Functional Analysis of Amino-Terminal Domains of the Photoreceptor Phytochrome B1[C][W]

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At the core of the circadian network in Arabidopsis (Arabidopsis thaliana), clock genes/proteins form multiple transcriptional/translation negative feedback loops and generate a basic approximately 24-h oscillation, which provides daily regulation for a wide range of processes. This temporal organization enhances the fitness of plants only if it corresponds to the natural day/night cycles. Light, absorbed by photoreceptors, is the most effective signal in synchronizing the oscillator to environmental cycles. Phytochrome B (PHYB) is the major red/far-red light-absorbing phytochrome receptor in light-grown plants. Besides modulating the pace and phase of the circadian clock, PHYB controls photomorphogenesis and delays flowering. It has been demonstrated that the nuclear-localized amino-terminal domain of PHYB is capable of controlling photomorphogenesis and, partly, flowering. Here, we show (1) that PHYB derivatives containing 651 or 450 amino acid residues of the amino-terminal domains are functional in mediating red light signaling to the clock, (2) that circadian entrainment is a nuclear function of PHYB, and (3) that a 410-amino acid amino-terminal fragment does not possess any functions of PHYB due to impaired chromophore binding. However, we provide evidence that the carboxyl-terminal domain is required to mediate entrainment in white light, suggesting a role for this domain in integrating red and blue light signaling to the clock. Moreover, careful analysis of the circadian phenotype of phyB-9 indicates that PHYB provides light signaling for different regulatory loops of the circadian oscillator in a different manner, which results in an apparent decoupling of the loops in the absence of PHYB under specific light conditions.

Circadian clocks regulate many rhythmic cellular and physiological processes and allow a wide range of organisms to adapt to the predictable daily changes in the environment (e.g. day/night cycles; Harmer, 2009). At the core of the clock, components of the central oscillator (clock genes/proteins) mutually regulate their expression/activity via multiple feedback loops, which results in an autonomous, self-sustained approximately 24-h oscillation. The core oscillation is relayed to diverse clock-controlled processes (gene expression, physiology, behavior) via the output pathway (Mas and Yanovsky, 2009). As the period length of the core oscillation always deviates from 24 h, the clock must be resynchronized to the environmental cycles regularly in order to provide precise temporal information. In nature, this resetting occurs on a daily basis in response to periodic environmental cues (e.g. changes in temperature and light conditions). Light is absorbed by specialized photoreceptors, and signals are forwarded by the input pathway to modulate the pace and the phase of the oscillator (Kozma-Bognár and Kaldi, 2008). The Arabidopsis (Arabidopsis thaliana) circadian oscillator is supposed to consist of three interlocked feedback loops (Locke et al., 2006). In the first loop (central or coupling loop), the morning-expressed CIRCADIAN CLOCK ASSOCIATED1 (CCA1)/LATE ELONGATED HYCOCOTYL (LHY) transcription factors inhibit the expression of the TIMING OF CAB EXPRESSION1 (TOCI) gene; conversely, the evening-expressed TOCI positively regulates the transcription of CCA1/LHY (Alabadi et al., 2001). In the second loop (evening loop), the predicted factor Y induces TOCI expression during the afternoon/evening, while TOCI represses Y during the night (Locke et al., 2005). It has

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been demonstrated that GIGANTEA (GI) functions as a component of Y (Locke et al., 2006). In the so-called morning loop, CCA1/LHY up-regulate the PSEUDO RESPONSE REGULATOR7/9 (PRR7/9) genes (homologs of TOC1) in the morning, and PRR7/9 proteins down-regulate CCA1/LHY expression during the day (Farre et al., 2005). The coordinated function of the three loops is required to generate the approximately 24-h basic oscillations in Arabidopsis (Locke et al., 2006).

Light signals are perceived and transduced to the clock by specialized photoreceptors, including members of the blue light-absorbing cryptochrome (CRY) and the red/far-red light-sensing phytochrome (PHY) families (Lin and Todo, 2005; Franklin and Quail, 2010). The Arabidopsis PHY photoreceptor family consists of five members (PHYA to PHYE), which function as molecular light switches. In the dark, PHYs are present in their inactive red light-absorbing (Pr) form (Amax = 660 nm). After capturing a photon by the covalently bound linear tetrapyrrole chromophore, they are converted to the active far-red light-absorbing conformer (Pfr), which initiates downstream signaling events in the cytosol or in the nucleus. The active Pfr form is converted to Pr by far-red light (Amax = 730 nm; Rockwell et al., 2006). The Pfr conformers of PHYs are translocated to the nuclei, where they form characteristic nuclear bodies (Kircher et al., 2002). The exact composition and function of nuclear bodies are not yet known, but they may represent multiprotein complexes, where PHYs interact with transcription factors and other regulatory proteins to control the expression of light-induced genes (Bae and Choi, 2008).

The involvement of PHYA, PHYB, PHYD, and PHYE has been demonstrated in the function of red light input to the clock (Somers et al., 1998; Devlin and Kay, 2000; Yanovsky et al., 2001). PHY signaling affects transcription rate and mRNA and protein turnover of several clock components, although the signal transduction pathways linking these events with light-activated PHYs are largely unknown (Kozma-Bognar and Kaldi, 2008). On the other hand, several reports have demonstrated that PHY-mediated continuous red light signals inversely affect the free-running period length in a fluence rate-dependent manner (parametric entrainment), and discrete red light pulses elicit characteristic phase shifts of the clock and overt circadian rhythms (nonparametric entrainment; Millar, 2003). The role of PHYB has been seen in both processes. It has been shown by several research groups that the absence of PHYB function (i.e. in phyB mutants) results in a long period of the rhythmic transcription of the CHLOROPHYLL A/B BINDING2 (CAB2) gene at higher fluences of continuous red light (Somers et al., 1998; Devlin and Kay; 2000; Hall et al., 2002), whereas no period alterations were observed in phyB mutants in continuous white light (Hall et al., 2002; Salome et al., 2002). Interestingly, phyB-9 but not phyB-1 displayed an early phase of CAB2:LUC expres-
GUS protein, a well-defined short dimerization domain of the transcription factor CPRF4 (Kircher et al., 1998) was fused to the PHYB fragments to facilitate the formation of homodimers. Subcellular localization was controlled by short NLS or nuclear exclusion signals (NES). All fusion proteins carried the yellow fluorescent protein (YFP) for easy detection of the fusion proteins in vitro and in vivo. Here, we demonstrate that both the 651- and 450-amino acid N-terminal fragments of PHYB mediate entrainment of the circadian clock. We show that this function of PHYB requires nuclear localization of the receptor. We provide evidence that the 410-amino acid N-terminal fragment lacks any physiological functions of PHYB, because it fails to autoligate the chromophore required for light absorption. Moreover, detailed quantitative analysis of core clock gene expression in phyB-9 indicates that PHYB affects the pace of the morning and the evening loops to different extents, which results in the apparent decoupling of the two loops under specific light conditions.

RESULTS

Structure, Expression, and Subcellular Localization of N-Terminal PHYB Fragments in Transgenic Plants

To test the functions of PHYB derivatives of different lengths, truncated versions containing 651, 450, and 410 amino acid residues were expressed in fusion with the YFP, a dimerization domain, and a NLS in the phyB-9 mutant background (B561-NLS, B450-NLS, and B410-NLS, respectively). The wild-type full-length PHYB protein tagged with YFP was used as a positive control (BFL), whereas B651-NES, which contains a NES instead of the NLS, was generated to test the effect of subcellular localization of PHYB on its function. Figure 1A shows the structures of the fusion proteins and the characteristic domain structure of the wild-type PHYB receptor. The approximately 120-kD PHYB folds into two major domains, the N-terminal and the C-terminal parts, which are separated by a flexible hinge region. The very N-terminal end of phytochromes plays a role in regulating the stability of the Pfr conformer (Sweere et al., 2001; Trupkin et al., 2007). The bilin-lyase motif carries the covalently bound tetrapyrole chromophore phytochromobilin (Wu and Lagarias, 2000). The phytochrome domain (PHY) contributes to the integrity of Pfr (Montgomery and Lagarias, 2002). The first half of the C-terminal part, called the PAS-related domain, contains two PAS-like subdomains, a core regulator region termed the Quail box, and dimerization motifs. A His kinase-related domain showing homology to bacterial two-component sensor kinases is located at the C terminus of the protein (Rockwell et al., 2006). Expression of the different PHYB fragments was tested by western-blot analysis (Fig. 1B). B450-NLS was expressed at a slightly higher level; BFL, B651-NLS, B651-NES, and B410-NLS showed comparable expression levels in transgenic lines selected for further experiments. Subcellular localization of the fusion proteins was tested by confocal laser scanning and epifluorescence microscopy. Confocal images shown in Figure 2 report YFP fluorescence data from 1-µm optical sections of cells and convincingly demonstrate that the vast majority of NLS- or NES-tagged PHYB derivatives are localized within or outside the nucleus, respectively. The NLS-tagged forms showed diffuse distribution in the nucleus, whereas BFL formed nuclear bodies in the nuclei of etiolated plants illuminated with red light for 24 h, in agreement with previously published results. Epifluorescence images shown in Supplemental Figure S1 demonstrate that nuclear translocation and the formation of nuclear bodies by BFL are light dependent, whereas the intracellular distribution pattern of B651-NLS or B651-NES was unaffected by the light conditions. These results verify the correct expression and subcellular compartmentalization of the truncated PHYB fragments.
Differential Complementation of Photomorphogenic and Flowering Time Phenotypes of \textit{phyB-9} by \textit{PHYB} Derivatives

To test the effectiveness of the different PHYB fusion proteins to inhibit hypocotyl elongation, seedlings were grown at different fluence rates of red light for 4 d and hypocotyl lengths were measured. Figure 3A shows that BFL, B651-NLS, and B450-NLS could rescue the long-hypocotyl phenotype of \textit{phyB-9}, whereas B651-NES and B410-NLS were ineffective. BFL was hypersensitive compared with ecotype Columbia-0 (Col-0) at all fluence rates. B651-NLS and B450-NLS showed hypersensitivity at medium, but especially at low, fluence rates. Similar results were obtained from plants grown in continuous white light or in light/dark cycles (data not shown). Mutations in the \textit{PHYB} gene cause early flowering under long-day (Neff and Chory, 1998), short-day, or even continuous light conditions (Endo et al., 2005). To test if the truncated PHYB fragments are able to restore the early-flowering phenotype of \textit{phyB-9}, plants were grown in long-day (16-h-white light/8-h-dark) conditions and flowering time was determined as the number of rosette leaves at the time of bolting. Figure 3B demonstrates that BFL
and B651-NLS flowered as the wild type, but B651-NES, B450-NLS, and B410-NLS were indistinguishable from phyB-9.

These results indicate that the 651-amino acid N-terminal fragment of PHYB acts as a functional photoreceptor if it is localized in the nucleus, whereas B410-NLS is not sufficient to regulate the light-dependent physiological processes tested. Interestingly, B450-NLS could mediate signaling to inhibit hypocotyl elongation, but it was unable to function in flowering time determination.

**The phyB-9 Mutation Affects Rhythmic Transcription of Central Clock Components in a Different Manner**

It has been demonstrated that loss of PHYB function results in a long-period phenotype of the CAB2:LUC marker at medium and high fluence rates of continuous red light (Somers et al., 1998). However, it was presumed, but not directly tested, that rhythmic expression of core clock components is affected in the same way. To test this, CAB2:LUC+, CCA1:LUC+, TOC1:LUC+, GI:LUC+, and PRR9:LUC+ markers were introduced in Col-0 and phyB-9 backgrounds, and free-running period length and circadian phase of the expression of these markers were tested at different fluence rates of red light. Figure 4 shows that, according to Aschoff’s rule, period length of all markers decreased with increasing light intensity and that the phyB-9 mutation did not eliminate this response. phyB-9 plants showed longer periods at most fluence rates tested. However, two classes of genes could be established based on the actual effect of the phyB-9 mutation on the period of their rhythmic transcription. CCA1:LUC+ and PRR9:LUC+ showed relatively large period differences (approximately 1 h) between Col-0 and phyB-9 over the entire fluence rate range (Fig. 4, B and C). In contrast, TOC1:LUC+ and GI:LUC+ displayed smaller, but significant, period differences (approximately 0.5 h), mainly at medium fluence rates (Fig. 4, D and E). Interestingly, in the case of both markers, Col-0 and phyB-9 plants showed identical periods at the lowest fluence rate tested (10 \( \text{\mu mol m}^{-2} \text{s}^{-1} \)). Moreover, the period of GI:LUC+ was not lengthened in the phyB-9 mutant at the highest fluence rate. TOC1:LUC+ showed longer, although less significant, periods in phyB-9 under these conditions. The period length of the circadian output marker CAB2:LUC+ showed characteristics similar to those of CCA1:LUC+ and PRR9:LUC+ (Fig. 4A). The phyB-9 mutation also affected the circadian phase of rhythmic expression of marker genes differently. PRR9:LUC+ and TOC1:LUC+ showed 1- to 2-h phase delays over the entire fluence rate range tested (Fig. 4, H and I). In contrast, CAB2:LUC+, CCA1:LUC+, and GI:LUC+ showed no significant phase changes in their expres-

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**Figure 4.** Analysis of the circadian phenotype of phyB-9 in continuous red light. Wild-type Col-0 (white squares) and phyB-9 (black squares) plants expressing CAB2:LUC+ (A and F), CCA1:LUC+ (B and G), PRR9:LUC+ (C and H), TOC1:LUC+ (D and I), or GI:LUC+ (E and J) were grown in 12-h-white-light/12-h-dark cycles for 7 d and transferred to different fluence rates of continuous red light. The free-running period length (A–E) and phase (F–J) of luminescence rhythms were determined and plotted as a function of fluence rate. Phase values were converted to circadian time (CT). Asterisks indicate statistically significant differences between Col-0 and phyB-9 as determined by Student’s two-tailed heteroscedastic t test: ** \( P < 0.01 \), * \( P < 0.05 \).
sion patterns at any fluence rate (Fig. 4, F, G, and J). These results indicate that PHYB provides different quality and/or quantity of input signals to the morning (CCA1, PRR9) and the evening (TOC1, GI) loops of the plant circadian oscillator in a fluence rate-dependent manner, which results in an apparent decoupling of the two loops at certain light intensities.

Complementation of the Circadian Period Phenotype of phyB-9 by PHYB Derivatives in Continuous Red Light

To test the ability of different PHYB fragments to restore the period phenotype of phyB-9, CCA1 and TOC1 mRNA rhythms were monitored by quantitative reverse transcription (qRT)-PCR assays in the different complemented lines. A medium fluence rate of continuous red light (50 μmol m⁻² s⁻¹) was used as a condition for free running, since all clock genes showed significant period lengthening in their transcription in this condition (Fig. 4, B–E). Tissue samples were harvested over a 72-h time course starting 60 h after the transfer to continuous red light. Figure 5 clearly shows that CCA1 and TOC1 expression had longer periods in phyB-9 in these conditions, which verifies the previous results (Fig. 4, B and D). The data also demonstrate that loss of PHYB function resulted in a lower amplitude of rhythmic CCA1 expression, but the amplitude of TOC1 expression was apparently unaffected. BFL, B651-NLS, and B450-NLS fully restored the period phenotype for both CCA1 and TOC1, but B651-NES and B410-NLS retained the phyB-9 phenotype. The ectopic expression of the functional PHYB fragments has not resulted in period lengths visibly shorter than that of the wild type. However, the amplitude of CCA1 and TOC1 expression was significantly increased in BFL plants (Fig. 5, A and F), whereas B651-NLS showed a similar increase in the amplitude of CCA1 expression only (Fig. 5B). The amplitude of rhythmic mRNA accumulation was unaffected by other PHYB derivatives. Since it is difficult and inaccurate to estimate period length from rhythmic mRNA data, the CCR2:LUC+ marker was introduced in the various backgrounds in order to obtain quantitative period length estimates in the complementing lines. Supplemental Figure S2 shows period estimates of CCR2:LUC+ expression from plants assayed in continuous red light at 50 μmol m⁻² s⁻¹. The period length of CCR2:LUC+ expression was approximately 1 h longer in phyB-9 compared with the wild type. Consistent with the mRNA data, BFL and B651-NLS complemented this phenotype, whereas B410-NLS did not. These results demonstrate that (1) N-terminal PHYB fragments containing the first 450 amino acids of PHYB are fully active in light signaling to

Figure 5. Complementation of the long-period phenotype of phyB-9 by PHYB fragments in continuous red light. Wild-type Col-0, phyB-9, and transgenic seedlings expressing BFL (A and F), B651-NLS (B and G), B651-NES (C and H), B450-NLS (D and I), or B410-NLS (E and J) in the phyB-9 background were grown in 12-h-white-light/12-h-dark cycles for 7 d and transferred to continuous red light at 50 μmol m⁻² s⁻¹ fluence rate. Samples were harvested at 3-h intervals starting at 60 h after the transfer for 3 d. CCA1 (A–E) and TOC1 (F–J) mRNA levels were determined by qRT-PCR. Data normalized to TUBULIN2/3 levels are shown. Col-0 (green lines) and phyB-9 (red lines) data are shown on each graph. Data from complementing lines are shown by blue lines.
the clock in continuous red light conditions, and (2) nuclear localization of PHYB is essential for this function.

**B651-NLS Mediates Resetting of the Clock by Red Light Pulses**

The circadian clock free running in darkness responds with characteristic phase shifts to light pulses. The magnitude and direction (delay or advance) of the phase shift depends on the actual phase of the clock at the time when the light pulse is given. Generally, light pulses applied during the first or second half of the subjective night trigger phase delays or advances, respectively (Covington et al., 2001). The role of PHYB in resetting the phase of leaf-movement rhythms in response to red light pulses has been suggested (Yanovsky et al., 2001). To test the function of PHYB fragments in this particular response, plants expressing CCR2:LUC+ in different genetic backgrounds were transferred to constant darkness at dusk (Zeitgeber time 12 [ZT12]) after 7 d of entrainment, and rhythmic luminescence was monitored. ZT0 is defined as the time of the last dark/light transition before transfer to constant conditions. A single red light pulse (100 μmol m⁻² s⁻¹ for 1 h) was applied to half of the plants after 29 h in constant dark, when the largest phase delays are expected (Kevei et al., 2007). Phase shifts were calculated by comparing the phase of CCR2:LUC+ expression in the induced plants with those of the noninduced control plants. Supplemental Figure S3 shows that wild-type plants produced approximately 6-h phase delays under these conditions, whereas the response was significantly reduced, but not eliminated, in the phyB-9 mutant. Similar to complementation of the long-period phenotype, BFL and B651-NLS were able to restore the wild-type response, but B410-NLS was not functional at all. These data suggest that N-terminal fragments of PHYB can mediate not only the effect of continuous light on the pace of the clock but also the effect of short light pulses on the phase of the clock.

**The Effect of PHYB on the Clock in Continuous White Light**

It has been demonstrated earlier that absence of PHYB function does not change the period of CAB2:LUC+ expression in continuous white light, which was explained by the contribution of all the other PHY and CRY photoreceptors to entrainment under this condition (Hall et al., 2002; Salome et al., 2002). These reports have also demonstrated that phyB-9 but not phyB-1 mutants show an early phase of CAB2:LUC+ expression in continuous white light. To investigate the role of PHYB in circadian entrainment in white light, the period and phase of core clock gene expression were analyzed in Col-0 and phyB-9 plants transferred to continuous white light. Surprisingly, with the exception of PRR9, transcription of all clock genes showed a short-period phenotype in phyB-9 (Fig. 6A).

The phase of CCA1:LUC+ and TOC1:LUC+ remained unchanged, but GI:LUC+ and PRR9:LUC+ showed an advanced phase of expression in phyB-9 (Fig. 6B). CAB2:LUC+ showed no period change but a slight early-phase phenotype in phyB-9, which is consistent with earlier results (Salome et al., 2002). In contrast, expression of another clock output marker, CCR2:LUC+, displayed significant period shortening and slight phase advance in the mutant plants. These data dem-

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**Figure 6.** Circadian phenotypes of phyB-9 in continuous white light. Seedlings expressing different luciferase reporters and/or N-terminal derivatives of PHYB as indicated were grown in 12-h-white-light/12-h-dark cycles for 7 d and transferred to continuous white light at 60 μmol m⁻² s⁻¹ fluence rate. A, Period estimates for CCA1:LUC+, TOC1:LUC+, GI:LUC+, PRR9:LUC+, CAB2:LUC+, and CCR2:LUC+ expression in Col-0 (black bars) and phyB-9 mutant (white bars) seedlings. B, Phase estimates for CCA1:LUC+, TOC1:LUC+, GI:LUC+, PRR9:LUC+, CAB2:LUC+, and CCR2:LUC+ expression in Col-0 (black bars) and phyB-9 mutant (white bars) seedlings. C, Period estimates for CCR2:LUC+ expression in Col-0, phyB-9, BFL, B651-NLS, and B410-NLS seedlings. Asterisks indicate statistically significant differences between Col-0 and phyB-9 as determined by Student’s two-tailed heteroscedastic t test: **P < 0.01, *P < 0.05.
onstrate that the absence of PHYB function has markedly different effects on the circadian oscillator in white and red light conditions, which suggests a complex interaction between the red and blue light input pathways to the circadian clock. To identify the particular domain of PHYB responsible for this functional interaction, CCR2:LUC+ rhythms were monitored in transgenic phyB-9 lines expressing different truncated derivatives of PHYB. The data presented in Figure 6C show that only the full-length PHYB, but none of the N-terminal derivatives, was able to restore the wild-type period of CCR2:LUC+. These results indicate that, in contrast to red light, the C-terminal domain of PHYB is required for normal entrainment in white light conditions.

DISCUSSION

PHYB is the dominant red/far-red light-absorbing photoreceptor in light-grown Arabidopsis plants. PHYB provides signals to regulate photomorphogenesis, to control flowering, and to entrain the circadian clock. Consequently, phyB mutants display long hypocotyls, accelerated flowering, and long periods of circadian rhythms under specific light conditions (Reed et al., 1993; Somers et al., 1998; Endo et al., 2005). The PHYB receptor folds into two main domains: the chromophore-bearing photosensory N-terminal domain and the C-terminal domain containing a His kinase-like subdomain and protein motifs for dimerization and nuclear translocation. It has been demonstrated that dimers of the N-terminal domain of PHYB possess full PHYB function regarding photomorphogenesis and flowering time determination if they are targeted to the nucleus by added foreign NLS motifs (Matsushima et al., 2003). In other words, the His kinase domain is dispensable for these functions of PHYB, and the main role of the C-terminal part is to provide a platform for dimerization and to control the entry of the protein in the nucleus.

To test the function of the N-terminal domain to provide light signaling to the circadian clock, we expressed the 651-amino acid N-terminal fragment fused to YFP, dimerization domains, and NLS or NES protein motifs (B651-NLS and B651-NES, respectively) in the phyB-9 background, which lacks any PHYB functions (Reed et al., 1993). The fusion proteins, including the YFP-tagged full-length PHYB control (BFL), were expressed at comparable levels (Fig. 1B) and showed the expected subcellular localization (Fig. 2; Supplemental Fig. S1). The light input pathway to the circadian oscillator mediates two modes of entrainment depending on the duration of illumination. In constant light conditions, the activity of the input pathway shortens the free-running period length with increasing light intensity (parametric entrainment). In constant darkness, however, short light pulses elicit discrete phase advances or delays of the oscillator and overt circadian rhythms (nonparametric entrainment; Devlin and Kay, 2001). Red light signals absorbed by PHYB contribute to both types of entrainment (Somers et al., 1998; Yanovsky et al., 2001). Figures 4 and 5 show that rhythmic expression of core oscillator genes has a longer period in phyB-9 plants compared with the wild type in continuous red light at 50 μmol m⁻² s⁻¹ fluence rate. Supplemental Figure S3 shows that in the absence of PHYB function, red light pulses trigger smaller phase shifts of the oscillator. Our results demonstrated that B651-NLS was as effective at restoring these phenotypes of the phyB-9 mutant as the full-length PHYB. Interestingly, BFL (and B651-NLS) showed significant increases in the amplitude of mRNA rhythm of CCA1 but did not cause shorter periods. It has been reported that overexpression of PHYB results in a short period of CAB1:LUC expression in continuous red light (Somers et al., 1998; Hall et al., 2002) and in early flowering under both short-day and long-day conditions (Bagnall et al., 1995). The lack of circadian and flowering-time phenotypes of BFL can be explained by similar PHYB levels in BFL and wild-type plants, in contrast to lines used in the cited studies, which show at least 15-fold overexpression of PHYB protein. In contrast to BFL and B651-NLS, B651-NES was unable to restore PHYB function to regulate rhythmic CCA1 or TOC1 expression (Fig. 5, C and H). These results demonstrate that the N-terminal domain of PHYB is able to mediate all aspects of entrainment of the circadian clock, and this function requires nuclear localization of PHYB. The data presented here also verify previous reports, since B651-NLS but not B651-NES restored the photomorphogenic and flowering phenotypes of phyB-9.

The molecular mechanism by which nucleus-localized PHYB mediates entrainment is unclear, but most likely it involves interaction of PHYB with regulatory proteins, which indirectly or directly affect the oscillator. Among such clock-affecting proteins, ZEITLUPE (ZTL) and EARLY FLOWERING3 (ELF3) have been shown to interact with PHYB in yeast or in vitro (Jarillo et al., 2001; Liu et al., 2001). ZTL is an F-box protein that targets TOC1 proteins for ubiquitination by Skp1/Cullin/F-box-type E3 ubiquitin ligases and for subsequent degradation (Mas et al., 2003) and severely affects the free-running period length (Somers et al., 2000; Kevei et al., 2006). ELF3 is a clock-controlled nuclear protein that attenuates light-induced resetting of the clock (Covington et al., 2001). However, both ZTL and ELF3 bind to the C-terminal domain of PHYB, which is dispensable for circadian function in red light according to our data; therefore, these interactions have no relevance to entrainment under these conditions.

In fact, PIF3 has been shown to interact with the N-terminal domain of PHYB Pfr (Shimizu-Sato et al., 2002) and also with TOC1 (Yamashino et al., 2003). Since PIF3 potentially binds to the G-box element located in the promoter of CCA1 and LHY genes, the PHYB-PIF3-TOC1 complex could provide a mechanism for directing regulatory light signals to certain
core oscillator genes (Martinez-Garcia et al., 2000). It has been shown that PIF1, PIF4, and PIF5 also interact with TOC1 (Yamashino et al., 2003). However, misexpression of PIF3 or PIF5 does not affect entrainment of the plant circadian clock (Fujimori et al., 2004; Viczian et al., 2005). Although the lack of such phenotypes could be explained by possible redundant action of several PIF3-like transcription factors, PIFs are probably not the terminal components of PHYB-mediated red light input to the clock but rather represent components of clock-controlled output processes (e.g. rhythmic hypocotyl elongation; Nozue et al., 2007).

Oka et al. (2004) reported that a short fragment containing 450 amino acids of the N-terminal domain of PHYB mediates photomorphogenic responses in the nucleus. We created phyB-9 plants expressing this PHYB derivative in fusion with dimerization and NLS signals (B450-NLS). Analysis of these transgenic lines verified that the PHY subdomain of the N-terminal part of PHYB is not required for the inhibition of hypocotyl elongation (Fig. 3A) and revealed that this subdomain is also dispensable for the regulation of period length. Figure 5, D and I, demonstrates that B450-NLS rescued the wild-type expression pattern of CCA1 and TOC1. Unlike the photomorphogenic and circadian phenotypes, the early-flowering phenotype of phyB-9 was not restored by B450-NLS. Figure 3B shows that B450-NLS plants grown in long days (16 h of light/8 h of dark) flowered at the same time as phyB-9 mutants. This result is consistent with those reported earlier (Oka et al., 2004). This observation indicates that the PHY subdomain may provide a binding site for a yet unidentified factor that is necessary to transmit signals eventually destabilizing CO protein and delaying flowering. The facts that PHYB acts on CO protein levels and that the flowering phenotype of phyB mutants is independent of photoperiod demonstrate that the early flowering of phyB mutants does not arise from altered clock function. Our results further support this conclusion, because B450-NLS plants generate wild-type circadian rhythms yet display the early-flowering phenotype of phyB-9.

To define the minimal N-terminal PHYB fragment, which is still functional in the nucleus, we created transgenic plants expressing a 410-amino acid derivative of PHYB (B410-NLS). Despite the fact that this version is only 40 amino acids shorter than B450-NLS, B410-NLS was unable to complement any phenotype of phyB-9 in any assays. It must be noted that the effect of both B651-NLS and B450-NLS was light dependent, and phenotypes indicating constitutive light-independent signaling were not observed in these plants in darkness (Matsushita et al., 2003; Oka et al., 2004; data not shown). This means that light-induced conformational change is a prerequisite of the biological function of these derivatives. To undergo light-induced Pr ↔ Pfr conversions, the chromophore must be attached to the PHYB apoprotein. Zinc blots were used to test the chromophore-binding capability of the different PHYB derivatives (Oka et al., 2004). Supplemental Figure S4 demonstrates that B651 and B450 were able to autoligate the chromophore. In contrast, B410 failed to incorporate the chromophore, indicating that the deletion probably affected the function of the bilin lyase domain.

Quantitative analysis of clock gene expression in the phyb-9 mutant revealed an unexpected complexity in the function of the light input to the circadian oscillator. The plant oscillator consists of three interlocked transcriptional/translational feedback loops. The “morning” loop is operated by interactions among CCA1/LHY and PRR7/9 genes, the “evening” loop is based on the cross-regulation between TOC1 and Y/GI, and the two loops are coupled by a central loop formed by CCA1/LHY, TOC1, and a yet unidentified X factor. In wild-type plants, these loops are coupled together, which is illustrated by stable and constant phase relationships of expression of these genes under free-running conditions. Theoretically, the morning and evening loops could be uncoupled from each other in certain conditions, for example, as a result of the misfunction of X. Data in Figure 4 demonstrate that loss of PHYB function affects the pace of the morning and evening loops differently. Expression of CCA1 and PRR9, components of the morning loop, shows significant long-period rhythm in phyb-9 over a wide range of fluence rates of continuous red light, but TOC1 and GI expression displays less significant period lengthening. Importantly, at lower fluences of red light, the pace of the evening loop is not affected by the lack of phyb-9, whereas the morning loop runs at a lower pace in the same conditions. These data demonstrate that, in these conditions, the coupling between the morning and evening loops has been weakened in phyb-9. Although the exact molecular mechanism of this phenomenon and the role of PHYB in the apparent decoupling remain unclear, our results shed light on the complexity of the function of the light input to the plant circadian oscillator. This complexity was further expanded by the analysis of circadian gene expression in the phyb-9 mutant in continuous white light (Fig. 6). Our data showed that PHYB has nearly opposite effects on the pace of the clock in red and white light, since circadian period lengths in the phyb-9 mutant are generally lengthened or shortened in continuous red or white light conditions, respectively. On the other hand, it has been demonstrated that the phyb-1 mutation has no effect on period length in monochromatic blue light (Devlin and Kay, 2000). The cool-white fluorescent tubes used in this experiment provide light enriched in blue light but still emit red light capable of PHYB activation. This is indicated by the long hypocotyls of phyb-9 seedlings under this condition (data not shown). It follows that the short-period phenotype phyb-9 is apparent upon simultaneous activation of the red and blue light signaling pathways to the clock, which could be explained by a negative effect of PHYB on blue light signaling. The unexpected short-period phenotype in continuous white light
has been observed for leaf-movement rhythms in the quintuple phy mutant (Strasser et al., 2010), also suggesting that blue light input to the clock is attenuated by the action of phytochrome(s). Since PHYB appears to have a positive effect on CRY2 signaling (Mas et al., 2000), other routes of blue light input may be affected. However, molecular details of this particular functional interaction between red and blue light signaling to the clock remain to be elucidated. We showed that BFL, but not B651-NLS, was able to complement the short-period phenotype of phyB-9 in white light (Fig. 6C). Since the BFL and B651-NLS proteins were expressed at similar levels (Fig. 1B), the lack of complementation by B651-NLS is most probably due to absence of the C-terminal domain of PHYB.

In conclusion, our data suggest that the nucleus-localized N-terminal domain of PHYB is fully functional in regulating photomorphogenesis, flowering time, and red light-dependent entrainment of the circadian clock. However, the C-terminal domain is essential for proper circadian entrainment in white light conditions. To explain the requirement of the C-terminal domain for this process, we propose that this domain mediates integration of the blue and red light signaling pathways to the clock.

MATERIALS AND METHODS

Plant Materials and Growth Conditions

Wild-type and phyB-9 mutant Arabidopsis (Arabidopsis thaliana) plants in the Col-0 background were used in this study. Surface-sterilized seeds were grown in 12-h-white-light (50 μmol m⁻² s⁻¹) / 12-h-dark cycles at 22°C (MLR-350; Sanyo) for 7 d before being transferred to continuous darkness at ZT12 or to continuous light at ZT0, and all measurements were carried out at constant 22°C. ZT0 is defined as the time of the last dark/light transition before transfer to constant conditions. Illumination was provided by cool-white fluorescent tubes or monochromatic light-emitting diode red light sources (max = 667 nm; Quantum Devices).

Construction of Transgenic Plants

The CAR2:LUc+, CCR2:LUc+, CCA1:LUc+, and TOC1:LUc+ constructs have been described (Toth et al., 2001; Doyle et al., 2002; Locke et al., 2006). GI and PR9 promoter fragments were obtained by PCR performed on genomic DNA of wild-type plants. Unique restriction sites were designed at the 5′ and 3′ ends of the promoter fragments to facilitate cloning in the pPCV-LUC+ binary vector (Toth et al., 2001). The amplified fragments contained the entire 5′ untranslated region but not the ATG of the corresponding genes. Fragment lengths and restriction sites at the 5′ and 3′ ends were as follows: GI, 2,749 bp; XbaI-BamHI; PR9, 1,330 bp; SalI-BamHI.

DNA fragments encoding the SV40 NLS motif (Kalderon et al., 1984) and the dimerization domain of CPRF4 (Kircher et al., 1998) were the same as described (Bevan et al., 1990). The DNA fragment encoding the NES motif (LALKLAGLDINKTGG) from the heat-stable inhibitor (PKI; Wen et al., 2000) and the dimerization domain of CPRF4 (Kircher et al., 1998) were the same as described previously (Toth et al., 2001; Doyle et al., 2002; Locke et al., 2006). To select transformants, seeds were selected on Murashige and Skoog medium supplemented with 15 μg mL⁻¹ hygromycin. Ten to 15 independent transformants for each construct were self-fertilized, and individuals of the T2 or the homozygous T3 progeny were used for luminescence or complementation assays, respectively. To create lines coexpressing the different PHYB derivatives and the CCR2:LUc+ marker, the CCR2:LUc+ construct was transferred in the pPCV812 vector, which carries a BASTA resistance marker (Bauer et al., 2004). Selected homozygous lines expressing different N-terminal fragments of PHYB were transformed with the CCR2:LUc+ pPCV812 construct, and transformants were selected based on their resistance to BASTA herbicide.

Analysis of Luminescence Rhythms

Luciferase activity was determined by measuring single seedlings with an automated luminometer (TopCount NXT; Perkin-Elmer) for 2 to 7 d as described previously (Kevei et al., 2006). For fluence rate curves, circadian periods of luminescence rhythms were measured in seedlings transferred to constant illumination of red or white light at the fluence rates indicated. All rhythm data were analyzed with the Biological Rhythms Analysis Software System (available at http://www.amillaro.org) running fast Fourier-transform nonlinear least-squares estimation. Variance-weighted mean periods within the circadian range (15–40 h) and si values were estimated as described from 10 to 36 traces per genotype. Phase values were determined as the time of the first full peak of luminescence rhythms in continuous light. Phase values were normalized to free-running period length and are shown as circadian time (Salome et al., 2002). Statistical analysis was done using Student’s two-tailed heteroscedastic t test. Experiments were repeated three or four times, and representative data sets are shown. For each genotype, at least three independent transgenic lines were analyzed.

For phase-shift experiments (Supplemental Fig. S3), plants expressing CCR2:LUc+ were grown in 12-h-white-light/12-h-dark cycles for 7 d and transferred to darkness at the last light-to-dark transition. CCR2:LUc+ luminescence was monitored at 1-h intervals for 5 d. After 29 h in constant dark, half of the plants were treated with a 1-h red light pulse at 100 μmol m⁻² s⁻¹ fluence rate. Phase values were determined as the time of the second peak after the light pulse and were normalized to the free-running period as above. Light-induced phase shifts were calculated by comparing the phase of CCR2:LUc+ rhythm in the induced and noninduced plants.

Analysis of Gene Expression

Total RNA extraction, cDNA preparation, and qRT-PCR assays were carried out as described (Kevei et al., 2007). qRT-PCR primers have been described (Edwards et al., 2005). All graphs show mRNA levels relative to the TUBULIN2/3 mRNA transcript (Endo et al., 2007). Total protein extraction, western-blot analysis, and detection of YFP fusion proteins were done essentially as described (Kevei et al., 2007). The assays were repeated two or three times, and representative data are shown.

Chromophore-Binding Assay (Zinc Blot)

For in vitro reconstitution, 4 μM phycocyanobilin as chromophore was incubated with purified recombinant GST-B651, GST-B450, or GST-B410 protein as described (Lagaris and Lagarias, 1989). Each apoprotein was expressed in Escherichia coli and purified on a glutathione-Sepharose 4B column. Zn²⁺-induced fluorescence of chromophore-bound proteins was visualized under UV light after SDS-PAGE separation according to Oka et al. (2004).

Measurement of the Hypocotyl Length

For hypocotyl length measurements, seeds were sown on wet filter paper and incubated in the dark for 48 h at 4°C. Cold-treated seeds were then irradiated with 6 h of white light and transferred to 22°C and dark for an additional 18 h. After this treatment, seedlings were grown at different fluence rates of light for 4 d. Measurement of the hypocotyl length was performed using MetaMorph Software (Universal Imaging). Hypocotyl lengths of light-grown seedlings were normalized to the corresponding dark-grown hypocotyl length. Fluence rate curves for hypocotyl elongation were obtained by plotting relative hypocotyl lengths against the light intensities used in the experiment displayed on a logarithmic scale. Subcellular distribution of YFP fusion proteins was analyzed in 4-d-old seedlings as described previously (Bauer et al., 2004).
Epifluorescence and Confocal Laser Scanning 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, 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.

For epifluorescence and light microscopy, seedlings were transferred to glass slides under dim-green safelight and analyzed with an Axioskop microscope (Zeiss). Excitation and detection of the YFP fluorophore were performed with YFP filter set (AHF Analysentechnik). Each experiment was repeated at least two times using at least five seedlings by analyzing the upper third of the hypocotyl. Representative cells were documented by photography with a digital Axioscam camera system (Zeiss). Documentation of cells was performed during the first 60 s of microscopic analysis to prevent the impact of microscopic light on localization patterns. Photographs were processed for optimal presentation using the Photoshop 7.0 (Adobe Systems Europe) and MS Office (Microsoft) software packages.

For confocal laser scanning microscopy, seedlings were transferred to glass slides under dim-green safelight and analyzed with an Axioskop microscope (Zeiss). Excitation and detection of the YFP fluorophore were performed with YFP filter set equipped with the META device (Zeiss).

Measurement of Flowering Time

Seeds were sown on soil and incubated for 2 d in darkness at 4°C. They were subsequently transferred to long-day (16-h-white-light/8-h-dark) conditions. Light sources were fluorescent (cool-white) tubes producing a fluence rate of 60 μmol m−2 s−1. Flowering time was recorded as the number of rosette leaves at the time when inflorescences reached 1 cm in height. The experiment was repeated twice using 30 to 40 plants per genotype.

Supplemental Data

The following materials are available in the online version of this article.

Supplemental Figure S1. Subcellular localization of PHYB fusion proteins.

Supplemental Figure S2. Quantitative analysis of complementation of the long-period phenotype of phyB−9.

Supplemental Figure S3. The 651-amino acid N-terminal fragment of PHYB is sufficient to mediate red light-induced resetting of the circadian clock.

Supplemental Figure S4. Chromophore does not incorporate into the recombinant protein GST-B410.

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LITERATURE CITED


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