Review

Electron transfer in photosystem I

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Abstract

This mini-review focuses on recent experimental results and questions, which came up since the last more comprehensive reviews on the subject. We include a brief discussion of the different techniques used for time-resolved studies of electron transfer in photosystem I (PS I) and relate the kinetic results to new structural data of the PS I reaction centre. ß 2001 Elsevier Science B.V. All rights reserved.

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1. Introduction

After the purple bacterial reaction centre [13], photosystem I (PS I) is the second photosynthetic reaction centre for which a highly resolved X-ray structure has been obtained [17,30]. An important difference from the purple bacterial reaction centre is that PS I contains a chain of four low potential (E_m < −500 mV versus NHE) secondary electron acceptors, the phylloquinone A1 and three [4Fe-4S] clusters named FX, FA and FB (see [7,21] for more comprehensive reviews on PS I). Relating structure and function of PS I may hence provide an original and essential contribution to the understanding of the principles of solar energy conversion developed by nature and – more generally – of electron transfer processes in proteins (note that photosynthetic electron transfer can be triggered by a short flash of light, allowing kinetic studies with high time-resolution).

The physiological function of PS I is to catalyse light-driven transfer of electrons from reduced plastocyanin or cytochrome c_6 located in the lumen to ferredoxin in the stroma. As oxidation and reduction of the redox partners occur on opposite sides of the membrane, PS I has to provide an efficient electron transfer pathway across the membrane. This electron transfer pathway is realised by a chain of cofactors bound to the reaction centre protein at different levels within the membrane. From a functional point of view, light absorption leads to formation of the excited state of the primary donor, P700, followed by electron transfer to a primary acceptor, A_0. The electron is then further transferred along a series of secondary electron acceptors, A_1, FX, FA, and FB, thereby crossing the membrane.

Fig. 1 presents the arrangement of the cofactors susceptible to be involved in electron transfer in PS I. P700 is a chlorophyll (Chl) dimer composed of a Chl a' (eC-A1) and Chl a (eC-B1). Two nearly sym-
metric branches of cofactors are present between P700 and F X, each composed of two chlorophylls (eC-B2/eC-A2 and eC-A3/eC-B3) and a phylloquinone (Q K-A/QK-B). Whether the two branches are used with similar probability or whether one of them is clearly dominating is presently debated (see Section 3.1). The first spectroscopically resolved electron acceptor, A0, is assumed to be represented by chlorophyll eC-A3 and/or eC-B3. Chlorophyll eC-B2 and/or eC-A2 is presumably in some way involved in electron transfer from excited P700 to A0 (see Section 3.3). These six chlorophylls, the two phylloquinones and the \[4Fe^4S\] cluster FX are bound by two large (\(\approx 83\) kDa), highly homologous subunits, PsaA and PsaB, while the two terminal electron acceptors, the \[4Fe-4S\] clusters FA and FB, are bound by the small (\(\approx 9\) kDa), stromal subunit PsaC.

A simplified scheme of the kinetics and energetics of electron transfer in PS I is presented in Fig. 2. The primary electron transfer steps (from excited P700 via A0 to A1) are ultrafast processes (picosecond time scale) with a considerable driving force. Electron transfer from A1 via FX to FA and FB has presumably a lower driving force (note that the reduction potential of A1 is not well established; see Section 3.2). The reoxidation of A1 occurs in tens to hundreds of nanoseconds; the further transfer to FA and FB is only poorly characterised (see Section 2.2).

Several techniques have been applied to study kinetics of electron transfer in photosystem I. The most abundant kinetic information was obtained by absorption change measurements. These measurements are based on the characteristic difference spectra of reduced and oxidised forms of the cofactors in the visible, UV, and near-IR spectral region. The accessible time window is very large, ranging from the sub-ps to seconds time scale. Limitations of this technique are mainly due to overlapping or similar absorption features of different cofactors, which are the same chemical species. This makes measurements of electron transfer from the P700 to the primary acceptors (all partners are chlorophyll molecules) or between the iron-sulphur centres difficult. Another difficulty is connected with the strong background absorption due to the presence of a large number of antenna chlorophylls in normal PS I preparations. Absorption changes due to singlet and triplet excited states of antenna pigments can be superimposed to the absorption changes due to electron transfer. A novel spectrophotometric technique [1] permits detection of small flash-induced absorption changes down to a nanosecond time scale even in intact cells [29].

Transient electron paramagnetic resonance (EPR) spectroscopy has been applied to follow the spin-polarised radical pair P700\(^+\)A1\(^-\) on a time scale from 10 ns to several microseconds (at room temperature), and to characterise magnetic interactions in this pair (see review by van der Est [68] in this issue). An advantage of this technique is its selectivity for paramagnetic species, so that antenna pigments do normally not interfere.

In oriented samples transmembrane electron transfer can be detected by measurements of the kinetic changes in the membrane potential (photovoltage). Unlike spectroscopic techniques, these measurements cannot provide direct information on the chemical nature and interactions of the molecules involved in the electron transfer reaction. However, besides the kinetic information, photovoltage measurements pro-
vide the electrogenicity of individual reaction steps. The electrogenicity is proportional to the dielectrically weighted transmembrane distances over which electron transfer occurs, and therefore contains structural information about the relative transmembrane distances between cofactors (when a homogeneous dielectric environment is assumed). The transmembrane organisation (see Fig. 1b) of the electron transfer pathway in PS I makes this system particularly suited for this kind of measurements. Advantages of photovoltage measurements are their inherent selectivity for electrogenic events, a high time resolution and an excellent signal-to-noise ratio. On the other hand, it should be kept in mind that electrogenic relaxations might contribute to a measured potential change. Another difficulty of the photovoltage measurements is related to the fact that different techniques are necessary for measurements on a sub-μs time scale [66] and for slower measurements (see, e.g., [65]), and that the time windows of both techniques do hardly overlap.

2. Recent data on electron transfer in PS I

2.1. Primary electron transfer (from excited P700 via \(A_0\) to \(A_1\))

The intrinsic time constant of the first step of charge separation in PS I is difficult to determine, because the intact PS I RC contains about 100 antenna chlorophylls that absorb photons and transfer excitation energy amongst them and to P700 in a few picoseconds. Hence, the apparent rate of charge separation (in the order of 30 ps) is slower than the intrinsic rate that would be observed in the absence of antenna chlorophylls upon excitation of P700. In addition, estimations of the intrinsic rate of charge separation from measured rates are dependent on the kinetic limitations, i.e., whether the excited state decay kinetics is trap limited or transfer limited. The model of excitation trapping in PS I is, however, still debated (see review by Gobets and van Grondelle [19] in this issue). Recent simulations of time-re-
solved absorption and fluorescence data yielded an intrinsic time constant of 770 fs [2] or 620 fs [18] or 500 fs [12] for electron transfer from excited P700 to the first electron acceptor, which is faster than previous estimates of 1–3 ps (reviewed in [7]). In contrast, from an analysis of the calculated difference between kinetics at 690 nm in open centres (P700 reduced prior to excitation) and in closed centres (P700 oxidised prior to excitation), a time constant of 9–10 ps was attributed to electron transfer from excited P700 to A0 in PS I from _Synechocystis_ sp. PCC 6803 [56]. A critical condition for the validity of this analysis is that the energy transfer processes in open and closed centres are identical, so that the difference signal represents exclusively electron transfer processes. As the absorbance changes due to excitation and energy transfer between antenna pigments were more than 10 times bigger than those attributed to electron transfer [56], the result of the subtraction might be affected already by subtle differences in energy transfer between open and closed centres. In [56], the overall excitation energy trapping times at P700 (~24 ps) were found to be very nearly the same in open and closed centres, justifying the subtraction procedure. Another study, however, reported a 10% increase in excitation energy lifetime upon closing the PS I reaction centre of _Synechococcus elongatus_ [12]. Summarising, the intrinsic rate of primary charge separation in PS I appears to be still uncertain.

To our knowledge, there is no direct kinetic evidence that a chlorophyll-like species other than P700 and A0 functions as a real intermediate in charge separation, although the arrangement of the cofactors suggests such a role for the chlorophyll eC-B2 and/or eC-A2 (see Section 3.3).

The time constant of electron transfer from A0 to A1 had been estimated previously from kinetic absorption spectroscopy monitoring the reoxidation of A0 to \( \tau \approx 30 \) ps (reviewed in [7]), which is close to the value of about 50 ps deduced from photovoltage measurements [25]. A recent paper [11] reported direct monitoring of the reduction of the phylloquinone A1 by ultrafast spectroscopy in the near-UV absorption band of A1 around 380 nm. The flash-induced absorption transients averaged over 380 to 390 nm increased with a time constant of 30 ps. The spectrum of the absorption changes reached at 150 ps after excitation (measured between 380 and 480 nm) resembles the spectrum at 2 ns after excitation, obtained by conventional kinetic absorption spectroscopy and attributed to the state P700\(^+\)A1\(^-\). It was concluded that electron transfer from A0 to A1 occurs directly and completely with \( \tau \approx 30 \) ps.

In another ultrafast study [46], a 28 ps phase observed in the blue spectral region was interpreted as due to overall excited state decay by electron transfer forming the state P700\(^+\)A1\(^-\). A striking result of this study is that the spectrum of the absorbance changes remaining after completion of the 28 ps phase shows a bleaching in the near-UV (measured down to 380 nm). This feature is in contrast to the ultrafast study [11] mentioned above and to several studies in nanosecond and microsecond time scales [7–10,29,39,45,63] where formation of the state P700\(^+\)A1\(^-\) was characterised by a broad absorption increase around 380 nm, largely due to the absorption of the phyllosemiquinone anion A1\(^-\). As the near-UV bleaching reported in [46] challenges the previous absorption studies on electron transfer involving A1, it would be important to check whether there is an important spectral evolution up to the nanosecond range under the experimental conditions used in [46]. Such a study might also give hints whether species not involved in electron transfer (e.g., singlet excited state of uncoupled antenna chlorophylls) contribute to the spectra on the picosecond time scale.

The same study [46] presents an (A1\(^-\)A1) difference spectrum in the 380 to 490 nm range obtained by subtracting a (P700\(^+\)P700) difference spectrum (measured on a 100-ms time scale using a conventional transient absorption set-up) from the spectrum of the absorbance changes remaining after completion of the 28 ps phase. The resulting spectrum shows a pronounced absorption increase around 430 nm and a bleaching around 400 nm, and has generally the shape of an inverted (P700\(^+\)-P700) spectrum. While a bleaching around 400 nm is in contrast to previous work (see above), the (A1\(^-\)A1) difference spectrum around 430 nm was previously not well established [6,7,10]. Because of the dominating bleaching due to oxidation of P700 around 430 nm, (A1\(^-\)A1) spectra obtained by subtraction of the (P700\(^+\)+P700) spectrum from spectra for the formation of the state P700\(^+\)A1\(^-\) are very sensitive to proper normalisation. On the other hand, electron trans-
fer from $A_1^\rightarrow$ to $FX$ ($t_{1/2} = 10–200$ ns; see Section 2.2) should be accompanied by a pronounced bleaching around 430 nm if the ($A_1^\rightarrow$–$A_1$) spectrum were to show a pronounced absorption increase ($FX$ bleaches around 430 nm upon reduction [21]). However, only a weak bleaching with kinetics in the 10 to 200 ns range has been observed around 430 nm [6,8,29,63]. Hence, unless the unusual ($A_1^\rightarrow$–$A_1$) spectrum in [46] should be due to a poor normalisation of the two experimental spectra subtracted, there would have to be considerable spectral evolution between about 100 ps and a few nanoseconds after excitation, a time range that has not yet been directly examined in PS I.

2.2. Secondary electron transfer (from $A_1^\rightarrow$ to the iron–sulphur clusters $FX$, $FA$ and $FB$)

Pathway, kinetics and energetics of secondary electron transfer in PS I are still a matter of debate. While the reoxidation of $A_1^\rightarrow$ could be studied by time-resolved absorption and EPR spectroscopy as well as photovoltage, data on the subsequent electron transfer between $FX$, $FA$, and $FB$ are scarce and mostly based on more indirect approaches.

For PS I complexes isolated from spinach, flash induced absorbance changes in the near-UV, attributed to the reoxidation of $A_1^\rightarrow$, showed biphasic kinetics with $t_{1/2} = 20$ ns and 150 ns (amplitude ratio about 2:1) [63]; transient EPR of a spin-polarised signal attributed to the state $(P700^+A_1^\rightarrow)$ resolved the 150 ns phase, but not the 20 ns phase, presumably because of insufficient time-resolution [5,67]. As a possible explanation for the biphasic reoxidation of $A_1^\rightarrow$, it was suggested [63] that the free energies of the states $P700^+A_1^\rightarrow$ and $P700^+FX$ are close to each other, i.e., the forward and backward electron transfer rates $k_1$ and $k_{-1}$ (see Scheme 1) are similar. In this model, establishment of a quasi-equilibrium between $P700^+A_1^\rightarrow$ and $P700^+FX$ gives rise to a fast phase of $A_1^\rightarrow$ reoxidation, and depopulation of this quasi-equilibrium by electron transfer from $FX$ to the next acceptor (now known to be $FA$; see below) gives rise to the slower phase.

For PS I isolated from cyanobacteria, only a single phase of $A_1^\rightarrow$ reoxidation $(t_{1/2} = 200$ ns) had been observed [6,63]. More recently, however, with an improved time resolution of 2 ns, a phase with $t_{1/2} = 7$ ns (representing about one third of the total decay of $A_1^\rightarrow$) was discovered in PS I from the cyanobacterium Synechocystis sp. PCC6803 [8]. In terms of the $A_1^\rightarrow$–$FX$ equilibrium model outlined above, these data imply that for PS I from Synechocystis, the state $P700^+FX$ would be about 20 meV higher in free energy than the state $P700^+A_1^\rightarrow$, whereas in PS I from spinach, $P700^+FX$ would be about 40 meV below $P700^+A_1^\rightarrow$ [8]. As the PS I complexes from spinach used in these studies were isolated by a rather harsh treatment, it was speculated that the pronounced 20 ns phase in these samples might result from a modification of the PS I complex during isolation [63].

Using a novel spectrophotometric technique, Joliot and Joliot [29] managed to follow the reoxidation of $A_1^\rightarrow$ in whole cells of a Chlorella sorokiniana mutant lacking most of photosystem II and of the peripheral antennae. They observed two phases of about equal amplitude with half-times of about 18 and 160 ns, respectively, which is close to the kinetics observed in PS I isolated from spinach and does not support the idea of a preparation artefact being responsible for the biphasic kinetics in spinach PS I.

In order to verify the $A_1^\rightarrow$–$FX$ equilibrium model, Joliot and Joliot [29] studied the effect of the membrane potential (varied by the presence or absence of uncoupler) on the reoxidation kinetics of $A_1^\rightarrow$. The rationale was that a change in the membrane potential should affect the equilibrium between $P700^+A_1^\rightarrow$ and $P700^+FX$ and hence – in the framework of the $A_1^\rightarrow$–$FX$ equilibrium model – the relative amplitudes of the two phases of $A_1^\rightarrow$ reoxidation. The absence of any significant effect of the membrane potential on the observed kinetics of $A_1^\rightarrow$ led Joliot and Joliot to consider the $A_1^\rightarrow$–$FX$ equilibrium model as unlikely and to assume $k_1 \gg k_{-1}$, i.e., that $P700^+A_1^\rightarrow$ is well above $P700^+FX$ in free energy [29]. As alternative explanations for the biphasic reoxidation of $A_1^\rightarrow$, they suggested that either PS I is present under two conformational states which differ by the reoxidation rate of $A_1^\rightarrow$, or that two phylloquinones (most likely

\[
(P700^+A_1^\rightarrow) \xrightarrow{k_1}{k_{-1}} (P700^+FX^\rightarrow) \xrightarrow{k_2} (P700^+FA^\rightarrow)
\]

Scheme 1.
Q$_K$-A and Q$_K$-B, see Fig. 1) are involved in electron transfer to F$_X$ with different rates (see Section 3.1).

The slower one of the two phases of A$_1$ reoxidation has been studied as a function of temperature in PS I from *S. elongatus* [57]. Between 300 and 200 K, the rate slowed down according to an activation energy of 220 meV; below 200 K, forward electron transfer from A$_1^-$ via F$_X$ to F$_A$ and F$_B$ was partially replaced by charge recombination in the pairs P700$^+$A$_1^-$ and P700$^+$F$_X^-$, and a kinetic heterogeneity of PS I became obvious below 150 K: in about 45% of the PS I complexes, forward electron transfer was blocked at the level of A$_1$, in about 20% at the level of F$_X$, and in only about 35% reduction of F$_A$ or F$_B$ was possible. Possible origins of this behaviour, including the idea that conformational substates are frozen below the liquid-to-glass transition of the medium (at around 180 K in the presence of 65% glycerol) were discussed extensively [7,57].

A$_1^-$ reoxidation in membrane fragments from *Chlamydomonas reinhardtii* as monitored by pulsed EPR was recently reported to occur with a time constant of 422 ns in a frozen suspension at 260 K (the EPR was recently reported to occur with a time constant of 422 ns in a frozen suspension at 260 K). By contrast, an intrinsic rate faster than (25 ns)$^{-1}$ for PS I from *Synechococcus* and (25 ns)$^{-1}$ for PS I from spinach [8]. By contrast, an intrinsic rate faster than (25 ns)$^{-1}$ for PS I from *Synechococcus* and (25 ns)$^{-1}$ for PS I from spinach [8].

An interesting case are the *menA* and *menB* mutants of *Synechocystis* sp. PCC 6803 that cannot synthesise phylloquinone, but recruit a foreign quinone Q, most likely plastoquinone-9, into the A$_1$ site [28,60,72]. PS I complexes isolated from these mutants showed efficient forward electron transfer to F$_A$/F$_B$. The reoxidation of Q$^-$ was biphasic with lifetimes of about 15 µs and 250 µs (minor phase) [60]. Based on an analysis of the electron transfer kinetics and on in vitro reduction potentials of quinones, it was concluded that electron transfer from Q$^-$ to F$_X$ is thermodynamically unfavourable in this mutant, but that overall transfer from Q$^-$ to F$_A$/F$_B$ is exergonic [60].

With respect to electron transfer between the iron–sulphur clusters F$_X$, F$_A$, and F$_B$, it has long been debated which of the two high electron densities in the PsAC subunit observed in the X-ray diffraction of PS I crystals corresponds to F$_A$ and which to F$_B$. Recent kinetic and spectroscopic data from different labs converge to the conclusion that the cluster located closer to F$_X$ is F$_A$, and that the distal cluster is F$_B$ [14,16,20,35,42,64,69], in line with a conclusion from the 2.5 Å crystal structure [17,30]. The electron transfer pathway should hence be F$_X$→F$_A$→F$_B$.

Concerning the kinetics of these two electron transfer steps (see [7] for a review of data up to 1996), it became clear from studies on ferredoxin reduction by PS I, involving F$_B$ [14,16,42,69] and occurring to a large extent in about 500 ns [16,38,61,62] that neither electron transfer from F$_X$ to F$_A$, nor from F$_A$ to F$_B$ can be slower than about 500 ns. In the framework of the A$_1$–F$_X$ equilibrium model (which is contested; see above), the intrinsic rate $k_2$ of electron transfer from F$_X$ to F$_A$ can be estimated from the measured biphasic reoxidation kinetics of A$_1^-$ (see above), yielding $k_2$≈80 ns$^{-1}$ for PS I from *Synechococcus* and $k_2$≈170 ns$^{-1}$ for PS I from spinach [8]. Based on the observation that the amplitude of a 25 ns phase of absorption changes attributed to a change of the membrane potential (electrochromic bandshift) was larger than expected for electron transfer from A$_1^-$ to F$_X$.

In principle, time resolved photovoltage measurements (see Introduction) should be able to monitor all transmembrane electron transfer steps, including that from F$_X$ to F$_A$. Experiments performed with a time resolution of 5–7 ns on PS I membranes from *Synechocystis* sp. PCC 6803 could be simulated by two phases: a fast instrument limited rise (attributed to the formation of the state P700$^+$A$_1^-$ [25]), and a slower monoexponential rise with $t_1/2$≈150 ns (attributed to overall electron transfer from A$_1^-$ via F$_X$ to F$_A$ and possibly to F$_B$) [14,37]. Removal of F$_A$ and F$_B$ by urea treatment decreased the relative amplitude of the latter phase more than two-fold, but

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1 A 7 ns phase as observed in transient absorption measurements of A$_1^-$ reoxidation in PS I from *Synechocystis* (see above) was not detected in these photovoltage measurements. This could be due to the fact that the expected electrogenicity for the establishment of the equilibrium (7 ns phase) is weak compared to the electrogenicity of the 150 ns phase, which is due to overall electron transfer from A$_1^-$ to F$_A$ or F$_B$. 

did not significantly alter its kinetics. It was concluded that the overall electron transfer from A\textsubscript{1} to F\textsubscript{A} or F\textsubscript{B} is limited by the step from A\textsubscript{1} to F\textsubscript{X}, and that the intrinsic rate of electron transfer from F\textsubscript{X} to F\textsubscript{A} must be faster than (50 ns\textsuperscript{-1}) [14,37]. A difficulty in these measurements was, that the directly measured signal is a convolution of the photovoltage kinetics generated by PS I with the ionic relaxation of the medium around the membrane fragments which had a characteristic time of only about 50 ns. Hence, to obtain the kinetics cited above, a convolution procedure had to be applied, which relies on the assumption that the ionic relaxation is homogeneous. In an alternative approach, 66% (v/v) glycerol was added which slowed the ionic relaxation down to about 10 \mu s so that it should have only a minor effect on photovoltage kinetics in the sub-microsecond range. Surprisingly, the second phase of the photovoltage rise was found to be slower (t\textsubscript{1/2} \approx 350 ns) than in the absence of glycerol (t\textsubscript{1/2} \approx 150 ns), although the reoxidation of A\textsubscript{1\textsuperscript{-}} as monitored by transient absorption was not affected by glycerol [37]. Unless there is a technical problem with the measuring cell in the presence of glycerol, this result appears to indicate that electron transfer beyond F\textsubscript{X}, which makes the main contribution to the electro-genic signal, is slowed down by glycerol [37]. It is of note that the amplitude of the 150 ns phase of the photovoltage rise was only about 35–40% of the total amplitude, whereas the transmembrane distance between A\textsubscript{1} and F\textsubscript{A} is about 50% of the distance between P700 and F\textsubscript{X} (see Fig. 1b). This deviation may be due to the fact that the 7 ns phase of A\textsubscript{1} reoxidation was not resolved in the photovoltage measurements and/or to a higher effective dielectric constant of the environment of the electron transfer path from A\textsubscript{1} to F\textsubscript{A} compared to that of the path from P700 to A\textsubscript{1} (note that the photovoltage amplitudes due to a charge displacement are inversely related to the dielectric constant).

With respect to electron transfer between F\textsubscript{A} and F\textsubscript{B}, it is of note that thermodynamically slightly favourable (\text{\text{\text{-}}}\Delta G^\circ \approx 0–80 \text{ meV}) transfer between the two [4Fe-4S] clusters of ferredoxins similar in structure to PsaC was found by NMR to occur in about 300 ns [34]. Hence, a time constant \leq 500 ns for electron transfer from F\textsubscript{X} to F\textsubscript{B} (as concluded from the reduction kinetics of ferredoxin; see above) seems reasonable, provided that the reduction potential of F\textsubscript{X} is lower or close to that of F\textsubscript{B}, at least when ferredoxin is bound. For references and a discussion on the energetics of electron transfer between the three iron–sulphur clusters of PS I, see [7].

2.3. Charge recombination reactions

Charge recombination between P700\textsuperscript{+} and the reduced form of any one of the electron acceptors can be observed when forward electron transfer to the subsequent acceptor is blocked, e.g., by prereduction or extraction of the latter (see [7] for a review of data up to 1996). Typical recombination times are some tens of nanoseconds for P700\textsuperscript{+}A\textsubscript{1\textsuperscript{-}}, 10–250 \mu s for P700\textsuperscript{+}A\textsubscript{1\textsuperscript{-}}, about 1 ms for P700\textsuperscript{+}F\textsubscript{X}, and some tens of milliseconds for P700\textsuperscript{+}F\textsubscript{A} and P700\textsuperscript{+}F\textsubscript{B\textsuperscript{\text{-}}}\. Recombination kinetics can be affected by the reduction state of the electron acceptors not directly involved in the recombination. Thus, P700\textsuperscript{+}A\textsubscript{1\textsuperscript{-}} recombination was found to be accelerated to t\textsubscript{1/2} \approx 250 ns when all three iron–sulphur clusters were reduced; this acceleration was attributed to an up-shift of the free energy of P700\textsuperscript{+}A\textsubscript{1\textsuperscript{-}} due to the electrostatic interaction between A\textsubscript{1\textsuperscript{-}} and F\textsubscript{X}, F\textsubscript{A\textsuperscript{-}}, and F\textsubscript{B\textsuperscript{-}} that favours a fast recombination of P700\textsuperscript{+}A\textsubscript{1\textsuperscript{-}} via thermally activated repopulation of the primary pair P700\textsuperscript{+}A\textsubscript{0\textsuperscript{-}} [50].

New data became available for the charge recombination reactions involving the terminal acceptors F\textsubscript{A} and F\textsubscript{B}. In intact PS I from S. elongatus, recombination between P700\textsuperscript{+} and a single electron equilibrated between F\textsubscript{A} and F\textsubscript{B} (the equilibrium ratio of F\textsubscript{A} and F\textsubscript{B} is a matter of debate [7]) occurs with t\textsubscript{1/2} = 80 ms at room temperature; it has an activation energy of (220 \pm 10) \text{meV} (measured between 275 and 306 K) which is consistent with a recombination pathway via thermally activated repopulation of the pair P700\textsuperscript{+}A\textsubscript{1\textsuperscript{-}} [31]. With two electrons on the F\textsubscript{A}\textendash{}F\textsubscript{B\textsuperscript{\text{-}}} complex, the recombination was accelerated to t\textsubscript{1/2} \approx 10 ms at room temperature. A redox titration of the recombination kinetics yielded midpoint potentials of (\approx 440 \pm 10) mV and (\approx 465 \pm 10) mV for the first and second reduction of the F\textsubscript{A}\textendash{}F\textsubscript{B\textsuperscript{\text{-}}} complex, respectively [31]. For unknown reasons, these potentials are considerably higher than those established previously by titrations with detection of the reduction state of the iron–sulphur clusters by low-temperature EPR (approximately \text{\text{-}540 and \text{-}590 \text{mV, re-}}
spectively; reviewed in [7]). A difference of about 50 mV between these two potentials would be consistent with the observed ratio of about eight between the recombination rates (see above), provided that both recombinations proceed by thermally activated repopulation of the same state (presumably (P700$^+$A$_1^-$); see above).

The recombination between P700$^+$ and F$_A^-$ after destruction of F$_B^-$ by treatment of cyanobacterial PS I with HgCl$_2$ was recently studied by two groups. The reported recombination rates are (90 ms)$^{-1}$ for a PS I preparation from Synechocystis sp. PCC 6803 [14] and (40.7 ms)$^{-1}$ for a PS I preparation from Synechococcus sp. PCC 6301 [64].

3. Some controversial points

3.1. Are both electron transfer branches active?

As outlined in Section 2.2, Joliot and Joliot [29] suggested that electron transfer from excited P700 to FX proceeds with similar probabilities along the two branches shown in Fig. 1, but that the last steps (Q$_X$A$^-$ to FX and Q$_X$B$^-$ to FX) have different rates. This model provides a simple explanation for the observed bi-phasic reoxidation ($t_{1/2}$ $\approx$ 10–20 ns and 200 ns) of the reduced secondary acceptor A$_1$. The authors considered the previously suggested A$_1$–FX equilibrium model (assuming similar free energy levels for the states P700$^+$A$_1^-$ and P700$^+$FX in a single active branch; see Section 2.2) as unlikely as they could not observe a significant change of the reoxidation kinetics of A$_1$ upon collapse of the membrane potential [29]. Taking into account the different levels of A$_1$ and FX within the membrane (see Fig. 1), it was estimated [29] that the collapse of the membrane potential should have increased the driving force of electron transfer from A$_1^-$ to FX by about 13 meV, and hence the equilibrium constant between the states P700$^+$A$_1^-$ and P700$^+$FX by a factor of 1.7. In the framework of the A$_1$–FX equilibrium model, this effect should have increased the amplitude ratio of the fast and slow phase of A$_1^-$ reoxidation by a similar factor, which should have been easily detectable. We would like to point out, however, that also in the two-branch model the kinetics of A$_1^-$ reoxidation should be affected by the membrane potential. For a 13 meV increase in driving force from the phylloquinones to FX, and assuming that the reorganisation energy is large compared to the driving force, Marcus theory [43,44] predicts an acceleration by a factor of 1.3 for both phases of A$_1^-$ reoxidation which was not reported by Joliot and Joliot [29]. Hence we feel that the lack of a significant membrane potential effect does not strongly favour the two-branch model over the A$_1$–FX equilibrium model. Nevertheless, we find the two-branch model very interesting as it may account for some hitherto unexplained features of PS I. In the following, we discuss some observations and reflections, which are relevant in this context.

1. In a PS I core preparation devoid of the terminal electron acceptors FX, FA, and FB, charge recombination between P700$^+$ and A$_1^-$ was biphasic with halftimes of about 10 $\mu$s and 110 $\mu$s [9]. A straightforward explanation would be that both branches are used and that the recombination rates of Q$_X$A$^-$ and Q$_X$B$^-$ with P700$^+$ are different. As the absorption difference spectra of the two phases of forward electron transfer from A$_1^-$ are significantly different between 450 and 480 nm [8,29,63], one would expect that the two recombination phases show a similar difference. However, at room temperature, such a difference was not observed [9]. This spectral analysis does hence not support the idea that the biphasic forward electron transfer and the biphasic recombination are in the same way related to the use of the two electron transfer branches. This is only a weak argument against the two-branch model, however, as the signal to noise ratio of the 110 $\mu$s recombination phase was rather poor. Interestingly, at cryogenic temperature (10 K) where the recombination between A$_1^-$ and P700$^+$ was found to be biphasic as well (halftimes of 15 $\mu$s and 150 $\mu$s), the spectra of the two phases were different [9]. Their difference correlates with the difference between the spectra of the two phases of forward electron transfer in a way that would be consistent with the view that the fast phase of forward electron transfer and the slow phase of recombination arise from one branch, and the slow forward and the fast recombination phases from the other branch.
2. Studies of secondary electron transfer by transient EPR spectroscopy at a time resolution of about 50 ns [5,67] could resolve only the slower one \( (t_{1/2} \approx 150 \text{ ns}) \) of the two phases of \( \text{A}_1^- \) reoxidation observed by optical spectroscopy. The spin-polarised EPR spectrum detected at the beginning of the 150 ns phase was similar (although not identical) to the spectrum observed in a PS I core preparation devoid of \( \text{F}_X, \text{F}_A, \) and \( \text{F}_B \) [67], and was attributed to the spin-correlated, weakly coupled radical pair \( \text{P}_700^\ast \text{A}_1^- \) created by fast forward electron transfer from a singlet precursor [5,67]. Within the framework of the \( \text{A}_1^- \text{F}_X \) equilibrium model, the slower phase of the reoxidation of \( \text{A}_1^- \) would be preceded by forth and back electron transfer between \( \text{A}_1 \) and \( \text{F}_X \) (see Section 3.2). The latter process is expected to diminish the spin polarisation of the excess electron on the acceptor side because \( \text{F}_X \) has a very fast spin relaxation rate [55]. Although we are not aware of any simulation of this scenario, it seems likely, that the transient residence of the excess electron on \( \text{F}_X \) would strongly modify the EPR spectrum of the pair \( \text{P}_700^\ast \text{A}_1^- \), in contrast to the observed similarity with the \( \text{P}_700^\ast \text{A}_1^- \) spectrum in PS I devoid of \( \text{F}_X, \text{F}_A, \) and \( \text{F}_B \). This problem is overcome in the two-branch model, because both \( \text{A}_1^- \) reoxidation phases are due to forward electron transfer from the initially created pairs \( \text{P}_700^\ast \text{Q}_X^- \text{A}^- \) and \( \text{P}_700^\ast \text{Q}_A^- \text{B}^- \).

3. A single active branch was strongly favoured by reports that one out of the two phylloquinones present in intact PS I could be extracted with hexane, and that PS I retaining only one phylloquinone was fully active in forward electron transfer [4,41]. If both branches were active with similar yields in untreated PS I, removal of one phylloquinone should block forward electron transfer at the level of \( \text{A}_0^- \) and give rise to charge recombination in the primary pair \( \text{P}_700^\ast \text{A}_0^- \) in about half of the centres, in contrast to observation [4]. It is of note, however, that in a more recent study, PS I extracted with hexane according to the same protocol retained approximately two phylloquinones per \( \text{P}_700 \) [59], questioning the above evidence for a single active branch.

4. EPR studies have established that a phylloquinone anion radical can be trapped (in addition to re-duced \( \text{F}_A, \text{F}_B, \) and \( \text{F}_X \)) by illuminating PS I at 200–230 K under reducing conditions ([24,70], and references cited in [7]). According to a recent spin-quantification, not more than one phylloquinone radical could be accumulated in that way [40,70]; it was concluded that only one of the two phylloquinones is redox active [70]. In previous work, however, a second phylloquinone radical could be trapped under harsher conditions than those necessary to trap the first one [24]. The latter observation may be explained in the framework of the two-branch model by assuming that the two phylloquinone anion radicals recombine with \( \text{P}_700^\ast \) at rather different rates (see also (1)); as trapping of a reduced acceptor requires rereduction of \( \text{P}_700^\ast \) by an external reductant prior to charge recombination, the phylloquinone with the faster recombination rate would be more difficult to be trapped in the reduced state.

5. The \( \text{A}_1^- \text{F}_X \) equilibrium model was based, among others, on the observation that the slower phase \( (t_{1/2} \approx 150 \text{ ns}) \) of forward electron transfer from \( \text{A}_1^- \) in spinach PS I was replaced by a 250 \( \mu \text{s} \) recombination with \( \text{P}_700^\ast \) upon prereduction of \( \text{F}_A \) and \( \text{F}_B \), while the fast phase \( (t_{1/2} \approx 25 \text{ ns}) \) of forward electron transfer was conserved [63]. This behaviour is expected if the 150 ns phase reflects electron transfer from the equilibrium between \( \text{A}_1^- \) and \( \text{F}_X \) to \( \text{F}_A/\text{F}_B \). To explain the same behaviour in the framework of the two-branch model, one would have to assume that prereduction of \( \text{F}_A \) and \( \text{F}_B \) blocks electron transfer to \( \text{F}_X \) from one of the phylloquinones, but not from the other one. A possible mechanism would be the Coulomb interaction between the transferred electrons and \( \text{F}_A/\text{F}_B^{-} \) that tends to decrease the driving force of electron transfer from the phylloquinones to \( \text{F}_X \). Due to the asymmetric positions of \( \text{F}_A \) and \( \text{F}_B \) (see Fig. 1), this effect is most likely more pronounced for the phylloquinone designated \( \text{Q}_A^- \text{A}^- \) than for \( \text{Q}_B^- \text{B}^- \). In addition, due to differences in the surroundings of the two phylloquinones, the driving forces may be different already under normal conditions (\( \text{F}_A \) and \( \text{F}_B \) oxidised), so that the Coulomb interaction with \( \text{F}_A^-/\text{F}_B^- \) might render electron transfer from the phylloquinones to \( \text{F}_X \) uphill in one branch and leave it still downhill in the other branch.
6. Within the framework of the $A_1^-$–$F_X$ equilibrium model, variations of the relative amplitudes of the two phases of $A_1^-$ reoxidation between different PS I preparations could be explained by slight variations of the free energy difference between the states $P700^+A_1^-$ and $P700^+F_X^-$. If instead the two phases were due to the two electron transfer branches, amplitude variations should be due to variations in the relative yield of charge separation in the two branches.

Concluding this discussion, none of the arguments against the two-branch model appears to be ‘waterproof’. The same is true for the arguments against the two-branch model.

3.2. The reduction potential of the phylloquinone $A_1$

Since the discovery of phylloquinone as secondary electron acceptor $A_1$, it has been recognised that its one-electron reduction potential should be at least as low as about $-700$ mV versus normal hydrogen electrode (NHE) in order to allow for efficient forward electron transfer to the terminal acceptors $F_A$ and $F_B$ ([7] and references therein). Two important points are still subject to controversial discussions:

- The reduction potential of $A_1$ relative to that of $F_X$, or more precisely, the free energy of the state $P700^+A_1^-$ relative to that of the state $P700^+F_X^-$ (see Eq. 1 in the legend of Fig. 2).
- The reason(s) why phylloquinone in PS I has a much lower reduction potential than the structurally similar menaquinone-9 as secondary electron acceptor $Q_A$ in the bacterial reaction centre from *Rhodopseudomonas viridis* ($E_m \approx -150$ mV [52]).

Different approaches have been used to estimate the reduction potential of $A_1$ or the free energy of the state $P700^+A_1^-$:

1. Interpreting the two phases of $A_1^-$ reoxidation in the framework of the $A_1^-$–$F_X$ equilibrium model (see Section 2.2), it was estimated that the state $P700^+A_1^-$ is about 40 meV above $P700^+F_X^-$ in the PS I-β preparation from spinach [63,8], and about 20 meV below $P700^+F_X^-$ in PS I from *Synechocystis* sp. PCC 6803 [8]. Obviously, these estimations are not valid if it will be confirmed that the two kinetic phases are due to the two electron transfer branches (see Section 3.1).

2. From a study of the efficiency of forward electron transfer to $F_A/F_B$ after reconstitution of diethyl ether extracted PS I with quinones with different in vitro reduction potentials, the potential of native phylloquinone in the $A_1$ site was estimated to be 50–80 mV more negative than that of $F_X$ [26]. As pointed out previously [7], the kinetic modeling of the data was based on a very slow rate of electron transfer from $F_X^-$ to $F_A/F_B$ ($4.6 \times 10^3$ s$^{-1}$) which is inconsistent with data on intact PS I (see Section 2.2).

3. Assuming that charge recombination of the pair $P700^+(F_A/F_B)^-$ ($t_{1/2} \approx 50$ ms at room temperature) proceeds via thermal repopulation of the pair $P700^+A_1^-$ and direct charge recombination in the latter pair with $t_{1/2} \approx 150$ μs, a free energy difference of 150 mV between $P700^+A_1^-$ and $P700^+(F_A/F_B)^-$ is obtained from a simple equilibrium consideration [7]. Unfortunately, the free energy difference between $P700^+F_X^-$ and $P700^+(F_A/F_B)^-$ is not well established. A value of 150–190 meV can be estimated from redox titrations of the iron–sulphur clusters in intact PS I (see [7] for references). This would imply that $P700^+F_X^-$ lies slightly above $P700^+A_1^-$ in free energy, but the free energy of $P700^+F_X^-$ may be overestimated because $F_A$ and $F_B$ were prereduced in the redox titration of $F_X^-$.

A lower free energy of $P700^+F_X^-$ (about 100 meV above $P700^+F_X^-$) was estimated from a comparison of the recombination time constants for $P700^+F_X^-$ in PS I devoid
of $F_A$ and $F_B$ ($\tau \approx 0.85$ ms) and $P700^+F_A$ in PS I devoid of $F_B$ ($\tau \approx 40.7$ ms) [64]. An uncertainty in this estimate is that the free energy levels and the recombination rates in these modified PS I complexes may be different from those in intact PS I (see also [60]). The same objection can be raised against an estimation of the free energy change for electron transfer from $A_1^-$ to $F_X$ ($-28$ meV) based on an analysis [7] of kinetic and spectroscopic data in PS I devoid of $F_A$ and $F_B$. As a more general problem we would like to mention that these approaches probe the energetics on a time scale that is much slower than forward electron transfer. Energy minimising conformational changes of the protein in response to changes of the charges on the cofactors are expected to occur on multiple time scales. Hence, the free energy levels of the different charge separated states during forward electron transfer may deviate significantly from those estimated from slow recombination reactions or redox titrations.

4. In the study suggesting that the two phases of $A_1^-$ reoxidation are due to the participation of two phylloquinones ($Q_K$-A and $Q_K$-B), the absence of an effect of the membrane potential on the relative amplitudes of the two phases was taken as evidence that electron transfer from each of the two phylloquinones to $F_X$ is much faster than the reverse reaction [29], i.e., $P700^+Q_K$-$A^-\ \text{and} \ P700^+Q_K$-$B^-\ \text{are well above} P700^+F_X^-$ in free energy. It appears to us, that the amplitude ratio of the two phases in the two-branch model should depend on the yields of primary charge separation in the two branches and is unlikely to be affected by the membrane potential, whatever the reaction free energy $\Delta G^0$ for electron transfer from the phylloquinones to $F_X$. Hence, accepting the two-branch model, the observation cited above cannot be used to estimate $\Delta G^0$.

Despite these uncertainties about the reduction potential of $A_1$, there is no doubt that it must be well below that of the terminal acceptor complex $F_A/F_B$ ($E_m \approx -530$ mV versus NHE), and is most likely close to or below that of $F_X$ ($E_m \approx -700$ mV). To explain such a low potential of phylloquinone, it was suggested that the environment of $A_1$ is aprotic and may provide electron repulsive groups [10,58]. Refer-

ring to data by Prince and coworkers [51], two papers [27,60] assumed that the half-reduction potential of phylloquinone (being the same as that of menaquinone) in the aprotic solvent dimethylformamide (DMF) is $-465$ mV versus NHE; the much lower potential of phylloquinone in the $A_1$ site was attributed to an extremely low acceptor property of the $A_1$ site. It appears to us that the cited potential of phylloquinone and menaquinone in DMF is not consistent with the original data [51], where the potentials were measured versus a standard calomel electrode (SCE); listed potentials for menaquinones range from $-746$ mV (menaquinone-0) to $-705$ mV (menaquinone-7) [51]. Apparently, the value of $-465$ mV versus NHE in [27,60] was obtained by adding 244 mV to the potential versus SCE measured in DMF. This conversion would be valid in aqueous solution, but may fail in aprotic solvents because of the junction potential between the aqueous reference electrode and the bulk solution [53]. Instead, one may use the half-reduction potential of ferrocene as a reference that was measured to be $+524$ mV versus SCE under the conditions of the quinone measurements [51]. The formal potential of ferrocene is assumed to be independent of the solvent [33], and was reported to be $+400$ or $+425$ mV versus NHE in water [33,71]). In a first approximation, one may therefore assume that potentials referred to NHE are about 100 mV lower than those versus SCE measured in [51]. This approach would yield potentials of about $-850$ to $-800$ mV versus NHE for different menaquinones in DMF, well low enough for what is required for phylloquinone in the $A_1$ site.

The 2.5-Å structure of the PS I complex from $S.\ elongatus$ should allow to analyse the differences in the quinone environments that might explain the strikingly different reduction potentials of menaquinone-9 in the $Q_A$ site of the reaction centre from $Rp.\ viridis$ ($E_m \approx -150$ mV [52]) and phylloquinone in the $A_1$ site of PS I ($E_m \approx -700$ mV). We would like to mention a few points that may be relevant in this respect:

- Amongst the main factors governing the redox potential of cofactors in situ are differential solvation interactions of the oxidised and reduced form with the protein environment. A favourable interaction of the protein with the negatively charged, reduced...
species can strongly increase the redox potential. In the case of the reaction centre of *R. viridis*, QA is situated in a highly polar environment with many polar groups and internal water molecules. Interaction with these groups will decrease the desolvation penalty for the (negatively) charged quinone and hence increase its reduction potential [36]. In PS I, a highly nonpolar environment could preserve the low reduction potential of A1. It is worth mentioning in this context that, in contrast to the quinone acceptor complex of the purple bacterial reaction centres, no proton transfer has to be coupled to electron transfer in PS I.

- The positive charge on the non-heme iron on the acceptor side of the purple bacterial reaction centre raises the potential in the quinone region [36] and may contribute to the higher reduction potential of QA compared to A1. Similarly, the presence of two negative charges on FX (including the cysteine ligands) may decrease the potential of A1.

- For both phylloquinones in PS I, only one of the carbonyls accepts a hydrogen bond (from a backbone NH group of a leucine residue) [17,30]. In *R. viridis*, there is an additional hydrogen bond, from a histidine residue to the second carbonyl of QA, that should stabilise the semiquinone anion state and hence increase the reduction potential of QA compared to that of A1 in PS I.

- As pointed out by Iwaki and Itoh [26], π–π interactions between a quinone ring and an aromatic residue should destabilise the reduced form of the quinone. Such an effect is probably exerted by TrpA679 and TrpB677 situated at plane-to-plane distances of 3.0–3.5 Å from the phylloquinones in PS I [17,30]. In *R. viridis*, π–π stacking between QA and TrpM250 is much weaker (plane-to-plane distance, 4.7 Å [47]).

### 3.3. How do reported electron transfer rates compare with the cofactor distances?

The relation between electron transfer rates and distances between the cofactors in PS I has been discussed previously ([7,32] and references therein) on the basis of the 4.5-Å and the 4-Å structure of PS I. Most of these discussions used a simple semi-empirical relation for the rate $k_{et}$ (in s$^{-1}$) of intraprotein electron transfer proposed by Moser and Dutton [48]:

$$\log k_{et} = 15 - 0.6 R - 3.1(\Delta G^0 + \lambda)^2/\lambda$$  \hspace{1cm} (2)

where $R$ is the shortest edge-to-edge distance (in Å) between donor and acceptor, $\Delta G^0$ the standard reaction free energy (in eV), and $\lambda$ the reorganisation energy (in eV). The ‘optimal’ (maximal) rate for a given distance is obtained for the special case $-\Delta G^0 = \lambda$. Except for primary charge separation (see (1) below) and charge recombination in the pair P700$^+$A1$^-$ (see (2) below), no obvious violation of Eq. 2 has been reported. A more refined analysis will be possible on the basis of the 2.5 Å structure of PS I [17,30]. At the present stage (the co-ordinates of the 2.5 Å structure were not yet released when we finished this manuscript), we restrict ourselves to a few comments:

1. Primary charge separation (electron transfer from excited P700 to A0 over an edge-to-edge distance of about 13 Å (according to the 4-Å structure [32])) is much faster ($k_{et}$ in the order of 10$^{12}$ s$^{-1}$; see Section 3.1) than the optimal rate of about 10$^7$ s$^{-1}$ calculated from Eq. 2 for $-\Delta G^0 = \lambda$. As pointed out previously [7,32], this discrepancy strongly suggests that an ‘accessory’ chlorophyll eC-B$_2$ and/or eC-A$_2$ is involved in this electron transfer.

2. The rate of charge recombination in the pair P700$^+$A1$^-$ (presumably about 4×10$^3$ s$^{-1}$ in intact PS I [7] and up to 10$^6$ s$^{-1}$ in PS I devoid of FX, FA, and FB [9]) exceeds the optimal rate of 5×10$^2$ s$^{-1}$ calculated from Eq. 2 with $-\Delta G^0 = \lambda$ and $R = 20.5$ Å [32]. One may consider two possibilities: (a) The electronic coupling between A1 and P700$^+$ is enhanced by the chlorophylls located in between (see Fig. 1), including the possibility of some charge delocalisation between A1$^-$ and A0 or between P700$^+$ and an ‘accessory’ chlorophyll eC-B$_2$ and/or eC-A$_2$. (b) The protein structure between P700 and A1 provides particularly favourable electron transfer pathways (see [3] for an approach to establish and to treat such pathways).

3. According to the 2.5-Å structure [17,30], the distance between the centre of FX and the midpoint of the two oxygen atoms of phylloquinone QK-A and of phylloquinone QK-B are 14.2 and 14.1 Å,
respectively. The benzylic rings of both phylloquinones are oriented towards FX, thus minimising the edge-to-edge distances to FX, as suggested previously from an electron transfer study [57]. Taking into account also the orientation of FX, the shortest edge-to-edge distance to the phylloquinones should be in the order of 8 Å for both branches (for FX, we considered the sulphur atoms of the cysteine ligands as edge atoms, because they receive a considerable electron density upon reduction of [4Fe^4S] clusters [7,49]). Inserting this distance in Eq. 2, the faster phase (k^W_4 \approx 10^7 s^{-1}) of A^\circ reoxidation (see Section 2.2) can be obtained with a small driving force and an unspectacular reorganisation energy (e.g., ΔG^0 = -0.1 eV and λ = 1 eV, or ΔG^0 = 0 and λ = 0.8 eV). If the slower phase of A^\circ reoxidation (k^W_3 \approx 10^6 s^{-1}) reflects electron transfer from the other phylloquinone to FX (see Section 3.1), the slower rate may be due to a higher reorganisation energy and/or a lower driving force, rather than to a larger distance between the cofactors.

4. Note added in proof

After submission of the manuscript, several articles appeared which are relevant to the question whether two electron transfer branches are active in PS I. Guergova-Kuras et al. [22] studied the effect of mutations of tryptophan residues supposed to be close to the phylloquinones on the in vivo kinetics of A^\circ reoxidation in whole cells of C. reinhardtii. While the mutation PsaA-W693F slowed specifically the slower phase (from t_{1/2} \approx 150 ns to \approx 400 ns), the mutation PsaB-W673F slowed specifically the faster phase (from t_{1/2} \approx 13 ns to \approx 70 ns). The double mutation PsaA-W693F/PsaB-W673F resulted in a slowdown of both phases. As pointed out in [22], these effects could be by far most easily explained with the assumption that the faster phase of A^\circ reoxidation reflects electron transfer in the PsaB-branch, and the slower one in the PsaA-branch, and that mutation of either tryptophan slows down reoxidation of the close by phylloquinone.

Hastings and Sivakumar [23] reported a Fourier transform infrared absorption spectrum associated with photoaccumulation of A^\circ in samples from Synechocystis sp. PCC 6803 devoid of Fe-S centres FA, FB, and FX. The complexity of the spectra in various spectral regions was taken as indication that two structurally distinct phylloquinones were photoaccumulated, which could correspond to the phylloquinones on the PsaA and PsaB branches.

Burton and coworkers [54] investigated the environment of the phylloquinone acceptors by site-directed mutagenesis of two tryptophan residues (W693 and W702) in the PsaA subunit of PS I from C. reinhardtii. Replacement of W693 with either histidine or leucine altered the electronic structure of the photoaccumulated A^\circ radical and slowed forward electron transfer (by a factor of \approx 3) as measured by the decay of the electron spin-polarised signal from the radical pair P700^+A^\circ at 260 K (the same mutations of W702 had no effect). The authors pointed out that the limited time resolution of this technique only allows to detect the slow phase of A^\circ reoxidation, which, in agreement with the results of Guergova-Kuras et al. [22], is proposed to occur in the PsaA-branch.

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References


