

A constitutively monomeric UVR8 photoreceptor confers enhanced UV-B photomorphogenesis

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The plant ultraviolet-B (UV-B) photoreceptor UVR8 plays an important role in UV-B acclimation and survival. UV-B absorption by homodimeric UVR8 induces its monomerization and interaction with the E3 ubiquitin ligase COP1, leading ultimately to gene expression changes. UVR8 is inactivated through redimerization, facilitated by RUP1 and RUP2. Here, we describe a semidominant, hyperactive allele, namely *uvr8-17D*, that harbors a glycine-101 to serine mutation. UVR8^{G101S} overexpression led to weak constitutive photomorphogenesis and extreme UV-B responsiveness. UVR8^{G101S} was observed to be predominantly monomeric in vivo and, once activated by UV-B, was not efficiently inactivated. Analysis of a UVR8 crystal structure containing the G101S mutation revealed the distortion of a loop region normally involved in stabilization of the UVR8 homodimer. Plants expressing a UVR8 variant combining G101S with the previously described W285A mutation exhibited robust constitutive photomorphogenesis. This work provides further insight into UVR8 activation and inactivation mechanisms and describes a genetic tool for the manipulation of photomorphogenic responses.

UV-B | photoreceptor | acclimation | signal transduction | abiotic stress

Plant growth and development rely on appropriate responses to the light environment. Plants have evolved different photoreceptor families that respond to photons of specific wavelengths: red/far-red light-sensing phytochromes (phyA–E); blue light-sensing cryptochromes (cry1 and cry2), phototropins (phot1 and phot2), and Zeilupe family proteins (ztl, fkl1, and lkp1); and the ultraviolet (UV)-B-sensing UV RESISTANCE LOCUS 8 (UVR8) (1–4). Photoreceptors shape plant development throughout the entire life cycle. At the seedling stage, photoreceptor-mediated light perception initiates processes such as inhibition of hypocotyl elongation, cotyledon expansion, biogenesis of the photosynthetic machinery, and synthesis of photoprotective pigments (5–7). Notably, phytochrome, cryptochrome, and UVR8 pathways converge on the CONSTITUTIVELY PHOTOMORPHOGENIC 1–SUPPRESSOR OF PHYA-105 (COP1–SPA) E3 ubiquitin ligase complex, which is responsible for degradation of photomorphogenesis-promoting factors and is inactivated by photoreceptors under light (8, 9). Perception of UV-B and blue light through UVR8 and cry1, respectively, is crucial for UV-B protection and survival in the field (10).

UVR8 is a β -propeller protein that exists in a homodimeric ground state held together by an intricate network of salt-bridge interactions (4, 11–13). Absorption of UV-B photons by specific tryptophan residues that provide chromophore function, of which W285 plays a prime role, disrupts the electrostatic interactions stabilizing the homodimer, resulting in UVR8 monomerization. The active UVR8 monomer interacts with the WD40 domain of COP1 using a cooperative binding mechanism involving the core β -propeller domain of UVR8 together with its disordered C terminus, which harbors a so-called valine-proline (VP) motif (14–17). Interaction of UVR8 with COP1 inhibits COP1 activity by direct competition with COP1 substrates and remodeling of COP1 E3 ligase complex composition (6, 17, 18).

The COP1 substrate ELONGATED HYPOCOTYL 5 (HY5), a crucial positive regulator of light signaling (19), is induced transcriptionally and stabilized posttranscriptionally and is necessary for the regulation of most UVR8-induced genes (14, 20–24). Additionally, UVR8 regulates the stability or DNA-binding activity of a set of transcription factors affecting UV-B responses (25–32).

REPRESSOR OF UV-B PHOTOMORPHOGENESIS 1 (RUP1) and RUP2 are among early UV-B-induced genes and encode components of a negative feedback loop that down-regulates UVR8 signaling (33). RUP1 and RUP2 physically interact with UVR8 and promote UVR8 redimerization back to its inactive state (33, 34). Additionally, a role for RUP1 and RUP2 as components of an E3 ligase complex mediating HY5 degradation has recently been suggested (35).

To dissect UVR8 function, several studies have made use of targeted mutagenesis to assess the role of individual tryptophan residues in photoreception and the importance of electrostatic interactions in mediating dimerization (4, 11, 12, 36–39). For example, UVR8^{W285F} is locked in its inactive homodimeric form and is thus unresponsive to UV-B (4, 11, 12, 37). Several other mutant forms, including UVR8^{W285A} and UVR8^{D96N,D107N}, display a constitutive interaction with COP1, whereas their downstream effects differ (37–39). Notably, UVR8^{W285A} is a weak UVR8 dimer that does not respond to UV-B activation (4, 11, 37). Overexpression of UVR8^{W285A} leads to constitutive

Significance

Coping with UV-B is crucial for plant survival in sunlight. The UV-B photoreceptor UVR8 regulates gene expression associated with photomorphogenesis, acclimation, and UV-B stress tolerance. UV-B photon reception by UVR8 homodimers results in monomerization, followed by interaction with the key signaling protein COP1. We have discovered a UV-B hypersensitive UVR8 photoreceptor that confers strongly enhanced UV-B tolerance and generated a UVR8 variant based on the underlying mutation that shows extremely enhanced constitutive signaling activity. Our findings provide key mechanistic insight into how plants respond and acclimate to UV-B radiation.

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photomorphogenesis, likely through inhibition of COP1 (37). On the other hand, UVR8^{D96N,D107N} is constitutively monomeric; however, despite its constitutive interaction with COP1, UV-B exposure is required to induce physiological responses (11, 39).

We present here *uvr8-17D*, a semidominant, hypersensitive *uvr8* mutant allele that harbors the UVR8^{G101S} protein variant. We found that UVR8^{G101S} is predominantly monomeric in vivo, requires UV-B for activation, and confers an exaggerated UV-B response when expressed at wild-type levels, which is attributed to impaired redimerization and sustained UVR8 activity. Moreover, we describe the engineered UVR8^{G101S,W285A} variant with amazingly strong constitutive activity in vivo.

Results

***uvr8-17D* Exhibits Enhanced UV-B-Induced Photomorphogenesis.** To identify regulators of the UVR8 pathway, we screened an ethyl methanesulfonate (EMS)-mutagenized *Arabidopsis* population (Col-0 accession) for mutants with an aberrant hypocotyl phenotype when grown for 4 d under weak white light supplemented with UV-B. Several mutants with long hypocotyls contained mutations in the *UVR8* coding sequence; we thus named these alleles *uvr8-16* and *uvr8-18* to *uvr8-30* (SI Appendix, Fig. S1A). Conversely, several of the mutants displaying short hypocotyls in the screen conditions contained mutations in *RUP2*; we named these alleles *rup2-3* to *rup2-10* (SI Appendix, Fig. S1B). However, one mutant with enhanced UV-B responsiveness did not carry a *RUP2* mutation and showed a semidominant hypocotyl phenotype under UV-B (SI Appendix, Fig. S2A). Interestingly, in this mutant we found a G-to-A transition in the *UVR8* locus that results in a glycine-101 to serine (G101S) amino acid change (Fig. 1A); we thus named this allele *uvr8-17D*.

uvr8-17D seedlings exhibited shorter hypocotyls and elevated anthocyanin levels under UV-B compared with wild type, whereas under white light there were only minor phenotypic differences (Fig. 1B–D). Hypocotyl lengths were indistinguishable in dark-grown seedlings, suggesting that *uvr8-17D* does not exhibit significant constitutive activity (SI Appendix, Fig. S2B and C). Consistent with the UV-B-hypersensitive phenotype, CHALCONE SYNTHASE (CHS) and HY5 accumulated to higher levels in *uvr8-17D* than in wild type (Fig. 1E and SI Appendix, Fig. S2D). *uvr8-17D* seedlings also showed enhanced UV-B acclimation and stress tolerance (Fig. 1F). Compared to wild type, *uvr8-17D* plants showed greater dwarfing at rosette stage under UV-B, whereas they grew normally in the absence of UV-B (Fig. 1G). Altogether, the *uvr8-17D* phenotype and UV-B responsiveness resembled that of UVR8 overexpression (UVR8-OX) lines (SI Appendix, Fig. S3A–C) (14); however, UVR8^{G101S} levels in *uvr8-17D* were comparable to endogenous UVR8 levels in wild type (Fig. 1H and SI Appendix, Fig. S3C).

We tested whether UVR8^{G101S} is affected in its homodimeric ground state or in UV-B-activated monomerization by SDS-polyacrylamide gel electrophoresis (SDS-PAGE) analysis without sample heat denaturation, as previously described (4). In this assay, UVR8^{G101S} migrated as a constitutive monomer, whereas wild-type UVR8 derived from seedlings grown in the absence of UV-B or irradiated with saturating UV-B migrated as a homodimer or a monomer, respectively (Fig. 1I). Moreover, in contrast to wild-type UVR8, no UVR8^{G101S} homodimers could be detected in vivo by dithiobis(succinimidyl propionate) (DSP) cross-linking, even in conditions without UV-B (Fig. 1J). However, when assessing the oligomeric state of recombinant UVR8^{G101S} in vitro, we found through analytical size-exclusion chromatography experiments that the UVR8^{G101S} mutant protein purified from insect cells behaved similarly to wild-type UVR8, in that it eluted as a dimer under –UV-B conditions and as a monomer after UV-B treatment (Fig. 1K). We thus conclude that UVR8^{G101S} is a hypersensitive UVR8 variant, which is very likely associated with its monomeric state in vivo.

UVR8^{G101S} Overexpression Leads to Extreme UV-B Photomorphogenesis.

UVR8^{G101S} expression at levels comparable to a previously described UVR8-OX line (14) (Fig. 2A) resulted in strikingly enhanced anthocyanin accumulation under UV-B (Fig. 2B and C). Interestingly, hypocotyl elongation was strongly reduced in UVR8^{G101S}-OX lines when compared with wild type and UVR8-OX, even in the absence of UV-B (Fig. 2D and E). To determine whether the latter was due to the extreme sensitivity of UVR8^{G101S} to very low levels of UV-B from the fluorescent tubes in this light field (37) or to the intrinsically constitutive activity of overexpressed UVR8^{G101S}, we measured the hypocotyl length of seedlings grown in the dark or in monochromatic red or blue light. Although UVR8^{G101S}-OX seedlings displayed only a slightly altered hypocotyl-growth phenotype in darkness, some of the lines showed open cotyledons, suggesting that UVR8^{G101S}-OX induces weak constitutive photomorphogenesis (SI Appendix, Fig. S4A and B). More strikingly, comparable to their phenotype described above (Fig. 2D and E), UVR8^{G101S}-OX seedlings were indeed shorter in continuous red or blue conditions (SI Appendix, Fig. S4C–F), suggesting that UVR8^{G101S} broadly sensitizes photomorphogenesis. Consistently, mature UVR8^{G101S}-OX plants exhibited dwarfing and early flowering even in the absence of UV-B, with dwarfing further exaggerated under UV-B (Fig. 2F). In agreement with this enhanced photomorphogenesis, UVR8^{G101S}-OX lines showed a constitutively enhanced UV-B tolerance that was even further enhanced upon UV-B acclimation (Fig. 2G).

UVR8^{G101S} Shows No Increased Binding Affinity for COP1 under UV-B.

In agreement with the constitutive activity of UVR8^{G101S} when overexpressed, yeast two-hybrid (Y2H) analysis showed that UVR8^{G101S} interacts weakly with COP1 in the absence of UV-B (SI Appendix, Fig. S5A). UV-B irradiation of the yeast strongly enhanced the UVR8^{G101S}-COP1 interaction to a level comparable to the UVR8-COP1 interaction under UV-B, which, however, is strictly UV-B dependent (SI Appendix, Fig. S5A) (4). Similarly, in comparison to UVR8, recombinant UVR8^{G101S} showed enhanced in vitro binding to the COP1 WD40 domain in the absence of UV-B as shown through quantitative grating-coupled interferometry (GCI)-binding assays (SI Appendix, Fig. S5B). UV-B enhanced the affinity of both UVR8 and UVR8^{G101S} for COP1, but the effect was much stronger for wild-type UVR8 (SI Appendix, Fig. S5B). Importantly, the G101S mutation did not increase the affinity of UV-B-exposed UVR8^{G101S} for COP1 in vitro. This suggests that intrinsically increased affinity of UVR8^{G101S} for COP1 does not underlie the enhanced activity of UVR8^{G101S} in planta. However, it should be noted that in addition to the intrinsic binding affinity, the in planta interaction of UVR8 with COP1 is affected by negative regulation through the activities of RUP1 and RUP2 facilitating UVR8 redimerization (34), which may be impaired in the constitutively monomeric UVR8^{G101S}. Indeed, coimmunoprecipitation of COP1 with YFP-UVR8 and YFP-UVR8^{G101S} in transgenic lines not only confirmed the weak constitutive interaction of UVR8^{G101S} with COP1, but also revealed an enhanced association of UVR8^{G101S} with COP1 under UV-B (SI Appendix, Fig. S5C). In agreement with increased COP1 interaction (40), nuclear accumulation of UVR8^{G101S} was enhanced compared to UVR8 (SI Appendix, Figs. S5 and S6). Mechanistically, COP1 activity under UV-B is mainly inhibited through the C-terminal VP motif of UVR8 (15–17), and, in agreement, we observed that CRISPR/Cas9-generated mutations causing truncation of the UVR8 C terminus suppressed the UV-B-hypersensitive phenotype of *uvr8-17D* (SI Appendix, Fig. S7A–D).

UVR8^{G101S,W285A} Shows Strong Constitutive Photomorphogenesis.

We hypothesized that combining the G101S mutation conferring UV-B hypersensitivity with the W285A mutation conferring weak constitutive activity (37) would create a strong constitutive

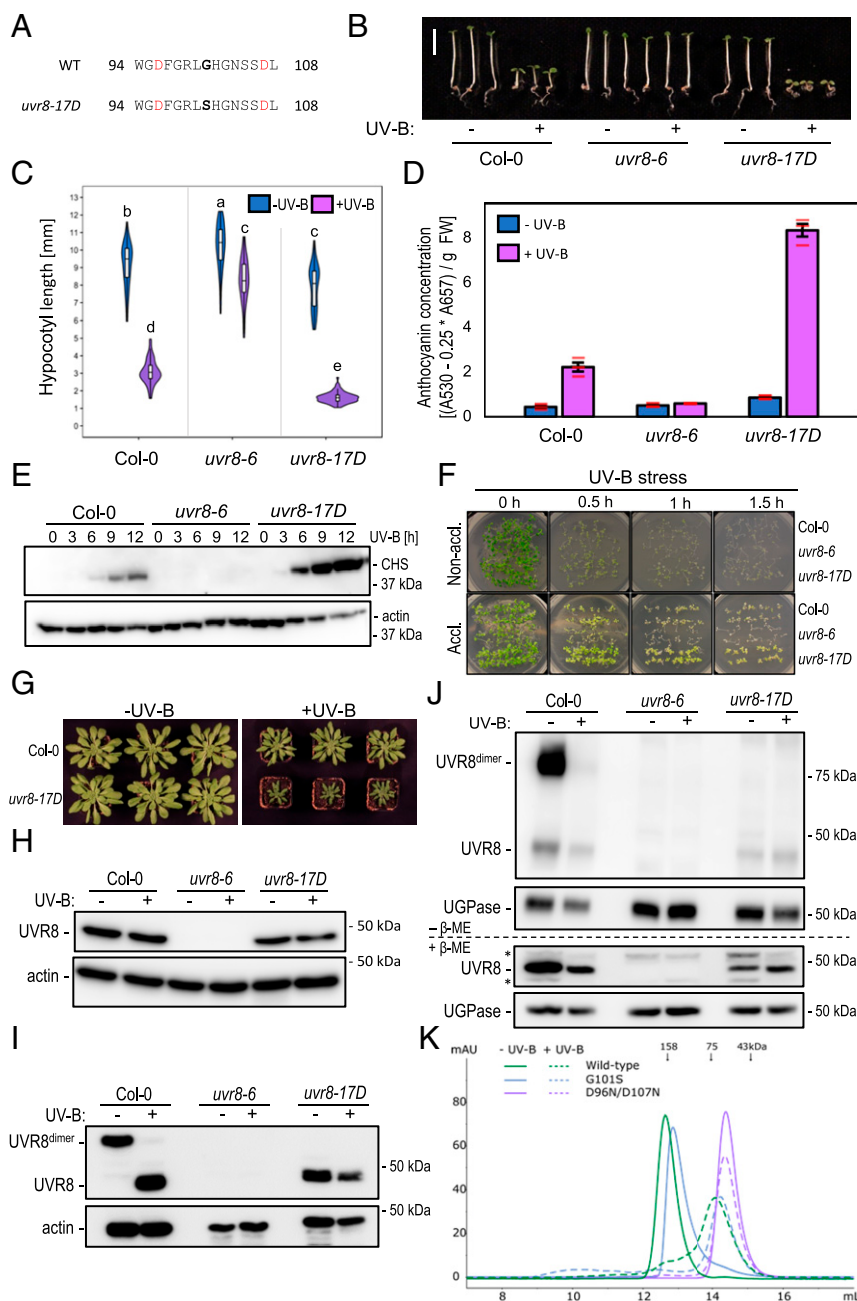


Fig. 1. Enhanced UV-B photomorphogenesis in plants expressing constitutively monomeric UVR8^{G101S}. (A) UVR8 amino acid sequence of residues 94 to 108 in wild type (WT) and *uvr8-17D*. The G101S mutation in *uvr8-17D* is indicated in bold. Asp-96 and -107 involved in dimer interaction are indicated in red. (B) Representative images of wild-type (Col-0), *uvr8-6* null mutant, and *uvr8-17D* seedlings grown in white light or white light supplemented with UV-B. (Scale bar, 5 mm.) (C) Quantification of hypocotyl length ($n > 60$); shared letters indicate no statistically significant difference in the means ($P > 0.05$). (D) Anthocyanin concentration; values of independent measurements (red bars), means, and SEM are shown ($n = 3$). (E) Immunoblot analysis of CHS and actin (loading control) protein levels in Col-0, *uvr8-6*, and *uvr8-17D* grown in white light for 4 d and then white light supplemented with UV-B for 0 to 12 h. (F) Survival of Col-0, *uvr8-6*, and *uvr8-17D* seedlings after UV-B stress. Seedlings were grown for 7 d in white light (nonacclimated) or white light supplemented with UV-B (acclimated), then exposed to varying durations (0 to 1.5 h) of broadband UV-B stress. Pictures were taken after a 7-d recovery period. (G) Rosette phenotype of Col-0 and *uvr8-17D* grown for 56 d under short-day conditions in white light or white light supplemented with UV-B. (H) Immunoblot analysis of UVR8 and actin (loading control) protein levels in 7-d-old Col-0, *uvr8-6*, and *uvr8-17D* seedlings. (I) UVR8 dimer/monomer status in Col-0 and *uvr8-17D*. Protein samples were extracted from dark-grown seedlings either exposed or not exposed to 15 min of saturating UV-B and analyzed using immunoblot analysis of samples separated through SDS-PAGE without prior heat denaturation. Actin is shown as loading control. (J) Immunoblot analysis of UVR8 in DSP cross-linked extracts (Top, -β-ME) of dark-grown Col-0, *uvr8-6*, and *uvr8-17D* seedlings either exposed or not exposed to 15 min of saturating UV-B. Cross-linking was reversed by treating with 5% β-ME (Bottom, +β-ME). UGPase is shown as loading control. β-ME, β-mercaptoethanol. (K) Size-exclusion chromatography of recombinant UVR8, UVR8^{G101S}, and UVR8^{D96N/D107N} proteins purified from Sf9 insect cells, with and without UV-B treatment.

UVR8 variant. Thus, we created UVR8^{G101S,W285A}, which, in contrast to UVR8^{W285A} and UVR8^{G101S} that were still able to form homodimers in vitro, appeared fully monomeric in vitro as

well as in the cross-linking assay in vivo (Fig. 3A and *SI Appendix, Fig. S84*). We observed a strongly elevated constitutive interaction of UVR8^{G101S,W285A} with COP1 compared to the

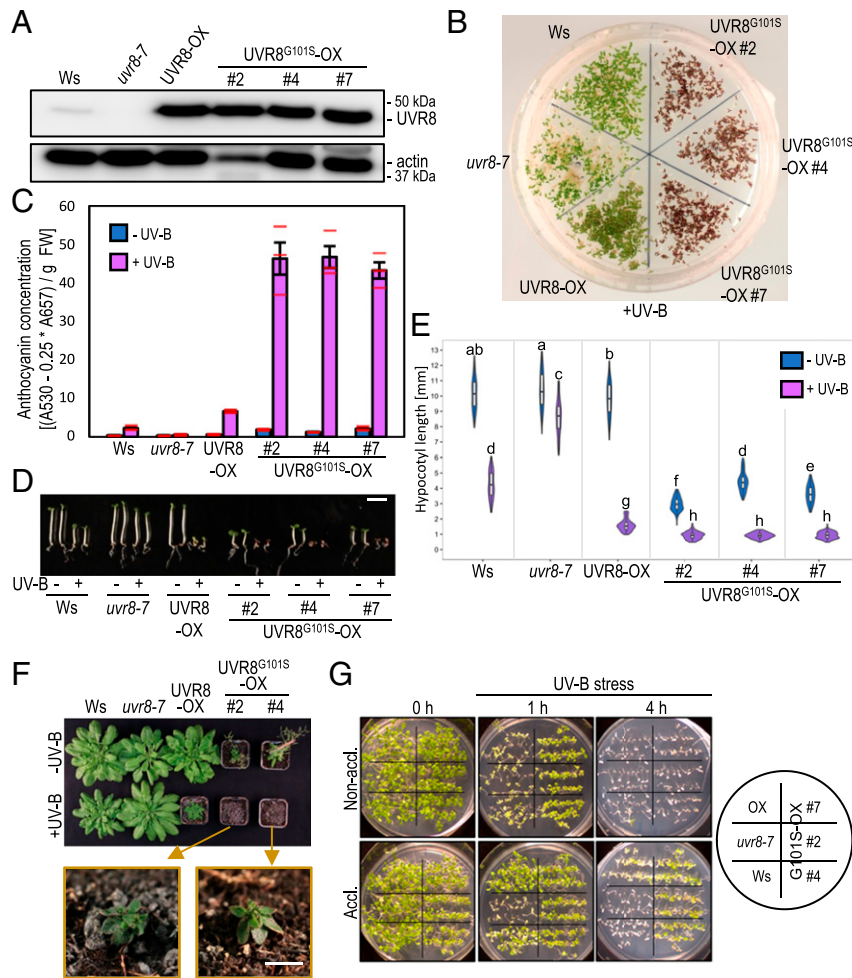


Fig. 2. Overexpression of UVR8^{G101S} results in strongly enhanced UV-B photomorphogenesis. (A) Immunoblot analysis of UVR8 and actin (loading control) protein levels in wild type (Ws), *uvr8-7*, *uvr8-7/Pro₃₅₅:UVR8* (UVR8-OX), and three independent *uvr8-7/Pro₃₅₅:UVR8^{G101S}-OX* (UVR8^{G101S}-OX #2, #4, and #7) lines. (B) Representative image of plate-grown seedlings grown under UV-B. (C) Anthocyanin concentration of seedlings depicted in B. Values of independent measurements (red bars), means, and SEM are shown ($n = 3$). (D) Representative images of seedlings grown in white light or white light supplemented with UV-B. (Scale bar, 5 mm.) (E) Quantification of hypocotyl length of seedlings depicted in D ($n > 60$). Shared letters indicate no statistically significant difference in the means ($P > 0.05$). (F) Rosette phenotype of representative plants grown for 56 d under short-day conditions in white light or white light supplemented with UV-B. The closeups show UV-B-grown UVR8^{G101S}-OX plants. (Scale bar, 5 mm.) (G) Survival of seedlings grown for 7 d in white light (nonacclimated) or white light supplemented with UV-B (acclimated) then exposed for varying durations (0 to 4 h) to UV-B stress. Pictures were taken after a 7-d recovery period.

COP1-UVR8^{G101S} or COP1-UVR8^{W285A} interaction in the Y2H system (Fig. 3B) as well as in GCI assays (Fig. 3C and see also *SI Appendix, Fig. S5B*). When we attempted to generate transgenic UVR8^{G101S,W285A}-OX lines (Pro₃₅₅:UVR8^{G101S,W285A}), we noticed seedling lethality in the T1 generation, strongly resembling *cop1* null alleles (41). We thus generated lines expressing UVR8^{G101S,W285A} under control of the estradiol-inducible XVE system or the *UVR8* promoter. Indeed, upon estradiol treatment, XVE:UVR8^{G101S,W285A} lines showed an extreme photomorphogenic phenotype in darkness (Fig. 3D and E). In lines with weak expression of UVR8^{W285A} and UVR8^{G101S,W285A} under the native *UVR8* promoter (Fig. 3F), the former (Pro_{UVR8}:UVR8^{W285A}) showed only a very minor phenotype in the absence of UV-B, similar to previous reports (36, 37), whereas the latter (Pro_{UVR8}:UVR8^{G101S,W285A}) showed a striking constitutive photomorphogenic phenotype resembling that associated with UVR8^{W285A}-OX (Fig. 3F-H and *SI Appendix, Fig. S8B and C*). Of note, this constitutive photomorphogenesis was despite UVR8^{G101S,W285A} protein levels being lower than endogenous UVR8 in wild type (Fig. 3F). We conclude that the combination of G101S and W285A in UVR8

yields a fully monomeric variant with very strong constitutive activity in vivo.

The Combination of G101S and W285F Confirms a Basal Constitutive Activity for UVR8^{G101S}. We further generated and characterized a UVR8^{G101S,W285F} variant. The W285F mutation is known to abolish all UV-B responsiveness (4, 37), whereas, as described above for UVR8^{G101S}-OX (Fig. 2C-E), the G101S mutation results in weak constitutive activity in conditions devoid of UV-B and a constitutively monomeric conformation in vivo. Indeed, integrating the G101S mutation into the UVR8^{W285F} variant caused strongly reduced dimer stability in UVR8^{G101S,W285F}, mimicking the conformation of UVR8^{G101S} (*SI Appendix, Fig. S8A*). As further predicted, UVR8^{G101S,W285F} retained a basal interaction with COP1 in the Y2H system, similar to the COP1-UVR8^{G101S} interaction in the absence of UV-B; however, by contrast, the COP1-UVR8^{G101S,W285F} interaction was not further enhanced under UV-B (*SI Appendix, Fig. S9A*). UVR8^{G101S,W285F}-OX lines exhibited a phenotype in both the absence and presence of UV-B resembling the phenotype of UVR8^{G101S}-OX lines in -UV-B conditions (*SI Appendix, Fig.*

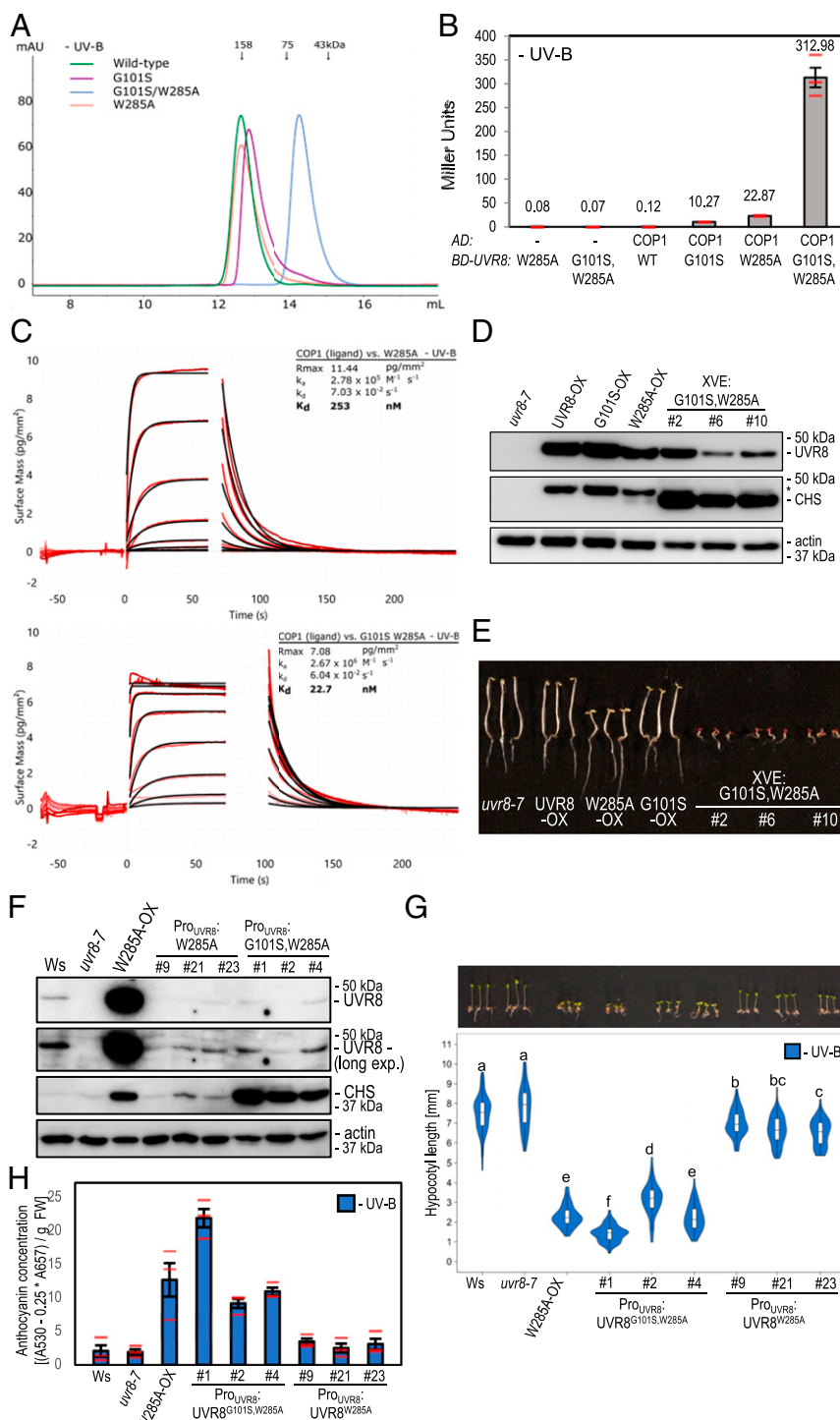


Fig. 3. Low expression of UVR8^{G101S,W285A} is sufficient for strong constitutive photomorphogenesis. (A) Size-exclusion chromatography of recombinant UVR8 (wild-type), UVR8^{G101S} (G101S), UVR8^{W285A} (W285A), and UVR8^{G101S,W285A} (G101S/W285A) proteins purified from Sf9 insect cells. (B) Quantitative Y2H analysis of the interaction between COP1 and UVR8 (WT), UVR8^{G101S}, UVR8^{W285A}, and UVR8^{G101S,W285A} in the absence of UV-B. AD, activation domain; BD, DNA-binding domain. (C) Binding kinetics of the full-length UVR8^{W285A} and UVR8^{G101S,W285A} versus the COP1 WD40 domain obtained by GCI experiments. Sensorgrams of protein injected are shown in red, with their respective heterogeneous ligand binding model fits in black. The following amounts were typically used: ligand – COP1 (2,000 pg/mm²); analyte – UVR8 (2 μM highest concentration). k_a , association rate constant; k_d , dissociation rate constant; K_d , dissociation constant. (D) Immunoblot analysis of UVR8, CHS, and actin (loading control) protein levels in seedlings of *uvr8-7*, *uvr8-7/Pro^{35S};UVR8* (UVR8-OX), *uvr8-7/Pro^{35S};UVR8^{G101S}* (#2 (G101S-OX)), *uvr8-7/Pro^{35S};UVR8^{W285A}* (W285A-OX), and three independent lines of *uvr8-7/XVE:UVR8^{G101S,W285A}* (XVE:G101S,W285A #2, #6, and #10) grown on 5 μM estradiol. Asterisk shows residual UVR8 signal after stripping of the polyvinylidene difluoride (PVDF) membrane. (E) Representative image of seedlings described in D grown for 4 d in darkness on plates supplemented with 5 μM estradiol. (Scale bar, 5 mm.) (F) Immunoblot analysis of UVR8, CHS, and actin (loading control) protein levels in wild type (Ws), *uvr8-7*, *uvr8-7/Pro^{35S};UVR8^{W285A}*, and three independent lines of each of *uvr8-7/Pro^{UVR8};UVR8^{W285A}* (#9, #21, and #23) and *uvr8-7/Pro^{UVR8};UVR8^{G101S,W285A}* (#1, #2, and #4). (G) Representative seedling images and quantification of hypocotyl length ($n > 60$) for seedlings described in F. Shared letters indicate no statistically significant difference in the means ($P > 0.05$). (Scale bar, 5 mm.) (H) Anthocyanin concentration of seedlings described in F. Values of independent measurements (red bars), means, and SEM are shown ($n = 3$).

S9 B–E), confirming that the constitutive phenotype induced by UVR8^{G101S} overexpression in the absence of UV-B is not linked to W285-mediated photoactivation.

The G101S Mutation Distorts a Critical Loop Interaction in the UVR8 Dimer. We next investigated the effects of the G101S mutation in structural detail. No crystals could be obtained for the UVR8^{G101S} protein, yet crystals developed for UVR8^{G101S,W285A}, diffracting to 1.75-Å resolution (SI Appendix, Table S1). The structure revealed a nonsymmetric dimeric arrangement of UVR8^{G101S,W285A} in the crystal lattice, as the relative position of the monomers are rotated ~10° with respect to the previously determined symmetric UVR8 wild-type dimer (SI Appendix, Figs. S10 and S11) (11). The altered dimeric assembly of UVR8^{G101S,W285A} contains a unique salt-bridge/hydrogen bond network between the two monomers with a reduced number of interactions compared to the wild-type dimer (SI Appendix, Fig. S11 and Table S2). The G101S mutation maps to a loop that forms part of the crystallographic UVR8 dimer interface (Fig. 4A and SI Appendix, Fig. S12). A structural superposition with wild-type UVR8 revealed that the position of the loop in UVR8^{G101S,W285A} is distorted by the G101S mutation (Fig. 4B–D). Importantly, the slightly larger Ser side chain cannot be accommodated within the loop. This forces a movement of the loop that affects the position of Asp-96 and -107, the mutation of which to Asn has been previously shown to impair UVR8 homodimerization (11, 39). The W285A mutation results in rearrangements of adjacent Trp side chains reminiscent of the movements observed in the UVR8^{W285A} crystal structure (12), highlighting the potential changes upon UV-B irradiation (Fig. 4C). We also obtained crystal structures of UVR8^{D96N,D107N} and UVR8^{D96N,D107N,W285A} variants that revealed no large structural rearrangements in the loop (Fig. 4E and F, and SI Appendix, Fig. S13). Together, our structural analysis reveals that the G101S mutation affects the conformation of a loop region involved in UVR8 homodimer stabilization.

Enhanced Activity of UVR8^{G101S} Is Caused by Impaired Inhibition through RUP1 and RUP2. Mutations in *RUP1* and *RUP2* enhance the UV-B photomorphogenic phenotype, as active monomeric UVR8 is more prevalent in *rup1 rup2* (33, 34). We hypothesized that monomeric UVR8^{G101S} enhances UV-B signaling due to its inability to dimerize, thus mimicking a *rup1 rup2* mutant after UV-B exposure. We first tested the interaction of UVR8 and UVR8^{G101S} with RUP2 in the Y2H system. Overall, the UVR8–RUP2 interaction was only slightly affected in yeast strains containing UVR8^{G101S} compared to UVR8 (Fig. 5A).

We then constructed higher-order mutant combinations between *uvr8-17D*, *rup1*, and *rup2*. Under UV-B, RUP2 was detected at an elevated level in *uvr8-17D* and *uvr8-17D rup1* when compared to that in wild type and *rup1*, respectively (Fig. 5B). Interestingly, *uvr8-17D* was similar to *rup1 rup2* in terms of hypocotyl length and flavonol accumulation (Fig. 5C and D), whereas *uvr8-17D* seedlings displayed an intermediate phenotype between a *rup2* and a *rup1 rup2* mutant regarding anthocyanin accumulation (Fig. 5E). Together with the observation that the anthocyanin phenotypes of *uvr8-17D rup2* and *uvr8-17D rup1 rup2* are stronger than that of the single *uvr8-17D* mutant (Fig. 5E), this suggests that RUP2 is still able to partially inhibit UVR8^{G101S}.

Interestingly, in the presence of functional RUP1 and RUP2, the UVR8^{G101S} mutation significantly enhances the UV-B phenotype (*uvr8-17D* versus wild type; Fig. 5C and E); however, in their absence, UVR8^{G101S} does not induce stronger UV-B signaling compared to UVR8 (*uvr8-17D rup1 rup2* versus *rup1 rup2*; Fig. 5C and E). Together, this demonstrates that, whereas RUP1 and RUP2 proteins retain some ability to negatively regulate UVR8^{G101S}, the enhanced activity of this variant is only evident in a background where RUP1 and RUP2 are present. We

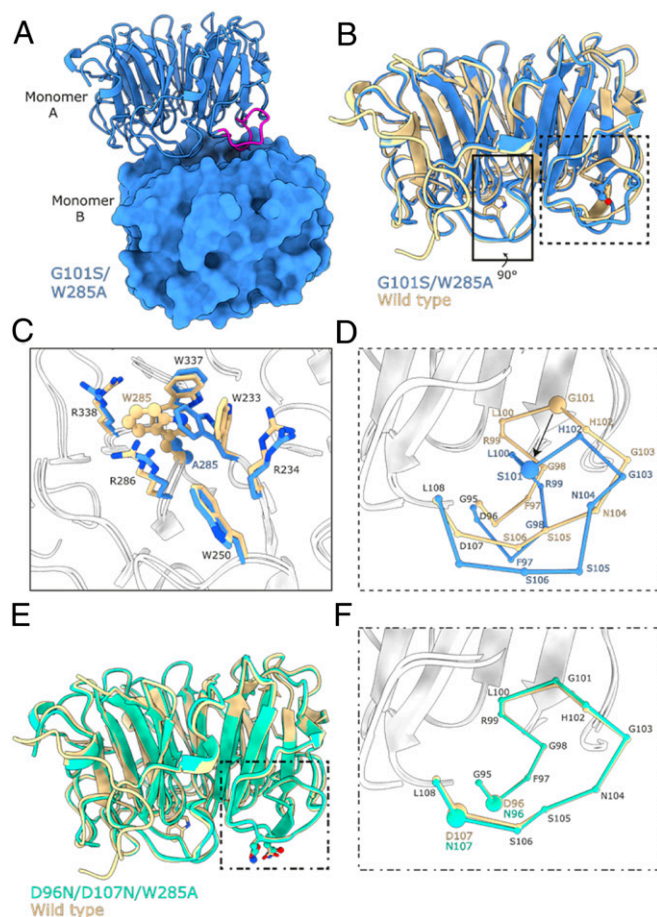


Fig. 4. UVR8^{G101S,W285A} distorts a critical interaction loop at the dimer interface. (A) Ribbon and surface representations of the UVR8^{G101S,W285A} dimer. Highlighted in magenta is the critical loop containing the G101S mutation. (B) Superposition of UVR8^{G101S,W285A} (blue) with a wild-type UVR8 (PDB:4D9S, yellow) in ribbon representation. The site of mutation, residue 101, is represented in a ball-and-stick manner and colored by their atomic identity. Residue 285 is represented by sticks. (C) Zoomed-in view of the site of the W285A mutation. The site of mutation is represented in a ball-and-stick manner and the surrounding residues are shown as sticks. (D) Zoomed-in view of the loop containing the G101S mutation. The loop is represented with each ball corresponding to a C_α carbon to highlight the structural rearrangements of the loop. (E) Superposition of UVR8^{D96N,D107N,W285A} (green) with UVR8 represented as in B. (F) Zoomed-in view of the loop containing G101 represented as in D.

conclude that the enhanced activity of UVR8^{G101S} under UV-B is due to impaired redimerization by RUP1 and RUP2, as UVR8^{G101S} shows a strongly reduced ability to dimerize in vivo.

Discussion

uvr8-17D is a UV-B-hypersensitive UVR8 photoreceptor allele that is linked with a single glycine-101 to serine amino acid change. The UV-B hypersensitivity of *uvr8-17D* is associated with its monomeric conformation in vivo, suggesting that redimerization facilitated by RUP1 and RUP2 is impaired and that UVR8^{G101S} therefore remains active much longer than wild-type UVR8 (Fig. 5F). Our work suggests that UVR8 dimerization is not absolutely required to keep UVR8 inactive, but it is important to make UVR8 inactivatable after activation by UV-B. In agreement, *uvr8-17D* phenotypes under UV-B resemble those of *rup1 rup2* double mutants, further supporting the prime importance of UVR8 redimerization to optimally balance UV-B-induced

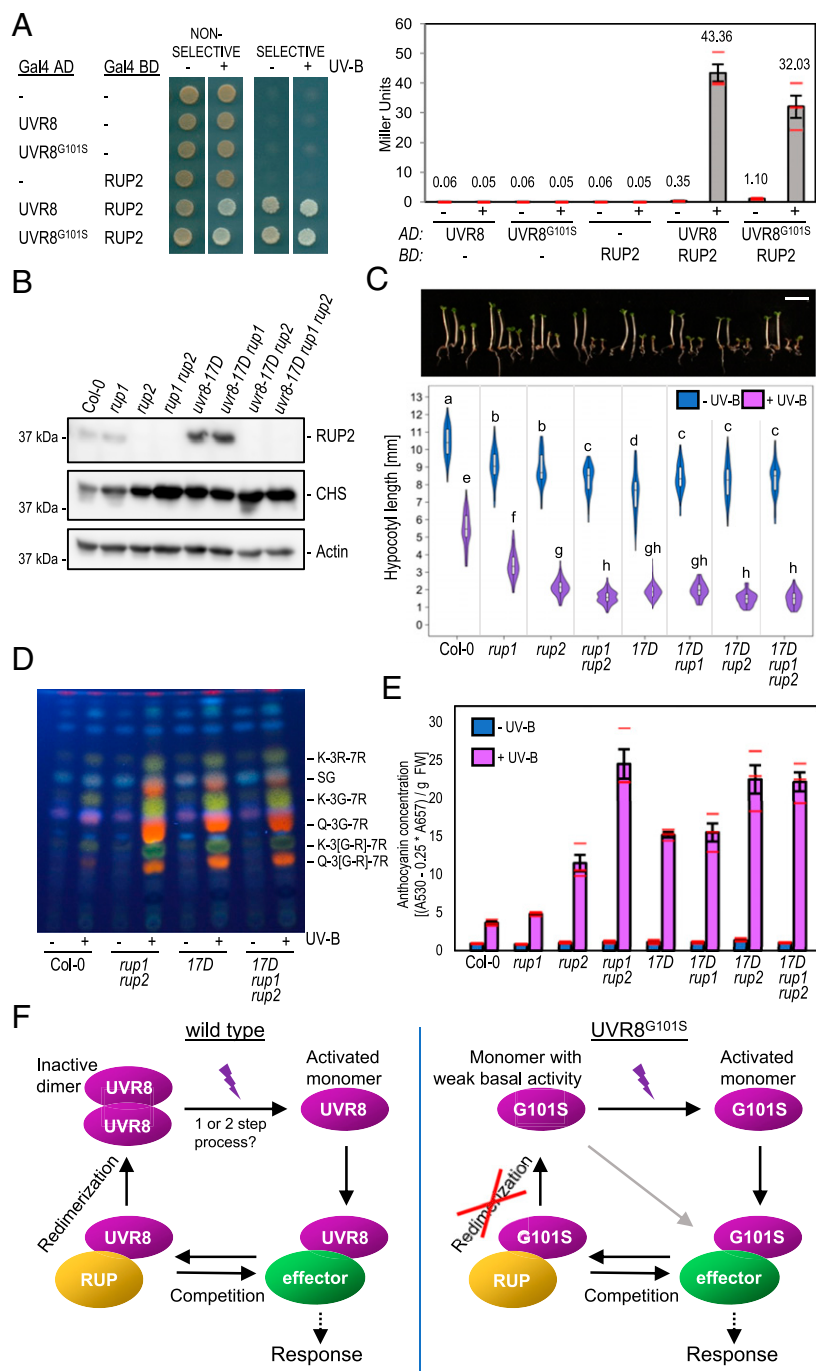


Fig. 5. The enhanced activity of UVR8^{G101S} depends on the presence of RUP1 and RUP2. (A) Y2H analyses of the interactions of RUP2 with UVR8 and UVR8^{G101S} in the presence or absence of UV-B. *Left*: growth assay on selective SD/-Trp/-Leu/-His medium. *Right*: quantitative β -galactosidase assay. AD, activation domain; BD, DNA-binding domain. (B) Immunoblot analysis of RUP2, CHS, and actin (loading control) protein levels in seedlings of wild type (Col-0), *rup1*, *rup2*, *rup1 rup2*, *uvr8-17D*, *uvr8-17D rup1*, *uvr8-17D rup2*, and *uvr8-17D rup1 rup2* grown for 4 d in weak white light supplemented with UV-B. (C) Representative images of seedlings described in B (*17D*, *uvr8-17D*) and quantification of hypocotyl length ($n > 60$). Shared letters indicate no statistically significant difference in the means ($P > 0.05$). (Scale bar, 5 mm.) (D) High-performance thin layer chromatography (HPTLC) analysis of the flavonol glycoside levels in 4-d-old seedlings of wild type (Col-0), *rup1 rup2*, *uvr8-17D*, and *uvr8-17D rup1 rup2* grown in white light or white light supplemented with UV-B. Identified compounds include: K-3R-7R, kaempferol-3-O-rhamnoside-7-O-rhamnoside; SG, sinapoyl glucose; K-3G-7R, kaempferol-3-O-glucoside-7-O-rhamnoside; Q-3G-7R, quercetin-3-O-glucoside-7-O-rhamnoside; K-3[G-R]-7R, kaempferol-3-O-[rhamnosyl-glucoside]-7-O-rhamnoside; Q-3[G-R]-7R, quercetin-3-O-[rhamnosyl-glucoside]-7-O-rhamnoside. (E) Anthocyanin concentration of seedlings described in B. Values of independent measurements (red bars), means, and SEM are shown ($n = 3$). (F) Working model of the UVR8 photocycle in the case of wild type (*Left*) and G101S-mutated UVR8 (*Right*). In the wild type, UV-B induces monomerization of dimeric UVR8 in a one- or two-step photon absorption process. The activated monomer then interacts with effector proteins such as COP1 and transcription factors to induce a photomorphogenic response. RUP1/RUP2 (RUP) proteins compete with these effectors to abrogate signaling activity. Afterward, RUP-bound UVR8 undergoes redimerization and is brought back to its initial inactive state. UVR8^{G101S} exists as a monomer in vivo and exhibits weak constitutive activity in the absence of UV-B, as seen in overexpression lines. UV-B absorption fully activates the UVR8^{G101S} monomer and RUP proteins still compete with other UVR8 signaling effectors. However, the full cycle cannot be completed because redimerization is intrinsically impossible. This results in sustained UVR8 signaling, leading to enhanced UV-B photomorphogenesis.

photomorphogenesis with plant growth and development (33, 34, 42–44).

Intriguingly, UVR8^{G101S} appeared monomeric in vivo, whereas in vitro it appeared monomeric in a gel-based assay and dimeric in a size-exclusion chromatography experiment (Fig. 1K and *SI Appendix*, Fig. S14). The SDS-PAGE assay is not sensitive enough to recognize weak dimeric variants such as UVR8^{W285A}, which appears monomeric (4, 11, 12, 37). However, in cross-linking and size-exclusion chromatography assays, UVR8^{W285A} is indeed dimeric, whereas UVR8^{G101S} cross-linked as a dimer only poorly and only when overexpressed. UVR8^{G101S} was however dimeric in size-exclusion chromatography assays, which could be linked to the different concentrations of UVR8 protein used and the different chemical environments between the in vitro assay and the in vivo cellular environment.

The monomeric nature of UVR8^{G101S} in vivo is reminiscent of the UVR8^{D96N,D107N} variant previously generated based on structural information (*SI Appendix*, Fig. S15A) (11, 39). Similar to UVR8^{G101S}, UVR8^{D96N,D107N} also shows weak constitutive COP1 binding (*SI Appendix*, Fig. S15 B–E) and requires UV-B for functional activation (39). By contrast though, no UV-B hypersensitivity was previously reported for transgenic lines expressing UVR8^{D96N,D107N} (39). However, in our phenotypic assays, transgenic UVR8^{D96N,D107N}-OX lines phenocopied the UV-B-hypersensitive phenotype of UVR8^{G101S}-OX (*SI Appendix*, Fig. S15 F and G), suggesting that UVR8^{D96N,D107N} and UVR8^{G101S} indeed have similar enhancing effects on UV-B responsiveness in vivo, consistent with our conclusion that the *uvr8-17D* phenotype is directly linked to its monomeric conformation in vivo. In support, our atomic structure of UVR8^{G101S,W285A} suggests a direct mechanistic link between the mutation of G101 and the misorientation of the D96 and D107 residues, likely underlying the impaired homodimer formation and weak constitutive interaction with COP1. In addition, the constitutive interaction of UVR8^{G101S} and UVR8^{D96N,D107N} with COP1 may sensitize other photomorphogenic pathways by sequestering COP1, although this effect is only apparent when these UVR8 variants are strongly overexpressed.

The effects of UVR8^{G101S} and UVR8^{D96N,D107N} thus suggest that monomeric UVR8 variants require UV-B to trigger the UV-B signaling pathway, confirming that engineered constitutive UVR8 monomers are not sufficient for strong constitutive UVR8 activity. This confirms that monomerization and activation of UVR8 by UV-B can clearly be uncoupled, at least for mutated UVR8 variants (39). However, in these cases, the mutant variants are simply monomeric versions that have not been UV-B activated. Whether wild-type UVR8 proteins that monomerize in response to UV-B—a process associated with broader structural changes that likely also affect the VP-containing C terminus (13, 16, 17, 45, 46)—can or even need to be similarly UV-B activated remains to be shown. Nonetheless, it may be that the activation of wild-type UVR8 is a two-step process, with photon absorption events required both for chromophore excitation that leads to monomerization and for functional activation. An engineered monomeric UVR8 mutant may be primed for activation by bypassing the need for monomerization prior to activation. Alternatively, one photon absorption event may be sufficient for both monomerization and activation. In this case, monomerization and activation by UV-B would be intrinsically linked in wild-type UVR8. Another possibility is that monomeric wild-type UVR8 absorbs additional UV-B photons as an efficient way to keep the photoreceptor active, without going through redimerization and reactivation by monomerization. Conclusive evidence to address these possibilities will require structural determination of full-length UV-B-activated monomer versus an engineered monomeric UVR8.

Despite remaining monomeric in vivo, UVR8^{G101S} retains an interaction with and remains weakly negatively regulated by RUP1

and RUP2. The UVR8–COP1 interaction may be inhibited by direct interaction of RUP proteins with the VP-containing C terminus of UVR8, which is the same domain that UVR8 uses to bind both COP1 and direct transcription factor targets (15–17, 25, 26, 47). This implies that UVR8 redimerization may proceed via two steps: 1) RUPs outcompete COP1 and other VP-domain interactors, separating them from UVR8; and 2) RUPs facilitate UVR8 redimerization. As redimerization occurs at a normal rate in a *cop1* background (34), competitive removal of COP1 alone is evidently not sufficient to induce redimerization. In *uvr8-17D*, step 1 could explain the remaining partial activity of RUPs on UVR8^{G101S}, whereas step 2 is abolished.

Importantly, the *uvr8-17D rup1 rup2* triple mutant did not show phenotypic differences compared with *rup1 rup2*, suggesting the mechanisms underlying UVR8^{G101S} hyperactivity are linked to RUP1/RUP2 activity. As a result of the more prevalent active UVR8 monomer, the increased signaling induced by UVR8^{G101S} originates early and upstream in the pathway and all subsequent UVR8 signaling mechanisms, including UVR8 nuclear accumulation, interaction with COP1, inhibition of its E3 ligase activity, and inhibition of transcription factor activity, are likely to be indirectly promoted by the relatively increased activity of UVR8^{G101S}.

UVR8 is widely conserved across the green lineage, and several studies have confirmed its role in mediating UV-B responses in a diverse range of species (48–56). G101 is conserved in UVR8 orthologs, as are residues involved in photoreception and dimer stability (4, 52, 57) (*SI Appendix*, Fig. S16). The phenotype of *uvr8-17D* holds the exciting possibility that a homologous glycine-to-serine mutation leads to enhanced UV-B responses in other species as well. This may eventuate as a convenient way to generate UV-B-hypersensitive phenotypes, including enhanced acclimation and UV-B tolerance, made possible by a single nucleotide substitution. Additionally, expressing UVR8^{G101S,W285A} at very low levels is sufficient to produce a striking constitutive photomorphogenic phenotype. Notwithstanding the use of UVR8^{G101S} in other plant species, both for fundamental research and potential biotechnological applications, *uvr8-17D* may serve as a genetic tool to further investigate UVR8 activity regulation and the effects of enhanced UVR8 signaling in *Arabidopsis*.

Methods

Hypocotyl length measurements, extraction, and analysis of anthocyanins and flavonoids, immunoblot analysis, protein cross-linking, Y2H analysis, protein purification from Sf9 cell cultures, analytical size-exclusion chromatography, in vitro methylation, protein crystallization and data collection, crystallographic structure solution and refinement, and GCI are described in *SI Appendix*, *Materials and Methods*.

Plant Material. *Arabidopsis thaliana* mutants *uvr8-6* (14), *cop1-4* (58), *hy5-215* (59), *rup1-1*, *rup2-1*, and *rup1-1 rup2-1* (33) are all in the Columbia (Col) accession. *uvr8-7*, *uvr8-7/Pro_{35S}:UVR8* line #2 (UVR8-OX) (14), *uvr8-7/Pro_{35S}:UVR8^{W285A}* line #3, and *uvr8-7/Pro_{35S}:UVR8^{W285F}* line #1 (37) are in the Wassilewskija (Ws) accession. *uvr8* and *rup2* alleles identified in this work stem from an EMS-mutagenized population in the Col accession.

Combinatorial mutants *uvr8-17D rup1-1*, *uvr8-17D rup2-1*, and *uvr8-17D rup1-1 rup2-1* were generated by genetic crossing. Genotyping of *rup1-1* and *rup2-1* alleles was performed as described (33), and *uvr8-17D* genotyping was through PCR amplification of a genomic fragment with a forward (5'-TCG GGA TGA GAT GAT GAC-3') and a reverse primer (5'-TAG ACC CAA CAT TGA CCC-3') followed by digestion with *HaeIII* (NEB) yielding diagnostic fragments of 344/173/145 bp for wild type and 489/173 bp for *uvr8-17D*.

To generate transgenic lines expressing UVR8 variants, mutations corresponding to G101S, W285A, W285F, and D96N/D107N were introduced into the UVR8 coding sequence in pDONR207 (Thermo Fisher Scientific) by site-directed mutagenesis. UVR8^{G101S}, UVR8^{G101S,W285A}, UVR8^{G101S,W285F}, and UVR8^{D96N,D107N} sequences were inserted into the Gateway-compatible binary vector pB2GW7 (60). UVR8^{G101S,W285A} was also inserted into the Gateway-compatible binary vector pMDC7 allowing estradiol-inducible expression (61). To generate lines expressing YFP-tagged constructs, UVR8 and

UVR8^{G101S} were inserted into the Gateway-compatible binary vector pB7WGY2 (60). The control line expressing Pro_{35S}:StreptII-3xHA-YFP in a Col background has been previously described (62).

To generate transgenic lines expressing UVR8 under its own promoter, the UVR8 promoter (1,809 bp upstream of the translational start ATG) was respectively cloned upstream of the UVR8^{W285A} and UVR8^{G101S,W285A} coding sequences and introduced into the Gateway-compatible binary vector pGWB501 (63).

The CRISPR/Cas9 system was used to delete the UVR8 C terminus in wild type and *uvr8-17D*. An sgRNA directed against the UVR8 sequence was inserted into the pHEE401E vector (64) using overlapping complementary oligos 5'-ATT GGC GAC ACC CAG CTT TTC CC-3' and 5'-AAA CGG GAA AAG CTG GGT GTC GC-3', and mutants were identified in T1 after sequencing the corresponding part of the UVR8 locus.

Arabidopsis plants were transformed using the floral dip method (65) and transgenic lines with a single insertion locus (75% resistance in T2) were selected for homozygosity in T3.

Growth Conditions. For hypocotyl length measurements, quantification of anthocyanins, analysis of flavonol glycosides, and acclimation assays, seedlings were grown under aseptic conditions on half-strength Murashige and Skoog (MS) medium (Duchefa) supplemented with 1% (wt/vol) sucrose (Applchem). For immunoblot analysis, seedlings were grown on sterile half-strength MS medium (Duchefa) without sucrose. For induction of estradiol-inducible expression under the XVE system, the growth medium was supplemented with 5 μ M of β -estradiol (Sigma).

After sowing, seeds were stratified for 2 d at 4 °C in the dark before the start of the light treatments, which were performed at 22 °C as described before (14, 22). White light (3.6 μ mol m⁻² s⁻¹) was supplied by Osram L18W/30 tubes, which were supplemented with narrowband Philips TL20W/01RS

tubes (1.5 μ mol m⁻² s⁻¹) for UV-B treatment. To analyze etiolated growth in darkness, seeds were exposed to 6-h white light (approximately 60 μ mol m⁻² s⁻¹) to induce germination before being transferred back to darkness. For monochromatic light treatments, a light-emitting diode (LED) chamber (floraLEDs, CLF Plant Climatics) was used with 5 μ mol m⁻² s⁻¹ of blue light or 30 μ mol m⁻² s⁻¹ of red light.

For analysis of vegetative growth, plants were grown in short days (8-h/16-h light/dark cycles) in UV-B lamp-containing GroBanks (CLF Plant Climatics), under white light (120 μ mol m⁻² s⁻¹) or white light supplemented with UV-B (1.5 μ mol m⁻² s⁻¹) (66).

For broadband UV-B treatments (acclimation assays, analyses of dimer/monomer status), Philips TL40W/12RS UV-B tubes were used (4, 14).

Data Availability. The atomic coordinates of complexes have been deposited with the following Protein Data Bank (PDB) accession codes: UVR8^{D96N,D107N}, 6XZL (<https://www.rcsb.org/structure/6XZL>), UVR8^{D96N,D107N,W285A}, 6XZM (<https://www.rcsb.org/structure/6XZM>), and UVR8^{G101S,W285A}, 6XZN (<https://www.rcsb.org/structure/6XZN>).

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