# Pigment-Pigment Interactions in PCP of Amphidinium carterae Investigated by Nonlinear Polarization Spectroscopy in the Frequency Domain

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ABSTRACT Peridinin-chlorophyll a-protein (PCP) is a unique antenna complex in dinoflagellates that employs peridinin (a carotenoid) as its main light-harvesting pigment. Strong excitonic interactions between peridinins, as well as between peridinins and chlorophylls (Chls) a, can be expected from the short intermolecular distances revealed by the crystal structure. Different experimental approaches of nonlinear polarization spectroscopy in the frequency domain (NLPF) were used to investigate the various interactions between pigments in PCP of Amphidinium carterae at room temperature. Lineshapes of NLPF spectra indicate strong excitonic interactions between the peridinin's optically allowed  $S_2$  (1Bu<sup>+</sup>) states. A comprehensive subband analysis of the distinct NLPF spectral substructure in the peridinin region allows us to assign peridinin subbands to the two Chls a in PCP having different  $S_1$ -state lifetimes. Peridinin subbands at 487, 501, and 535 nm were assigned to the longer-lived Chl, whereas a peridinin subband peaking at 515 nm was detected in both clusters. Certain peridinin(s), obviously corresponding to the subband centered at 487 nm, show(s) specific (possibly Coulombic?) interaction between the optically dark  $S_1(2A_g^-)$  and/or intramolecular charge-transfer (ICT) state and  $S<sub>1</sub>$  of Chl a. The NLPF spectrum, hence, indicates that this peridinin state is approximately isoenergetic or slightly above  $S_1$  of Chl a. A global subband analysis of absorption and NLPF spectra reveals that the Chl a  $Q_v$ -band consists of two subbands (peaking at 669 and 675 nm and having different lifetimes), confirmed by NLPF spectra recorded at high pump intensities. At the highest applied pump intensities an additional band centered at  $\leq 60$  nm appears, suggesting—together with the above results—an assignment to a low-dipole moment  $S_0 \to S_1/ICT$  transition of peridinin.

### INTRODUCTION

Dinoflagellates possess a unique water-soluble peripheral antenna complex, peridinin-chlorophyll a-protein (PCP). The carotenoid peridinin is the main light-harvesting pigment in PCP. The crystal structure of PCP from Amphidinium (A.) carterae reveals a trimeric organization of the complex (1). Within each monomeric subunit, pigments are organized in two clusters related by a pseudo-symmetry. Each cluster contains one chlorophyll (Chl) a surrounded (within van der Waals distances) by four peridinins (1). The closest distances between peridinins and Chls, as well as between peridinins in one cluster, are  $\sim$ 4 Å. The center-to-center distance between the two Chls *a* in a PCP monomer is rather large ( $\sim$ 17 Å; see Hofmann et al. (1)). Thus, whereas pronounced peridinin-

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peridinin  $(2-6)$  as well as peridinin-Chl *a* interactions can be expected in PCP, interaction between the Chls appears to be negligible.

Peridinin is an unusual carotenoid, which has a C-37 backbone instead of the usual C-40 carbon skeleton. Peridinin also contains an allene moiety, a lactone ring in conjugation with the  $\pi$ -system and acetate and epoxy groups on the opposite  $\beta$ -rings. These features strongly influence the photophysical properties; in nonpolar solvents, peridinin shows a typical carotenoid vibrational substructure of the optically allowed  $S_0 \rightarrow S_2$  transition in the 350–550 nm range. (Note that the following nomenclature for carotenoid singlet states is used:  $S_0 = 1Ag^-, S_1 = 2Ag^-,$  and  $S_2 = 1Bu^+$ ; the recently experimentally detected state with  $1B_{\rm u}^-$  character (7) is assumed to lie above  $S_2$  in peridinin (8)). The peridininabsorption spectrum in polar solvents, as well as in the PCP protein environment, is rather unstructured (Fig. 1). In particular, the presence of a carbonyl group was suggested to result in a broadening of the absorption bands and a pronounced solvent-dependence of the  $S_1$ -state lifetime (9,10). The latter behavior has been attributed to the formation of an intramolecular charge-transfer state (ICT), which mixes with the  $S_1$  state (11,12).

Peridinin is generally assumed not to contribute to PCP absorption in the Chl  $a Q_y$  region (Fig. 1). However, the energetic location as well as the dipole strength of the  $S_0 \rightarrow$ 

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FIGURE 1 Absorption spectra of the peridinin-chlorophyll *a*-protein  $(PCP)$  and Chl  $a$  (in diethyl ether; shaded, red-shifted by 6 nm to match Chl a-absorption in PCP).

 $S_1$  transition of peridinin—in particular, in the proteinaceous environment in PCP—is a matter of controversy (e.g.,  $(6,8,10,13-15)$ ). For a recent critical review of the issue of optically dark states of carotenoids and a thorough discussion of the various experimental approaches to detect these states, see Polivka and Sundström (16).

Amino-acid sequences of the C- and N-terminal regions of the PCP apoprotein are only  $\sim$  56% identical (1). Hence, the site energies of the attached chromophores (peridinins as well as Chls *a*) may differ in the C- and N-terminal clusters. The site energies of the four peridinins within one cluster are also expected to vary as a consequence of their different attachment sites in the protein matrix. In the  $Q_y$ -absorption region of PCP from A. carterae, two subbands (related to the two Chls of one monomer) have been partly resolved at cryogenic temperatures (5,17). Moreover, two different  $C=O$  stretching modes (1697 and 1681 cm<sup>-1</sup>) were observed by fluorescence line-narrowing spectroscopy, suggesting that there are two different emitting Chls  $a$  (5). However, Kleima et al. (18) as well as Ilagan et al. (17) state that the two Chls are spectrally indistinguishable at room temperature (RT). A Gaussian deconvolution of the RT fluorescence emission spectrum of PCP from Symbiodinium microadriaticum revealed two well-separated emission bands centered at 675.5 and 679 nm (19,20). An assignment of the spectroscopically different Chls  $a$  to the C- or N-terminal clusters has not been achieved, and a comprehensive spectral subband analysis of the PCP Soret/ $S_2$  region absorption reflecting all known structural information is not yet possible. An attempt of fitting circular dichroism (CD) and absorption spectra involving all possible inter- and intracluster pigment combinations resulted in only a moderately successful match between experimental and calculated spectra (3). However, four peridinin subbands were identified and excitonic coupling between peridinins was estimated to be of the order of 100–300  $\text{cm}^{-1}$  (3). By combining information from absorption, linear dichroism, CD, and triplet-minus-singlet spectra of PCP with the absorption spectrum of peridinin in solution, four

peridinin subbands were also resolved (5) with similar wavelength maxima to those reported in Carbonera et al. (3). So far, none of the above-mentioned peridinin subbands have been assigned to either the N- or C-terminal cluster or to the corresponding Chl a.

In addition, modes and routes of excitation energy transfer (EET) from peridinin to Chl  $a$  after excitation via the allowed  $S_0 \rightarrow S_2$  transition are not yet firmly established, and the proportion of peridinin  $\rightarrow$  Chl EET that proceeds via the optically dark state(s) is a matter of debate. Several studies (12,13,21) suggest that excitation energy is transferred directly (with an efficiency of 25–50%) from the peridinin  $S_2$ state to the Chl a  $Q_x$  state in addition to a route via the  $S_1$  and/ or ICT state(s). In contrast, different workers (4,6,22) favor the notion that EET proceeds exclusively via the  $S_1$  state after rapid internal conversion from  $S_2$ .

Fluorescence excitation spectra indicate a high efficiency of EET from peridinin to Chl  $a$  in PCP (4,17,23). Wavelengthindependence of fluorescence excitation efficiency in the peridinin-absorption region was taken as being consistent with fast EET between the peridinins. This observation would also be in line with strong excitonic coupling among peridinins as suggested by CD spectra (2,24) and structure-based calculations (6). In contrast, Akimoto et al. (22) found a significant wavelength-dependence of the fluorescence excitation efficiency for wavelengths shorter than 520 nm in PCP from Alexandrium cohorticula. Moreover, from unaltered  $S_2$  lifetimes in PCP (as compared to peridinin in solution), these authors concluded that EET between peridinin  $S_2$  states in PCP is unlikely. However, the fluorescence excitation spectrum alone does not provide information about the EET routes and their relative proportions in PCP. To investigate these issues in more detail, it is necessary to elucidate the photophysical properties of each individual peridinin, i.e., site energy (subband), excited-state lifetime, and the mode of coupling between the pigments. Indeed, spectral substructure models for the peridinin-absorption band have been proposed (3,6,8). Damjanovic et al. (6) have suggested that one peridinin per cluster (per612 or per622, in their notation) does not undergo direct EET to Chl  $a$ , but can only transfer energy to a neighboring peridinin at the  $S_2$  level. Overall peridinin to Chl EET rates have been derived from femtosecond-transient absorption and fluorescence measurements without being able to differentiate between individual peridinin species (4,12,13,22).

Apparently, the above issues are either not or only partially addressable by conventional spectroscopic approaches. Thus, nonlinear polarization spectroscopy in the frequency domain (NLPF; see Fig. 2) was applied to study pigmentpigment interactions in PCP at RT. The aims were: 1), to determine the mode of coupling between peridinins in the  $S_2$ state; 2), to resolve the spectral substructure of the peridinin  $S_2$  band and Chl a  $Q_y$ -absorption regions, as well as to differentiate the routes of EET from peridinin to Chl  $a$ ; and 3), to elucidate interaction between the peridinin  $S_1$  state and Chl a, as well as the energetic location of the former.



FIGURE 2 Principle of nonlinear polarization spectroscopy in the frequency domain (NLPF). Pump, probe, and signal fields  $(E)$  are distinguished by the indices  $p$ ,  $t$ , and  $s$ , respectively, as well as the corresponding wave number vectors  $(k)$ . The NLPF signal is the component of the signal field perpendicular to the incident probe field.

## MATERIALS AND METHODS

#### Sample preparation

PCP was isolated from A. carterae as described (1) and diluted before the experiments in 25 mM Tris-HCl/10 mM KCl, pH 7.5. The sample (OD  $\cong$ 1.1 at 670 nm) was contained in a 5-mm quartz cell (Hellma, Müllheim, Germany), which does not show detectable birefringence.

## Experimental setup

NLPF spectra were recorded using a  $90^{\circ}$  setup as described previously (25). Pump  $(\lambda_n)$  and probe  $(\lambda_t)$  beams, characterized by a spectral linewidth of  $\sim$ 0.05 cm<sup>-1</sup>, are obtained from dye lasers (DCM in DMSO, or Coumarin 47/307 in methanol, for the Chl  $Q_v$  and Soret/allowed peridinin-absorption regions, respectively), synchronously pumped by an excimer laser with a pulse duration of 15 ns. The absolute wavelengths, as well as spectral linewidths of the pump and probe laser beams, were assessed with picometer precision using a Wavemeter (Model #450A, Burleigh Instruments, Victor, NY). NLPF spectra were obtained by tuning the pump laser across the spectral region under investigation with an increment of 0.2 nm. The  $\chi^2$ approach for description of the NLPF spectra is justified for pump-beam intensities of  $\sim 0.7 - 4 \times 10^{22}$  photons cm<sup>-2</sup> s<sup>-1</sup>. Intensity fluctuations of the pump and probe lasers were on the order of 5%. To obtain a better S/N ratio, 10 repetitive scans were averaged. To compare NLPF signal amplitudes it was essential to keep intensities of the pump and probe fields constant over the entire tuning range. The NLPF signal amplitude was corrected for the dependence on the optical density of the sample at the respective  $\lambda_t$ . To measure strong-field NLPF spectra, pump-beam intensities were varied by a neutral glass filter set from  $1.5 \times 10^{22}$  to  $4.5 \times 10^{24}$ photons  $\text{cm}^{-2} \text{ s}^{-1}$ . Absorption, CD, and fluorescence spectra were measured before and after the NLPF experiments to monitor sample integrity. All measurements were done at RT.

#### Information contents of NLPF spectra

The absorption of each individual pigment in a pigment-protein complex is shaped by interaction with its environment. This means that resolution of the spectral substructure of the overall absorption band in connection with the spectroscopic characterization of individual pigments reveals information about pigment-pigment as well as pigment-protein interactions. It has been shown previously that NLPF is a powerful method to elucidate spectral substructure and pigment-pigment interactions (26–28). NLPF is a degenerate four-wave mixing technique. A detailed description of the theoretical background of NLPF spectroscopy, together with a thorough discussion of the information content in NLPF lineshapes, is given in Beenken and Ehlert

(29). Briefly, the NLPF-signal results from nonisotropic bleaching of an isotropic sample by a polarized pump laser field (see Fig. 2). The bleaching is probed by a second laser field, whose polarization plane is oriented at 45° with respect to the pump field. The NLPF signal is measured in the propagation direction of the probe beam, but in the polarization plane perpendicular to that of the probe field and has the same wavelength. Conventionally, a NLPF spectrum is measured at a fixed probe wavelength  $\lambda_t = 2\pi c/\omega_t$  by tuning the pump wavelength  $\lambda_p = 2\pi c/\omega_p$  across the absorption band under investigation. Two measuring regimes (with regard to pump intensity) providing different information can be distinguished.

## Low pump intensity  $(x^{(3)}$  approach)

In the following, the characteristics of a NLPF spectrum of a homogeneously broadened optical transition (corresponding to an isolated two-level system) will be discussed exemplarily. The theoretical lineshape  $S(\omega_{p})$  of the NLPF spectrum measured at  $\lambda_t$  is given in the limits of a  $\chi^{(3)}$  approach by the square module of the lineshape function  $s(\omega_p)$  of the nonlinear response to pump and probe frequency (29),

$$
\begin{split} \left| s(\omega_{\text{p0}}, \omega_{\text{t0}}) \right|^2 \\ &= \left| \frac{i}{\Gamma - i\omega_{\text{t0}}} \left( \frac{2}{\gamma} \times \frac{2\Gamma}{\Gamma^2 + (\omega_{\text{p}} - \omega_{\text{0}})^2} + \frac{2}{\gamma + i(\omega_{\text{p}} - \omega_{\text{t}})} \right) \right. \\ &\times \left( \frac{1}{\Gamma - i\omega_{\text{t0}}} + \frac{1}{\Gamma + i(\omega_{\text{p}} - \omega_{\text{0}})} \right) \bigg)^2, \end{split} \tag{1}
$$

where  $\Gamma = 1/T_2$  is the dephasing rate and  $\gamma = 1/T_1$  is the energy relaxation rate.

The center frequency of the absorption profile is represented by the transition frequency  $\omega_0 = 2\pi c/\lambda_0$  of the underlying two-level system. The spectral displacement between pump and probe beam as well as the center frequency are represented by the terms  $\omega_{p0} = \omega_p - \omega_0$  and  $\omega_{t0} = \omega_t - \omega_0$ . The absolute signal amplitude scales with a factor  $\alpha(1/\omega_{t0})$ . In general, the NLPF-spectrum exhibits, besides a narrow  $T_1$ -peak, a spectrally broad  $T_2$ peak. The T<sub>1</sub>-peak is observed exclusively in the vicinity of  $\omega_n \approx \omega_t$ , and its spectral width is determined by the energy relaxation rate,  $\gamma$ . At room temperature,  $\gamma$  is much smaller than the dephasing rate  $\Gamma$  ( $\Gamma \gg \gamma$ ), which determines the width of the T<sub>2</sub>-peak. Therefore, the T<sub>2</sub>-peak for  $\omega_p - \omega_t \gg \gamma$ can be approximated by the first term of the lineshape function and results in a broad spectrum centered at  $\lambda_0$  with a lineshape corresponding to the square module of the absorption profile. The amplitude of the NLPF signal scales with the square module of the pump intensity and excited-state lifetime  $T_1 =$  $1/\gamma$  and, hence, correlates with ground-state depletion. Absorption bands of identical pigments at energetically nonequivalent binding sites are superimposed, resulting in a heterogeneous substructure of the overall absorption band. This situation can be described by an ensemble of  $(\nu)$  two-level systems, with different transition frequencies  $\omega_{\nu}$ , dephasing, and energy relaxation rates. Such a heterogeneous substructure results in a lineshape function, which is the sum of the individual lineshape functions weighted by the strength of the transition dipole moment  $\vec{\mu}_{v}$  and the concentration  $n_{\text{mol},v}$  of individual pigments combined into a weighting factor  $\alpha_{\nu}$ :

$$
S(\omega_{\mathrm{p}}) = |\sum \alpha_{\nu}^{2} s_{\nu}(\omega_{\mathrm{p}})|^{2}
$$
  
\n
$$
\alpha_{\nu}^{2} \propto \sqrt{n_{\mathrm{mol},\nu}} |\vec{\mu}_{\nu}|^{2}.
$$
\n(2)

Thus, the lineshape of the overall NLPF-spectrum does not necessarily conform to the square module of the absorption profile, and may contain a series of more or less distinguishable  $T_2$ -peaks. By virtue of this, an underlying substructure may become discernible with a resolution exceeding that of conventional spectroscopy (29). Moreover, besides a possible subband analysis, the lineshape of the NLPF spectrum may reveal the mode of pigment-pigment interaction.

Two basic cases of pigment-pigment interaction can be distinguished readily in NLPF spectra (27–29):

- 1. Weak dipole-dipole interaction gives rise to incoherent EET. The subbands represent independent species that are connected via Förstertype EET. Since the probe field inspects ground-state depletion, the lineshape of the NLPF spectrum is essentially determined by the excitedstate lifetimes (29). In the case of weak coupling, excitation would not be shared instantaneously between pigments; excitation occurs according to the spectral overlap of each individual subband with the pump field. The NLPF signal is enhanced for the subband(s) in which  $\lambda_t$  is located. Pigments with longer excited-state lifetimes contribute decisively to the overall NLPF signal. Hence, the NLPF lineshape is  $\lambda_t$ -dependent and structured NLPF spectra will be expected. Probing in a higher-energy subband (relative to the pump field) results in almost the same NLPF lineshape as in the case of a heterogeneous substructure. Probing in a lower-energy subband, the NLPF spectra are distinct; the peak in the spectral region of the higher-energy subband grows with increasing EET rate.
- 2. Strong dipole-dipole interaction leads to excitonic coupling which may result in splitting of the transition energies, redistribution of dipole strengths and delocalization of excitation energy. Strongly coupled pigments share a common ground state: a homogeneous substructure emerges. The energy-level scheme is characterized by a common ground state of the coupled pigments. The excitonic states are connected by the exciton downscattering rate. Peaks of nonprobed subbands are considerably more pronounced in NLPF spectra in the case of strong coupling. This may lead—in particular, for multiple coupled pigments—to a uniform/unstructured NLPF lineshape. Additionally, even if there is no spectral overlap between the subbands—the signal corresponding to the nonprobed subband does not disappear. This effect originates from the common ground state of the interacting species.

#### High pump intensity

The  $\chi^{(3)}$  approach to describe the NLPF spectrum is not applicable at strong pump intensities (30). Above a certain pump intensity, the amplitude of the NLPF signal no longer follows the square module of the pump intensity, and saturation effects influence the NLPF spectrum. Moreover, the lineshape of the NLPF spectrum broadens, whereas center wavelengths remain unchanged. The onset of these effects is determined by strength of the transition dipole moment  $\vec{\mu}$ , energy relaxation, and dephasing rates. Consequently, the NLPF spectrum differs for subbands representing pigments at nonequivalent binding sites. Thus, a subband with a small transition dipole moment, which would not be detectable in NLPF spectra at low pump intensities, may become obvious at higher pump intensities.

Summarizing, NLPF offers experimental approaches to elucidate spectral substructure and the mode of pigment-pigment coupling in pigment-protein complexes: 1), by measuring NLPF spectra at low intensities (pumping and probing in the same and/or different spectral regions (27,28)); and 2), by measuring NLPF spectra at high pump intensities, NLPF may also reveal subbands with very low transition dipole moment, e.g., the  $S_1$  state of peridinin.

#### RESULTS AND DISCUSSION

## Mode of coupling between peridinins in the  $S_2$  state

To establish the prevailing mode of coupling in the peridinin  $S_2$ -absorption band, NLPF spectra were measured by pumping and probing in the respective region.

The RT-absorption spectrum of PCP is shown in Fig. 1. The Chl-absorption regions are indicated (shaded). Peridinin has a broad ( $S_0 \rightarrow S_2$ ) absorption band between ~350 and

560 nm. The Chl a Soret band absorbs in the same region at wavelengths shorter than 470 nm. The Chl  $a Q_v$ -absorption band peaks at 670 nm. It is generally assumed that peridinin does not absorb in the vicinity of the Chl  $a Q_y$  band. The overall absorption of the eight peridinins in the PCP monomer is rather featureless. There is considerable spectral overlap of the constituent subbands as well as no significant difference between the spectral widths of the peridinins (8).

NLPF spectra of PCP pumped exclusively in the peridinin  $S_0 \rightarrow S_2$ -absorption region (between 480 and 545 nm) were measured by probing either in the peridinin-alone region or in the region where Chl  $a$  also absorbs (schematically shown in Fig. 3 A). These spectra are displayed in Fig. 3, B and C. Two main features are immediately obvious:

1. NLPF spectra for  $\lambda_t$  in the range 500–530 nm (where exclusively peridinin absorbs) are uniform, and reveal no



FIGURE 3 NLPF experiment to elucidate the mode of coupling between peridinins in the  $S_2$  states. (A) Schematic: PCP is pumped in the peridinin  $S_2$ -absorption region (485–540 nm) and probed in the peridinin  $S_2$ /Chl Soret region. The shaded arrow symbolizes overall EET from peridinin to Chl a. (B) NLPF spectra obtained for probe wavelengths in the ''exclusive peridinin  $S_2$ -absorption'' region. Solid lines represent smoothed data. (C) NLPF spectra obtained for probe wavelengths in the peridinin  $S_2$ /Chl Soret region.

substructure (Fig. 3 B). The spectra in Fig. 3 B show typical characteristics of excitonically coupled pigments. Strong excitonic coupling creates a common ground state. Excitation is shared instantaneously among the pigments and in the case of spectrally strongly overlapping subbands, the resulting NLPF spectrum should have a smooth, more-or-less uniform lineshape—independent of  $\lambda_t$ . Apparently, this is the case for all four peridinins within one cluster in PCP.

2. NLPF spectra for  $\lambda_t$  in the range 455–470 nm, where both peridinin and Chl a absorb, become nonuniform and show distinct structures characterized by a shoulder at  $\sim$ 510 nm and rise steeply at wavelengths shorter than 490 nm (Fig. 3 C). These features represent a NLPF signal due to additionally probing Chl. Since peridinins, exclusively, are pumped in this experiment, these signals originate from depletion of the ground states of the two Chls due to EET from peridinin. As mentioned above, the fluorescence-excitation spectrum of PCP shows no wavelength dependence in this pump region, indicating equal EET efficiency from all peridinins to Chl. On the other hand, the additional Chl-derived NLPF signal is substructured, suggesting that the two acceptor Chls must have different excited-state lifetimes.

Apparently, the enhanced Chl-related NLPF signal observed upon pumping between 490 and 520 nm is due to peridinins transferring their energy preferentially to the Chl  $a$  with the longer  $S_1$  lifetime. These observations show that the peridinins in the two clusters are spectrally distinguishable; and that the two Chls a are not photophysically identical. However, the question of whether peridinin-to-Chl EET proceeds directly from  $S_2$  to  $Q_x$ , or via the dark state(s) of peridinin to  $Q_y$ , cannot be decided from the above experiment.

### Peridinin spectral substructure and EET to Chls

A more straightforward way to assign peridinin subbands to the two Chls in the PCP monomer is to monitor Chl  $a$  groundstate depletion in a spectral region with no peridinin absorption after initial peridinin excitation. This was accomplished by tuning  $\lambda_p$  across the peridinin-absorption region (460– 540 nm) and positioning  $\lambda_t$  in the  $Q_y$ -absorption region of Chl  $a$  (645–680 nm) (Fig. 4 A). In the case of identical peridinin subbands in both clusters, identical NLPF lineshapes, resembling the squared peridinin-absorption profile, independent of  $\lambda_t$ , would be expected—provided that EET from all peridinins to Chl is uniform. On the other hand, in the case of nonidentical peridinin spectral forms in both clusters and different Chl  $S_1$  lifetimes, peridinins transferring energy to the longer-lived Chl are expected to be enhanced in the NLPF spectrum, and vice versa.

NLPF spectra obtained for  $\lambda_t$  from the blue (645 nm) to the red (685 nm) edge of the Chl  $a Q<sub>y</sub>$  band clearly do not reproduce the squared peridinin-absorption profile—but,



FIGURE 4 NLPF experiment for subband analysis of the peridinin  $S_2$ absorption band in PCP. (A) Schematic: PCP was pumped in the peridinin  $S_2$ -absorption region and probed in the Chl a  $Q_y$  region. The shaded arrow symbolizes overall EET from peridinin to Chl a. (B) NLPF spectra for probe wavelengths between 645 and 685 nm (and pump wavelengths in the 450– 540 nm region). Note that the spectra are composed of two data sets due to tuning-range limitations of the pump lasers. Solid lines represent smoothed data. Individual NLPF spectra are displaced by a constant, offset for better representation.

instead, display a significant substructure (Fig. 4 B). Nevertheless, NLPF signals obtained for  $\lambda_p > 470$  nm are exclusively due to peridinin excitation.

Although the NLPF signal is enhanced for  $\lambda_p > 480$  nm, it is diminished—but clearly distinct from zero at shorter pump wavelengths (see also Fig. 5 B). Three peaks at  $\sim$  490, 500, and 540 nm are apparent in the former region—irrespective of  $\lambda_t$ (Fig. 4 B). There is also a local minimum at  $\sim$  515 nm, being most pronounced for  $\lambda_t$  being located at the blue edge of the  $Q_v$ band. This minimum is gradually leveled out for  $\lambda_t > 670$  nm. The latter observation indicates the existence of two subbands centered at  $\sim$  515 nm being associated with the two different Chls. The 515-nm peridinin feature associated with the longerlived (red-shifted) Chl is increased with shifting  $\lambda_t$  away from the blue (short-lived) Chl, gradually filling up the initial gap. The steep rise of the NLPF signal at the short-wavelength edge of the pump region can be assigned to direct pumping of the Chl  $B_x$  absorption (being negligible at  $\lambda > 470$  nm).

To extract quantitative spectral subband information, a combined analysis of NLPF spectrum at  $\lambda_t = 645$  nm and the



FIGURE 5 NLPF spectrum probed at 645 nm. (A) Model underlying the fit. See text for further details.  $(B)$  Fit of the NLPF spectrum.  $(C)$  Fit of the PCP-absorption spectrum in the NLPF pump region. Open circles represent the measured data, solid lines represent the best fits; the resolved absorption subbands are additionally shown in panel C.

absorption spectrum (between 450 and 560 nm) was performed. The analysis is based on the model shown in Fig. 5 A. Four excitonically coupled peridinins are associated with each Chl. The two Chls have different  $S_1$  lifetimes (and different site energies, corresponding to  $\sim$  5 nm; see below). EET between the Chls is allowed in both directions. Inhomogeneous broadening of the peridinin subbands was not taken into account. We do not differentiate between the two possible EET pathways (via peridinin  $S_2$  and Chl  $Q_x$  or the dark  $S_1$  and  $Q_y$ ) finally populating Chl  $Q_y$ , but instead combine them in one rate (assumed to be equal for all peridinins). The latter assumption is based on the fact that fluorescence excitation efficiency is uniform over the entire peridininabsorption range.

No lifetime values for the two Chl  $a$  subbands can be deduced from these data because these NLPF spectra represent only the broad-band term of the NLPF lineshape formula as shown in Eq. 1. A quantitative determination of the exact ratio would be only amenable by  $T_1$  peak (high-resolution) NLPF measurements (29). Nevertheless, lifetime ratios of different components are reflected in broad-band NLPF spectra of multipigment systems; in this particular experiment, subbands transferring energy to the longer-lived Chl-species would appear enhanced in the NLPF spectra (29).

Fits based on the assumption of equal  $S_1$  lifetimes for the Chls consistently failed. The model used to evaluate the lineshapes of the NLPF spectra in Figs. 4 and 5 implies that the two Chls  $a$  have different  $S_1$  lifetimes, in accordance with the other experimental observations (in particular, the subband analyses of the Chl  $Q_y$  band; see below). Our data indicate a signal ratio of  $\sim$ 2–3 upon probing in the blue edge of the Chl  $a Q<sub>v</sub>$  band, which is consistent with a lifetime ratio for the Chls of  $\sim$ 1.5 (as exemplarily used in the fit in Fig. 5, B and  $C$ ).

The decline of the NLPF signal at  $\lambda_p$  470 nm can have two different causes. According to earlier observations of Song et al. (24), the peridinin  $S_2$ -absorption band of native PCP from Amphidinium rhyncocephaleum splits into two components, a blue- and a red-shifted part. The more blue-shifted peridinins transfer their excitation energy apparently to the shorter-lived Chl whereas the red-shifted peridinins belong to the cluster serving the longer-lived Chl. Higher vibrational bands of the red-shifted peridinins tail into the spectral region ,480 nm. Vibrational bands generally give considerably lower contributions to the overall NLPF spectra. Superposition of the contributions from the blue-shifted peridinins and the vibrational bands leads to loss of NLPF substructure in the spectral region  $\leq 480$  nm.

The fit of the NLPF spectrum is shown as a solid line in Fig. 5 B. Three peridinin subbands (centered at  $\sim$  535, 501, and 487 nm) can be assigned to the cluster of the longer-lived Chl. A subband centered at  $\sim$  515 nm is associated with the shorter-lived Chl a. The  $\lambda_t$ -dependence of the NLPF spectra also reveals that the cluster associated with the longer-lived Chl must contain a peridinin species peaking at  $\sim$ 515 nm (see Fig. 4  $B$ ).

The peridinin subbands obtained by the fit are indicated with respect to the PCP-absorption spectrum in Fig. 5 C. The subbands partially correspond to those previously reported: Carbonera et al. (3) identified peridinin subbands at 543, 534, 518, and 485 nm and assigned them to per2, per3, per4 and per1, respectively, in the PCP structure. The peridinins were not, however, assigned to the clusters, since the authors assumed that site energies of Chls and peridinins at homologous positions in both clusters are identical. A 543-nm subband was not observed in our study due to measuring-range limitations. A pronounced subband at 501 nm—missing in the analysis in Carbonera et al. (3), as evident in the deviation between their fitted and measured CD spectra in the 500 nm region—was identified in this work. Kleima et al. (5) identified at least four peridinin subbands centered at 520, 529 (or 537), 546, and 555 nm, but found no evidence for a band at 487 nm.

### A peridinin response upon Chl  $Q_v$  excitation

It has been shown repeatedly that peridinin undergoes very efficient ( $\sim$ 90%) EET to Chl *a* after excitation of the S<sub>2</sub> state (e.g., (17)). To date, however, it is still a matter of debate whether EET proceeds directly from  $S_2$  of peridinin to Chl  $Q_x$ , via the dark state(s) of peridinin to  $Q_y$  of Chl, or to what extent via both channels. Moreover, the mode of interaction (Coulombic versus Dexter mechanism) is not yet established. Furthermore, the possibility of "mixing" between the Chl  $S_1$ state and low-lying states of peridinin has been raised (31).

Interactions between  $Q_y$  of Chls and peridinin dark states should be apparent in a ground-state depletion of the respective peridinin after excitation of the Chls. By pumping in the Chl  $a \, Q<sub>v</sub>$  region and probing at different wavelengths in the peridinin-alone region (shown schematically in Fig. 6 A), such interaction should generate a NLPF signal corresponding to the  $S<sub>2</sub>$  energies of interacting peridinins. A comparable approach has been successfully applied by the authors recently to provide evidence for excitonically coupled Chls  $a$  and  $b$  in the antenna complexes CP29 (27) and LHC II (28).

Fig. 6 B shows a quasi-three-dimensional plot of the NLPF spectra obtained by pumping between 650 and 685 nm (in the Chl  $a Q<sub>v</sub>$  region) and probing between 450 and 530 nm (Chl a Soret/peridinin  $S_2$ -absorption region). The NLPF signal for  $\lambda_t$  in the Chl Soret/peridinin-absorption region would be expected to decrease in parallel to the Chl  $a B_x$  absorption, declining to baseline at wavelengths  $>470$  nm. However, an additional NLPF signal is observed for  $\lambda_t$  around 490 nm and can be attributed to a depletion of the peridinin ground state due to Chl *a*-peridinin interaction. Under our experimental conditions, intersystem crossing into the Chl triplet-manifold after Chl  $Q<sub>y</sub>$  excitation would be possible, in principle. Triplet EET from Chl  $a$  to peridinin was shown to be very effective in PCP with a peridinin triplet-absorption rise-time of  $17 \pm 7$  ns (4,5). Rapid equilibration of triplet excited states among the peridinins of one cluster was demonstrated for PCP by

ODMR (2). However, if a peridinin triplet population were the origin of the NLPF signal, it should be observable in the entire  $\lambda_t$ -range. This is clearly not the case; in particular, no NLPF signal is detectable for  $\lambda_t$  > 510 nm (see also Fig. 6, B and C).

Lineshapes of the NLPF spectra probed in the immediate vicinity of 490 nm (in contrast to all other spectra) resemble the squared Chl  $a Q<sub>v</sub>$  absorption. We assign these NLPF signals to the 487-nm peridinin species associated with the longer-lived Chl identified by the NLPF measurements described in ''Peridinin spectral substructure and EET to Chls'', above. The mechanism of interaction between the excited Chl and peridinin cannot be readily deduced on the basis of the current experiment. However, it has been suggested from structurebased calculations that an electron exchange (Dexter-type) mechanism is far less effective than the Coulomb mechanism in PCP (6,32). In addition, breaking of the  $C_{2h}$  symmetry by substitution of the polyene chain in peridinin is expected to result in a significant dipole moment of the  $S_0 \rightarrow S_1$  transition (amounting to  $\sim$  5% of that of  $S_0 \rightarrow S_2$ ) (33). The latter prediction is supported by the observation of weak fluorescence from  $S_1$  of peridinin in solution (11,34). Moreover, distortion of the linear structure of the peridinins upon binding to the apoprotein may further contribute to  $C_{2h}$ -symmetry breaking (1,8).

Assuming that a Coulombic mechanism was operative, a NLPF signal is expected in the spectral region where donor fluorescence and acceptor-absorption spectra overlap. Thus, viewing along the  $\lambda_p$  axis in Fig. 6 B (the lineshape of the observed NLPF signals will indicate the  $S_1$  energies of the(se) peridinin(s); see also Fig. 6 C), the maximum of the NLPF signal will be red-shifted or blue-shifted—if peridinin  $S_1$  were lower or higher energetic, respectively, than  $S_1$  of Chl *a*. Since the NLPF spectra probed at  $\sim$ 490 nm have maxima corresponding to the Chl  $a \ Q_{v}$  absorption, we suggest that the dark state is approximately isoenergetic to or slightly above  $S_{1,0}$  of Chl a.



FIGURE 6 NLPF experiment to elucidate interactions between Chl a and peridinin  $S_1$  states. (A) Schematic: PCP was pumped in the Chl a  $Q<sub>y</sub>$  region and probed in the Chl a-Soret/peridinin  $S_2$ -absorption region. The shaded arrow symbolizes EET from Chl a to peridinin. (B) Quasi-three-dimensional representation of the (smoothed) NLPF spectra. (C) Intensity courses (along the probe wavelength range) of the NLPF signals pumped at  $\sim$  670 nm (solid) and 650 nm (shaded).

In addition to the features indicating interaction between  $S_1$  of Chl *a* and a dark state of peridinin, another NLPF signal (peaking at  $\sim$  500 nm) is observed upon pumping at 650 nm (Fig. 6 C). Fig. 6 C compares the two cross-sections (for  $\lambda_p =$ 650 and 670 nm) along the  $\lambda_t$  range. The steep slope at the blue edge of the curve obtained for  $\lambda_p = 670$  nm suggests an assignment to a NLPF signal due to pumping and probing Chl. Upon shifting  $\lambda_p$  to 650 nm, this feature is clearly lacking, indicating that Chl is not pumped. Obviously, a low dipole moment transition of peridinin rather than Chl is directly pumped at  $\sim 650$  nm.

Such a peridinin  $S_1/ICT$  energy would be lower than any value reported so far. In a recent article, Shima et al. (8) locate the  $S_{1,0}$  energy for peridinin dissolved in  $CS_2$  at 16,200 cm<sup>-1</sup> ( $\sim$  617 nm) in accordance with previous studies (11,34). Zimmermann et al. (15) deduced from two-photon fluorescence excitation experiments that  $S_1$  of peridinin in benzene as well as in PCP is located even more to the blue (peaking at  $\sim$ 530 nm), mixing with the  $B_{\rm u}^+$  state. Their assignment is apparently in contradiction to one-photon fluorescence studies ((11,34); see also (8)). Shima et al. (8) conclude that the two-photon excitation spectra in Zimmermann et al. (15), as well as their own, are essentially due to the  $S_2$  =  $1Bu<sup>+</sup>$  state carrying a significant two-photon character in peridinin. Furthermore, it was concluded that the  $S_0 \rightarrow S_2$ transitions of peridinins Per612 and Per622 are considerably blue-shifted as compared to the other peridinins and characterized by a  $2Ag^-$  state lying above  $S_2$  (8). The latter prediction is hard to reconcile with the experimental results presented here. Apparently, at least one of the blue-shifted peridinin species, peaking at  $\sim$ 495 nm, shows unique interaction with the  $S_1$  state of Chl a mediated via the carotenoid's low-lying  $S_1/\text{ICT}$  state.

### Spectral substructure of the Chl  $a Q<sub>v</sub>$  absorption

As mentioned above, the distance between the two Chls  $a$  in a PCP monomer is fairly large and interaction between the Chls appears to be weak. Consequently, Chl site energies are determined essentially by pigment-protein interactions. The Chl  $a$   $Q_v$ -absorption region was investigated by positioning the probe beam at different  $\lambda_t$  (640–680 nm) and tuning  $\lambda_p$  at low intensity across the same region (exemplary spectra are shown in Fig. 7). Two main features are observable in these spectra:

- 1. The lineshape of the measured NLPF spectra does not correspond to the square module of the Chl  $a \, Q_{v}$ -absorption profile (exemplarily shown for  $\lambda_t = 670$  nm, Fig. 7 C).
- 2. Lineshapes of the NLPF spectra for  $\lambda_t$  < 665 nm (Fig. 7 D) deviate from those for  $\lambda_t$  between 665 and 680 nm. An additional signal component is observed at the blue edge (exemplarily shown in Fig. 7 D).

The observation that the NLPF spectra do not correspond to the squared absorption profile indicates spectral heterogeneity of the  $Q_y$  band of PCP (meaning that two subspecies exist). However, these subbands are not immediately obvious.

To further characterize the underlying substructure, a global line-shape analysis comprising a fit of the set of NLPF spectra (obtained at different  $\lambda_t$ ) and the absorption spectrum was performed. The procedure is described in detail in Voigt et al. (26).

Fits with only one Chl a  $Q_y$  band yielded an  $\chi^2 = 4.46$ and consistently failed. However, a model with two Chl a-subbands results in an  $\chi^2 = 2.5$  and was sufficient to describe all NLPF spectra for  $\lambda_t$  in the range from 665 to 680 nm and the



FIGURE 7 Spectral substructure analysis of the Chl a  $Q_v$ -absorption region. Exemplary spectra are shown: (A) NLPF spectrum obtained at  $\lambda_t = 680$  nm. (B) NLPF spectrum obtained at  $\lambda_1 = 660$  nm. (C) The NLPF spectrum obtained at  $\lambda_1 = 670$  nm is compared to the squared absorption spectrum (*dashed line*). (D) NLPF spectrum obtained at  $\lambda_1 = 650$  nm, comparison of two and three subband fits. Solid lines indicate fits assuming two subbands as a result of a global analysis of the spectra obtained for  $\lambda_1 > 665$  nm, including the absorption spectrum. The shaded line represents a fit with an additional (third) subband.

 $Q_v$ -absorption spectrum of PCP simultaneously. An F-test revealed a probability  $\gg$ 99.9% that the two-subband approach represents an improved model as compared to that of a single band. The best global fit was obtained with center wavelengths of 669 and 675 nm and widths of 320 and 200  $\text{cm}^{-1}$  (full-width at half-maximum) for the two Chl  $a Q<sub>v</sub>$  subbands, respectively. The shorter-wavelength band will consequently have a shorter effective excited-state lifetime. The latter requirement is in line with the results of NLPF experiments to elucidate EET from peridinin to Chl  $a$  (obtained by pumping in the allowed peridinin band and probing in the Chl  $a Q_v$  band; compare to Fig. 4 B).

When attempting to fit the NLPF spectra obtained at  $\lambda_t =$ 650 and 655 nm, a third subband had to be taken into account. The fit, however, did not identify the peak wavelength of the additional band as precisely as the other two. Nevertheless, the third band must have a center wavelength of  $\sim$  660 nm, but cannot be interpreted as a further Chl- $Q<sub>v</sub>$ -absorption subband, since only two (noninteracting) Chls are found in a PCP monomer.

A distinct NLPF signal originating from peridinin groundstate depletion was observed upon pumping in the Chl  $Q_v$ region and probing in a spectral region  $($ >470 nm) where exclusively peridinin absorbs (see previous section). On the basis of the above results, it cannot be decided whether this peridinin ground-state depletion is a result of a direct excitation of a low dipole moment  $S_0 \rightarrow S_1/ICT$  transition of certain peridinins and/or due to EET from a neighboring Chl.

In any case, the(se) peridinin(s) must have a non-negligible  $S_0 \rightarrow S_1/\text{ICT}$  transition dipole moment and show considerable spectral overlap with the Chl  $a Q_v$  band. An  $S_1$  transition dipole moment of up to five Debye (depending on parameterization) has been inferred previously from theoretical considerations of breaking of the  $C_{2h}$  symmetry for peridinins in PCP (6). Moreover, Krueger et al. (13) have deduced a peridinin  $S_1$  transition dipole moment of approximately three Debye in PCP from transient absorption measurements. However, a dipole moment of that size should be detectable in absorption—which is in contradiction to experimental results with peridinin in solution and the interpretation of the PCP  $Q_v$ region absorption spectrum by previous workers (e.g., Kleima et al. (5)). Nevertheless, a low-dipole moment peridinin transition may become visible in NLPF spectra, measured at strong pump intensities exceeding the range of validity of the  $\chi^{(3)}$ approach for the dominant Chls (see also Beenken and May (30)).

## Additional evidence for spectral heterogeneity of the Chl  $Q_y$  region

Another approach to reveal spectral heterogeneity and (hidden) low dipole-moment species in complex spectra is provided by the pump-intensity dependence of the NLPF spectral profiles. In the case of Chl spectral heterogeneity (including different  $S_1$  lifetimes of the subbands), contributions of the (two) spectral subbands to the overall NLPF spectrum are expected to vary with pump intensity (30). With increasing pump intensity, the subband with the longest  $S_1$  lifetime (usually the lowest energetic band) becomes progressively saturated, and contributions from other, shorter-lived and/or low dipole-moment bands may become apparent (see also Schubert et al. (35)). Hence, a blue-shift of the overall NLPF spectrum, in addition to an increase in amplitude, would be expected. Both phenomena are observed with PCP; Fig. 8 A shows NLPF spectra of PCP obtained at pump intensities between  $1.5 \times 10^{22}$  and  $4.5 \times$  $10^{24}$  photons cm<sup>-2</sup> s<sup>-1</sup> (probed at  $\lambda_t = 660$  nm). Note that the spectra obtained at lower pump intensities (traces  $a-c$  in Fig. 8 A) are magnified 250 times. The NLPF spectra display a clear shift toward shorter wavelengths with increasing pump intensity. The dependence of the NLPF signal



FIGURE 8 Pump intensity dependence of NLPF spectra of PCP. (A) NLPF spectra probed at  $\lambda_1 = 660$  nm (smoothed). Actual data points are shown exemplarily in trace g (open circles, with typical SD). Spectra obtained at the three lowest pump intensities  $(a-c)$  are magnified 250-fold. Intensity variation was between  $1.5 \times 10^{22}$  (a) to  $4.5 \times 10^{24}$  (g) photons cm<sup>-2</sup> s<sup>-1</sup>. (B) Intensity dependence of the NLPF signal amplitude pumped and probed at 680 nm (*open*) symbols) and at 660 nm (solid symbols). Dotted line indicates strict quadratic intensity dependence. (C) Normalized NLPF spectra from A.

amplitude on pump intensity is shown for  $\lambda_t = 660$  and 680 nm (i.e., on the *blue* and *red edges* of the PCP  $Q_v$ -absorption region, respectively) in Fig. 8 B. The NLPF signal obtained at  $\lambda_t = 660$  nm displays later commencement of saturation effects (leading to deviation from the almost quadratic pump intensity dependence) than the signal at  $\lambda_t = 680$  nm, indicating heterogeneity of the  $Q_v$ -absorption region.

This blue-shift is even more obvious when the NLPF spectra are normalized (Fig. 8 C). The moderate shift (observed in the NLPF spectrum at intermediate intensity in trace d) is consistent with the existence of two subbands with different excited-state lifetimes and different (by a few nanometers) center-wavelengths. Traces  $f$  and  $g$  in Fig. 8  $C$ display a more pronounced blue-shift (toward  $\sim$  660 nm). A further blue-shift at even higher pump intensities is not expected since saturation of the NLPF signal is observed in this pump intensity range (Fig. 8 B). The shift, however, cannot be explained on the basis of the two Chl a subbands at 669 and 675 nm but requires a third, low dipole moment band centered at  $\sim 660$  nm. The blue-shift is also observed upon probing at 680 nm, but at this  $\lambda_t$  it is, however, less pronounced (data not shown).

The latter observation would be in line with the different slopes and saturation behavior of the NLPF signals observed in the blue and red edges of the  $Q_v$  band, and is consistent with the requirement of a third subband to fit the low pumpintensity NLPF spectra obtained at  $\lambda_t = 650$  and 655 nm. The  $\sim$ 660-nm band is tentatively assigned to the S<sub>1</sub>/ICTstate of peridinin(s) and most likely corresponds to the peridinin molecule with a blue-shifted  $S_2$  state (at 495 nm). Assignment of this subband to the previously resolved  $S_1 \rightarrow$  $S_n$  transient absorption (10) can be excluded by a comparison of the respective maximum positions ( $\sim$ 590 vs. 660 nm).

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