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Review

# Coordination of Cryptochrome and Phytochrome Signals in the Regulation of Plant Light Responses

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**Abstract:** In nature, plants integrate a wide range of light signals from solar radiation to adapt to the surrounding light environment, and these light signals also regulate a variety of important agronomic traits. Blue light-sensing cryptochrome (cry) and red/far-red light-sensing phytochrome (phy) play critical roles in regulating light-mediated physiological responses via the regulated transcriptional network. Accumulating evidence in the model plant *Arabidopsis* has revealed that crys and phys share two mechanistically distinct pathways to coordinately regulate transcriptional changes in response to light. First, crys and phys promote the accumulation of transcription factors that regulate photomorphogenesis, such as HY5 and HFR1, via the inactivation of the CONSTITUTIVE PHOTOMORPHOGENIC1/SUPPRESSOR OF PHYA-105 E3 ligase complex by light-dependent binding. Second, photoactive crys and phys directly interact with PHYTOCHROME INTERACTING FACTOR transcription factor family proteins to regulate transcriptional activity. The coordinated regulation of these two pathways (and others) by crys and phys allow plants to respond with plasticity to fluctuating light environments in nature.

**Keywords:** phytochrome; cryptochrome; de-etiolation; photoperiodic flowering; gene expression

## 1. Introduction

Because plant life relies on solar radiation as an energy source, plants have acquired a light-sensing mechanism to maximize the availability of light for photosynthesis [1,2]. Specifically, germination does not occur in deep subterranean darkness and occurs only when the plant senses a low amount of light under the ground at depths close to the surface of the soil. The post-germinative seedlings in the soil elongate the hypocotyl with a closed hook, and yellowish cotyledons move toward the surface of soil. Upon reaching the surface and absorbing light, seedlings start photomorphogenic development, in which hypocotyl elongation ceases, the hook and cotyledons start to open, and chloroplasts begin developing to maximize light capture and autotrophic growth. This developmental transition from a dark-grown seedling to a light-grown seedling is called de-etiolation. Even aboveground plants must sense the presence and proximity of neighboring plants and compete with them for solar energy through a response called Shade Avoidance Syndrome (SAS). Plants detect shade from neighboring plants as alterations in the light quantity and quality, and, in response, they elongate the hypocotyl, petiole, and internode to bring the photosynthetic organs to more favorable conditions for

photosynthesis. Plants also perceive the day length to control the transition from the vegetative phase to reproductive phase.

As exemplified above, light as an environmental signal regulates plant development throughout the plant life cycle. The blue (B) and red (R) light regions of the light spectrum are most efficiently utilized for the photosynthesis; thus, B light-sensing cryptochrome (cry) and R light-absorbing phytochrome (phy) play major roles in regulating plant light responses, such as light-dependent seed germination, de-etiolation, SAS and photoperiodic flowering [3–5]. Importantly, crys or phys also mediate important agronomic traits in crop species [4–6]. For example, rice mutants defective in all phy proteins set very few seeds [7]. Reduced expression of *CRY1a* and *CRY1b* in barley impairs seed dormancy [8]. A decreased level of *CRY2a* expression accelerates senescence in soybean [9]. All these examples indicate the importance of cry and phy action in regulating the growth and quality of crops. Great progress has been made in understanding the action mechanisms of crys and phys in the model plant *Arabidopsis* [5,10,11]. In this article, we review the mechanisms of action of cry and phy in *Arabidopsis* and discuss how cry and phy coordinate to regulate light responses. The findings on the coordinated action of crys and phys in *Arabidopsis* will provide insights to improve the yield and quality of crop species.

## 2. Physiological Functions and Actions of Phytochrome

Phys are a unique type of photoreceptor that display photoreversible conformational changes between two spectrally distinct forms: R light-absorbing Pr and far-red (FR) light-absorbing Pfr. R light transforms Pr into biologically active Pfr to induce phy-mediated responses, such as seed germination, de-etiolation, etc. Conversely, FR light inactivates phys by converting Pfr back into Pr. Phy proteins are synthesized as Pr and remain in the cytosol in the dark. Upon light absorption, phy proteins become Pfr and translocate into the nucleus, where phys regulate the transcription of a number of genes that mediate plant light responses [12,13]. Interestingly, phy proteins form granular structures called photobodies (originally called nuclear speckles) in the nucleus in response to light. Although photobody formation by phy has been shown to require HEMERA protein and the photobody is a presumed site for protein degradation [14,15], the functions of this structure are not fully understood.

The *Arabidopsis* genome encodes five phys (phyA–phyE) [16]. Of these, phyA and phyB play the most important roles. Accordingly, these two phys are found in all angiosperm species whose genomes have been sequenced to date, including crops such as rice and soybean [7,17,18]. Analyses of *phyA* and *phyB* mutants revealed that phyA and phyB regulate related responses, such as seed germination and de-etiolation, via different modes of action [3]. phyB is a major photoreceptor that mediates low light fluence-induced R/FR-reversible seed germination and hypocotyl elongation inhibition under continuous R light [19,20], whereas phyA is the sole photoreceptor mediating notably sensitive low light fluence-induced R/FR-irreversible seed germination and hypocotyl elongation inhibition under continuous FR light [20–22]. Moreover, because of the partially overlapping absorption spectra of Pr and Pfr, an equilibrium is established between Pr and Pfr even under continuous monochromatic light. Importantly, FR light is capable of establishing a 1% Pfr/ $P_{\text{total}}$  ratio at a photoequilibrium state, whereas R light establishes an 80% Pfr/ $P_{\text{total}}$  ratio [19]. Therefore, both phyA-mediated responses require a low amount of Pfr (<0.1% Pfr/ $P_{\text{total}}$  for seed germination and 1% Pfr/ $P_{\text{total}}$  at photoequilibrium for FR-induced inhibition of hypocotyl elongation), suggesting that phyA is a highly sensitive phy species.

The sensitive phyA-mediated responses are presumably enabled by the high levels of phyA in etiolated seedlings [23]. However, the overexpression of phyB at a similar level to phyA was capable of complementing the phyB-mediated hypocotyl response under R light but not the phyA-mediated hypocotyl response under FR light in a *phyAphyB* double mutant [24–26], suggesting that phyA acquires specific functions necessary for sensitive responses. The nuclear localization of phyA and phyB is required for FR- and R-mediated inhibition of hypocotyl elongation, respectively [27–30]. Accordingly, a sensitive response can be observed in the nuclear accumulation of phyA but not phyB under FR [31]. This FR-induced nuclear accumulation of phyA is mediated by two cargo proteins,

FAR-RED ELONGATED HYPOCOTYL1 (FHY1) and its homolog FHY1-LIKE, which specifically bind with the Pfr form of phyA but not that of phyB [30,32–34]. However, this sensitive nuclear translocation mechanism is not sufficient to explain the observed difference in sensitivity between phyA and phyB because nuclear-targeted phyB does not mediate de-etiolation under FR [24,28]. Therefore, phyA must have a specific unidentified mechanism by which it exerts a sensitized response in the nucleus. In contrast, upon the light exposure that transforms Pr into Pfr, the phyA-mediated response is attenuated by rapidly removing the Pfr of phyA through proteasome-mediated degradation [35–37], whereas phyB remains relatively stable under such conditions [38]. This desensitization mechanism is at least partially mediated by CONSTITUTIVE PHOTOMORPHOGENIC1 (COP1), which constitutes the E3 ubiquitin ligase subunit [39].

The light-labile property of phyA restricts the action of phyA to seedling development that occurs under subterranean darkness to the light transition or under dense canopies in the natural environment [3,40]. Because plant tissue absorbs R light but reflects or transmits FR light, a plant canopy provides low R/FR conditions. Under a dense canopy where the R/FR ratio is low enough to inactivate phyB and stabilize phyA, sensitive phyA functions antagonize the loss of phyB activity, thereby resulting in modest hypocotyl elongation [40]. Modest phyA-mediated hypocotyl elongation may prevent detrimental excessive elongation under such conditions in nature. The vital importance of phyA is highlighted under a dense canopy [41]. However, an intermediate R/FR ratio, which is encountered by proximal neighbor plants, degrades phyA and shifts the photoequilibrium of phyB toward reducing the ratio of active Pfr so that plants elongate the hypocotyl robustly to compete with their neighboring plants [40]. This response is known as low R/FR-induced SAS [42,43]. Low R/FR also accelerates petiole and internode elongation by inactivating phyB. In the high R/FR condition, where the plant density is low in an open habitat, phyA is degraded and phyB plays critical roles in regulating seedling development. In such conditions, phyB also regulates a wide range of responses, such as the inhibition of the petiole and internode elongation, promotion of stomatal development and determination of flowering time [3]. It should be noted here that the phyA level is not zero, and, even in such conditions, it maintains certain functions, albeit marginal compared with those of phyB.

### 3. Physiological Functions and Actions of Cryptochrome

*Arabidopsis* has two members of the cry family that have B light photoreceptor functions: cry1 and cry2 [5,44,45]. The other angiosperm species whose genomes have been sequenced to date also have at least two *CRY* genes each [4,46]. Both crys undergo B light-dependent phosphorylation, and the phosphorylation of the crys is closely correlated with their functions [47–49]. Homodimerization is required for the phosphorylation and physiological activity of the crys [50–52]. Recently, it was found that cry2 forms a homodimer in response to B light [53]. Thus, these results indicate that B light-dependent homodimerization is the elementary process of cry2 photoactivation. The B light inhibitors of cryptochromes (BIC) 1 and 2 were identified as negative regulators of cry1 and cry2, and they interact with cry2 to suppress the B light-dependent dimerization, phosphorylation, and physiological activities. Therefore, BIC1 and BIC2 constitute the desensitizing mechanism of both crys to sustain the homeostasis of physiologically active cry [53]. However, whether BICs inhibit cry1 dimer formation remains to be tested. In addition to this, cry2 activity is attenuated by B light-dependent rapid protein degradation, at least in part via a COP1-mediated pathway [54–57].

cry2 localizes exclusively in the nucleus [55]. Although cry1 localizes equally in the nucleus and cytoplasm [58,59], the nuclear-targeted cry1 regulates most cry1-mediated responses [58]. In addition, thousands of genes are regulated by both cry1 and cry2 in response to B light during the de-etiolation process [53,60–63]. Thus, it is generally accepted that photoactivated cry1 and cry2 transduce light signals to downstream components, primarily in the nucleus. Different from cry1 but similar to phys, photoactivated cry2 forms photobodies in the nucleus [56,64,65]. Photobody formation is closely correlated with cry2 function and degradation. However, the function and mechanism of cry2 photobody formation are not fully understood, which is similar to the case of phy photobody

formation. Interestingly, cry2 photobodies partly overlap with phyB photobodies, suggesting that these photobodies are the site for cry2–phyB interactions [65].

Mutant analyses of *cry1* and *cry2* mutants have revealed that *cry1* is the primary photoreceptor mediating the B-induced inhibition of hypocotyl elongation [44]. *cry1* mediates de-etiolation in a wide range of B light intensities. However, *cry2* regulates the hypocotyl response preferentially under weak B light because of the B light fluence-rate-dependent degradation [54]. However, compared with phyA, *cry2* does not show a sensitive light response, although both phyA and *cry2* are rapidly degraded by light exposure.

*Arabidopsis* is a long-day (LD) plant that flower under LD conditions. *cry2* mutants exhibit a robust late-flowering phenotype only in LD conditions [45], whereas *cry1* mutants do not show a late-flowering phenotype [66–68], indicating that *cry2* is the predominant B light receptor regulating photoperiodic flowering. However, certain experiments have shown that *cry1* single mutants flower late under LD conditions, although their late-flowering phenotype is not as obvious as that of *cry2* mutants [69,70]. Two dominant *cry1* alleles harboring either L407F or G389R flower early in LD conditions [71,72]. Moreover, compared with any of the *cry1* or *cry2* single mutants, the *cry1cry2* double mutant flowered later than the wild type in continuous B light [73], suggesting that *cry1* acts redundantly with *cry2* to promote flowering in this condition. Thus, the evidence suggests that *cry1* also partially functions in the regulation of photoperiodic flowering time. Therefore, *cry1* and *cry2* have their specific roles in the regulation of hypocotyl elongation and photoperiodic flowering, and they partially redundantly regulate these responses.

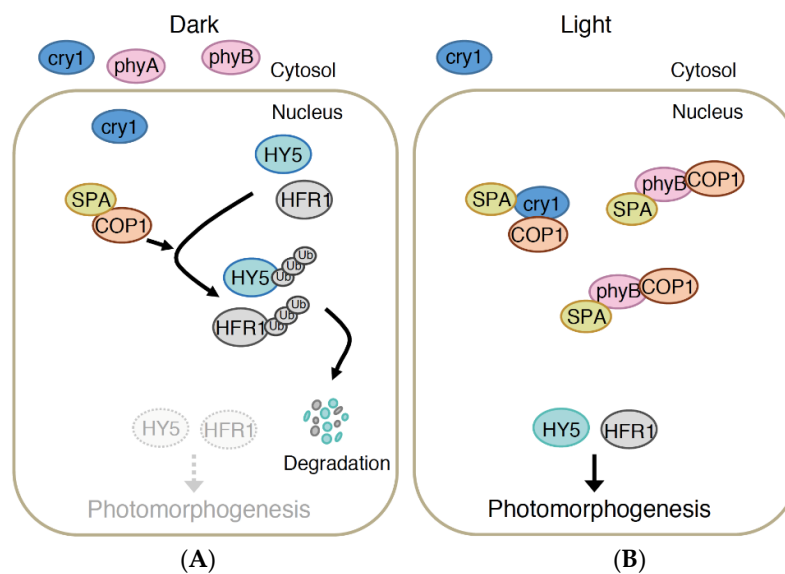
Although B light-mediated inhibition of hypocotyl elongation and photoperiodic flowering are two major phenotypes analyzed as functions of cry, the protein also mediates several other B light responses [4,5]. For example, recent analyses have highlighted the functions of cry in the control of SAS [74–76]. Shading by neighboring plants attenuates R light as well as B light. Simply reducing B light induces similar morphological changes to low R/FR-induced SAS, including the elongation of the hypocotyl, petiole, and internode in *Arabidopsis*. This low B light (LBL)-induced SAS has been shown to be regulated by crys [74–76]. *cry1cry2* double mutant seedlings display a significantly reduced response to LBL in the regulation of hypocotyl elongation, whereas both *cry1* and *cry2* mutant seedlings are still responsive to LBL to a large extent, suggesting that both *cry1* and *cry2* regulate the effect of LBL-induced SAS on hypocotyl elongation. However, the *cry1* mutant elongates the petiole irrespective of the unshaded or LBL condition, whereas the *cry2* mutant remained responsive to LBL. Thus, *cry1* functions become more dominant in controlling LBL-induced SAS in adult plants.

#### 4. Signal Transduction Pathways of Cryptochromes and Phytochromes

Crys and phys mediate related responses through regulated transcription, although they work under differing wavelengths of light. Accordingly, recent analyses have revealed that crys and phys share two common signaling pathways, and they both bind with the COP1/SUPPRESSOR OF PHYA-105 (SPA) complex, which targets certain sets of transcription factors for degradation [77–81], and a subset of bHLH transcription factors, PHYTOCHROME INTERACTING FACTORS (PIFs) [76,82,83], to control transcription. In addition to these common signaling partners, *cry2* also binds with CRYPTOCHROME-INTERACTING BASIC HELIX-LOOP-HELIX (CIB) proteins, another set of bHLH transcription factors that directly bind to the promoter of the flowering inducer gene *FLOWERING LOCUS T* (*FT*) and promote its transcription [84,85]. Phys also interact with other potential signaling partners, such as PHYTOCHROME KINASE SUBSTRATE (PKS) family proteins [86]. However, PKS family proteins appear to function more in linking phy signals to phototropic responses, which are regulated by another set of B light photoreceptors, phototropins, rather than cry-mediated responses [87,88]. Therefore, we have omitted these proteins from the discussion in this article.

#### 4.1. COP1/SPA Pathway

The *cop1* mutant displays a constitutively photomorphogenic (*cop*) phenotype, in which dark-grown *cop1* mutant seedlings mimic light-grown wild-type seedlings [89]. The *spa1* mutant was initially identified as the suppressor mutant of *phyA-105*, a weak *phyA* allele that promotes a long hypocotyl under continuous FR light [90,91]; however, subsequent evidence showed that a higher-order combination of *spa1* and mutants of other SPA genes (SPA2–4), e.g., *spa1spa2spa3spa4*, displayed strong *cop*-like phenotypes [92,93]. This genetic evidence indicates that COP1 and SPAs are the negative regulators of photomorphogenesis. Compared with the *cop1* and *spa* mutants, a mutant of *ELONGATED HYPOCOTYL 5 (HY5)* was identified as a long-hypocotyl mutant under white light but not in the dark, thus indicating that the *HY5* gene encodes the positive regulator of photomorphogenesis [89,94]. Indeed, *HY5* has been shown to act as the bZIP transcription factor regulating the transcription of a number of light-responsive genes [95,96]. Consistent with these mutant phenotypes, COP1 and SPA proteins together with CULLIN4 (CUL4) constitute the E3 ubiquitin ligase complex, which ubiquitinates *HY5* proteins for degradation through the 26S proteasome pathway to suppress photomorphogenesis in the dark [97] (Figure 1). Upon light exposure, *cry*s and *phys* inactivate the COP1/SPA complex, thereby leading to the accumulation of *HY5* protein in the nucleus to induce photomorphogenesis [89,94]. The COP1/SPA complex targets other positive regulators of photomorphogenesis, such as LONG HYPOCOTYL IN FAR-RED1 (HFR1), *HY5* HOMOLOGUE, and LONG AFTER FAR-RED LIGHT1, and a key regulator of flowering, *CONSTANS (CO)*, in the dark [98–103].



**Figure 1.** Hypothetical model depicting the mechanism how *cry*s and *phys* regulate the COP1/SPA pathway. (A) In the dark, COP1 and SPAs constitute an active E3 ligase complex that ubiquitinates the transcription factors, such as *HY5* and *HFR1*, which are positive regulators of photomorphogenesis. Poly-ubiquitinated *HY5* and *HFR1* are degraded through 26S proteasome pathway. (B) Upon light exposure, active *cry1*, *phyA* and *phyB* bind with SPAs in the nucleus and disrupt the interaction between COP1 and SPAs. This disruption inactivates the COP1/SPA complex, and therefore, *HY5* and *HFR1* accumulate to promote photomorphogenesis. *cry2* also binds with SPAs in response to blue light. The binding of *cry2* to SPAs does not disrupt the interaction between COP1 and SPAs but strengthens the binding of *cry2* to COP1. This enhanced interaction between *cry2* and COP1 inactivates the activity of COP1 to promote accumulation of *CO* that initiates flowering. The exact mechanism by which enhanced interaction between *cry2* and COP1 regulates the COP1 activity remains unknown. COP1, CONSTITUTIVE PHOTOMORPHOGENIC1; *cry1*, cryptochrome 1; *HFR1*, LONG HYPOCOTYL IN FAR-RED1; *HY5*, ELONGATED HYPOCOTYL 5; *phyA*, phytochrome A; *phyB*, phytochrome B; SPA, SUPPRESSOR OF PHYA-105; Ub, ubiquitin.

Cry1 and cry2 reportedly bind with SPA proteins in response to B light [77–79]. Moreover, phyA and phyB have been shown to interact with SPA proteins in a Pfr conformer-specific manner [80,81]. Consistent with the observations that COP1 requires a SPA1 association to exert E3 ligase activity in vitro and COP1 and SPA1 are dissociated by light exposure in plant cells [100,104], the binding of all these photoreceptors (except for cry2) to SPA proteins weakens the connection between COP1 and SPA proteins [77,78,80,81]. Hence, light-activated cry1, phyA, and phyB bind to SPAs to dissociate COP1 from SPAs, thus inhibiting COP1 activity and leading to the accumulation of HY5 and possibly other transcription factors, which promotes de-etiolation. However, B light-dependent cry2 binding to SPA proteins does not alter the affinity of COP1 for SPA proteins but strengthens the binding of cry2 to COP1 [79]. Although the mechanism remains unknown, this enhanced binding between cry2 and COP1 may attenuate COP1 activity to promote the accumulation of CO, which binds to the promoter of *FT* and induces its expression, thereby resulting in floral initiation. In addition to the light-dependent COP1 inactivation mechanism, these photoreceptors sequester the COP1 protein from transcription factors by depleting it from the nucleus under prolonged light exposure [105]. However, this nuclear exclusion mechanism is not fully understood.

The overlapping functions of cry1 and cry2 in regulating de-etiolation and photoperiodic flowering are at least partially explained by the COP1/SPA pathway (and PIF pathway) shared by these two B light receptors. However, the different modes of action of cry1 and cry2 to regulate COP1 activity may determine the substrate specificity of COP1 in the generation of functional differences between cry1 and cry2. The different affinities of cry1 and cry2 for SPA proteins may also account for the functional specificities because SPA proteins have diverse roles in plant development to a certain extent. However, the largest difference in signal transduction mechanisms between cry1 and cry2 is that only cry2 can bind with CIBs, which are specialized to function in photoperiodic flowering, in a B light-dependent manner [84,85]. Namely, the COP1/SPA or the PIF pathway likely represents the overlapping functions between cry1 and cry2, and the CIB pathway may confer on cry2 its more specific roles in the regulation of photoperiodic flowering [106]. However, the mechanism by which the COP1/SPA pathway and CIB pathway, which both regulate *FT* expression, converge downstream of cry2 is still unknown.

#### 4.2. PIF Pathway

PIF3 was first identified as a protein that binds to phys in a Pfr-specific manner [107,108]. Of the bHLH subfamily 15, to which PIF3 belongs, 7 PIFs (PIF1, PIF3, PIF4, PIF5, PIF6, PIF7 and PIF8) have been demonstrated to bind to Pfr of phyB through the Active Phytochrome B Binding (APB) motif [83]. PIF1 and PIF3 have also been shown to bind to phyA. Accordingly, PIF1 and PIF3 possess the Active PhyA Binding (APA) motif, which is necessary for phyA binding, in addition to the APB motif and the DNA binding bHLH domain. PIF1, PIF3, PIF4 and PIF5 are phosphorylated and degraded rapidly by R light exposure [109–114]. The R light-induced rapid phosphorylation and degradation of PIFs is largely dependent on phyA and phyB. Furthermore, the *pif1pif3pif4pif5* (*pifQ*) mutant exhibited a cop-like phenotype, albeit weak [115,116], suggesting that PIFs repress photomorphogenesis in the dark and that phyA and phyB induce photomorphogenesis by removing the PIF-imposed repression of photomorphogenesis via the rapid degradation of PIF1, PIF3, PIF4 and PIF5 in response to R light (Figure 2). However, phyB levels increase more in higher-order *pif* mutants under continuous R light [117,118]. The gentle degradation of phyB is dependent on binding to the APB motif of PIF3 and possibly other PIFs [119], which suggests a negative feedback circuit in which the binding of active phyB to PIFs not only promotes photomorphogenesis but also attenuates phyB signaling by reducing active phyB. The gentle degradation of phyB is partially mediated by COP1 [38]. In addition, the E3 ligase complex composed of CULLIN 3 and Light-Response Bric-a-Brac/Tramtrack/Broad (CUL3<sup>LRB</sup>) was found to recognize the phyB–PIF3 complex for its ubiquitination and proteasomal degradation under prolonged continuous R light [120] (Figure 2).

Interestingly, PIF1 is degraded under continuous FR light or by exposure to a small amount of R light [113,121]. This sensitive PIF1 degradation is mediated by phyA via binding to APA. Thus, one

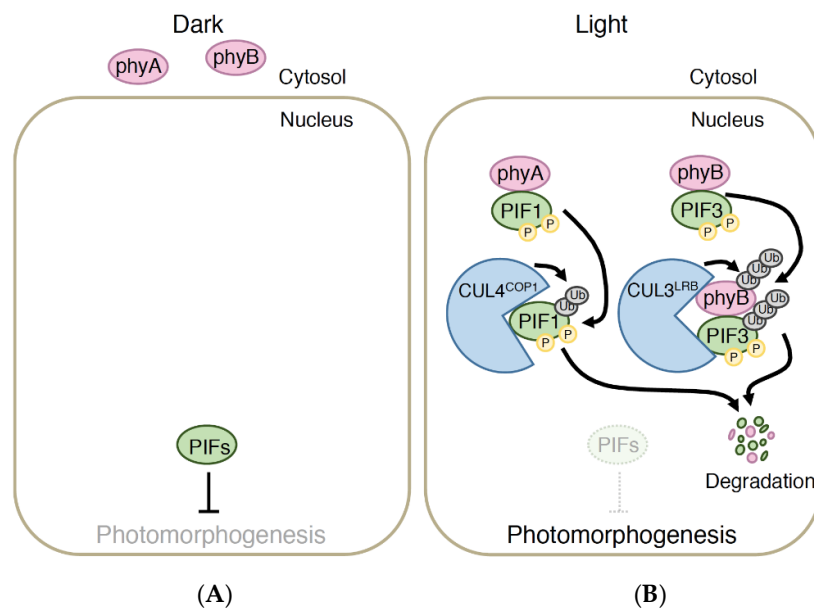
may expect that the binding of phyA to PIF1 is the sensitive reaction. However, the binding affinity to PIF1 in vitro does not differ between phyA and phyB [122]. Nevertheless, PIF1 has the highest affinity to phyA among all PIF family proteins tested to date. In addition, the CUL4<sup>COP1/SPA</sup> E3 ligase complex has recently been shown to promote phyA-induced rapid PIF1 degradation [123] (Figure 2). Mutations in *CUL3* do not affect phyA-induced rapid PIF1 degradation, thus defining the difference between CUL3<sup>LRB</sup>-mediated, phyB-dependent PIF3 degradation under prolonged R light conditions and CUL4<sup>COP1/SPA</sup>-mediated, phyA-induced PIF1 degradation. CUL4<sup>COP1/SPA</sup>-mediated sensitive PIF1 degradation provides clues to understanding the mechanism underlying the phyA-mediated sensitive response in the nucleus. Further analysis regarding the specificity or kinetics of these E3 ligase complexes will be necessary to fully understand the different modes of action of phyA and phyB.

The phyB regulation of PIF activity is accomplished not only of R-induced rapid degradation but also of other mechanisms. Specifically, the N-terminal photosensory domain of phyB is capable of binding to PIFs and mediating de-etiolation [28,124–126], although it does not degrade PIF3 [127]. Instead, PIF3, which co-immunoprecipitates the *RGA* and *PIL1* promoter DNA sequences, no longer binds with these promoter sequences under R light in transgenic plants expressing the N-terminal photosensory domain of phyB [127]. Both full-length phyB and the N-terminal photosensory domain of phyB inhibit the DNA binding of PIF1 and PIF3 under R light but not in the dark in vitro, suggesting that phyB promotes de-etiolation not only by degrading PIFs but also by sequestering PIFs from their target DNA sequences in response to R light [127]. Another example is that PIF7, which is phosphorylated but not degraded by R light, regulates de-etiolation and low R/FR-induced SAS [117,128]. However, how PIF7 activity is regulated remains unknown.

Recently, B light-dependent binding of crys to PIFs has been demonstrated [76,82]. To date, cry1 has been shown to bind to PIF3, PIF4 and PIF5 and cry2 has been shown to bind to PIF4 and PIF5, although not all combinations have been tested. These interactions regulate the B light-mediated inhibition of hypocotyl elongation, LBL-induced SAS and B light inhibition of warm temperature-induced cell growth. Similar to phys, crys are suggested to inhibit the activity of PIF4 and PIF5, although crys do not induce the degradation of PIF4 and PIF5 [76,82]. Whether crys sequester PIF4 and PIF5 from their DNA binding sites in a similar manner to phyB has not been tested. Thus, the mechanisms by which cry regulates the activity of PIF4 and PIF5 must be elucidated in the future.

Although most of the PIFs appear to have considerable functional redundancy in regulating de-etiolation, PIF family member proteins have distinct control functions in the diverse array of light-regulated physiological responses [83]. For instance, PIF1 has a dominant role in regulating seed germination [129] and PIF6 regulates seed dormancy [130]. PIF4, PIF5 and PIF7 promote SAS, with PIF4 and PIF5 acting more in LBL-induced and cry-mediated SAS and PIF7 mediating low R/FR-induced and phyB-mediated SAS [76,128]. Interestingly, phyB has recently been shown to act as a thermosensor [131,132]. Accordingly, PIF4 regulates temperature-dependent responses [133]. In addition, PIF4 is also known to regulate stomatal development [134]. However, neither the *pif* single nor the *pifQ* mutant shows a clear flowering phenotype in the normal temperature [116,135], implying that cry-PIF may not provide the mechanisms underlying the functional specificity between cry1 and cry2. In addition to the other diverse roles of PIFs, this family integrates light signals via endogenous signals, such as phytohormone signals and the circadian clock, thus highlighting the importance of PIFs in coordinating the transcriptional network to regulate plant development [83].





**Figure 2.** Hypothetical model depicting the mechanism by which phyA and phyB regulate the degradation of PIFs. (A) In the dark, the inactive phyA and phyB are localized in the cytosol. On the other hand, the nuclear localized PIFs regulate the expression of their target genes to repress photomorphogenesis. (B) In response to light, phyA and phyB translocate from the cytosol into the nucleus and bind with PIFs. The binding of phyA or phyB to PIFs initiates the phosphorylation and subsequent degradation of PIFs. phyA and phyB mediate the degradation of PIFs via distinct mechanisms. phyB bound PIF3 is recognized, ubiquitinated by CUL3<sup>LRB</sup> E3 ligase complex and degraded through 26S proteasome pathway. phyA mediates the phosphorylation of PIF1 in response to very low amount of light and the CUL4<sup>COP1</sup> E3 ligase complex recognizes the phosphorylated PIF1 for the ubiquitination and subsequent degradation through 26S proteasome pathway. The rapid degradation of PIFs trigger photomorphogenesis by removing the PIF-imposed repression of photomorphogenesis. COP1, CONSTITUTIVE PHOTOMORPHOGENIC1; CUL, cullin; LRB, Light-Response Bric-a-Brac/Tramtrack/Broad; P, phosphate group; phyA, phytochrome A; phyB, phytochrome B; PIF, PHYTOCHROME INTERACTING FACTOR; Ub, ubiquitin.

#### 4.3. Interaction between the COP1/SPA Pathway and PIF Pathway

As described above, the CUL4<sup>COP1/SPA</sup> E3 ligase complex directly targets PIF1 for light-induced rapid degradation [123]. In addition, the accumulation of PIF3 proteins is considerably reduced in the *cop1* mutant, even in the dark [110], suggesting that COP1 indirectly promotes the accumulation of PIF3, which likely occurs via the degradation of an as yet unknown negative regulator of PIF accumulation in the dark. Therefore, the COP1/SPA pathway directly and indirectly regulates the accumulation of PIFs in the dark or under light conditions.

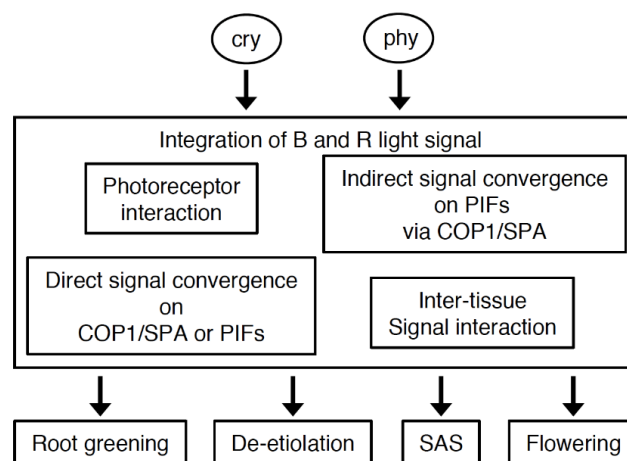
HY5 and PIFs (at least PIF1 and PIF3) have been shown to physically bind to form an antagonizing module that participates in the regulation of photosynthetic gene and reactive oxygen species-responsive gene transcription [136,137]. Although HY5 and PIF3 independently mediate the hypocotyl response under continuous R or FR light, PIF3 function is dependent on HY5 in the regulation of FR light-induced anthocyanin accumulation [138]. These results indicate the complex interaction between HY5 and PIFs downstream of photoreceptors. Consistent with the observation that the function of HFR1, which promotes phy-mediated de-etiolation and SAS, depends on PIF4 and PIF5, HFR1 inhibits the activity of PIF4 and PIF5 by binding to them to form non-functional heterodimers, which no longer bind to their target DNA sequences [139,140]. The *hfr1* mutant was initially identified as the long-hypocotyl mutant under continuous FR light [141]. However, subsequent reports have indicated that a phyA-independent but cry1-dependent long-hypocotyl phenotype occurs

under continuous B light [142] and HFR1 regulates the transcription of more than 70% of B light- and cry1-mediated genes [143], indicating that HFR1 acts downstream of cry1 signal transduction. Moreover, at least PIF4 (and one of PIF1, PIF3 and PIF5) has been reported to regulate the cry1-mediated hypocotyl response under continuous B light [82] and that PIF4 and PIF5 regulate cry-mediated and LBL-induced SAS [74–76,144]. The involvement of both HFR1 and PIF4/5 in cry-mediated responses suggests that HFR1 also forms non-functional heterodimers with PIF4 and PIF5 downstream of crys. Therefore, the regulation of PIF functions by the COP1/SPA complex involves at least two separate mechanisms: One that directly or indirectly regulates PIF accumulation and the other that regulates PIF activity by regulating the accumulation of its target proteins, such as HY5 and HFR1. However, the dependency and specificity of these interactions may differ across responses or light environments.

In the regulation of de-etiolation in the dark, *cop1* and *pif1* have a synergistic effect. In addition, HY5 protein accumulates in *pifQ*, even in the dark. According to these observations, PIF1 binds with COP1 to enhance the HY5 recognition of COP1 and promote the ubiquitination of HY5 [145]. Therefore, the COP1/SPA pathway and the PIF pathway affect each other, thus representing the complexity of downstream cry and phy signal transduction.

### 5. Coordination of Cryptochrome and Phytochrome Signal Transduction

Compared with the monochromatic light conditions in the laboratory, both crys and phys regulate related responses via shared signaling pathways under solar radiation in nature. Understanding how these photoreceptors act in natural or near-natural conditions is a challenging topic because of the complexity. Cry and phy independently incorporate B and R/FR light information, converge signals by photoreceptor interactions, and converge signals on common intermediates. Alternatively, they transduce the signals to selective intermediates and converge downstream (Figure 3). Here, we dissect the complex relations between cry and phy and examine the convergence of cry and phy signaling in the regulation of light responses.



**Figure 3.** Integration of blue and red light signals.

Cry and phys converge blue and red light signals at different levels to co-regulate physiological responses, such as root greening, de-etiolation, SAS, photoperiodic flowering, etc. The co-action mechanism includes photoreceptor interactions, direct signal convergence on common intermediates, indirect signal convergence on PIFs downstream of COP1/SPA, inter-tissue signal interactions, etc., although the role of photoreceptor interactions and the mechanism responsible for the direct convergence of cry and phy signals on the COP1/SPA complex or PIFs remain elusive. The types of co-action may be different depending on the responses. Indirect signal convergence on PIFs via COP1/SPA and inter-tissue signal interaction are evident in the regulation of SAS and photoperiodic flowering, respectively.

### 5.1. Observed Co-Action between Cryptochromes and Phytochromes in Physiological Responses

The *cry1cry2* double mutant responds normally to R light [73]. Conversely, the *phyAphyB* double mutant and the *phy* mutant, which lacks all five *Arabidopsis* phy species, respond to B light [146,147], although their responses are impaired to a certain extent, likely because phys absorb B light. These results clearly indicate that crys and phys do not require each other to exert their substantial signaling functions. Nevertheless, classical photobiological analyses have indicated co-action between crys and phys under certain conditions [148]. Specifically, B light exposure enhanced the effect of phy-mediated R light induction of the accumulation of glyceraldehyde-3-phosphate dehydrogenase in mlo seedlings. phy-mediated anthocyanin accumulation in mlo required B light exposure. In addition, an analysis of *Arabidopsis* photoreceptor mutants has provided genetic evidence of co-action between crys and phys. The most prominent synergistic effect has been observed in root greening [149]. B light induces stronger root greening than R light. This strong effect of B light on root greening can be observed in the *phyA* and *phyB* single mutants but is almost completely defective in the *cry1* mutant and the *phyAphyB* double mutant, indicating that *cry1* and *phyA* or *phyB* promote B light-induced root greening in a coordinated manner. The synergism between *cry* and *phy* mutations can also be seen in the regulation of de-etiolation. *phyB* mutation diminishes the function of *cry1*, and *cry1* mutation does so for the function of *phyB* in the regulation of hypocotyl response under simultaneous R and B irradiation [150]. However, these effects can be observed only under a short period of dichromatic irradiation with B and R light, not under continuous dichromatic irradiation, suggesting that *cry1* and *phyB* coordinate the inhibition of hypocotyl elongation under suboptimal light conditions. Such synergism is not observed between *cry1* and *phyA*. In addition, *phyB* and *cry2* mutations displayed a synergistic effect, albeit weakly, rather than an additive effect in the inhibition of hypocotyl elongation under continuous dichromatic irradiation with B and R light [73]. Thus, *phyB* partially co-acts with *cry1* or *cry2* in the regulation of hypocotyl response. The co-action between *phyB* and *crys* can also be observed in SAS because LBL enhances the effect of low R/FR-induced, *phyB*-mediated SAS [144].

In contrast, antagonistic effects have been reported between *cry* and *phy* mutations. For example, the *cry1* mutant has a substantially elongated hypocotyl compared with the wild type under continuous white light; however, the hypocotyl length of *cry1phyA* is significantly shorter than that of *cry1* [151]. A similar antagonistic effect between *cry1* and *phyA* is also observed in cotyledon expansion and chlorophyll accumulation under continuous white light [151]. The antagonistic effect between *cry* and *phy* is more evident in the regulation of photoperiodic flowering [73]. Namely, the *cry2* mutant shows a late-flowering phenotype specifically under LD conditions, but the effect of *cry2* mutation is considerably reduced in the *phyB* mutant, which flowers early regardless of day length.

### 5.2. Possible Roles of Photoreceptor Interactions and Direct Signal Convergence on Common Intermediates in the Co-Action between Cryptochromes and Phytochromes

*Cry1* physically binds with *phyA* and *phyB* in vitro, and *cry2* directly binds with *phyB* in plant cells [65,152]. Particularly, interactions between *phyB* and *cry2* have been observed only in certain photobodies [65]. These photoreceptor binding interactions may represent the underlying mechanism for the physiological co-action of crys and phys. *Avena phyA*, which displays kinase activity, directly binds with and phosphorylates *cry1* in vitro [152]. However, the phosphorylation of *cry1* occurs normally in *Arabidopsis phyA* or other *phy* mutants [48], suggesting that the observed in vitro *cry1* phosphorylation catalyzed by *phyA* exerts marginal effects on physiological responses. Hence, the effects of photoreceptor interactions on their physiological co-action remain unknown. Given that both *phyB-PIF3* and *cry2* undergo light-dependent phosphorylation that triggers polyubiquitination and protein degradation, it would be interesting to examine whether related protein kinases may act as the common regulators for the two distinct photoreceptors.

Shared signaling partners between crys and phys may provide clues to the molecular mechanism underlying the co-action of crys and phys. The interaction of crys with the COP1/SPA complex or PIFs does not require phys and vice versa because they bind to these signaling partners in yeast cells or

in vitro systems where other photoreceptors are not observed [76–82,108,153]. However, the binding of cry to these signaling partners may modify their affinity for phy and vice versa, the binding of cry may change the binding of phy to these signaling partners and vice versa, and the photoreceptor interactions may accelerate or decelerate binding to signaling partners. These possibilities should be experimentally addressed in the future, and the binding sites of these photoreceptors on their signaling partners should be a focus of such investigations. Crys and phys share a binding site on SPA proteins in certain cases; however, they recognize different structures in other cases [77–81]. Therefore, crys and phys possibly form a large protein complex or compete for the same structure of SPAs.

Crys and phyB recognize the different structures of PIF4/5 because crys bind with PIF4/5<sup>mAPB</sup> that harbor a mutation on the APB motif, which abolishes phyB binding [76]. In addition, phyB indeed binds with PIF4/5 under LBL conditions, in which crys can interact with PIF4/5 [76], suggesting that a trimeric complex can be formed among phyB, crys, and PIF4/5. Interestingly, the expression of PIF4/5<sup>mAPB</sup> does not fully induce hypocotyl elongation in response to LBL [76], suggesting that phyB binding to PIF4/5 promotes PIF4/5 activity to enhance hypocotyl growth in response to LBL. This promotion of PIF4/5 activity by phyB under LBL conditions is in striking contrast to the function of phyB in repressing PIF4/5 activity in R light [83], implying that crys may possibly reverse the function of phyB in the trimeric cry-PIF4/5-phyB complex under these light conditions.

Accordingly, B light-induced degradation of PIFs is also of great interest [154,155]. The B light-activated phyA mediate the rapid degradation of PIF1 and PIF3, whereas phy-mediated PIF1 degradation is enhanced in a *cry1cry2* double mutant under B light [154], indicating that crys antagonize phyA-mediated PIF1 degradation. Cry-phy binding may inhibit phyA to bind to PIF1, thereby protecting PIF1 proteins from degradation. Because crys bind to PIF3, PIF4 and PIF5 [76,82], crys may also interact with PIF1. Therefore, the binding of crys to PIF1 may reduce the affinity of PIF1 for phyA or impair the function of PIF1-bound phyA in a trimeric cry-PIF1-phyA complex. Moreover, cry1 and cry2 may function indirectly to antagonize phyA functions via the regulation of COP1/SPA activity because COP1 plays a role in the degradation of PIF1 [123]. Thus, the mechanism underlying the cry-mediated stabilization of PIFs, which represents the antagonism between crys and phyA, should be carefully examined in the future.

### 5.3. Indirect Signal Convergence Downstream of Cryptochromes and Phytochromes

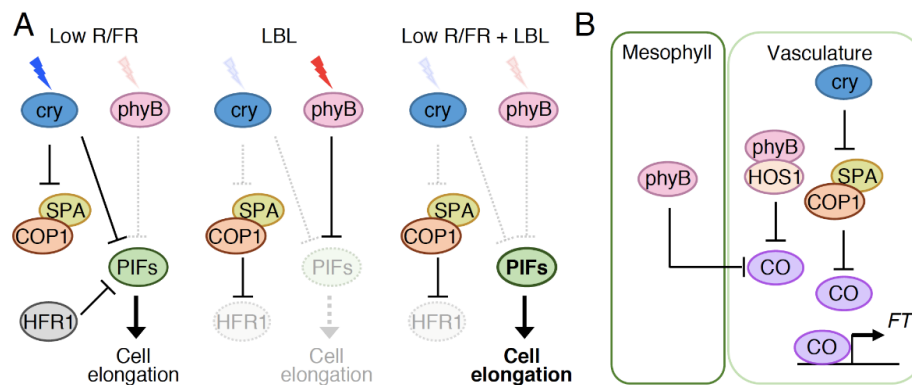
Regardless of the common intermediates, cry and phy may have preferential inputs and interact downstream. A transcriptome analysis comparing seedling response to continuous R light and 3 h of B light included with the continuous R light, in which cry1 and phyB coordinate to regulate de-etiolation, revealed that only 6% of B light-regulated, cry1-mediated genes are regulated by phyB under background R light conditions [63], suggesting that cry1 and phyB regulate transcription mostly via independent pathways. In addition, the expression of certain B light-regulated, cry1-mediated genes persists in R light even in the absence of B light, and this expression contributes to promoting the co-action of R and B light effects in the de-etiolation process, suggesting that phyB incorporates the independent, cry1-mediated transcriptional network into its signaling pathway to enhance light responses. The co-action between crys and phyB in the control of de-etiolation requires SPA1, SPA4, HY5, and HYH [63].

The preferential roles of common intermediates downstream of cry and phy are more evident in the regulation of SAS. phyB and crys bind to PIFs to regulate low R/FR-mediated and LBL-mediated SAS, respectively [76,114,128]. Because low R/FR reduces the Pfr/P<sub>tot</sub> ratio, it also impairs the binding of phyB to PIFs, thereby increasing the active PIF level to promote cell elongation. Although low R/FR increases the abundance of PIF4/5 [114], phyB favors PIF7 rather than PIF4/5 to regulate low R/FR-mediated SAS primarily by regulating the expression of auxin biosynthesis genes and auxin-responsive genes [128]. Instead, crys favor PIF4/5 to regulate LBL-induced SAS. PIF4/5 preferentially regulate the transcription of genes involved in cell wall modifications to facilitate cell elongation under LBL [76]. It should be noted here that the transcription of the cell wall modifying

genes is frequently regulated by auxin and LBL-induced and cry-mediated SAS is partly dependent on auxin [75]. Thus, the mechanism how crys regulate the transcription of the cell wall modifying genes in LBL via PIF4/5 should be carefully examined in the future.

Moreover, the function of the COP1/SPA pathway that positively regulates SAS is of great interest for understanding the complex interactions between phy and cry. COP1 enhances the effects of PIF3/4/5 on SAS under shade conditions with both LBL and low R/FR ratios [156], although whether it enhances the effects of PIF7 under such conditions is currently unknown. The function of COP1 in the regulation of SAS is largely dependent on HFR1. Interestingly, COP1 decreases the stability of the HFR1 protein but increases the abundance of the HY5 protein, which likely antagonizes the function of PIFs to avoid excess elongation under shade conditions [156], although the methods by which COP1 differentially recognize HY5 and HFR1 under these conditions are currently unknown. As described above, HFR1 and PIF4/PIF5 form non-functional heterodimers and the functions of HFR1 under shade conditions are dependent on PIF4/5 [139]. Therefore, under shade conditions, the COP1/SPA complex promotes the removal of HFR1-mediated repression, which represents the mechanism by which PIF4/5 exert their functions and elongate the hypocotyl. In addition, the stepwise application of low R/FR and LBL differentiated the roles of photoreceptors and enabled the description of more specific roles [144]. Namely, LBL enhances the effect of low R/FR via PIF4/5/7 in SAS-mediated petiole and hypocotyl elongation. First, LBL enhances the low R/FR-mediated abundance of PIF5. Notably, LBL combined with low R/FR decreases HFR1 abundance, although low R/FR alone stabilizes HFR1. Therefore, LBL enhances the effect of low R/FR not only by increasing the accumulation of PIF5 but also by promoting the proteasomal degradation of HFR1, thus highlighting the role of crys in regulating COP1/SPA activity in response to shade. Taking this evidence together, shade likely activates the function of PIFs by reversing phyB and cry binding, and in addition, the inactivated crys remove the HFR1-mediated inhibition of PIF activity via the regulated COP1/SPA pathway, thereby facilitating PIF-mediated cell growth [144,156] (Figure 4). Interestingly, DELLA proteins, which are the negative regulators of gibberellin signaling and the inhibitors of PIF functions, are degraded under shade conditions [157], thus representing another mechanism underlying the regulation of PIF activity by photoreceptors.

Other types of photoreceptor signaling interactions were also found in the regulation of photoperiodic flowering. As mentioned earlier, cry2 antagonizes phyB in the regulation of photoperiodic flowering [73]. The accumulation of CO, which promotes the transcription of *FT* in the vascular tissue to initiate flowering, represents this antagonistic effect between cry2 and phyB [158]. Namely, cry2 stabilizes CO, whereas phyB degrades it. As described above, cry2 stabilizes CO by inactivating the COP1/SPA complex [79,98,103]. However, the phyB-mediated degradation of CO is COP1 independent [98]. PFT1, a component of phyB signal transduction, promotes earlier flowering, although it regulates CO expression rather than protein stability [159,160]. PHL, a phyB binding protein that regulates flowering time, binds to CO protein at dusk not at dawn when phyB promotes CO degradation [161]. Whether PHL affects the stability of CO is also unclear. HOS1, a phyB binding E3 ubiquitin ligase, mediates the degradation of CO in the phloem companion cells [162,163]. However, the mechanism underlying phyB-mediated CO degradation is not fully understood. Interestingly, the tissue-specific expression of photoreceptors demonstrated that phyB expressed in mesophyll cells regulates flowering time [163,164], although phyB expressed in vascular tissue partly complement the early flowering *phyB* mutant phenotype [163]. On the contrary, cry2 promotes photoperiodic flowering solely in vascular tissue [165], thus suggesting that cry2 and phyB antagonize each other to control photoperiodic flowering partly via inter-tissue signaling between vascular tissue and mesophyll cells. The substance of inter-tissue signaling that is generated by phyB in mesophyll cells and suppresses flowering by promoting the degradation of CO via a COP1-independent mechanism in vascular tissue should be carefully investigated to fully understand the co-action between photoreceptors. Given that both of crys and phys are expressed throughout plant body, it would be interesting to examine whether this type of non-cell-autonomous interaction among photoreceptors regulate other light responses.



**Figure 4.** Indirect signal convergence downstream of crys and phyB. **(A)** cry- and phyB-mediated SAS. In the regulation of SAS, PIFs integrate cry and phyB signals via the direct binding of crys and phyB to PIFs and via cry-mediated inactivation of COP1/SPA. Under LBL conditions, inactivated crys relieve their direct inhibition of PIF activity and COP1/SPA activity, thereby leading to the degradation of HFR1 proteins. Pfr of phyB degrades PIF4/5 and phosphorylates and subsequently inactivates PIF7. Compared with non-shade conditions, LBL conditions induce PIF5 accumulation; however, active phyB may sequester PIFs from their target sequence under non-shade conditions. Low R/FR inactivates phyB, which promotes PIF4/5 accumulation and PIF7 dephosphorylation. In addition, crys bind to at least PIF4/5 to negatively regulate their activity and they inactivate the COP1/SPA complex, thereby promoting HFR1 accumulation to form non-functional heterodimers with PIF4/5. Therefore, in low R/FR conditions, although active PIFs accumulate, their activity is inhibited by the binding of crys and HFR1, which leads to the moderate activity of PIFs. Under low R/FR combined with LBL conditions, accumulated PIFs are released from direct inhibition of crys and HFR1 and exert full activity. Low R/FR-induced *HFR1* transcription is also inhibited by LBL. Moreover, HY5 may be stabilized under low R/FR combined with LBL. HY5 accumulations presumably inhibit detrimental excessive elongation. The molecular mechanism by which crys and phyB generate these different assignments remains unknown. **(B)** Antagonistic effect of phyB and cry2 on the regulation of photoperiodic flowering. Cry2 promotes flowering, whereas phyB inhibits flowering. cry2 promotes CO accumulation via the inactivation of the COP1/SPA complex in vascular tissue, and CO directly regulates *FT* expression to induce flowering. However, phyB promotes CO degradation via a COP1-independent mechanism. phyB binds with HOS1, an E3 ubiquitin ligase, to promote CO degradation in vascular tissue. In addition, phyB decelerates flowering in mesophyll cells. Therefore, phyB is also likely to promote CO degradation via inter-tissue signaling, although this signal has not been identified.

## 6. Conclusions and Future Outlook

Crys and phys act under different colors of the light spectrum to regulate related developmental processes via two independent but shared signaling pathways. These two common signaling pathways and direct cry-phy interactions provide clues to understand the molecular mechanisms underlying the co-action of crys and phys. However, as exemplified by cry- and phy-mediated SAS, cry and phy differentially regulate light responses via the preferential selection of signaling pathways that lead to the regulation of PIF activity. The mechanisms underlying the preferential selection of signaling pathways by crys and phys regardless of the binding of both photoreceptors to common signaling intermediates must be addressed to fully understand plant light responses. Crys and phys have complicated interactions including direct binding, signal convergence on or downstream of their common intermediates, and inter-tissue signal transduction, thus providing the mechanism by which plants are capable of responding plastically to fluctuating light environments in nature. These interactions likely provide general regulation of cell growth and photoperiodic flowering in many crop species. In addition, photoreceptors regulate many important agronomic traits that are likely species specific. Therefore, understanding the cry-phy interactions in individual crop species will provide insights for improving the yields and quality of crops.

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