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Spectrally decomposed dark-to-light transitions in *Synechocystis* sp. PCC 6803

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Abstract

Photosynthetic activity and respiration share the thylakoid membrane in cyanobacteria. We present a series of spectrally resolved fluorescence experiments where whole cells of the cyanobacterium *Synechocystis* sp. PCC 6803 and mutants thereof underwent a dark-to-light transition after different dark-adaptation (DA) periods. Two mutants were used: (i) a PSI-lacking mutant (ΔPSI) and (ii) M55, a mutant without NAD(P)H dehydrogenase type-1 (NDH-1). For comparison, measurements of the wild-type were also carried out. We recorded spectrally resolved fluorescence traces over several minutes with 100 ms time resolution. The excitation light was at 590 nm so as to specifically excite the phycobilisomes. In ΔPSI, DA time has no influence, and in dichlorophenyl-dimethylurea (DCMU)-treated samples we identify three main fluorescent components: PB–PSII complexes with closed (saturated) RCs, a quenched or open PB–PSII complex, and a PB–PSII ‘not fully closed.’ For the PSI-containing organisms without DCMU, we conclude that mainly three species contribute to the signal: a PB–PSII–PSI megacomplex with closed PSII RCs and (i) slow PB → PSI energy transfer, or (ii) fast PB → PSI energy transfer and (iii) complexes with open (photochemically quenched) PSII RCs. Furthermore, their time profiles reveal an adaptive response that we identify as a state transition. Our results suggest that deceleration of the PB → PSI energy transfer rate is the molecular mechanism underlying a state 2 to state 1 transition.

Keywords  Cyanobacteria · Spectrally resolved fluorometry · Singular value decomposition · Time-resolved spectroscopy · Plastoquinone pool · Cyclic electron flow

Abbreviations

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Definition</th>
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<tbody>
<tr>
<td>ATP</td>
<td>Adenosine triphosphate</td>
</tr>
<tr>
<td>CEF</td>
<td>Cyclic electron flow</td>
</tr>
<tr>
<td>COX</td>
<td>Cytochrome <em>c</em> oxidase</td>
</tr>
<tr>
<td>DA</td>
<td>Dark adaptation</td>
</tr>
<tr>
<td>DCMU</td>
<td>Dichlorophenyl-dimethylurea</td>
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<tr>
<td>EET</td>
<td>Excitation energy transfer</td>
</tr>
<tr>
<td>FDP</td>
<td>Flavodiiron proteins</td>
</tr>
<tr>
<td>FNR</td>
<td>Ferredoxin–NADP + reductase</td>
</tr>
<tr>
<td>Flv1/3 (2/4)</td>
<td>Flavodiiron protein 1/3 (2/4)</td>
</tr>
<tr>
<td>$F_M$</td>
<td>Maximum fluorescence during a saturating pulse</td>
</tr>
<tr>
<td>$F_S$</td>
<td>Steady-state fluorescence at the end of background illumination</td>
</tr>
<tr>
<td>$F_V$</td>
<td>Variable fluorescence</td>
</tr>
<tr>
<td>LEF</td>
<td>Linear electron flow</td>
</tr>
<tr>
<td>HiP</td>
<td>High-light inducible protein</td>
</tr>
<tr>
<td>NADPH</td>
<td>Nicotinamide adenine dinucleotide phosphate</td>
</tr>
<tr>
<td>NDH-1</td>
<td>NAD(P)H dehydrogenase type-1</td>
</tr>
<tr>
<td>PB</td>
<td>Phycobilisome</td>
</tr>
<tr>
<td>PC</td>
<td>Plastocyanin</td>
</tr>
<tr>
<td>PMF</td>
<td>Proton-motive force</td>
</tr>
<tr>
<td>PQ</td>
<td>Plastoquinone</td>
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<td>PS</td>
<td>Photosystem</td>
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Alonso M. Acuña and Pascal van Alphen have contributed equally to this work.

Electronic supplementary material  The online version of this article (https://doi.org/10.1007/s11120-018-0505-0) contains supplementary material, which is available to authorized users.

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Introduction

Cyanobacteria possess a remarkable thylakoid membrane that contains both photosynthetic and respiratory subunits. Photosynthetic activity is best described by the so-called Z scheme (Govindjee and Shevela 2011; Björn and Nickelsen 2013; Stirbet and Govindjee 2011). Input light interacts with pigment–protein complexes, the phycobilisomes (PB), that absorb and transfer excitation energy downhill to a series of cofactors until it reaches a particular pair of chlorophylls (Chls), the special pair, located in the core pigment–protein complexes of photosynthesis: the photosystems (PS) I and II. The special pair is excited to a charge transfer state (van Grondelle et al. 1994) from which an electron is released and, after being itself transferred through a series of cofactors, including primary electron acceptor $Q_A$, it is expelled from PSII in the form of the twice reduced secondary electron acceptor $Q_B$ (PQH$_2$) that physically wanders to the cytochrome $b_{6}f$ (cyt $b_{6}f$). From there, electrons are shuttled to a small copper containing protein, the plastocyanin (PC) that can deliver the electron to PSI. This, so-called, linear electron flow (LEF) finalizes with the electron transfer from PSI to NADPH (van Thor et al. 1999).

Remarkably, in cyanobacteria respiratory processes share some of the same components, as the plastoquinone (PQ) pool, the PC pool, the cyt $b_{6}f$ and the cytochrome oxidase participate both in photosynthetic and respiratory activities (Liu et al. 2012; Liu 2015; Lea-Smith et al. 2016; Schmetterer 1994; Jeanjean et al. 1993). The respiratory subunits providing the electron input to the PQ pool are the type-1 and -2 NADPH dehydrogenase, respectively, NDH-1 and NDH-2, and the succinate dehydrogenase (SDH). NDH-1 and SDH have been reported to be the main sites where electrons are donated (Liu et al. 2012; Cooley and Vermaas 2001). As for LEF, PQH$_2$ delivers the electrons to cyt $b_{6}f$ from where they either continue their way to the cytochrome oxidase via the PC pool and accomplish respiration or, alternatively, get shuttled to PSI and participate in cyclic electron flow (CEF) around PSI, a mechanism by means of which the cell adjusts the NADPH/ATP production ratio. Thus, as light input drops and photosynthetic electron flow stops, the PQ pool still serves as a vehicle for electron transfer related to ongoing respiratory activity.

Furthermore, it has been reported that the flavodiiron proteins (FDPs) provide alternative electron transfer pathways (Shimakawa et al. 2015; Ermakova et al. 2016; Zhang et al. 2009; Allahverdiyeva et al. 2013) and are involved in photoprotection (Allahverdiyeva et al. 2015; Zhang et al. 2009; Bersanini et al. 2017) although the specific function of these heterodimers remains a controversial topic. The cyanobacterium *Synechocystis* sp. PCC 6803 (*Synechocystis* hereafter) has four FDPs: Flv1–4. While the pair Flv1/3 is involved in photoreduction of $O_2$ (Allahverdiyeva et al. 2013) after accepting electrons from PSI, Zhang et al. (2009) reported plummeting levels of evolved oxygen in a mutant of *Synechocystis* lacking Flv2/4 and exposed to high-light, making them more susceptible to photoinhibition, and therefore, suggesting that Flv2/4 can supply PSI with an additional channel for electron transfer to alleviate excitation pressure. Crucially, Bersanini et al. (2014) found that overexpression of the flv4-2 operon in *Synechocystis* led to improved PSI photochemistry and kept the PQ pool in a predominantly oxidized redox state, providing unequivocal evidence that Flv2/4 acts as an important electron sink at the PSI acceptor side. The authors also report that a fully assembled phycobilisome is required for the stable expression of the electron sink mechanism.

Extended periods (several minutes) of darkness are known to drive the cyanobacterial cell towards a low fluorescence state denominated state 2 (Mullineaux and Allen 1986, 1990; Papageorgiou 1996; Campbell et al. 1998). This state is characterized by an increase of the effective antenna size of PSI. Exposing dark-adapted cells to light drives a state 2 to state 1 transition, typically within 10–30 s (Jallet et al. 2012; Kirilovsky 2007; Kaňa et al. 2009, 2012). State 1 is characterized by an increase in the effective antenna size of PSII, and hence, state 1 has a higher fluorescence yield (Kirilovsky et al. 2014). The cell is able to reversibly transition between state 1 and state 2 (state transitions) depending on the light conditions, in an effort to re-distribute excitation energy between PSI and PSII. See Mullineaux and Emlyn-Jones (2005) for an in-depth review on state transitions.

This fluorescence study is an effort in disentangling these heavily intertwined energy and electron transfer pathways. After exciting PB with 590 nm light, spectrally resolved fluorescence traces were recorded over several minutes with 100 ms time resolution. The following mutants of *Synechocystis* were used in comparison to the wild-type (WT): (i) a PSI-lacking mutant (ΔPSI hereafter); and (ii) a mutant without NDH-1 (M55 hereafter). In the absence of PSI, the cell lacks linear electron flow and, consequently, cannot produce NADPH, thus requiring glucose to enable growth (Vermaas et al. 1994). It also presumably possesses additional photoprotective mechanisms related to overexpression of the high-light inducible protein (HiIP) HiID (Sinha et al. 2012; Chidgey et al. 2014). By definition, transitions do not take place in ΔPSI. The M55 mutant, lacking the *ndhB* gene, does not assemble the NDH-1 complexes (Ogawa 1991).

<table>
<thead>
<tr>
<th>SAS</th>
<th>Species-associated spectrum</th>
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<tbody>
<tr>
<td>SDH</td>
<td>Succinate dehydrogenase</td>
</tr>
<tr>
<td>SVD</td>
<td>Singular value decomposition</td>
</tr>
<tr>
<td>WT</td>
<td>Wild-type</td>
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This is a highly disruptive mutation because these complexes have been reported to be involved in both respiratory and photosynthetic activities (Mi et al. 1995; Schmetterer 1994). Of particular interest is the disruption of respiratory activity in this mutant which causes a decreased input of electrons into the PQ pool in the dark. Since NDH-1 serves as an electron acceptor for ferredoxin (Fd), it is also involved in CEF, which helps increase the PMF, leading to increased synthesis of ATP. The M55 mutant retains the capacity to undergo state transitions (Schuurmans et al. 2015). Furthermore, this mutant reportedly has impaired CO₂ uptake (Ohkawa et al. 2000). In addition to the genetic disruptions, some samples have been treated with dichlorophenyl-dimethylurea (DCMU), a common inhibitor known for closing the PSII RCs as it blocks electron transfer from $Q_A$ to $Q_B$.

In an effort to anticipate from which pigment-protein complexes the fluorescence signal will originate we present in Fig. 1 a cartoon overview of putative fluorescent species that may play a role in the experiments reported in “Results” section. Beyond the evident species, i.e., the PB–PSII complex, a functional PB–PSII–PSI megacomplex (Fig. 1a, e, g) and the CpcG2-type of PB (Fig. 1b) are presented. The PB–PSII–PSI megacomplex has been isolated by Liu et al. (2013) and Kondo et al. (2009), and Gao et al. (2016) have reported on the CpcG2 type of PB. Chukhutsina et al. (2015) claim to observe such antenna in ultra-fast time-resolved fluorescence experiments. Additionally, we explore the possibility of a PB–PSII dimer complex with a single closed RC while the other remains open (see Fig. 1d, f).

### Materials and methods

#### Cell cultures

Wild-type *Synechocystis* was obtained from D. Bhaya (Stanford). Wild-type cells were grown in a photobioreactor in turbidostat mode (growth rate ca. 0.05 h⁻¹) at an optical density at 730 nm (OD₇₃₀) of 0.40 ± 0.01, inoculated in modified BG-11 supplemented with 10 mM NaHCO₃ and bubbled with 1% (v/v) CO₂ in N₂. The culture was illuminated with 636 nm light intensity of 80 µmol photons m⁻² s⁻¹ as measured outside the reactor, opposite and centered relative to the light panel.

The PSI-deficient mutant of *Synechocystis* (Shen et al. 1993) was obtained from C. Funk (Umeå University, Sweden). Prior to preparing a liquid culture, cells were streaked on BG-11 agar plates from a stock stored at − 80 °C in 15% glycerol as described before (Acuña et al. 2018a). Subsequently, the plates were placed in an incubator (Versatile Environmental Test Chamber MLR-350H, Sanyo) at 30 °C with a humidified atmosphere with increased CO₂ (2% v/v). The plates were covered with layers of paper to reduce incident light intensity to below 5 µmol photons m⁻² s⁻¹. For...

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**Fig. 1** Visualization of putative fluorescent species in *Synechocystis* ordered along the scale of their expected fluorescence quantum yield $\phi_f$. Yellow arrows represent EET from the PB core to PS I or II, their width indicates how efficient EET is from the PB to the respective PS. An “X” stands for a closed PSII RC. a PB–PSII–PSI megacomplex efficiently quenched by both PSI I and II (open RC); b CpcG2 type of PB transferring energy to PSII; c PB–PSII complex with open RCs; d PB–PSII–PSI megacomplex with a single RC closed and efficiently quenched by PSI and PSII; e PB–PSII–PSI megacomplex with fast energy transfer to PSI and closed PSII RCs; f PB–PSII complex with a single closed RC; g PB–PSII–PSI megacomplex with slow energy transfer to PSI and closed PSII RCs; h PB–PSII complex with closed RCs; i functionally uncoupled PB.
liquid cultures, 25 mL modified BG-11 (BG-11-PC, van Alphen et al. manuscript in preparation, substituting NaNO₃ for 10 mM NH₄Cl) was inoculated and supplemented with 10 mM glucose, 5 µg mL⁻¹ chloramphenicol, and 25 mM 1,4-piperazinediopropanesulfonic acid (PIPPS)–KOH buffered at pH 8.0 in a 100 mL flask (FB33131, Fisherbrand). In similar fashion to the agar plates, flasks were wrapped in layers of paper to reduce the incident light intensity to below 5 µmol photons m⁻² s⁻¹. The flasks were placed in a shaking incubator (Innova 44, New Brunswick Scientific) at 120 rpm and 30 °C, equipped with a custom LED panel containing orange-red and blue LEDs of 632 and 451 nm (blue, both 8 nm full width at half maximum), respectively.

The M55 mutant was a gift from T. Ogawa (Nagoya University, Japan) and was grown in a chemostat kept at 30 °C, with a dilution rate of \( D = 0.03 \) h⁻¹, illuminated by 635 nm light intensity of 40 µmol photons m⁻² s⁻¹ and bubbled with 1% (v/v) CO₂ in N₂.

The abundance of Synechocystis sp. PCC6803 ‘wild types’ provides an additional dimension of complexity, as there may be significant differences that are not readily apparent in routine culturing (Zavřel et al. 2017). We have attempted to account for these differences by subjecting the different wild-type strains to similar conditions as the derived mutant strains since introducing all mutations in a single wild-type strain was not successful. However, no spectral difference was observed between the ‘Stanford’ and ‘Vermaas’ wild-types in either photoautotrophic conditions as used for WT and M55, or heterotrophic conditions as used for the PSI-deficient mutant. All strains used, i.e., WT, M55, and ΔPSI, derive from the same branch of glucose-tolerant, non-motile Synechocystis strains, i.e., the ATCC branch (Trautmann et al. 2012; Zavřel et al. 2017).

**Spectrally resolved fluorescence induction**

The multiple LED set-up described in Lambrev et al. (2010) and Acuña et al. (2016) was placed in the vicinity of where the Synechocystis cells were grown, ensuring that the cells were in well-defined physiological conditions with minimal time between sampling and measuring. For every strain, the illumination protocols (Fig. S1) have been applied to whole cells harvested from the same batch, diluted to an OD₇₃₀ of 0.4 in fresh medium. In all cases, the background light (590 nm) intensity was 450 µmol photons m⁻² s⁻¹ except during the saturating pulses where it was 1300 µmol photons m⁻² s⁻¹. Each pulse (1 s duration) is analyzed with 10 fluorescence emission data points with a time resolution of 100 ms. Cells have undergone varying DA times after harvesting prior to application of the illumination protocol (Fig. S1).

As reported previously (Acuña et al. 2018a), the last saturation pulse is applied in darkness to test the reversibility of the signal and to ensure that we can rule out any light-induced damage, which we can for all the results reported here.

In experiments where DCMU was used, DCMU (10 µM) was added to the sample before the experiment was conducted. It is well established that this concentration completely blocks the \( Q_A \rightarrow Q_B \) electron transfer see e.g., Kaňa et al. (2012) and Kirillovsky et al. (1990).

**Additional elements of data analysis**

Here, we describe an extension of the spectral decomposition methods that have been described in Acuña et al. (2016). Acquired fluorescence spectra (\( p \) wavelengths) at \( m \) time points can be represented by an \((m \times p)\) data matrix \( \Psi \). The Singular Value Decomposition (SVD) procedure (Golub and Van Loan 1996) decomposes \( \Psi \) according to

\[
\Psi = U \cdot S \cdot V^T
\]  

Into an \((m \times m)\) matrix \( U \), where the \( m \) columns are called the left singular vectors \((lsv)\); the \((m \times p)\) diagonal matrix \( S \) whose diagonal elements \( (s_1, s_2, s_3...) \) are called the singular values and the transpose of the \((p \times p)\) matrix \( V, V^T \), where the \( p \) rows are called the right singular vectors \((rsv)\). Time-dependent characteristics are contained in the \( lsv \), while the \( rsv \) are a linear combination of the species associated spectra (SAS). The original matrix \( \Psi \) can be satisfactorily reconstructed by means of the \( n \) most significant singular vectors yielding a \((nt \times nλ)\) matrix \( \Psi_n \). In Acuña et al. (2016) data matrices of rank \( n = 2 \) were analyzed. While the core of the method remains the same, we also present data matrices that are of rank \( n = 3 \), meaning that the transformation matrix applied to the singular vectors had to be correspondingly expanded.

We seek a mathematical transformation \( \Psi_n = U \cdot S \cdot (A^{-1} \cdot A) \cdot V^T \) to resolve the SAS and their time-dependent concentrations. For a rank \( n = 3 \) matrix, \( A \) has the following form:

\[
A = \begin{pmatrix}
1 & a_{12} & a_{13} \\
& a_{21} & a_{22} & a_{23} \\
& & a_{31} & a_{32} & a_{33}
\end{pmatrix}
\]  

and the final decomposition of \( \Psi_{(n=3)} \) is expressed as a linear combination of three nt \( \times 1 \) vectors corresponding to the concentrations and the 1 \( \times nλ \) vectors corresponding to the SASₙₜₐ:

\[
\Psi_{n=3} = c_{11}SAS_{1,1} + c_{12}SAS_{2,2} + c_{13}SAS_{3,3}.
\]  

Hereafter, we will refer to these vectors using the simplified notation: \( c_1, c_2, c_3, SAS_{1,1}, SAS_{2,2}, \) and \( SAS_{3,3} \). Furthermore, a number of criteria are followed to judge whether these transformed vectors are biophysically meaningful.

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With the expansion to a rank \( n = 3 \) matrix, the number of parameters to be estimated increases; this is why, additionally to the criteria presented in our previous work (Acuña et al. 2016), we have added, for each SAS, SAS smoothness–penalties:

\[
\text{penalty}_{\text{smoothness}} = \sum_j w_j \left| \int \frac{\partial^2}{\partial \lambda^2} \text{SAS}_j(\lambda) \right|. \tag{4}
\]

This is one among other contributions to the objective function that we seek to minimize. See the SI for the explicit objective function used in this work.

Finally, each concentration value \( c_{ti} \) output at the position \( t \) that corresponds to a saturating pulse is multiplied by a parameter \( p_f \) that accounts for the difference in light intensity between the background illumination and the saturating pulse. Thus, relative concentrations that are independent of the light intensity are estimated.

**Results**

In the first section, we discuss results obtained following different protocols. First, we show results of the photosynthetically “simpler” system, ΔPSI (which does not contain any of the species depicted in Fig. 1a, d, e, g) after a series of DA periods. Then, we move on to the PSI-containing samples of the strains M55 and WT, first showing results of DCMU-treated samples with two different DA periods, then, a series of several DA periods with intact cells (without DCMU).

**ΔPSI cells**

We report analyses of fluorescence data acquired from ΔPSI cells that have undergone different dark-adaptation periods. The rank of the data matrices for all DA periods is at least three (one example is shown in Fig. S2). A decomposition analysis using the main three components has been performed. In Fig. 2, we show results for the two DA periods 1 and 34 min. The full series of six DA periods is shown in Fig. S3. Based on visual inspection of the F660/F680 ratio

![Fig. 2](image-url) Spectral decomposition of two independent experiments carried out on ΔPSI cells. c shows the first 2 min of the light protocol for the two dark-adaptation (DA) periods 1 and 34 min. For each condition, a zoom view of the first (last) pulse is depicted in a and b (d and e). The corresponding SASs (shown as insets) are depicted on c. The colored bar on top illustrates the light regime: 450 µmol photons m\(^{-2}\) s\(^{-1}\) of 590 nm light (orange) or darkness (black). In gray, the deviation in the sum of concentrations from unity. For the full data matrices and the complete series of DA periods, see Fig. S3.
and the relative amplitudes (fluorescence quantum yield) of the SASs, they are interpreted as follows: SAS2 (green) as a PB–PSII complex with fully closed RCs (f-closed) (Fig. 1h); SAS2 (red) as a PB–PSII with partially closed PSII RCs (p-closed) (Fig. 1f) and, SAS1 (black) as a PB–PSII complex with PSII RCs open (Fig. 1c) or PSII quenched by HliP (Acuña et al. 2016).

Most probably the SAS of a PB–PSII complex with PSII RCs open or quenched by HliP differ slightly, leading to a data matrix of rank 4 (see Fig. S2). Since the analysis is performed using the main three components, it is not possible to describe the data matrix fully. This manifests in the sum of concentrations shown in gray on the insets of Fig. 2c. For short DA periods, the sum of concentrations is close to flat. As DA increases, however, it becomes increasingly difficult to obtain a fully flat pattern, especially during initial phases. Also, while SAS2 and SAS3 give robust results, SAS1 because of its low $\phi_p$ provides the highest uncertainty. Even though a quenched component is consistently required to minimize the residuals, determining the exact shape and the F660/F680 ratio becomes non-trivial due to only subtle changes in the sum of concentrations profile. Certainly, the optimization routine may be compensating for the lack of the fourth component, in which case SAS1 is expected to be a mixture of the two quenched components. Assuming so, this would explain that the $c_1$ (black) decreases during the saturating pulses, as expected for open complexes (Fig. 1c), while tending towards a baseline level of around 40% (cf. Fig. 2, Table S2). These results suggest the existence of two quenched components of similar SAS, one of which is photochemically, the other non-photochemically quenched. Judging from the deviations in the sum of concentrations, the photochemically quenched component plays a role predominantly during the first 30 s.

The rationale behind the p-closure interpretation (SAS2) lies in the decreasing $c_3$ (red) during the saturating pulses, as opposed to the rising $c_3$ (green), which is formed as more light is shone onto the sample. Indeed, despite the strong Chl a signature of SAS2, the organism demonstrates the ability to close more RCs as $c_2$ decreases. Indeed, $c_2$ behaves as a partially ‘open’ species. We will discuss some possible origins of the p-closed RCs in the “Discussion” section.

As for the concentration profiles shown in Fig. 2c, it is interesting to note that the onset of the background illumination triggers a slow (several seconds) population exchange, predominantly between species SAS1 and SAS3 while SAS2 remains fairly flat. Relatively constant fluorescence levels are observed for times $t > 40$ s. Furthermore, contrary to previous observations in rank 2 analyses of ΔPSI data (Acuña et al. 2018a), the concentration profiles show no systematic changes which may be unequivocally related to the DA periods. This is probably due to a crucial difference in the growth conditions of the different batches: while in Acuña et al. (2018a) NaNO3 was used as the nitrogen source in the medium, here we have used NH4Cl. The former may result in a deficiency in nitrogen assimilation, as the reduction of NO3$^-$ to NH3 requires the direct output of PSI, i.e., reduced ferredoxin. Thus, in the absence of PSI, this reaction is hampered. Additionally, NH4Cl has a lower ‘electron cost’ than NaNO3 since NH4$^+$ can be directly used for metabolic reactions unlike NO3$^-$ which has to be reduced to NH3 first, requiring a total of eight electrons. This may lead to less of an overload in the electron transfer pathways of this mutant and, as a result, the cell may need to rely on less regulatory mechanisms during darkness explaining why different DA periods do not result in the systematic patterns observed by Acuña et al. (2018a). Nevertheless, the response is highly dynamic with none of the three concentration profiles standing out for being the predominant one.

### M55 and WT cells pre-treated with DCMU

The data matrices of experiments carried out on WT and M55 cells are of rank 3. The cells were pre-treated with DCMU and underwent two different DA periods as shown in Fig. 3i, j. The three SASs are shown in the insets. A quenched species, SAS1 (black), is present in all samples. The interpretation of SAS2 (green) is a PB–PSII–PSI complex with f-closed RCs. Here again, SAS2 (red) has a quenched 660 nm emission relative to SAS3 and this is interpreted as a PB–PSII–PSI complex with p-closed RCs. Saturating pulses are still able to close a certain fraction of PSII RCs as manifested in the decrease of both $c_1$ and $c_2$. Interestingly, the fraction that can be closed in M55 is smaller than in WT (cf. $F_S$ values in Table S1), suggesting that a fully assembled NDH-1 unit (in the presence of DCMU) contributes to efficiently alleviating the excitation pressure on PSI. As for the time profiles, we read the following: for 8-min dark-adapted samples, $c_1$ shows a slight increase suggesting that formation of the quenched species (black) is favored in darkness. This is valid for both M55 (cf. Fig. 3a, c) and WT (cf. Fig. 3e, g). Also visible on these panels is the fact that WT forms more of the quenched species as compared to M55. Note that $c_1 < c_2$ systematically for M55 as background light sets in. This ratio is reversed for the WT. Moreover, while M55 very quickly finds a steady-state level that varies little over time, WT shows an equilibration phase of ca. 10 s during which the quenched species decreases and then levels off.

### M55 cells without DCMU

Fluorescence signals from M55 cells that have undergone DA periods of 1 and 34 min are shown in Fig. 4. The full DA series of analyses of M55 cells is shown in Fig. S4, with a zoom-view of the first 35 s in Fig. S5. The SASs
are interpreted as follows: SAS3 (green) as a PB–PSII–PSI complex with moderate EET from PB to PSI and \( f \)-closed PSII RCs (see Fig. 1g), SAS2 (red) as a PB–PSII–PSI mega-complex with fast energy transfer to PSI and with \( p \)-closed RCs (see Fig. 1e), and, SAS1 (black) as a PB–PSII(–PSI) complex with \( f \)-open PSII RCs (see Fig. 1a). EET from PB to PSI most probably occurs via ApcD (Dong et al. 2009) and depending on whether it is slow or fast, more or less 680 nm light is able to escape. This would explain the strong decrease in the F660/F680 ratio, which serves as a proxy for how much excitation light was ultimately trapped by PSI. The corresponding concentration profiles also show interesting dynamics: unlike the DCMU-treated samples, the shape of the pulses resembles much less that of a step function (cf. Figs. 3a, 4a); take for instance \( c_3 \) which after reaching a maximum within the first 200–300 ms, decreases as much as 10% within the duration of the pulse. This suggests efficient electron transfer from PSII to the PQ pool and subsequent re-opening of PSII RCs. The behavior of \( c_2 \) is of particular interest: while for short DA periods, during the first pulse, \( c_2 \) tends towards zero. For DA = 8 min, \( c_2 \) averages to 5% and it increases to 14 and 19% for DA of 21 and 34 min, respectively (see Table S2). Hence, darkness favors the formation of the species SAS2. Seemingly, during the first 20 s of background light this occurs at the expense of the species SAS3 only. Thus, for longer DA periods, \( c_2 \) (\( c_3 \)) has a larger (smaller) initial concentration and, as background light sets in, it continues to gradually increase (decrease) during the first 10–20 s resulting, for long DA periods, in a momentarily larger concentration of SAS2 than SAS3. Then, \( c_3 \) tends to

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**Fig. 3** Spectral decomposition of two independent experiments with two distinct dark-adaptation (DA) periods carried out on M55 (left) and WT (right) cells with DCMU. The a and b depict a zoom view into the first and last pulse of M55 with DA none; c and d M55, DA=8 min; e and f WT, DA = none; g and h WT, DA = 8 min. The i and j show the full time profiles. The corresponding SASs and the deviation in the sum of concentrations from unity are shown as insets. The colored bar on top illustrates the light regime: 450 µmol photons m\(^{-2}\) s\(^{-1}\) of 590 nm light (orange) or darkness (black).
a $F_S$ level of ca. 40% and it restores that level quite consistently regardless of the initial conditions. We attribute this behavior to a state 2 to state 1 transition. Furthermore, the $c_1 F_S$ value is also ca. 40% (see Table S2) in all experiments i.e., the background light is only able to close 60% of the PSII RCs. In both WT and M55, DA favors the formation of a megacomplex in which fast energy transfer from PB to PSI (see Fig. 1e) is ensured (Liu et al. 2013), consequently increasing its antenna size and ultimately affecting the ratio CEF/LEF.

**WT cells without DCMU**

Fluorescence signals from WT cells that have undergone the two different DA periods 1 and 34 min are shown in Fig. 5. The full DA series of analyses of WT cells is shown in Fig. S6, with a zoom-view of the first 35 s in Fig. S7. The SASs are interpreted as in the previous section. Also in this series of experiments, the initial $c_2$ (red) increases with longer DA periods, and, as visible from the first pulse of every experiment (see Fig. S6a–f), it reaches up to 40% for DA = 34 min. As background illumination sets in, the levels of open and closed complexes (black and green) are similar. For all experiments, the concentration $c_1$ i.e., the open complexes, also shows a relatively steady level throughout the whole period of background illumination only decreasing during saturating pulses. Similarly to the M55 mutant, the initial $c_2$ and $c_3$ depend on DA.

**Discussion**

**Partial closure of PSII RCs**

As there is no energetic interaction between PSII reaction centers (Mullineaux and Holzwarth 1993), we can assume that PSII fluorescence is directly proportional to the fraction of $Q_A$; therefore, partial closure of PSII RCs means that the light pressure does not suffice to completely depopulate the PQ pool. The question is what is/are the underlying mechanism(s) that allow the PQ pool to cope with the excitation pressure during the background illumination phase?

One possibility is that a small fraction of PB–PSII–PSI complexes have one of the RCs of the PSII dimer closed
while the other remains open. In this case, the \( p \)-closed contribution, in the ΔPSI experiments, would be attributed to the species depicted in Fig. 1f; and to the species in Fig. 1d, for the PSI-containing samples.

A major conflict with this interpretation is provided by the DCMU experiments, where \( p \)-closure cannot be ascribed to PSII RCs that would have remained open. The capacity to keep PSII RCs open would have to stem from an additional pool electrons can escape to, for example, a pool of Flv2/4 proteins. Consequently, the very definition of ‘open’ would have to be extended; while hitherto ‘open’ exclusively referred to the PQ-pool ability to accept electrons from the PSII special pair, there would be two types of mechanisms that contribute to alleviating the excitation pressure on PSII RCs: \( pq \)-open when related to the capacity of the PQ-pool and \( flv \)-open when related to that of the Flv2/4 heterodimer. Thus, we hypothesize that DCMU ensures that all RCs are \( pq \)-closed, but that the alternative electron transport via the Flv2/4 pathway would however remain accessible.

Although the exact binding site and rate are still unknown for Flv2/4, the assumption in this case would have to be that \( Q_B \) is not involved in the electron transfer. Instead, we observe a quenched 660 nm emission which would align well with the observations by Bersanini et al. (2014) that point towards Flv2/4-related activity with concomitant PB co-operation. Indeed, even the properties of a PB “super core” as recently proposed by Zlenko et al. (2017) might be required. Furthermore, as manifested in the lower \( F_S \) levels of \( c_i \) in WT (Table S1), the amount of \( flv \)-closure is ca. 20% larger (for both DA periods) in the WT than it is in strain M55. This directly points at a role of NDH-1 in keeping the PQ-pool reduced (Chen 2017). Certainly, there must be additional acceptors to Flv2/4 that explain why, despite NDH-1 dysfunction, the RCs are not fully \( flv \)-closed in M55.

Spectrally resolving the fluorescence signal of DCMU pre-treated cells of a Δ\( flv2/4 \) mutant as well as measurements of expression levels of these proteins will have to support such an interpretation and test this hypothesis.

**Time dependence in M55 and WT cells without DCMU**

Without DCMU i.e., with unblocked quinone acceptors, the F660/F680 of SAS\(_2\) greatly increases, evidencing the

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**Fig. 5** Spectral decomposition of two independent experiments carried out on WT cells without DCMU. \( e \) shows the first 2 min of the light protocol for the two dark-adaptation (DA) periods 1 and 34 min. For each condition, a zoom view of the first (last) pulse is depicted in a and b (d and e). The corresponding SASs (shown as insets) are depicted on e. The colored bar on top illustrates the light regime: 450 \( \mu \)mol photons m\(^{-2}\) s\(^{-1}\) of 590 nm light (orange) or darkness (black). In gray, the deviation in the sum of concentrations from unity. For the full data matrices and the complete series of DA periods, see Fig. S6.
efficient re-opening of RCs. Also, then the time profiles, especially $c_2$ and $c_3$, are much more dynamic. Here, we lay the focus of the discussion on three different aspects: (i) the overall behavior of the $F_S$ fluorescence levels, (ii) some observations specific to the $F_M$ levels, and (iii) DA dependence of points (i) and (ii). To conclude, we will compare the results obtained with the M55 mutant and the WT.

Figures 6 and 7 show $c_2$ and $c_3$, respectively, during the first minute of the light protocol. In Fig. 6, the zoom view of $c_2$, which corresponds to the concentration of a PB–PSII–PSI

**Fig. 6** Zoom-view into the first minute of the light protocol for WT and the M55 mutant. The profile $c_2$, corresponding to megacomplexes with fast PB to PSI energy transfer is shown for all DA periods. Background illumination sets in at $t=7$ s. Immediately afterwards, three intervals are indicated: a first 300 ms after onset of background illumination; b $7.3 < t < 13$ s and c $13 s < t$. a is characterized by a rapid population increase and decrease presumably due to closure of a single RC followed by a population of $c_3$ (Fig. 7), b is characterized by a steady increase reaching a maximum and during c the profiles decline steadily again. In the case of M55 cells, DA times of 21 and 34 min display a longer period of steady increase, $B_2$, reaching a maximum value at $t=20$ s

**Fig. 7** Zoom-view into the first minute of the light protocol for WT and the M55 mutant. The profile $c_3$, corresponding to megacomplexes with slow PB to PSI energy transfer, is shown for all DA periods. The decrease in the initial concentration values correlates with increasing DA periods. This is interpreted as a state 1 to state 2 transition. Background illumination triggers a state 2 to state 1 transition.
complex with fast energy transfer to PSI, reveals *grosso modo* three different phases of the $F_S$ levels: as background illumination sets in, there is a quick equilibration that happens within 300 ms and that we refer to as phase A. During this phase, the $c_2$ profile quickly decreases until a minimum value is reached while $c_1$ shows a correlated increase, just as observed during the first data points of the saturating pulse applied in darkness (cf. Figs. S5 and S7). We conclude that this is due to opening/closing of RCs, meaning that SAS$_2$ has a hybrid nature: the PB–PSII–PSI megacomplex with fast energy transfer to PSI and $p$-closed (Fig. 1e) gets to the $f$-closed state within these first 300 ms of illumination. Thus, we see both the complexes with open RCs being closed as a fast (within ms) decrease in $c_2$ and the steady $F_S$ level (changing only in the scale of seconds) of the complexes with already closed RCs. The phase B ($7.3 < t < 13$ s) is characterized by a progressive increase of the $F_S$ level, which we interpret as the cell first adopting a configuration in which the energy transfer to PSI is increased, presumably to decrease the excitation pressure on PSII. After reaching a maximum, $c_2$ enters a phase C of steady decline attributed to a state 2 to state 1 transition, and which leads to an increase in the number of PB–PSII–PSI complexes with slow energy transfer to PSI (see Fig. 1g). Note, however, that while the phase B in the WT consistently lasts ca. 5 s for all DA periods, in M55, DA periods of 21 and 34 min result in a delayed state 2 to state 1 transition (see Fig. 6). For DA of 1, 3, 5, and 8 min the phase $B_1$ is as long as the phase B observed in WT, but for DA periods of 21 and 34 min (orange and red in Fig. 6), the phase $B_2$ extends until $t=18$ s. Seemingly, with a lacking NDH-1 unit, which impairs electron transfer into the PQ pool and CEF, the cell needs a longer time, during which the amount of megacomplexes with fast PB → PSI energy transfer further accumulates, before triggering a state 2 to state 1 transition.

Another instructive aspect of the time profiles is the behavior of the $F_M$ levels during the 1000 ms (10 × 100 ms) the saturation pulse lasts. Note that the $c_2$ profiles (Fig. 6, especially the M55 traces) display a sharp decrease during the first 100 ms of each pulse, supporting the idea of rapid closure of PSII RCs followed by an $F_M$ level that remains fairly constant during the last 900 ms—the level used to give an estimate of the $F_V$ values in Table S3. This average $F_M$ level during the pulse is in turn attributed to the relative concentration of the species. Thus, given $F_V$ values that progressively decrease, we infer indeed that the concentration of the species effectively decreases over time. Take the case for the M55 mutant after 34 min DA (red in Fig. 6) as an example: notice how the effect of the saturation pulses on the sample evolves from (times $t < 50$ s) raising the $F_M$ level above the $F_S$ level (positive $F_V$ value) to (times $t > 50$ s) not having virtually any other effect than the rapid closure of RCs within the first 100 ms of the pulse, after which the fluorescence level during saturation matches that of the $F_S$ levels during background illumination ($F_V$ value is virtually zero). Furthermore, the $F_V$ decrease we have just described is accompanied by an $F_V$ increase in the $c_2$ profile, as visible in Fig. 7 (concentration profiles of PB–PSII–PSI complexes with slow PB → PSI energy transfer): note how the DA = 34 min profile (red) systematically has lower $F_M$ levels for times $t < 50$ s but gradually increases until it, for times $t > 50$ s, has reached the same $F_M$ levels as the rest of the traces—the DA = 21 min profile (orange) being an interesting exception.

Therefore, although the $F_S$ levels always correspond to a mixture of hybrid species, the $F_M$ levels allow one to make a statement on the relative concentration of the species with closed RCs (Fig. 1e, g). Considering that the $F_S$ levels of $c_1$ remain relatively constant throughout the experiment with average $F_V$ values of $-0.35$ for both M55 and WT (cf. Table S3), we conclude that a population exchange exclusively exists between the species with fast and slow EET to PSI.

Finally, this effect on the $F_V$ values also clearly depends on the length of the DA periods. Compare, for instance, in the WT traces in Fig. 6, the $F_V$ values of the second pulse for DA = 1 and DA = 34 min: 0.04 for the former, 0.14 for the latter (cf. $F_{V1}$ values in Table S3). This means that a difference of ca. 30 min in DA has led to 10% more of the megacomplex with fast PB → PSI EET being formed. The formation of these complexes, we propose, is at the core of state transitions. Figure 7, showing the $c_1$ concentration profiles may help visualizing this idea: the first peak illustrates the decreasing concentration of megacomplexes with slow EET to PSI, which we attribute to the state 1 to state 2 transition in darkness, related to respiratory activity. Based on the stoichiometry published by Moal and Lagoutte (2012), who suggest a dense packing of pigment-protein complexes in the thylakoid membranes of *Synechocystis*, we speculate that a conformational change occurs during darkness, which would yield an even denser packing, favoring fast energy PB → PSI transfer. Physical re-distribution of the complexes could result in slowing down the EET rate to PSI.

Finally, we come to compare the M55 mutant and the WT on the basis of a summary figure, that we will briefly describe first: Fig. 8 shows, for a selected DA time (34 min), the scaled SASs (Fig. 8a) of WT and M55 as well as the corresponding time traces (Fig. 8b), accompanied by the species that we conclude that the signal originates from. For comparison, SAS$_3$ from ΔPSI is also shown, where the small F660/F680 ratio is due to the EET from PB to PSI exclusively, with the closed PSII RC responsible for the intense Chl $a$ emission that peaks at 680 nm. The spectra in Fig. 8a have been normalized relative to the 640 nm phy-cocyanin emission, which is expected to be the same in all samples. The resulting ΔPSI-SAS confirms the analyses of
in independent measurements in the ΔPSI mutant carried out in a set-up in Třeboň, CZ (see Fig. 7 in Acuña et al. 2016). As for the time profiles in Fig. 8b, we would like to point at a key difference between the M55 mutant and the WT based upon the two following observations: (i) in c2, the FV values for times t > 50 s are close to zero for the M55 (orange), whereas the WT (red) still has a positive value and (ii) in c3, the FM values of the first saturation pulse reveal that the WT (dark green) keeps a considerably lower amount of megacomplexes with slow PB → PSI energy transfer than the M55 (light green) mutant does. In other words, in darkness, the M55 mutant forms less megacomplexes with fast PB → PSI energy transfer than the WT, while it also, during background illumination, exchanges its population for slow PB → PSI transferring megacomplexes more quickly than the WT does. It is almost as if the M55 mutant resisted to having to form the megacomplex with fast PB → PSI EET even though it is capable of doing so. This would be consistent in that, given a dysfunctional NDH-1 and consequently an impaired CEF around PSI and lower input of electrons into the PQ pool, the cell would favor a lower PSI activity by keeping a higher population of PB–PSII–PSI complexes with low EET to PSI (Fig. 1g) instead of the species with high PB-to-PSI EET rate (Fig. 1e). From the use of the M55 mutant, we conclude that the decreased rate of reduction of the PQ-pool—due to the dysfunctional NDH-1 complex—has a clear effect on the state of PSII (pq-vs. flv-closure) and shows the entanglement between respiration and photosynthesis in the cyanobacterial thylakoid membrane.

Ultimately, each SAS that we have resolved in this study should be explainable as the steady-state emission spectrum of a complex from Fig. 1. Recently, we have been able to reconstruct the SASs of the ΔPSI mutant from the spectral properties of the chromophores, the energy transfer kinetics and the photochemical quenching in the PBS–PSII complex (Acuña et al. 2018b). This is a first step towards linking the here observed SASs to molecular events and structures.

**Concluding remarks**

We have presented a series of experiments during which *Synechocystis* WT and mutants thereof undergo a dark-to-light transition after being prepared under different conditions. The data matrices have been analyzed as rank 3 systems. Two major findings are presented: first, from the DCMU experiments we conclude the existence of a pool of PSII electron acceptors additional to the PQ pool and postulate that the Flv2 and Flv4 proteins could build such a pool. Thus, a PSII RC would be fully open (closed) when it is both pq- and flv-open (closed). In the DCMU experiments, the FS levels suggest PSII RCs that are in the state ‘pq-closed/flv-open’ (Fig. 3). Moreover, the PSII → Flv2/4 electron transfer mechanism (i) may require a high energy back transfer rate to the PB; and (ii) increases its efficiency with a well-functioning NDH-1 complex; which points to a possible (so far unknown) electron acceptor to the Flv2/4 heterodimer. A fundamental assumption in this reasoning implies that electrons transferred to the Flv2/4 pool bypass the secondary quinone Qb. Second, we postulate that SAS2 in Fig. 5 originates from a PB–PSII–PSI megacomplex with fast EET to PSI and pq-closed. A slow EET to PSI results in
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