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Molecular basis for turnover inefficiencies (misses) during water oxidation in photosystem II†

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Photosynthesis stores solar light as chemical energy and efficiency of this process is highly important. The electrons required for CO₂ reduction are extracted from water in a reaction driven by light-induced charge separations in the Photosystem II reaction center and catalyzed by the CaMn₄O₅-cluster. This cyclic process involves five redox intermediates known as the S₀–S₄ states. In this study, we quantify the flash-induced turnover efficiency of each S state by electron paramagnetic resonance spectroscopy. Measurements were performed in photosystem II membrane preparations from spinach in the presence of an exogenous electron acceptor at selected temperatures between –10 °C and +20 °C and at flash frequencies of 1.25, 5 and 10 Hz. The results show that at optimal conditions the turnover efficiencies are limited by reactions occurring in the water oxidizing complex, allowing the extraction of their S state dependence and correlating low efficiencies to structural changes and chemical events during the reaction cycle. At temperatures 10 °C and below, the highest efficiency (*i.e.* lowest miss parameter) was found for the S₁ → S₂ transition, while the S₂ → S₃ transition was least efficient (highest miss parameter) over the whole temperature range. These electron paramagnetic resonance results were confirmed by measurements of flash-induced oxygen release patterns in thylakoid membranes and are explained on the basis of S state dependent structural changes at the CaMn₄O₅-cluster that were determined recently by femtosecond X-ray crystallography. Thereby, possible “molecular errors” connected to the e[–] transfer, H⁺ transfer, H₂O binding and O₂ release are identified.

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Introduction

Oxygen is an essential part of our atmosphere and a major component of cellular respiration. It appeared about 2.7 billion years ago as a by-product of oxygenic photosynthesis evolved in cyanobacteria.¹ Now oxygenic photosynthesis is present in all three kingdoms which possess photosynthetic organisms such as cyanobacteria, algae and plants. Together, these organisms provide the chemical energy for essentially all life in the biosphere. Oxygenic photosynthesis is driven by two photosystems, PSII and PSI, that are both localized in the thylakoid membrane and work in sequence to utilize sun light and to provide reducing equivalents for the CO₂ fixation.

Photosystem II (PSII) is a multisubunit pigment–protein complex that forms the starting point of the photosynthetic electron flow by catalysing the light-induced oxidation of water

and the reduction of plastoquinone (PQ)^{2–5} (Fig. 1). The first step of the photosynthetic water splitting is light excitation of the primary electron donor chlorophylls (Chl), P₆₈₀. After the initial charge separation between P₆₈₀ and pheophytin, e[–] is transferred *via* the two quinone acceptors, Q_A and Q_B, to the PQ pool to be utilized in subsequent reactions in the thylakoid

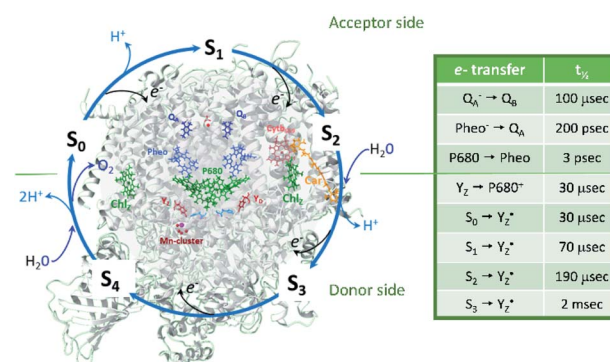


Fig. 1 Protein and cofactor organization of PSII based on 6DHE pdb file, S₁-enriched state,⁵² S cycle and sequence of events involving e[–], H⁺ and H₂O leading to oxygen evolution in the WOC. The half-times of the individual electron transfer steps are indicated in the table and are reviewed in ref. 2–4, 6, 7 and 9.

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membrane. Q_B serves as an exchangeable carrier and is reduced by two electrons (e^-) and two protons (H^+) before exchange with another plastoquinone from the PQ pool. These electron transfer reactions are usually referred to as the acceptor side reactions in PSII (Fig. 1).^{2-4,6-8}

P_{680}^+ is a highly oxidizing species ($E_m = +1.35$ V) and extracts an electron from the nearby redox-active tyrosine residue, Y_Z . Y_Z together with the $CaMn_4O_5$ -cluster and its water and protein environment compose the water oxidizing complex (WOC), the catalytic site where oxidation of water occurs. Y_D subsequently oxidizes the $CaMn_4O_5$ -cluster with bound water molecules. The reactions in the WOC are referred to as the donor side reactions (Fig. 1).^{4,9-11}

The quest to solve the photosynthetic water oxidation mechanism started with the famous Joliot's experiment where O_2 release, for the first time, was studied under a train of short (10 μ s) light flashes.¹² When applied to the dark-adapted algal suspension, the first O_2 release peak appeared on the 3rd flash with the next maxima appearing after every 4 flashes until these oscillations were damped and oxygen yields became equal on each flash. These flash-induced oxygen oscillation patterns (FIOPs) were explained by Kok and coworkers in a model, which postulates that during water oxidation, the WOC cycles through five intermediate redox states, collectively called the S states, labelled S_0 – S_4 (Fig. 1).^{13,14} S_0 is the most reduced state while S_1 , S_2 and S_3 represent sequentially higher oxidation states in the WOC. S_1 is the dominating state in the dark while the S_2 and S_3 states are metastable and decay back to the S_1 state in a few minutes at room temperature.¹⁵⁻¹⁸ In addition, the S_0 state can be slowly (in tens of minutes) oxidized to the S_1 state by Y_D , the second redox active tyrosine in PSII.^{19,20} O_2 is released during the $S_3 \rightarrow [S_4] \rightarrow S_0$ transition, where S_4 is a transient state (Fig. 1).^{4,9} Thus, four consecutive charge separations and accompanying electron transfer events are necessary in PSII to oxidize two water molecules to molecular oxygen and four protons (Fig. 1), which accounts for the period of four oscillations in FIOPs.^{9,13,14}

In Kok's S state cycle model, the dampening of the FIOP oscillations with the increasing flash number is explained by "double hits" and "misses". The double hit parameter accounts for centers that perform two consecutive charge separations and S state transitions as a result of a single flash excitation.^{13,14} Double hits are produced by the long flash tails from Xe flash-lamps and broad LEDs light pulses (tens to hundreds of μ s long) and can be eliminated by using short nanosecond laser flashes which produce only a single charge separation in PSII.²¹ The *miss parameter* is the most important factor in understanding the S state cycle advancement and dampening of the FIOPs. It represents a probability of the WOC of not advancing to the next S state when the PSII center is exposed to a single flash, and is connected to the turnover efficiency by the formula: $\text{miss} = 1 - \text{turnover efficiency}$. Misses are routinely used in the analysis of the S state cycle, mostly in the analysis of FIOPs, where they were first introduced.^{13,14,17,18,22-29} However, this concept has also been used in the analysis of S state cycle intermediates studied by almost any technique, such as variable and delayed fluorescence,^{25,30-36} transient optical,³⁷ EPR,³⁸⁻⁴² FTIR⁴³⁻⁴⁵ and EXAFS spectroscopies.⁴⁶⁻⁵⁰ These analyses were done on many types of

photosynthetic species with different degree of purification ranging from intact leaves and cells to PSII core complexes.

The first crystal structure of dark adapted PSII from thermophilic cyanobacteria has been solved almost two decades ago and the highest resolution available now is at 1.9 Å.⁵¹ Recently, Kok's model was used in analysis of the crystal structures obtained from the advanced S states of the cycle by serial femto-second X-ray crystallography.⁵² In this study, changes in the WOC structure were superimposed with the S states of the full S cycle for the first time.

Under saturating light excitation, the origin of misses in PSII could be acceptor or donor side related. A limitation on the acceptor side electron transfer arises from blocked or slow oxidation of Q_A^- by Q_B (or Q_B^-). The presence of Q_A^- at the time of light excitation results in the failure of charge separation. A small contribution to misses may also arise under such conditions from a charge recombination between Q_A^- and P_{680}^+ that occurs in the 100 microseconds time range before P_{680}^+ could be reduced by Y_Z .^{53,54} More interesting and mechanistically more important are the donor side induced misses, that originate from the molecular chemistry of the S state transitions. These "actual misses" are in the focus of our present study.

In most studies, the miss parameter was obtained by measurements of only one probe, such as oxygen release during the $S_3 \rightarrow [S_4] \rightarrow S_0$ transition^{17,18,22-28} or the S_2 multiline EPR signal during the $S_1 \rightarrow S_2$ transition.^{19,39,41,42} This type of analysis, however, has the internal limitation that it allows only determination of the average miss parameter for all S state transitions per cycle. Determination of the misses specific to the individual S state transition in this case is not trivial and requires additional measurements of the S state decay times, estimation of contribution from the secondary electron donors in PSII such as Y_D and Cytochrome b_{559} and extensive modelling.^{29,55-58}

In our previous studies, we developed an alternative approach where the individual misses were determined from the exact distribution of S states in the PSII sample after each laser flash advancement.^{21,40} For this purpose we used well-known, S state-specific electron paramagnetic resonance (EPR) probes such as split S_1 , S_3 and S_0 signals and the S_2 state multiline signal. We found that the misses during the water oxidation process in PSII are S state-dependent.⁴⁰ Here we extend the previous study and determined the individual misses over a wide temperature range and at the different frequencies of the advancing (turnover) laser flashes. We also compare our EPR data to FIOPs data obtained under similar conditions and conclude the analysis by relating the origin of the S state dependent misses to events at the $CaMn_4O_5$ -cluster elaborated by femtosecond X-ray crystallography.⁵² This allowed us, for the first time, to pinpoint possible "molecular errors" that cause the S state dependent misses during the water oxidizing Kok cycle.

Experimental procedures

Preparation of thylakoid and PSII membranes

Spinach thylakoid membranes were isolated as described in ref. 20 and PSII-enriched membrane fragments (BBY-type) as in ref. 86 and stored at -80°C until used.



FIOP measurements

The FIOP experiments were performed with an unmodulated in-house build Joliot-type electrode embedded in the thermostated, buffer flowing stainless steel cell as described in ref. 20. 10 μL of thylakoid membranes at concentration of 2.4 mg Chl per mL in a measuring buffer, containing 25 mM 2-(*N*-morpholino)ethanesulfonic acid (Mes) – NaOH (pH 6.3), 400 mM sucrose, 5 mM MgCl_2 and 10 mM NaCl, were given 2 saturating pre-flashes at frequency of 1 Hz and dark adapted for 15 min at 20 °C. No artificial electron acceptors were present. Thereafter, 16 saturating flashes were given at indicated temperature (–10 °C, 1 °C, 10 °C and 20 °C) and flash frequency (1.25 and 5 Hz) and FIOPs were recorded. Flashes were provided by a Nd:YAG G100 laser from Spectra Physics (6 ns, 100 mJ, 532 nm). FIOPs analysis was done by using in-house developed software routine.

EPR sample preparation

PSII membranes were diluted to 2 mg Chl per mL in the measuring buffer and filled into the calibrated EPR tubes of 4 mm outer diameter. The EPR samples were exposed to room light at 20 °C for 5 minutes to fully oxidize Y_D , and were then dark adapted for 15 min. Thereafter, PSII in the samples was synchronized to the S_1Y_D^+ state by the application of two saturating pre-flashes given at 1 Hz frequency followed by a dark adaptation for 30 min at 20 °C.^{21,40} Then PpBQ in dimethylsulfoxide (DMSO) was added to a final concentration of 0.5 mM. 30 s after the addition of PpBQ, samples were transferred to an ethanol bath at the indicated temperature (–10 °C, 1 °C, 10 °C and 20 °C). After sample temperature equilibration for 3 min, the samples were immediately exposed to saturating turnover flashes (from 0 to 6) at indicated flash frequency (1.25, 5 or 10 Hz) and frozen within 1–2 s in an ethanol–dry ice bath at 198 K and then transferred to liquid N_2 . Flashes were provided by a Nd:YAG G200 laser from Spectra Physics (6 ns, 840 mJ, 532 nm).

EPR spectroscopy

EPR measurements were performed with a Bruker BioSpin GmbH (Germany) ELEXYS E500 spectrometer with a SuperX ER049X microwave bridge and high Q ER4122SHQE-LC cavity. The spectrometer was equipped with an ESR 900 cryostat and ITC 503 temperature controller from Oxford Instruments, UK. The Y_D^+ radical signal and the S_2 state multiline signal were measured directly after the flashing. Then illumination into the EPR cavity at 5 K with visible light (160 W m^{-2} , white light lamp projector, 4 min) to induce the split S_1 , S_3 and S_0 EPR signals, and with 830 nm light (280 W m^{-2} , LQC830-135E laser diode, Newport, USA, 10 min) to induce only the split S_3 EPR signal, were carried out as described in.^{21,40} In addition, after the initial measurements, each sample was illuminated at 198 K in dry ice/ethanol bath (white light, 4 min) and the S_2 state multiline was recorded again. Each experiment of the flash series at a defined temperature and flash frequency was carried out at least twice with deviation of the determined S state distributions by less than 5%.

Quantification of the S state distribution after turnover flashes

Quantification of the S state distribution after the application of turnover flashes was done as in ref. 21 and 40. All EPR data were normalized to the amplitude of the fully induced Y_D^+ radical signal of each sample. The fraction of centers in the S_1 , S_2 , S_3 and S_0 states in the samples exposed to 0–6 flashes, were determined from the split S_1 , S_2 multiline, split S_3 , and split S_0 EPR signals (Fig. 2) correspondingly. In the dark sample, after application of our pre-flash protocol, all PSII centers stay in the S_1 state. Only the split S_1 signal was observed and thus, its intensity represents 100% of the PSII centers. Our early results have shown that all centers in the S_1 state are turned over to the S_2 state by one saturating laser flash provided at 1 °C. Thus, the S_2 multiline signal intensity induced by a single flash at 1 °C also represents 100% of the PSII centers. The application of two flashes at 1 °C resulted in the appearance of S_3 state but with some PSII centers remaining in the S_2 state. The population of the S_3 state was determined by measuring the split S_3 EPR signal induced by near-infrared (NIR) or visible light. Similarly, the amplitude of the split S_0 EPR signal induced by visible light was defined by the PSII centers in S_0 state in three-flash sample. Illumination at 198 K allowed estimation of PSII centers

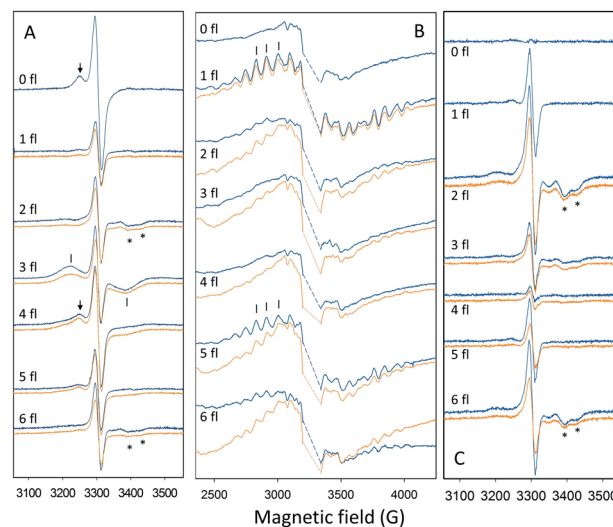


Fig. 2 EPR spectra used to quantify the fraction of PSII centers in the different S states after 0–6 saturating flashes with 1.25 Hz frequency at –10 °C (orange) and 10 °C (blue). (A) Flash-dependent oscillation of the split S_1 , split S_3 and split S_0 EPR signals. EPR signals were induced by illumination by visible light for 4 min at 5 K. The spectra are light minus dark difference spectra. (B) Flash-dependent oscillation of the S_2 multiline EPR signal. The large intensity from Y_D^+ in the center has been removed for clarity (dashed line). (C) Flash-dependent oscillation of the split S_3 EPR signal induced by illumination by NIR light for 10 min at 5 K. The spectra presented are light minus dark difference spectra. Peaks used to quantify the EPR signals are indicated by arrows (S_1 , A), bars (S_0 , A and S_2 , B) and stars (S_3 , A and C). EPR conditions: for (A and C): microwave power 25 mW, microwave frequency 9.27 GHz, modulation amplitude 10 G, temperature (T) 5 K; for (B): microwave power 10 mW, microwave frequency 9.27 GHz, modulation amplitude 20 G, T 10 K.



remaining in the S_1 state and to ensure the maximal size of the S_2 state multiline EPR signal in each sample. It should be noted that in samples that give rise to an EPR spectrum with mixed split EPR signals, a weighted deconvolution of these as described in ref. 21 and 40 was used to obtain a pure split EPR signals from the different S states to enable the analysis. Finally, the fraction of PSII centers in each S state was defined by the corresponding size of respective EPR signal, making possible the quantification of S state distribution in all samples. The procedure described here is used for analysis of experiments with turnover flashes given at 1 °C but was also applied for experiments at other temperatures. All spectral analysis was done by Xepir 2.6b Bruker BioSpin software.

Analysis of the changes in the CaMn_4O_5 -cluster during the S cycle

Changes in the CaMn_4O_5 -cluster and its immediate surrounding during the S state cycle were visualized by using 6DHE, 6DHF, 6DHO and 6DHP pdb files from the Protein Data Bank.⁵² Molecular visualization was done with YASARA view program (YASARA Biosciences GmbH).

Results

S state distribution after the application of turnover flashes at different temperatures

It is well-known that the S state transitions during the water oxidation cycle are strongly temperature dependent.^{15,18,28,40,47,59} Previously, we used EPR spectroscopy to determine the S state dependence of the misses at two temperatures, 1 °C and 20 °C.⁴⁰ In this study, we have now extended these data to two other temperatures: −10 °C and 10 °C. The resulting EPR spectra and corresponding S state distribution after each turnover flash are shown in Fig. 2 and Table 1. From these data the miss parameter after each flash was determined directly from the fraction of PSII centers that did not advance to the next S state as

described in ref. 40 and presented in Table 2. Table 2 also shows our earlier data obtained at 1 °C and 20 °C for comparison.

We observed no misses in the S_1 to S_2 transition at −10 °C. This is similar to the situation at 1 °C. The miss factor for the S_2 to S_3 state transition at −10 °C was found to be very high, 38%. For the S_3 to S_0 and S_0 to S_1 transitions the miss parameter was also found to be higher than at higher temperatures – 16% and 17% respectively (Tables 1 and 2).

At 10 °C, the miss in the S_1 to S_2 transition was no longer zero, instead a very small miss (3%) was found. For the rest of the S state transitions the miss factor was smaller than at lower temperatures. The miss parameter in the S_2 to S_3 state transition, 19%, was again highest if compared to the other transitions at the same temperature. At 10 °C, the miss parameters for the S_3 to S_0 and S_0 to S_1 transitions were 5% and 7%, respectively (Tables 1 and 2).

The temperature dependence of the miss parameter for each individual transition in the S cycle between −10 °C and 20 °C from the present and previous study⁴⁰ are compiled in Table 2 and Fig. 3. Two different trends can be observed. First, the S_1 to S_2 transition showed an increase in the miss parameter with increase of the temperature. No miss was observed in the S_1 to S_2 transition at −10 °C and 1 °C. Above this temperature this transition was less effective, the miss parameter reaching 10% at 20 °C (Fig. 3).⁴⁰

The S_2 to S_3 and S_3 to S_0 state transitions in the S cycle were different. The highest miss factor in the entire temperature range studied was found for the S_2 to S_3 transition at −10 °C – almost 40% (Table 2). It gradually decreased with increasing temperature reaching 16% at 20 °C (Fig. 3). The miss parameter in the S_3 to S_0 transition was lower than in the S_2 to S_3 transition but its temperature dependence was very similar. At −10 °C it was found to be 16% and decreased to only 3% at 20 °C (Table 2, Fig. 3). In the S_0 to S_1 transition, the miss is first decreased from 17% to 7% when temperature was changed from −10 °C to 10 °C and then slightly increased to 11% at 20 °C (Table 2, Fig. 3).

It is also worth to mention that under our experimental conditions the overall S state turnover was found to be most

Table 1 Distribution of the different S states (% of the total PSII amount) in EPR samples after the application of 0–6 turnover flashes given at a flash frequency of 1.25 Hz at −10 °C and 10 °C^a

Temp.	Fl. no.	S_1	S_2	S_3	S_0	$S_1^{(2nd)}$	$S_2^{(2nd)}$	$S_3^{(2nd)}$	Total
−10 °C	0	100							100
	1		100						100
	2		38 ± 2	62 ± 1					100
	3		18 ± 1	30 ± 1	52 ± 1				100
	4		13 ± 1	12 ± 1	32 ± 1	43 ± 1			100
	5			5 ± 1	15 ± 1	25 ± 3	54 ± 2		99
	6				12 ± 1	5 ± 1	42 ± 2	39 ± 1	98
10 °C	0	100							100
	1	3 ± 1	97 ± 1						100
	2		21 ± 1	79 ± 1					100
	3		4 ± 1	21 ± 1	75 ± 1				100
	4		3 ± 1	6 ± 1	20 ± 2	70 ± 1			99
	5			3 ± 1	9 ± 2	19 ± 1	70 ± 1		101
	6				4 ± 1	8 ± 1	33 ± 1	56 ± 1	101

^a The fraction of the S_i state was determined from the EPR spectra as described in the text and in ref. 40.



Table 2 Miss parameters in each S state transition determined from the results in Table 1 (at 1.25 Hz flash frequency)^a

Temp.	$S_1 \rightarrow S_2$	$S_2 \rightarrow S_3$	$S_3 \rightarrow S_0$	$S_0 \rightarrow S_1$	$S_1 \rightarrow S_2^{(2nd)b}$	$S_2 \rightarrow S_3^{(2nd)b}$	Total ^c	Average ^d
−10 °C	0	38	16	17	nd	nd	71	17.75
1 °C ^e	0	23	7	10	0	20	40	10
10 °C	3	19	5	7	0	20	34	8.5
20 °C ^e	10	16	3	11	8	15	40	10

^a The miss factor is given in % of total PSII in the corresponding S_i state that didn't proceed to the next S_{i+1} state after the flash. ^b Miss parameters of the transitions in the second turnover of S cycle. ^c Sum of misses for all S transition of the first turnover of the S cycle (total miss). ^d Average miss for single transition of the first turnover of the S cycle. ^e Data from ref. 40. nd - not determined. Accuracy is <5% (standard error).

effective at 10 °C (Table 2, Fig. 3). The sum of the miss factors for the entire S cycle (the total miss) was the lowest at this temperature (34%) and the average miss factor was 8.5%. At 1 °C and 20 °C temperature, the total and average misses were slightly higher than those at 10 °C. However, at −10 °C, the total miss was more than 70%, mostly due to the contribution of the high miss during the S_2 to S_3 state transition (Table 2, Fig. 3).

FIOP measurements at different temperatures

In order to verify our EPR data we performed classical Joliot-type experiments by measuring FIOPs at different temperatures. These measurements were performed on the thylakoid membranes which retain an active PQ-pool. There are two differences in the experimental conditions in these measurements if compared to EPR: (i) more intact PSII containing membranes were used (ii) in the absence of exogenous electron acceptor. These differences were necessary in order to perform measurements on the bare electrode and to obtain lasting O_2

release oscillations. The rest of the conditions were similar to our EPR measurements and the data are shown in Fig. 4 and Table 3. It is known that FIOPs show significant temperature dependence.^{18,28} It is clear that at 1.25 Hz flash frequency oscillations were running deep in the temperature range between 1 °C and 20 °C, reflecting good WOC turnover (Fig. 4A–C). At −10 °C oscillations were damped after the second cycle (first 8 flashes), indicating high misses during S state turnover at this temperature (Fig. 4D). This tendency was even more pronounced in measurements at flash frequency of 5 Hz (Fig. 4E–G). With this higher flash frequency at −10 °C, oscillations were damped already after the first cycle (first 4 flashes, Fig. 4G).

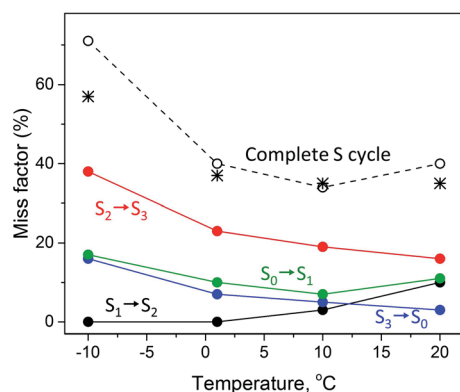


Fig. 3 Temperature dependence of the miss factor occurring in the $S_1 \rightarrow S_2$ (black), $S_2 \rightarrow S_3$ (red), $S_3 \rightarrow S_0$ (blue) and $S_0 \rightarrow S_1$ (green circles) state transitions as determined for a flash frequency of 1.25 Hz by EPR. The miss factor is the fraction of PSII centers that did not advance to the next S state after a turnover flash and was calculated from the data on the S state composition provided in Table 2 and ref. 40. The black dotted line (open black circles) represents the accumulated total miss of the complete S state cycle, i.e. is the sum of the misses during the individual S state transitions at the corresponding temperature. The black stars represent the total miss obtained with FIOP measurements (Fig. 4, Table 3).

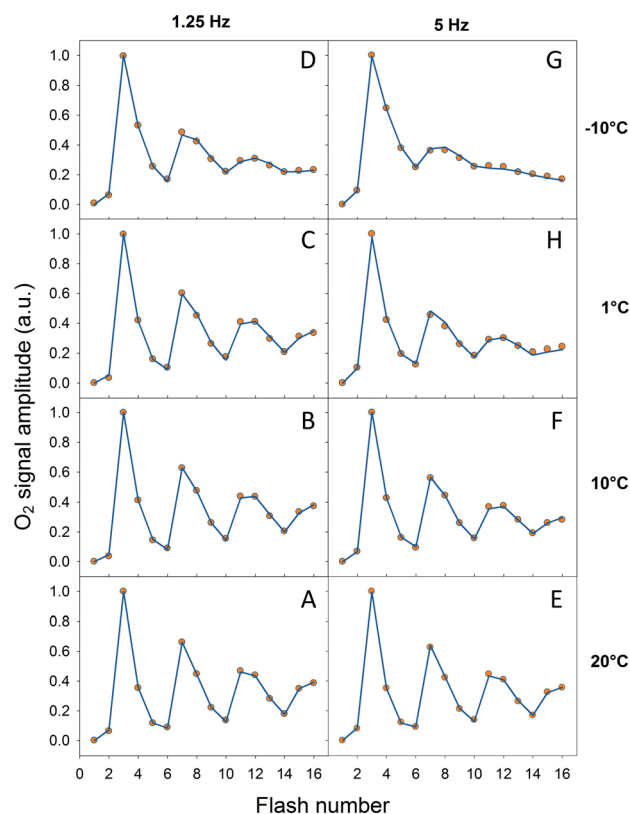


Fig. 4 Normalized FIOPs obtained from the thylakoid membranes at different temperatures (−10 °C, 1 °C, 10 °C and 20 °C) and flash frequencies (1.25 Hz and 5 Hz). Measured oxygen yield is shown by orange circles and simulation by blue lines.



Table 3 Miss factors determined from the FIOP data shown in Fig. 4^a

Temp.	Fl. freq.	Total ^b	Average ^c	Total EPR ^d	Average EPR ^e
−10 °C	1.25	57 ± 4	14.4	71	17.75
	5	69 ± 5	17.4	nd	nd
1 °C	1.25	37 ± 2	9.3	40	10
	5	44 ± 3	11.0	61	15.25
10 °C	1.25	35 ± 2	8.8	34	8.5
	5	37 ± 3	9.2	nd	nd
20 °C	1.25	35 ± 1	8.8	40	10
	5	35 ± 1	8.8	50	12.5

^a Flash frequencies are given in Hz. ^b Sum of misses for all S transition of the S cycle (total miss). ^c Average miss for single transition of the S cycle. ^d Total miss obtained from EPR measurements (Tables 2 and 6). nd – not determined. ^e Average miss obtained from EPR measurements (Tables 2 and 6). nd – not determined.

It was not possible to determine the miss factor for each individual S state transition with our FIOP measurements, however our analysis allowed us to determine the average and total miss for each measurement (Table 3). The average miss at 1.25 Hz was found to be about 9% at 1 °C, 10 °C and 20 °C, while at −10 °C it was about 14%. Correspondingly, the total miss of the complete S state turnover was almost 60% at −10 °C, while at the higher temperatures it was only 35–37% (Table 3). In addition, the steady-state O₂ yield during flash sequence at both frequencies declined by almost 20% when temperature was decreased from 20 °C to −10 °C (Fig. 4).

The temperature dependence of average miss parameter and total miss determined from the FIOP measurements was found to be very similar to the ones determined by EPR measurements (Fig. 3, Table 3). The only discrepancy we observed at −10 °C (almost 15% difference). This lower FIOP miss most probably reflects a combination of few factors, the most important of which is the more intact acceptor side in PSII in thylakoids. Otherwise, the similarity is much better at higher temperatures

and the overall trend in both EPR and FIOP measurement is the same (Fig. 3 and Table 3). This confirms our EPR measurements and quantitative analysis of the miss parameter in each individual S transition.

S state distribution after the application of turnover flashes at different frequencies

EPR experiments described at first were performed at flash frequency of 1.25 Hz. We performed another set of the experiments using the flash frequencies of 5 and 10 Hz. The experiments were performed at two temperatures, 1 °C and 20 °C, and the results are shown in Tables 4–6, Fig. 5 and ESI Fig. 1 and 2.[†]

Application of flashes with higher frequency had a profound effect on the miss parameter at 1 °C (Table 4, Fig. 5). For the S₂ to S₃ transition, the miss parameter rose from 23% at 1.25 Hz to 35% at 5 Hz and to more than 40% at 10 Hz. The effect on the S₃ to S₀ and S₀ to S₁ transition was similar – more than two times increase in the miss parameter with increasing frequency of the applied flashes (Table 6 and Fig. 5).

At 20 °C (Table 5, Fig. 5), there was a similar increase of the miss factor with higher flash frequency (1.25 Hz vs. 10 Hz) for the S₂ to S₃ and S₃ to S₀ transitions, in this case from 16% to 29% and from 3% to 11%. In contrast to the results at 1 °C, the increase of the miss parameter during the S₀ to S₁ transition was almost negligible at 20 °C (Table 6, Fig. 4).

Thus, the miss parameter in each S state transitions was not only temperature but also frequency dependent. There was a profound increase in the miss parameter with the increasing frequency of the applied turnover flashes at 1 °C as determined by both EPR and FIOP measurements (Table 6). The effect was also present at 20 °C. Largest increase was again observed for the S₂ to S₃ transition. It should be noted that determination of the miss parameter based on the FIOP measurements at 10 Hz flash frequency was not possible due to the overlap of the O₂ release peaks (not shown).

Table 4 Distribution of the different S states (% of the total PSII amount) in EPR samples after the application of 0–6 turnover flashes given at frequency of 5 Hz and 10 Hz at 1 °C (see ESI Fig. 1)^a

Fl. freq.	Fl. no.	S ₁	S ₂	S ₃	S ₀	S ₁ ^(2nd)	S ₂ ^(2nd)	S ₃ ^(2nd)	Total
5 Hz	0	100							100
	1		100						100
	2		35 ± 1	65 ± 1					100
	3		16 ± 2	29 ± 2	57 ± 1				102
	4		11 ± 2	13 ± 1	26 ± 1	49 ± 1			99
	5			6 ± 2	15 ± 1	28 ± 1	53 ± 3		102
	6				6 ± 1	10 ± 1	44 ± 2	40 ± 1	100
10 Hz	0	100							100
	1		100						100
	2		43 ± 2	57 ± 1					100
	3		24 ± 2	28 ± 1	48 ± 1				100
	4		17 ± 1	13 ± 2	33 ± 3	37 ± 3			100
	5			6 ± 1	22 ± 1	23 ± 3	46 ± 1		97
	6				14 ± 1	14 ± 1	41 ± 1	29 ± 3	98

^a The fraction of the S_i – state was determined from the EPR spectra as described in the text and in ref. 40.



Table 5 Distribution of the different S states (% of the total PSII amount) in EPR samples after the application of 0–6 turnover flashes given at frequency of 5 Hz and 10 Hz at 20 °C (see ESI Fig. 2)^a

Fl. freq.	Fl. no.	S ₁	S ₂	S ₃	S ₀	S ₁ ^(2nd)	S ₂ ^(2nd)	S ₃ ^(2nd)	Total
5 Hz	0	100							100
	1	10	90						100
	2		29 ± 1	71 ± 2					100
	3		11 ± 3	23 ± 1	66 ± 1				100
	4		10 ± 3	10 ± 1	22 ± 2	58 ± 1			100
	5			4 ± 1	9 ± 1	26 ± 1	62 ± 1		101
	6				6 ± 1	7 ± 1	38 ± 1	46 ± 3	97
10 Hz	0	100							100
	1	10	90						100
	2		33 ± 2	64 ± 1					97
	3		17 ± 2	26 ± 1	57 ± 1				100
	4		15 ± 1	7 ± 1	28 ± 1	49 ± 2			99
	5			3 ± 1	9 ± 1	30 ± 2	58 ± 2		100
	6				3 ± 1	14 ± 1	39 ± 2	42 ± 1	99

^a The fraction of the S_i – state was determined from the EPR spectra as described in the text and in ref. 40.

Table 6 Miss parameters in each S transition determined after application of turnover flashes given at different flash frequencies (in Hz)^a

Temp.	Fl. freq.	S ₁ → S ₂	S ₂ → S ₃	S ₃ → S ₀	S ₀ → S ₁	Total ^b	Average ^c	Total FIOP ^e	Average FIOP ^f
1 °C	1.25 ^d	0	23	7	10	40	10	37	9.3
	5	0	35	12	14	61	15.25	44	11
	10	0	43	16	23	82	20.5	nd	nd
20 °C	1.25 ^d	10	16	3	11	40	10	35	8.8
	5	10	21	7	12	50	12.5	35	8.8
	10	10	29	11	14	64	16	nd	nd

^a The miss factor is given in % of total PSII in the corresponding S_i state that didn't proceed to the next S_{i+1} state after the flash. ^b Sum of misses for all S transition of the first turnover of the S cycle (total miss). ^c Average miss for single transition of the first turnover of the S cycle. ^d Data from ref. 40. ^e Total miss obtained from FIOP measurements (Table 3) is shown for comparison. ^f Average miss obtained from FIOP measurements (Table 3) is shown for comparison. nd – not determined. Accuracy is <5% (standard error).

Discussion

Miss parameter originates from the WOC

The discovery of the period-four oscillation of the oxygen evolution by PSII and the introduction of the S cycle concept laid important foundation for understanding of photosynthetic water splitting.^{12–14} The dampening of the oscillation with increasing flash number is routinely explained with misses and double hits. Double hits, which normally originate from double turnover in the S cycle during a long flash, currently are easily eliminated by using shorter nanosecond laser flashes thus, limiting the charge separations in PSII to one per flash.²¹ Our measurements confirm this since we did not observe, for example, any EPR signals from the S₃ state after one flash, or from the S₀ state after two flashes, and further on (Tables 1, 4 and 5 and Fig. 2, ESI Fig. 1 and 2†).

At saturating flash excitation, the miss parameter may have many components.⁹ As we will outline below, two of the trivial ones we have excluded for the optimal conditions. Firstly, not all Q_A[•] could be oxidized between the flashes. In this situation, no stable charge separation can be obtained and thus a miss would be the consequence. Secondly, at very long times between

flashes, especially if reduced Y_D is present, charge recombination between already formed S₂ and S₃ states can occur and thereby increase the miss parameter. The remaining “actual misses” are a consequence of Y_Z[•] reduction by components other than the CaMn₄O₅-cluster. This most commonly happens by electrons from the acceptor side, but also other PSII cofactors may occasionally be oxidized, or reactive oxygen species may be formed. The following discussion outlines why we can exclude the trivial reasons for misses, and then aims to clarify the reasons for why the recombination reaction can win over the forward reaction more frequently in the S₂–S₃ transition than in all other transitions.

In our EPR experiments, both trivial reasons for misses were, in most cases, completely eliminated by the use of excess concentration of the exogenous electron acceptor PpBQ. Addition of 0.5 mM PpBQ efficiently keeps the Q_A–Fe–Q_B non-heme iron quinone complex in PSII completely oxidized, effectively eliminating recombination reactions with the S₂ and S₃ states. In the same way, it eliminates also the presence of Q_A[•] at the time of the flash and overcomes the acceptor side limitations of non-Q_B reducing PSII centers.^{53,54,60} This is confirmed by the maximal induction of the S₂ state multiline signal and similar



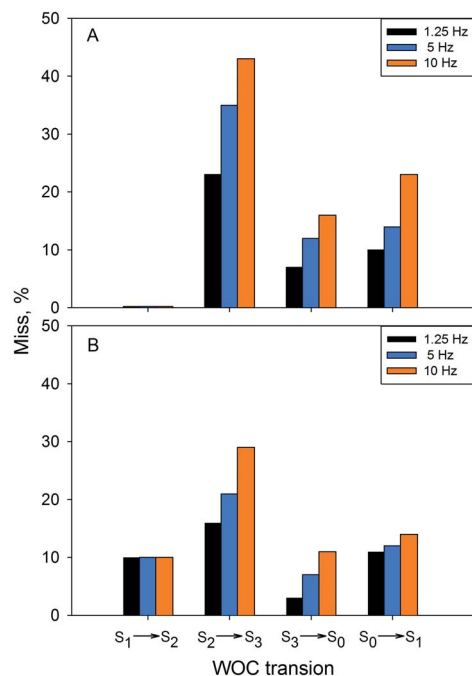


Fig. 5 The miss occurring in the $S_1 \rightarrow S_2$, $S_2 \rightarrow S_3$, $S_3 \rightarrow S_0$ and $S_0 \rightarrow S_1$ transitions at 1 °C (A) and 20 °C (B) after application of the turnover flashes at different frequencies: 1.25 Hz (black bars), 5 Hz (blue bars) and 10 Hz (orange bars). The miss was calculated from the fraction of PSII centers that did not advance to the next S state after a turnover flash.

misses observed during the second S cycle (Tables 1 and 2), as well as the deep oscillation of the S_2 multiline signal with almost no S_2 state population after flashes 3 and 4 (Table 1, 10 °C). The only exception were measurements at low temperature where electron transfer to PpBQ was less efficient and some blocked centers may occur (less than 5%).¹⁶ This can be seen as a partial contribution to the higher miss at 10 °C (Table 2, Fig. 3).

Thus, it is clear that at 1.25 Hz flash frequency the acceptor side of PSII kept efficiently oxidized (Tables 1 and 2, also see ref. 40). However, our experiments at 5 and 10 Hz frequency of the turnover flashes, especially at lower temperatures, indicate that some limitations appear on the acceptor side of PSII. At these conditions there is not enough time to complete the electron transfer from Q_A^- to PpBQ before the next flash arrives, and increase in misses during all S transitions is observed (Tables 3 and 6). It is worth to mention that the complete S cycle turnover takes place within 1–2 ms (Fig. 1 and see ref. 4 and 9 for the review). The highest frequency we used, 10 Hz or one flash per 100 ms is far too low to exert any limitations on these transitions. Therefore, it is very likely that the effect we observe in this experiment originates from the lingering electron on the acceptor side of PSII (most likely Q_A^-) at least in a fraction of the PSII centers (so-called closed or inactive PSII centers) making charge separation and therefore, the next state transition impossible. This is in agreement with the earlier results that demonstrated that the reactions at the PSII acceptor side are rate limiting at high flash frequencies.⁶¹ There are indications,

however, that the turnover efficiency of PSII is not affected by flash frequency up to 30 Hz when PpBQ was used as exogenous electron acceptor at 10 °C.⁶²

As mentioned above, another potential factor which can contribute to the increased miss parameter is the redox state of secondary tyrosine, Y_D . If present, reduced Y_D , can be oxidized by the S_2 and S_3 states in the dark in the pH dependent manner.^{17,56,58} Due to these reactions, if a flash is given in the S_1 or S_2 state in the presence of reduced Y_D , the transition to the S_2 and S_3 state is reversed in a fraction of PSII centers within 1 s between the flashes.⁵⁶ However, our pre-flash treatment kept Y_D fully oxidized and thus its interference with produced S states in the time frame of our experiments (*i.e.* sample freezing, 1–2 s) is negligible. The latter argument also holds for Q_B^- , Cyt b_{559} or other electron donor/acceptor cofactors in PSII, for all of which the redox reactions with the S states are even slower (Fig. 1).⁵⁷

Thus, taking into account implementation of the pre-flash protocol in the presence of PpBQ in EPR measurements, our starting sample was 100% in the S_1 state with 100% oxidized Y_D (Y_D^+). During S state advancement there was thus no contribution from recombination reactions with the S_2 or S_3 states or of blocked centers to the miss parameter. Therefore, we can safely conclude that at our measuring conditions, at least at low flash frequencies and temperatures above 0 °C, the misses we observed in PSII membrane samples in the EPR and FIOPs measurements are actual misses originated from the molecular events in the WOC during S state transitions. While simple kinetic competition between forward and charge recombination reactions certainly also contribute to misses,^{35,36} we propose that the actual misses are mainly a consequence of “molecular miss” events at the water oxidation (donor) side of PSII that do not allow an S state dependent fraction of centers to advance to the next S state, thereby leading to reduction of Y_Z by Q_A^- or Q_B^- .

The highest miss was found in the $S_2 \rightarrow S_3$ transition

Understanding nature of miss factor during S state transitions is important not only for elucidating the reaction mechanism of water oxidation, but also for analysis of spectroscopic, electronic and geometric structure of the WOC. Our current and previous EPR study⁴⁰ and more recent investigation,⁵⁵ where global fit analysis of the FIOPs was used and where life-times of the all metastable S states and redox states of Q_A , Q_B and Y_D were considered, showed that the misses during water oxidation are S state-dependent and the highest miss was found in the $S_2 \rightarrow S_3$ transition, whereas the $S_3 \rightarrow S_0$ transition, where O_2 formation and release occur, showed a relatively low miss factor (Table 2, Fig. 3).

However, these results are in contradiction with studies by de Wijn *et al.*²⁵ and by Suzuki *et al.*⁴³ On the basis of the fluorescence studies on the thylakoid membranes from spinach, de Wijn and van Gorkom reported a lack of misses in the S_0 to S_1 transition, whereas the largest miss was found for the $S_3 \rightarrow S_0$ transition.²⁵ Similarly, the largest miss in the $S_3 \rightarrow S_0$ transition was also reported by Suzuki *et al.* after using FTIR difference spectroscopy in the PSII core complexes from



Thermosynechococcus elongatus and PSII membranes from spinach.⁴³ Both these studies relied on the acceptor side components for dissemination of the individual miss factors. In the first study the flash dependent Chl fluorescence transient yield, which depends on the redox state of Q_A was used as a probe to deconvolute the miss parameters. In the second study, the electron flow in PSII detected by monitoring the CN stretching bands of ferri/ferrocyanide (used as an exogenous electron acceptor), was used with the same purpose.

In contrast, our measurements only rely on the signals originated from the WOC. We directly measure the S state distribution after the flash by EPR spectroscopy. Each EPR signal we measured, originates from the CaMn_4O_5 -cluster and is easily distinguishable from others (Fig. 2). Moreover, neither inactive PSII centers or acceptor side reactions or other components in PSII interfered with in our experiments. In addition, measurements at low temperature, where misses are amplified significantly, support our finding that the highest miss during the S cycle takes place during the $S_2 \rightarrow S_3$ transition. These facts are now well-established and clearly observable (Table 2, Fig. 3).

WOC miss parameters vary with temperature

It is known that temperature has a profound effect on the S state transitions by tuning of electron and proton transfer.^{15,18,28,35,40,47,59} Our measurements show different temperature behavior of misses during these transitions (Table 2, Fig. 3). Two trends were observed. The $S_1 \rightarrow S_2$ transition proceeds with a 100% efficiency at low temperatures but occurred with 10% misses at 20 °C, *i.e.* misses were increasing with increasing temperature (Table 2, Fig. 3). The miss parameter in the $S_0 \rightarrow S_1$, $S_2 \rightarrow S_3$ and $S_3 \rightarrow S_0$ transitions shown the opposite effect, *i.e.* they decrease with increasing temperature. The only exception is $S_0 \rightarrow S_1$ transitions which increased by 4% from 10 °C to 20 °C (Table 2, Fig. 3). Typical difference in the miss factor was about 10% in the investigated temperature range, except for the $S_2 \rightarrow S_3$ which was more than twice higher.

Our understanding is that these two trends reflect the different nature of molecular events during the S state cycle. The $S_1 \rightarrow S_2$ transition is unique among the reactions in the S state cycle in many aspects. It is the only transition which involves only an electron transfer step, *i.e.* it is not coupled to a proton release from the WOC^{63,64} and is thus largely pH independent.^{37,39} It is also operational at much lower temperatures than all other transition.¹⁵ At our measured low temperatures (−10 °C to 10 °C), the $S_1 \rightarrow S_2$ transition proceeds without any miss with transition efficiency of almost 100%. Only at higher temperatures misses started to occur. This reflects the fact that the WOC is unaltered and the surrounding protein requires only a small readjustment during this transition. We thus propose that the misses at 20 °C at least partially originate from the higher disorder of the protein matrix between the CaMn_4O_5 -cluster and Y_Z .

The opposite temperature dependence of misses in the other S state transitions reflects their more complicated chemical nature. They all include de-protonation events, show strong pH dependence in the physiological region and exhibit a much

higher temperature limit.¹⁵ In addition, the $S_2 \rightarrow S_3$ and $S_3 \rightarrow [S_4] \rightarrow S_0$ transitions involve water binding events and the later one also the O_2 formation and release event (Fig. 1). Decrease of miss factors with increased temperature during these transitions indicate that these events are promoted at high temperatures. We are addressing the molecular nature of these events in the section below.

Molecular origin of the miss factor at the WOC during the S cycle

Much spectroscopic, biophysical and biochemical information that has been accumulated over the years allow to certain extent understand the chemical nature of S state transitions and to pinpoint some possible missteps from which misses could be originated. We have attempted this exercise in our first publication.⁴⁰ Much more changed since then. First of all, the first high resolution structure of PSII and the CaMn_4O_5 -cluster became available.⁵¹ More importantly, with the development of the femtosecond XFEL methods it became possible to obtain structures of all four stable and metastable S states of the WOC.⁵² This allows us to discuss the origin of the misses based on the molecular structures and most feasible mechanism of water oxidation presently available.

The electron transfer only event in the $S_1 \rightarrow S_2$ transition implies no significant structural change in the CaMn_4O_5 -cluster. This is corroborated by a number of EXAFS spectroscopy studies.^{47,65–69} More recent structural studies also show no fundamental change upon this transition.⁵² The CaMn_4O_5 -cluster remained in the open, non-cubane geometry corresponding to the low spin configuration (Fig. 6).^{70,71} Change of Mn4(III) to Mn4(IV) oxidation state might induce small structural readjustments in the first coordination sphere, which in our opinion could contribute to the appearance of the miss at high temperature (20 °C). Otherwise, there is no chemical reason for miss in this transition.

The situation is dramatically changed during the $S_2 \rightarrow S_3$ transition where we found the lowest transition efficiency at all temperatures. This transition is pH dependent and shows the highest deuterium isotope effect on the kinetics for electron transfer from Y_Z .^{31,72,73} Both effects reflect large proton movements around Y_Z and from the CaMn_4O_5 -cluster to the bulk water. In addition, EXAFS studies have revealed major structural rearrangements in the CaMn_4O_5 -cluster involving a shift in coordination number for one of the Mn atoms.^{47,65–69,74} This was recently confirmed by XFEL experiments that show that a new oxygen (Ox), likely from the Ca-bound water (W3), is inserted and forms an additional hydroxo-bridge between Ca and Mn1, which is oxidized from Mn1(III) to Mn1(IV) (Fig. 6). It is suggested that the insertion is accompanied by the removal of the first proton from W3 and binding of a new water to Ca to fill the empty W3 binding site.^{52,75} This well-orchestrated sequence of events requires a structurally well-defined hydrogen and/or water network around the CaMn_4O_5 -cluster, especially surrounding the Ca, Y_Z and Mn1 site (the O1 channel^{52,75,76}) and immediate protein ligands. This correlates with changes in the Mn1 coordination and distances to Mn3 and Mn4. Even a small



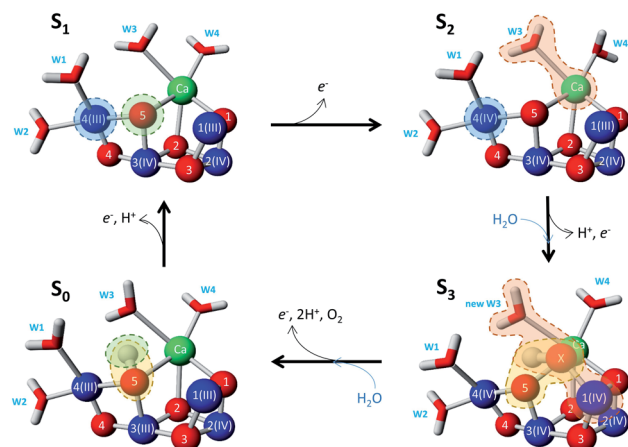


Fig. 6 Possible changes in the structure of the CaMn_4O_5 -cluster, e^- transfer, H_2O binding, H^+ and O_2 release during the S cycle. Sites of the miss origin during $\text{S}_1 \rightarrow \text{S}_2$, $\text{S}_2 \rightarrow \text{S}_3$, $\text{S}_3 \rightarrow \text{S}_0$ and $\text{S}_0 \rightarrow \text{S}_1$ transitions are correspondingly highlighted by blue, orange, yellow and green colored areas.

hindrance of any of these many steps during the $\text{S}_2 \rightarrow \text{S}_3$ transition will result in either complete failure or at least slowness of the transition and as a result in an increased miss parameter (charge recombination of Y_Z with acceptor side). Our results clearly suggest that this complicated chemistry is reflected in a lowered transition efficiency.

It must be mentioned that alternative conformations of the S_2 state and possibly S_3 state were proposed to exist and are manifested, for example, by the high spin EPR signal in the S_2 state. This structure has been proposed to represent a closed cubane in contrast to the low spin open cubane structure shown in Fig. 6. In an alternative scenario for water insertion during the $\text{S}_2 \rightarrow \text{S}_3$ transition, the so-called pivot or carousel pathways, the dominant, low-spin conformation of the S_2 state, often referred to S_2A , needs to convert after Y_Z oxidation into the closed cube, high-spin S_2B conformation, in which Mn4 attains the Mn(III) oxidation state instead of Mn1, before water can be inserted at Mn4(III) and the S_3 state is reached by Y_Z reduction.^{71,77–79} This scenario involves even more steps and sensitive equilibria, fully consistent with our finding that the $\text{S}_2 \rightarrow \text{S}_3$ transition, which has a low-driving force, has the highest miss. Both conformations were proposed to be interconvertible and energetically similar (discussed in ref. 78).

Very large chemical changes and protein structural rearrangements take place during the $\text{S}_3 \rightarrow [\text{S}_4] \rightarrow \text{S}_0$ transition, involving the final step in the water oxidation cycle – the formation and release of O_2 molecule. This final step in water oxidation is pH dependent,^{39,43} involves two deprotonation steps^{63,64} and binding of the second water molecule in the cycle (Fig. 6). EXAFS studies have also revealed that the coordination chemistry around the Mn1 atom that was altered in the $\text{S}_2 \rightarrow \text{S}_3$ transition is returned to its original state.^{47,65–69}

These data correlate with the latest structural study.⁵² According to one possibility,⁸⁷ O5 and Ox form dioxygen and concomitantly with O_2 release the new water molecule binds under deprotonation so that it binds as hydroxo at the O5

binding site. Thereby, the CaMn_4O_5 -cluster returns to the open cubane configuration and attains its lowest valence state (Fig. 6). We observed a quite low transition efficiency at -10°C , but not at higher temperatures in this transition (Fig. 3). Our understanding is that this final step in water oxidation to molecular oxygen (O_2) has a strong driving force after a critical, highly reactive intermediate (S_4) is formed from S_3Y_Z .^{80,81} Thus, O_2 release and the binding of the second water likely do not contribute to the miss under normal conditions. However, the formation of the S_4 state involves a deprotonation of the catalytic site, possibly of Ox, and the oxidation of the Mn_4CaO_5 -cluster, to form either a Mn-oxyl radical or a $\text{Mn(V)}\text{-oxo}$.^{82–84} Furthermore, additional structural changes cannot be excluded. Thus, all molecular misses connected to the $\text{S}_3 \rightarrow \text{S}_4 \rightarrow \text{S}_0$ transition are expected to come from these initial transformations leading to the formation of the S_4 state (Fig. 6).

We also observed relatively low miss parameter in the $\text{S}_0 \rightarrow \text{S}_1$ transition, especially at -10°C (Fig. 3). The $\text{S}_0 \rightarrow \text{S}_1$ transition is pH dependent^{39,85} and involves the release of one proton.^{37,64} The structural changes at the CaMn_4O_5 -cluster, however, are not large and are considered to involve deprotonation of a μ -oxo-bridge between two of the Mn atoms^{64,66} and it is reasonable to suggest that it takes place at the O5 position (Fig. 6). This deprotonation event is the one that governs misses during this transition and is different from the change in Mn-coordination and water binding occurring in the previous transitions thus, resulting in the altered temperature dependence.

Conclusions

In this work we correlate the actual misses or transition efficiencies in the S state cycle with different degrees of failure of the specific molecular reactions during the WOC advancement. The absence of misses (100% transition efficiency) during the $\text{S}_1 \rightarrow \text{S}_2$ transition reflects the only electron transfer event at the Mn4. The transition efficiency during the $\text{S}_2 \rightarrow \text{S}_3$ transition and its strong temperature dependence reflect deprotonation steps, water binding and insertion of new hydroxo resulting in significant structural rearrangements at the CaMn_4O_5 -cluster that need to be performed with a low driving force. The intermediate transition efficiency in the $\text{S}_0 \rightarrow \text{S}_1$ and $\text{S}_3 \rightarrow \text{S}_0$ transitions and their slight temperature dependence instead reflects that these are governed by electron and proton transfer steps that involve only minor structural changes at the WOC. Details of these reactions are depicted in Fig. 6.

Data availability

The data supporting this article are available in the ESI.†

Author contributions

F. M. and S. S. conceived and designed experiments. G. H., P. C., and M. F. performed experiments. G. H., P. C., S. S., J. M. and F. M. analyzed data. F. M. and J. M. wrote the manuscript.



Conflicts of interest

There are no conflicts to declare.

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