Mixed Exciton–Charge-Transfer States in Photosystem II: Stark Spectroscopy on Site-Directed Mutants

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ABSTRACT We investigated the electronic structure of the photosystem II reaction center (PSII RC) in relation to the light-induced charge separation process using Stark spectroscopy on a series of site-directed PSII RC mutants from the cyanobacterium *Synechocystis* sp. PCC 6803. The site-directed mutations modify the protein environment of the cofactors involved in charge separation (P$_{D1}$, P$_{D2}$, Chl$_{D1}$, and Phe$_{D1}$). The results demonstrate that at least two different exciton states are mixed with charge-transfer (CT) states, yielding exciton states with CT character: (P$_{D2}$$^{+}$+P$_{D1}$$^{−}$−Chl$_{D1}$)$^{+673}$nm and (Chl$_{D1}$$^{+}$+Phe$_{D1}$$^{−}$)$^{+681}$nm (where the subscript indicates the wavelength of the electronic transition). Moreover, the CT state P$_{D2}$$^{+}$P$_{D1}$$^{−}$ acquires excited-state character due to its mixing with an exciton state, producing (P$_{D2}$$^{−}$P$_{D1}$$^{+}$)$^{−684}$nm. We conclude that the states that initiate charge separation are mixed exciton-CT states, and that the degree of mixing between exciton and CT states determines the efficiency of charge separation. In addition, the results reveal that the pigment-protein interactions fine-tune the energy of the exciton and CT states, and hence the mixing between these states. This mixing ultimately controls the selection and efficiency of a specific charge separation pathway, and highlights the capacity of the protein environment to control the functionality of the PSII RC complex.

INTRODUCTION

The primary photochemical event in photosynthesis is light-induced charge separation. This process takes place in the membrane-bound, photochemically active reaction center (RC), a pigment-protein complex found in nonoxygenic photosynthetic bacteria and oxygen-evolving organisms (cyanobacteria, algae, and higher plants). The charge-separated state created in the RC of photosystem II (PSII) drives water splitting and generates an electrochemical gradient across the membrane that is used to generate ATP and reduce NADP$^+$. Understanding the molecular mechanisms that lead to charge separation is crucial because the success of photosynthesis depends on the efficiency of this process.

Primary charge separation in PSII can be studied in isolated PSII RC complexes that comprise the D$_1$-D$_2$-Cyt b$_{559}$ subunits, as well as in larger core complexes. Structural studies on PSII core complexes (1–5) have confirmed that four chlorophyll (Chl), two pheophytin (Phe), and two quinone (Q) molecules bound to the D$_1$-D$_2$ heterodimeric RC complex are arranged in two symmetric branches (D$_1$ and D$_2$) across the membrane in the center of the complex (Fig. 1). Two additional Chl molecules, Chl$_{Z1}$, are located at opposing sites on the periphery of the D$_1$-D$_2$ complex, with two β-carotene molecules situated between the peripheral Chl$_{Z2}$ and the center of the complex. The spatial distribution of the central cofactors (Phe$_{D1}$, Chl$_{D1}$, P$_{D1}$, P$_{D2}$, Chl$_{D2}$, and Phe$_{D2}$ (where the subscript indicates the protein branch where the cofactors are localized)) gives rise to a system of coupled pigments that interact, creating collective excited (exciton) states with contributions from various cofactors. In addition, the system is not static. Fast nuclear motions (intra- and interpigment vibrations and protein vibrations) and slow conformational motions of the protein produce homogeneous and inhomogeneous broadening, respectively, of the electronic transitions. Therefore, after a photon is absorbed by the RC, the excitation energy is distributed among the cofactors (depending on the protein conformation or disorder) and converted into a charge-separated state after a series of energy and electron transfer reactions. This means that charge separation can be initiated from different combinations of cofactors (6,7) giving rise to different charge separation pathways (8–13).

Theoretical (11,13) and experimental (12) studies have demonstrated the presence of at least two different low-energy exciton states—(Chl$_{D1}$Phe$_{D2}$)$^{+}$ and (P$_{D2}$P$_{D1}$Chl$_{D1}$)$^{+}$—in the PSII RC of higher plants, leading to two different charge separation pathways (note that the cofactors sequence corresponds with their order of participation in the excitonic wavefunction). The choice of a certain charge separation pathway by a single RC complex in the ensemble is strongly dependent on disorder (protein conformation). The sequences of states for the two charge separation pathways, the so-called Chl$_{D1}$ and P$_{D1}$ paths, are as follows:

Chl$_{D1}$ path : (Chl$_{D1}$Phe$_{D1}$)$^{+}$$\rightarrow$Chl$_{D1}$$^{+}$Phe$_{D1}$$^{+}$$\rightarrow$P$_{D1}$$^{+}$Phe$_{D1}$$^{-}$

P$_{D1}$ path : (P$_{D1}$P$_{D2}$Chl$_{D1}$)$^{+}$$\rightarrow$P$_{D2}$$^{+}$P$_{D1}$$^{-}$$\rightarrow$P$_{D1}$$^{+}$Chl$_{D1}$$^{-}$$\rightarrow$P$_{D1}$$^{+}$Phe$_{D1}$$^{-}$

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The Stark spectrum \( \text{Abs}_{\text{Field on}} \) minus \( \text{Abs}_{\text{Field off}} \) for isolated absorption bands in a nonoriented, immobilized sample is described by the Liptay formalism as a linear combination of the zeroth, first, and second derivatives of the absorption spectrum \( \text{Abs} \). The molecular parameters \( \Delta \alpha \) and \( \Delta \mu \) scale with the first and second derivatives, respectively, of the absorption spectrum (the classic Stark effect). The shape of the Stark spectrum of PSII RC from higher plants was shown in a previous study \((17)\) to be dominated by the second derivative of the absorption spectrum (2DerAbs) and therefore by \( \Delta \mu \). However, in that work deviations from the classic Stark effect were found at wavelengths > 675 nm where the Stark spectrum was red shifted with respect to the 2DerAbs (the nonclassic Stark effect, which is not included in the Liptay formalism). This highlights the limitations of the Liptay formalism when applied to the PSII RC: it cannot describe either the presence of excitonic interactions or the mixing of excitons and CT states (nonclassic Stark effects). To investigate these nonclassic effects and their consequences on the Stark line shape and amplitude, a more sophisticated theoretical model is needed. To that end, Novoderezhkin et al. \((11)\) developed an exciton model based on a quantitative simultaneous fit of the absorption, linear dichroism, circular dichroism, steady-state fluorescence, triplet-minus-singlet, and Stark spectra together with the spectra of pheophytin-modified RCs, and so-called RC5 complexes that lack one of the peripheral Chls. The line-shape functions were expressed in terms of the modified Redfield approach. In this model, the excited-state manifold consists of the excited states of the eight chlorine cofactors (PheD1, ChlD1, Pd1, Pd2, ChlD2, PheD2, ChlZD1, and ChlZD2) and one CT state. Of most importance, excitonic interactions, exciton-CT mixing, and coupling to phonons (protein motions) are explicitly included in the model, which allows a consistent modeling of the whole set of experimental data using a unified physical picture. As a result, by including or excluding the excitonic interactions and the mixing of excitons and CT states, the authors were able to investigate the consequences of these nonclassic Stark effects on the line shape and amplitude of the Stark spectrum. To summarize, the fitting of the Stark PSII RC spectrum according to this model shows that 1), excitonic interactions produce band shifts in the Stark spectrum; and 2), exciton-CT mixing produce increased Stark band amplitudes when compared with the 2DerAbs \((11)\). Therefore, comparing the 2DerAbs with the Stark spectrum is a useful tool for qualitatively interpreting the Stark spectrum. Novoderezhkin et al. \((11)\) also showed that the Stark spectrum line shape depends on the energy and configuration of the CT states.

The information contained in the Stark spectra of PSII is difficult to extract due to the spectral congestion in the Qy absorption region. To overcome this problem, we chose to use a combined approach: Stark spectroscopy applied to site-directed mutants. In such systems, a specific amino
acid mutation alters the protein environment of a cofactor, i.e., the energy of the exciton and primary RPs states in which the cofactor participates, as well as the exciton-CT mixing. As a result, the line shape and amplitude of the Stark spectra of the mutants changes with respect to the wild-type (WT). These changes can be directly correlated with the energetic changes induced by the mutation as observed in the absorption spectrum, allowing a band assignment that finally leads to a description of the charge separation mechanisms. The mutations studied modify the protein environment of the cofactors involved in charge separation (PheD1, P1D1, P2D2, and ChlD1).

Mutants created at residue D1-130 alter the strength of the hydrogen bond to the 13′-keto carbonyl of PheD1 (Fig. 1), which leads to significant effects on the quantum yield of primary RP formation (ΦRp) in the isolated PSII RC from Syn. In Syn. the residue in position D1-130 is naturally a glutamine (Gln) residue, whereas in higher plants it is a glutamate (Glu). Transient absorption measurements by Giorgi et al. (18) indicated that ΦRp higher plants > ΦRp D1-Gln130Glu mutant > ΦRp Syn. WT. The authors, by considering the ΦRp higher plant equal to unity, showed that ΦRp D1-Gln130Glu mutant ≈ 0.75 and ΦRp Syn. WT ≈ 0.55. The complete removal of the hydrogen-bond interaction in the D1-Gln130Leu results in an even lower ΦRp (≈0.35).

Due to the chemical structure of these residues (Gln contains an amide and Glu contains a carboxylic acid), the strength of the hydrogen bond to the 13′-keto carbonyl of PheD1 is stronger in higher plants than in cyanobacteria, as shown by Shibuya et al. (19) using FTIR spectroscopy. These authors also showed that the redox potential (Ered) of Phe with the 13′-keto carbonyl hydrogren-bonded with Glu is less negative compared with Gln by 84 mV (19). This means that the strengthening of the hydrogen-bond interaction facilitates the reduction of Phe. In the study presented here, the absorption and Stark spectra of the cyanobacterium Syn. PSII RC isolated from the WT and D1-Gln130Glu mutant strains were compared with those of the PSII RC isolated from a higher plant (spinach).

We investigated the role of the Chls involved in charge separation (P1D1, P2D2, and ChlD1) in seven site-directed mutants that modify the protein environment of P1D1 (D1-His198Gln, D1-His198Asn, and D1-His198Ala), P2D2 (D2-His197Gln, D2-His197Asn, and D2-His197Ala), and ChlD1 (D1-Thr179His; see Fig. 1). All of the introduced mutations allow RC assembly and complete chromophore binding (18,20,21). For the P1D1 and P2D2 mutants, the histidine (His) that axially coordinates P1D1 and P2D2 is exchanged by glutamine (Gln), asparagine (Asn), or alanine (Ala). The electronic polarizability of the π-system of the imidazole moiety of His stabilizes the excited states of nearby cofactors by dispersive interactions. Therefore, in these mutants, the site energy of P1D1 or P2D2 and consequently the exciton states in which P1D1 and/or P2D2 participate are expected to shift to the blue (20). In addition, the D2-His197 mutation is also likely to perturb the energy of the exciton states containing ChlD1 because the His residue removed is indirectly hydrogen bonded to the 13′-keto carbonyl of ChlD1 through a water molecule (5). The introduced residues, Gln and Asn, contain an amide moiety that could, directly or indirectly through a water molecule, coordinate the central magnesium of P1D1 and P2D2, or create hydrogen-bond interactions with nearby carbonyls, whereas this is not possible for the alkyl side chain of the Ala residue. For the D1-Thr179His mutant, the replacement of threonine (Thr), which axially coordinates ChlD1 via a water molecule (5), with His is expected to stabilize the ChlD1 site energy and consequently the exciton states in which ChlD1 participates, producing a red shift in the absorption spectrum of the mutant (21). Previous spectroscopic analyses of these mutants (20,21) led to the assignment of 1), the 683 nm transition to the lowest exciton state in the RC localized mainly on ChlD1; and 2), the 673 nm transition to the low-energy exciton state formed mostly by P1D1 and P2D2.

We report here the comparison of the low-temperature (77 K) absorption and Stark spectra of WT and eight site-directed mutants. The results show that the electronic states that initiate charge separation in PSII RC are mixed exciton-CT states: (P D1+P D2−ChlD1)+673 nm− (ChlD1+PheD1−)+681 nm−, and (P D2+P D1−ChlD1)+684 nm− (where the subscript indicates the wavelength of the electronic transition).

**MATERIALS AND METHODS**

**Sample preparation**

PSII RC samples from *Synechocystis* sp. PCC 6803 were prepared as described previously (18). PSII core samples from *Synechocystis* sp. PCC 6803 were prepared as described previously (20,22). The PSII RC isolated from spinach was prepared as described previously (23). The monomeric PSII core samples were analyzed and purified by fast-phase liquid chromatography using a size-exclusion column to eliminate possible PSI contamination and contributions of free Chls and carotenoids to the spectra. The samples were PSI-free, but some contained a significant amount of carotenoid aggregates. The purified samples were diluted in a buffer containing 50 mM MES pH 6.5, 20 mM CaCl2, 5 mM MgCl2, 100 μM K3Fe(CN)6, 0.03% β-DM, and 57% glycerol (v/v). K3Fe(CN)6 was added to oxidize Qa to the neutral state and hence keep the RCs open for charge separation. The PSII RC samples were diluted in a buffer containing 20 mM Bis-Tris pH 6.5, 20 mM NaCl, 5 mM MgCl2, 0.09% β-DM, and 57% glycerol (v/v). The addition of K3Fe(CN)6 is not necessary for the RC samples because Qa is lost during the isolation procedure.

**Spectroscopy**

Absorption and Stark spectra were simultaneously recorded at 77 K in a home built setup as described previously (24,25). The optical density of the samples in the Stark cell (thickness 80 μm) was ≈0.7 cm−1 at 675 nm. The Stark cell was immersed in liquid nitrogen in an Oxford liquid nitrogen cryostat. The Stark cell was rotated 45° with respect to the propagation direction of the horizontally polarized measuring light to set the angle between the electric field component of the light and the externally applied electric field, χ, to the magic angle (54.7°). The Stark spectra
The spinach RC, which contains a Glu at position D1-130, the maximum of the Phe QX band peaks at 543.5 nm. The amplitude of the Phe QX band is very similar in the three RC preparations, indicating that the normalization procedure is correct.

**Stark spectra**

The differences in composition between the Syn. RC complexes and between Syn. and spinach RC have a minor effect on the Stark spectra due to the small contribution to the Stark spectra of the CP47 antenna complex in the absorption regions of interest (with respect to the contribution to the Stark spectra of the RC, see Fig. S1 in the Supporting Material). The Stark spectra of the three RC samples resemble the second derivative of the absorption spectra (2DerAbs; Fig. S2), indicating that, to a first approximation, these Stark spectra are dominated by a change in dipole moment, $\Delta \mu$, between the ground and excited states associated with the electronic transitions. A similar Stark effect was observed in the bacterial RC (26–28) and in the previously reported Stark spectrum of higher plant PSII RC (spinach) (17). In the PSII RC, the Stark spectra are dominated by the excitonically coupled (and coupled to CT states) cofactors located in the active branch (P51, ChlD1, and PheD1) and inactive branch (P52). The monomeric Chls$_{Z}$, ChlD$_{2}$, and PheD$_{2}$ cofactors are not expected to give a significant Stark response (17) because they do not substantially contribute to the collective exciton states (11) that initiate charge separation, i.e., they are not supposed to be mixed with CT states.

The Stark spectrum of the Syn. RC WT in the QY absorption region is composed of two negative bands at 672.5 and 680.5 nm, three maxima at 677 and 685 nm, and a shoulder at 689.5 nm. For the D1-Gln130Glu (spinach-like) mutant, the positions of the Stark bands are virtually identical to those of the WT. The main difference is the twofold increase in amplitude of the negative band at 680.5 nm in the mutant. The Stark spectrum of the spinach RC WT in the QY absorption region is composed of two negative bands at 670.5 and 681 nm, and three maxima at 663, 676.5, and 687.5 nm, as reported previously (17). The Stark band shifts in the spinach RC with respect to the Syn. samples may be due to small differences in the energy of the exciton states between the two organisms. The amplitude of the 670.5 nm Stark minimum in the spinach RC is similar to its Syn. counterparts, whereas the amplitude of the 681 nm minimum is similar to the 680.5 nm minimum in the D1-Gln130Glu mutant. It is interesting to note that upon replacement of Gln with Glu in the D1-Gln130Glu mutant, the amplitude of the 680.5 nm Stark minimum resembles the amplitude of the spinach RC, which also contains a Glu in position D1-130. In the Phe QX absorption region, an amplitude increase of the PheD1 Stark minima is also clearly observed for the D1-Glu130 samples (Syn. mutant and spinach RC WT) with respect to the

**RESULTS AND DISCUSSION**

**PheD1 mutant**

The simultaneously recorded 77 K absorption and Stark spectra of the Syn. PSII RC WT and D1-Gln130Glu mutant, and of spinach PSII RC WT (D1-Glu130) normalized to the RC content are shown in Fig. 2. A comparison of these spectra provides information about the effect of the hydrogen-bond interaction between the residue at position D1-130 and the 13$^\text{1}$-keto carbonyl of PheD1 on the efficiency of the charge separation process.

**Absorption spectra**

The amplitude of the Syn. RC samples’ absorption is higher with respect to the spinach RC because the Syn. RC samples are not completely pure. They contain approximately one CP47 antenna complex per two RC complexes, i.e., they are RC-enriched preparations (hereafter called RC complexes). The small differences in the antenna content of the Syn. RC complexes hinder the comparison of the WT and mutant absorption line shape. In the Phe QX absorption region, the replacement of Gln in WT for Glu in the D1-Gln130Glu (spinach-like) mutant produces a 2.5 nm red shift of the Phe QX band at 540 nm in WT to 542.5 nm in the mutant, as observed previously (18). For
D1-Gln130 sample (Syn. RC WT). For the Syn. RC WT, the PheD1 Qx Stark minimum is at 540 nm (i.e., no shift in position between Stark and absorption bands) and its amplitude is $5 \times 10^{-6}$ ($\Delta Abs$/a.u.), whereas for the D1-Gln130Glu mutant and the spinach RC WT, the PheD1 Qx Stark minima are at 542.5 and 543.5 nm (0.5 nm red shift of the Stark spectrum with respect to the absorption band) and their amplitudes are $13 \times 10^{-6}$ and $15 \times 10^{-6}$ ($\Delta Abs$/a.u.), respectively.

When interpreting these results, it is important to remember that the replacement of Gln at position D1-130 by Glu in the D1-Gln130Glu mutant increases the quantum yield of RP formation (18). Remarkably, the increase of $\Phi_{RP}$ correlates with the increase of the Stark minimum amplitude at 680.5–681 nm. In addition, we observe a similar trend when we compare the Stark spectra with the 2DerAbs spectra (Fig. S2): the larger the Stark amplitude with respect to the Stark spectra with respect to that of the RC is approximately $X$ ($\Delta Abs$/a.u.). This spectral shape corresponds to new absorption bands that appear to the red edge of the 680.5–681 nm Stark minimum. It is important to remember that the replacement of Gln at position D1-130 by Glu in the D1-Gln130Glu mutant increases the quantum yield of RP formation (18). Remarkably, the increase of $\Phi_{RP}$ correlates with the increase of the Stark minimum amplitude at 680.5–681 nm. In addition, we observe a similar trend when we compare the Stark spectra with the 2DerAbs spectra (Fig. S2): the larger the Stark amplitude with respect to that of the RC is approximately $X$ ($\Delta Abs$/a.u.).

The presence of two states contributing to the 680.5–681 nm Stark minimum is further supported by the 2DerAbs spectra in which two separated minima are present (Fig. S2). According to our model (11), the CT state $P_{D2}^{+}P_{D1}^{-}$ is strongly mixed with the $(P_{D2}^{-}P_{D1}^{+})^{*}$ exciton state. In this case, the CT state borrows dipole strength from the exciton, producing a red-most state $(P_{D2}^{-}P_{D1}^{+})^{*}$ with an absorption centered around 684 nm (where $\delta*$ indicates the fraction of dipole strength acquired by the CT state). Due to its CT character, this state is expected to contribute to the Stark spectra, which will be dominated by $\Delta u$ and therefore will resemble the second derivative of the absorption spectra. Thus, we suggest that the $(P_{D2}^{-}P_{D1}^{+})^{*}$ state contributes to the red edge of the 680.5–681 nm Stark minimum. It appears as a positive shoulder at 689.5 nm in the Syn. RCs, whereas it appears as a negative shoulder around 684 nm in the spinach RC.

**Pd1, Pd2, and Chld1 mutants**

The simultaneously recorded 77 K absorption and Stark spectra of Syn. core complex for the WT, D1-His198Gln, and D2-His197Gln mutants; WT, D1-His198Asn, and D2-His197Asn mutants; WT, D1-His198Ala, and D2-His197Ala mutants; and WT and D1-Thr179His mutant in the Qy region are shown in Fig. 3.

The effect of these mutations on the Phe Qx region is shown and discussed in the Supporting Material (see “The Phe Qx absorption band” section, Fig. S3, and Table S1).

The PSII core complex analyzed in this section contains the photochemically active RC as well as the core antenna proteins CP43 and CP47. In higher plants (spinach) for which the RC, CP43, and CP47 isolation procedures are well established (23,29–32), the amplitude of the antenna Stark spectra with respect to that of the RC is approximately equal around 670 nm and presents an $\approx$4-fold decrease around 680 nm (Fig. S1). Taking this finding and the fact that the mutations only affect the cofactors in the RC into account, we will discuss the core complex absorption and Stark spectra in terms of changes within the RC.

**Absorption spectra**

As a first approximation, we assume that the mutations influence only the energy of the electronic transition of a certain exciton state, and not the transition dipole moment strength or direction. To examine the energy shifts, we normalized the absorption spectra of WT and mutants to the cofactor content and calculated the absorption difference spectra ($Abs_{\Delta \text{dif}} = Abs_{\text{mutant}} - Abs_{\text{WT}}$; Fig. 3). In the Abs$_{\Delta \text{dif}}$ spectra, the negative bands correspond to absorption bands that disappear in the WT upon mutation, and the positive bands correspond to new absorption bands that appear upon mutation.

Three different mutations were constructed at positions D1-His198 and D2-His197, which axially coordinate $P_{D1}$ and $P_{D2}$, respectively. The absorption changes induced
by the exchange of His for the amide residues, Gln and Asn, are similar for the D1 branch mutants, as well as for the D2 branch mutants (Fig. 3). In the four mutants (D1-His198Gln, D1-His198Asn, D2-His197Gln, and D2-His197Asn), two minima at around 675 and 683 nm, and one maximum at around 666 nm with different relative amplitude and width are observed in the absorption difference spectra (Fig. 3). For the two D1-His198 mutants, the main change upon mutation is an apparent blue shift of the 675 nm band to 666 nm, in agreement with the assignment of a band at 673 nm to the low-energy exciton state formed predominantly by PD1 and PD2 (PD1PD2 exciton; see below) (20,21,33), which shifts to the blue upon His replacement. For the D2-His197 mutants, the amplitude of the 675 nm minimum is significantly smaller, suggesting that PD2 participates less than PD1 in the PD1PD2 exciton. The observed absorption loss at 683 nm is similar for both the D2-His197 mutants and is assigned to a secondary effect on ChlD1.

This secondary effect on ChlD1 has several possible origins: the loss of the hydrogen bond to the 131-keto carbonyl makes the ChlD1 π-conjugated system less asymmetric and/or the disappearance of the dispersive interactions provided by His (which also stabilizes the exciton states in which ChlD1 participates). Both effects are expected to induce a blue shift in the absorption spectrum. Alternatively, the mutation may increase the disorder in the system. In this case, a narrow localized state may become more delocalized to compensate for the higher disorder of the system, i.e., the narrow band broadens. The third explanation is more in line with the second derivative line shape of the Absdif spectra (Fig. 3). When we compare the effect of the Gln and Asn mutations in the active and inactive branches, it is interesting to note that the main absorption loss has its counterpart, with smaller amplitude, in the other branch mutant, highlighting the presence of excitonic coupling between PD1 and PD2.

**FIGURE 3** Simultaneously recorded 77 K absorption and Stark spectra of the *Synechocystis* PCC 6803 core complex in the QY absorption region for the WT, D1-His198Gln, and D1-His197Gln mutants (top left panel); WT, D1-His198Asn, and D2-His197Asn mutants (top-right panel); WT, D1-His198Ala, and D2-His197Ala mutants (bottom-left panel); and WT and D1-Thr179His mutant (bottom right panel). The absorption and Stark spectra are normalized to the cofactor content. The absorption difference spectra have been multiplied by 5 to facilitate visualization. The Stark spectra were recorded at \( \chi = 54.7^\circ \) and an external electric field strength of \( 2.25 \times 10^5 \) V cm\(^{-1}\).
For both of the D1 and D2 Ala mutants, the absorption difference spectra have similar spectral shapes. The minimum at 683 nm is similar in both mutants; the main difference is found in the 675 nm band, which is broader and has higher amplitude in the D1 mutant. In the Ala mutants, the site-directed mutation (Fig. 1) affects both sides in a similar way, with the D1 side being the most affected. These observations support the notion that Pd1 and Pd2 are energetically coupled, i.e., they contribute to the same exciton state: the low-energy state that is located around 675 nm (with higher participation from Pd1) (21,34). The high-energy component of this exciton state is located around 660 nm (34). It has a lower transition dipole moment strength and is therefore not observed in the Absdiff spectra.

The mutation constructed at position D1-Thr179 (Thr axially coordinates ChlD1 via a water molecule (5); Fig. 1) gives rise to a different absorption difference spectral shape with respect to the Pd1 and Pd2 mutants. In this case, an asymmetric minimum with contributions from 681, 677, and 671 nm components (in decreasing order of amplitude) and two maxima at 667 and 687 nm are observed. This spectral shape can be understood as a band broadening (the asymmetric minimum broadens to a band extending from 660 to 700 nm with ≈25 nm full width at half-maximum (FWHM)) or as a band shift (taking into account the amplitude of the maxima and minima, the 681 nm band red shifts to 687 nm, and the 677 and 671 nm bands blue shift to 666 nm). Recent experimental (21) and theoretical (33) studies on this mutant that showed the red shift of the ChlD1 exciton upon Thr exchange by His support the second interpretation. For this mutant, the red shift of the main 681 nm band and the blue shift of the minor 671 and 677 nm bands indicate that ChlD1 participates in two excitions: the ChlD1 exciton and, to a lesser extent, the Pd1Pd2 exciton. In summary, the results are consistent with the presence of two different low-energy exciton states in the ensemble of PSII RCs (11–13,20,21,35), the Pd1Pd2ChlD1 exciton (the low-energy exciton state that absorbs around 673 nm) and the ChlD1PheD1 exciton state (see “PheD1 mutant” section above, “The Phe QX absorption band” section in the Supporting Material, Fig. S3, and Table S1) with absorption around 681 nm, in agreement with the literature. (Note that the order of the cofactors in the exciton nomenclature, Pd1Pd2ChlD1, reflects a decreasing extent of participation in the exciton state.).

**Stark spectra**

The interpretation of the Stark spectra is based primarily on the comparison of the absorption and Stark spectra between WT and mutants (Fig. 3) in terms of band shifts and differences in band amplitudes, and secondarily on the comparison of the Stark spectra with the second derivative of the absorption spectra (2DerAbs; Fig. S4).

The Stark spectra of the PSII cores, WT, and mutants resemble the second derivative of the absorption spectra (2DerAbs; Fig. S4), indicating that to a first approximation, these Stark spectra are dominated by a change in dipole moment, Δμ, between the ground and excited states associated with the electronic transitions. A similar Stark effect was observed in the bacterial RC (26,27) and the higher plant PSII RC (17) (Fig. 2).

The Stark spectrum of the WT is composed of two main minima at 673.25 and 687.75 nm and three maxima located at 661, 677, and 686.25 nm. In the second derivative of the absorption spectrum (Fig. S4), three minima are present at 670, 677.5, and 682.5 nm, and two maxima are present at 661 and 687 nm. The Stark spectrum presents shifts of 1–3 nm (nonclassic Stark effect) and a threefold amplitude increase with respect to the 2DerAbs, which indicates that exciton couplings among the cofactors and exciton-CT mixing, respectively, are present in the WT core complex. For all mutants, band shifts in the Stark spectra with respect to the 2DerAbs are observed, indicating that in the mutants the exciton couplings are also present. However, for all mutants (with the exception of D2-His197Gln), the amplitudes of the minima of the Stark spectra are of the same order as those of the 2DerAbs spectra, indicating that the mixing with CT states is reduced or absent (Fig. S4). The most significant Stark band positions and amplitudes in the Qy region are listed in Table S2.

Thus far, the Stark spectra have confirmed the assignments from the absorption and the literature (20,21). At this point, we will examine the potential of the Pd1Pd2ChlD1 exciton state to initiate charge separation. This potential depends on the degree of mixing between the exciton and CT states: the stronger the mixing, the larger the Stark amplitude and the higher the potential to initiate charge separation (11). The changes in band position and amplitude for the Pd1 and Pd2 mutants show a correlation between the blue shift and the amplitude decrease of the 673.25 nm minimum: the stronger the shift, the larger the Stark amplitude reduction (Fig. S5). This correlation indicates that the absorption band shifts induced by the mutations decrease the exciton-CT mixing present in the WT, i.e., in the WT the (Pd1Pd2ChlD1)* exciton absorbing at 673.25 nm is mixed with the Pd2*/Pd1– CT state (due to the strong overlap between the electronic wavefunctions of these states) (11), yielding the (Pd2*/Pd1–ChlD1)* state. Thus, we conclude that the mixed (Pd2*/Pd1–ChlD1)* state initiates charge separation following the Pd1 path.
(PD2_D1_PheD1)* \rightarrow PD2_D1 ChlD1 \rightarrow PD1_D1 ChlD1 \rightarrow PD1_D1 ChlD1\dash PheD1 (12)) in a fraction of the RCs in which this mixed exciton-CT state is the lowest-energy state, and that the relative contribution of this path is strongly reduced in the PD1 and PD2 mutants.

Concerning the red minimum, at 681.75 nm in the WT, the mutants can be divided into two groups: 1), the PD1 and PD2 mutants with a blue shift (more significant for the PD2 mutations) and decrease in amplitude (similar for PD1 and PD2 mutants) depending on the specific mutation; and 2), the ChlD1 mutant with a red shift and a significant amplitude decrease. Considering the contribution of two exciton-CT states to the red Stark minimum observed in the RC samples, the blue shift present in the PD1 and PD2 mutants can be understood as a loss of a red component in the red Stark minimum. As discussed above, the PD1 and PD2 mutations induce a decrease in the (PD1PD2ChlD1)* ↔ PD2PD1 mixing that reduces the Stark amplitude of the 673.25 nm Stark minima (the (PD2_D1_PheD1)* mixed state has less CT character). In turn, the decrease of the mixing reduces the transition dipole moment of the red-most PD2_PD1 CT state. Therefore, the decrease of the (red) contribution of PD2_D1 ChlD1 CT state to the Stark spectra appears as a blue shift of the red Stark minimum. The fact that the decrease in Stark amplitude is the same in both PD1 and PD2 mutants can be explained by considering that the PD1 and PD2 cofactors contribute equally to the PD2PD1 CT state. In the case of the 673.25 nm minimum, the mutation effect is more pronounced for the PD1 mutants because the contribution of the PD1 cofactor to the (PD1PD2ChlD1)* exciton is larger than the contribution of PD2.

In the D1-Thr179His mutant, the absorption red shift of the exciton with participation of ChlD1 produces a substantial amplitude decrease (1/10 of the value for WT) and red shift (3.25 nm with respect to WT) of the red Stark minimum. These observations could be interpreted as a decrease of the ChlD1 exciton-CT mixing due to the red shift of the ChlD1 exciton absorption. However, the fact that two exciton-CT states contribute to the red Stark minimum and the comparison of the D1-Thr179His Stark spectra with the 2DerAbs indicate that the situation is more complex. This comparison shows that the 677.75 nm minimum in the 2DerAbs is almost completely absent in the Stark spectrum, and the 684 nm minimum in the 2DerAbs overlaps with the Stark minimum (Fig. S4). These observations indicate that 1), the red shift of the ChlD1 exciton absorption in the mutant disrupts the mixing with a CT state present in WT (as suggested by the absence of the 677.75 nm Stark minimum); and 2), the 684 nm minimum corresponds to the Stark contribution of the (PD2_D1)* state (in agreement with the interpretation of the Stark spectra presented in the “PheD1 mutant” section; Fig. 2). The fact that the 684 nm Stark minimum has the same amplitude as the 2DerAbs (and not a higher one, as would be expected for a state with significant CT character) suggests that the mutation in the environment of ChlD1 also affects the electronic properties of the (PD2_D1)* state.

Taken together, these results lead us to propose that the Stark minimum at 681.75 nm in WT arises from the combination of two different states. On the one hand, the (ChlD1_D1_PheD1)* state (produced by the mixing of the (ChlD1_PheD1)* exciton with the ChlD1_D1_PheD1 CT state) contributes to the blue edge of the 681.75 nm Stark minimum (∼681 nm; see also “The Phe QX absorption band” section in the Supporting Material, Fig. S3, and Table S1). On the other hand, the (PD2_D1)* state (formed by the mixing of the PD2_D1 CT state with the (PD1PD2ChlD1)* exciton) contributes to the red edge of the 681.75 nm Stark minimum (∼684 nm). Thus, we conclude that the (ChlD1_D1_PheD1)* state leads to charge separation via the ChlD1 path (12) whenever this exciton is the lowest-energy state of the RC. In addition, direct excitation of the (PD2_D1)* state leads to charge separation via the PD1 path (11).

In summary, we conclude that the electronic states that initiate charge separation are mixed exciton-CT states, not pure exciton states. This finding leads to the following refined charge separation pathways:

**P1 path**

\[
(PD2_D1_PheD1)* \rightarrow PD2_D1 \rightarrow PD2_D1 ChlD1 \rightarrow PD2_D1 PheD1 \rightarrow PD1_D1 PheD1
\]

**ChlD1 path**

\[
(ChlD1_D1_PheD1)* \rightarrow ChlD1_D1_PheD1 \rightarrow PD1_D1 PheD1
\]

**PD2 path**

\[
(PD2_D1)* \rightarrow PD2_D1 \rightarrow PD2_D1 ChlD1 \rightarrow PD2_D1 PheD1
\]

where the subscripts indicate the approximate center wavelength of the electronic transition for each of the mixed states.

This conclusion is in excellent agreement with the results from a high-resolution fluorescence spectroscopy (fluorescence line-narrowing) study by Peterman et al. (36). The authors studied the electronically excited state of PSII RC using fluorescence line-narrowing spectroscopy at 5 K, and compared the obtained spectral features with those obtained earlier for the primary electron donor (37). Their results show that there is a striking resemblance between the emitting and charge-separating states, including a very similar shape of the phonon wing, almost identical frequencies of a number of vibrational modes, a very similar double-Gaussian shape of the inhomogeneous distribution.
function, and relatively strong electron-phonon coupling for both states. All of these observations are explained by the fact that the excited states that give rise to charge separation in the PSII RC are mixed exciton-CT states; for instance, strong electron-phonon coupling is expected for states with CT character. It is also interesting to note that the best numerical simulation of the absorption, nonselectively, and selectively excited emission spectra was obtained with an inhomogeneous distribution function consisting of two Gaussians, one peaking at 680.6 nm (width 80 cm⁻¹, relative area 0.91) and the other peaking at 683.6 nm (width 70 cm⁻¹, FWHM) (36). These values are in excellent agreement with the properties and center wavelengths for the (ChlD₁[δ⁺PheD₁δ⁻])₆₈₁nm and (P₂D₂[δ⁺P₁δ⁻])₆₈₄nm states, respectively, found in our study. The state simulated by a Gaussian band centered at 680.6 nm corresponds to the mixed exciton-CT state (ChlD₁[δ⁺PheD₁δ⁻])₆₈₁nm, which carries most of the oscillator strength due to its major exciton character. The state simulated by a Gaussian band centered at 683.6 nm corresponds to the mixed CT-exciton (P₂D₂[δ⁺P₁δ⁻])₆₈₄nm state with lower oscillator strength due to its minor exciton character (acquired by borrowing oscillator strength from the (P₂D₂P₁)(ChlD₁)⁺ state).

CONCLUSIONS

The results demonstrate that the states that initiate charge separation are mixed exciton-CT or CT-exciton states, not pure exciton states as previously believed. The degree of mixing between exciton and CT states determines the quantum yield of RP formation and therefore the efficiency of charge separation (the higher the mixing, the higher the efficiency of charge separation). The exciton-CT states that initiate charge separation are (P₂D₂[δ⁺P₁δ⁻])₆₇₃nm and (ChlD₁[δ⁺PheD₁δ⁻])₆₈₁nm (where the subscript indicates the approximate center wavelength of the electronic transition for each mixed state). Moreover, the CT state (P₂D₂[δ⁺P₁δ⁻]) acquires excited state character due to its mixing with an exciton, producing the (P₂D₂[δ⁺P₁δ⁻])₆₈₄nm CT-exciton state, which is also able to initiate charge separation.

In addition, the results show that the pigment-protein interactions fine-tune the energy of the exciton and CT states, and hence the mixing between these states, which ultimately controls the selection and efficiency of a specific charge separation pathway.

SUPPORTING MATERIAL

Two tables, five figures, and references are available at http://www.biophysj.org/biophys/supplemental/S0006-3495(12)00683-2.

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