Exchange of a single amino acid residue in the cryptophyte phycobiliprotein lyase

GtCPES expands its substrate specificity

Natascha Tomazic1, Kristina E. Overkamp1, Marco Aras1,#, Antonio J. Pierik2, Eckhard Hofmann3, and Nicole Frankenberg-Dinkel1

1Abteilung Mikrobiologie, Fachbereich Biologie, Technische Universität Kaiserslautern, Germany
2Abteilung Biochemie, Fachbereich Chemie, Technische Universität Kaiserslautern, Germany
3Proteinkristallographie, Fakultät für Biologie und Biotechnologie, Ruhr-Universität Bochum, Germany

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# present address: Lehrstuhl Technische Biochemie, Technische Universität Dortmund, Germany

*To whom the correspondence should be addressed: Nicole Frankenberg-Dinkel, Abteilung Mikrobiologie, Fachbereich Biologie, Technische Universität Kaiserslautern, 67663 Kaiserslautern, Germany; nfranken@bio.uni-kl.de; Tel. +49 631 205 2353; Fax: +49 631 205 3799.

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ABSTRACT

Cryptophyte algae are among the few eukaryotes that employ phycobiliproteins (PBP) for light harvesting during oxygenic photosynthesis. In contrast to the cyanobacterial PBP that are organized in large membrane-associated super complexes, the phycobilisomes, those from cryptophytes are soluble within the chloroplast thylakoid lumen. Their light-harvesting capacity is due to covalent linkage of several open-chain tetapyrrole chromophores (phycobilins). Guillardia theta utilizes the PBP phycoerythrin PE545 with 15,16-dihydrobiliverdin (DHBV) in addition to phycoerythrobilin (PEB) as chromophores. Thus far, the assembly of cryptophyte PBPs is not yet completely understood but involves the action of PBP-lyases as shown for cyanobacterial PBP. PBP-lyases facilitate the attachment of the chromophore in the right configuration and stereochentistry. Here we present the functional characterization of eukaryotic S-type PBP lyase GtCPES from G. theta. We show GtCPES mediated transfer and covalent attachment of PEB to the conserved Cys82 of the acceptor PBP β-subunit (PmCpeB) of Prochlorococcus marinus MED4. Based on the previously solved crystal structure, the GtCPES binding pocket was investigated using site-directed mutagenesis. Thereby, amino acid residues involved in phycobilin binding and transfer were identified. Interestingly, exchange of a single amino acid residue Met67 to Ala extended the substrate specificity to phycoerythrobilin (PCB) likely by enlarging the substrate-binding pocket. Variant GtCPES_M67A binds both PEB and PCB forming a stable, colorful complex in vitro and in vivo produced in Escherichia coli. GtCPES_M67A is able to mediate PCB transfer to Cys82 of PmCpeB. Based on our data we postulate that a single amino acid residue determines the bilin-specificity of phycoerythrin S-type lyases but that additional factors regulate hand over to the target protein.
Phycobiliproteins (PBPs) are the photosynthetic light-harvesting structures of cyanobacteria, red algae and cryptophytes and are used to absorb regions of the visible light spectrum that are poorly covered by chlorophylls. Consequently, PBPs efficiently absorb light in the spectral region between 430 and 680 nm (1). They consist of apo-proteins with covalently linked open-chain tetrapyrrole molecules termed phycobilins. While PBPs in cyanobacteria and red algae are organized in large light-harvesting structures called phycobilisomes, those of cryptophyte algae are soluble in the thylakoid lumen of the chloroplast (2-4). Cryptophycean PBPs occur in large concentrations and are organized in (αβ)(α′β′)-heterodimers (4). In contrast to cyanobacteria, cryptophytes only use a single type of PBP. The model species *Guillardia theta* utilizes phycocyanin 545 (PE545) with an absorption maximum at 545 nm (5). It consists of the α-subunit CPEA with bound 15,16-dihydrobacteriopheophytin (DHPV) and the β-subunit CpeB loaded with three molecules of phycocyanobilin (PCB) (6,7). The *G. theta* GtCPES mediates transfer of PEB to Cys 82 of PmCpeB apo-protein and we were not able to solve this issue until now (14). Likely, the lyase is not catalyzing the transfer reaction itself, but rather assists in the handover of the phycobilin in the correct configuration to allow stereospecific autocatalytic attachment.

In this study, we demonstrate the specific ability of GtCPES to support the transfer of PEB to Cys 82 of CpeB from the cyanobacterium *Prochlorococcus marinus* MED4, whereas CpeB can be purified in adequate amounts to test chromophore transfer employing GtCPES (17). To verify whether GtCPES is sufficient for correct bilin addition to *PmCpeB* we compared spontaneous and lyase mediated attachment of 3(Z)-PEB to *PmCpeB* by recording the fluorescence emission of assembled holo-CpeB. Covalent attachment was further confirmed by zinc-blot analysis. *PmCpeB* lacking the cysteine 

**RESULTS**

*GtCPES mediates transfer of PEB to Cys 82 of PmCpeB*. Although we have previously shown that GtCPES binds PEB and 15,16-DHBV with high affinity, an involvement in the chromophorylation of the *G. theta* PBP β-subunit at β-Cys 82 has not yet been shown. This was mainly due to solubility issues of the recombinant *G. theta* CpeB apo-protein and we were not able to solve this issue until now (14). Furthermore, it is still unclear whether chromophorylation of PBP has to follow a specific order. In this regard, it has previously been reported that β-Cys 155 is the first site of chromophorylation in phycocyanin (18). Therefore, we took another approach to investigate whether GtCPES is able to support PEB attachment to the PE β-subunit. We made use of β-PE (CpeB) from the cyanobacterium *P. marinus* MED4 (*PmCpeB*). This β-subunit is the only PBP in this organism and is highly degenerated, as it only possesses a single chromophorylation site at Cys 82 (19,20). Recombinant *PmCpeB* can be purified in adequate amounts to test chromophore transfer employing GtCPES (17). To verify whether GtCPES is sufficient for correct bilin addition to *PmCpeB* we compared spontaneous and lyase mediated attachment of 3(Z)-PEB to *PmCpeB* by recording the fluorescence emission of assembled holo-CpeB. Covalent attachment was further confirmed by zinc-blot analysis. *PmCpeB* lacking the cysteine
attachment site (i.e. \textit{PmCpeB\_C82A}) served as a negative control (Figure 1). In all experiments, the lyase was used in excess to prevent the presence of free bilin in the sample. Spontaneous and lyase-mediated attachment resulted in the formation of covalent fluorescent phycobiliprotein complexes (Figure 1AB). Reaction products (i.e. holo-\textit{PmCpeB}) were stronger and displayed a more stable fluorescence in presence of \textit{GtCPES} with an emission maximum at 568 nm ($\lambda_{em}=550$ nm). Addition of free 3(Z)-PEB to apo-\textit{PmCpeB} resembled spontaneous attachment as described before (17). These measurements are in agreement with data observed for the native \textit{Prochlorococcus} lyase CpeS (17) and confirm the activity of \textit{GtCPES} in mediating the correct attachment of PEB to $\beta$-Cys82. Interestingly, when we employed an \textit{in vivo} assembled \textit{GtCPES:PEB} complex (obtained by coexpressing \textit{GtCPES} with PEB biosynthesis genes), we obtained a fluorescent product that resembled even better the native holo-\textit{PmCpeB} with a fluorescence emission at 571 nm (data not shown). Our negative control, \textit{PmCpeB\_C82A}, displayed basal interaction with free 3(Z)-PEB visualized by an initial increase but subsequent decrease in fluorescence emission at 615 nm ($\lambda_{em}=550$ nm) (Figure 1C). Employing the PEB-loaded \textit{GtCPES} lyase, no transfer was observed indicated by negligible fluorescence emission at 616 nm ($\lambda_{em}=550$ nm) (Figure 1D). As expected, no covalent attachment was observed in zin-co-lot analysis (Figure 1E). Although we have previously shown that \textit{GtCPES} can also bind the semi reduced intermediate 15,16-DHBV (14), DHBV-loaded lyase did not result in transfer of DHBV to \textit{PmCpeB} (data not shown). As previously shown, DHBV is also not able to spontaneously attach to \textit{PmCpeB} (17).

\textbf{Two glutamate residues are important for PEB binding.} Now that we established the assistance (chaperon) function of \textit{GtCPES} during PEB transfer, we went ahead to perform a rigorous analysis to identify amino acid residues involved in substrate binding and transfer. Despite several approaches for co-crystallization of the lyase with its substrate PEB, no co-crystal structure was obtained. However, one molecule of 1,6-hexanediol was found within the barrel in the original structure giving a hint for the putative substrate binding site and allowing the initial identification of two residues involved in substrate binding (14). We now used the crystal structure and an amino acid sequence alignment of S-type lyases to identify ten individual amino acid residues potentially involved in PEB binding and transfer (Figure 2, Figure S1, (14)). These residues were exchanged to alanine residues using site directed mutagenesis. The following variants were generated: R18A, H21A, N22A, W69A, W75A, E136A, R146A, R148A, S150A and E168A (Figure 2). Residues E136, R146, and R148 are highly conserved among S-type lyases and are in close proximity to the 1,6-hexanediol molecule within the crystal structure of \textit{GtCPES} and likely involved in substrate binding/coordination (14). Most of the other amino acid residues are located within the barrel and have potential influence on PEB binding. W75, which is located at the top of the lyase barrel, almost like a lid over the barrel, has putative influence on the interaction with the apo-protein CpeB (Figure 2). All variants were produced and purified in sufficient amounts and tested for their ability to bind PEB. In addition, thermal shift assays were performed to evaluate the effects of sequence variations on protein integrity and stability (21).

Upon binding to the wt \textit{GtCPES} lyase, the absorbance maximum of PEB is red shifted from 535 nm to 598 nm with a significant increase in intensity with a shoulder at 557 nm (14). Five of the investigated \textit{GtCPES} variants (R18A, N22A, W75A, R148A and S150A) showed almost identical binding spectra with a long wavelength major peak and a shoulder at a shorter wavelength (Table 1). These variants were considered wt-like. The remaining five variants displayed moderate to considerable changes from the wt \textit{GtCPES:PEB} spectrum (Figure 3). The spectrum of variant W69A together with PEB still displayed two absorption maxima. However, what appears as a shoulder in the spectrum of the wt lyase, is a more distinct peak at 549 nm in this variant (Figure 3). The variants H21A and E168A also displayed two absorption peaks when incubated with PEB. However, there appears to be a large proportion of unbound PEB in these samples due to the absence of an increase in extinction coefficient of PEB upon incubation with the variants. In addition, the peak maxima were rather broad suggesting that the chromophore is less stretched, again suggesting a larger proportion of free PEB in the sample (22). In contrast, variants E136A and R146A did not bind PEB as a spectrum almost identical to that of free PEB was observed (Figure 3). The lack of PEB binding of the R146A variant is likely due to (partly) misfolded/aggregated protein. No discrete melting point between 47 and 58 °C was obtained, as contrary to other mutants. The latter was also true for variant H21A. Therefore, both amino acid residues are likely important for structural integrity of the protein. Both variants were not investigated further. Interestingly, all variants that showed changes in their absorption properties with bound PEB, also
displayed fluorescence emission at ~638 nm when excited at 550 nm (Table 1). This is likely due to unspecific interaction of PEB with the lyase variants and differs from the wt protein where no fluorescence upon PEB binding was observed. In conclusion, both glutamate residues, E136 and E168 are strongly involved in PEB binding, while W69 has only minor influence on binding the substrate. H21 and R146 are important for structural integrity.

**A tryptophan residue is important for PEB transfer to PmCpeB.** We next tested whether the generated GtCPES variants that still showed PEB binding were also able to transfer PEB to apo-PmCpeB. Transfer of PEB was monitored using fluorescence spectroscopy as described above. Most of the variants that did show wt-like binding of the substrate PEB, did also display this behavior for the transfer reaction (Figure 4, S2). Interestingly, variant W69A that only showed minor absorption changes upon PEB binding (see paragraph before) did show abnormal transfer behavior that resembled that of a non-lyases mediated spontaneous reaction. Such reactions are indicative of initial increase but subsequent decrease of fluorescence emission at 565 nm with an additional fluorescence emission maxima observed at longer wavelength. Therefore, W69 likely is involved in PEB binding and its exchange leads to a large proportion of unbound PEB, which is then transferred in a spontaneous, unspecific reaction to PmCpeB. The same was observed for variant E168A that already showed only weak to negligible PEB binding. This variant already displayed fluorescence emission when incubated with PEB (Emax= 642 nm) which decreased upon incubation with PmCpeB. Transfer is observed but based on the fluorescence emission, which increased and decreased, it appears that there is a large proportion of spontaneous transfer taking place. Variant S150A showed a normal PEB binding spectrum but reduced transfer indicative of a low fluorescence intensity increase upon incubation with PmCpeB. The most important residue for PEB transfer is a tryptophan located at the upper rim of the barrel. Variant W75A, although still being able to bind PEB like the wt lyase, is not able to facilitate the transfer of PEB to PmCpeB. In order to test whether tight binding and therefore reduced release rates of PEB might cause this observation; binding affinities of PEB to wt GtCPES and the W75A variant were determined. Interestingly, GtCPES_W75A is able to bind PEB more than 10-fold tighter than the wt protein suggesting that this tight binding might prevent the release and transfer of PEB (Table 2).

**A single methionine residue determines substrate specificity.** GtCPES possesses a narrow binding pocket harboring amino acid residues with bulky side chains. As a consequence, GtCPES has a very high substrate specificity towards PEB. Nevertheless it is also able to bind the biosynthetic precursor 15,16-DHBV (14). Binding of the PEB-isomer phycocyanobilin (PCB) was not observed likely due to a rigid double bond between C15-C16 resulting in a missing flexibility of D-ring to fit into the binding pocket (Figure 5C). In order to characterize the molecular determinants of the substrate specificity, we compared both binding pocket and sequence of GtCPES with the PCB-specific S-type lyase CpcS from *Thermosynechococcus elongatus* (*Te* CpcS) (Figure 5A, S1). Based on a comparison of the two available crystal structures of *Te*CpcS and *Gt*CPES, it became obvious that *Te*CpcS possesses a binding pocket with amino acid residues with less bulky side chains (14,23) We therefore went ahead to mutagenize the binding pocket of GtCPES with the aim to widen it. Subsequently, the variant proteins were tested for their ability to bind PCB as well. We started to exchange amino acid residues around position 67 because these residues specifically constrict the binding pocket (Figures 2, 5). Overall, methionine 67 indeed seems to be highly specific for determining the substrate specificity. M67A and M67V variants of GtCPES were still able to bind PEB as indicated by binding spectra of PEB that resembled the wt situation (Table 3). In addition, we tested whether these variants were still able to transfer PEB to PmCpeB. Both methionine variants and the I64A variant displayed specific transfer of 3(Z)-PEB (Figure 6). Interestingly, the M67A variant had the strongest affinity towards PEB of all measured GtCPES variants (Table 2) indicating that the affinity alone does not determine whether the chromophore is released from the binding pocket of the lyase or not. Interestingly, both methionine variants were also able to bind PCB. Incubation with PCB resulted in a shift of the PCB absorption maximum to higher wavelengths (639 nm) together with an increase of extinction reflecting phycocyanin binding (Figure 5). PCB binding furthermore also resulted in a fluorescent complex formation with an emission maximum at 662 nm (Table 3). As both methionine variants displayed the same spectroscopic behavior, all following experiments were only conducted employing the M67A variant.

**GtCPES_M67A binds PCB with high affinity in vitro and in vivo.** With respect to the gained PCB binding ability of the M67A variant, we next wanted to determine the affinity towards PCB. One way to do so is to investigate whether the CPES variant will form a complex with PCB when all necessary
genes are coexpressed in *E. coli* (14,24). Coexpression resulted in intensely colored cells for *Gt*CPES*M67A* with PEB as well as PCB (Figure 7). This complex remained stable through the affinity chromatography indicating strong binding of PCB. The latter was confirmed by determining the binding affinities for 3(Z)-PEB and 3(E)-PCB. The calculated Kd values showed strong binding with 0.044 µM and 7 µM for PEB and PCB, respectively (Table 2). Wild type *Gt*CPES on the other hand only formed an *in vivo* complex with PEB but not PCB ((14) and Figure 7). During the subsequent purification process of *Gt*CPES, the phycobilin remained bound to M67A as monitored by UV/vis-spectroscopy (Figure 7B). Absorption spectra of the elution fraction from affinity chromatography showed binding spectra with maxima similar to *in vitro* binding of 3(E)-PCB and 3(Z)-PEB by *Gt*CPES*M67A* (Table 3). Moreover, binding of PCB resulted in the formation of a fluorescent complex with *Gt*CPES*M67A*. This is in agreement with the complex formed *in vivo* (Table 3). Extinction coefficients for formed complexes with *Gt*CPES*M67A* were determined assuming that all PBP-lyase molecules were loaded with PCB or PEB. Resulting coefficients are 29.56 mM⁻¹ cm⁻¹ (*Gt*CPES*M67A*:PEB) at 600 nm and 15.23 mM⁻¹ cm⁻¹ (*Gt*CPES*M67A*:PCB) at 640 nm in sodium phosphate buffer. Additionally, *in vivo* formed *Gt*CPES*M67A*:PEB complex was also able to transfer loaded PEB to *Pm*CpeB detected by fluorescence spectroscopy (data not shown). Here, emission maxima were similar to *in vitro* formed complex mediated transfer.

**Gt*CPES*M67A transfers PCB to Cys82 of *Pm*CpeB but not to CpcB.** Subsequently, we were interested whether the *Gt*CPES variant M67A exhibited the ability to mediate the transfer of 3(E)-PCB to Cys82 of *Pm*CpeB. At first, spontaneous attachment of 3(E)-PCB was monitored by fluorescence spectroscopy (Figure 8A). Here, fluorescence spectra revealed slow but continuous emission increase over time with maximum wavelengths of 640 nm (*Pm*CpeB), 642 nm (*Pm*CpeB_C82A) corresponding to unspecific interaction of PCB with the apo-PBP. Lyase mediated product formation was detected after adding PBP subunit (*Pm*CpeB, *Pm*CpeB_C82A) to *in vitro* formed complex of *Gt*CPES*M67A* and 3(E)-PCB, as described before. All spectra differed from spontaneous attachment with the emission intensity of the first spectrum after 1 min reaction time representing the fluorescent complex between 3(E)-PCB and *Gt*CPES*M67A* with maximum around 663 nm (Figure 8D and Table 2). Over time, this emission maximum shifted to shorter maxima wavelengths (651 nm) indication transfer of PCB to the apo-protein. Similar fluorescence emission maxima (Emax =647 nm) were observed using CpcS lyase mediated PCB transfer to the CpcB subunit (Figure S2). The control experiment employing CpeB_C82A did not show any changes in absorption or fluorescence and only displayed the specific fluorescence background of *Gt*CPES:PCB (Figure 8B,E). Covalent attachment of PCB to the apo-CpeB was further confirmed using zinc blot analysis (data not shown) (25). Unfortunately, attempts to detect chromopeptides using mass spectrometry failed.

In a final experiment, we asked the question whether the PCB-binding *Gt*CPES*M67A* variant would be able to transfer PCB to the phycoecyanin β-subunit (CpeB) of *Synechococcus* sp. PCC7002. Unfortunately, here the results were not conclusive. The presence of the lyase led to an attachment product that showed fluorescence emission at 659 nm, only ~3-4 nm different to the spontaneous attachment (Figure 8C,F). However, the emission maximum also differed significantly from that mediated by the CpcS lyase, which resulted in a CpeB with a fluorescence emission at 647 nm (Figure S3). We conclude that *Gt*CPES*M67A* is not able to quantitatively transfer PCB to CpeB. Whether small amounts of PCB were correctly transferred to CpeB has to be determined.

**DISCUSSION**

We have previously initially characterized and crystallized the PBP lyase CPES from *G. theta*. PBP lyases are proteins that assist the correct attachment of light harvesting phycobilin pigments to conserved cysteine residues within a phycobiliprotein (15). Once all phycobilins are attached, the individual phycobiliproteins assemble into larger structures, either into phycobilisomes (cyanobacteria and rhodophytes) or into soluble heterodimers (cryptophytes). *Gt*CPES is an S-type PBP lyase and was postulated to attach PEB to Cys82 of the β-subunit CpeB. Although we were not able to show the transfer to the PBP subunit from *G. theta*, a transfer was observed using CpeB from the cyanobacterium *Prochlorococcus marinus* MED4. Both proteins share 55% sequence homology. We therefore conclude that *Gt*CPES is a highly specific lyase for the attachment of 3(Z)-PEB also to β-Cys82 of CpeB of *G. theta*. This β-subunit has in total three PEB molecules bound: an additional one with a double linkage at Cys50,61 and one at Cys155 (6,7). Accordingly, it is reasonable that *Gt*CPES is highly specific and is
Phycobiliprotein lyase substrate specificity

only able to transfer 3(Z)-PEB but not 15,16-DHBV, although binding of both bilins is observed. *Gt*CPES shows a very narrow substrate specificity to the phycobilin it is transferring. This is mainly due to a very confined binding pocket, which is characterized by several bulky amino acid residues that restrict binding to bilins containing a C15-C16 double bond like PCB or phytochromobilin (PΦB) (14). However, S-type lyases from certain cyanobacteria that only possess the phycobilin PCB, i.e. CpcS, appear to have a broader substrate specificity. The CpcS lyase from *Thermosynechococcus elongatus* for instance is an universal lyase being able to bind and transfer PEB, PCB and PΦB (23). Within this current study, we identified several amino acid residues that are important for phycobilin binding, transfer and substrate specificity. PEB binding is primarily mediated by two glutamate residues, E136 and E168. Both residues are facing the interior of the pocket and are likely involved in coordinating two of the pyrrole nitrogens of the bilin. E136 is highly conserved in the family of S-type lyases. W75 is also highly conserved within the S-type lyases and located at the upper rim of the barrel. Here, an involvement in the interaction with the apo-PBP is postulated. Interestingly, the *Gt*CPES_W75A variant had a significantly higher binding affinity to the substrate. We therefore initially postulated that a high affinity retains the bilin within the lyase and therefore prevents the transfer to the apo-protein. However, this is in contrast to the data observed for the *Gt*CPES_M67A variant that displayed an even higher binding affinity towards PEB (0.044 µM for M67A vs. 0.86 µM for W75A) implying that the affinity to the substrate does not determine whether the bilin is transferred or not. We rather postulate that a correct interaction with the target PBP must be present for a correct transfer to occur. This postulate is in conclusion with our data showing that the *Gt*CPES_M67A variant is unable to transfer the bound PCB to the CpcB subunit, but only to the CpeB subunit. There might be smaller structural differences that determine the specific interaction of the CPES lyase with its corresponding (or in our case homolog) apo-protein.

Finally, we identified a crucial amino acid residue for substrate specificity. M67 determines the narrow substrate specificity of PEB-specific S-type lyases. This residue is highly conserved among CpeS lyases and is only substituted by an isoleucine in some cases (Figure S1). Within the PCB specific S-type lyases, this position is occupied by valine. With its shorter side chain, a valine residue at this position widens the pocket and allowing the more rigid PCB to enter the pocket. The same hold true for our M67A and M67V variants. When the long side chain of methionine is modified to either alanine or valine, the substrate specificity is broadened and the protein can bind both, PEB and PCB. We therefore hypothesize a crucial function for amino acid residue at position 67 in determining the substrate specificity of S-type lyases. Overall, our data provide very strong support for a dual function of the lyase both as bilin-selective binder and bilin-prepositioning chaperon for the hand-over to a dedicated target.
EXPERIMENTAL PROCEDURES

Materials
All chemicals were American Chemical Society grade or better unless specified otherwise. Expression vector pCOLAduet-1 was obtained from Novagen; pASK-IBA7+ was from IBA Life Sciences, pET-Duet1 from Merck KgaA, and pGro7 was from TaKaRa. Strep-Tactin®-Sepharose from IBA, and TALON® metal affinity resin from Clontech were used. HPLC-grade acetone, acetonitrile, formic acid, and spectroanalytical grade glycerol were obtained from J.T. Baker. Sep-Pak cartridges were obtained from Waters.

Construction of expression plasmids and site directed mutagenesis
For construction of pCOLA_cpeB a synthetic gene codon optimized for E. coli K12 (GENEius algorithm, MWG Eurofins Operon) encoding the cpeB gene from Prochlorococcus marinus MED4 was PCR amplified with primers (Table 4) encompassing selected recognition sites (EcoRI, HindIII) for cloning into pCOLAduet (Novagen).
For construction of pCOLA_cpcB the plasmid pBS150v_7002_cpcB was used (26) This construct was a gift of W. M. Schlucher (University of New Orleans). The cpcB gene from Synechococcus sp. PCC7002 was PCR amplified (Table 4) with same recognition sites as for cpeB for following cloning into pColaDuet.
The construction of the plasmids pGtCPES, pE136A, pR146A, pR148A, pC149A, pETGtCPES, pTDho1pcyA and pTDho1pebS were described before (14,24,27) All additional site-directed variants of GtCPES and PmCpeB were generated from pGtCPES or pCOLACpeB using the QuikChange® site-directed mutagenesis kit (Stratagene) with help of primers listed in Table 4 (only the forward primer is shown, the reverse primer is the complement, codon changes are underlined). The resulting plasmids were verified by sequencing.

Production and purification of recombinant proteins
GtCPES and all variants as well as CpcS were produced and purified as described before (14,28) PmCpeB and CpcB were produced expressing pCOLAcpeB, pCOLAcpeB_C82A or pCOLAcpeB in E. coli BL21(DE3)-RIL. Cells were grown at 37 °C in LB medium supplemented with 50 µg/ml kanamycin, and for cpeB with 100 mM D-sorbitol and 2.5 mM betaine to OD 578 nm of 0.5 - 0.6. Subsequently, cells were induced by isopropyl-β-D-thiogalactopyranoside (IPTG; 0.5 mM) and incubated overnight at 17 °C. Cells were harvested by centrifugation, resuspended in lysis buffer (50 mM Tris-HCl pH 7.5, 100 mM NaCl, 10% Glycerol) and lysed by two or three passages through LM10 Microfluidizer® High Shear fluid homogenizer (Microfluidics) (cpeB, cpcB, gtCPES constructs) at 19.000 psi or by three times 3 min sonification (50%, KE76, Sonopuls HD6600, Bandelin) (only gtCPES constructs).
PmCpeB, its variant PmCpeB_C82A and CpcB were purified by affinity chromatography using TALON® metal affinity resin (Clontech) and purification was carried out according to the manufacturer’s instructions based on sodium phosphate buffer (60 mM, 300 mM NaCl, pH 7.5). For imidazole removing a dialysis against 100fold volume of sodium phosphate buffer (60 mM sodium phosphate, 300 mM NaCl, pH 7.5) over night at 4 °C (120 rpm) was performed.

Determination of protein concentration
Purified proteins were concentrated using Amicon® Ultra-4 Ultracel®-10K (MWCO: 10.000 Da; Merck). Protein concentrations were quantified using the calculated molar extinction coefficient ε 280 (29).

Coproduction of phycobilins and PBP-lyases
Heterologous coproduction of GtCPES or variant GtCPES_M67A and PEB biosynthesis enzymes was performed in E. coli BL21(DE3) containing pET-constructs of gtceS and gtceS_M67A (pET_gtCPES; pET_M67A) and pho1pebS or pTDho1pcyA. Cultures were grown in LB medium supplemented with 100 mM sorbitol and 2.5 mM betaine at 37 °C, 100 rpm to an OD 578 nm of 0.6 prior to induction with 0.1 mM IPTG and incubated overnight at 17 °C (~16 h). After cell harvesting by centrifugation, cells were resuspended in lysis buffer (60 mM sodium phosphate, 300 mM NaCl, pH 7.5), and disrupted by pressure based homogenizer (LM10 Microfluidizer®, Microfluidics), preferably, or sonification. Purification process was described before (14)

Determination of extinction coefficient for PBP-lyase:phycobilin complex
Extinction coefficients were determined for in vivo produced and purified PBP lyase:phycobilin-complexes. Here, 30 µM of PEB or PCB were added to 20 µM of GtCPES or variant GtCPES_M67A,
total volume was 550 µl. Phycobilin excess was removed by gel filtration (PD Minitrap G25 Sephadex column, GE healthcare) and performed according to the manufacturer’s instructions with spin protocol based upon centrifugation (3000 rpm, 2 min). PBP-lyase concentration was determined by UV/vis-spectroscopy between all steps. Assuming that all PBP-lyase molecules were loaded with phycobilins, extinction coefficient for PBP-lyase: PEB complex at 600 nm and PBP-lyase:PCB complex at 640 nm were calculated with aid of Lambert-Beer law.

Determination of extinction coefficient for that complex (ε600 = 27.79 mM⁻¹ cm⁻¹) enabled the calculation of complex bound protein concentration under the assumption that all GtCPES molecules were present in complex with PEB.

**Thermal shift experiments**
The temperature of the unfolding transition midpoint, for excluding effect due to denaturation of GtCPES variants, was determined in thermal shift experiments (30,31) For determination fluorescent protein stain SYPRO® Orange (5000x, Sigma-Aldrich) and CFX Connect Real-Time PCR Detection system (Bio-Rad) were used. Microplates were obtained from Biozym. With exclusion of light 50 µl total volume composed of 5 µl protein (2 mg ml⁻¹), 5 µl SYPRO® Orange (100x) and sodium phosphate buffer (60 mM sodium phosphate, 300 mM NaCl, pH 7.5) were tested in triplicate. Controls were conducted without protein component. Melting temperatures were determined automatically by software CFX Maestro.

**Preparation of phycobilins**
PCB (3(E)-PCB) was isolated from Spirulina cells as described previously (32). Production, purification and preparation of PEB were described before (14) but PEB was concentrated by Advance Alpha 2–4 LSCplus freeze dryer (Christ). The vacuum was set to 0.04 mbar with ice condenser set at -40 °C. Phycobilins were resuspended in an appropriate amount of DMSO before use. The isolated phycobilins were analyzed in terms of isomers via HPLC using Luna 5µ C18 column (Phenomenex) and stored at -20 °C on silicagel orange in the dark. If isomer ratio showed strong excess of E or Z-isomer for PEB or PCB, concentrations were determined using ε571: 46.9 mM⁻¹ cm⁻¹ (3(E)-PEB) in MeOH/5% HCl and ε685: 37.15 mM⁻¹ cm⁻¹ (3(E)-PCB) in MeOH/2.5% HCl (33). Due to the absence of a reported extinction coefficient for 3(Z)-PEB ε571 of the related 3(E)-PEB was used. 3(Z)-PEB was directly applied in experiments without separation of isomers and further purification because of Z-isomer excess.

**Phycobilin binding and transfer by GtCPES variants**
For binding studies excess of GtCPES or GtCPES variant (20 µM), respectively, and phycobilin (5 µM) were mixed in sodium phosphate buffer, pH 7.5. Absorption spectra (Agilent 8453 UV-visible spectrophotometer) and in case of transfer studies fluorescence emission spectra (Series 2, FA-256, Aminco Bowman) of the mixture and free phycobilins in the same buffer were detected. Measurements were done at 60% sensitivity and 975 V. For transfer studies 20 µM PmCpeB, PmCpeB C82A or CpcB was added 2 min after GtCPES (variant):phycobilin-complex formation and fluorescence emission was measured after 1, 5, 10, 15, 20, 30 and 45 min (Exc. 550 nm PEB; Exc. 600/620 nm PCB). All samples were incubated and measured at room temperature. After an incubation time of 45 min all sampled were prepared for and separated by SDS-PAGE on a 12.5% gel. Proteins were transferred to PVDF membrane that was subsequently incubated in 1.3 M zinc acetate for 1 h at 4°C, and afterward zinc-enhanced fluorescence of proteins covalently associated with bilins was visualized under UV light (312 nm) (25)

**Spectroscopic analysis of phycobilin binding**
Increasing amounts of GtCPES (variant) (2-15 µM) were added to 10 µM (final concentration) 3(Z)-PEB or 3(E)-PCB in a final volume of 200 µl of sodium phosphate buffer, pH 7.5, under exclusion from light. After incubation for 2 min absorbance spectra were recorded using Agilent 8453 UV-visible spectrophotometer. The method was described by Frankenberg and Lagarias (34) and was adapted, here. Analysis of spectra was performed using Microsoft Excel.

To obtain phycobiliprotein lyase dissociation constants, absorbance differences (ΔA) at the λmax of each complex were plotted as a function of GtCPES concentration with aid of Microsoft Excel. Dissociation constants KD were obtained by fitting the parameters of the equation (1) to the experimental data, where [Ptotal] describes the total concentration of GtCPES, [L-total] of phycobilin and [PL] is the concentration of complex.

\[
K_D = \frac{([P_{total}] - [PL])([L_{total}] - [PL])}{[PL]}
\] (1)
Data and image processing
Spectra were generated in Origin® data analysis/graphic software, models of protein structure or ligand binding pocket were edited with aid of published GiCPES crystal structure (PDB code 4TQ2) by Pymol (35). Cavity model of GiCPES binding pocket was displayed by surface cavity mode of Pymol.

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CONFLICT OF INTEREST
The authors declare that they have no conflicts of interest with the contents of this article.

AUTHORS CONTRIBUTION
NT, KEO, AP, EH and NFD designed the research, NT, KEO, MA performed the experiments, AP helped analyzing data, NT, KEO, NFD analyzed data and wrote the manuscript. All authors approved the manuscript.
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TABLE 1. Binding and transfer of 3(Z)-PEB after incubation with different GrCPES variants compared to the wild type.
Melting points (in °C) of proteins determined in triplicate and absorbance/emission maxima of binding/transfer are given. Complex $E_{\text{max}}$; detected emission maximum after adding PBP-lyase to free PEB. $E_{\text{max}}$: emission maximum after 45 min incubation time of GrCPES (variant):bilin with $Pm$CpeB. Excitation wavelength was 550 nm.

| GrCPES (variant) | Tm [°C] | Binding PEB | $\lambda_{\text{max}}$ [nm] | Complex $E_{\text{max}}$ [nm] | Transfer PEB | $E_{\text{max}}$ [nm] |
|------------------|---------|-------------|-----------------|---------------------------|-------------|-----------------|
| WT               | 56.0 ± 0.0 | yes      | 554, 600       | ---                      | yes         | 568             |
| R18A             | 48.0 ± 0.0 | yes      | 552, 593       | ---                      | (yes)$^a$   | 567             |
| N22A             | 55.0 ± 0.0 | yes      | 555, 598       | ---                      | yes$^a$     | 568             |
| W69A             | 48.0 ± 0.0 | yes      | 549, 596       | 637                      | no          | 565             |
| W75A             | 58.0 ± 0.0 | yes      | 554, 599       | ---                      | no          | n.d.            |
| E136A            | 47.5 ± 0.0 | no       | 539            | 638                      | no          | 565             |
| R148A            | 56.0 ± 0.0 | yes      | 552, 600       | ---                      | yes         | 566             |
| S150A            | 48.2 ± 0.3 | yes      | 554, 600       | ---                      | (yes)$^a$   | 568             |
| E168A            | 49.0 ± 0.0 | (no)$^b$ | 541, 597       | 642                      | (no)        | 568             |

$^a$ Less increase of fluorescence emission compared to wtGrCPES. $^b$ Binding spectra differs from wtGrCPES:PEB spectra.

TABLE 2. Binding affinities for complex formation with PEB and PCB.
Phycobilin binding analysis was performed as described were tested by titration described in the experimental procedures. $K_D$ values were calculated from equation (1) by using solver function of Microsoft Excel. Results were averaged over two measurements. n.b., no binding.

| GrCPES (variant) | PEB $K_D$ [µM] | PCB $K_D$ [µM] |
|------------------|----------------|----------------|
| WT               | 14.2 ± 2.6     | n.b.           |
| M67A             | 0.044 ± 0.003  | 7.0 ± 0.4      |
| W75A             | 0.86 ± 0.06    | ---            |
TABLE 3. *In vitro* binding and transfer to *Pm*CpeB of PEB and PCB after incubation with different *Gt*CPEs variants.

Melting points (in °C) of proteins determined in triplicate and absorption/emission maxima of binding/transfer are presented. (Exc. 550 nm PEB; Exc. 600/620 nm PCB)

| GtCPEs (variant) | Tm [°C] | 3(Z)-PEB | 3(E)-PCB |
|------------------|---------|----------|----------|
|                  |         | Binding  | A_max [nm] | E_max [nm] | Transfer | E_max [nm] | Binding | A_max [nm] | E_max [nm] | Transfer | E_max [nm] |
| I65A             | 47.5 ± 0.5 | yes | 555, 599 | --- | yes | 568 | no | 616 | --- | no | 640 |
| M67A             | 53.3 ± 0.3 | yes | 553, 598 | 635b | yes | 569 | yes | 639 | 662 | yes | 651 |
| M67V             | 52.3 ± 0.4 | yes | 555, 597 | --- | yes | 567 | yes | 638 | 663 | no | 640 |

*a*, Absorbance spectrum differs from wtGtCPEs: bilin concerning extinction. *b*, fluorescence only detected for *in vivo* binding. ND means not determined, n.d., not detectable.

TABLE 4. Constructed expression plasmids. Forward primers for site-directed mutagenesis are shown, reverse primer is the complement. Changed codons are underlined.

| Plasmid | Host vector | Recognition sites | Primers (5’ - 3’) |
|---------|-------------|-------------------|------------------|
| pCOLA_cpeB | pCOLADuet-1 | EcoRI | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pCOLA_cpcB | pCOLADuet | HindIII | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pR18A | pGtCPEs | EcoRI | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pH22A | pGtCPEs | HindIII | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pN23A | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pI65A | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pM67A | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pM67V | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pW69A | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pW75A | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pE136D | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pS150A | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pE168A | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pI65A_M67A | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pI65A_M67V | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pI65A_M67V_L89P | pGtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |
| pET_M67A | pET-GtCPEs | GCCGAATTCTATGACAGTTTCAAAAGGTAAT |

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FIGURES

FIGURE 1. *Gt*CPES mediated and spontaneous PEB chromophorylation of *Pm*CpeB or *Pm*CpeB_C82A and covalent binding formation of *Pm*CpeB with PCB or PEB visualized by zinc-enhanced fluorescence. The apo-PBPs *Pm*CpeB or *Pm*CpeB_C82A were incubated with PEB (A, C) or wt*Gt*CPES:PEB complexes (B, D). Emission spectra (λ<sub>ex</sub> = 550 nm) of 1, 5, 10, 15, 20, 30, and 45 min after addition of *Pm*CpeB or *Pm*CpeB_C82A, respectively, were detected. First and last spectra are shown by bold lines. Emission maxima are given and the course of fluorescence emission changes is indicated by arrows. After spontaneous and PBP-lyase mediated transfer studies of PEB and PCB, samples were used for SDS-PAGE and blotted onto PVDF membrane (E). Membrane was incubated for 1 h at 4 °C in 1.3 M zinc acetate and detected under UV-light (312 nm). Fluorescence signals based on zinc-phycobilin complexation describe covalent binding of phycobilins to *Pm*CpeB (with his-tag: 21.4 kDa). Absent signals indicate missing binding.
FIGURE 2. Location of amino acid residues of GtCPES investigated within this study. The structure of GtCPES (PDB code 4TQ2) is shown in cartoon representation. Residues discussed in the main text are shown as sticks with color coding representing the effect of a site specific exchange on GtCPES function: white – no change; green – structural defect; purple – bilin binding; pink – bilin transfer and selectivity. Figure generated with Pymol (35).
FIGURE 3. UV-Vis spectra of 3(Z)-PEB bound to GtCPES and variants. Free PEB spectra are shown as solid lines, the spectra of lyase variants with PEB are shown as dotted lines. Peak maxima are given and the respective amino acid exchange.
FIGURE 4. Transfer of 3(Z)-PEB to PmCpeB by GtCPES variants. Apo-PmCpeB was incubated with GtCPES variants preincubated with PEB and. Emission spectra ($\lambda_{ex} = 550$ nm) at 1, 5, 10, 15, 20, 30, and 45 min after addition of PmCpeB were taken. First and last spectra are shown by bold lines. Emission maxima are given and the course of fluorescence emission changes is indicated by arrows.
FIGURE 5. Transformation of PEB specific lyase into universal lyase. Comparison of a cavity model of the phycobilin binding pocket of GtCPES (A) and GtCPES_M67A (B). Model was generated using Pymol with aid of surface cavity mode (35). C. Absorption spectra of free 3(E)-PCB (solid line) compared with absorption spectra after addition of wt GtCPES (left panel) and GtCPES_M67A (right panel) (dashed line). Absorbance maxima are indicated.
FIGURE 6. Transfer of 3(Z)-PEB to PmCpeB by GtCPES variants with larger binding pocket. Apo-PmCpeB was incubated with GtCPES variants preincubated with PEB. Emission spectra ($\lambda_{ex} = 550$ nm) at 1, 5, 10, 15, 20, 30, and 45 min after addition of PmCpeB were taken. First and last spectra are shown by bold lines. Emission maxima are given and the course of fluorescence emission changes is indicated by arrows.
FIGURE 7. In vivo complex formation of GiCPES_M67A with PCB and PEB. Coproduction of wtGiCPES and variant GiCPES_M67A with synthesis enzymes for PCB (Ho1, PcyA) and PEB (Ho1, PebS) were conducted in E. coli BL21(DE3). Formed GiCPES_M67A:bilin complexes were purified by affinity chromatography and detected by color formation (A) and UV/vis-spectroscopy (B; GiCPES_M67A:PCB). Absorption maxima in elution fraction are labelled. Coproduction of wtGiCPES, Ho1 and PcyA shows no color (control).
**FIGURE 8.** Spontaneous and *GtCPES_M67A* mediated PCB chromophorylation of *PmCpeB, PmCpeB_C82A* and CpcB. The apo-PBPs *PmCpeB, PmCpeB_C82A* and CpcB were incubated with PCB (- no lyase) or *GtCPES_M67A*:PCB complexes. Emission spectra ($\lambda_{ex} = 600$ nm (A,B,D,E); $\lambda_{ex} = 620$ nm (C and F)) of 1, 5, 10, 15, 20, 30, and 45 min after addition of apo-PBPs were detected. First and last spectra are shown by bold lines. Emission maxima are given and the course of fluorescence emission changes is indicated by arrows.