Formation of tyrosine radicals in photosystem II under far-red illumination

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Abstract Photosystem II (PS II) contains two redox-active tyrosine residues on the donor side at symmetrical positions to the primary donor, P_{680}. Tyr\textsubscript{Z}, part of the water-oxidizing complex, is a preferential fast electron donor while Tyr\textsubscript{D} is a slow auxiliary donor to P_{680}. We used PS II membranes from spinach which were depleted of the water oxidation complex (Mn-depleted PS II) to study electron donation from both tyrosines by time-resolved EPR spectroscopy under visible and far-red continuous light and laser flash illumination. Our results show that under both illumination regimes, oxidation of Tyr\textsubscript{D} occurs via equilibrium with Tyr\textsubscript{Z}\textsuperscript{•} at pH 4.7 and 6.3. At pH 8.5 direct Tyr\textsubscript{D} oxidation by P_{680}\textsuperscript{+} occurs in the majority of the PS II centers. Under continuous far-red light illumination these reactions were less effective but still possible. Different photochemical steps were considered to explain the far-red light-induced electron donation from tyrosines and localization of the primary electron hole (P_{680}\textsuperscript{+}) on the Chl\textsubscript{D1} in Mn-depleted PS II after the far-red light-induced charge separation at room temperature is suggested.

Keywords Photosystem II · Tyrosine Z and D · Electron transfer · Far-red light

Abbreviations

Chl Chlorophyll
Car Carotenoid
DPC Diphenylcarbazide
EPR Electron paramagnetic resonance
Ferri Potassium ferricyanide K\textsubscript{3}[Fe(CN)\textsubscript{6}]
PS II Photosystem II
Pheo Pheophytin
QA and Q\textsubscript{B} Primary and secondary plastoquinone acceptor in PSII
Tyr\textsubscript{Z} Tyrosine 161 on the D1 protein
Tris Tris(hydroxymethyl)aminomethane
Tyr\textsubscript{D} Tyrosine 160 on the D2 protein

Introduction

Solar energy is successfully utilized by plants, algae, and bacteria in the process called photosynthesis. In oxygenic photosynthesis, solar energy is converted to chemical energy in the form of carbohydrates and O\textsubscript{2} is released as a byproduct (Kern and Renger 2007; Renger and Renger 2008; Vinyard et al. 2013). The initial reaction of photosynthesis takes place in photosystem II (PS II), a multicomponent Chl protein complex embedded in the thylakoid membrane of chloroplasts and cyanobacteria. The active PS II complex is made from 25 protein subunits and host a chain of the redox-active cofactors involved in the key water oxidation reaction and subsequent electron transfer (Umena et al. 2011; Wei et al. 2016). These cofactors are bound by the PS II central core which is composed of the D\textsubscript{1} and D\textsubscript{2} proteins, the inner pigment–protein antenna complexes CP43 and CP47, Cyt b_{559}, and several low molecular weight essential subunits (Danielsson et al. 2006; Umena et al. 2011; Suga et al. 2015). On the luminal side, water-oxidizing complex of...
plants and algae is shielded by three extrinsic proteins PsbO, PsbP, and PsbQ (Bricker et al. 2012).

The sequence of electron transfer reactions leading to the oxidation of water occurs in the following order. After light absorption by antenna, P_680 is excited and rapidly loses an electron to the nearby primary electron acceptor Pheo. Reduced Pheo\(^-\) passes an electron to the bound plastoquinone Q_\(A_1\), forming Q_\(A_1\)^\(-\) which in turn transfers an electron to Q_\(B\) (Renger and Renger 2008). All these cofactors are single electron carriers while exchangeable plastoquinone Q_\(B\) can accept two electrons and then become double protonated upon reduction. Reduced Q_\(B\)H_2 diffuses from the Q_\(B\)-pocket and is replaced by another plastoquinol from the membrane PQ pool (Renger and Renger 2008; Barber 2016). On the donor side of PS II, the water-oxidizing complex is composed of Mn_4CaO_5 cluster and redox-active tyrosine D1-161 (TyrZ), and mostly bound by the D1 protein. P_680\(^+\) is a strong oxidant with redox potential of 1.25 V, high enough to drive water oxidation reaction via TyrZ (Grabolle and Dau 2005; Cardona et al. 2012). Water oxidation occurs at the Mn_4CaO_5 cluster which goes through S-cycle to oxidize water to a molecular O_2 and four protons by transferring four electrons to P_680\(^+\) via TyrZ (Rappaport et al. 2002; Renger and Renger 2008; Vinyard et al. 2013).

In intact oxygen-evolving PS II Tyr\(Z^*\) oxidation has half-time in nsec–µsec range (Brettel et al. 1984; Renger 2012). In the absence of Mn_4CaO_5 the half-time became by 2–3 orders of magnitude slower (Babcok and Sauer 1975a; Brettel et al. 1984). PS II contains another redox-active tyrosine D2-160 (TyrD), which is located symmetrically to TyrZ on the D2 protein (Styring et al. 2012). Contrary to Tyr\(Z^*\), Tyr\(D^*\) is very stable and stays oxidized in the dark for minutes to hours (Babcok and Sauer 1973; Styring and Rutherford 1987; Vass and Styring 1991). Due to the slow oxidation under physiological pH, TyrD is not competitive to TyrZ as an electron donor to P680\(^+\). However, at elevated pH with a pK_a \sim 7.6, TyrD becomes a very efficient donor, with half-times comparable to those seen for Tyr\(Z\), \(t_{1/2} = 190\) ns (Faller et al. 2001, 2002). The difference in the environment of two tyrosines is the reason for their difference in oxidation kinetics (Umena et al. 2011). TyrD is in relative hydrophilic environment, deeply buried in the protein interior. On the contrary, TyrZ is in more hydrophilic surrounding with a cluster of water molecules nearby (Ferreira et al. 2004; Umena et al. 2011; Suga et al. 2015). Interestingly, TyrD has only a single water molecule nearby which can take two positions (2.6–3.1 and 4.3–4.5 Å) (Umena et al. 2011; Saito et al. 2013; Suga et al. 2015; Sjöholm et al. 2016; Ahmadova et al. 2017).

The primary donor in PS II, P_680 consists of four Chl molecules bound by the D1/D2 heterodimer denoted as P_D1, P_D2, Chl_D1, and Chl_D2, (see Scheme 1). The distance from reaction center Chl P_D1 and P_D2 to the Tyr\(Z\) and Tyr\(D\), respectively is 9.1–9.3 Å. Both tyrosines can be oxidized by P_680 entity after light-triggered primary charge separation formed P_680\(^+\) + Pheo\(^-\) pair. It is still under debate which Chl in P_680 entity forms the primary donor (Rappaport and Diner 2008). Interestingly, apart from visible-light-driven charge separation, the far-red light-driven charge separation up to 800 nm was reported in PS II (Thapper et al. 2009). Taking into account the lower energy of the far-red photons an alternative primary charge separation event was proposed (Thapper et al. 2009). It was also shown that at low temperature the primary charge pair formed under the far-red light illumination is ChlD1\(^+\) Pheo\(^-\) (Hughes et al. 2006; Romero et al. 2012; Mokvist et al. 2014; Novoderezhkin et al. 2016). This conclusion was made based on the different yields of the electron donor pathways in PS II at 5 K (Mokvist et al. 2014). The situation at physiological temperatures is unclear (Reimers et al. 2016).

The slow oxidation behavior of TyrD makes it possible to study by the conventional EPR spectroscopy while TyrZ oxidation kinetics is too fast to be followed. Different biochemical treatment such as Tris and NH_2OH washing can remove the Mn_4CaO_5 cluster and extrinsic proteins, leaving PS II core exposed to the lumen (Boussac and Etienne 1982b; Gadjieva et al. 1999; Mamedov et al. 2007). This makes TyrZ oxidation to slow down by two or three orders of magnitude \(t_{1/2} = 20–600\) ms (Buser et al. 1990). In this case both tyrosine donors can be accessed by EPR spectroscopy. Symmetrically situated at the different sides of P_680, they constitute a useful, simplified system to study the far-red-induced photochemistry at room temperature (Scheme 1).

In the present work we investigate the primary charge separation through oxidation of two tyrosines under two different excitation wavelengths. We have studied tyrosine
oxidation kinetics in the Mn-depleted PS II membranes at different pH values. The difference in the oxidation efficiency of TyrZ and TyrD allowed us to suggest localization of the primary electron donor Chl in P680 after far-red illumination at physiological temperature.

Materials and methods

Sample preparation

PS II-enriched membranes (BBY type) were isolated from hydroponically grown spinach (Spinacia oleracea) by the method of Berthold et al. 1981 with some modifications according to Völker et al. (1985). The samples were resuspended in a 25 mM MES buffer, pH 6.1, 400 mM sucrose, 15 mM NaCl, and 3 mM MgCl2 at a Chl concentration of 5–6 mg/mL and stored at −80 °C until use. All sample preparations were performed in darkness or under the dim green light.

The Mn4CaO5 cluster and extrinsic subunits were removed by the Tris washing (Gadjieva et al. 1999, Mamedov et al. 2007). PS II membranes were resuspended in 1.0 M Tris buffer at pH 9.1 with Chl concentration 1 mg/mL. They were stirred at 4 °C for 30 min under room light. After centrifugation, the pellet was washed twice with a low molar buffer containing 2 mM MES–NaOH pH 6.1, 300 mM sucrose, 10 mM NaCl, 3 mM MgCl2, and stored at −80 °C. This treatment washed away >90% of bound Mn and all three extrinsic proteins (Gadjieva et al. 1999). TyrD reduction in Tris-washed PS II samples was obtained by incubation samples at a concentration of 4–5 mg of Chl/mL in the dark for 10 h at room temperature (21 °C). The incubation treatment reduced about 95% of the TyrD (Fig. 1).

For the experiments samples were diluted to 2 mg Chl/mL by addition of the appropriate amount of high molar measuring buffer, containing 300 mM sucrose, 10 mM NaCl, and 3 mM MgCl2 with 150 mM of either glutamic acid (pH 4.7), MES (pH 6.3), or glycylglycine (pH 8.5). The final buffer concentration after addition of high molar buffer was 25 mM.

Steady-state oxygen evolution activity was measured with a Hansatech Clark-type electrode at 20 μg of Chl/mL in a measuring buffer at pH 6.1. 2 mM potassium ferricyanide K3[Fe(CN)6] and 0.5 mM PpBQ were used as electron acceptors. The activities of the PSII membrane preparation was ∼550 μmol of O2× (mg of Chl)−1 × h−1 (pH 6.3), while Tris-washed samples did not show any oxygen evolution.

EPR measurements

Room temperature EPR measurements were performed with ELEXSYS E500 spectrometer (Bruker Biospin GmbH) equipped with a SuperX bridge and a super high Q SHQE4122 cavity. The measurements were done in a 250 μL quartz flat cell at a sample concentration of ~2 mg of Chl/mL. Steady-state TyrZ/D oxidation was monitored after induction with LED setup (white or far-red light, see below) mounted at the EPR cavity window at field position of 3465 G. In addition, TyrZ/D oxidation kinetics was triggered with a 6 ns, 100 mW, 523 or 732 nm laser flash given to the sample. Data analysis was performed with the Bruker Xepr 2.1

![Fig. 1](image)

Fig. 1 EPR spectra of the Tyr radicals from the Tris-washed PS II membranes at pH 8.5 induced by white light illumination (A, black spectra) or far-red light illumination (B, red spectra). Spectra shown are after 10 h dark incubation at room temperature (dotted line spectra), during continuous illumination (dashed line spectra) and after 5 min of dark incubation (solid line spectra). The arrows indicate the field position (3465 G) for the kinetics measurements. EPR conditions: microwave frequency 9.75 GHz, microwave power 8 mW, modulation amplitude 5 G, temperature 293 K
software. The standard error in the signal amplitude estimation in our EPR measurements was less than 5%.

**Light sources**

Samples were illuminated with two kinds of LED: white and far-red light. White light LED had one major emission peak at 450 nm and two smaller peaks at 549 and 600 nm. Far-red light LED had an emission peak at 732 nm. Two cut-off Schott filters CC4 and RG9 were used with far-red LED to ensure no visible-light contamination (see Supplementary Fig. 1S for the actual output spectra of used LED light sources). LEDs were set up at the EPR cavity window. 532 and 732 nm laser flashes (20 mJ, pulse duration 6 ns, pulse bandwidth ±0.1 nm) were provided by the Quanta-Ray MOPO-730 optical parametric oscillator, driven by Nd: YAG laser (Spectra Physics, USA).

**Results**

TyrD• is much less stable in the Mn-depleted PS II than in the active PS II samples with the full O2-evolving activity, where reduction of TyrD• requires application of the reducing agents (Vass and Styring 1991; Sjöholm et al. 2016; Ahmadova et al. 2017). The dark incubation of our Tris-washed PS II membranes for 10 h at room temperature resulted in the reduction of 90–95% of TyrD• (Fig. 1, dotted line spectra). Under illumination, both tyrosines can be observed by the conventional EPR spectroscopy; however, their different kinetic properties allow to clearly distinguish between TyrD• and TyrZ• after light is switched off. Moreover, similar EPR properties of both radicals in the absence of the Mn cluster allow to quantify TyrZ• on the basis of TyrD• radical (Boussac and Etienne 1982a, 1984; Roffey et al. 1994).

Illumination of the sample for 2 min with white light at pH 8.5 and subsequent 5 min dark adaptation resulted in full oxidation of TyrD• (Fig. 1A, solid line spectrum). Any induced TyrZ• has decayed within a few minutes of dark adaptation (decay half-time of TyrZ• is ca 600 ms in Tris-washed PS II at this pH, Table 1) and any decay of TyrD• is negligible. Thus, the resulted spectrum is taken for 100% of TyrD• and was used in further quantifications. When measurements were performed under illumination conditions, tyrosine radical spectrum arose from TyrD• (100%) and the additional intensity, which is attributed to the TyrZ• radical (75%, Fig. 1A, dashed line spectrum). When measurements were done under far-red illumination, the additional intensity from TyrZ• amounted to only 15% (Fig. 1B, dashed line spectrum).

The kinetics of TyrD• and TyrZ• oxidation were measured by monitoring the EPR signal induction at 3465 G (arrow in Fig. 1) under continuous illumination at three different pH values (Fig. 2). The field position chosen for kinetic measurement is lying outside of the magnetic field range in which signals from Chl and Car cations could contribute (Visser et al. 1977; Hanley et al. 1999).

**Tyrosine oxidation under continuous white or far-red light illumination**

Steady-state oxidation of TyrD• and TyrZ• residues in the Mn-depleted PS II membranes under continuous illumination with white or far-red light are shown in Fig. 2. The oxidation kinetics was accelerated towards the higher pH values under both illumination conditions. The total yield of tyrosine radical formation was also pH dependent and increased towards high pH. The maximum formation of tyrosine radicals was observed at pH 8.5 (Fig. 2C).

Under white light illumination, the steady-state level of tyrosine oxidation reached maximum at any given pH quite fast within ca 50 s (Fig. 2, black traces). The maximum corresponds to 65% at pH 4.7, 100% at pH 6.3, and 195% at pH 8.5. Almost 200% of oxidation indicates that full induction of both TyrZ• and TyrD• was achieved at high pH. Some decay of the steady-state level of the signal was observed during illumination after 50 s at pH 6.3 (~5%) and 100 s at pH 8.5 (~10%) (Fig. 2B, C, black traces). This decay is due to the full reduction of the acceptor side which leads to the backflow of electrons from QA− or QB− to TyrZ• and/or photoinhibition. Such decay was not observed in the presence

| Table 1 | Fitted half-times and amplitudes of TyrZ• exponential decay after induction by 532 and 732 nm laser flash in the Tris-washed PSII membranes without any additions or in the presence of 2 mM ferricyanide ( Ferri) and 2 mM ferricyanide and 1 mM DPC ( Ferri+DPC) |
|--------|------------------------------------------------------------------------------------------------|
| t½ (ms) (Ampl., %) | pH 4.7 | pH 6.3 | pH 8.5 |
| No add. | Ferri | Ferri+DPC | No add. | Ferri | Ferri+DPC | No add. | Ferri | Ferri+DPC |
| 532 nm | 22 ± 2 ms (34%) | 22 ± 2 ms (85%) | 10 ± 1.5 ms (42%) | 222 ± 25 ms (43%) | 118 ± 15 ms (95%) | 17 ± 2 ms (55%) | 557 ± 53 ms (48%) | 436 ± 49 ms (109%) | 59 ± 7 ms (25%) |
| 732 nm | 21 ± 2 ms (18%) | 24 ± 2 ms (20%) | 6 ± 1 ms (8%) | 170 ± 21 ms (19%) | 105 ± 13 ms (27%) | 24 ± 2 ms (25%) | 596 ± 53 ms (20%) | 338 ± 37 ms (37%) | 76 ± 7 ms (16%) |

The standard error in the signal amplitude estimation in our EPR measurements was <5%.
of exogenous acceptor (see below). Interestingly, after the light was turned off and 5 min of dark incubation of white light illuminated samples, no decay of tyrosine amplitude was observed at pH 4.7 (Fig. 2A, black dot). While at higher pH, the amplitude of tyrosine signal decreased to 67% at pH 6.3 and to 105% at pH 8.5 (Fig. 2B, C, black dots).

The kinetics of tyrosine oxidation under the far-red illumination was different from the kinetics under the white light illumination at all three pH values investigated. Oxidation of tyrosine was much slower at pH 4.7 and 6.3 (Fig. 2A, B, red traces). However, the sharp rise has been observed at pH 8.5 and was comparable to the white light oxidation rise (Fig. 2C, red trace). The total yield of Tyr$^\bullet$ under far-red light illumination was 47% at pH 4.7, 60% at pH 6.3, and 105% at pH 8.5. At pH 4.7 and 6.3 Tyr oxidation never reached the steady-state level even after illumination for 325 s (Fig. 2A, B, red traces), whereas at pH 8.5, the oxidation reached half of the level of tyrosine oxidation under white light illumination very fast within 50 s and was continued to slowly rise afterwards (Fig. 2C, red trace). Incubation of far-red illuminated samples for 5 min resulted in the additional rise of tyrosine amplitude at pH 4.7 and 6.3 to 67% (Fig. 2A and B, red dot). In contrast, at pH 8.5 decrease of tyrosine amplitude to 91% was observed (Fig. 2C, red dot).

To conclude this part, continuous illumination of the reduced Mn-depleted PSII membranes with white light resulted in the formation of stable Tyr$_Z^\bullet$ radical at pH 4.7 and 6.3 as could be judged from its post-illumination stability. The formation was much slower and less effective under far-red light illumination if compared with white light illumination. In contrast, at pH 8.5 illumination with white light resulted in the formation of both Tyr$_Z^\bullet$ radical and Tyr$_D^\bullet$ radical, while illumination with far-red light resulted in albeit fast but only Tyr$_D^\bullet$ formation. Thus, the only conditions which resulted in the observation of semi-stable Tyr$_Z^\bullet$ were high pH and white light.

**Flash-induced tyrosine signal formation**

In order to further investigate the sequence of tyrosine radical formation in PS II, the monochromatic laser flashes at 532 and 732 nm were used. Figure 3A-C show the kinetic of Tyr$_Z^\bullet$ and Tyr$_D^\bullet$ formation and decay after five consecutive flashes separated by 5 s interval (indicated by arrows). Tyr$^\bullet$ was formed after each flash is divided into the decaying and non-decaying tyrosine signal. According to the decay half-times of Tyr$_Z^\bullet$ and Tyr$_D^\bullet$ in the Mn-depleted PSII preparations, the decaying part of tyrosine signal is attributed to Tyr$_Z^\bullet$ and non-decaying part to Tyr$_D^\bullet$ (Babcock and Sauer 1975a; Vass and Styring 1991). At pH 4.7, with green and far-red flashes, Tyr$_Z$ and Tyr$_D$ oxidation took place in 5–10% of PS II centers (Fig. 3A). Similarly, very small induction of Tyr$_Z$ and Tyr$_D$ was observed at pH 6.3 with far-red flashes (15–20%, Fig. 3B, red trace). With green flashes at pH 6.3 however, both tyrosines were induced in the substantial number of PS II centers, resulting in 41% of non-decaying Tyr$_D$ (black trace). At pH 8.5 significant amount of both tyrosines was formed at both wavelengths as could be judged from the decaying and non-decaying parts similarly to what was observed under continuous illumination. Again the final amplitude of Tyr$_D$ induced by green flashes (117%) was higher than induced by far-red flashes (72%, Fig. 3C).

Contribution of Tyr$_Z$ to the total oxidation process was studied in more detail in the same samples after Tyr$_D$ oxidation was completed (Fig. 3D-F). After accumulation, it is clear that Tyr$_Z$ signal was inducible at all three pHs by both 532 and 732 nm wavelengths. The amplitude of Tyr$_Z$ was twice higher after green flashes if compared to far-red
flashes. Similarly to the continuous illumination experiments, TyrZ oxidation was pH dependent and was small at pH 4.7 and increased towards pH 8.5. This difference was pronounced with green flash induction (Fig. 3D-F, black traces). TyrZ• signal decay was also pH dependent and slowed down towards higher pHs with both wavelengths, from 21 ms at pH 4.7 to 596 ms at pH 8.5 (Fig. 3D-F; Table 1).

Influence of exogenous electron acceptor

Better oxidation of tyrosines could be achieved by addition of ferricyanide which also prevents the loss of charge separation by recombination of QA− TyrZ• and QB− TyrZ• states (Bishop and Spikes 1955; Delrieu and Rosengard 1989). We have measured oxidation of tyrosine in the presence of ferricyanide in order to further understand the tyrosine oxidation in Mn-depleted PS II under continuous illumination with white and far-red light. Figure 4 (green traces) show the oxidation kinetics in the presence of ferricyanide at pH 4.7, 6.3 and 8.5 respectively. At pH 4.7 the amplitude of tyrosine oxidation was twice higher in the presence of ferricyanide if compared to a trace without any addition (Fig. 4, green traces). However, under far-red illumination, the kinetics was slower, but more efficient, and still rising after 325 s of illumination (Fig. 4B, green trace).

The kinetics of tyrosine oxidation measured at pH 6.3 with the ferricyanide addition under white light was very similar to the kinetics obtained without any additions (100% oxidation), except for the absence of the small decay after the maximum amplitude was reached (Fig. 4C, green and black traces). This is reasonable since the presence of the acceptor is preventing decay to the steady-state equilibrium between tyrosine oxidation and recombination reactions. Under far-red light however, the oxidation kinetics were much slower and rose to the higher level in the presence of ferricyanide than without any additions and reached the same amplitude (100%) as under white light illumination (Fig. 4D, green trace). At higher pH, under white light illumination, the oxidation kinetics and amplitude were again very similar in the presence or absence of ferricyanide reaching ca 200% indicating full oxidation of both TyrZ and TyrD (Fig. 4E, green and black traces). Interestingly, under far-red light at pH 8.5, the amplitude was higher than without acceptor almost reaching 142% (Fig. 4F, green trace) if compared to the black trace without additions (100%). This indicates that in addition to the full induction of TyrD•, 50% of TyrZ• was induced by

![Fig. 3 TyrZ and TyrD oxidation kinetics at 3465 G in the reduced Mn-depleted PS II membranes, induced by a train of five 532 nm (black traces) or 732 nm laser flashes (red traces) at pH 4.7 (A), pH 6.3 (B) and pH 8.5 (C). Each trace represents an average of four single measurements in independent samples. TyrZ oxidation kinetics induced by single 532 nm (black traces) or 732 nm laser flash (red traces) at: pH 4.7 (D), pH 6.3 (E) and pH 8.5 (F). Each trace represents an average of 199 flashes. EPR conditions are the same as in Fig. 2](image-url)
far-red light at pH 8.5 in the Mn-depleted PS II centers. The rise of the signal was also fast and comparable to the white light induction at these conditions.

**Influence of exogenous electron donor**

To distinguish between direct (by P$_{680}^+$) or indirect (via equilibrium with Tyr$_Z^*$, see Scheme 2) oxidation of Tyr$_D$, DPC as an exogenous electron donor to Tyr$_Z^*$ was added to the samples before illumination. It is known that addition of DPC accelerates the Tyr$_Z^*$ lifetime and this makes it less available for Tyr$_D$ oxidation (Babcock and Sauer 1975b; Yerkes and Babcock 1980; Roffey et al. 1994). Figure 4A–D (pink traces) show oxidation kinetics under continuous white and far-red light illumination in the presence of DPC at pH 4.7 and 6.3, respectively. It is clear that addition of DPC significantly inhibited tyrosine oxidation in the majority of PS II centers under both illumination conditions. At pH 6.3 oxidation was slightly better than at pH 4.7; however, at both pHs tyrosine oxidation did not reach the complete equilibrium and the kinetics were still rising after 325 s of both white and far-red light illumination.
illumination (Fig. 4A-D, pink traces). Interestingly, at pH 8.5 the tyrosine induction was better and both white and far-red light illumination have the same effect on oxidation which reached 70% of the PS II centers (Fig. 4E, F, pink traces).

We also performed these measurements in the presence of both exogenous donor and acceptor, DPC and ferricyanide, to see how the steady-state equilibrium between tyrosine oxidation and recombination reaction will be affected (Fig. 4, blue traces). The effect was very pH dependent. At pH 4.7 the presence of both DPC and ferricyanide resulted in higher tyrosine amplitude than in the presence of only DPC under both light conditions (Fig. 4A, B, blue traces). It restored amplitude to the level which was achieved in the measurements without any additions (compare to black and red traces). At pH 6.3 the effect was bigger and final amplitude of Tyr• was significantly higher than in the presence of only DPC and 25–35% higher than in a sample without any additions (Fig. 4, compare black, pink, and blue traces (C) and red, pink, and blue traces (D). The oxidation kinetics under white light was however slowed down (Fig. 4C, blue trace). At pH 8.5 addition of ferricyanide had no effect and the final oxidation level was very similar to the level obtained only in the presence of DPC (Fig. 4E, F, pink and blue traces).

Scheme 2 Electron transfer events and redox equilibria leading to TyrD and TyrZ oxidation in the Mn-depleted PS II centers

![Scheme 2](image)

Fig. 5 TyrZ and TyrD oxidation kinetics in the reduced Mn-depleted PS II membranes, induced by a train of five 532 nm (A, black traces) or 732 nm laser flashes (B, red traces) at pH 4.7 in the presence of 2 mM ferricyanide (solid line) or in the presence of 2 mM ferricyanide and 2 mM DPC (dotted line). TyrZ oxidation kinetics induced by single 532 nm (C, black traces) or 732 nm laser flash (D, red traces) at pH 4.7 in the presence of 2 mM ferricyanide (solid line) or in the presence of 2 mM ferricyanide and 2 mM DPC (dotted line). EPR conditions are the same as in Fig. 3

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It should be noted that only TyrD• was formed in the presence of DPC or DPC and ferricyanide under both white and far-red light as can be estimated from the residual signal after the light was switched off (not shown). We were not able to observe any fast decaying TyrZ• and the signal was never higher than 100% at all pH values measured. Our conclusion from these measurements is that DPC is an effective donor to TyrZ• at all three pH values and the addition of ferricyanide only eliminates the recombination reaction from the acceptor side of PS II which takes place at low pH values (Ahmadova et al. 2017).

Flash-induced tyrosine signal formation in the presence of donor and acceptor

Figures 5, 6, and 7 show the kinetics of TyrZ• and TyrD• formation and decay after five consecutive laser flashes separated by 5 s each (indicated by arrows) in the presence of electron donor and acceptor. In the presence of only ferricyanide we observed TyrZ• oxidation at pH 4.7 and 6.3 from the first given green flash (Figs. 5A, 6A, black solid traces). At pH 8.5 oxidation was very efficient and resulted in complete oxidation of TyrZ with consequent decay and of TyrD (Fig. 7A, black solid trace). Interestingly, no fast decay was observed after the first flash, similar to measurements without any additions (Fig. 3C, black trace).

Interestingly, we did not detect TyrZ oxidation with single far-red flashes in the presence of ferricyanide and final TyrD oxidation was less than 12% at pH 4.7 or at pH 6.3 (Figs. 5B, 6B, red solid traces). At pH 8.5 with far-red flashes, we observed almost full oxidation of TyrD after five flashes (80%, Fig. 7B, red solid trace). Unlike in oxidation with green flashes, the fast decay kinetics (TyrZ•) was observed on the third flash and onward but seemingly the first two flashes induced only TyrD oxidation in the majority PS II centers at high pH (48%, Fig. 7B, red solid trace).

Accumulated TyrZ• signal induced by green or far-red flashes in the presence of ferricyanide is shown in Figs. 5, 6, and 7C and D, solid traces. With 532 nm induction, the amplitude of TyrZ• slightly increased towards high pH and corresponded to 85% at pH 4.7, 95% at pH 6.3, and 109% at pH 8.5. With 732 nm induction, we observed only 20% at pH 4.7, 27% at pH 6.3, and 37% at pH 8.5. The decay half-times were also pH dependant and corresponded to 22 ms.
at pH 4.7, 118 ms at pH 6.3, and 436 ms at pH 8.5, Figs. 5, 6, and 7C and D, solid traces, Table 1. The decay half-time was independent of the induction wavelength.

We also investigated the kinetics of flash-induced tyrosine signal formation in the presence of both DPC and ferricyanide (Figs. 5, 6, 7A and B, dotted traces). The addition of DPC almost completely inhibited the tyrosine formation at pH 4.7 and 6.3. We did not observe any TyrZ* formation in the presence of DPC and the final TyrD* formation was less than 5% at these two pH values (Figs. 5, 6A and B, dotted traces). At pH 8.5, the TyrZ* signal was still unresolvable in the presence of DPC but TyrD oxidation occurred in the majority of the PS II centers with 532 nm flashes (>75%, Fig. 7A, black-dotted trace) and in less centers with 732 nm flashes (45%, Fig. 7B, red-dotted trace). Figures 5, 6, and 7C and D show that in the presence of DPC the TyrZ* induction is strongly inhibited and very short lived (ca 20 ms or less, Table 1) at low pH values, while at high pH the half-time was less affected and significant loss of TyrZ* amplitude could be attributed to competitive reduction of P_{680}^+ by TyrD (Scheme 2) (Fig. 7C, D, dotted traces).

**Discussion**

The special Chl molecules, P_{680}, serve as a primary electron donor in PS II. P_{680} is a tetrameric pigment entity which comprises four Chl molecules. The central Chl pair, P_{D1} and P_{D2} are weakly excitonically coupled and situated at 30° angle to the horizontal plane. The other two Chls, Chl_{D1} and Chl_{D2} are occupying symmetrical positions at 10 Å center-to-center distance each from P_{D1} and P_{D2}, respectively (Scheme 1) (Umena et al. 2011; Suga et al. 2015; Wei et al. 2016). The localization of excitation energy in P_{680} and the first Chl* electron donor formed after the “standard” charge separation conditions (visible-light excitation) have been extensively studied by different spectroscopic methods (Zech et al. 1997; Diner et al. 2001; Groot et al. 2005; Holzwarth et al. 2006; Romero et al. 2010, 2012). The primary hole is consensually placed on the P_{D1} Chl (P_{D1}^+) (Scheme 1) (Hillmann et al. 1995; Diner et al. 2001; Schlodder et al. 2008b; Cardona et al. 2012) although some groups reported that weak spectral differentiation among all four Chls might lead to a distribution of the excitation energy at ambient temperature (Romero et al. 2010, 2012). There are also reports...
that the reduction of Pheo$_{D1}$, which is the primary acceptor in PS II (Scheme 1) occurs prior to the oxidation of P$_{D1}$/P$_{D2}$ (Groot et al. 2005; Holzhwarth et al. 2006).

The far-red photochemistry at low temperature has been suggested to induce different primary charge pair, Chl$_{D1}^+$ Pheo$^-$ (Mokvist et al. 2014). This was based on the different donation efficiency to P680$^+$ from Tyr$_Z$ and Cyt b$_{559}$/Chl$_Z$ pathways under green and far-red illumination at 5 K, in the so-called product analysis of the charge-separated state (Mokvist et al. 2014). It seems that nature of the primary electron hole in P$_{680}^+$ varies depending on the temperature and excitation wavelength (Raszewski et al. 2008). In this paper, we investigate if the similar effect of excitation wavelength (visible vs. far-red light) on the primary donor occurs at physiological conditions. Tyr$_Z$ and Tyr$_D$, which are symmetrically positioned at about 9.2 Å distance from the central P$_{D1}$ and P$_{D2}$ Chls, respectively, (Scheme 1) were used as competing electron donors to elucidate the nature of the primary charge-separated state at room temperature.

The task is complicated by the fact that there are two electron transfer pathways for Tyr$_D$ to be oxidized in the Mn-depleted PS II. The first one is occurring via Tyr$_Z^*$ which is formed rapidly after oxidation by P$_{680}^+$. Tyr$_D^+$ with its lower redox potential, is then slowly oxidized in the following reaction (Boussac and Etienne 1982c, 1984; Faller et al. 2001):

$$\text{Tyr}_D\text{Tyr}_Z \xrightarrow{\text{light}} \text{Tyr}_D\text{Tyr}_Z^* \rightarrow \text{Tyr}_D^+\text{Tyr}_Z$$  \hspace{1cm} (1)

This oxidation pathway occurs at low and middle pH values. However, at high pH (above pK$_a$ of Tyr$_D$ (Vass 1991; Faller et al. 2001; Ahmadova et al. 2017)), direct Tyr$_D^+$ oxidation by P$_{680}^+$ was reported in the Mn-depleted preparation in at least half of the PS II centers (Faller et al. 2001). As a result, partial localization of the electron hole on P$_{D2}$ Chl was suggested (Faller et al. 2001).

Our data show that under continuous illumination we mostly observed oxidation of Tyr$_D$ (except white light illumination at high pH where both Tyr$_Z^*$ and Tyr$_D^*$ were formed (Fig. 2)). The pH dependence of Tyr$_D^*$ induction was similar to what was reported on green flash-induced Tyr$_D^*$ formation in intact PS II (Vass and Styring 1991; Sjöholm et al. 2016; Ahmadova et al. 2017). In our Tris-washed PSI membranes, under continuous white light illumination Tyr$_D^+$ oxidation takes place in 65 and 100% at pH 4.7 and 6.3, respectively (Fig. 2A, B, black traces). With far-red light illumination (732 nm LED light), oxidation at these pH values was much slower and less effective (Fig. 2A, B, red traces). The diminished formation of Tyr$_D^+$ under far-red light could be due to either less effective charge separation or due to the different nature of the primary donor which makes the far-red photochemistry less effective.

The first hypothesis seems unlikely because under 732 nm light most of the PS II centers undergo charge separation. As it was shown by (Thapper et al. 2009) illumination with 730 nm light resulted in P$_{680}^+$ Pheo$^-$ primary charge pair formation in the vast majority of the PS II centers. This is by far in more centers than the decreased tyrosine radical formation under similar 732 nm illumination (Fig. 2A, B). Thus, the possibility of the second hypothesis cannot be ruled out. At pH 8.5 continuous white light illumination induces both tyrosines. Interestingly, illumination with 732 nm light resulted in only Tyr$_D^*$ formation (Fig. 2C, red trace). Full induction of Tyr$_D^*$ (100%) also indicates that far-red light excites 100% of the PS II centers, as mentioned above. The difference between tyrosine formation at pH 8.5 under visible and far-red illumination could originate either from different tyrosine oxidation pathways (Tyr$_D$ vs. Tyr$_Z$) or different localizations of the primary donor. Here our data with the addition of exogenous donor and acceptor to regulate the “redox pressure” in and out from Tyr$_Z$ (Scheme 2) will help to answer these questions.

The addition of ferricyanide increased oxidation amplitude of Tyr$_D$ at low pH by efficiently preventing recombination from the acceptor side of PS II (Fig. 4, green traces). At pH 6.3 and 8.5, the effect of ferricyanide addition was only observable under the far-red illumination (Fig. 4D, F). Since we also observed that the addition of ferricyanide also increased the amplitude of Tyr$_Z^*$ induction (Table 1), it is clear that final oxidation of Tyr$_D$, at pH 4.7 and 6.3, takes place via Tyr$_Z^*$ as described in reaction (1). More importantly, this indicates that recombination reaction was more efficiently prevented by the ferricyanide addition under far-red light illumination. This implies different recombination partners formed under the far-red light on the donor side of PS II. Since Tyr$_Z^*$ is the same at both illumination conditions, the only difference could be assumed at the P$_{680}^+$ entity.

The most important and informative results were obtained when measurements were done in the presence of DPC. DPC is known to be an efficient electron donor to Tyr$_Z^*$ in the Mn-depleted PS II preparations (Babcock and Sauer 1975b; Yerkes and Babcock 1980; Roffey et al. 1994). In our case, both Tyr$_D^+$ and DPC are competing for the Tyr$_Z^*$ reduction and in the presence of DPC the decay half-time of Tyr$_Z^*$ is significantly decreased (Table 1); thus, effectively blocking indirect Tyr$_D$ oxidation (Scheme 2). The addition of DPC severely inhibited Tyr$_D^*$ formation at pH 4.7 and 6.3 under both white and far-red illuminations (Fig. 4A-D, pink traces). DPC inhibition of the Tyr$_D^*$ formation was more effective under far-red light illumination (only 18 and 19% formation at pH 4.7 and 6.3, respectively). Interestingly, even under white light illumination at pH 8.5 only Tyr$_D^+$ oxidation was observed in the presence of DPC and no extra intensity could be attributed to the Tyr$_Z$ oxidation.
(Fig. 4E, F, pink traces). This is another indication for the direct oxidation of TyrD by P_{680}^{+} at high pH (Scheme 2) (Faller et al. 2001).

In the presence of both DPC and ferricyanide, the oxidation kinetics was found to be similar to what was found in the absence of any additions (Fig. 4A-D, blue traces). In this case, availability of the electron donor (DPC) and acceptor (ferricyanide) to and from TyrZ^{*} allowed the same final steady-state oxidation level, thus, effectively restoring the original “redox pressure” on TyrZ^{*}.

Flash-induced TyrZ^{*} oxidation was observed at all pHs and under both 532 and 732 nm laser flash in our Tris-washed PS II membranes. It was both pH dependent (as was reported before (Boska et al. 1983)) and wavelength dependent (Table 1; Figs. 3E, D, and 5, 6, 7C and D). The decay half-time of TyrZ^{*} was pH dependent (Babcock and Sauer 1975a; Shigemori et al. 1997) but wavelength independent (Table 1). This indicates that after the TyrZ^{*} formation (if any), its consequent reduction either from TyrD^{*} or DPC (when present) or by recombination from the acceptor side (Scheme 2) was independent on the way of how the primary charge separation occurred. At pH 4.7 and 6.3 the presence of DPC decreased the amplitude of TyrZ^{*}, especially under 732 nm flash (to final 3% and 25%, respectively if compared to 532 nm flash (Figs. 5, 6C and D; Table 1)). This resulted in significantly less TyrD^{*} formation, especially after far-red flashes (Figs. 5, 6A and B). In contrast at high pH 8.5, even if the effect of DPC addition on TyrZ^{*} kinetics was similarly dramatic, a significant amount of TyrD^{*} was formed. The absence of the fast decay on the first flash in Fig. 7A, B points out to the direct oxidation of TyrD^{*} by P_{680}^{+} under both green and far-red (although much less efficiently) flashes.

Thus, our data indicate that different photochemistry is involved in oxidation of two tyrosines in the Mn-depleted PS II membranes. The far-red light-induced photochemistry is taking place in the majority of the PS II centers. At normal pH values, it results in the decrease of TyrZ^{*} formation and correspondingly higher recombination rate under the far-red light illumination. Recombination reaction takes place between the acceptor side of PS II (Q_{A}^{-} or Q_{B}^{-}) and P_{680}^{+} which prevents TyrZ^{*} formation. The reason for this could be a different primary charge separation event and correspondingly the localization of the electron hole in P_{680}^{+} (Scheme 1).

The central Chl pair, P_{D1} and P_{D2} is excitonically weakly coupled. The greater physical separation, the slight differences in tetrapyrrole ring orientation, and the smaller dipole strength of the Q_{y} transition cause weaker electronic interaction between special Chl pair (Diner and Rappaport 2002; Raszewski et al. 2005; Schlodder et al. 2008a). This is why the Chl pair, P_{D1} and P_{D2} do not represent the lowest energy sink for the excitation energy (Diner and Rappaport 2002; Raszewski et al. 2005; Schlodder et al. 2008a). Whereas the monomeric Chl_{D1} has the lowest site energy because of the absence of such coupling (Schlodder et al. 2008a). The far-red light bears low excitation energy if compared to white light. Therefore, far-red light-induced excitation migration among four Chls in P_{680} would be an energetically less favorable process. It is more likely that under the far-red light the excitation localized on Chl_{D1}, closer to the primary electron acceptor, Pheo (Scheme 1) was shown to take place at very low temperatures (Mokvist et al. 2014).

Localization of the electron hole on Chl_{D1} is not an ideal situation since recombination from the acceptor side Chl_{D1}^{+} Q_{A}^{-} is faster and efficiently quenches productive charge separation. On the other hand, TyrZ is still in the close distance to both Chl_{D1} and P_{D1} to have an efficient donation in the centers where Chl_{D1}^{+} is still available. Moreover, at physiological temperatures, as soon as Chl_{D1} loses an electron, the hole can migrate to the neighboring P_{D1} and P_{D2}. After the hole jumps on P_{D1}, the localization of the hole on P_{D1} or P_{D2} becomes a very random process. This why at high pH direct oxidation of TyrD by P_{680}^{+} (via P_{D2}^{+} as was suggested by Faller et al. 2001) is possible at least in part of the PS II centers under the far-red light illumination.

Thus, our results indicate that in the Mn-depleted PS II at room temperature the primary charge separation pathway under the far-red excitation occurs via Chl_{D1}^{+} Pheo_{D1}^{−} primary pair, similar to what was reported for the active PS II at ultra-low temperature (Mokvist et al. 2014). The question if the same reaction occurs in the active PS II under physiological conditions requires further investigations.

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References

Ahmadova N, Ho FM, Styring S, Mamedov F (2017) Tyrozine D oxidation and redox equilibrium in photosystem II. Biochim Biophys Acta 1858(6):407–417. doi:10.1016/j.bbabio.2017.02.011

Babcock GT, Sauer K (1973) Electron paramagnetic resonance signal II in spinach-chloroplast. 1. Kinetic analysis for untreated chloroplasts. Biochim Biophys Acta 325(3):483–503. doi:10.1016/0005-2728(73)90209-6

Babcock GT, Sauer K (1975a) A rapid, light-induced transient in electron paramagnetic resonance signal II activated upon inhibition of photosynthetic oxygen evolution. Biochim Biophys Acta 376(2):315–328. doi:10.1016/0005-2728(75)90024-9
Raszkewski G, Diner BA, Schloder E, Renger T (2008) Spectroscopic properties of reaction center pigments in photosystem II core complexes: revision of the multimer model. Biophys J 95(1):105–119. doi:10.1529/biophysj.107.123935

Renger G (2012) Mechanism of light induced water splitting in photosystem II of oxygen evolving photosynthetic organisms. Biochim Biophys Acta 1817(8):1164–1176. doi:10.1016/j.bbabio.2012.02.005

Reimers JR, Biczysko M, Bruce D, Coker DF, Frankcombe TJ, Hashimoto H, Hauer J, Jankowiak R, Kramer T, Linnanto J, Mamedov F, Müh F, Rätspe M, Renger T, Sjöholm J, Wan J, Wang Z, Wang-Otomo Z-Y, Weng Y-X, Yang C, Zhang J-P, Freiberg A, Krausz E (2016) Challenges facing an understanding of the nature of low-energy excited states in photosynthesis. Biochim Biophys Acta 1857(9):1627–1640

Renger G, Renger T (2008) Photosystem II: the machinery of photosynthetic water splitting. Photosynth Res 98(1–3):53–80. doi:10.1007/s11120-008-9345-7

Roffey RA, van Wijk KJ, Sayre RT, Styring S (1994) Spectroscopic characterization of tyrosine-Z in histidine 190 mutants of the D1 protein in photosystem II (PS II) in Chlamydomonas reinhardtii. Implications for the structural model of the donor side of PS II. J Biol Chem 269(7):5115–5121

Romero E, van Stokkum IJM, Novoderezhkin VI, Dekker JP, van Grondelle R (2010) Two different charge separation pathways in photosystem II. Biochemistry 49(20):4300–4307. doi:10.1021/bi1003926

Romero E, Diner BA, Nixon PJ, Coleman WJ, Dekker JP, van Grondelle R (2012) Mixed excitation–charge-transfer states in photosystem II: stark spectroscopy on site-directed mutants. Biophys J 103(2):185–194. doi:10.1016/j.bpj.2012.06.026

Saito K, Rutherford AW, Ishikita H (2013) Mechanism of tyrosine D oxidation in photosystem II. Proc Natl Acad Sci USA 110(19):7690–7695. doi:10.1073/pnas.1300817110

Schlodder E, Coleman WJ, Nixon PJ, Cohen RO, Renger T, Diner BA (2008a) Site-directed mutations at D1-His198 and D1-Thr179 of photosystem II in Synechocystis sp. PCC 6803: deciphering the spectral properties of the PS II reaction centre. Philos Trans 363(1494):1197–1202

Schlodder E, Renger T, Raszkewski G, Coleman WJ, Nixon PJ, Cohen RO, Diner BA (2008b) Site-directed mutations at D1-Thr179 of photosystem II in Synechocystis sp. PCC 6803 modify the spectroscopic properties of the accessory chlorophyll in the D1-branched reaction center. Biochemistry 47(10):3143–3154. doi:10.1021/bi0702059f

Shigemori K, Mino H, Kawamori A (1997) pH and temperature dependence of tyrosine Z’ decay kinetics in tris-treated PS II particles studied by time-resolved EPR. Plant Cell Physiol 38(9):1007–1011. doi:10.1093/oxfordjournals.pcp.a029264

Sjöholm J, Ho FM, Ahmadova N, Brinkert K, Hammarström L, Mamedov F, Styring S (2016) The protonation state around TyrD/TyrD’ in photosystem II is reflected in its biphasic oxidation kinetics. Biochim Biophys Acta 1858:147–155

Styring S, Rutherford AW (1987) In the oxygen evolving complex of photosystem II the S0 state is oxidized to the S1 state by D+ (signal II slow). Biochemistry 26(9):2401–2405. doi:10.1021/bi00383a001

Styring S, Sjöholm J, Mamedov F (2012) Two tyrosines that changed the world: interfacing the oxidizing power of photosynthesis to water splitting in photosystem II. Biochim Biophys Acta 1817:76–87

Suga M, Akita F, Hirata K, Ueno G, Murakami H, Nakajima Y, Shimizu T, Yamashita K, Yamamoto M, Ago H, Shen JR (2015) Native structure of photosystem II at 1.9 Å resolution viewed by femtosecond X-ray pulses. Nature 517(7532):99–U265. doi:10.1038/nature13991

Thapper A, Mamedov F, Mokvist F, Hammarström L, Styring S (2009) Defining the far-red limit of photosystem II in spinach. Plant Cell 21(8):2391–2401. doi:10.1105/tpc.108.064154

Umema Y, Kawakami K, Shen JR, Kamiya N (2011) Crystal structure of oxygen-evolving photosystem II at a resolution of 1.9 Å. Nature 473(7345):55–U65. doi:10.1038/nature09913

Vass I, Styring S (1991) pH-dependent charge equilibria between tyrosine-D and the S states in photosystem II. Estimation of relative midpoint redox potentials. Biochemistry 30:830–839

Vinary DJ, Ananyev GM, Dismukes GC (2013) Photosystem II: the reaction center of oxygenic photosynthesis. In: Kornberg RD (ed) Annual review of biochemistry, vol 82. Annual Reviews, Palo Alto, pp 577–606. doi:10.1146/annurev-biochem-070511-100425

Visser JWM, Rijgersberg CP, Gast P (1977) Photooxidation of chlorophyll in spinach chloroplasts between 10 and 180 K. Biochim Biophys Acta 460(1):36–46. doi:10.1016/0005-2728(77)90149-9

Völker M, Ono T, Inoue Y, Renger G (1985) Effect of trypsin on PSII particles: correlation between Hill-activity, Mn-abundance and peptide pattern. Biochim Biophys Acta 806:25–34

Wei X, Su X, Cao P, Liu X, Chang W, Li M, Zhang X, Liu Z (2016) Structure of spinach photosystem II–LHCII supercomplex at 3.2 Å resolution. Nature 534 (7605):69–74. doi:10.1038/nature18020. http://www.nature.com/nature/journal/v534/n7605/abs/nature18020.html#supplementary-information

Yerci CT, Babcock GT (1980) Photosystem II oxidation of charged electron donors. Surface charge effects. Biochim Biophys Acta 690(3):360–372. doi:10.1016/0005-2728(80)90207-8

Zech SG, Kurreck J, Eckert H-J, Renger G, Lubitz W, Bittl R (1997) Pulsed EPR measurement of the distance between P680+ and QA− in photosystem II. FEBS Lett 414(2):454–456. doi:10.1016/S0005-2728(97)01054-5