REVIEW

Chlorophyll fluorescence emission spectroscopy of oxygenic organisms at 77 K

J.J. LAMB*, G. RØKKE**, and M.F. HOHMANN-MARRIOTT**+:

Department of Electronic Systems & ENERSENSE, NTNU, Trondheim, Norway*
Department of Biotechnology & CenTroN for Synthetic Biology, NTNU, Trondheim, Norway**

Abstract

Photosynthetic fluorescence emission spectra measurement at the temperature of 77 K (−196°C) is an often-used technique in photosynthesis research. At low temperature, biochemical and physiological processes that modulate fluorescence are mostly abolished, and the fluorescence emission of both PSI and PSII become easily distinguishable. Here we briefly review the history of low-temperature chlorophyll fluorescence methods and the characteristics of the acquired emission spectra in oxygen-producing organisms. We discuss the contribution of different photosynthetic complexes and physiological processes to fluorescence emission at 77 K in cyanobacteria, green algae, heterokont algae, and plants. Furthermore, we describe practical aspects for obtaining and presenting 77 K fluorescence spectra.

Additional key words: fluorescence; low temperature; photosynthesis.

Introduction

Historical background

“Chlorophyll fluorescence is red and beautiful” (Govindjee 1995) and has fascinated researchers for a long time (see reviews by Govindjee 1995, 2004). Chlorophyll (Chl) extracts prepared from leaves possess a very intense red fluorescence that caught the attention of Brewster in the 1830's (Brewster 1834). Correlations between the weaker in vivo Chl fluorescence emission and photosynthetic performance have been suggested (Müller 1887, as referenced by Govindjee 1995, 2004), but Kautsky and Hirsch were the first to unambiguously relate Chl fluorescence yield to the rate of photosynthesis (Kautsky and Hirsch 1931). The utility of Chl fluorescence for exploring photosynthesis was consequently demonstrated in many different photosynthetic organisms. Many breakthroughs in our understanding of photosynthesis are the result of researchers’ creativity to develop techniques to resolve the spectral and temporal characteristics of Chl fluorescence in intact photosynthetic systems and by purifying individual components thereof.

One Chl fluorescence-based technique, which has been widely adopted, was developed by Seymour Steven Brody while working on his Ph. D. in the Rabinowitch laboratory at the University of Illinois at Urbana Champaign (Brody 1958, Rabinowitch and Govindjee 1969). It had been previously demonstrated that lowering the temperatures sharpens spectral characteristics, such as absorption and fluorescence, due to the loss of intramolecular vibrations. However, the improved spectral resolution was not the primary motivation for Brody to investigate fluorescence characteristics of the green algae Chlorella at liquid nitrogen temperatures. Brody wanted to eliminate biochemical and physiological processes (Hirsch and Rich

Received 27 October 2017, accepted 10 January 2018, published as online-first 8 February 2018.
*Corresponding author; e-mail: martin.hohmann-marriott@ntnu.no

Abbreviations: Chl – chlorophyll; LHC – light-harvesting complex; PSI – photosystem I; PSII – photosystem II.

This review is dedicated to Govindjee. In addition to Govindjee’s original contributions to the field, we, the authors, are also very thankful to Govindjee for sharing historical context and personal connections, which contributed to researchers making their discoveries. In the case of 77 K fluorescence, the topic of this review, Govindjee was not only witness to its first implementation, but also went on to refine and extend the interpretation of this powerful measuring technique. The following short historical introduction into 77 K fluorescence often refers to information obtained from publications by Govindjee and coworkers.

Acknowledgements: Martin Hohmann-Marriott acknowledges support from the Research Council Norway (grant number 240741). Gunvor Røkke’s research was supported by a PhD fellowship from the NT faculty of the Norwegian University of Science and Technology – NTNU. Jacob Lamb acknowledges the support from the ENERSENSE research initiative, and his research was supported by a postdoctoral fellowship from the Norwegian University of Science and Technology – NTNU.

© The Author(s). This article is published with open access at link.springer.com
2010), and thereby gained direct insights into excitation and electron transfer processes involved in photosynthesis using Chl fluorescence as a reporter. Around the same time during which Brody investigated fluorescence emission by the green alga *Chlorella* (Brody 1958), Litvin and Krasnovsky (1957) investigated fluorescence emission of plant material at 77 K.

One immediate finding of these measurements was that in addition to the main Chl fluorescence band at 685 nm, which is readily observed at room temperature, a second emission band with peak intensity at 720 nm becomes more prominent at lower temperatures (Fig. 1) (Brody 1958). Frederick Cho and Govindjee demonstrated that at even lower temperatures - at liquid helium temperatures (4 K) (Rabinowitch and Govindjee 1969, Cho 1970a, b, Cho and Spencer 1966) a third emission band at 695 nm can be separated (Fig. 2).

Fig. 1. Chlorophyll fluorescence emission by *Chlorella* cells at room temperature and –193°C. The original figure legend: “Fluorescence spectra of *Chlorella* at room temperature (crosses) and –193°C (open circles). The fluorescence intensities indicated are the same for both curves. The decrease in fluorescence yield at 690 mµ is probably due to the increased scattering of the exciting and fluorescent light”.

Remarks: The fluorescence yield at 690 nm is expected to be about two times larger at –196°C compared to room temperature fluorescence yield, as suggested by Brody in the original figure legend. The term mµ is a historical notation that is equivalent with nanometer (nm).

This publication represents the first report of the increased long-wavelength chlorophyll fluorescence emission at 77 K, which has been established to be mainly associated with PSI. Data was digitized from Brody (1958).

Fig. 2. Chlorophyll fluorescence emission of *Chlorella pyrenoidosa* as a function of temperature
(A): Chlorophyll fluorescence emission at specific wavelengths as a function of temperature in *Chlorella pyrenoidosa* as a function. The excitation light wavelength was 400 nm. The original figure legend: “Emission at 685 mµ (F685), at 696 mµ (F696), and at 760 mµ (F738; also referred to as F720 in the text) as a function of temperature (~–196 to 20°C).” Panel A has been digitized from Cho et al. (Cho et al. 1966).

(B): Chlorophyll fluorescence emission spectra of *Chlorella pyrenoidosa* as a function of temperature (~–269, –247, –233, –218, –196°C). Excitation light wavelength was 485 nm. The original figure description: “Fig. 1. shows the emission spectra measured in the 680–720 mµ range for several temperatures (~–269, –247, –233, –218, and –196°C). Upon warming the sample, a shift from 695 mµ to 699 mµ in the location of the peak of the “F697.5” band is noticeable. As the temperature decreases from -196 to -269°C, the fluorescence increases steadily. The total intensity at –269°C is about 2 times that at –196°C. The profile of the fluorescence spectrum at –196°C shows clearly the F689, F697.5, and F725 bands; at –269°C, the F689 appears as a very sharp band and it dominates both F697.5 and 725, the F697.5 band shows only as a shoulder at –269°C.” Panel A has been digitized from Govindjee and Yang (1966).

Remarks: The term mµ is a historical notation that is equivalent with nanometer (nm). This figure was used to demonstrate that the 738 nm (PSI) emission band is not a reabsorption artifact.

Associating these fluorescence emission bands with the concept of two types of photosystems (PSI and PSII) (Emerson 1957, Govindjee et al. 1960, Govindjee 1963) as part of the Z-scheme of photosynthesis, was achieved by the work of a community of researchers (reviewed by Govindjee and Björn 2017). So it was recognized that in red algae, which have a spectrally distinct light-harvesting system, preferentially exciting these light-harvesting systems, the phycobilisomes, increases the fluorescence emission at 685/695 nm (Krey 1966). Exciting Chl b in higher plants also increased the fluorescence emission at 685/695 nm, thus suggesting that Chl b-containing light-harvesting
systems are associated with PSII (Govindjee and Yang 1966, Rabinowitch and Govindjee 1969). Fractionation of spinach thylakoids into PSII-enriched and PSI-enriched fractions confirmed the assignments of Chl fluorescence emission bands at 685/695 nm to PSII and the emission band at 720 nm to PSI (Boardman et al. 1966).

That the number of light-absorbing pigments associated with PSI and PSII can be modulated in a physiologically relevant manner was shown by Murata (1969, 1970), in red algae using 77 K Chl fluorescence measurements. These “state transitions,” are not unique to red algae, but have also been observed in other organism groups. Bonaventura and Myers (1969) showed dynamic adjustment of energy distribution between the photosystems in a green alga using room temperature fluorescence paired with oxygen measurements. However, 77 K fluorescence emission at –196°C was used more frequently in the past, including “Chl fluorescence at liquid nitrogen temperatures” and “Chl fluorescence at –196°C”.

We have limited this review to steady-state fluorescence emission at 77 K and discuss the underlying physical processes, spectra, and its components. In addition to steady-state Chl fluorescence measurements at 77 K, many powerful techniques that use low-temperature Chl fluorescence emission have been developed, including time-resolved techniques (Strasser et al. 2004). Our focus here is on the physiological interpretation of information obtained from 77 K Chl fluorescence spectra. For this, we provide a visual overview of spectral features and physiological responses in different organism groups.

Physical background

Excitation of molecules

When molecules interact with light, the energy contained in the photons can be used to transfer an electron to an energetically higher orbital, thus generating a molecule in an excited state. Molecules with an extended conjugated system of bonds are likely to interact with photons in the visible spectrum. All Chls possess extended conjugated bond systems and interact with photons centered at two wavelengths that represent the first and second excited state of the Chl. The absorption band at higher energy is often termed “the blue absorption band”, “B band” (consisting of several states) or “Soret band”, whereas the lower energy absorption band can be referred to as the “red absorption band” or the “Q band” (consisting of two states, Qx and Qy) (Gouterman et al. 1963) (Fig. 3).

An excited Chl can return to the ground state through dissipating the energy difference between excited state and ground state through different modes, which compete with one another. The second excited state of Chls is converted efficiently to the first excited state, and the energy difference is released as heat. The energy of the first excited state is either transferred to another pigment, or converted into: (1) heat, (2) chemical energy by driving charge separation, (3) a long-lived triplet state through a reversal of the spin of the electron, or (4) the emission of a photon. The energy of the emitted photon corresponds to the energy difference between the lower vibronic sublevels of the first excited state and ground state. This “Stokes-shifted” photon is the fluorescence that is the basis of many spectroscopic techniques, including 77 K fluorescence emission analysis.

The fluorescence characteristics of photosynthetic machinery are dependent on the emission characteristics of individual Chls and the cooperative excitation-coupling network these Chl form. A single Chl molecule possesses a fluorescence emission band, which reflects the transition from the first excited state to the ground state (Clayton 1980). This fluorescence band around 685 nm is broadened at room temperature as vibronic sublevels with higher energy are populated. Energy levels of excited states of individual Chls are also influenced by the protein and lipid environment that a Chl experiences. In vivo, the second broad fluorescence emission band around 735–740 nm is thought to be due to an increased emission wavelength through self-absorption at shorter wavelengths (Franck et al. 2002).

The overall emission of photosynthetic systems is a combination of main and vibronic sublevel emission, environment of individual Chls, and the energetic landscape of the Chl collective. For example, in plants, Chls associated with the PSII reaction center fluoresce at 685 and 695 nm – due to the main transition emission, and around 735–740 nm – due self-absorption enhanced fluorescence in this region. In intact PSII, Chls in the peripheral antennae, when isolated fluoresce at 680 nm, are coupled efficiently to the Chls within the PSII reaction center. Thus, only a small amount of 680 nm fluorescence emission is observed in intact systems, as the low energy Chls within the PSII reaction center (with a main emission at 685 and 695 nm) emit fluorescence. Furthermore, fluorescence emission from vibronic sublevels of Chls within the peripheral antennae and reaction center is emitted as a broad band centered at 730 nm. PSI has additional fluorescence emission that combines with that of PSI, light-harvesting complexes, and vibronic sublevel emission to generate the combined fluorescence emission pattern of an organism (Fig. 4).
Fig. 3. Diagram relating the energy levels with the absorption spectrum and the fluorescence of chlorophyll (Chl) a. The data on energy levels in Chl a was obtained from Frank et al. (1994), absorption data for Chl a was obtained from Chen and Blankenship (2011), and the fluorescence data was retrieved from http://omlc.org/spectra/PhotochemCAD/html/122.html.

Table 1 gives an overview of major emission bands observed in various photosynthetic materials.

Excitation transfer
In whole, intact photosynthetic systems, Chl fluorescence represents a tiny fraction of all excitation energy captured by pigments in intact photosynthetic organisms, as energy is efficiently channeled into charge separation (Hillier and Babcock 2001). The bulk of the light-absorbing molecules in known photosynthetic organisms – except Heliobacteria – are not positioned within the photosynthetic reaction centers, but in separate peripheral light-harvesting systems. Light energy, absorbed by pigments in these light-harvesting systems is transferred via other pigments within the light-harvesting system ( Förster 1965, Şener et al. 2011) to the reaction center antennae and finally to the reaction center core, where the excitation energy is used to accomplish a charge separation event.

Pigments
Oxygenic organisms employ three main classes of pigments for light harvesting: Chls, phycobilins, and carotenoids.

Chls and phycobilins possess a large absorption cross-section and are very fluorescent – features that make them well suited to participate in the Förster-type exchange of excitation. Indeed, the consortia of Chls (i.e., the light-harvesting systems of plants) and phycobilins (i.e., phycobilisomes, the light-harvesting system of cyanobacteria) are very efficient in channeling excitation to the Chl-containing reaction centers.

Carotenoids absorb light efficiently between 450–550 nm and display tiny fluorescence yields (Frank et al. 2006). Thus, the efficient excitation transfer between carotenoids and Chls cannot be understood by a Förster-type excitation transfer theory. Instead, energy transfer between carotenoids and Chls may be understood to be mediated by the singlet–singlet excitation energy transfer process (Owens 1992, Young and Frank 1996). Table 2 gives an overview of common pigments found in various organisms.

Fluorescence emission spectroscopy
A way to assess the interaction of light-harvesting systems with the reaction centers is to expose a sample to monochromatic light centered on a specific pigment absorption maximum and monitor the fluorescence emission. In cyanobacteria and red algae, phycobilins have an absorption spectrum that is distinct from the Chls of the reaction centers. For this reason, many early insights into
photosynthesis were obtained in phycobilin-containing organisms. Alga and plants expand the range of absorbed photons with Chls b and c, which are distinct from the Chl a of the reaction centers. The association of light-harvesting systems with the reaction centers can, therefore, be probed by exciting the light-harvesting system pigments and monitoring the fluorescence emission that is specific for PSII and PSI.

**Room temperature fluorescence emission of plants**

In whole, intact photosynthetic systems, only 1–2% of excitation energy captured is lost as fluorescence (Maxwell and Johnson 2000). At room temperature, PSII emits 90% of fluorescence while PSI contributes the remaining 10% (Govindjee 1995). Due to the low room temperature fluorescence emission of PSI compared to PSII, the overlapping fluorescence spectra of the main fluorescence emission band of Chls associated with PSI, and vibronic emission of Chls associated with PSII around 720–730 nm (Franck et al. 2002), it is difficult to assess the magnitude of PSI fluorescence emission. Another factor that makes the assignment of fluorescence to PSI and PSII difficult at room temperature is the modulation of fluorescence yield due to the reduction state of the electron acceptors within the photosystems. The prime modulator of overall Chl fluorescence at room temperature is the reduction state of the first stable electron acceptor of PSII, a quinone named Qa (Duysens and Sweers 1963). In contrast, there is almost no modulation of PSI fluorescence yield by the reduction state of the electron acceptors of PSI at room temperature, which is due to the efficient quenching of excitation by P700+ (Dau 1994a, b).

**Fluorescence at low temperatures**

Fluorescence emission spectra at 77 K offer a key advantage over room temperature measurements, as modulation caused by physiological acclimations and biochemical reactions are eliminated. Compared to room temperature, the 77 K fluorescence yield of the PSII core complex is about two times higher, and fluorescence yield of PSI increases by a factor of around 20 (Mukerji and Sauer 1988, Dekker et al. 1995), thus PSII and PSI fluorescence signatures become discernable from each other.

Electron transport reactions, apart from those involved in primary charge separation and charge stabilization within the photosystems, are inhibited at 77 K. Electrons accumulate on the acceptor side of PSII, and on the acceptor side of a fraction of PSI (Sétil et al. 1984, Schlodder et al. 1998) after brief illumination, and remain at the electron acceptors, even during low-intensity illumination, such as applied during the collection of fluorescence emission spectra. However, when samples are dark-adapted and remain unexposed to light after being frozen freezing at 77 K, fluorescence yield changes upon illumination at 77 K reflecting the electron transport within the reaction centers (Ley and Butler 1980). Lowering the temperature to 77 K also leads to a decrease in energy and thus electrons occupy lower vibrational levels, reflected by a sharpening of the fluorescence emission band compared to room temperature. Excitations are more likely trapped on the longer wavelength Chls, as the energy of the vibrational ground state is higher in the shorter-wavelength Chls. Thus, at temperatures lower than 77 K, more fluorescence within PSII is emitted at 695 nm (in addition to 685 nm), PSII and PSI fluorescence emission shifts to longer wavelengths, and emission of additional low energy pigments increase. For example, in plants fluorescence of Chls in LHCII complexes (Rabinowitch and Govindjee 1969, Cho and Spencer 1966) are thought to trap excitation below liquid nitrogen temperature increasingly, thereby contributing to fluorescence emission at 680 nm (Rijgersberg et al. 1979).

Having a discernible PSI-specific fluorescence signal enables the investigation of the association of light-harvesting systems with PSI or PSII, by exciting pigments (such as Chl b, c, carotenoids, and phycobilins) that are preferentially located in the peripheral light-harvesting systems. Furthermore, direct excitation of Chl a and the resulting fluorescence emission patterns can be used to determine the stoichiometries of the photosystems (Murakami 1997), and has provided major insights into physiological adaptations of photosynthetic organisms.

**Technical and practical aspects for 77 K fluorescence measurements**

Several methodologies have been developed to obtain 77 K fluorescence spectra with a variety of instruments and sample preparation procedures. The following section contains an overview of these techniques and procedures.

**Fluorometer and measurements**

In a fluorometer that is capable of quantifying fluorescence emission, monochromatic light is used to excite a sample at the desired wavelength. The fluorescence is detected at an angle of 90° to the incident excitation beam. The illumination and detection wavelength can be modulated by optical filters in combination with monochromators. For obtaining 77 K fluorescence emission and excitation spectra, instruments that use monochromators in combination with full spectrum excitation light sources and photomultiplier tubes have traditionally been used (Hipkins and Baker 1985). However, for obtaining fluorescence emission measurements, a less costly instrument that uses LEDs as an illumination source and a CCD-array spectrometers can also be employed (Lamb et al. 2015; Fig. 5).

To obtain useful spectral data that lacks artifacts, care must be taken to avoid spectral distortions by self-shading (see Fig. 6) (Govindjee and Yang 1966, Weis 1985). A dilution series of the sample can be used to find the concentrations where spectral distortions occur. Comparing these dilution spectra reveals that self-shading
Table 1. Assignment of fluorescence emission signals at 77 K.

<table>
<thead>
<tr>
<th>Band assignment</th>
<th>Origin of 77 K fluorescence</th>
<th>Key reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>F640</td>
<td>phycoerythrin</td>
<td>Sobiechowska-Sasim et al. 2014</td>
</tr>
<tr>
<td>F645</td>
<td>phycocyanin</td>
<td>Sobiechowska-Sasim et al. 2014</td>
</tr>
<tr>
<td>F660</td>
<td>allophycocyanin</td>
<td>Sobiechowska-Sasim et al. 2014</td>
</tr>
<tr>
<td>F680</td>
<td>LCM, Lhca, Lhcb, Lhef (Lhev), Lhx, RedCLH</td>
<td>Rijgersberg et al. 1979</td>
</tr>
<tr>
<td>F685</td>
<td>PSII core: CP43, IsiA</td>
<td>Andrizhiyevskaya et al. 2005</td>
</tr>
<tr>
<td>F695</td>
<td>PSII core: CP47</td>
<td>Andrizhiyevskaya et al. 2005</td>
</tr>
<tr>
<td>F710–720</td>
<td>PSI reaction center antenna (cyanobacteria)</td>
<td>Yamamoto et al. 2013</td>
</tr>
<tr>
<td>F720–760</td>
<td>PSI peripheral antenna (plants) chlorophyll vibronic sublevels</td>
<td>Karapetyan et al. 2014</td>
</tr>
</tbody>
</table>

Table 2. Distribution of photosynthetic pigments. ¹Many cyanobacterial species contain Chl a as their only chlorophyll-type pigment. ²Heterokont algae.

<table>
<thead>
<tr>
<th>Blins</th>
<th>Chl a</th>
<th>Chl b</th>
<th>Chl c</th>
<th>Chl d,f</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cyanobacteria</td>
<td>√</td>
<td></td>
<td>√</td>
<td></td>
</tr>
<tr>
<td>Red algae</td>
<td>√</td>
<td></td>
<td></td>
<td>√</td>
</tr>
<tr>
<td>Diatoms²</td>
<td></td>
<td></td>
<td>√</td>
<td></td>
</tr>
<tr>
<td>Eustigmatophyta²</td>
<td>√</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Brown algae²</td>
<td></td>
<td>√</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dinoflagellates²</td>
<td></td>
<td>√</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cryptomonads</td>
<td></td>
<td></td>
<td>√</td>
<td></td>
</tr>
<tr>
<td>Green algae</td>
<td>√</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Plants (Viridiplantae)</td>
<td>√</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Fig. 5. Custom-built fluorometer. Images of a custom-built instrument setup published by Lamb et al. (2015). During data acquisition the instrument is covered by a black cloth, which has been omitted to show the setup. A: Dewar that holds the samples immersed in liquid nitrogen. The dewar part that orientates the sample in regards to the LED and detection fiber is supported by a white piece of plastic at the bottom, B: a 3D-printed housing, C: an excitation LED, D: a long-pass filter, and E: an optical fiber detector (at right angle to excitation LED). (Lamb et al. 2015).

leads to decrease in fluorescence yield of the bands associated with PSII (685/695 nm) compared to the PSI band, and a red shift of fluorescence emission spectra. With the sensitivity of modern instruments, compromises between spectra quality and signal-to-noise levels are unlikely to occur, but strategies and models for compensating for unavoidable self-absorption artifacts in tissues have been developed (Cordón and Lagorio 2006).

Optical filters can be employed to avoid spectral artifacts that are inherent to the optical components of the instruments and the characteristics of the sample. A (narrow) band pass filter can be used to eliminate stray light that has been transmitted by the excitation monochromator. Another artifact that can be removed by inserting a (narrow) band pass filter is the elimination of photons due to harmonic transmission by the excitation monochromators (i.e., photons with 1.5-times the excitation wavelength and 2-times the excitation wavelength may also pass the monochromators). This 1.5-times wavelength harmonic is of relevance when Chl is excited by 435-nm light (resulting in an artefactual peak at 652.5 nm), whereas the 2-times wavelength transmission artifact is of relevance if Chl is excited in the UV part of the spectrum.

Furthermore, a long-pass filter can be used to avoid excitation light, which is scattered by the sample, from entering the detector (Lakowicz 1983a, b).

Data processing of fluorescence spectra

Before measurements are reported, it is necessary to correct the spectra, and explicitly state the measuring configuration used. Corrections to spectra are necessary due to spectral features of the illuminating light source, as well as the spectral response of the detector used. Useful instructions and background for obtaining corrected spectra have been provided before (Lakowicz 1983a,b; Hofstraat et al. 1992).

A feature of most Chl fluorescence data is that they are reported in “relative fluorescence units.” The reason that most of the time non-quantified spectra, which report the fluorescence yield per absorbed photon, are reported, is the impracticality imposed by the lack of suitable equipment, as well as differences in the optical properties of samples, including scattering. Other factors that limit quantitative reporting of Chl fluorescence are changes in optical properties of the samples due to freezing, and difficulty in attaining reproducible orientation of the sample within the
The 738 nm (PSI) emission band is not a reabsorption artifact. This figure was used to demonstrate that reabsorption within a single chloroplast fragment.

Further information from the text explaining the data: “The concentration is expressed on the graphs in per cent absorption at 680 μ, ranging from about 100°-~ to 2%. These experiments show that reabsorption of the 696 μ and 685 μ emission bands is insignificant at less than 5% absorption at 680 μ, but becomes significant at the higher concentrations. Thus, the 738 μ band must be a "real" band (i.e. not due to reabsorption of the main band). (Further quantitative investigation is, however, required because our present data have not been corrected for reabsorption within a single chloroplast fragment.)”

This figure has been digitized from Govindjee and Yang (1966). Remarks: The term μ is a historical notation that is equivalent with nanometer (nm). This figure was used to demonstrate that the 738 nm (PSI) emission band is not a reabsorption artifact.

Dewars and cryostats

The dewar is at the heart of 77 K fluorescence measurements as it maintains the sample at low temperatures. Named after its inventor Sir James Dewar, a dewar is a specialized vacuum flask that houses low-temperature liquids, such as nitrogen or helium. For 77 K fluorescence measurements, the dewar takes the form of an open vessel, where a vacuum separates two glass walls. The dewar consists of a larger reservoir that houses the majority of liquid nitrogen and a thinner extension, which orients the sample within the fluorometer. Samples that have already been frozen in liquid nitrogen are transferred to the dewar and maintained at a temperature of 77 K for the duration of the measurement. Dewars that have been designed to hold tubes for EPR experiments and other spectrophotometric measurements are commercially available. However, researchers who are lucky enough to have access to a good glassblower may find custom-designed dewars to be a cheaper alternative.

In addition to the relatively simple dewar-based setup, which can be adapted to many fluorometers, highly sophisticated instruments that interface with a defined suite of devices is also available. These instruments use cryostats, which allow precise temperature control down to liquid helium temperatures (4°C) and have optical interfaces for specific instruments or fiber optics.

Sample tubes

Unlike the situation for spectrophotometers, no standardized optical path length is used for 77 K experiments. In some laboratories, dewars made for EPR experiments are used, and thus matching EPR tubes with an inner diameter of 3–5 mm are used for measurements. An alternative to these commercial, closed-end tubes is an open-ended tube design, which provides a very cost-effective and practical alternative. The liquid samples can enter the tube by simply lowering the tube into the sample liquid. Once the tube is filled with the desired amount, the user seals the tube using their thumb and then freezes the sample in liquid nitrogen. The frozen liquid may escape from an open tube during measurements, and it is good practice to measure Chl fluorescence without a sample to assess contamination. In our experience, contamination does not occur even after measuring up to one hundred tubes. The real advantage of the open tube design is the ease of cleaning the tube so they can be reused continuously. When tubes are stored in a liquid nitrogen dewar for a long time, a film of ice can form, which should be removed with tissue paper, before inserting the sample into the dewar for measurements.

Fluorescein

An approach to compare the amplitude of fluorescence spectra is to introduce a fluorescent molecule to the sample at a known concentration. This allows the spectra from different samples to be normalized irrespective of their sample preparation properties, thus providing a quantitative insight into the changes between samples. One such molecule that can be included in the sample is fluorescein (Sjöback et al. 1995). When light is used to excite Chl a, it has an excitation maximum between 435–460 nm, and this light also excites fluorescein. The emission maximum of fluorescein is then observed at 508 nm (El Bissati et al. 2000), 545 nm (Walters and Horton 1991), or 535 nm (Krause et al. 1983, Krause and Weis 1984).

Glycerol

Glycerol has frequently been added to samples before freezing to 77 K as a cryoprotectant. This treatment reduces the formation of ice crystals during the freezing
procedure, therefore reducing damage to the sample and preventing light scattering. Unfortunately, glycerol treatment has been found to alter 77 K spectra in cyanobacteria (Mullineaux 1994) by reducing the affinity of phycobilisomes to PSII, resulting in increased fluorescence emission by the phycobilisomes (Mullineaux 1994). A detailed study on the effect of glycerol on the photosynthetic machinery of the cyanobacterium Spirulina platensis (Li et al. 2007) revealed that glycerol weakens energy transfer of the terminal phycobilisome emitter (L_Ca) to the reaction center of PSII as well as interfering with energy transfer between phycobilins.

**Sample preparation**

Samples for measurement of fluorescence at 77 K can either be suspensions, such as algae, extracted thylakoids, and chloroplasts, or photosynthetic tissues, such as whole leaves. Dependent on the sample type, different sample preparation methods for measurements at 77 K have been developed. For standardization, samples are diluted or concentrated to the same Chl concentration. This concentration must be low enough to avoid self-shading of excitation and re-absorption of the emission light, yet high enough to obtain a high signal to noise ratio.

**Cell suspensions and thylakoids**

The fluorescence emission of many single-celled organisms can be measured in their respective growth medium. However, some cyanobacteria and algae accumulate fluorescent molecules within their growth medium, and thus washing with fresh culture medium may be required. When cells are spun down for washing with the fresh medium, it is crucial to develop a rapid procedure to do so, as prolonged exposure to new conditions, such as the absence of light within the centrifuge or changes in pH may elicit a physiological response. A physiological response that has severe consequences for Chl fluorescence within a short period is anaerobic incubation response that has severe consequences for Chl fluorescence (Hohmann-Marriott and Demmig 1987). Therefore, a tissue sample must have a known orientation that is stable during measurements. Due to the high Chl concentration, there is substantial self-shading in most intact photosynthetic tissues. To overcome self-shading, Egelbert Weis developed a method that dilutes the Chl concentration within tissues to a lower concentration (Weis 1985). For this, the tissue sample is ground up in a liquid nitrogen-cooled mortar, and water (Pfundel and Pfeffer 1997) or quartz (Weis 1985) is added to dilute the sample. The ground sample is then inserted into a closed, cooled glass tube with a small diameter. The original description of the method for “diluted leaf powder” (Weis 1985) also demonstrates the spectral shifts of fluorescence emission spectra that occur due to self-shading.

**Processing of 77 K fluorescence emission data**

Recorded raw 77 K fluorescence spectra are usually processed, before presenting them in publications, in order to remove artifacts. After compensating spectra for light source and instrument response curves (see “Data processing section”), remaining artifacts arise from the fluorescence of the medium the samples are suspended in, and scattering. Recording of media fluorescence emission spectra can be used to compensate for these artifacts. Spectral features of the medium that fall outside the fluorescence spectra of the sample can be used to scale the medium fluorescence and subtract it from the sample spectrum. Some bacteria and algae produce fluorescent molecules that accumulate in the medium. The fluorescence of these molecules together with scattering artifacts can form a broad spectral band that extends into the Chl fluorescence spectra. This spectral band can often be effectively subtracted from the Chl fluorescence spectrum by an exponential or Gaussian function. These functions can be anchored at a wavelength were no, or very little Chl fluorescence is expected, which is usually at 800 nm or 850 nm in oxygenic organisms. Adjusting the fluorescence level at 850 nm to zero, even without the subtraction of a logarithmic function for artifact removal, is often performed to display data in figures.
77 K fluorescence emission of isolated complexes and organisms

Photosystems
The dominant room temperature fluorescence emitted by PSII, together with its variability, has been an early focus of spectral analysis in photosynthesis research. The insight that two photosystems are working in concert in oxygenic photosynthesis (Rabinowitch and Govindjee 1969) lead to a hunt for the fluorescence signal that is emitted by the second type of reaction center. Fluorescence measurement at 77 K played a crucial part in establishing the identity of the second type of photosystem we now know as PSI. The discovery and mis-assignments of Chl fluorescence signals have been reviewed by Govindjee (2004) and Strasser et al. (2004).

Photosystem II
In this section, we focus on the fluorescence emitted by PSII. As PSII is associated with different light-harvesting systems in different organisms, we only discuss the fluorescence characteristics of the PSII reaction center antennae that contain Chl \( a \), which are well established. There remains uncertainty about a functional assignment within PSII in Chl \( d \)-containing \( Acaryochloris \) species, as well as Chl \( d \)- and \( f \)-containing cyanobacteria capable of red light photoacclimation (FaRLiP) (Gan et al. 2015). The 77 K spectra of Chl \( d \)- (Miyashita et al. 1996) and Chl \( f \)- (Chen et al. 2012) containing PSII are discussed in details in the original literature.

PSII is composed of the reaction center core and the reaction center antenna. The PSI reaction center core houses 6 molecules of Chl \( a \), and 2 \( \beta \)-carotenes, 2 phaeophytins, and 2 quinones are associated with D1 (PsbA) and D2 (PsbD). Four of these Chls are part of the special set of Chls that mediates charge separation, while the remaining 2 Chls energetically couple the special set of Chls to the reaction center core antenna. The 32 Chl \( a \) molecules, and 6 \( \beta \)-carotenes, which make up the core antennae pigments, are housed in the proteins CP43 (PsbC) and CP47 (PsbD). The absorption and fluorescence spectra of isolated CP43 and CP47 (Fig. 7A,B) and isolated PSII (Fig 7C,B) reaction centers of plants and cyanobacteria are very similar indicating a very stringent conservation through evolution.

At room temperature, PSII emission originates predominantly from Chls fluorescing at 695 nm. Upon further cooling to 77 K, a distinct fluorescence at 685 nm also gains prominence. The emission at 695 nm was early on correctly assigned to be emitted by Chls within PSII (reviewed by Govindjee 2004). The fluorescence emission at 685 nm, however, was first thought to emanate from the antenna complexes of plants (reviewed by Strasser et al. 2004). This fluorescence emission was later also assigned to the PSI reaction center antenna proteins CP43 (Rijgersberg et al. 1979) and CP47 (Nakatani et al. 1984). More recent studies on isolated plant PSII particles (Andrizhiyevskaya et al. 2005) confirm the association of 695 nm emission with CP47, but indicate that the 685 nm emission arises from fluorescence emitted by both CP47 and CP43 at 77 K. Compared to room temperature, the fluorescence emission by PSII increases by a factor of about two when cooled to temperature of 77 K. At this temperature, PSII still contributes to long wavelength fluorescence, which overlaps with PSI fluorescence emission at 720/740 nm (Butler 1977).

PSII assembly and repair
The de novo assembly of PSII subunits is a fascinating topic (Eaton-Rye and Sobotka 2017) that also informs our understanding of the evolution of photosynthesis (Cardona 2016). In both cyanobacteria and plants, the assembly of PSII begins with binding of cytochrome \( b_{599} \) to the D2 subunit in the thylakoid membrane, forming the D2 pre-complex (Komenda et al. 2012, Nickelsen and Rengstl 2013). The D1 pre-complex is then bound, resulting in the heterodimeric reaction center pre-complex. Addition of the CP47 pre-complex to the heterodimeric reaction center pre-complex forms the RC47 complex (Boehm et al. 2012). The CP43 core antenna also forms a pre-complex with other subunits that bind to the RC47 complex.
Repair of damaged PSII shares common features with the assembly of PSII (Järvi et al. 2015). Damage to PSII reaction centers by light often affects D1 and results in the removal and consequent degradation of the damaged D1 protein (Komenda et al. 2012, Mulo et al. 2012). The repair of the damaged PSII involves the synthesis and insertion of a new D1 polypeptide into the PSII complex (Kyle et al. 1984, Järvi et al. 2015). This repair mechanism allows minimal energy expenditure during this process (Nixon et al. 2005, Takahashi and Badger 2011). Subcomplexes can be characterized by the presence or absence of fluorescence signal specific for the CP43 and CP47. In addition, increased fluorescence yield of peripheral light-harvesting complexes indicates that the assembly is not well coordinated or that PSII is damaged. Fluorescence spectroscopy at 77 K has provided valuable insights into the assembly and repair of the photosynthetic machinery (McCormac et al. 1996, Mysliwa-Kurdziel et al. 1997, Komenda et al. 2012, van Wijk et al. 1995).

Photosystem I

The discovery and assignment of the fluorescence emission at 720/730 nm to PSI is closely linked to the establishment of the two photosystem model for oxygenic photosynthesis (reviewed by Govindjee 2004). PSI complexes show a wide variety of emission spectra depending on the species and physiological conditions, but are located between 720–730 nm in oxygenic phototrophs, with some cyanobacteria having emission bands centered at wavelengths as long as 760 nm (Karapetyan et al. 2014).

The PSI reaction center of cyanobacteria (Fig. 8A), green algae, and plants (Fig. 8B) consists of two proteins (PsaA and PsaB) which together house around 85 Chls (Jordan et al. 2001). In addition, several smaller protein complexes contain about 10 Chl a molecules in cyanobacteria and plants. Cyanobacteria – which can form PSI trimers – are known to expand the trimeric PSI in stress conditions by circular antenna system that contributes an additional 180 (Boekema et al. 2001a) to 218 (Bibby et al. 2001a) Chl a molecules.

The absorption cross-sections of PSI of green algae, red algae, heterokont algae, and plants are further expanded by light-harvesting systems that belong to the three trans-membrane helix family of light-harvesting complexes (Busch et al. 2010). In plants, four of these light-harvesting complexes (Lhca) are associated with PSI, adding 52 Chl a and 9 Chl b molecules. An additional 20 Chl a molecules interface the Lhca with the reaction center (Mazor et al. 2015). An unknown number of light-harvesting systems are associated with the PSI of red algae and heterokont algae.

In cyanobacteria and plants, fluorescence around 720 nm is emitted by Chls within the reaction center antenna (Karapetyan et al. 2014). Plants also possess an additional pool of long wavelength emitters located in the Lhca antenna (Morosinotto et al. 2003), specifically within Lhca3 and Lhca4 (Wientjes et al. 2011). This assignment is in agreement with previous greening studies, which indicated that the fluorescence emission at 736 nm is emitted by peripheral antenna complexes (containing Chl b), while the emission around 724 nm is emitted by the PSI reaction center antenna (Mullet et al. 1980a,b).

In plants, the fluorescence emission of PSI at 77 K is not modulated by the reduction state of P700. However, in red algae (Ley and Butler 1977) and cyanobacteria, (Karapetyan et al. 2014), modulation of the fluorescence yield dependent on the reduction state of P700 have been reported, but during steady-state fluorescence emission measurements, this modulation can be neglected.

![Fluorescence emission from isolated PSI and PSI super complexes from cyanobacteria and plants. The fluorescence emission spectra (excited at 440 nm) of isolated PSI (black), isolated CP43’ and (blue) isolated CP43’-PSI supercomplex (red) of the cyanobacterium Synechocystis PCC 6803 are shown in panel A. The fluorescence emission spectra (excited at 440 nm) of isolated PSI-Lhca-super complex (black), Lhcb-trimers (blue) PSI-Lhca-(Lhcb-trimer) super complex (green) of the plant Arabidopsis thaliana is shown in panel B. The emissions in both panels are normalized to the maximum of the Soret band. The fluorescence spectra in panel A have been digitized from Bibby et al. (2001a).

The fluorescence spectra in panel B have been digitized from Galka et al. (2012).]
Interpretation of 77 K fluorescence in different organism groups

Cyanobacteria and red algae

Cyanobacteria are a diverse group of oxygen-producing organisms, which share many features of their photosynthetic machinery with their ancestors, which gave rise to the chloroplasts of photosynthetic eukaryotes (Hohmann-Marriott and Blankenship 2011). This evolutionary relationship is reflected in the photosynthetic machinery of red algae, which uses similar pigments and protein structures as cyanobacteria. Several cyanobacteria have achieved model status, including unicellular *Synechocystis* and *Synechococcus* species, as well as multicellular, nitrogen-fixing *Anabaena* and *Nostoc* species, while the unicellular *Cyanidioschyzon* and multicellular *Porphyridium* species are red algal model systems. In the following section, we mainly discuss the well-characterized photosynthetic machinery of cyanobacteria with the inference that the photosynthetic machinery of red algae is similar.

Peripheral light-harvesting systems and supercomplexes

What makes cyanobacteria a rewarding species for spectroscopic studies are their light-harvesting pigments, the phycobilins, which are covalently linked within phycobiliproteins. The phycobiliproteins can be assembled into large structures, the phycobilisomes (de Marsac 2003, Marx et al. 2014). Phycobilins absorb light within the “green gap” between the two main absorption bands of Chl a. Phycobilisomes are in many conditions primarily associated with PSII, but as discussed in detail later, can dynamically (Mullineaux 2014) adjust this association and can form supercomplexes with both photosystems (Liu et al. 2013). There are several types of phycobilins; including phycoerythrobilin, phycocyanobilin, and allophycocyanobilin, but not all species of cyanobacteria contain all of these phycobilins. For example, *Synechocystis* sp. PCC 6803 contains phycocyanin, allophycocyanin, and phycoerythrin, while the latter is absent in *Synechococcus* sp. PCC 7002. Phycoerythrin absorbs excitation between 475−575 nm (maximum 565 nm), and fluoresces maximally at 640 nm; phycocyanin absorbs excitation between 525−635 nm (maximum 620 nm), and fluoresces at maximally at 644 nm; allophycocyanin absorbs between 550−665 nm (maximum 650 nm), and fluoresces maximally around 670 nm. As the number of Chl per reaction center is higher in PSI (96 Chl a molecules) (Jordan et al. 2001) than that in PSII (35 Chl a molecules) (Umema et al. 2011), exciting Chls at 435 nm preferentially excites PSI, while photons with a wavelength of 590/635 nm preferably excite PSII in cyanobacteria. The PSI/PSII ratio in cyanobacteria is much higher than 1, and consequently more than 90% of Chls can be associated with PSI.

Some cyanobacteria can utilize other Chls, in addition to Chl a, to perform oxygenic photosynthesis. A polyphyletic group of organisms, the prochlorophytes, possess a membrane-embedded antenna system that houses Chl b along with Chl a, which is related to the stress-induced antenna protein isiA and the PSII-subunit CP43 (La Roche et al. 1996), with similar fluorescence emission spectra. Chl d is used by the *Acaryochloris marinus*, and Chl d and Chl f (Li and Chen 2015) are used by a diverse group of cyanobacteria that take advantage of environments rich in far-red photons.

Photosystem I

Cyanobacterial PSI has species-specific long wavelength-emitting Chls ranging from 727−760 nm with 2-7 Chls estimated to contribute to the emission (Karapetyan et al. 2014). There is also a spectral range of emission in red algal PSI, ranging from 708 nm for *P. cruentum* to 728 nm for *C. caldarum*. Trimeric PSI complexes are commonly found in cyanobacteria under a variety of conditions (Fig. 8A) (Boekema et al. 1987, 2001; Garczarek et al. 1998) and *Synechocystis* sp. PCC 6803 (Bibby et al. 2001a,b). Some cyanobacteria can form monomeric, trimeric, and tetrameric PSI, with each possessing different optical characteristics including Chl fluorescence emission bands at 77 K (monomer at 725 nm, trimer at 730 nm, and tetramer at 715 nm) (Li et al. 2014). In red algae, however, PSI appears to be monomeric (Gardian et al. 2007).

Red algae have light-harvesting complexes related to the Lhca and Lhcb of alga and plants, which are absent in cyanobacteria (Busch et al. 2010). These three-transmembrane helix proteins, named Lhcr, are exclusively associated with the PSI of red algae. As in cyanobacteria (Kondo et al. 2007, Watanabe et al. 2014), there is good evidence that a specific pool of phycobilisome proteins is associated with PSI in red algae also (Busch et al. 2010).

State transition

State transitions are dynamic adjustments of the
435 nm (D). Phycobilisomes donate more energy to PSII in state 1. Uncoupler FCCP. The plant DCMU under the light, and in the state 2 by treatment with the were locked in the state 1 by application of the PSII inhibitor is observed. Fore no fluorescence emission characteristic of phycoerythrin while genes for phycoerythrin synthesis are missing, and there- C. reinhardtii can synthesize phycocyanin and allophycocyanin, thus panel 6301 can synthesize phycocyanin and allophycocyanin, and can function as a PSI antenna system under stress and can function as a PSI antenna system under stress. The Chl a in PSI and PSII, or energy transfer occurs between Chl a in PSII and PSI, in a process called “spillover” (Biggins and Bruce 1989), was debated for a long time (McConnell et al. 2002, Li et al. 2004). However, there is now good evidence that the movement of phycobilisomes is a critical part of state transitions (Joshua and Mullineaux 2004), while spill-over appears not to be the major route for transferring energy from the phycobilisomes to PSI (Mullineaux 2008). A recent detailed study on the movement of phycobilisomes in different red algae (Kaňa et al. 2014) found that phycobilisomes have very limited mobility in the thermophilic red algae C. caldarium, while the mesophilic red algae P. cruentum exhibited phycobilisome movement in analogy to state transitions observed in cyanobacteria.

**Stress-induced adaptations**

Iron limitation induces changes in the composition of the photosynthetic machinery in cyanobacteria. Under iron limitation, proteins encoded by the isiAB operon are expressed (Pakrasi et al. 1985a,b, Laudenbach et al. 1988, Riehman and Sherman 1988, Burnap et al. 1993). While isiB codes for a thioredoxin that can functionally replace ferredoxin, IsiA possesses homology to the Chl a-binding PSI protein CP43 (Burnap et al. 1993, Falk et al. 1995), and can function as a PSI antenna system under stress conditions. During iron limitation, 18 IsiA from an antenna ring around trimeric PSI in Synechococcus sp. PCC 7942 (Boekema et al. 2001) and Synechocystis sp. PCC 6803 (Fig. 8A) (Bibby et al. 2001a,b). It was also suggested that IsiA could replace CP43, thus acting as an alternative antenna complex for PSI (Pakrasi et al. 1985b), or as an excitation energy dissipater, with the ability to protect PSI from photoinhibitory damage during iron starvation (Park et al. 1999). The induction of the IsiA protein under strong light, even in the presence of iron, confirmed its photoprotective role (Havaux et al. 2005). The presence of IsiA can be discerned by an increase in 685 nm emission (Burnap et al. 1993, Falk et al. 1995, Park et al. 1999), and an increase in energy partitioning to PSI. The latter is reflected by a more prominent fluorescence emission by the red Chls of the PSI reaction center antenna Chls around at 720 nm.

**Green algae and plants**

Green algae and plants share a common ancestor and consequently share features of their photosynthetic machinery that modulate the distribution of absorbed light energy between PSI and PSII (Fig. 9A)(Mullineaux 2014). 77 K fluorescence studies in cyanobacteria and red algae (Murata et al. 1966), played a crucial role in establishing the concept of state transition, as changes in energy distribution could be easily investigated by preferentially illuminating the phycobilins while monitoring the fluorescence emission bands specific for PSI and PSII (Murata 1969). These studies indicate that both PSI and PSII receive excitation that was absorbed by phycobilins (Mullineaux 1992). Whether state transitions are based on a physical movement of phycobilisomes between PSI and PSII, or energy transfer occurs between Chl a in PSII and PSI, in a process called “spillover” (Biggins and Bruce 1989), was debated for a long time (McConnell et al. 2002, Li et al. 2004). However, there is now good evidence that the movement of phycobilisomes is a critical part of state transitions (Joshua and Mullineaux 2004), while spill-over appears not to be the major route for transferring energy from the phycobilisomes to PSI (Mullineaux 2008). A recent detailed study on the movement of phycobilisomes in different red algae (Kaňa et al. 2014) found that phycobilisomes have very limited mobility in the thermophilic red algae C. caldarium, while the mesophilic red algae P. cruentum exhibited phycobilisome movement in analogy to state transitions observed in cyanobacteria.

![Fig. 9. Fluorescence emission characteristics of state transition in cyanobacteria, green algae, and plant during state transition. State 1 is indicated by the black trace while state 2 is indicated by the red trace. The cyanobacterium Synechococcus 6301 was excited at 600 nm (A) and 430 nm (B). Synechococcus cells were locked in the state 1 by treating the cells with far-red light, which preferentially excites PSI, while the cells investigated in the state 2 were dark-acclimated. The green algae Chlamydomonas reinhardtii was excited at 455 nm (C). Chlamydomonas cells were locked in the state 1 by application of the PSII inhibitor DCMU under the light, and in the state 2 by treatment with the uncoupler FCCP. The plant Arabidopsis thaliana was excited at 435 nm (D). To lock the Arabidopsis cells in different states, excitation light favoring PSII and PSI, respectively, was utilized. Spectra were and normalized to the Soret emission band for all figures.](image-url)
organism for photosynthesis research, such as toolboxes for genetic manipulation, and the ability to grow non-photosynthetically. However, *Chlamydomonas* possesses many features that are altered or unique compared to plants (Erickson *et al.* 2015).

**Peripheral light-harvesting systems and supercomplexes**

In plants and green algae, the absorption cross-section of the reaction center is extended by membrane-embedded light-harvesting systems. In addition to Chl a that is also found in the reaction center core and reaction center antenna, plants and green algae also synthesize Chl b that is associated with peripheral membrane-bound antenna complexes (LHCs). These antenna complexes and non-pigment-containing linker complexes form different supercomplexes with PSI and PSII, some of which can be isolated and structurally and spectroscopically characterized (Fig. 10) (Tokutsu and Minagawa 2013, Wei *et al.* 2016).

![Fig. 10. Fluorescence emission spectra of photosynthetic components of the green algae *Chlamydomonas reinhardtii*. (A) The 77 K fluorescence emission spectrum (excitation wavelength 440 nm) of trimeric LHCII complexes isolated from *C. reinhardtii* (Natali and Croce 2015). (B) The 77 K fluorescence emission spectra of an isolated His-tagged PSII core complex (black) (excitation wavelength 435 nm) (Sugiura and Inoue 1999), and the 77 K fluorescence emission (excitation wavelength 440 nm) of PSII-LHCII super complex (red) (Drop *et al.* 2014). (C) The 77 K fluorescence emission spectra of LHCI-PSI super complex (excitation wavelength is 435 nm) (Kargul *et al.* 2003). (D) The 77 K fluorescence emission spectrum of a whole cell in state 1 (black) and state 2 (red) (excitation wavelength 440nm) (Iwai *et al.* 2008). Panel A was digitized form Natali and Croce (2015). Panel B was digitized from Sugiura and Inoue (1999) and Drop *et al.* (2014). Panel C was digitized from Kargul *et al.* 2003 (2003). Panel D was digitized from Iwai *et al.* 2008 (2008).

**Major and minor light-harvesting complexes**

The protein complexes that house the pigments of the peripheral antennae can be grouped into monomeric Chl-binding complexes often called minor antennae complexes and trimer-forming LHCs (Busch *et al.* 2010) that are predominantly associated with PSII (Lhcb) or PSI (Lhca). Some Lhcb proteins in plants and algae can change the association between PSII and PSI during state transitions. Lhcb can form homo- and hetero-trimers that house 24 Chl a molecules, 18 Chl b molecules, and 12 carotenoids, with isolated trimers emitting fluorescence maximally between 678–680 nm at 77 K (Standfuss and Kühlbrandt 2004). Modeling in combination with mutational deletion of Chl ligands (Novoderezhkin *et al.* 2005) identified a spatially clustered group of three Chls as the final emitters, *i.e.*, these Chls are likely to donate excitation to the PSI reaction center. These Chls are likely responsible for fluorescence emission at 680 nm (F680). Lhcb can also form aggregated states that may be involved in photoprotection through excitation quenching, and exhibit a red-shifted fluorescence emission at 77 K (700–715 nm) (Ruban *et al.* 1997, 2012).

**Photosystem II**

The PSII of green alga and plants is organized in a dimer and Lhcb trimers can interface with both reaction centers of this dimer. Excitation transfer from the trimer to the reaction center is accomplished via three pigment-containing protein subunits (CP29, CP26, and CP24) associated with each reaction center monomer. Structural studies show that the subunit CP29 contains 10 Chl a molecules, 3 Chl b molecules, 3 carotenoids, and CP26 contains 9 Chl a molecules, 3 Chl b molecules, 3 carotenoids (Wei *et al.* 2016). Mutational analysis and modeling suggest that CP24 contains 5 Chl a molecules, 5 Chl b molecules, and two carotenoids (Passarini *et al.* 2009). These minor Chl-containing complexes are not known to contribute to 77 K fluorescence in intact systems largely.

**Photosystem I**

In contrast to PSI of cyanobacteria, the green algal and plant PSI possess tightly associated Lhca (Ben-Shem *et al.* 2003). The PSI supercomplex of *Pisum sativum* contains the reaction center proteins (PsaA and PsaB) as well as four Lhca (Lhca1–4) that form the peripheral antennae as a “dimer of dimers” (Mazor *et al.* 2015). Around 20 additional Chls energetically link the PSI reaction center with the peripheral antennae (Ben-Shem *et al.* 2003). The fluorescence emission of plant PSI is mainly due to Chl a in the Lhca antenna (Croce *et al.* 1998). Specifically, it is thought that one Chl a dimer within each Lhca (Qin *et al.* 2015) is the emitter of long wavelength fluorescence, whereas the reaction center core is only a minor contributor to this emission. Interestingly, no long wavelength emission was observed in PSI isolated from a prasinophyte green algae (Swingley *et al.* 2010).
State transition
The term “state transition” and its historical development has been introduced in the section on cyanobacteria. It became apparent that algae and plants also perform state transitions (Bonaventura and Myers 1969). As in cyanobacteria, the physiological basis for state transitions in green algae and plants appears to be the imbalance of electrons produced by PSII and utilized by carbon fixation (Minagawa 2011). However, in green algae and plants, the excitation coupling of Lhcb, not the excitation coupling of phycobilisomes to the two photosystems, is modulated. In the presence of a (partly) oxidized PQ pool, Lhcb are tightly excitationally coupled to PSII. Upon reduction of the PQ pool, some Lhcb are phosphorylated (Allen 1992) and consequently more Lhcb donate excitation to PSI. Whether much physical movement of the Lhcb is required for state transition is still unclear. It seems unlikely that a substantial mass migration of Lhcb between PSII-rich grana and PSI-rich stroma occurs, but instead, Lhcb may make additional contact with PSI. State transitions can be readily observed (Fig. 9B) by exiting samples at either the Chl a (430/440 nm) or Chl b (450/455 nm) Soret absorption band, while observing the emission bands of PSI (685/695 nm) and PSI (720/740 nm), with Chl b excitation providing a more specific signal for changes in excitation distribution.

Heterokont algae
Heterokonts comprise many photosynthetic algae including diatoms (Bacillariophyceae), brown algae (Phaeophyceae), and Eustigmatophyceae. Four membranes surround the chloroplast of these algae, indicating a secondary (Cavalier-Smith 1999) or even more complex endosymbiotic history (Keeling 2013).

Phaeodactylum
Major and minor light-harvesting complexes
Phaeodactylum tricornutum is a pennate diatom that, despite several clade-unspecific features regarding its life cycle and morphology, has achieved model status. The photosynthetic machinery of Phaeodactylum, however, shows features that are consistent and typical for most diatoms. In addition to light absorbed by Chl a, diatoms use alternative pigments, in particular, c-type Chls (c1-3) and carotenoids, with fucoxanthin being the most common. This composition of pigments was used to name the light-harvesting systems that are typical of many heterokont algae, the fucoxanthin-Chl-binding proteins (FCP) (Gundermann and Büchel 2014). The peripheral light-harvesting systems of most heterokont algae can be preferentially excited using the Soret band of Chl a, which are unlikely to be at homologous locations to the terminal emitter in the Lhca and Lhcb of plants (Gundermann and Büchel 2014). The absence of emission bands associated with Chl c species indicates that all Chl c is efficiently coupled to Chl a within the FCPs.

FCP is related to the three transmembrane-helix LHCs of plants and algae. The FCP can be grouped into three fractions: (1) FCPs, which are unique to heterokont algae (Lhcf), (2) FCPs, which are related to red algal Lhca (Lhcr), and (3) FCPs, which related to Lhcsr, a protein complex characterized in the green algae Chlamydomonas and the moss Physcomitrella (Lhcx) (Gundermann and Büchel 2014). Homology modeling in combination with spectroscopic and biochemical characterization suggest that each Lhcf can bind 6 Chl a molecules, 4 Chl c molecules, and 5–6 carotenoids, which in addition to fucoxanthin may also include lutein, diadinoxanthin, and diatoxanthin (Gundermann and Büchel 2014). In addition to Lhcf, another distantly related three-transmembrane helix protein belonging to the “red lineage Chl a-binding-like proteins” (RedCAP) is also present in heterokont algae.

Lhcf can form trimers (Lepetit et al. 2007, Nagao et al. 2013) analogous to the Lhcb trimers in plants and algae. While some uncertainty about the association of Lhcf to PSI to heterokont algae remains, there is good evidence that Lhcf associates predominantly with PSI (Juhas and Büchel 2012).

The fluorescence emitted by PSI-associated FCP has two prominent bands at 685 nm and 697 nm (Fig. 11A). These fluorescence bands, however, are emitted by a group of Chl a, which are unlikely to be at homologous locations to the terminal emitter in the Lhca and Lhcb of plants (Gundermann and Büchel 2014). The absence of emission bands associated with Chl c species indicates that all Chl c is efficiently coupled to Chl a within the FCPs.

Photocystem II
PSII core complexes form dimers in diatoms, such as Phaeodactylum, and exhibit a fluorescence emission band with a maximum at 692 nm (Fig 10B) (Yokono et al. 2015). This emission band appears to be a combination of a 692 nm and a 684 nm emitter located in CP47 (Yokono et al. 2015), while CP43 fluorescence emission is absent at 77 K. A different 77 K fluorescence emission pattern is observed in monomeric PSII, where the maximum fluorescence is emitted around 687 nm (Yokono et al. 2015). There is little biochemical evidence that any FCPs are associated with PSII to form stable super complexes in heterokont algae (Grouneva et al. 2011) or at least not into very stable super complexes (Nagao et al. 2010). However, spectroscopy-based studies reveal that the functional antenna size of PSII can be dynamically adjusted, thus suggesting a functional association between PSII and some FCPs (Miloslavina et al. 2009).

Photosystem I
Isolated PSI of diatoms has a long wavelength emission at
Fig. 11. Fluorescence emission spectra of photosynthetic components of heterokont diatom algae (*Phaeodactylum* and *Chaetoceros gracilis*). (A) The 77 K fluorescence emission spectrum (excitation wavelength 435 nm) of FCP from *Phaeodactylum tricornutum* (Litvín et al. 2016). (B) The 77 K fluorescence emission spectrum (excitation wavelength 425 nm) of PSII monomer (red) and dimer (black) from *Phaeodactylum tricornutum*, excited at 425 nm (Yokono et al. 2015). (C) The 77 K fluorescence emission spectrum (excitation wavelength 425 nm) of FCP-PSI super complexes from *Chaetoceros gracilis*, excited at 450 nm (Ikeda et al. 2008). (D) The 77 K fluorescence emission spectra (excitation wavelength 425 nm) of whole cells of *Phaeodactylum tricornutum* with different levels of non-photochemical quenching (Lavaud and Lepetit 2013).

Panel A was digitized from Litvín et al. (2016). Panel B was digitized from Yokono et al. (2015) Panel C was digitized from Ikeda et al. (2008) Panel D was digitized from Lavaud and Lepetit (2013).

715–720 nm (Fig. 11C) (Berkaloff et al. 1990, Veith and Büchel 2007, Ikeda et al. 2008). In addition, there is also a broad emission centered at 740–750 nm, which is a combination of vibronic sub-bands of all Chls within the sample.

**Nannochloropsis**

*Nannochloropsis* species are heterokont algae belonging to the Eustigmatophyceae family. *Nannochloropsis oceanica* and *gaditana* have received increased attention as they can accumulate high amounts of lipids. The architecture of the photosynthetic machinery of *Nannochloropsis* species has recently been investigated in detail (Herbstová et al. 2015, Litvín et al. 2016).

Eustigmatophyceae are unique among the heterokont algae in lacking Chl c and fucoxanthin in their LHCs. Thus, Chl a serves as the only Chl of both PSI and PSII, as well as in the peripheral LHCs. *Nannochloropsis* species use predominantly the carotenoids violaxanthin and vaucheroxanthin as light-harvesting pigments and for photoprotection. Chl a and the carotenoids are housed in LHCs, called VCP for viola-/vaucheriaxanthin Chl protein (Fig. 12A) (Litvín et al. 2016).

**Light-harvesting complexes**

*Nannochloropsis* species possess LHCs that can be clustered according to their evolutionary origin as diatoms including Lhcx and Lhcβ – the *Eustigmatophycean* version of the diatom Lhcf, and Lhcr. Furthermore, RedCLH, a light-harvesting protein first identified in *Chromera velia*, is also present (Litvín et al. 2016). Biochemical isolation suggests that VCP can form trimers (Litvín et al. 2016). VCP seems to be very rich in carotenoids, with about one carotenoid per two Chls, with violaxanthin and vaucheroxanthin being the dominant carotenoids (Litvín et al. 2016). Until high-resolution structural data of Lchv is available, it may be assumed that VCPs bind pigments analogous to Lhca, Lhcb (in green algae and plants), and Lhcf (in diatoms).
Photosystem II
Biochemical separation procedures suggest that the PSII complexes can contain about 4–5 VCPs per PSII core (Litvin et al. 2016) that belong to the Lhcr family. Interestingly, 77 K fluorescence spectra of isolated PSII particles show no fluorescence emission band at 685 nm and PSII that lack CP43 are present when PSII is isolated (Fig 11B) (Litvin et al. 2016). Whether this 77 K fluorescence emission pattern in combination with the CP43-lacking PSII complexes indicate that PSII has structural features, which are different from green algae and plants, needs to be resolved in the future.

Open Access This article is distributed under the terms of the Creative Commons Attribution License which permits any use, distribution, and reproduction in any medium, provided the original author(s) and the source are credited.

References
Choi F.: Low-temperature (4–77° K) spectroscopy of Chlorella,
temperature dependence of energy transfer efficiency. – BBA-Bioenergetics 216: 139-150, 1970b.


Kautsky H., Hirsch A.: [New attempts for carbon dioxide assimilation]. – Naturwissenschaft 19: 964-964, 1931. [In German]


Liu H., Roose J.L., Cameron J.C., Pakrasi H.B.: A genetically tagged Psb27 protein allows purification of two consecutive


Müller N.: [Relationships between assimilation, absorption and fluorescence in the chlorophyll of the living leaf.] – Jahrb. Wiss. Bot. 9: 42-49, 1887. [In German]


Park Y.I., Sandström S., Gustafsson P., Öquist G.: Expression of the isiA gene is essential for the survival of the cyanobacterium Synechococcus sp. PCC 7942 by protecting photosystem II.