New mechanism revealed for light-state transition in cyanobacterium Arthrospira platensis according to 77-K fluorescence kinetics

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The mechanisms of oxygen evolution and carbon fixation in oxygenic organisms depend on the equal distribution of excitation energy to photosystems I and II, which is regulated by a mechanism referred to as light-state transition. In this work, a novel mechanism, energy spillover from PS I to PS II referred to as “inverse spillover”, was revealed besides “mobile phycobilisome (PBS)” and the “spillover” of energy from PS II to PS I in cyanobacteria. Under continuous illumination with blue light, time-dependent 77-K fluorescence spectra demonstrated heterogeneous kinetics for the PBS and photosystem components, indicating that inverse spillover and mobile PBS work successively to regulate the excitation to a balanced distribution in cyanobacterial cells under blue light. Inverse spillover and mobile PBS occur under both 100 and 300 µmol m⁻² s⁻¹ blue-light conditions but they are accelerated under the latter.

cyanobacteria, light-state transition, blue light, 77-K fluorescence kinetics, mobile phycobilisome, inverse spillover

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induced transition from state 2 to state 1 in cyanobacterium *Arthrospira platensis* cells. The heterogeneous kinetics of PBS and photosystem components, independent of the light intensity of 100 or 300 µmol m⁻² s⁻¹, indicated a novel mechanism, energy spillover from PS I to PS II referred to as “inverse spillover”, involved in the state transition besides PBS mobility. Spectral analysis confirmed that inverse spillover was a natural mechanism and not an artificial result arising from spectral normalization.

1 Materials and methods

1.1 Culture and growth conditions

*Arthrospira platensis* FACHB-900 cells were cultured in AB medium (pH 9.0, 1 L) at 25°C with continuous shaking and illumination by three 40-W fluorescence lamps. Ten-day cultures were harvested by centrifugation, washed and resuspended in fresh growth medium. Cells of the same generation were used in all experiments.

1.2 State transitions

For transition from state 2 to state 1, the cells were previously induced to state 2 by orange light (Ditric Optics 580-nm long-pass and 600-nm short-pass filter) of 20 µmol m⁻² s⁻¹ and were then immediately brought to state 1 by blue light (Ditric Optics 410-nm long-pass and 460-nm short-pass filter) of 100 or 300 µmol m⁻² s⁻¹. Afterward, the cells were immediately put into liquid nitrogen for measurement of 77-K fluorescence spectra. The light intensity was measured by an ultraviolet radiometer.

1.3 Spectral measurements

Fluorescence emission spectra at a temperature of 77 K were recorded by an F7000 spectrofluorometer (Hitachi, Japan). The spectra presented in this work are the averages of five independent measurements. The excitation and emission slit widths were set to 5 nm. The chlorophyll concentration of the cells used for fluorescence measurement was adjusted to no more than 5 µg (Chl a) mL⁻¹ estimated from the absorbance at 665 nm in methanol extracts [14].

2 Results and discussion

The 77-K fluorescence difference spectra were recorded over time when the state-2 cells were continuously illuminated by blue light of 100 µmol m⁻² s⁻¹, as shown in Figure 1(a). From the spectra, the time-dependent fluorescence amplitudes for PBS rods (C-phycocyanin; C-PC), PBS core (allophycocyanin; APC) and PS I and PS II components were derived, as shown in Figure 1(b). A remarkable feature is that the PBS fluorescence components do not always keep pace with the fluorescence components of the photosystems, which completely differs from the case for orange-light-induced transition from state 1 to state 2 [13]. Figure 1(b) shows that the transition from state 2 to state 1 took 8 min, at which time the PS I fluorescence and PS II fluorescence became saturated. However, the PBS components became invariable at 4 min, at about half way into the transition to state 1, suggesting that PBSs did not contribute further after that time. Apparently, the state transition undergoes two successive sub-processes. In the first 4 min, the increase in C-PC and PS II fluorescence and the decrease in APC and PS I fluorescence suggests that PBSs move to PS II, which is a typical feature of transition from state 2 to state 1 [11,12], and the homogeneous kinetics of the PBS and photosystem fluorescence indicate that the state transition was regulated by mobile PBS. After 4 min, PS II and PS I components increased and decreased respectively, independent of the PBS components, suggesting energy spillover from PS I to PS II, which is referred to as inverse spillover in this work.

![Figure 1](attachment:image_url)
In the case that the cyanobacterium cells were illuminated by blue light of 300 µmol m⁻² s⁻¹, the time-dependent difference 77-K fluorescence spectra and the component amplitudes are shown in Figure 2. The heterogeneous kinetics for PBS and photosystem components is similar to those in Figure 1. However, quantitatively, PS I and PS II fluorescence components became saturated at 2 min while the PBS components became saturated at 1 min, suggesting that the state transition was accelerated compared with that under 100 µmol m⁻² s⁻¹ illumination. It was observed that the frequency of fluorescence oscillation or the state transition time for cyanobacterium *Synechocystis* PCC6803 cells was a function of orange-light intensity, which is ascribed to the acceleration of PBS movement at higher intensity [13]. The kinetics demonstrated in Figures 1 and 2 suggest that not only PBS movement but also inverse spillover was accelerated at higher light intensity, in turn suggesting that the state transition rate is a function of light intensity irrespective of whether a single mechanism or multiple mechanisms are involved. Importantly, this implies that cyanobacteria sense any imbalance of the excitation distribution and respond to it via state transition. Previously, it was proposed that state transition was a natural behavior of cyanobacteria under solar irradiation and not short-term acclimation under an artificial light condition [13], which is further confirmed by this work.

The fluorescence kinetics in Figures 1 and 2 suggest that mobile PBS and inverse spillover worked successively during the state transition. However, it might be argued that the heterogeneous kinetics were an artificial result of the normalization to 712 nm and not a natural behavior of the photo-functional entities. To clarify this, the spectra at 100 µmol m⁻² s⁻¹ were alternatively normalized to several other wavelengths, for which the difference spectra and the time-dependent amplitudes are shown in Figure 3.

In Figure 3, the normalization wavelengths were selected quite arbitrarily for emission peaks (648 nm for C-PC or 730 nm for PS I), a valley (674.8 nm) and a position between a peak and valley (688 nm). Noticeably, the fluorescence kinetics reproduce the basic feature in Figure 1, confirming that heterogeneous kinetics reveal a novel mechanism and are independent of normalization. Normalization is generally acceptable for 77-K fluorescence spectra because it eliminates linearly correlated fluctuations and retains linearly independent “signals”. In this sense, normalization to any wavelength in a spectrum is theoretically reasonable. However, normalization to a peak will conceal some components and amplify others, as shown in Figure 3(a)–(c). Consequently, normalization to a valley wavelength is a better choice.

Until now, it has been believed that mobile PBS and spillover of energy from PS II to PS I were involved in light-state transitions in cyanobacteria, whereas inverse spillover, the energy transfer from PS I to PS II, was revealed as an additive mechanism in the current work. In principle, light-state transition is a mechanism of inter-regulation between energetic and electronic states in oxygenic photosynthesis. At the molecular level, a complex mechanism is required to explain energy transfer from not only the photosynthetic antenna to the photosystems but also from one photosystem to another [15,16] and electron transfer within PS II or PS I and between them. Specifically, under blue light, it is PS I that is overexcited while PS II is less excited; therefore, it is reasonable that there is energy spillover from PS I to PS II according to the energy-balance rule. Considering that the PS I energy level is lower than the PS II energy level, how the “uphill” transfer take place remains unknown. In fact, blue light resulting in overexcitation of PS I leads to excess oxidized reaction-center P700⁺ owing to a lack of PS II electrons, which may correlate to the inverse spillover in some way. It has been proposed that light-state transition in cyanobacteria might involve the movement of membrane complexes and/or changes in the oligomerization states of the complexes.

![Figure 2](image-url)  
**Figure 2**  
Difference fluorescence spectra (a) and plots of amplitudes of the four components against time (b) for state-2 cells illuminated by blue light of 300 µmol m⁻² s⁻¹, as derived from 77-K fluorescence emission spectra (inset). Results are obtained for excitation at 580 nm and are normalized to 712 nm.
(including PS II and PS I) [17,18]. It was found previously that PS I oligomerization states are reversibly variable under light-to-dark and dark-to-light transitions [12,19]. Apparently, either the movement of PS I or PS II or dissociation of PS I trimer into monomers may shorten the distance between PS I and PS II, amplify the effective concentration of PS I and enhance the spectral overlap of the PS I emission with PS II absorption, which certainly enhances the probability of uphill energy transfer. However, the molecular mechanism that triggers inverse spillover has yet to be revealed.

3 Conclusion

In the current work, a novel mechanism, energy spillover from PS I to PS II referred to as inverse spillover, was revealed for blue-light-induced state transition in cyanobacterium *Arthrospira platensis* FACHB-900 cells. Under blue light, 77-K fluorescence kinetics suggest that mobile PBS and inverse spillover worked successively and both were accelerated under a light condition of 300 µmol m⁻² s⁻¹ compared with those under a light condition of 100 µmol m⁻² s⁻¹. The result suggests that cyanobacteria are able to balance the excitation distribution under various light conditions but the mechanisms involved may be multiple. Further research is necessary to clarify how inverse spillover occurs under blue light.

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