Alteration of the Axial Met Ligand to Electron Acceptor A₀ in Photosystem I: Effect on the Generation of P700⁺A₁⁻ Radical Pairs as Studied by W-band Transient EPR

Anton Savitsky · Oksana Gopta · Mahir Mamedov · John H. Golbeck · Alexander Tikhonov · Klaus Möbius · Alexey Semenov

Received: 12 June 2009 / Revised: 29 June 2009 / Published online: 17 November 2009 © The Author(s) 2009. This article is published with open access at Springerlink.com

Abstract Photosystem I (PS I) mutants from the cyanobacterium Synechocystis sp. PCC 6803 bearing point mutations to the axial ligands of A₀A (M688N_PsaA) and A₀B (M668N_PsaB) were studied by high-field W-band electron paramagnetic resonance (EPR) spectroscopy. It was found that the EPR observables of PS I from the M668N_PsaB mutant were virtual identical to that of the wild type (WT), and are clearly distinct from the M688N_PsaA mutant. In particular, the P700⁺ decay kinetics in the M688N_PsaA mutant is significantly slower than in the WT or the M668N_PsaB mutant. The analysis of the out-of-phase electron–electron dipolar electron spin echo envelope modulation shows that in the M668N_PsaB mutant, the estimated distance of 26.0 ± 0.3 Å agrees well with the 25.8 Å distance for the P700⁺A₁⁻ radical pair measured in the X-ray crystal structure. In the M688N_PsaA mutant, two populations are found with estimated distances of 26.0 ± 0.3 and 25.0 ± 0.3 Å in a ratio of 0.7–0.3, which agree well with the 25.8 Å distance for the P700⁺A₁⁻ radical pair in the X-ray crystal structure.

A. Savitsky · K. Möbius (✉)
Department of Physics, Free University of Berlin, Berlin, Germany
e-mail: moebius@physik.fu-berlin.de

J. H. Golbeck (✉)
Department of Biochemistry and Molecular Biology, The Pennsylvania State University, University Park, PA, USA
e-mail: jhg5@psu.edu

A. Tikhonov (✉)
Department of Biophysics, Faculty of Physics, Moscow State University, Moscow, Russia
e-mail: an_tikhonov@mail.ru
pair and the 24.6 Å distance for the \( P_{700}^+A_{1B}^- \) radical pair measured in the X-ray crystal structure. The data confirm that under the experimental conditions employed in this work, which involve dark-adapted samples without the pre-reduction of the iron–sulfur clusters, electron transport in cyanobacterial PS I is asymmetrical at 100 K, with the majority of electron transfer taking place through the A-branch of cofactors.

1 Introduction

Photosystem I (PS I) of cyanobacteria, algae, and higher plants is a membrane-bound chlorophyll (Chl)–protein complex that mediates light-induced electron transfer across the lipid bilayer from plastocyanin (Pc) to ferredoxin (Fd). PS I is a so-called type I photosynthetic reaction center (RC), which employs low-potential iron–sulfur clusters (\( F_A/F_B \)) as the terminal electron acceptors. The three-dimensional structure of PS I from the thermophilic cyanobacterium *Thermosynechococcus elongatus* has been solved by X-ray diffraction to a resolution of 2.5 Å [1]. The monomeric PS I complex (≈300 kDa) contains 12 different protein subunits, nine of which (PsaA, PsaB, PsaF, PsaI to PsaM and PsaX) are membrane-bound, and three of which (PsaC, PsaD and PsaE) are membrane-extrinsic. The membrane-embedded core of PS I is a heterodimer of the two largest subunits, PsaA and PsaB, which along with PsaF, PsaI to PsaM and PsaX, binds 96 molecules of Chl \( a \), 2 molecules of \( \beta \)-carotene, two molecules of phylloquinone (PhQ), and an interpolypeptide \( [4Fe–4S] \) cluster \( F_X \). The terminal \( [4Fe–4S] \) clusters, \( F_A/F_B \), are bound to the ≈9 kD peripheral stromal subunit, PsaC.

The arrangement of the electron transport cofactors in the PsaA, PsaB, and PsaC subunits is shown in Fig. 1. The primary electron donor, \( P_{700} \), is comprised of a molecule of Chl \( a \) and a molecule of Chl \( a' \) (Chl1A/Chl1B), the porphyrin planes being parallel to one another (interplanar distance, 3.6 Å) and perpendicular to the membrane plane. The X-ray diffraction analysis reveals additional domains of high electron density which are attributed to two Chl \( a \) molecules in positions corresponding to the accessory bacteriochlorophyll molecules in the RC of purple bacteria (Chl2A/Chl2B), two Chl \( a \) molecules in positions corresponding to the bacteriopheophytin molecules in the RC of purple bacteria [2–4] (Chl3A and Chl3B), and two PhQ molecules (Q\(_A\) and Q\(_B\)). Pairs of Chl and PhQ molecules are located in nearly \( C_2 \)-symmetric electron transport branches A and B associated with subunits PsaA and PsaB, respectively. Branch A incorporates Chl molecules Chl1A, Chl2A, Chl3A, and PhQ \( A_{1A} \), whereas branch B incorporates Chl molecules Chl1B, Chl2B, Chl3B, and PhQ \( A_{1B} \). The spectroscopically detected PS I cofactors include \( P_{700} \) (the special pair of molecules Chl1A and Chl1B), \( A_0 \) (one or two pairs of Chl molecules denoted as Chl2A/Chl3A and Chl2B/Chl3B), \( A_1 \) (one of the two molecules of phylloquinone (\( A_{1A}/A_{1B} \)), and \( F_X \), \( F_A \), and \( F_B \) (the three \([4Fe–4S]\) clusters) (see Fig. 1).

According to the currently accepted paradigm, the primary electron donor \( P_{700} \) is promoted to the excited singlet state \( P_{700}^* \) after absorbing a light quantum. This event is followed by the primary act of charge separation between \( P_{700}^* \) and the primary
acceptor $A_0$, forming the radical pair $P_{700}^+A_0^-$. An alternative scheme has been proposed in which the initial charge-separated state occurs between Chl2A and Chl3A and between Chl2B and Chl3B followed by migration of the hole from Chl2A and Chl2B to $P_{700}$ [5]. It should be noted, however, that this scheme has been recently challenged on the basis of electrostatic calculations of the midpoint potentials of the relevant cofactors [6]. At room temperature, the electron is transferred to the PhQ(s) $A_1$ within 50 ps and then to the iron–sulfur clusters $F_X$ and $F_A/F_B$ in $\leq$200 ns. In the absence of an exogenous electron acceptor, the electrons on $F_X^-$ and $[F_A/F_B]^{-\cdots}$ recombine with $P_{700}^+$ with lifetimes of 0.5–5 and 30–100 ms, respectively (for review, see Ref. [7]). The symmetric arrangement of the cofactors, as well as the convergence of the A- and B-branches at the $F_X$ cluster, suggest that both branches may be active in electron transfer. Interestingly, in the other main two types of photosynthetic RCs, namely bacterial RCs and photosystem II (PS II), electron transfer proceeds unidirectionally along the analogous A-branch. In these two RCs, the unidirectionality of electron transfer is functionally justified, because in contrast to PS I, the primary quinone $Q_A$ serves as a single electron acceptor, while the secondary quinone $Q_B$ serves as a gate to accept sequentially two electrons (and two protons).

The issue of whether electron transfer in PS I utilizes the A-branch of cofactors, the B-branch of cofactors, or both, has been a topic of lively debate for the last decade (reviewed in Ref. [8]). Two different approaches have been taken to investigate this issue. One involves altering amino acids, pigments, and cofactors in the vicinity of the $A_{1A}$ and $A_{1B}$ PhQs, and the other involves altering the H-bond and the axial ligand to the $A_{0A}$ and $A_{0B}$ Chls. The mutant PS I complexes generated near the $A_1$ and $A_0$ sites have been studied primarily using time-resolved optical

![Diagram of electron transfer cofactors in PS I from *Thermosynechococcus elongatus*](image)
spectroscopy and electron paramagnetic resonance (EPR) spectroscopy. The advantage of EPR is its ability to reveal spin density distributions of the transient states of the electron transfer cofactors, which reflect the electronic structure and the distances between the donor–acceptor radical pairs. The disadvantage is that to obtain information from dipolar EPR experiments [9], frozen solutions must be used at temperatures well below physiological conditions.

In *Synechocystis* sp. PCC 6803, site-symmetric mutations (Met → Leu) of the Chl3A/Chl3B (A₀) axial ligands affect almost exclusively the A-branch electron transfer when analyzed by femtosecond to picosecond time-resolved optical spectroscopy [10], by microsecond to millisecond optical spectroscopy and by X-band (9.5 GHz) time-resolved EPR (TREPR) spectroscopy at room and low temperatures [11], and by Q-band (35 GHz) TREPR spectroscopy at low temperatures [12]. The results showed that the M688L PsaA mutant exhibits: (i) slower growth rates, higher light sensitivity, and reduced amounts of PS I; (ii) a reduced yield of electron transfer from P700 to FA/FB at room temperature; (iii) an increased formation of the 3P700 excited triplet state due to P700²⁺A₀⁻⁻ recombination; (iv) a change in the intensity and shape of the electron polarization patterns of the consecutive radical pair states the consecutive radical pair states P700/C1²⁺A₁⁻⁻; and P700/C1²⁺FX⁻⁻ (v) a slowing of the A₀A to A₁A electron transfer step from 30 to 200 ps at room temperature. Except for an accelerated rate of charge recombination between P700²⁺ and [F₆/F₇]⁻⁻, PS I from the M688L PsaB mutant was similar to PS I from the wild-type (WT). Site-symmetric mutations (Met → Asn) of the Chl3A/Chl3B (A₀) axial ligands were also prepared and studied by femtosecond to picosecond time-resolved optical spectroscopy [10]. The lifetime of P700²⁺ in the M688N PsaA and M688N PsaB mutants was found to be similar to that in the M688L PsaA and M668L PsaB mutants.

To further characterize the M688N PsaA and M668N PsaB mutants, high-field W-band EPR experiments were carried out. By extending standard X-band (9.5 GHz) EPR to high-field/high-frequency techniques, significant advantages in terms of resolution and sensitivity are realized [9]. In particular, W-band EPR spectroscopy provides an excellent opportunity to distinguish and follow separately EPR signals from the ion-radicals P700²⁺ and A₁⁻⁻ (see Fig. 2). Moreover, electron–electron dipolar experiments, for instance, out-of-phase electron spin echo envelope modulation (ESEEM) on spin-correlated radical pairs [13] performed at high magnetic fields, allow one to significantly simplify the data analysis and to improve the interpretation of the results. Data analysis of standard X-band ESEEM spectroscopy must include a multiparameter fit of detected Fourier-transformed Pake-type spectra [14]. Such an analysis includes possible contributions to the electron-dipolar ESEEM spectra originating from nuclear modulations, for instance, from the ¹⁴N nuclei (nuclear Larmor frequency νₙ of 1.0 MHz at 9.5 GHz EPR) of P700²⁺, as well as artifacts due to incomplete excitation of the EPR spectra of both radicals in the pair. High-field EPR is virtually free of contributions from nuclear modulations (νₙ(¹⁴N) = 10 MHz at 95 GHz). Furthermore, the spectral resolution in combination with the structure of the radical pair allows one to selectively detect the perpendicular dipolar frequency, which can be analyzed by including two different dipolar contributions stemming from different radical pair geometries.
2 Materials and Methods

2.1 Cyanobacteria and Mutants

The M688NPsaA and M668NPsaB mutants were constructed in a *Synechocystis* sp. PCC 6803 preparation similar to that described for the M688LPsaA and M668LPsaB mutants [11]. The mutant cells were grown photomixotrophically with 5 mM glucose in liquid culture under conditions of light-activated heterotrophic growth [15]. Growth was monitored turbidometrically at 730 nm using a Cary 14 spectrophotometer modified for computerized data acquisition. The relevant sections of the *psaA* and *psaB* genes were resequenced after every large-scale growth to confirm that the Met residue had been replaced by an Asn residue. PS I trimers were isolated from thylakoid membranes using *n*-dodecyl-β-D-maltopyranoside (DM) and purified on two successive sucrose density gradients. The trimeric PS I complexes were present in the bottom green band and were resuspended in 50 mM Tris, pH 8.0, containing 0.03% DM and 15% glycerol at a Chl concentration of 3 mg/ml, and stored at −80°C until required.

2.2 W-band EPR Measurements

High-field EPR measurements were performed on a homebuilt W-band (95 GHz/3.4 T) multipurpose EPR spectrometer described previously [9, 16]. The spectrometer was equipped with a TE_{011} optical transmission cavity with unloaded quality factor $Q_U = 5000$ (empty). The sample solutions were placed in a quartz capillary (inner diameter, 0.6 mm) transferred to the EPR probe head and after dark adaptation at room temperature for 10 min cooled down to 100 K. Light-induced electron transfer was initiated by light pulses at 532 nm (Nd:YAG laser, 5 ns pulse width,
0.5 mJ on the sample surface) or by continuous illumination at 690 nm using continuous-wave (cw) diode laser (25 mW output, ca. 10 mW on the sample surface) guided to the center of the EPR cavity through a quartz fiber with diameter of 0.8 mm.

The heterodyne microwave bridge allowed us to perform cw EPR with field modulation and lock-in detection, TREPR without field modulation but direct detection, as well as pulsed EPR experiments. The recombination kinetics was obtained by recording the short-lived EPR absorption after laser flash (532 nm) via lock-in transient detection with magnetic field modulation (30 kHz, 0.1 mT modulation amplitude). The time resolution of this detection technique was thus set to about 1 ms. The pulsed EPR measurements were performed using the Hahn-echo sequence \((t_p)\tau(2t_p)\tau\)-echo under cw light illumination or allowing for a time after laser flash, in the sequence \(T_{DAF}, h\nu-\tau(2t_p)\tau-\tau(t_p)\)-echo under pulsed light illumination. The \(t_p\) pulse length of the \(\pi/2\) microwave (mw) pulses was generally set to 30 ns. The phase of the first mw pulse was cycled between \(+x, -x\). The quadrature-detected echo traces, \((s_{-y}, s_y)\), were digitized with 500 megasamples/s (2 ns per point) by a digital oscilloscope and transferred to the computer for further evaluation. To obtain the in-phase field-swept EPR spectra, the \(s_{-y}\) echo response traces at \(\tau = 150\) ns, corresponding to the first pulse phase settings \(+x, -x\), were subtracted from each other, and the pure echo response, i.e., free of free-induction decay (FID) and cavity ringing signals, was integrated over the whole echo duration. The out-of-phase EPR decay profiles were evaluated from \(s_x\) echo traces by integrating the echo responses over the time window, centered at the \(s_{-y}\) echo maximum that covers 60% of the echo intensity, to optimize resolution and signal-to-noise ratio. The echo-decay traces were acquired stepping the pulse separation time, \(\tau\), with 10 ns starting from 60 ns. The delay-after-flash time \(T_{DAF}\) was generally set to 400 ns.

2.3 Low-Temperature Optical Measurements

Transient absorbance changes in the near-infrared were measured in PS I complexes using a laboratory-built double-beam spectrophotometer and a low-temperature cryostat (Janis Research, Wilmington, MA). The excitation beam was provided by a frequency-doubled (\(\lambda = 532\) nm), \(Q\)-switched Nd:YAG laser (Spectra Physics). The pulse width at half-maximum was 8 ns. The cw measuring beam was provided by a semiconductor diode laser operating at 820 nm (Melles Griot). PS I trimers isolated from Synechocystis sp. PCC 6803 were suspended at 50 \(\mu\)g/ml Chl in 25 mM Tris–HCl buffer, pH 8.3, 10 mM Na ascorbate, 4 \(\mu\)M 2,6-dichlorophenolindophenol (DPIP), and 0.04% \(n\)-dodecyl-\(\beta\)-d-maltoside. The sample was frozen in the dark before measurement.

3 Results

3.1 Decay Kinetics of Flash-Induced EPR Signals from \(P_700^{\ast+}\) and \(A_1^{\ast-}\)

Figure 3 compares the decay kinetics of the field-modulated EPR signal from \(P_700^{\ast+}\) generated by repetitive laser flashes (1 Hz repetition rate) in PS I complexes from
the WT and the M688N\textsubscript{PsaA} and M668N\textsubscript{PsaB} mutants. After a fast flash-induced generation of P\textsubscript{700}\textsuperscript{+} a fraction of the light-induced signal decays reversibly at 100 K (the irreversibly formed P\textsubscript{700}\textsuperscript{+} is subtracted and hence not represented here). The kinetics of the post-illumination reduction of P\textsubscript{700}\textsuperscript{+} (charge recombination) could be deconvoluted into at least three components: a fast kinetic phase (\(\approx 2\) ms), a slow kinetic phase (\(\approx 200\) ms), and a very slow kinetic phase (\(\approx 3\) s). The half-times and statistical weights of these components, which were detected in PS I from the WT and the M688N\textsubscript{PsaA} and M668N\textsubscript{PsaB} mutants, were determined from the multiexponential fitting of the P\textsubscript{700}\textsuperscript{+} decay curve, and are listed in Table 1. PS I complexes from the B-branch mutant (M668N\textsubscript{PsaB}) show practically the same kinetics of P\textsubscript{700}\textsuperscript{+} decay as that observed in PS I complexes from the WT. PS I complexes from the M688N\textsubscript{PsaA} mutant are, however, characterized by a markedly slower reduction of P\textsubscript{700}\textsuperscript{+}. In contrast to the WT and M668N\textsubscript{PsaB} mutant, in which the fast component of P\textsubscript{700}\textsuperscript{+} decay dominates (\(\geq 50\text{--}60\%\)), the fast component comprises only 25\% of the total amplitude in the M688N\textsubscript{PsaA} mutant.

It should be added that the fast component of the post-illumination reduction of P\textsubscript{700}\textsuperscript{+} (\(<1\) ms), which can also be observed by optical and electrometrical methods [17, 18], is likely to be underestimated in the EPR experiment due to the intrinsic time limitation of the technique [9]. We point out, however, that the measured EPR kinetics of P\textsubscript{700}\textsuperscript{+} decay is in very good agreement with the optical measurements. To illustrate this point, let us consider Fig. 4, in which we compare the semi-logarithmic plots of the post-illumination P\textsubscript{700}\textsuperscript{+} decay kinetics measured at 100 K in WT PS I complexes by both optical and EPR methods. The optically measured kinetic trace reveals a variety of exponential components. Specifically, after fast initial steps of P\textsubscript{700}\textsuperscript{+} decay (\(\approx 500\) \(\mu\)s, statistical weight about 60\%), slower components of P\textsubscript{700}\textsuperscript{+} decay are present, which are practically identical with the kinetics measured by EPR.
Note that the EPR-detectable portion of the post-illumination kinetics of reversible P700/C1? reduction comprises about 40% of the maximal concentration of P700/C1? determined from optical measurements (Fig. 4). The coincidence of the EPR-detectable kinetics of P700/C1? re-reduction with the relevant portion of optically measured kinetics shows that both methods provide consistent results.

The fast component of P700/C1? decay measured by EPR ($\tau_{1/2}^{(1)} = 2.0 \pm 0.4$ ms in the WT and M668N_PsaB mutant, and $\tau_{1/2}^{(1)} = 12 \pm 2$ ms in the M688N_PsaA mutant) can be attributed without ambiguity to charge recombination of the radical pair P700+/A1-. In Fig. 5, we compare the EPR-measured decay kinetics for the cation-radical P700+ and the anion-radical A1- in PS I from the M668N_PsaB mutant. One can see that the dominant fast component of the A1- decay (≥90%) has a half-time 1.7 ± 0.3 ms, which overlaps with the characteristic time of the fast component of P700+ decay. In PS I from the M688N_PsaA mutant, the first component of P700+ decay, attributed to recombination of the radical pair P700+/A1-, is markedly slower ($\tau_{1/2}^{(1)} = 12 \pm 2$ ms) than that in PS I from the WT and the M668N_PsaB mutant. In contrast to the WT and M668N_PsaB samples, in which the fast EPR-detectable component dominates

### Table 1 Parameters of P700+/ decay kinetics obtained from the analysis of the EPR signal changes in response to laser flashes

| PS I complex        | $\tau_{1/2}^{(1)}$ (ms)/weight (%) | $\tau_{1/2}^{(2)}$ (ms)/weight (%) | $\tau_{1/2}^{(3)}$ (ms)/weight (%) |
|---------------------|-----------------------------------|-----------------------------------|-----------------------------------|
| WT                  | 2.2 ± 0.5 50                      | 140 ± 40  30                     | >3000 ± 20  20                   |
| WT (FX core)        | 2.0 ± 0.4 66                      | 200 ± 20 34                      | – –                                             |
| M668N_PsaB          | 2.0 ± 0.4 58                      | 240 ± 20 24                      | 5000 ± 1000 17                           |
| M688N_PsaA          | 12.5 ± 2.0 25                     | 280 ± 20 59                      | >3000 ± 20 15                           |

![Fig. 4](image)

Fig. 4 Comparison of the decay kinetics of post-illumination reduction of P700+ in WT PS I complexes as measured by optical (dashed line) and W-band EPR (solid line) methods at 100 K.
in the M688NPsaA sample the fast component comprises only 25% of the total amplitude (note that due to the insufficient time response of the lock-in detection EPR technique, the amplitude and lifetime of the fast component may be underestimated in the EPR measurement of the WT and M668NPsaB mutant, see Fig. 4).

The slow EPR-detectable component of P700/C1? decay ($\tau_{1/2}^{(2)} \approx 140$ ms, statistical weight $\approx 30\%$) can be attributed tentatively to recombination of the radical pair P700/C1? -FX -$. This component was absent in the kinetics of the A1/C1 decay (Fig. 5), although a similar component of the P700/C1? decay ($\tau_{1/2}^{(2)} \approx 200$ ms, $\approx 34\%$) was observed in the P700–FX core preparation isolated from WT cells (data not shown, see Table 1 for kinetic parameters). The very slow component ($\tau_{1/2}^{(3)} \geq 3$ s; $\approx 20\%$ in the WT and $\approx 15–17\%$ in the M688NPsaA and M668NPsaB mutants) remains unassigned. Because the very slow component is absent in the P700–FX core preparation, it is tempting to assign it to a previously unrecognized albeit small amount of recombination from P700/FX -[FA/FB] $. However, this assignment requires confirmation by additional spectroscopic methods.

3.2 Comparison of EPR Spectra of PS I from WT, M688NPsaA and M668NPsaB Mutants

In this section, we compare field-swept pulse EPR spectra of dark-adapted PS I complexes illuminated with continuous laser light (690 nm). As shown in Fig. 6, there are no visible EPR signals attributed to P700/C1? in dark-adapted preparations of PS I from the WT and the M668NPsaB mutant. In similar PS I preparations from the M688NPsaA mutant, however, a small signal from P700/C1? is present. Illumination of
dark-frozen samples with saturating, continuous light induced a partly reversible oxidation of P\textsubscript{700}. After switching off the light, the intensities of the EPR signals from P\textsubscript{700}\textsuperscript{\textbullet} decreased, but did not reach the initial levels that were present prior to illumination. Thus, in each case, a significant portion of the electrons transferred at 100 K cannot recombine with P\textsubscript{700}\textsuperscript{\textbullet}, probably due to trapping by the terminal electron acceptors, e.g., F\textsubscript{A} and F\textsubscript{B} [18]. Note that PS I from WT cells shows a higher reversibility of the EPR signal from P\textsubscript{700}\textsuperscript{\textbullet} characterized by the ratio $D/P_0 = 0.70$ (see Fig. 6 for definitions) than the two mutants. For PS I complexes from the M688N\textsubscript{PsaA} and M668N\textsubscript{PsaB} mutants, reversible changes in the amplitude of the P\textsubscript{700}\textsuperscript{\textbullet} EPR signal were lower, $D/P_0 = 0.44$ and 0.57, respectively.

In addition to the signal from P\textsubscript{700}\textsuperscript{\textbullet} there are changes in the EPR spectrum due to the reduction of at least one other paramagnetic species. Figure 6 shows that the illumination of PS I with continuous red light at 100 K gives rise to signals in the low-field region that could belong to a semiquinone species. Analysis of the difference spectra (‘light on’ minus ‘light off’) allows one to extract information on possible sources of the reversible EPR signal in the low-field region. Figure 7

![Figure 6](https://example.com/fig6.png)
shows difference spectra normalized either to the amplitude of the signal from P$_{700}^+$ (a) or the low-field difference signal (b). As can be seen in Fig. 7b, the EPR spectra of the light-induced semiquinone-like radicals have the same shape, which are shape typical of the anion-radical A$_1^-$. As we will see below, the small amount of the A$_1^-$ signal in the M688N$_{PsaA}$ mutant correlates with the results of the measurement of the transient radical pair P$_{700}^+$A$_1^-$ generated by repeated laser flashes.

3.3 Out-of-phase ESEEM Study of PS I from the M688N$_{PsaA}$ and M668N$_{PsaB}$ Mutants

The out-of-phase ESEEM signal of the P$_{700}^+$A$_1^-$ radical pair provides information on the magnitude of the electron–electron dipolar coupling between the radicals, thus allowing the distance between P$_{700}^+$ and A$_1^-$, $r_{PA}$, to be determined (for recent reviews, see Refs. [9, 19, 20]). Figure 8a compares the out-of-phase ESEEM signals of the detected 2-pulse echo-decay in PS I from the M688N$_{PsaA}$ mutant (solid line) and M668N$_{PsaB}$ (dashed line) mutant recorded at the maximum of the EPR spectrum of P$_{700}^+$. It can be seen that in the M668N$_{PsaB}$ sample, the oscillations of the signal occur with a lower frequency and are less damped as compared to the M688N$_{PsaA}$ sample. This difference reflects structural differences between the radical pairs generated in the PS I complexes from the two mutants.

At W-band the modulation frequency observed in the out-of-phase detected echo-decay corresponds to the perpendicular dipolar coupling frequency, $v_\perp$, which value depends on the distance $r_{PA}$ as

$$v_\perp = 5.204 \times 10^{10} \times r_{PA}^{-3},$$

(1)

where $v_\perp$ is measured in s$^{-1}$ and $r_{PA}$ in units of Å. Here, we consider only the perpendicular dipolar coupling frequency $v_\perp$ and not the parallel component $v_\parallel$. This is because the dipolar vectors directed from P$_{700}^+$ to the semiquinones A$_{1A}^-$ and A$_{1B}^-$ are oriented almost perpendicularly to the z-axes of both quinones [21].

For a detailed analysis of the dipolar coupling, we turn to the frequency domain by performing a Fourier transformation (FT) of the zero-time reconstructed sine decay traces shown in Fig. 8a. Figure 8b compares the sine fast Fourier-transform (sin-FFT) amplitudes for PS I from the M688N$_{PsaA}$ and M668N$_{PsaB}$ mutants. There is a notable difference between samples; in particular, in the M688N$_{PsaA}$ mutant, the line at 3 MHz reveals a shoulder shifted towards higher frequencies. This could indicate that in the M688N$_{PsaA}$ mutant the radical pair P$_{700}^+$A$_{1B}^-$ makes a contribution to the overall signal. Further analysis of these data allowed us to evaluate the extent of this contribution.

Figure 9 shows the high-frequency regions of the sin-FFT amplitude spectra for the M668N$_{PsaB}$ mutant and the M688N$_{PsaA}$ mutant in the presence of dithionite, which was added to the sample to increase the signal amplitude (see Sect. 3.4). For the M668N$_{PsaB}$ sample (Fig. 9a), the line was fitted by a Lorentzian with a peak frequency $v_0 = 2.95$ MHz and a line width $\Delta v = 0.4$ MHz estimated as the half-width at half-maximum (HWHM). Figure 9b shows the deconvolution of the sin-FFT amplitude spectrum of the M688N$_{PsaA}$ sample into the sum of two Lorentzian
curves: one with spectral parameters of the M668NPsaB sample ($m_0 = 2.95 \text{ MHz}$ and $D_m = 0.4 \text{ MHz}$), and the other shifted to a higher frequency by 0.4 MHz ($m_0 = 3.35 \text{ MHz}$, $D_m = 0.4 \text{ MHz}$). The statistical weights of these components are 0.7 and 0.3, respectively. According to Eq. (1), spectral lines with frequencies $m_0 = 2.95 \text{ MHz}$ and $m_0 = 3.35 \text{ MHz}$ correspond to distances $r_{PA} = 26.0 \pm 0.3$ and $r_{PA} = 25.0 \pm 0.3$ Å, respectively, which are in a good agreement with the X-ray structural data (Fig. 10). Taking into account that the spin density distribution in the cation-radical $P_{700}$ is shifted towards the B-side Chl of the $P_{700}$ dimer [22, 23], and assuming that spin density in the anion-radicals $A_{1A}$ and $A_{1B}$ is located in the center of their quinone rings, the X-ray data suggest the following center-to-center distances: $r_{P_{700}A_{1A}} = 25.8 \text{ Å}$ and $r_{P_{700}A_{1B}} = 24.6 \text{ Å}$. These values are in good agreement with the experimental data (Fig. 10).
agreement with our estimates of the distances, 26.0 ± 0.3 and 25.0 ± 0.3 Å, by Eq. (1).

3.4 EPR Spectra of the Radical Pair \( P_{700}^+ A_{1}^- \)

Next we compare the transient spin-correlated radical pair (SCRP) signals of \( P_{700}^+ A_{1}^- \) generated by laser pulses (532 nm, 2.5 Hz repetition rate) in PS I from the WT and the M688N\textsubscript{PsaA} and M668N\textsubscript{PsaB} mutants. Figure 11 presents the SCRP signals detected 400 ns after the laser flash at 100 K. For the sake of comparison, all the spectra shown in Fig. 11 are normalized to the amplitude of the low-field component of each spectrum, which belongs to semiquinone anion radicals. As one can see in Fig. 11, the PS I complexes from the WT and the M668NPsaB mutant give rather intense signals, for which the line shapes are typical of the radical pair \( P_{700}^+ A_{1}^- \) \([9]\). In the case of PS I from the M688N PsaA mutant, a much smaller SCRP signal was observed. In the presence of dithionite, which reduces \( F_A \) and \( F_B \), the intensity of the signal increases by a factor of about three (not shown). This enhancement effect is probably due to an increased contribution of a B-side electron cycle as a result of the reduction of \( F_A/F_B \).

It should be noted that due to the large number of factors involved, the intensity of the SCRP signal from \( P_{700}^+ A_{1}^- \) cannot serve as a simple and unambiguous quantitative measure of the electron flux through any given branch. Nevertheless, the relatively low signal intensity in the M688N\textsubscript{PsaA} mutant, even in the presence of dithionite, is best explained either by a block in electron transfer beyond \( A_0 \) in the reversible (at 100 K) A-branch of electron transfer cofactors, or by a slowing down of the recombination of the radical pairs \( P_{700}^+ A_{1}^- \) and \( P_{700}^+ F_X^- \). In the latter case, a number of active reaction centers in the state \( P_{700} A_1 \), i.e., those PS I complexes that
would be in the ground state and, hence, ready for repetitive flash-induced generation of $P_700^*/C1^-$, would diminish.

The line shape of the low-field signal from $A_1^-/C1^-$ is the same in PS I from the WT and the M688NPsaA and M688NPsaB mutants, whereas the signal in the high-field region, which is dominated by $P_700^*/C1^-$, differs markedly. This situation is somewhat similar to that described by Poluektov and collaborators [21], who observed at 100 K two kinds of high-frequency/high-field EPR spectra of the radical pairs $P_700^*/A_1^-$ in PS I complexes from predeuterated cells of the cyanobacterium *Synechococcus lividus*.

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**Fig. 10** Mutual arrangement of the primary electron donor $P_{700}$ (the Chl1A and Chl1B special pair) and the phyloquinones $A_{1A}$ and $A_{1B}$ based on the X-ray crystal structure data [1] (PDB entry 1JBO). The distances between $P_{700}$ and $A_{1A}$ (25.8 Å) in the A-branch, and between $P_{700}$ and $A_{1B}$ (24.6 Å) in the B-branch, correspond to distances between the spin density center in $P_{700}$ (see Refs. [22] and [23]) and the geometrical centers of the benzoquinone fragments of $A_{1A}$ and $A_{1B}$.

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**Fig. 11** Field-swept 2-pulse W-band EPR spectra of the light-induced spin-correlated radical pairs $P_{700}^*/A_1^-$ recorded 400 ns after the laser flash at 532 nm in the PS I complexes from the WT cells, and the M688NPsaA and M688NPsaB mutants at 100 K. The spectra are normalized to the $A_1^-$ EPR signal.
One spectrum can be ascribed to the SCRP signal $P_{700}^+A_{1A}^-$ from the A-branch, while the other, observed under strongly reducing conditions, can be ascribed to the SCRP signal $P_{700}^+A_{1B}^-$ from the B-branch. We note, however, that the difference between the SCRP spectra, which we observed for PS I complexes from the WT and the M668N$_{PsaB}$ mutant and from PS I complexes from the M688N$_{PsaA}$ mutant, is not as dramatic as that reported in Ref. [21].

4 Discussion

The aim of the present study is to characterize PS I mutants, bearing point mutations to the axial ligands of $A_{0A}$ and $A_{0B}$, with the use of a variety of the W-band EPR techniques. They include cw EPR, field-swept ESE, TREPR, and out-of-phase ESEEM of the spin-correlated radical pair $P_{700}^+A_{1A}^-$. The use of W-band EPR afforded an opportunity to measure the kinetics of the $A_{1A}^-$ radical separately from the $P_{700}^+$ radical. The results are summarized below:

1. The flash-induced kinetics of $P_{700}^+$ decay coincides in PS I from the WT and M668N$_{PsaB}$ mutant, and differs significantly from the M688N$_{PsaA}$ mutant as do the relative amounts of the fast, slow and very slow components. The $P_{700}^+$ decay kinetics in the case of the M688N$_{PsaA}$ mutant is significantly slower than in the WT or the M668N$_{PsaB}$ mutant.

2. Under continuous illumination, there is a significant difference in the amount of reversibly formed semiquinone $A_{1B}^-$ between the WT, M688N$_{PsaA}$ and M668N$_{PsaB}$ samples. The amount of $A_{1B}^-$ was the largest in the WT, lower in the M668N$_{PsaB}$ mutant, and negligible in the M688N$_{PsaA}$ mutant.

3. The out-of-phase ESEEM study reveals a lower frequency of the spin-echo envelope modulations in the M668N$_{PsaB}$ mutant as compared to the M688N$_{PsaA}$ mutant. The analysis of these data allowed the distances between the $P_{700}^+A_{1A}^-$ radical pairs in the M688N$_{PsaA}$ and M668N$_{PsaB}$ mutants to be determined as 25.0 $\pm$ 0.3 and 26.0 $\pm$ 0.3 $\AA$, respectively. These distances are close to the distances estimated on the basis of the X-ray structure and spin density distribution in the $P_{700}$ dimer (25.8 $\AA$ for the $P_{700}^+A_{1A}^-$ radical pair and 24.6 $\AA$ for the $P_{700}^+A_{1B}^-$ radical pair).

4. The spin-correlated $P_{700}^+A_{1B}^-$ radical pair spectra were nearly identical for the WT and M668N$_{PsaB}$ samples, whereas for the M688N$_{PsaA}$ sample in the high-field region the shape of the spectrum was clearly different. In addition, the intensity of the signal in case of PS I from the M688N$_{PsaA}$ mutant was significantly lower than that of PS I from the WT and the M688N$_{PsaB}$ mutant.

The question is why the A- and B-side behaviors are so different in the two otherwise symmetrical mutants. An attempt of modeling the electron transfer rates suggests nearly equivalent midpoint potentials for $A_{1A}$ ($-671$ mV), $A_{1B}$ ($-696$ mV), and $F_X$ ($-680$ mV) [24]. In this model, electron transfer from $A_{1A}$ to $F_X$ would be endothermic by 9 mV and electron transfer from $A_{1B}$ to $F_X$ would be exothermic by 16 mV. The thermodynamically unfavorable electron transfer from $A_{1A}$ to $F_X$ at low temperature could explain the observation of the
TREPR-detected $P_{700}^{+}A_{1A}^{-}$ radical pair on repeated laser flashes. It should be noted that the absolute values of the midpoint potentials in this study are not supported by electrostatic calculations, in which the redox potentials for $A_{1A}$ and $A_{1B}$ were calculated to be $-531$ and $-686$ mV, respectively [25]. However, Ishikita and Knapp [25] did not take into account the difference between the reference redox potential of menaquinone in water and dimethylformamide, and therefore have overestimated the absolute values of midpoint potentials for $A_{1A}$ and $A_{1B}$. This error does not affect the estimation of a 155 mV energy difference between $A_{1A}$ and $A_{1B}$ because the error in the liquid junction potential is the same for $A_{1A}$ and $A_{1B}$. More recent electrostatic calculations of redox potentials, carried out within the framework of a semi-continuum dielectric approach, show that the midpoint potential of $A_{1A}$ is 170 mV more positive than that of $A_{1B}$, and an energy gap between $A_{1A}$ and $F_X$ of $\sim 80$ mV [6].

The decrease in the distance typical for the $P_{700}^{+}A_{1A}^{-}$ pair to the distance typical for the $P_{700}^{+}A_{1B}^{-}$ pair, as observed by Poluektov and coworkers [21] and by Santabarbara and coworkers [26, 27], may be explained by a block in electron transfer through the A-branch. This is probably a result of reduction of $A_{1A}$ with dithionite and illumination (note that the more reducing $A_{1B}$ cofactor remains oxidized). Similarly, the observation of a change in the high-field region along with the appearance of a higher frequency shoulder in the dipolar spectrum (this work) may be explained by a partial block in electron transfer through the B-branch. It remains to be explained how the otherwise fast electron transfer through the B-side cofactors can be observed within the limited EPR time window [20] even at 100 K, given that there is little or no activation energy associated with this electron transfer step [28]. Perhaps, because Asn can potentially form an H-bond to the second C=O of the quinone, the position of the A-branch quinone could have moved in some of the PS I complexes to a position closer to $P_{700}^{+}$. The spectra, however, argue against this possibility because the more equal H-bonding scheme would be expected to reduce the hyperfine coupling from the $-\mathrm{CH}_3$ fragment.

It is clear that in *Synechocystis* sp. PCC 6803, the measured parameters of PS I from the M668N$_{PsaB}$ sample are virtually identical to those of the WT, and are clearly distinct from those that of the M668N$_{PsaA}$ sample. Previously, mutant PS I complexes, in which M668 on the A-branch and M668 on the B-branch were replaced by Leu residues, were studied by time-resolved optical absorption spectroscopy at room temperature [10], by X-band TREPR at room and at low temperatures [11], and by Q-band TREPR at low temperatures [12]. The measured parameters of PS I from the M668L$_{PsaB}$ sample were almost identical to those that of the WT, and were clearly distinct from the M688L$_{PsaA}$ sample. In particular, the quantum yield of electron transfer was lower, and the rate of forward electron transfer from $A_{0A}$ to $A_{1A}$ was slowed by a factor of 3–10 in the M688L$_{PsaA}$ mutant [10]. Thus, these earlier results are consistent with the present study.

In contrast to *Synechocystis* sp. PCC 6803, the site-symmetric mutants M688H$_{PsaA}$ and M668H$_{PsaB}$ in *C. reinhardtii* showed similar effects when the decay of the electron spin-polarized signal from the $P_{700}^{+}A_{1A}^{-}$ radical pair was measured by low-temperature EPR. In particular, the M688H$_{PsaA}$ mutant showed no
evidence for electron transfer to A\textsubscript{1A} at low temperature, and the M668HPsaB mutant showed no evidence for electron transfer to A\textsubscript{1B} at low temperature [29]. In a point dipole approximation, the P\textsubscript{700}\textsuperscript{+}-to-A\textsubscript{1B} distance of the M688HPsaA mutant was estimated to be 24.3 Å, and the P\textsubscript{700}\textsuperscript{+}-to-A\textsubscript{1A} distance in the M668HPsaB mutant was estimated to be 25.4 Å. The difference in the two distances is in agreement with the difference in the P\textsubscript{700}-to-A\textsubscript{1A} and P\textsubscript{700}-to-A\textsubscript{1B} distances according to the X-ray diffraction data. An intermediate value of 25.0 Å was obtained in the WT membranes, which was offered as evidence for bidirectional electron transfer [26]. Ultrafast transient absorbance measurements showed that the A\textsubscript{0}\textsuperscript{−} minus A\textsubscript{0} difference signal accumulates in both the M688HPsaA and M668HPsaB mutants, indicating that electron transfer from A\textsubscript{0}\textsuperscript{−} to A\textsubscript{1} is blocked or slowed on the appropriate branch [30]. However, a recent spin-polarized EPR study shows a transient EPR spectrum, probably due to P\textsubscript{700}A\textsubscript{1A}\textsuperscript{−}, in the M688HPsaA variant along with no evidence for slowing of electron transfer to F\textsubscript{X} in the M668HPsaB variant [31, 32]. Indeed, a more recent time-resolved optical study of the M688HPsaA or M668HPsaA variants showed unchanged lifetimes of the fast and slow kinetic phases but an alteration in the ratio of fast to slow kinetic phases [33]. Thus, the situation in C. reinhardtii is more complex than originally thought. Further resolution of the differences and discrepancies between and among the C. reinhardtii and Synechocystis sp. PCC 6803 mutants will require additional study, possibly comparative studies of the mutants under similar conditions using the same methodologies. A paper by van der Est and colleagues [34] in this special issue further describes the low-temperature behaviour of the M688NPsaA and M668NPsaB mutants by multifrequency time-resolved and cw EPR.

Acknowledgments  We are grateful to Art van der Est for fruitful discussions and for a critical reading of the manuscript. This work was supported by the Deutschen Forschungsgemeinschaft (DFG) (MO 132/19-2), the DFG-Russian Foundation for Basic Research (Cooperation Project NNIO_a 08-04-91951), the Russian Foundation for Basic Research (Grants 09-04-01657-a, 09-04-00978-a and 07-04-01050), the Federal Agency for Science and Innovation (Grants 02.512.12.2005 and 02.512.11.2286), and the US National Science Foundation (MCB-0519743).

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