

# Crystal Structure of a Phycourobilin-Containing Phycoerythrin at 1.90-Å Resolution<sup>1</sup>

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## INTRODUCTION

The structure of R-phycoerythrin (R-PE) from the red alga *Griffithsia monilis* was solved at 1.90-Å resolution by molecular replacement, using the atomic coordinates of cyanobacterial phycocyanin from *Fremyella diplosiphon* as a model. The crystallographic R factor for the final model is 17.5% ( $R_{\text{free}}$  22.7%) for reflections in the range 100–1.90 Å. The model consists of an  $(\alpha\beta)_2$  dimer with an internal noncrystallographic dyad and a fragment of the  $\gamma$ -polypeptide. The  $\alpha$ -polypeptide (164 amino acid residues) has two covalently bound phycoerythroblins at positions  $\alpha 82$  and  $\alpha 139$ . The  $\beta$ -polypeptide (177 residues) has two phycoerythroblins bound to residues  $\beta 82$  and  $\beta 158$  and one phycourobilin covalently attached to rings A and D at residues  $\beta 50$  and  $\beta 61$ , respectively. The electron density of the  $\gamma$ -polypeptide is mostly averaged out by threefold crystallographic symmetry, but a dipeptide (Gly-Tyr) and one single Tyr could be modeled. These two tyrosine residues of the  $\gamma$ -polypeptide are in close proximity to the phycoerythroblins at position  $\beta 82$  of two symmetry-related  $\beta$ -polypeptides and are related by the same noncrystallographic dyad as the  $(\alpha\beta)_2$  dimer. Possible energy transfer pathways are discussed briefly. © 1999 Academic Press

**Key Words:** light-harvesting complexes; red algae; phycobiliproteins; protein structure; molecular replacement.

The process of photosynthesis converts light energy to chemical energy. For the absorption of light, cyanobacteria and red algae use water-soluble light-harvesting complexes, called phycobilisomes, which are attached to the stromal side of the thylakoid membrane. They have a molecular mass of approximately  $7\text{--}15 \times 10^6$  Da and transfer the absorbed energy with an efficiency of over 95% (Gantt and Lipschultz, 1973; Sauer, 1975; Glazer, 1989) to the photosynthetic reaction center.

Phycobilisomes are made up of two structural subunits: a core complex, located close to the reaction center, and rod-like segments that are attached to this core. Both subunits contain different phycobiliproteins in the form of stacked single discs. The various phycobiliproteins differ predominantly in the number and nature of the pigments they carry and therefore in their absorption spectra. Their arrangement is such that those absorbing in the blue/green region of the spectrum are located at the end of the rods, while those absorbing in the red region are located in the core complex.

The core complex consists predominantly of allophycocyanin (APC),<sup>4</sup> with an absorption maximum  $\lambda_{\text{max}}$  at approximately 650 nm, while the major constituents of the rods are phycocyanin (PC), with  $\lambda_{\text{max}}$  at 615–620 nm, and phycoerythrin (PE), with  $\lambda_{\text{max}}$  at 495–565 nm (MacColl and Guard-Friar, 1987).

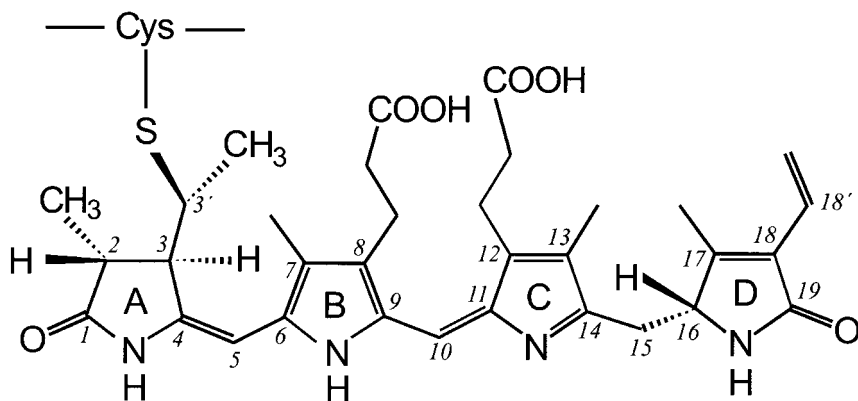
The polypeptide composition of cyanobacterial and algal phycobiliproteins is very similar. Two polypeptides of about 15 ( $\alpha$ ) and 17 kDa ( $\beta$ ) form an  $(\alpha\beta)$  heterodimer, which is the basic structural unit in most phycobiliproteins. Each polypeptide of the basic  $(\alpha\beta)$  heterodimer includes eight  $\alpha$ -helices (X, Y, A, B, E, F, G, and H). Their nomenclature points to the structural similarity of helices A to H to the globin fold (Schirmer *et al.*, 1985). The positions of these helices are so conserved within the three-dimen-

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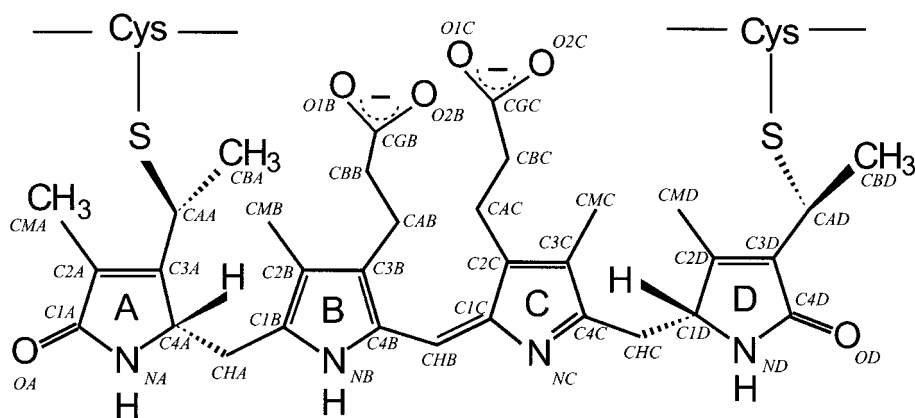
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<sup>4</sup> Abbreviations used: R-PE, R-phycoerythrin; B-PE, B-phycoerythrin; C-PC, C-phycocyanin; APC, allophycocyanin; PC, phycocyanin; PE, phycoerythrin; PCB, phycocyanobilin; PXB, phycobiliviolin; PEB, phycoerythrobin; PUB, phycourobilin; rms, root-mean-square; NCS, noncrystallographic symmetry.



PEB (Phycoerythrobilin)



PUB (Phycourobilin, doubly linked)

**FIG. 1.** Chemical structure of the bilin chromophores (top) PEB and (bottom) doubly linked PUB. The nomenclature of the atoms is indicated.

sional structures that most of the more recent phycobiliprotein structure determinations could be done using the method of molecular replacement. The  $(\alpha\beta)$  heterodimer is arranged around a crystallographic threefold axis to form an  $(\alpha\beta)_3$  trimer. With few exceptions (Schirmer *et al.*, 1985, 1987) this trimer is then related by a twofold noncrystallographic symmetry (NCS) to form the  $(\alpha\beta)_6$  hexamer. In addition, a third polypeptide of varying molecular mass is present and functions as a so-called linker protein (for reviews see Glazer, 1985, 1989; Huber, 1989; Grossman *et al.*, 1993). In the case of R-phycoerythrin (R-PE) and B-phycoerythrin (B-PE), this linker protein ( $\gamma$ -polypeptide) is pigmented, has a molecular mass of about 30 kDa, and is tightly bound to the  $\alpha$ - and  $\beta$ -polypeptides. This yields a stoichiometry of

$(\alpha\beta)_6\gamma$  and a molecular mass of about 240 kDa for both phycobiliproteins.

In R-PE and B-PE all pigments are open-chain tetrapyrrole chromophores (bilins), which are covalently bound to cysteines (Glazer *et al.*, 1976). There are, however, differences in pigmentation. B-PE carries five phycoerythrobilins (PEBs) per  $(\alpha\beta)$ , while in R-PE two phycoerythrobilins (see Fig. 1, top) are attached to the  $\alpha$ -polypeptide, and two phycoerythrobilins and one doubly linked phycourobilin (PUB, see Fig. 1, bottom) are bound to the  $\beta$ -polypeptide. The pigment composition of the  $\gamma$ -polypeptide varies. For B-PE four bilins (two PEBs and two PUBs) are described (Glazer, 1985; Ficner *et al.*, 1992) while for R-PE the amount of pigments varies from four to five bilins (PEB and PUB), depending on the organism.

Up to three different  $\gamma$ -polypeptides are observed for R-PE of some species (Apt *et al.*, 1993; Stadnichuk *et al.*, 1993).

In the past few years the structures of various phycobiliproteins could be solved, showing the conformation of the bound chromophores within the protein (for review see Betz, 1997). The conformation of phycocyanobilin (PCB) was determined by Schirmer *et al.* (1985, 1986, 1987), Duerring *et al.* (1991), and Brejc *et al.* (1995). The conformation of phycobiliviolin (PXB), phycoerythrobilin, and phycourobilin was determined by Duerring *et al.* (1990); Ficner *et al.* (1992), Ficner and Huber (1993); and Chang *et al.* (1996); respectively.

In this paper we report the solution of the structure by molecular replacement and refinement of R-PE from the red alga *Griffithsia monilis* at 1.90-Å resolution. The model consists of an  $\alpha$ -polypeptide carrying two PEBs at positions  $\alpha 82$  and  $\alpha 139$  and of a  $\beta$ -polypeptide carrying two PEBs at positions  $\beta 82$  and  $\beta 158$  and one PUB doubly-linked to residues  $\beta 50$  and  $\beta 61$ . The two polypeptides  $\alpha$  and  $\beta$  are arranged around a NCS dyad to form an  $(\alpha\beta)_2$  dimer as an asymmetric unit. In the following, residues on the NCS-related  $\alpha$ -polypeptides of the asymmetric unit are designated A and K, respectively. Similarly, residues on the  $\beta$ -polypeptide are named B and L, respectively.

Although the structure of R-PE from *G. monilis* is expected to be closely related to the structure of B-PE from *Porphyridium sordidum* and that of R-PE from *Polysiphonia urceolata*, it is of interest because it was refined to high resolution using the native sequence of *G. monilis* R-PE. For the refinement of the two other structures the sequences of *Porphyridium cruentum* and *Polysiphonia boldii* were used, respectively. Moreover, we were able to model three residues into the electron density that arises from the  $\gamma$ -polypeptide.

## MATERIALS AND METHODS

*Crystallization, data collection, and crystallographic computing.* The purification, crystallization, and preliminary crystallographic characterization of R-PE from the red alga *G. monilis* have been described elsewhere (Ritter *et al.*, 1997). For the solution of the structure at a resolution of 2.3 Å we collected data from only one crystal (data set A) with a Stoe Imaging Plate detector system (Stoe, Darmstadt, Germany) at 14°C. The X-ray source (CuK $\alpha$ ) was a rotating anode (Stoe) operating at 40 kV, 100 mA. For data reduction and determination of the space group ( $R3$  with  $a = b = 187.35$  Å,  $c = 59.31$  Å) the XDS program (Kabsch, 1993) was used. High-resolution data were collected at the EMBL beamline X11 at the Deutsches Elektronensynchrotron (DESY) in Hamburg, Germany, at 17°C, using a MarResearch Image Plate detector. Two crystals were used to obtain a complete data set extending to 1.82-Å resolution (data set B). Table I shows the statistics for both data sets. No indications for twinning, which is sometimes observed for crystals of phycobiliproteins, were found. All crystallographic computations, including molecular replacement, were carried out with X-PLOR 3.1 and 3.851 (Brünger, 1992, 1997).

*Sequence determination, model building, and refinement.* The sequence was obtained by genetic methods and was deposited with the EMBL database (Accession No. Z98528). An alignment with the sequences of R-PE from *Pol. boldii* (Roell and Morse, 1993), B-PE from *Por. cruentum* (Sidler *et al.*, 1989), and C-PC from *Fremyella diplosiphon* (Mazel *et al.*, 1988) is shown in Table II (for review see Apt *et al.*, 1995). The coordinates of the 1CPC model (Duerring *et al.*, 1991) were taken from the Brookhaven database and used for molecular replacement. Insertions and deletions relative to 1CPC were modeled with "O" (Jones *et al.*, 1991). For refinement, molecular dynamics and conventional energy restrained least-squares refinement procedures on positional parameters and  $B$  factors were used and water molecules were integrated into the model. A bulk solvent correction was included in the refinement procedure. The free  $R$  factor (Brünger, 1997) was used to optimize the refinement strategy.

## RESULTS

### Structure Solution by Molecular Replacement

As the unit cell parameters of R-PE resemble those of 1CPC, a similar packing was expected. This assumption was confirmed by a cross-rotation search. The  $(\alpha\beta)_2$  dimer of 1CPC was placed into a triclinic unit cell ( $a = 155$  Å,  $b = 90$  Å,  $c = 100$  Å,  $\alpha = 90^\circ$ ,  $\beta = 90^\circ$ ,  $\gamma = 90^\circ$ ) and its Patterson was generated with reflections from 15 to 4 Å. The 1000 highest Patterson vectors were selected within the radius 45–5 Å. The cross-rotation function was calculated in  $2.5^\circ$  steps ( $0 \leq \Theta_+ < 720^\circ$ ,  $0 \leq \Theta_2 \leq 180^\circ$ ,  $0 \leq \Theta_- \leq 120^\circ$ ; see Rao *et al.*, 1980) in Lattman (1972) angle space and gave two clear maxima of equal height at Eulerian angles  $-40^\circ$ ,  $0^\circ$ ,  $0^\circ$  and  $90^\circ$ ,  $180^\circ$ ,  $0^\circ$ . These maxima are related by a  $180^\circ$  rotation and correspond to the internal twofold symmetry of the model dimer. The position of this twofold axis was confirmed by a self-rotation function with reflections within the range 15–4 Å. The highest noise peak of the rotation function was at 95% of the height of the correct solutions but displayed a relative height of only 30% after Patterson correlation calculation.

As we anticipated from the packing constraints, the position corresponding to the highest peak ( $18\sigma$ ) of the translation function showed that the "hexamer" axis coincides with the crystallographic threefold axis. Placing the model into the unit cell of the crystal yielded an initial  $R$  factor of 48.2% (6–3 Å).

TABLE I

Data Statistics for the Observed Data Collected with the Stoe Image Plate (A) and at the EMBL Beamline X11 in Hamburg (B)

Data set	Resolution (Å)	Reflections	Unique reflections	$I_{\text{unique}} \geq 3\sigma$ (%)	Completeness (%)	$R_{\text{int}}^a$ (%)
A	10.00–2.30	62 202	33 603	75.8	96.0	6.9
	2.40–2.30	5 388	3 093	47.0	44.8	21.9
B	15.00–1.82	131 195	67 598	59.5	96.8	5.9
	2.00–1.82	28 679	15 857	21.1	92.0	31.3

$$^a R_{\text{int}} = \sum |I(h) - \langle I(h) \rangle| / \sum I(h).$$

TABLE II

Sequence Alignment between  $\alpha$ - and  $\beta$ -Polypeptides of R-PE from *Griffithsia monilis*, R-PE from *Polysiphonia boldii*, B-PE from *Porphyridium cruentum*, and C-PC from *Fremyella diplosiphon*

		(a) $\alpha$ -Polypeptide										
		10	20	30	40	50	60	70	80			
Sequence number:												
R-PE <i>G. monilis</i>		MKSVITTTISAADAAGRFPS	SSDLESIQGNIQRAAA	LEAAQKLSGNHEAVV	KEAGDACFAKYSYL	--KNAGEAGDSPEKIN						
R-PE <i>Pol. boldii</i>		.....Y..T...Q.V.....	.....E..GS.....	.....S..G...--P...ENQ....								
B-PE <i>Por. cruentum</i>		.....V.....N.....	.....S.....E..A.....	.....A...--P...ENQ....								
C-PC <i>F. diplosiphon</i>		..TPL.EAVA...SQ...L..	TEIQTAF.RFRQ.S.S.A.	.KA.TEKASSLASG.AN.	VYS.FP.TTSQ.GPNFASTQTGKD							
Secondary structure		hhhhhhhhhhht	hhhhhhhhhhhhhhhhhh	hhhhhhhhhhhhhhhhhh	hhhhhhhhhhhhhhhhhh	hhhhhhhhhhhhhhhhhh	hhh-h	tt	tt	hhhh		
		└helix X┐	└helix Y┐	└helix A┐	└helix B┐	└helix C┐						
		PEB $\alpha$ 82								PEB $\alpha$ 139		
Sequence number:		*	90	100	110	120	130	*	150	160		
R-PE <i>G. monilis</i>		KCYRDIDHYMRLINYS	LVVGGTGPVDEWGI	AGSREVVYRALN	LPGSAYIAAFTFR	DRDLRCVPRDMSSQ	AGVEFTSALDYVI					
R-PE <i>Pol. boldii</i>		.....T.....L.....	.....A.....T....	SA.....V.....I.....	.....CT....L.							
B-PE <i>Por. cruentum</i>		.....V.....V..D.....	.....L.....A.....	.....T.....T...V.SIAY.....	.....SAY...L.							
C-PC <i>F. diplosiphon</i>		..V...GY.L.MVT.C.....	..L.DYL.G.IA.IN.TFD.	SP.W.VE.LKYIKANHGL---	GDPA..AN.YI..A.							
Secondary structure		hhhhhhhhhhhhhhhhht	hhhhhhhhhhhhhhhhht	hhhhhhhhhhhhhhhh	hhhh	hhhhhhhhhhhhhhhh	hhhh	hhhhhhhhhhhhhhhh	hhhh	hhhhhhhhhhhhhhhh		
		└helix E┐	└helix F┐	└helix F┐	└helix G┐	└helix H┐	└helix H┐	└helix H┐	└helix H┐	└helix H┐		
Sequence number:		164										
R-PE <i>G. monilis</i>		NSLC	164									
R-PE <i>Pol. boldii</i>		...S	164									
B-PE <i>Por. cruentum</i>		.A.S	164									
C-PC <i>F. diplosiphon</i>		.A.S	162									
Secondary structure		htt										
		└┐										
		(b) $\beta$ -Polypeptide										
		10	20	30	40	PUB $\beta$ 50/61		70	80			
Sequence number:												
R-PE <i>G. monilis</i>		MLDAFSRVVVTSDAKA	AVVGGSDLQSLKSF	INDGNKRLDAVNY	IVSNASCIVSDAV	SGMICENPGLIAP	GGNCYTNRMA-					
R-PE <i>Poly. boldii</i>		.....N..S.....S.....	.....A..T.....S.....	.....I.....T.....								
B-PE <i>Por. cruentum</i>		.....N.....N.....	.....A.....A.....	.....S..A.....M.....	.....S.....							
C-PC <i>F. diplosiphon</i>		....AK...SQA...RGE.	LS...QIDA.SALVA....	M.V..R.TG.S.T..AN.	ARSLFA.Q.Q.....	A..S....-						
Secondary structure		hhhhhhhhhhhttt	hhhhhhhhhhhhhhhh	hhhhhhhhhhhhhhhh	hhhhhhhhhhhhhhhh	hhhhhhhhhhhhhhhh	hhhh	tt	tt	hhhh		
		└helix X┐	└helix Y┐	└helix A┐	└helix B┐	└helix C┐						
		PEB $\beta$ 82								PEB $\beta$ 158		
Sequence number:		*	90	100	110	120	130	140	150	*		
R-PE <i>G. monilis</i>		ACLDRGEIILRYVSY	ALLAGDSSVLDD	RCLNGLKETYIAL	GVPTASSSRAVS	IMKATATAFITNT	ASGRKVEVAAGD	CQA				
R-PE <i>Pol. boldii</i>		.....A...E.....	.....N.TV.....	A.VC..S...Q.....	I E...S.							
B-PE <i>Por. cruentum</i>		.....A...E.....	.....N.I.....	Q.V.....T.E..MSF.....	TS							
C-PC <i>F. diplosiphon</i>		...M.....T...F...A.....	.....L...T.GS.VAVG.	QK..EA.L.IAGD.N---	GITR...AS							
Secondary structure		hhhhhhhhhhhhhhhhht	hhhhhhhhhhhhhhhh	hhhhhhhhhhhhhhhh	hhhhhhhhhhhhhhhh	tt	hh					
		└helix E┐	└helix F┐	└helix F┐	└helix G┐	└helix G┐						
Sequence number:		170	177									
R-PE <i>G. monilis</i>		LQAEAAASYFDKVG	SSID	177								
R-PE <i>Pol. boldii</i>		.AS.V...C.R.VAAVS	177									
B-PE <i>Por. cruentum</i>		.AS.V....R..AA.S	177									
C-PC <i>F. diplosiphon</i>		.M..V.....AA.AVA	172									
Secondary structure		hhhhhhhhhhhhhhhh										
		└helix H┐										

Note. Dots represent identical amino acids, and deletions are marked by dashes; secondary structure elements were determined using DSSP (Kabsch and Sander, 1983). **h**,  $3_{10}$ -helix; **h**,  $\alpha$ -helix; **t**, turn. The nomenclature of the helices is according to Schirmer *et al.* (1985, 1987). An asterisk marks the position of the chromophores, \*==\* indicates a doubly bound chromophore. Sequence numbers refer to *G. monilis*.

### Model Building and Refinement

The asymmetric unit of R-PE from *G. monilis* contains an  $(\alpha\beta)_2$  dimer (ABKL) and on average  $1/3$  of the  $\gamma$ -polypeptide, yielding a stoichiometry of  $(\alpha\beta)_{2\gamma 1/3}$ . Before the first cycle of refinement, all side chains of the 1CPC model were truncated to yield a poly-Ala(Gly) sequence and the chromophores were omitted. After completion of the X-PLOR refinement protocol (Brünger, 1997), a  $(F_{\text{obs}} - F_{\text{calc}})$  map and a  $(2F_{\text{obs}} - F_{\text{calc}})$  map were inspected and used for inserting a tentative amino acid sequence into the model.

As the amino acid sequence of *G. monilis* R-PE was initially not available, we used the sequence of *Pol. boldii* R-PE (Roell and Morse, 1993) for structure determination.

For refinement at high resolution the sequence of *G. monilis* R-PE was used and the resolution was increased to 1.90 Å in a stepwise fashion. After inclusion of the majority of the water molecules, a bulk solvent correction was introduced, which resulted in an improvement of the value of  $R_{\text{free}}$  of approximately 1%. At this stage three residues of the

**TABLE III**  
Parameters of the Refined R-PE Model

Number of nonhydrogen atoms	5855
Number of chromophore atoms	430
Number of solvent molecules	333
$R_{\text{work}}$ (%)	17.5
$R_{\text{free}}$ (%)	22.7
Standard deviations from ideal values	
Bond length (Å)	0.004
Bond angles (°)	1.1
Temperature factors (Å <sup>2</sup> )	
Protein	22.1
Main chain atoms	20.8
Side-chain atoms	23.7
Chromophores	24.2
Solvent molecules	33.6

$\gamma$ -polypeptide were modeled into the electron density. Refinement of these residues indicated that they were fully occupied.

All residues of the  $\alpha$ - and  $\beta$ -polypeptides as well as the chromophores could be built into the electron density. There are only a few solvent-exposed residues (insertion at position  $\beta$ 147) with disordered side chains. The density for the expected  $\gamma$ -N-methylation of the asparagine residue at position  $\beta$ 72 was also very well defined. In contrast to the nomenclature used by Ficner and other authors (Schirmer *et al.*, 1985, 1986, 1987; Duerring *et al.*, 1990, 1991; Ficner *et al.*, 1992; Chang *et al.*, 1996) we used a continuous numbering of the amino acid sequence. The final model ( $R_{\text{work}} = 17.5\%$ ,  $R_{\text{free}} = 22.7\%$ ) included 333 water molecules; its quality is summarized in Table III.

The mean positional error of the atoms as esti-

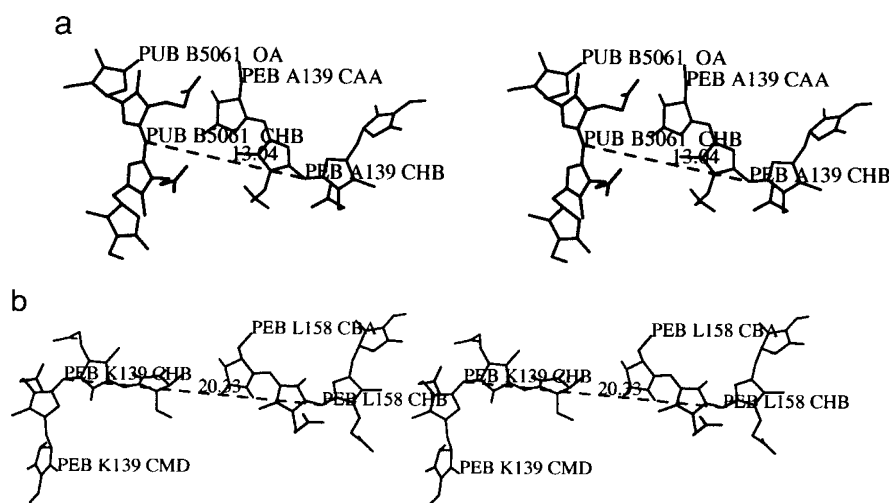
mated from a Luzzati (1952) plot is 0.27 Å. A Ramachandran (Ramachandran and Sasisekharan, 1968) plot shows that with the exception of the NCS-related residues Thr B75/L75 (near  $\phi = 88^\circ$ ,  $\psi = 148^\circ$ ) all other residues fall into allowed regions. The peptide nitrogen atoms of Thr B75/L75 are hydrogen bonded to oxygen atoms OD of symmetry-related PEBs at position  $\alpha$ 82. The electron densities of residues Thr B75/L75 are well defined and their conformation is highly conserved among all phycobili-protein structures (Schirmer *et al.*, 1985, 1986, 1987; Duerring *et al.*, 1990, 1991; Brejc *et al.*, 1995; Chang *et al.*, 1996).

The atomic coordinates of the refined model have deviations of bond lengths and bond angles from ideality of 0.004 Å and 1.1°, respectively. They are available from the Brookhaven Protein Data Bank (Entry No. 1B8D).

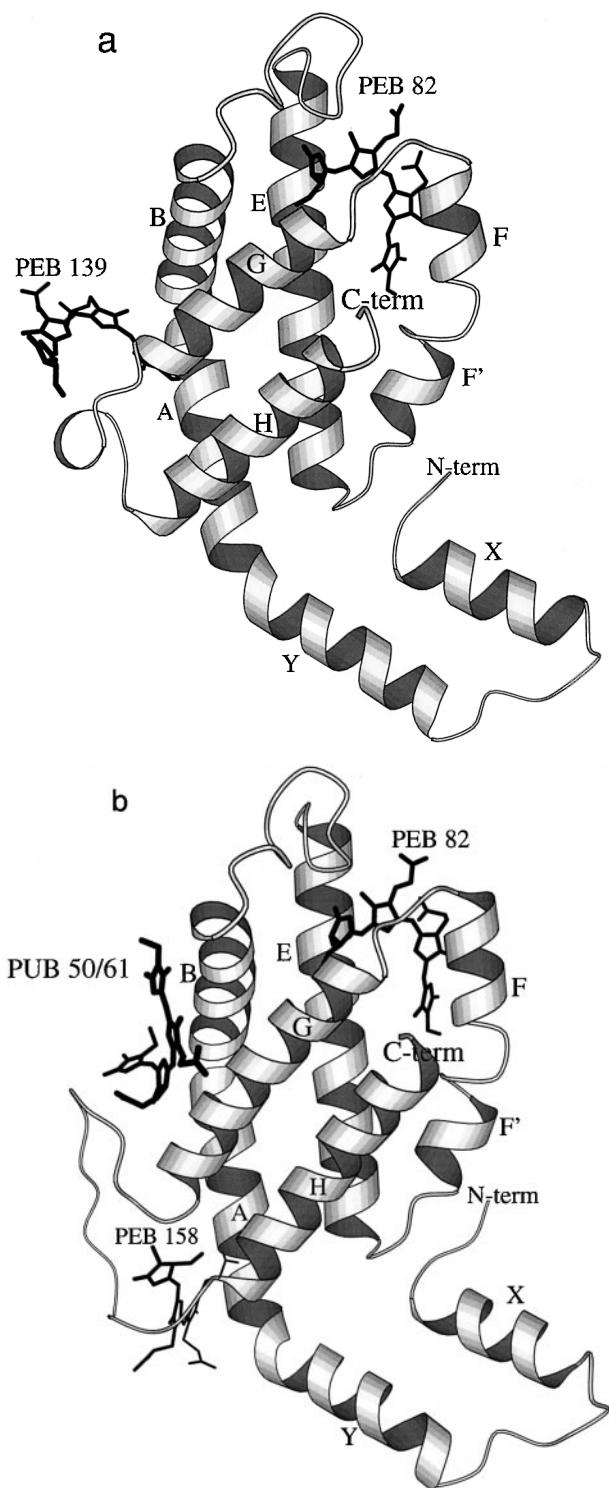
## DISCUSSION

### *Crystal Packing and Protein Contacts*

The structure of R-PE from the red alga *G. monilis* was solved by the method of molecular replacement, using 1CPC as a model, and refined at 1.90-Å resolution. Though the parameters of the unit cells are very similar and the positions of the helices within the proteins are almost identical, there is a striking difference in the packing of 1CPC and R-PE: the columns of hexamers are rotated by  $-40^\circ$  around the  $z$  axis, so that different intercolumn contacts are formed. Obviously, due to the approximate cylindrical symmetry of the columns, different packing arrangements can be accommodated within a similar unit cell. A special feature of the crystal packing in



**FIG. 2.** (a) Relative orientation (stereo view) of the chromophores PUB B50/61 and PEB A139 of a symmetry-related  $\alpha$ -polypeptide. The distance between the centers of gravity of these two chromophores is approximately 13.0 Å. (b) Relative orientation (stereo view) of the chromophores PEB K139 and PEB L158 of a symmetry-related  $\beta$ -polypeptide. The closest distance between these two chromophores is 3.4 Å, while the distance between their centers of gravity is 20.3 Å.



**FIG. 3.** Ribbon representation of (a) the  $\alpha$ -polypeptide and (b) the  $\beta$ -polypeptide. The chromophores are shown as thick lines. Plots were produced with Molscrip (Kraulis, 1991).

this R-PE structure is the chromophore–protein crystal contact between PEB K139 and the residues Cys L158 and Gln L159 on a symmetry-related  $\beta$ -polypeptide. The protein–protein contacts are predominantly made by the loops between helices G and H of the  $\alpha$ - and the  $\beta$ -polypeptides.

In R-PE the crystal packing makes it possible for PUB at position B50/61 on subunit B to come close to PEB at position A139 of a symmetry related  $\alpha$ -polypeptide. The distance between the centers of gravity of these two chromophores is approximately 13.0 Å, and their orientation relative to each other is almost perpendicular (Fig. 2a). There is also a short distance between chromophore PEB K139 and chromophore PEB L158 of a symmetry-related  $\beta$ -polypeptide. The closest distance between these two chromophores is 3.4 Å, while the distance between their centers of gravity is 20.3 Å, resulting from a parallel arrangement of these chromophores (Fig. 2b). Furthermore there is a close distance between PEB K139 and PUB L50/61 of a symmetry-related  $\beta$ -polypeptide. The distance between their centers of gravity is approximately 16.5 Å and their relative orientation is almost perpendicular. Thus, the crystal packing in R-PE could make an exciton interaction possible as an artifact of the crystalline state. Compared to the 1LIA entry in the Brookhaven database the arrangement of the chromophores within the crystal packing is the same for R-phycoerythrin from *Pol. urceolata*.

#### Molecular Structure

**$\alpha$ - and  $\beta$ -Polypeptides.** As in many other known phycobiliproteins the two polypeptides  $\alpha$  and  $\beta$  of R-PE from *G. monilis* are arranged as an  $(\alpha\beta)_6$  hexamer with the  $\gamma$ -polypeptide located in the central cavity. The  $\alpha$ -polypeptide contains 164 residues and the  $\beta$ -polypeptide contains 177 residues. The percentage of helix contents in the  $\alpha$ - and the  $\beta$ -polypeptides is 79 and 73%, respectively.

The main structural difference from 1CPC is a deletion of two amino acids at position  $\alpha$ 76 (C-terminus of helix C) and an insertion of four amino acids at position  $\alpha$ 138 (N-terminus of helix H', corresponding to position  $\alpha$ 140 in B-PE, including the attachment site of PEB  $\alpha$ 139) of the  $\alpha$ -polypeptide, as well as an insertion of five amino acids at position  $\beta$ 145 (corresponding to position  $\beta$ 147 in B-PE) of the  $\beta$ -polypeptide.

A comparison of the structure of R-PE from *G. monilis* with the 1LIA and the 1CPC entry shows root-mean-square (rms) deviations for  $C_{\alpha}$  atoms (using a distance cutoff of 3.8 Å) of 0.50 Å for 682  $C_{\alpha}$  atoms for 1LIA and 1.06 Å for 655 atoms for 1CPC.

Figure 3 shows the polypeptide chain of *G. monilis* R-PE as a ribbon trace together with the chromo-

**TABLE IV**  
Protein Environment of the Chromophores

Chromophore atom	PEB A82	PEB A139	PEB B82	PEB B158	PUB B50/61
OA	3.5 GlyA73-N 3.4 AlaA72-O		3.5 CysB73-SG 3.3 AsnB72-OD1	3.3 GlyB156-N 3.5 AlaB154-O	3.5 SerB147-OG 2.9 GlyB148-N
NA	3.5 AlaA72-O 3.4 CysA82-SG		2.9 AsnB72-OD1	2.9 AlaB154-O 3.7 CysB158-SG	
NB	2.7 AspA85-OD2	2.7 W383-OH2 3.1 ArgA142-NH1	2.7 AspB85-OD2	2.6 AspB39-OD2	2.8 AspB54-OD2
O1B		3.0 ArgA137-NH2	2.7 W548-OH2 3.5 ArgB77-NH1 3.0 ArgB77-NH2	2.9 W402-OH2 3.5 W482-OH2	3.6 SerB147-N
O2B	2.8 W549 OH2 3.0 SymW456-OH2	3.3 ArgA137-NH1 3.2 ArgA137-NH2	3.1 ArgB77-NH2 2.7 ArgB78-NE	2.6 W482-OH2	2.7 SerB147-OG
NC	2.9 AspA85-OD2 3.4 AspA85-OD1 3.4 ArgA84-NH2	3.0 W383-OH2 3.2 ArgA137-NH1	2.9 AspB85-OD2 3.2 AspB85-OD1 3.4 ArgB84-NH2	2.7 AspB39-OD2 3.6 W420-OH2	2.7 AspB54-OD2 3.3 W443-OH2
O1C	2.7 W345 OH2 2.8 ArgA84-NH1 2.7 ArgA84-NH2	2.8 ArgA142-NH2 2.6 W470-OH2	2.8 ArgB84-NH1 2.9 ArgB84-NH2		2.8 W467-OH2
O2C	2.8 LysA81-NZ 2.8 SymW456-OH2	3.2 ArgA142-NH2	2.8 W465-OH2 2.7 W548-OH2		3.4 ArgB129-NH1 3.5 W358-OH2
ND	3.0 W345 OH2	2.9 W470-OH2		3.0 AsnB35-OD1	2.7 W626-OH2
OD	2.9 W314 OH2 3.0 SymThrB75-N	2.9 AsnA47-ND2 3.1 W494-OH2		2.9 GlnA28-NE2	2.8 W555-OH2 3.3 CysB61-SG

Note. The distance is given in Å directly before the respective hydrogen-bond partner.

phores. A detailed description of the protein environment of the chromophores is given in Table IV.

**Chromophores.** R-PE has three PEBs ( $\alpha$ 82,  $\beta$ 82, and  $\beta$ 158) at equivalent chromophore positions as in 1CPC. Two additional chromophores, one PEB and one PUB, were found in the structure at equivalent

positions as in B-PE from *Por. sordidum* and R-PE from *Pol. urceolata*. The PEB chromophore is covalently bound to a Cys residue by ring A. It is located at position  $\alpha$ 139 on a solvent-exposed loop of the  $\alpha$ -polypeptide, protruding from the protein and connecting helices G and H'. The doubly linked PUB is bound to Cys  $\beta$ 50 on helix B of the  $\beta$ -polypeptide with ring A and to Cys  $\beta$ 61 on the same helix with

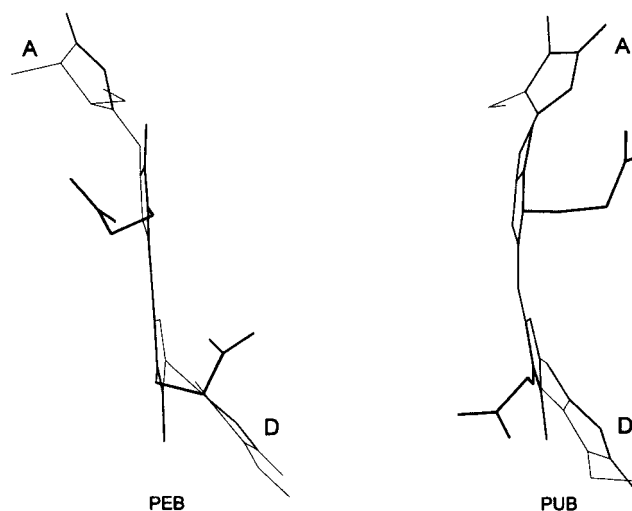
**TABLE V**

Stereochemistry and Conformation of the Chromophores of Subunits A and B of R-PE

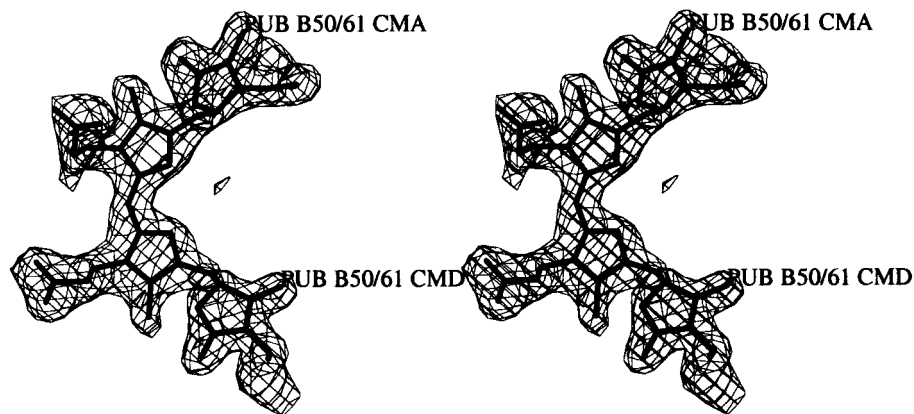
Chromophore	PEB A82	PEB A139	PEB B82	PEB B158	PUB B50/61
Chiral atoms					
C-2	R	R <sup>a</sup>	R	R	—
C-3	R	R	R	R	—
C-3'	R	R <sup>a</sup>	R	S	S
C-4	—	—	—	—	S
C-16	R	R	R	R	R
C-18'	—	—	—	—	R
Conformation of double bonds					
C <sub>4</sub> (ring A–B)	Z-anti	Z-anti	Z-anti	Z-anti	—
C <sub>10</sub> (ring B–C)	Z-syn	Z-syn	Z-syn	Z-syn	Z-syn

Note. The conformations of the chromophores on the related subunits K and L are the same. The nomenclature of the chromophore atoms is given in Fig. 1. The positions 3' and 18' indicate the binding of the chromophore to the protein. syn, synperiplanar; anti, antiperiplanar; —, not a chiral atom.

<sup>a</sup> For the determination of the stereochemistry PEB K139 was used, as the electron density of the methyl group of the chromophore PEB A139 was weak due to strong exposure of the chromophore to the solvent.



**FIG. 4.** Conformation of PEB  $\beta$ 82 and PUB  $\beta$ 50/61 chromophores. The conformation of all other PEB chromophores is very similar. Rings B and C of the chromophores point toward the reader, while rings A and D point in the opposite direction. The boat-like conformation of PUB can clearly be seen.

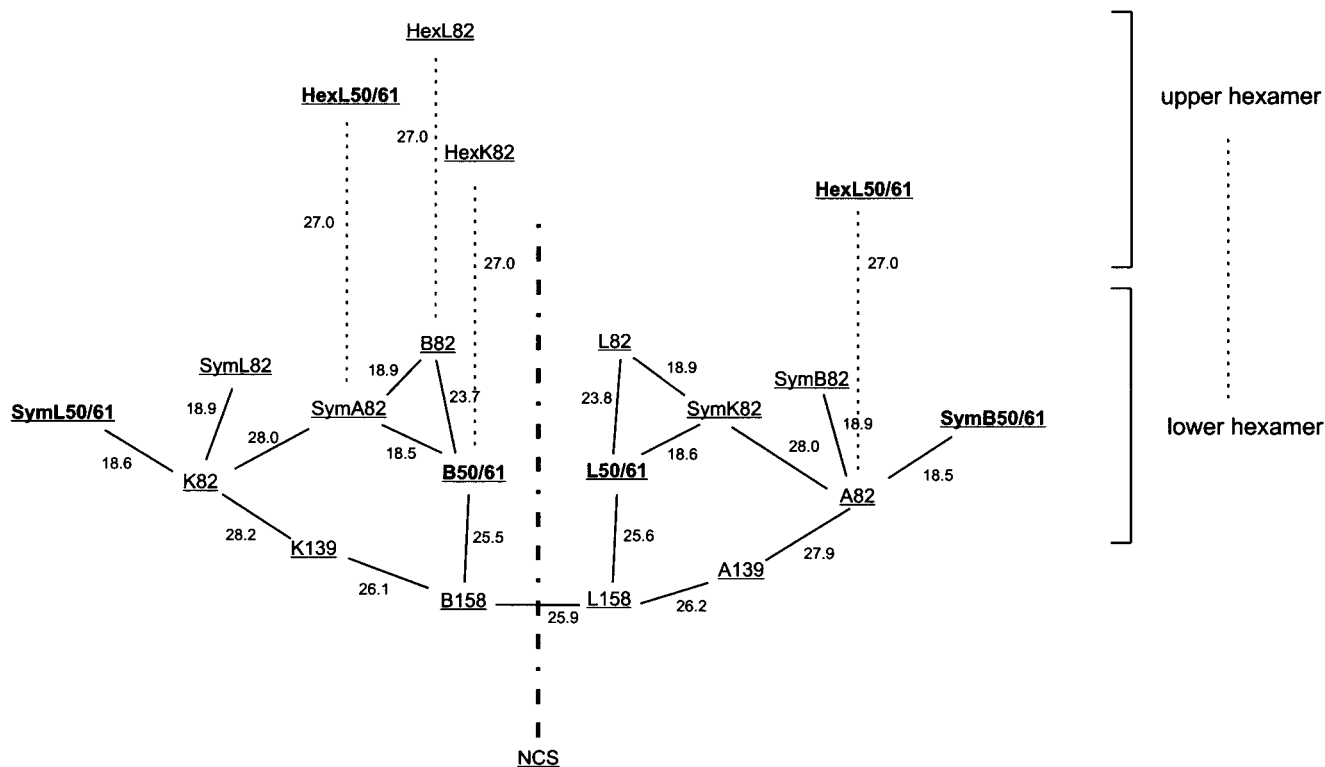


**FIG. 5.** Stereo view of chromophore PUB  $\beta 50/61$ . The  $2F_{\text{obs}} - F_{\text{calc}}$  density map is displayed at  $1.0\sigma$ . Ring A of the chromophore is on the top of the drawing.

ring D. The stereochemistry of the chromophores PEB  $\alpha 82$ ,  $\alpha 139$ ,  $\beta 82$ , and  $\beta 158$  is identical to the stereochemistry of the chromophores of B-PE from *Por. sordidum* (for the nomenclature of the chromophore atoms see Fig. 1). The stereochemistry of the atoms C-2 and C-3' of the chromophore PEB A139 was derived from the symmetry-related chromophore PEB K139, as the electron density of the

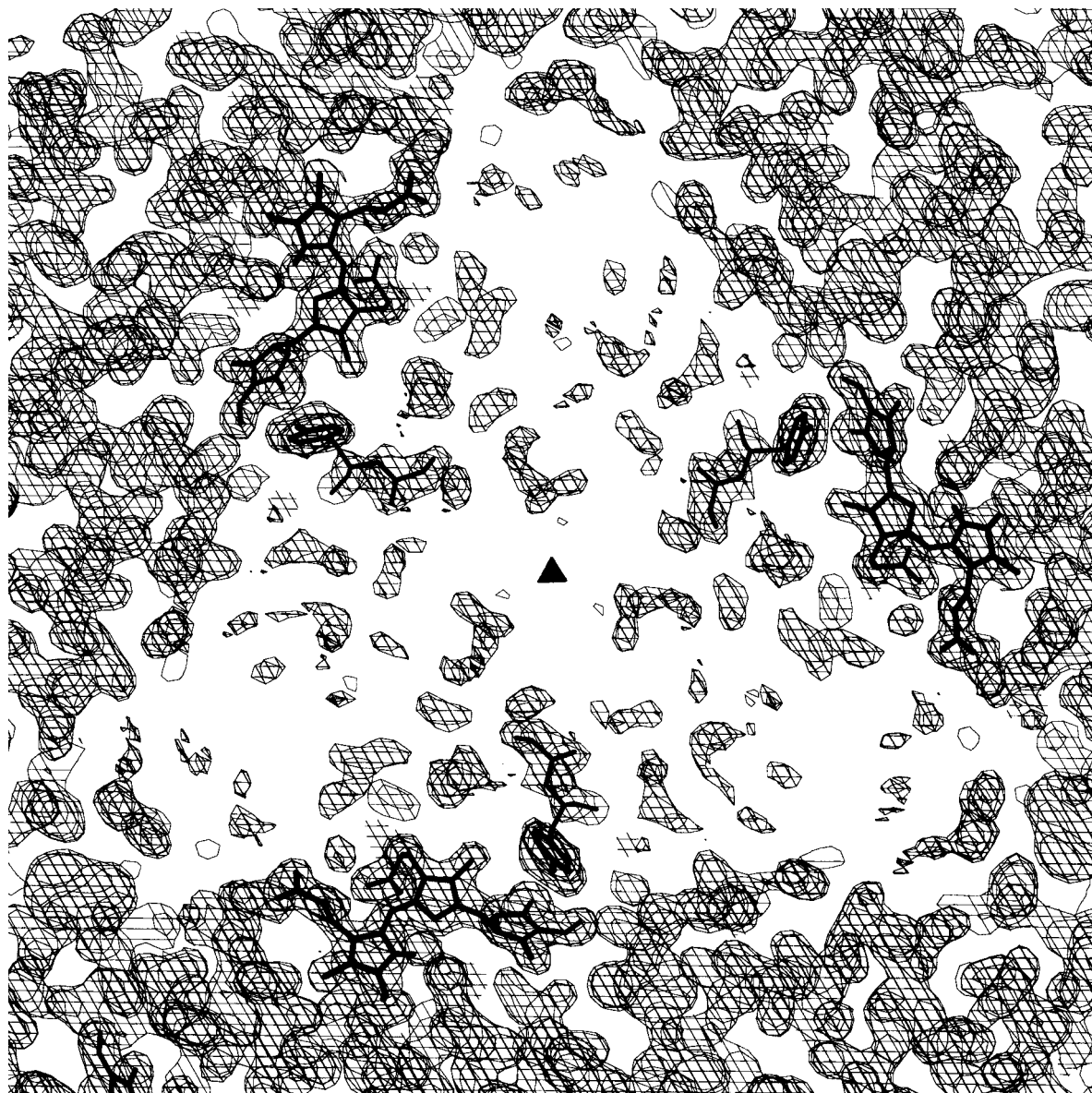
methyl group CMA of PEB A139 was weak due to the strong exposure to the solvent. The stereochemistry of the individual chromophores is given in Table V.

The conformation of the PUB chromophore differs strongly from the conformation of the PEB chromophores. In PUB, rings A and D are located on the same side of the chromophore plane built by rings B and C, whereas in all PEB chromophores these two



**FIG. 6.** Schematic presentation of the arrangement and distances of the chromophores in the asymmetric unit ABKL. The chromophores are projected into the  $xy$  plane, and the view is in  $z$  along the crystallographic threefold axis. The NCS symmetry of the asymmetric unit is shown. Distances are given in Å. Only distances  $<30$  Å are shown for selected chromophores. Sym, symmetry-related chromophores within the R-PE hexamer; Hex, symmetry-related chromophores of two stacked hexamers.



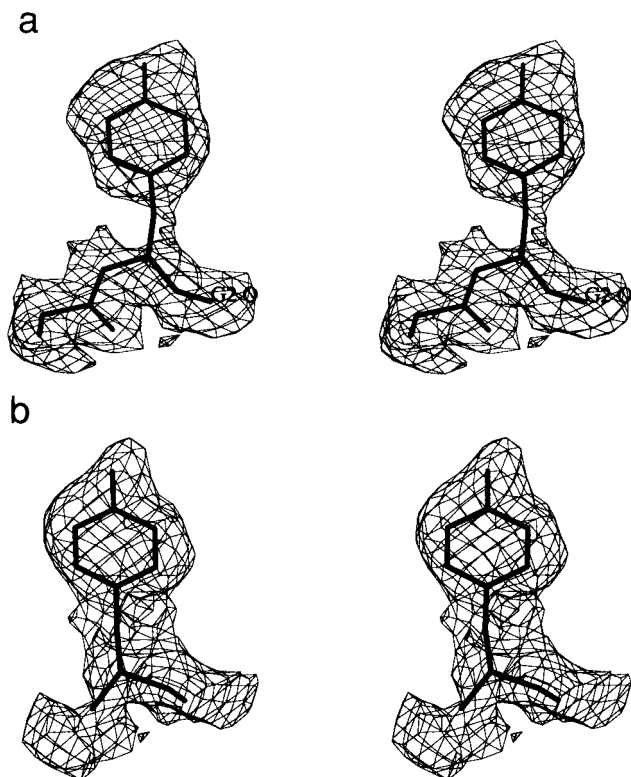


**FIG. 7.** Slab of the electron density in the central cavity of R-PE displayed at  $1\sigma$ . This segment of electron density encompasses the symmetry related chromophores PEB B82 together with dipeptides Gly G1-Tyr G2 (heavy lines) of the  $\gamma$ -polypeptide. The crystallographic threefold axis is indicated.

rings are found on different sides of the chromophore plane. Figure 4 shows the “boat-like” conformation of PUB together with the “z” conformation of the PEB chromophores. The stereochemistry of the PUB could be determined accurately (Table V) due to the high resolution of the structure and the very-well-defined electron density of the chromophore. Looking at the 1LIA entry, PUB in *Pol. urceolata* seems to be more evenly bent than the PUB in *G. monilis*, which makes an exact determination of the stereochemistry difficult and is probably an artifact of the lower resolution of that study (2.8 Å). Figure 5 shows the chromophore PUB B50/61 in R-PE from *G. monilis*

within a  $(2F_{\text{obs}} - F_{\text{calc}})$  density map displayed at  $1.0\sigma$ .

*Energy transfer.* Though the absorption maximum of PUB ( $\lambda_{\text{max}}$  at 498 nm) is at a much shorter wavelength than the absorption maximum of the PEB ( $\lambda_{\text{max}}$  at 535–567 nm) chromophores (Apt *et al.*, 1995), PUB is not isolated in the periphery of the protein, building the starting point of an energy transfer pathway in a single disc but is rather located in the vicinity of the chromophores PEB  $\beta$ 82 and PEB  $\beta$ 158 and a symmetry-related chromophore PEB  $\alpha$ 82 within the hexamer. It is also close to a symmetry-related PEB  $\alpha$ 82 in the hexamer stack



**FIG. 8.** (a) Stereo view of the dipeptide Gly G1-Tyr G2. The  $2F_{\text{obs}} - F_{\text{calc}}$  density map is displayed at  $0.8\sigma$ . (b) Stereo view of Tyr G3. The  $2F_{\text{obs}} - F_{\text{calc}}$  density map is displayed at  $0.8\sigma$ .

(Fig. 6). Assuming a suitable orientation of the transition dipole moments of the chromophores, it is possible that PUB not only absorbs light in the blue region of the spectrum, but also takes part in the energy transfer within a single disc as described for PEB  $\beta 50/61$  in the structure of B-PE from *Por. sordidum* (Ficner *et al.*, 1992). Figure 6 shows schematically the relative orientation of the chromophores in the asymmetric unit of R-PE and lists the shortest distances between them. In addition, some symmetry-related chromophores within the same hexamer and between a stack of hexamers are shown.

Looking at the distances ( $<30 \text{ \AA}$ ) between neighboring chromophores in the R-PE hexamer, we observed two prominent distances:  $18.5$  and  $26 \pm 2 \text{ \AA}$ . Ficner *et al.* (1992) observed similar distances for the chromophores in B-PE ( $19.5$  and  $25 \pm 3.5 \text{ \AA}$ ). Calculating the energy transfer rates in B-PE, they found that the transfer of energy is not primarily related to the distance between the chromophores as similar distances between chromophores did not account for similar energy transfer rates. This indicates that the orientation of the chromophores in phycobiliproteins is not only optimized for energy transfer to the photosynthetic reaction center but is also a prerequi-

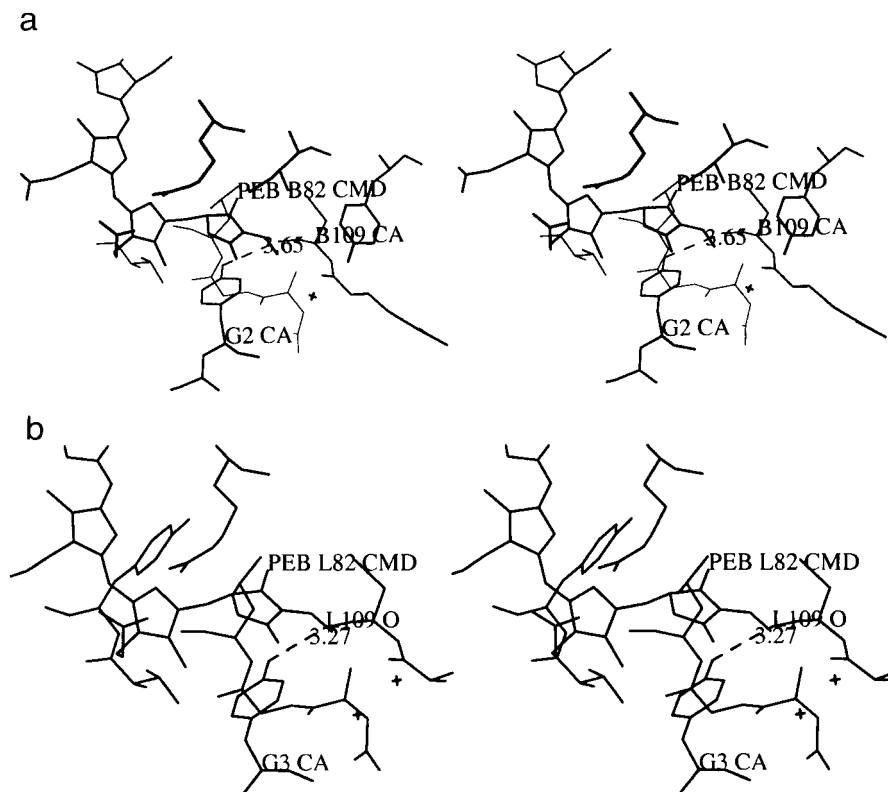
site for an effective capture of light energy and its interim storage in the antenna complex.

*$\gamma$ -Polypeptide.* SDS-PAGE indicates that there are at least two different  $\gamma$ -polypeptides present in the crystals of R-PE (Ritter *et al.*, 1997); however, only isolated spots of weak electron density showing a threefold symmetry could be observed in the center of the  $(\alpha\beta)_6$  hexamer (Fig. 7), so that the  $\gamma$ -polypeptide cannot be clearly seen in a  $(2F_{\text{obs}} - F_{\text{calc}})$  electron density map.

A careful inspection of a  $(F_{\text{obs}} - F_{\text{calc}})$  difference map displayed at  $4\sigma$ , however, revealed a strong difference in electron density inside the  $(\alpha\beta)_6$  hexamer in close proximity to the chromophores PEB B82 and PEB L82 that could not be explained by the initial model. We interpreted this electron density as deriving from amino acid residues of the  $\gamma$ -polypeptide and modeled one dipeptide (Gly G1-Tyr G2) and one single amino acid (Tyr G3) into well-defined regions of this density near chromophore PEB B82 and chromophore PEB L82, respectively. Both Tyr residues are related by the same NCS dyad as the chromophores. Figure 8 shows the residues with their  $(2F_{\text{obs}} - F_{\text{calc}})$  electron density, displayed at  $0.8\sigma$ .

In contrast to the residue Tyr G3, the electron density for Tyr G2 was not ideally defined as it was, in principle, possible to model either a Tyr or a Trp residue. In particular the density of the C $\beta$  atom of Tyr G2 is defined only at a low sigma cut ( $0.8\sigma$ ). However, as a Tyr residue fitted the density best and taking into account the fact that G2 and G3 are related by the NCS dyad, we modeled the difference electron density as Tyr G2. The side chains of Tyr G2 and Tyr G3 are both in close proximity ( $3.5$ – $3.7 \text{ \AA}$ ) to ring D of the chromophores PEB B82 and PEB L82, respectively (Fig. 9). Tyr G3 is hydrogen-bonded to the peptide oxygen of residue Cys L109 ( $3.3 \text{ \AA}$ ), while the corresponding distance between Tyr G2 and Cys B109 is about  $3.7 \text{ \AA}$ . This shows that the hydrophobic pockets of chromophores PEB B82 and PEB L82 are partly built by the  $\gamma$ -polypeptide rather than being completed by the  $(\alpha\beta)_6$  hexamer. As a consequence, both chromophores are shielded against the chromophores of the  $\gamma$ -polypeptide by the polypeptide backbone of these residues, which inhibits close contact between the chromophores of the  $\beta$ -polypeptide and the  $\gamma$ -polypeptide. This would make an exciton exchange rather improbable and would instead favor a Förster (1965) mechanism for energy transfer.

The fact that the  $\gamma$ -polypeptide cannot be seen in the electron density was also reported for B-PE (Ficner *et al.*, 1992) and R-PE (Chang *et al.*, 1996) and is due to the incompatibility of the  $(\alpha\beta)_6\gamma$  stoichiometry with the threefold symmetry of the lattice, as the  $\gamma$ -polypeptides show no threefold



**FIG. 9.** (a) Protein environment of the residues Gly G1-Tyr G2 of the  $\gamma$ -polypeptide. (b) Protein environment of Tyr G3 of the  $\gamma$ -polypeptide.

symmetry in their sequence (Apt *et al.*, 1993; Ficner and Huber, 1993). Thus, the  $\gamma$ -polypeptides are present in the crystal, but their arrangement within the hexamers is statistical, so that their electron density is averaged. Looking at the sequence of the  $\gamma$ -polypeptide of R-PE from *Aglaothamnion neglectum* (Apt *et al.*, 1993) we found 18 Tyr residues, 6 of which we would expect to be involved in the interaction with the PEB chromophores at position  $\beta$ 82 due to the occupancy refinement results.

### Conclusions

With the exception of the bilins occurring in cryptophytes, the structures of most other phycobilins (PCB, PXB, PEB, and PUB) have been solved. The future goal is to determine the structure of the linker proteins and complexes of phycobiliproteins together with linker proteins. In this respect, the structure of R-PE reported here could allow the modeling of a separately determined  $\gamma$ -polypeptide into the hexamer using the two symmetry-related Tyr positions reported here as markers.

We thank Frank Sharples for his expert assistance in culturing *Griffithsia*.

*Note added in proof.* The refinement of R-PE from *Polysiphonia urceolata* at 1.9 Å resolution has recently been reported (Jiang *et al.*, 1999).

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