive to low fluences of R light (Oka et al., 2004). The same fusion proteins, again with the exception of the truncated PHYE593-YFP-DD-NES, complemented the flowering time of the phyA/phyB/phyE triple mutant (Fig. 3c). The abundance of the native phyE and the various PHYE-YFP fusion proteins was down-regulated by irradiation with cR light (25 μmol m⁻² s⁻¹). Exposure to 120 h cR light reduced the abundance level of the native phyE c. two-fold (Fig. S6a) and that of the full-length and truncated PHYE-YFP fusion proteins c. four-fold (Fig. S6b). We also analyzed the cellular distribution of the full-length and truncated PHYE-YFP fusion proteins in this genetic background. The PHYE593-YFP-DD-NLS fusion protein was constitutively nuclear; its accumulation in the nucleus did not increase after prolonged exposure to R light (Fig. S7a–c). By contrast, PHYE593-YFP-DD-NES, as expected, was detectable only in the cytoplasm, independent of the light conditions (Fig. S7d–f). As for PHYE593-YFP-DD-NLS, short 5-min R light irradiation induced the formation of early NBs, but extended R treatment did not promote the appearance of late, stable NBs associated with this fusion protein (Fig. S7a–c). By contrast, we could not detect the formation of any NBs containing PHYE593-YFP-DD-NES. The PYHE-YFP fusion protein was detectable in the nuclei of etiolated seedlings, but a 24-h R light treatment clearly induced its nuclear abundance. In addition, R light also induced the appearance of early NBs, but did not promote the formation of late, stable PHYE-YFP-associated NBs (Fig. S7g–i). Quantitative analysis of the R light-induced accumulation of the PHYE-YFP fusion protein in the nucleus corroborated these observations (Fig. 3d).
Homodimers of PHYE-YFP are functional and imported into the nucleus at very low fluences of R light independent of phyA, phyB and phyD.

To test to what extent the functionality of PHYE-YFP depends on phyD, the PHYE-YFP transgene from the phyA/phyB/phyE mutant was introgressed into the phyA/phyB/phyD/phyE quadruple null background. Analysis of R light-induced hypocotyl growth inhibition demonstrated that PHYE-YFP is functional in the absence of phyA, phyB and phyD (Fig. 4a). This figure also shows that seedlings expressing PHYE-YFP display a characteristic over-expression phenotype when compared with the phyA/phyB/phyD/phyE mutant, and the physiological response in this genetic background also saturates at 0.01 μmol m⁻² s⁻¹ fluence rate of R light, and resembles the data obtained by the analysis of PHYE-YFP function in the phyA/phyB/phyE triple null background (Fig. 3a). The extreme sensitivity of these PHYE-YFP responses to R light prompted us to test whether the biological activity and nuclear accumulation of PHYE are mediated by FHY1/FHL proteins, shown to be essential for the translocation of phyA into the nucleus. To this end, we produced PHYE-YFP-expressing transgenic phyA-201 null mutant lines. Figure 4(b) illustrates that the PHYE-YFP-expressing seedlings are hypersensitive to low fluences of R light when compared with phyA-201 or Col-0 WT. Quantitative analysis of R light-induced accumulation of PHYE-YFP in the nucleus demonstrated that this process is saturated at 0.08 μmol m⁻² s⁻¹ fluence rate of R light in contrast with PHYB-GFP (Fig. 4c–d). We note that R light is ineffective in inducing the formation of early and/or late, stable PHYE-YFP NBs in the phyA/phyB/phyD/phyE background, whereas these nuclear structures were readily detectable in the phyA/phyB/phyE mutant (Table S4). To test whether the PHYE-YFP photoreceptor functions and is imported into the nucleus as a monomer or homodimer, we analyzed total protein extracts prepared from etiolated and R light-treated seedlings by native gel electrophoresis. Our data demonstrate that the PHYE-YFP fusion protein is detected nearly exclusively as a homodimer in these extracts, and that extended irradiation with R light reduces significantly the abundance of PHYE-YFP (Fig. 4e). In addition, we show that, similar to the phyA-201 null mutant, a short pulse of R light is not capable of inducing significant accumulation of PRR9, CAB2 or ELIP1 transcripts in transgenic phyA/phyB/phyD/phyE seedlings over-expressing PHYE-YFP (Fig. 5a). However, Fig. 5(b) indicates that over-expressed PHYE-YFP in the phyA/phyB/phyE and/or phyA/phyB/phyD/phyE background can restore the transcriptional regulation of EXPANSIN5 and 9, thought to regulate cell wall extension in an opposite fashion. Taken together, these data strongly suggest that PHYE-YFP can function as a homodimer.
independent of phyB, phyD and phyA, the nuclear import of phyE does not require functional FHY1/FHL proteins, the PHYE-controlled signaling cascade operates differently from that controlled by phyA at very low fluences of R light, and the formation of late, stable PHYE-YFP NBs requires functional phyB.

Discussion

In this work, we have demonstrated that homodimers of the N-terminal fragments of PHYC, PHYD and PHYE are biologically active and restore the photomorphogenic phenotypes of specific mutants lacking these photoreceptors. Our data show a good correlation between the expression levels and physiological activities of these truncated photoreceptors, suggesting that their C-terminal domains are dispensable for signaling. These data also indicate that import/accumulation of the photoreceptors to/in the nucleus is essential for biological activity. Moreover, our data demonstrate that exposure to R light modulates the abundance of these fusion proteins, similar to native phyC, phyD and phyE (Figs S2, S4, S6), and they uniformly fail to produce detectable late NBs (Figs S3, S5, S7). These observations are strikingly similar to the data reported for phyB N-terminal fragments (Matsushita et al., 2003; Oka et al., 2004; Palágyi et al., 2010), but, in contrast with phyB, the mode of action by which PHYC, PHYD and PHYE N-terminal fragments launch R/FR-dependent signaling is not understood. As for phyB and PHYB N-terminal fragments, it has been reported that they mediate light signaling by inhibiting the activity of PIFs via the release of these bHLH-type transcription factors from their DNA targets (Park et al., 2012). The inhibition requires interaction between the Pfr form of the photoreceptor and PIFs, and it has been concluded that this interaction competes directly with DNA binding.

At present, there are no data available to demonstrate that homodimers of full-length or N-terminal fragments of PHYC, PHYD and PHYE Pfr can bind to PIF3 or any other PIFs, either in vitro or in vivo. In this respect, we note that the data reported by Clack et al. (2009) show that heterodimers of full-length phyB/phyC and phyb/phyD can be co-immunoprecipitated with PIF3. However, we should emphasize that it is highly unlikely that PHYC, PHYD and PHYE N-terminal fragments would heterodimerize with full-length phyB and phyD, and, in addition, we demonstrate in this work that full-length PHYE-YFP is functional in the absence of phyB and phyD. Thus, we conclude that, in the absence of additional molecular details, the precise mechanism underlying R/FR reversible signaling by these phytochrome species remains elusive.

Independent of the molecular mechanism governing signaling by the PHYC, PHYD and PHYE N-terminal fragments, our
results clearly demonstrate that, pairwise, the truncated and full-length phyC, phyD and phyE photoreceptors regulate the same responses, which, in turn, display similar fluence rate dependencies. Close inspection of the data also reveal that the responses regulated by PHYE-YFP and N-terminal PHYE-YFP are not only initiated, but also saturated, at much lower R light intensities than those controlled by PHYC and PHYD. The data shown in Fig. 4(b) indicate that 8–10 times over-expressed PHYE could contribute to signaling at very low fluences of R light. The data reported by Hennig et al. (2002) also demonstrate a similar role for native phyE in controlling seed germination, and thus provide further support for the conclusion above. Microarray analysis has...
documented that phyA plays a prominent role in launching the R light-induced transcriptional cascade(s) (Tepperman et al., 2006), and the same authors showed that R light is ineffective in inducing the transcription of key regulatory transcription factors in the phyA/phyB mutant (Tepperman et al., 2004). These data suggest that the molecular mechanism by which PHYE regulates growth is substantially different from that used by phyA and phyB. Independent of the molecular mechanism mediating the phyE-controlled signaling cascade, our data show that R light-induced or repressed transcription of EXPANSIN5 and EXPANSIN9, respectively, is reconstituted in the PHYE-YFP-complemented mutant lines (Fig. 5b), and this process does not require functional phyB and phyD. PHYE593-YFP-DD-NLS, similar to PHYE-YFP, is also active in controlling the transcription of selected genes in the complemented transgenic plants (Fig. S8). As for the early steps of R light-induced signaling, it is generally accepted that light quality- and quantity-dependent translocation of phyA (Hilbrunner et al., 2005) and phyB (Kircher et al., 1999) into the nucleus is critical and rate limiting. The import of phyA into the nucleus is mediated by FHY1/FHL (Genoud et al., 2008). Here, we show that the fluence rate dependence of physiological responses and the nuclear accumulation of the PHYE-YFP fusion protein display excellent correlation. The accumulation of PHYE-YFP in the nucleus is induced by extremely low intensities of R light in WT, phyA/phyB/phyD and phy1/fhl mutant seedlings, and saturates at low intensities of R light (Fig. 4c). These data demonstrate that translocation of PHYE-YFP into the nucleus is not regulated by FHY1/FHL, and obligate heterodimerization of phyE with phyB (Clack et al., 2009) is dispensable for the regulation of this process, but does not exclude the possibility that phyB/phyE heterodimers are also imported into the nucleus in planta. The same authors also reported that over-expressed PHYE-MYC6 fusion protein is monomeric and is assumed to be inactive in R/FR light-induced signaling. By contrast, we detected PHYE-YFP mainly as homodimers in the transgenic phyA/phyB/phyD mutant, in both etiolated and R light-treated seedlings. We speculate that the apparent contradiction could be the result of the increased stability of PHYE-YFP homodimers relative to PHYE-MYC6 or, alternatively, the MYC6 tag may destabilize the inherently weak phyE homodimers. We also note that the abundance of the PHYE-YFP homodimers (Fig. 4e) appears to be more strongly reduced by extended irradiation of R light when compared with the total amount of PHYE-YFP (Fig. S6a,b). Whether the accelerated degradation of phyE homodimers plays a role in the saturation of phyE-controlled responses at low fluence rates of R light remains to be elucidated. Notwithstanding these as yet unresolved issues, our data demonstrate that over-expressed PHYE-YFP forms homodimers, PHYE-controlled physiological responses saturate at low fluence rates of R light, and the import of PHYE-YFP homodimers into the nucleus is mediated by as yet unknown molecular machinery that differs substantially from that described for phyA. When compared with phyE, the nuclear import of phyB displays a different fluence rate dependence, in that it is not saturated at low intensities of R light, and it has been shown that PIFs may be involved in the mediation of translocation of phyB into the nucleus (Pfeiffer et al., 2012). Thus, it is tempting to speculate that the molecular mechanism required for R-induced accumulation of phyE and phyB in the nucleus is also different; however, validation of this hypothesis requires additional molecular and genetic evidence. PhyB Pfr localized in the nucleus is associated with nuclear complexes, termed nuclear bodies, whose size and number appear to be modulated by the intensity and duration of exposure to R light (Kircher et al., 1999; Chen et al., 2004; Van Buskirk et al., 2012). It has been reported by Kircher et al. (2002) that PHYB-GFP, PHYC-GFP and PHYE-GFP fusion proteins are associated...
with stable NBs in white light-grown transgenic Ws seedlings. Here, we report that we were unable to detect late, stable NBs associated with the truncated PHYC, PHYD and PHYE-YFP fusion proteins (Table S4). By contrast, we observed the formation of such PHYC-YFP and PHYE-YFP NBs in transgenic seedlings expressing the native phyB photoreceptor, but not in transgenic phyA/phyB/phyD mutants, independent of the R light treatments applied (Table S4). These data suggest that the formation of stable phyC and phyE NBs under extended R irradiation requires either phyB or active signaling by phyB. We assume that the formation of stable PHYD-YFP NBs requires signaling by other photoreceptor(s), as it can only be detected in white light (Table S4). The formation of early, transient phyC, phyD and phyE NBs displays a more complex pattern. It appears that R light only induces the formation of transient PHYE593-YFP-DD-NLS NBs in the phyA/phyB/phyE mutant background (Figs S3, S5, S7). With regard to the R light-induced formation of transient PHYC-YFP and PHYE-YFP NBs, we have shown that such NBs can be detected in the WT Ws background. We have also demonstrated that native phyB is dispensable for the formation of transient phyE NBs, but the absence of both native phyB and phyD prevents the formation of such PHYE-YFP NBs (Table S4). By contrast, we found that PHYD-YFP fails to form R light-induced transient NBs in any genetic background investigated. On the basis of these data, we tentatively conclude that the formation of these transient NBs might be associated with heterodimerization, but this is dismissible, at least for PHYE-YFP signaling.

Taken together, our data suggest that PHYC-YFP, PHYD-YFP and PHYE-YFP fusion proteins, as well as their truncated N-terminal derivatives, are biologically active in the modulation of R light-regulated photomorphogenesis in the specific genetic backgrounds tested. The transgenic plants generated could be useful to gain more insight into the molecular mechanism by which these type II phytochromes signal. Furthermore, independent of the precise mode of action, our data collectively suggest that the truncated PHYC, PHYD and PHYE photoreceptors function as R/FR reversible switches in planta. N-terminal fragments of phyB have been successfully used to control cellular signaling processes in a R/FR reversible fashion in bacteria, yeast and mammalian cells (for a review, see Wang et al., 2012). Thus, we believe that it could be advantageous to explore the applicability of our chimeric photoreceptors to synthetic biology research approaches.

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References


Supporting Information

Additional supporting information may be found in the online version of this article.

**Fig. S1** Schematic illustration of the domain structures of the native Arabidopsis phytochrome proteins and chimeric PHYC, D and E N-terminal fusion proteins used in this study.

**Fig. S2** Immunoblot analysis of the expression level of the various PHYC-YFP fusion proteins compared with the native phyC in Ws wild-type.

**Fig. S3** Intracellular localization of the various PHYC-YFP fusion proteins determined in transgenic phyC/phyD seedlings.

**Fig. S4** Immunoblot analysis of the expression level of the various PHYD-YFP fusion proteins compared with the native phyD in Ler wild-type seedlings.

**Fig. S5** Intracellular localization of the various PHYD-YFP fusion proteins determined in transgenic phyA/phyB/phyD seedlings.

**Fig. S6** Immunoblot analysis of the expression level of the various PHYE-YFP fusion proteins compared with the native phyE in Ler wild-type seedlings.

**Fig. S7** Intracellular localization of the various PHYE-YFP fusion proteins determined in transgenic phyA/phyB/phyE seedlings.

**Fig. S8** PHYE regulates red light-induced gene repression.

**Table S1** Oligonucleotides used in this study

**Table S2** Summary of the generated transgenic lines and references to the mutant backgrounds used

**Table S3** Comparison of endogenous and transgenic PHYC, PHYD and PHYE protein levels in etiolated seedlings calculated on immunoblot assays

**Table S4** Red light-dependent formation of nuclear bodies

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