Heterogeneity of Photosystem II as it Occurs in Domain Specific Regions of the Thylakoid Membrane of Spinach (*Spinacia oleracea* L.)

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The end is apparent in every beginning, yet, only by knowing the whole story do we come to understand it. This is the pleasure of life.

David Weaver, 2000.
I. Abstract

Thylakoid membrane fractions were prepared from specific regions of thylakoid membranes of spinach (*Spinacia oleracea*). These fractions, which include grana (B3), stroma (T3), grana core (BS), margins (Ma) and purified stroma (Y100) were prepared using a non-detergent method including a mild sonication and aqueous two-phase partitioning.

The significance of PSIIα and PSIIβ centres have been described extensively in the literature. Previous work has characterized two types of PSII centres which are proposed to exist in different regions of the thylakoid membrane. α-centres are suggested to aggregate in stacked regions of grana whereas β-centres are located in unstacked regions of stroma lamellae.

The goal of this study is to characterize photosystem II from the isolated membrane vesicles representing different regions of the higher plant thylakoid membrane. The low temperature absorption spectra have been deconvoluted via Gaussian decomposition to estimate the relative sub-components that contribute to each fractions signature absorption spectrum. The relative sizes of the functional PSII antenna and the fluorescence induction kinetics were measured and used to determine the relative contributions of PSIIα and PSIIβ to each fraction. Picosecond chlorophyll fluorescence decay kinetics were collected for each fraction to characterize and gain insight into excitation energy transfer and primary electron transport in PSIIα and PSIIβ centres.

The results presented here clearly illustrate the widely held notions of PSII/PSI and PSIIα/PSIIβ spatial separation. This study suggests that chlorophyll fluorescence decay lifetimes of PSIIβ centres are shorter than those of PSIIα centres and, at F_m, the longer lived of the two PSII components renders a larger yield in PSIIα-rich fractions, but smaller in PSIIβ-rich fractions.
II. Acknowledgments

There are a number of people present at Brock University who deserve many thanks. The Brock Electronic and Machine Shops have been a constant source of solutions for the Bruce lab. This project was enhanced greatly for having their assistance. The same sort of thanks are deserved by my lab mates. To do no one any injustice, I will leave out all of their names. None of them are insignificant by any means however, as soon as I begin to list them I become flustered because they all deserve to be mentioned first and foremost. They are well aware of who they are, and I hope they are also aware of how important they have been for the last two years. I thank my examiners (both internal and external) and advisors for their time and guidance during this project.

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I can never be able to pay my parents back for their decision to have me stay around for the last two years to complete this very important job.

There are three more people that are involved in this project that pushed it towards the checkered flag. The first is not even aware that she is part of this project. Christine Dobbin now knows more about photosynthesis than any other person outside the field, and she doesn’t even know it! That poor girl put up with my random babbling about reaction centres, antennae, and broken down β-centres and appeared to enjoy it. She is bright and beautiful, a deadly combination!

The other two are of course Doug Bruce and Sergej Vasil’ev. In an effort to supervise a successful master’s degree candidate, these two men have somehow helped put me in a position to become one of them. Scary? Not really. I admire them for their wisdom, enthusiasm, generosity and most of all their positive role in my life, to help me figure out my own way in this exciting field of intriguing pursuits and bold discoveries.
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VI. List of Abbreviations

\( \alpha \)-DM - \( n \)-dodecyl-\( \alpha \),D-maltoside

a.u. - arbitrary units

B3 - grana thylakoid membrane fraction

BBY - photosystem II-enriched preparation made according to Berthold et al., 1981

BChl - bacteriochlorophyll

BS - grana core thylakoid membrane fraction

CCI - core complex I

CCII - core complex II

CF\(_0\) - a coupling factor

CF\(_1\) - a coupling factor

Chl \( a \) - chlorophyll \( a \)

Chl \( b \) - chlorophyll \( b \)

Chl \( z \) - accessory reaction centre Chl \( a \), bound to a tyrosine residue

cyt \( b_{f} / f \) - cytochrome \( b_{f} \)-cytochrome \( f \) complex

DAS - decay associated spectra

DCMU - 3-(3,4,-dichlorophenly)-1,1-dimethyurea

\( F_{0} \) - minimal level of fluorescence

\( F_{M} \) - maximal level of fluorescence

\( F_{SAT} \) - fluorescence yield after a single saturation flash

\( F_{V} \) - variable fluorescence (\( F_{M} - F_{0} \))

Fd - ferredoxin

FNR - ferredoxin-NADP\(^{+}\) reductase

HEPES - \( N \)-2-hydroxyethylpiperazine-\( N' \)-ethanesulfonic acid

LHCI - light-harvesting complex I

LHCII - light-harvesting complex II

Ma - grana margins thylakoid membrane fraction

OEE - oxygen-evolution enhancer

P680 - photosystem II reaction centre

P700 - photosystem I reaction centre

PC - plastocyanin

Pheo - pheophytin

PMF - proton motive force

PQ - plastoquinone

PSI - photosystem I

PSI\( \alpha \) - photosystem I alpha centre

PSI\( \beta \) - photosystem I beta centre

PSII - photosystem II

PSII\( \alpha \) - photosystem II alpha

PSII\( \beta \) - photosystem II beta

\( Q_{A} \) - first bound plastoquinone acceptor

\( Q_{B} \) - second semi-bound plastoquinone acceptor

qE - energy-dependent quenching

qI - photoinhibitory quenching

qN - nonphotochemical quenching

qP - photochemical quenching

qT - state transitions

\( R_{0} \) - critical distance for Förster transfer measured in \( \text{Å} \)

rubisco - ribulose bisphosphate carboxylase/oxygenase

T3 - stroma lamellae thylakoid membrane fraction

Y100 - purified stroma lamellae thylakoid membrane fraction
1. Introduction

Chloroplasts are the energy converting organelles of higher plants. Chloroplasts not only contain the earth's most abundant soluble enzyme, ribulose bisphosphate carboxylase/oxygenase (rubisco), but also the most abundant biological membrane, the photosynthetic membrane known as the thylakoid. Thylakoid membranes, like all biological membranes, are fluid-mosaic structures in which lipid bilayers act as two-dimensional fluids (Singer and Nicolson 1972). Proteins and protein complexes may diffuse freely within its structure, both rotationally and laterally. The light reactions of photosynthesis take place in and across the thylakoid membrane, The word thylakoid comes from the Greek, "ΘΨΛΑΚΟΕΙΔΗΣ", meaning sac-like. The sac-like grana region in higher plants occur in stacked, or appressed membrane regions joined by unstacked, or unappressed stroma lamellae. Four major components of the plant thylakoid membrane are the light-driven photosystems I and II (PSI and PSII), a cytochrome $b_6/f$ (cyt $b_6$-$f$) complex and an ATP synthase. Each of these components are intimately involved, in series, in the light reactions of electron transport and proton pumping in cooperation with mobile carriers to liberate electrons from water, which ultimately reduce NADP⁺, to form NADPH, stored chemical energy (Albertsson 2001; Barber 1990; Menke 1962).

Some contemporary insights into thylakoid membrane biomass have estimated that 1 µg of chlorophyll corresponds to 16.7 cm² of plant surface. This value may be highly variable and is indeed species specific (Barber 1990). The thylakoid membranes employed in a typical leaf occupy a total surface area 600 to 1000 times greater than this area. Barber states, "I think there is little doubt that the thylakoid membrane is the most abundant biologically active lipo-protein structure on our planet!" (Barber 1990).

Upon examination of the dynamic architecture of the thylakoid membrane, its mystery begins to reveal itself. About twenty years ago it was first noted that the two light driven pigment-protein complexes are spatially separated within thylakoid membranes of higher plants (Anderson and Melis 1983). This regional segregation surprised many researchers since these photosystems operated in series (Hill 1937; Hill and Bendall 1960).
Figure 1 - Electron micrograph of thylakoid membranes (80,000x). Grana appears as flattened stacked discs (GT), connected by unstacked stroma lamellae (ST) (Staehelin 1986).
Clearly, thylakoid membranes of green plants are some of the most complex and dynamic membrane systems in biology (Staehelin and van der Staay 1996).

Photosynthetic systems and assemblies are most commonly studied by one of three preparation types. Intact systems offer insight into real processes that may occur under physiological conditions. Commonly, these experiments employ cyanobacteria, green algae or intact chloroplasts. Intact chloroplasts may be within the leaf, in protoplasts or carefully isolated from the other plant material. These organisms are well suited for photosynthesis research on multiple levels because of the extensive homology that exists among cyanobacteria, green algae and higher plants. These similarities are likely the results of evolutionary endosymbiosis several millions of years ago.

Semi-intact systems are the most widely used preparations for biophysical studies of higher plants. Intact thylakoid membranes can very easily be harvested from whole leaves and assayed for photosynthetic activity in ways very similar to those employed in experiments utilizing intact systems. The variable fluorescence yield, a commonly measured character, collected from isolated thylakoid membranes may be somewhat smaller in magnitude, but nonetheless, the data obtained from this very abundant source of plant material remains tremendously valuable.

The third, and most labour intensive preparation type used to study photosynthesis involves the further dissection of semi-intact systems to their individual components. Some decades ago, researchers began to design specialized isolation procedures to effectively zoom in on specific pigment-protein complexes, or individual proteins, derived from intact thylakoid membranes, harvested from the chloroplasts of higher plants or cyanobacteria. The use of such systems is widespread in the field. These localized components have allowed investigators to answer very specific questions surrounding the activities, or inactivities of countless components essential to light-driven electron transport and chemical energy accumulation. In combination with recombinant DNA techniques, the knowledge of photosynthetic systems has met with unparalleled growth. Random knock-outs and site-directed mutagenesis have allowed researchers to probe specific genes that are intimately involved in photosynthetic processes. It is now possible to assay the
biophysical characteristics of photosynthesis in the absence and/or up-regulation of a wide variety of genes in model organisms such as *Synechocystis* sp. PCC 6803, a commonly used cyanobacterium (Kaneko et al. 1996; Tabata and Ikeuchi 2001; Vermaas 1998), *Chlamydomonas reinhardtii*, a single celled eukaryotic green alga, (Kindle 1998; Nedelcu and Lee 1998; Silflow 1998) and *Arabidopsis thaliana*, a higher plant (Arabidopsis Genome Initiative 2000; Barkan 1998; Jansson 1999; Wakasugi et al. 2001), where the entire genomes (including the chloroplasts genome where appropriate, see Hiratsuka et al. 1989 for example) has been sequenced and mapped.

This study involves a thorough biophysical study of the photosynthetic apparatus of the higher plant *Spinacia oleracea* L., commonly referred to as spinach. The method employs an advanced separation technique whereby chloroplast derived thylakoid membranes are subjected to a mild sonication followed by an aqueous two-layer partitioning (Albertsson 1986; Albertsson et al. 1994; Åkerlund and Albertsson 1994). The uniqueness of this technique lies in its detergent free nature. In the past, thylakoid fractions rich in one pigment-protein complex or another were prepared solely by detergent dependent isolations (Berthold et al. 1981). Certainty that the complexes of interest are maintained in their native form is reduced upon the introduction of detergents. Thylakoid membranes from *Nicotiana tabacum* (tobacco) chloroplasts (Gadjieva et al. 1999) and the green algae *Dunaliella salina* (Stefánsson et al. 1997) have also been successfully fractionated using the relatively non-invasive detergent-free technique. This exciting method is not limited to thylakoid membranes. It has also proven to be useful in the specific domain isolation of plasma membrane and smooth endoplasmic reticulum from mammalian liver cells (Gierow 1994; López-Pérez et al. 1981).

This study employed five unique thylakoid membrane fractions. These included grana, stroma lamellae, grana core, grana margins and a special purified stroma lamellae. The spatial distribution of different types of PSII reaction centres, hereon referred to as PSIIX and PSIỊβ centres (the foci of this study), is well documented. PSIIX centres are reported to aggregate within the stacked regions of grana, whereas PSIỊβ centres are located within the unstacked stroma
lamellae (Albertsson et al. 1990b; Albertsson et al. 1991; Albertsson et al. 1994). This allows, with considerable certainty, measurements to be obtained from functional PSII centres of either α- or β-variety separately and clearly.

This study is a multifaceted and comprehensive pursuit of the characterisation of PSII heterogeneity. It is unique in that the plant material employed in these experiments has been mechanically derived from the thylakoid membranes of spinach without using detergents (Albertsson 1986; Albertsson et al. 1990b; Albertsson and Svensson 1988; Svensson et al. 1990). This is one of the key elements to this study. The detergent free isolation procedure offers confidence in the native state of the preparations. For the first time, PSIIα from appressed grana regions and PSIIβ from unappressed stroma exposed regions of higher plant thylakoid membranes are studied using the techniques described here. Spatial separation of the two PSI and PSII, as well as PSIIα and PSIIβ will be demonstrated using a variety of biophysical techniques including: absorption spectroscopy, 77K fluorescence emission spectroscopy, fluorescence induction kinetics and PSII absorbance cross section analyses. The picosecond chlorophyll fluorescence decay kinetics and their global analysis provide insight into primary processes of photosynthesis with respect to PSII heterogeneity.
2. Literature Review

2.1. Photosynthesis

Photosynthesis is the process by which sunlight fuels a series of redox reactions to produce stored chemical energy where breathable oxygen is a byproduct. A very simplified summary of the entire process is demonstrated in Saussure's equation (1).

\[ 6CO_2 + 6H_2O \xrightarrow{h\nu} C_6H_{12}O_6 + 6O_2 \] (1)

The sunlight available to plants is approximately 2 mE·m\(^{-2}\)·s\(^{-1}\) considering only radiation involved actively in photosynthesis, a considerably small amount relative to the gross amount of solar radiation that is cast towards the earth (Mauzerall and Greenbaum 1989). The portion of this light employed by any given plant, or photosynthetic bacteria, is dependent upon the type(s) of pigments they harbour. Since each pigment type may preferentially absorb from different regions of the spectrum, an organism that employs a wider range of pigments may, hypothetically, make use of a larger portion of the radiant energy available.

2.1.1. The Light Reactions

Water is split by a light-catalysed process which occurs in the lumen of the thylakoid membrane. The liberation of an electron from water, coupled with the evolution of O\(_2\) by the Mn-cluster (Kok et al. 1970) is the primary step in oxygenic photosynthesis. The instability of the PSII reaction centre in its oxidized excited state offers it a unique ability. Simply put, PSII’s ability to oxidize water is a direct consequence of the high oxidizing potential of the oxidized reaction centre special pair, P680\(^+\) (Durrant et al. 1995). To reach P680\(^+\), the electron is passed by one other intermediate carrier, Y\(_z\) (a tyrosine radical), the primary donor. The special pair of chlorophylls, P680, is the terminal acceptor of absorbed excitation energy by the PSII-antenna in higher plants. It is here that the electron is excited. Excited P680\(^-\) performs a charge separation (P680\(^+\)•Pheo\(^-\)•Q\(_a\)) upon reduction of pheophytin (Pheo), a pigment molecule similar to Chl \(a\), without the central Mg atom:

\[ P680 + Pheo + h\nu \xrightarrow{3ps} P680^+ + Pheo^- \] (2)
followed by a charge stabilization, \((P680^+\text{Pheo} \cdot Q_a^-)\). Charge stabilization is achieved by the further movement one electron at a time to \(Q_a\). Both charge separation and charge stabilization are energetically favourable reactions however \(Q_a\) reduction is more stable (or less reversible) since the movement of the electron is further removed from \(P680^+\) (Hara et al. 1997). The two-electron gate allows \(Q_b\) to be reduced to the semiquinone in one step, then to plastoquinol on the second:

\[
2H_2O + 4 \text{photons} + 2PQ + 4H^+ \rightarrow O_2 + 4H^+ + 2PQH_2
\]

(3)

The \(Q_b\) site must stabilize the semiquinone since it is usually very unstable and reactive. Upon reduction by \(PQH_2\), the cytochrome \(b_6\) (also called \(cyt\ b 563\)) - cytochrome \(f\) (for \(frons\) or leaf) complex, abbreviated \(cyt\ b_6-f\), is then able to reduce a small copper-containing protein, plastocyanin (PC), a mobile carrier,

\[
2PQH_2 + 4PC(Cu^{2+}) \rightarrow 2PQ + 4PC(Cu^+) + 4H^+
\]

(4)

which is subsequently oxidized by the P700 reaction centre within the lumen of the thylakoid.

LHCl, the major antenna system of PSI, delivers absorbed excitation energy by inductive resonance from its 100 chlorophylls to the P700 reaction centre. Excited P700 reduces \(A_o\), a Chl a molecule;

\[
P700 + Chla + hv \rightarrow P700^+ + Chla^-
\]

(5)

followed by the reduction of \(A_1\), phylloquinone (vitamin \(K_1\)). \(F_x\), an iron-sulphur group (with a 4Fe-4S centre similar to the 2Fe-2S group of \(cyt\ b_6-f\)) is able only to pick up a single electron at a time, even though four centres are present. Electrons are transferred to a soluble form of Fe-S protein, ferredoxin (Fd):

\[
 hv + 4PC(Cu^+) + 4Fd(Fe^{3+}) \rightarrow 4PC(Cu^{2+}) + 4Fd(Fe^{2+})
\]

(6)

In the stroma ferredoxin-NADP\(^+\) reductase (FNR) catalyses the oxidation of mobile ferredoxin by NADP to form NADPH:
4Fd(Fe^{2+}) + 2NADP^+ + 2H^+ \rightarrow 4Fd(Fe^{3+}) + 2NADPH \quad (7)

ADP (+ Pi) is converted to ATP and H_2O by the coupling factor, ATPase. The CF_0 subunit extends through the hydrophobic region of the thylakoid and the CF_1 subunit is attached on the stromal side. Protons generated by the hydrolysis of H_2O and the oxidation of PQH_2 are pumped out of the lumen through CF_0, towards CF_1 where ADP is phosphorylated in the stroma. These reactions are summarized in the z-scheme diagram in figure 3 (Salisbury and Ross 1992). NADPH and ATP produced during these reactions are used to reduce CO_2 to carbohydrates in the subsequent dark reactions (figure 2).
Figure 2 - The relationship between the "light" reactions and the "dark" reactions of photosynthesis.
Figure 3 - Summary of the light-driven electron transport events within thylakoid membranes, commonly called the z-scheme. The reduction potential of each carrier indicates a higher affinity for electrons as it becomes increasingly positive (Zubay 1993).
2.2. Thylakoid Membrane Architecture

The thylakoid membranes within the chloroplasts of higher plants are functionally similar to those of cyanobacteria, which are suspected to be their ancestors via endosymbiosis many millions of years ago. Chloroplasts typically measure 5 μm in diameter and are 1 to 2 μm thick. In mature leaf cells these green photosynthetically active plastids occupy only 20% of the cellular volume but they may occupy up to 70% of the surface area (Ellis and Leech 1985). Like mitochondria, these organelles contain their own circular genome, however many of the genes coding for photosynthetic proteins have been translocated to the nucleus. Much of what is left functions primarily in fatty acid synthesis (Waller et al. 1998). Chloroplasts divide by binary fission, a process that closely resembles bacterial cell division morphologically and genetically (Leech 1976; Whatley 1988). The principal determinant leading to the number of chloroplasts that may occur in a leaf mesophyll cell is its size (Dean and Leech 1982). Furthermore, evidence suggests that chloroplasts may only appear once a critical cell size is attained (Ellis et al. 1983). A recent study with respect to chloroplast size revealed that employment of many small chloroplasts may be more advantageous to mature higher plants than fewer large ones. Small chloroplasts are more mobile within the cell than large chloroplasts. Movement within the cytoplasm of leaf cells offers 1) a means of protection from photodamage to the photosynthetic apparatus under excess light conditions and 2) enhances efficient utilization of low-incident photon flux densities (Jeong et al. 2002). Chloroplast movement, with respect to light conditions, is a process known as phototaxis.

Many components of the thylakoid membranes have been remarkably conserved between cyanobacteria and higher plants through evolution with respect to their structure and function. For this reason many studies regarding the biophysical and photochemical properties of the process of photosynthesis apply to both prokaryotic and eukaryotic systems, regardless of the organism used in the laboratory. The main difference lies in the relative organization of their macrodomains. Whereas many cyanobacteria arrange concentric layers of thylakoid membranes within their prokaryotic cellular interior, the thylakoid membranes within the chloroplasts of most eukaryotic higher plants are arranged in regions of stacked (or appressed) grana and unstacked (or
unappressed) stroma lamellae. The ratio of stacked to unstacked membrane regions is variable and depends primarily upon growth conditions (Staehelin 1986). Under low light conditions, grana have larger diameters and make up a larger proportion of the thylakoid membrane area. For example, it is reported that at higher light intensities, *Zea mays* mesophyll chloroplasts contain 61% stacked regions and an average granum diameter of 0.35 μm, however at lower light intensities the stacked regions make up 73% of the thylakoid architecture and grana measure 0.43 μm on average (Staehelin 1986).

The formation of grana has been well documented. Thylakoid membrane appressions occur between regions of the membrane where the Coulombic repulsive forces between two adjacent membrane surfaces are decreased (Dahlin *et al.* 1990). Coulomb’s law states that the magnitude of the electrostatic force exerted by one point charge on the other point charge is directly proportional to the magnitudes of the charges and inversely proportional to the square of the distance between them. At high salt concentrations (>5 mM) the electrostatic screening of surface negative charges is improved so reduced coulombic repulsion permits thylakoid membrane stacking and a concomitant phase separation of membrane complexes based on differential surface charge densities (Barber 1990). The method of thylakoid membrane vesicle separation used in this study takes advantage of this fundamental property of appressed grana and nonappressed stroma lamellae. Since one photosystem (and its associated antenna) has a greater net surface charge density than the other it is believed that the phenomenon of spatial separation may rest on this parameter. Reversible changes in the degree of stacking are induced by changes in cation concentration. Trivalent cations are more effective than divalent cations which are more effective than monovalent cations (Horton 1999). These changes greatly influence the chlorophyll fluorescence yield, even when photochemical activity is blocked by presence of a herbicide. At low salt levels, where the electrostatic screening is lowest, a reduction of the fluorescence yield occurs since lateral intermixing of all complexes allows efficient energy transfer between the photosystems (Armond *et al.* 1976; Barber 1990; Newell *et al.* 1987).

Membrane appression occurs not only among the stroma exposed surface of the grana
partition, but also within the lumen of the thylakoid membranes therein. Attractive forces between
two opposing inner surfaces keep them within close contact (Albertsson 1982). This space is
loaded with proteins, including lumenally exposed portions of intrinsic membrane proteins and
soluble ones as well (Bricker et al. 2001; Kieselbach et al. 1998). The functional role of grana
stacking is not completely understood. Some early studies suggested that formation of grana may
be required to activate photosystem II (PSII) (Arnzten and Briantais 1975). Evidence contrary to
these ideas was revealed when algal mutants, deficient in grana, where shown to be
photochemically competent (Goodenough et al. 1969; Goodenough and Staehelin 1971) and with
mutants of higher plants which demonstrated high PSII activity while maintaining relatively few
appressed membrane regions (Highkin et al. 1969; Keck et al. 1970; Armond et al. 1976). It is
suggested however, that the formation and maintenance of grana offer the thylakoid membranes
within chloroplasts functional flexibility in terms of a) maximizing the absorption cross-section by
densely packing Chl $a$ and $b$ in large numbers, b) ability to modify the cross-section in both short
term (Allen 1992), and long term, acclimation, situations and c) rapidly alter energy quenching upon
exposure to excess light to prevent photodamage (Aro et al. 1993; Horton 1999).

More than thirty years ago spatial separation of PSI and PSII was described within the
thylakoid membranes of higher plants (Anderson and Boardman 1966; Boardman and Anderson
1964). Functional PSII complexes are found predominantly in the appressed membrane regions
of grana, where the PSI complex occurs at its lowest frequency. The margins of the grana
surrounding the PSII-rich core houses a large proportion of PSI (Anderson and Goodchild 1987).
The unappressed stroma regions house different species of PSI and PSII. This special
organization of two photosystems is a universal feature of all granal chloroplasts (Arvidsson and
Sundby 1999; Garab and Mustárdy 1999; Gunning and Schwartz 1999; Horton 1999; Mehta et al.
1999) in spite of the relatively low viscosity of the glycerolipid phase of the thylakoid membrane
which one may suspect may confer a homogenous mixture of all thylakoid components (Graan and
Ort 1984; Voet and Voet 1995). The ATP synthase is restricted to stroma exposed regions of the
thylakoid membrane (Miller and Staehelin 1976). The size of the CF$_1$ subunit makes it virtually
impossible for this large coupling factor to squeeze within the appressed membranes of stacked
grana (Allen and Forsberg 2001). The advantages and disadvantages of spatial separation of the
photosystems will be addressed in section 2.2.3.

2.2.1. Photosystem I - the FeS type reaction centre

Photosystem I (PSI) is an evolutionary product of anoxygenic green photosynthetic bacteria
based on bacteriochlorophyll (BChl). The BChl antennae of green bacteria are substantially larger
than that of the purple bacteria and their reaction centres perform electron transport in a slightly
different way (figure 4). Not until the late 1980s were the fundamental differences revealed among
the purple, green and the newly discovered heliobacteria. The reaction centres of both the green
sulphur bacteria and the heliobacteria are less complex homodimeric homologs of PSI from higher
plants and cyanobacteria (Baymann et al. 2001). PSI centres in higher plants do not aggregate
into trimers, which they do in cyanobacteria. The Psal (and possibly Psal) gene product, which
manages this trimerization in cyanobacteria (Schluchter et al. 1996), may rather be part of light-
harvesting complex I (LHCl) stabilization. This is only one of the key differences surrounding PSI
structure between higher plants and cyanobacteria. The structure of PSI will not be discussed in
such great detail here (relative to that of PSII, section 2.2.2.), however several important features
are described below.
Figure 4 - Green sulphur bacterial photosynthesis. Electrons, originating from sulphur donors are used to reduce NAD+ via light driven ATP cyclic photophosphorylation.
Table 1 - Composition of core complex I and inner antenna of higher plants (Scheller et al. 2001).

<table>
<thead>
<tr>
<th>Gene</th>
<th>Subunit (cofactors)</th>
<th>Mass (kDa)</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>PsaA</td>
<td>PsaA (~96 Chl a, ~22 β-carotene, P700, A0)</td>
<td>83.2</td>
<td>Light-harvesting</td>
</tr>
<tr>
<td>PsaB</td>
<td>PsaB (A1, Fx)</td>
<td>82.4</td>
<td>Charge separation, e⁻ transport</td>
</tr>
<tr>
<td>PsaC</td>
<td>PsaC (FxA, FxB)</td>
<td>8.8</td>
<td>e⁻ transport</td>
</tr>
<tr>
<td>PsaD</td>
<td>PsaD</td>
<td>17.6</td>
<td>Binds ferredoxin, PSI-C</td>
</tr>
<tr>
<td>PsaE</td>
<td>PsaE</td>
<td>10.8</td>
<td>Binds ferredoxin and FNR, Cyclic e⁻ transport</td>
</tr>
<tr>
<td>PsaF</td>
<td>PsaF (Chl a)</td>
<td>17.5</td>
<td>Binds PC, LHCI-730</td>
</tr>
<tr>
<td>PsaG</td>
<td>PsaG (Chl a)</td>
<td>10.8</td>
<td>Binds LHCI-680</td>
</tr>
<tr>
<td>PsaH</td>
<td>PsaH (Chl a)</td>
<td>10.2</td>
<td>Binds LHClII (qT), Stabilizes PSI-D</td>
</tr>
<tr>
<td>Psal</td>
<td>Psal</td>
<td>4</td>
<td>Stabilizes PSI-L</td>
</tr>
<tr>
<td>PsaJ</td>
<td>PsaJ</td>
<td>5</td>
<td>Stabilizes PSI-F</td>
</tr>
<tr>
<td>PsaK</td>
<td>PsaK (Chl a)</td>
<td>9</td>
<td>Binds LHCl-680</td>
</tr>
<tr>
<td>PsaL</td>
<td>PsaL (Chl a)</td>
<td>18</td>
<td>Stabilizes PSI-H</td>
</tr>
<tr>
<td>PsaN</td>
<td>PsaN</td>
<td>9.8</td>
<td>Stabilizes PC</td>
</tr>
</tbody>
</table>

Table 2 - Light harvesting components of PSI (Scheller et al. 2001).

<table>
<thead>
<tr>
<th>Gene</th>
<th>Subunit</th>
<th>Mass (kDa)</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lhca1</td>
<td>Lhca1</td>
<td>22</td>
<td>Light-harvesting, LHCI-730</td>
</tr>
<tr>
<td>Lhca2</td>
<td>Lhca2</td>
<td>23</td>
<td>Light-harvesting, LHCI-680B</td>
</tr>
<tr>
<td>Lhca3</td>
<td>Lhca3</td>
<td>25</td>
<td>Light-harvesting, LHCl-680A</td>
</tr>
<tr>
<td>Lhca4</td>
<td>Lhca4</td>
<td>22</td>
<td>Light-harvesting, LHCI-730</td>
</tr>
</tbody>
</table>

* Cofactors bound are ~10 Chl a, ~2 Chl b and ~3 carotenoids in each subunit.
PSI consists of 17 proteins in higher plants. Those belonging to the core complex (CCI) form one group. The other group, LHCl, is intimately related to CCI. In higher plants the LHCl, consisting of four proteins act as the peripheral antenna, similar to the phycobilisomes present in cyanobacteria, with obvious differences. A summary of the subunits that compose PSI is found in tables 1 and 2.

The PsaL, Psal and PsaM proteins, located close to the three-fold axis of the PSI-trimer, far from lipid-exposed regions in cyanobacteria (Schmid et al. 2001) are the first group. Their function, structural in nature, was proposed several years ago during mutageneic studies, where trimers were unable to form in the absence of PsaL (Kuroiwa et al. 2000). PsaL is the primary link between the three monomers, harbouring some hydrophobic carotenoids and three chlorophylls. The Psal subunit, located between PsaL and PsaM, is involved in trimerization as well. No chlorophyll is bound to Psal, however it does form hydrophobic contacts with some carotenoids. The interactions between PsaL and Psal have been conserved over evolution and are maintained in plant systems which employ PSI as a monomer coupled to an elaborate accessory light-harvesting complex. PsaM is the smallest of all PSI subunits (3.4 kDa) and is found in cyanobacteria. Other than its presence as merely an open reading frame in the chloroplast genome of liverwort, PsaM is yet to be found in any plant PSI preparation (Fromme et al. 2001).

The three stromal subunits, like the name suggests, do not contain transmembrane helices. PsaC, PsaD and PsaE are involved in docking ferredoxin or flavodoxin. Homologus polypeptide sequences of these proteins occur in green sulphur bacteria and heliobacteria (Baymann et al. 2001).

Two large protein subunits, PsaA and PsaB make up the reaction centre of PSI. In green sulphur bacteria, a homodimer of the PscA protein forms the reaction centre. Similarly, a homodimer of the PshA protein forms the reaction centre of heliobacteria. The homology between these transmembrane proteins is well documented (Baymann et al. 2001). Along with PsaC, these three proteins are the most essential part of CCI. In the reaction centre of PSI, P700 resides in the PsaA/PsaB heterodimer. As well as the primary electron acceptor A_o (a Chl a molecule), A_1
(phytloquinone), and F\textsubscript{x} (a 4Fe-4S cluster). Upon excitation of P700 to its lowest excited state, P700\textsuperscript{−}, primary charge separation occurs in 1-3 ps as A\textsubscript{0} is reduced. The redox potential of P700\textsuperscript{−} is highly negative (~1.3 V) which makes it a strong reductant. Whether P700 occurred as a monomer or dimer in higher plants was the item of some debate for several years (Ikegami and Itoh 1988) (see (Sétif 1992) for detailed review). Similar to the bacterial reaction centre, the dimerization of the P700 reaction centre was confirmed. However, surprising to most researchers, the P700 special pair does not employ a Chl a homodimer, but rather a Chl ala' heterodimer. Chl a' is the C13' epimer of Chl a (Kobayashi et al. 1988; Webber and Lubitz 2001). These chlorophylls are separated by 6.3 Å (closer than the bacterial reaction centre special pair) and are found nearest the lumenal side of the thylakoid membrane. They are oriented parallel to each other, perpendicular to the membrane plane, with an interplanar distance of 3.6 Å. A weaker electronic coupling between the π electron systems in P700, relative to the bacterial reaction centre, is a product of only partial overlap of the heterodimeric chlorophyll ring structures (Fromme et al. 2001).

The terminal electron acceptors F\textsubscript{a} and F\textsubscript{b} are bound to PsaC (Scheller et al. 2001).

A subgroup of polypeptides make up the extrinsic subunits of CCI. These include gene products PsaD - PsaN. Their functions are listed in table 1. Most of the subunits of the extrinsic CCI are involved in binding and stabilization of other CCI subunits and/or subunits of the LHCI (Scheller et al. 2001).

The LHCI subunits are listed in table 2. The products of four nuclear genes, Lhca1-4, compose this peripheral antenna of PSI. Two others, Lhca5 and Lhca6, are present in some species in very low quantities. The LHCI binds about 70-110 chlorophylls with a Chl ala/b ratio of 2.5. The LHCI can be divided into two subcomplexes. The LHCI-730 subcomplex (characterized by its 77K fluorescence emission peak at 730 nm) is formed from the Lhca1/Lhca4 heterodimer. The LHCI-680 sub complex can be further fractionated into LHCI-680A (Lhca3) and LHCI-680B (Lhca2) (Scheller et al. 2001; Thornber et al. 1991).

Low energy pigments, relative to the reaction centre P700, are present in LHCI, namely those which occur at 735 nm absorbance maximum. Removal of the LHCI reveals a peak 720 nm
emission, attributed to Chl a present in the inner antenna of CCI. These chlorophylls act as a funnel for excitation energy towards P700 (Scheller et al. 2001).
Figure 5 - Model of the pigment-protein distribution of the PSI complex including the light-harvesting complex I (LHCI) and the core complex I (CCI). Molecular weights are shown within the structures, labelled by subunit and sum of the chlorophylls they contain is described beneath (adapted from (Thornber et al. 1991)).

Chl/P700 = 194
Chl \( a/b = 6.05 \)
Figure 6 - Simplified schematic of purple nonsulphur bacterial photosynthesis. The oxidation of the electron source succinate precedes the reduction of NAD⁺ via light driven electron transport.
Photosystem II (PSII) is an evolutionary product of anoxygenic purple photosynthetic bacteria (figure 6). PSII is, structurally, very similar to the bacterial reaction centre, but spectroscopically and functionally, remains very different (see figure 6). These differences may be correlated with its unique water-oxidizing capabilities and/or its photochemical (and non-photochemical) regulatory functions in order to minimize unwanted and damaging side reactions (Durrant et al. 1995). PSII performs a unique and dangerous function in nature. The generation of the strong oxidant, P680+, capable of reaping electrons from energy-poor H₂O sustains all life on earth. The pigment-protein complexes of PSII may be split into two related but functionally and spatially separated groups. The one group is comprised of those associated with the light-harvesting complex II and the other, those associated with the core complex II. The number of chlorophylls associated with PSII may vary, and is often species specific, however it is usually greater than 200 chlorophyll molecules make up an entire PSII complex, with a Chl a/b of 2.05. This Chl b content is relatively high, compared to the number of Chl b molecules associated with PSI. The vast majority of these chlorophyll molecules is associated with peripheral light-harvesting complex, the primary light capturing tool of PSII (Bassi et al. 1987b). The polypeptides of LHCII are immunologically distinct from those of LHCI associated with PSI (Melis 1991). Six pigment-protein complexes, Lhcb1-6, are aggregated to construct this very large antenna. LHCIib, the most abundant, harbours 40-45% of the total chlorophyll. 10-15% of the remaining chlorophyll is bound by the other three. Lhcb4 (LHClIa), encoded by the Lhcb4 gene, is located relatively close to the core complex and is comprised of at most, only 5% of the chlorophyll. This may include six chl a, two chl b (Pascal et al. 2000). It is present in all species of higher plants and some green algae with small variations in its size, usually ranging from 29 to 31 kDa. Its molecular weight gives it its alternative name, CP29, for chlorophyll-binding protein of 29 kDa. The absorbance spectra (at room temperature) of Lhcb4 has a maximum at 675 nm and a second peak at 645 nm. Xanthophylls, leutin and violoxanthin occur in nearly equal proportions but together, with some small amounts of neoxanthin, are enriched relative to other LHCII subunits (Thornber et al. 1991).
Lhcb4 has also been suggested to be the site of energy-dependent quenching (qE) at the E166 residue (Pesaresi et al. 1997).

Lhcb5 (LHClIc), is also called CP26 (a chlorophyll-binding protein of 26-29 kDa, depending on the species). The absorption maximum occurs at 671 nm at room temperature and a 77K fluorescence maximum at 680 nm. Pigment composition assays on LHClIc reveal that Chl a occurs twice as much as Chl b (Bassi et al. 1987a) and it is relatively enriched in xanthophyll lutein, neoxanthin with smaller amounts of violoxanthin (Thornber et al. 1991).

Lhcb6 (CP24) has been suggested to be a linker between the abundant LHClIb and CCII. Its absorption maximum occurs at 668 nm or 674 nm and has (Bassi et al. 1987a). Of the total chlorophyll present in chloroplasts, only 3% is bound to Lhcb6. The xanthophyll lutein is the most prevalent carotenoid with smaller amounts of neoxanthin (lowest of all LHClII subunits) and violoxanthin (Thornber et al. 1991).

The most abundant chlorophyll-binding protein of PSII is LHClIb, which accounts for approximately one-third of the total protein of the chloroplast and about 45% of the chlorophyll. LHClIb has received much attention since it not only serves as a large chlorophyll-binding protein, but also as a thylakoid membrane adhesion protein and as a mediator of energy transfer between PSII and PSI. LHClIb is nuclearly encoded by 3 to 16 gene families (species specific) (Thornber et al. 1991).

The involvement of LHClIb in cation-mediated grana stack formation is well documented (Armond et al. 1976). Not surprisingly, the appearance of grana in developing thylakoid membranes at various stages parallels the accumulation of LHClIb. The addition of cations (divalent more so than monovalent) to thylakoid membrane suspensions causes LHClIb in membrane vesicles to cluster within areas of membrane contact (Horton 1999; McDonnel and Staehelin 1980; Mullet and Arntzen 1980; Ryrie et al. 1980).

LHClIb mediated energy transfer between PSII and PSI is accomplished through phosphorylation/dephosphorylation of this complex. State 1-state 2 transitions (qT), are a photoprotective component of non-photochemical quenching (qN). Under light conditions favouring
PSII absorption, LHCIIb is found in the grana stacks, tightly coupled to the rest of LHCII and PSII, increasing its effective absorbance cross section (Yakovlev et al., 2002) (i.e. state 1). LHCIIb in this form is nonphosphorylated and promotes membrane adhesion. Upon preferential excitation of PSII, LHCIIb becomes phosphorylated by a membrane-bound kinase, consequently, the net negative surface charge on the protein increases. Higher coulombic repulsion causes a slight unstacking of the grana appressed membranes and the phospho-LHCIIb complexes (p-LHCIIb) migrate to nonappressed membrane regions, the stroma lamellae. Upon reversal of these conditions, the dephosphorylation of p-LHCIIb is performed by a membrane-bound phosphatase prior to return to appressed grana membranes and association with PSII. This well-orchestrated redistribution of absorbed excitation energy is reported to be regulated by the redox state of the plastoquinone pool (Allen et al. 1981) (see figure z-scheme) and by the cyt b_{6}/f complex (Bennett et al. 1988).

Recently, the \textit{cab80} gene from pea coding for Lhcb was inserted in the cyanobacteria, \textit{Synechocystis} sp. PCC 6803, via the \textit{psb3} promoter on the novel pA31lhcb plasmid. Since cyanobacteria do not employ the large extrinsic light-harvesting complexes similar to higher plants, the protein was very unstable in cyanobacterial thylakoid membranes and has a very short lifetime. Only fragments of the gene product can be see following the addition of xanthophylls (He et al. 1999).
Table 3 - Polypeptides of the inner and outer light-harvesting complex of PSII. All are nuclear-encoded (Erickson 1998).

<table>
<thead>
<tr>
<th>Gene</th>
<th>Subunit</th>
<th>Apparent MW (kDa)</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>psbR</td>
<td>R</td>
<td>10</td>
<td>Donor and acceptor side function</td>
</tr>
<tr>
<td>Lhcb1</td>
<td>LHCII</td>
<td>30</td>
<td>Light-harvesting*</td>
</tr>
<tr>
<td>Lhcb2</td>
<td>LHCII</td>
<td>31</td>
<td>Light-harvesting*</td>
</tr>
<tr>
<td>Lhcb3</td>
<td>LHCIIa</td>
<td>25</td>
<td>Light-harvesting*</td>
</tr>
</tbody>
</table>

Inner Antenna

| Lhcb4 | CP29   | 35                | Excitation energy transfer and dissipation* |
| Lhcb5 | CP26   | 36                | Excitation energy transfer and dissipation+ |
| Lhcb6 | CP24   | 18                | Excitation energy transfer and dissipation+ |

+ All light absorptive protein in the LHCII employ chl a, chl b, lutein, neoxanthin, violaxanthin.

Table 4 - Polypeptides of the oxygen-evolving complex of PSII. All are nuclear-encoded (Erickson 1998).

<table>
<thead>
<tr>
<th>Gene</th>
<th>Subunit</th>
<th>Apparent MW (kDa)</th>
<th>Function</th>
</tr>
</thead>
<tbody>
<tr>
<td>psbO</td>
<td>OEE1</td>
<td>30 (33)</td>
<td>Stabilizes Mn, binds Ca^{2+} and Cl^-</td>
</tr>
<tr>
<td>psbP</td>
<td>OEE2</td>
<td>20 (23)</td>
<td>Binds Ca^{2+} and Cl^-</td>
</tr>
<tr>
<td>psbQ</td>
<td>OEE3</td>
<td>18 (16)</td>
<td>Binds Cl^-</td>
</tr>
</tbody>
</table>
The core-complex II (CCII) is comprised of four pigment-proteins which can be equally subdivided into two groups. The first subgroup consists of two pigment-proteins known as CCIIa and CCIIb, the *psbB* and *psbC* gene products respectively. The CCIIa subunit binds ~15 Chl *a* molecules and has an apparent molecular mass of 47 kDa (CP47), however the calculated mass of 56 kDa has been reported for spinach (Barber *et al.* 2000). Two to three β-carotenes are also associated with CP47. Of the 508 amino acid residues that make up the ultimate structure, 200 are present in lumenally exposed loops. An even larger portion of the polypeptide spans the thylakoid membrane six times, where the large hydrophilic loop joins at helices 5-6. The remaining portions occur as two small hydrophilic loops in the stroma that join to helices 2-3 and 4-5 and the stroma exposed N- and C-terminal ends. Evidence that suggests that the large extrinsic protein region binds chlorophyll is poor. Its function is more likely to involve water-splitting processes.

The second subgroup of CCII pigment-proteins make up the PSII reaction centre. The *psbA* and *psbD* gene products, better known as D1 (32 kDa) and D2 (30 kDa). Along with the 9 kDa and 4 kDa polypeptides that assemble to form the cytochrome *b*$_{559}$ (cyt *b*$_{559}$) (Stewart and Brudvig 1998) the reaction centre is complete and houses the ‘special pair’ chlorophylls, P680.

A third Chl *a* binding subunit of CCII is the *psbS* gene product. This 22 kDa protein (CP22) absorbs well at 674 nm and twice in the blue, at 440 and 468 nm. Some reports indicate that in may also bind Chl *b* with a chl *alb* of 1.6 - 4.3 (Funk *et al.* 1994). 77K fluorescence emission spectra of CP22 reveals a peak at 675 nm. Its role as a PSII subunit may be one of minor light-harvesting. Or, it may bind chlorophylls scavenged from other degraded components of the PSII-supercomplex. Recently, CP22 has been reported to be intimately involved in energy dependent quenching, qE, a component of nonphotochemical quenching, qN (Li *et al.* 2000). Most interestingly, this protein can be immunologically detected in the cyanobacteria, *Synechocystis* sp. PCC 6803 and *Phormidium laminosum*. Some differences among the number of transmembrane helicities, and the presence/absence of chl *b* with respect to species raise some interesting evolutionary questions (Funk *et al.* 1994).

Products of the *psbE* and *psbF* genes are the α and β subunits of the cyt *b*$_{559}$ complex
respectively. The ultimate function of this complex remains obscure, however it is well known that absence of one or both the subunits inhibits the assembly of PSII. Likely proposals suggest that cyclic electron transfer around PSII may be orchestrated by cyt $b_{559}$ in some way, protecting PSII from photoinhibition (Arnon and Tang 1988; Aro et al. 1993; Thompson and Brudvig 1988). Some authors suggest that cyt $b_{559}$ may also be able to donate an electron to $Y_z^+$ in the dark, to reduce this dangerous oxidant (Canaani and Havaux 1990). Cross-linking studies have demonstrated the close association of cyt $b_{559}$ to the D1 subunit of the PSII reaction centre (Barbato et al. 1992; Stewart and Brudvig 1998). More recently however, high resolution x ray crystal structure suggest that the location of this subunit may be on the opposite side of the reaction centre, nearer the D2 protein (Zouni et al. 2001).

The similarities that exists between the P680 (due to a characteristic bleaching observed at 680 nm upon oxidation of this species) reaction centre, the primary electron donor, of higher plants (and cyanobacteria) and the bacterial reaction centre have been well documented (Rochaix et al. 1984; Youvan et al. 1984). Unlike the reaction centre of purple bacteria, the PSII reaction centre, P680, the chlorin rings of the two special pair Chl $a$ molecules ligated to HIS198 on the D1 and D2 proteins may not be parallel to each other. Less overlap between the chlorin rings, at an angle of 30°, causes exciton coupling to be relatively weak, however, charge separation is twice as efficient. The consequence of weak exciton coupling is actually two-fold. First, P680 is a relatively shallow trap, since the free energy of the charge separated state is very near that of the of excited state. Compared to the bacterial primary electron donor, P870/P960, where coupling of the excited state is severely red-shifted ($V = 550$ cm$^{-1}$ for P870 and 950 cm$^{-1}$ for P960) relative to other reaction centre pigments, P680 possesses a very small red shift relative to other reaction centre pigments, hence, the shallow trap. Secondly, charge separation, although efficient, is reversible. The exciton-radical pair equilibrium (Leibl et al. 1989; Schatz et al. 1988) illustrated in equation 8, demonstrates how a charge recombination between P680$^+$ and Pheo$^-$ may follow primary charge separation and render an exciton in the antenna rather than pushing on towards charge stabilization (Karukstis et al. 1990).
One can see that a photon may visit the P680 reaction centre many times until it is ultimately trapped and used for photochemistry or dissipated from the antenna (see figure 7). An alternative would be a diffusion limited system, whereby the orientations of pigments and their ability to funnel absorbed excitation energy towards the reaction centre complex is the limiting factor towards primary charge separation (Vasil'ev et al. 2001).

The excitonic coupling of pigments depends largely on their relative distances and mutual orientations. Recently, the crystal structure of the PSII reaction centre from the cyanobacteria, Synechococcus elongatus, had been resolved to 8 Å (Rhee et al. 1997; Rhee et al. 1998) and then to 3.8 Å (Zouni et al. 2001). These structures are useful in determining the relative distances and mutual orientations of the Chl a present in the core complex (Vasil'ev et al. 2001). Until these precise structures had been released, researchers often found themselves making bold predictions regarding these parameters based on models derived from electron microscopy of 2D crystals (Barber and Kühlbrandt 2000; Lyon 1998; Marr et al. 1996; Mayanagi et al. 1998; Nakazato et al. 1996).

P680+ has an enormous redox potential of about 1.17 V compared to a value of only 0.4 - 0.6 V for P700+, the oxidized form of the primary donor of PSI. P680+ is thought to be the most oxidizing species found in living organisms. Such a high oxidizing potential affords P680+ the ability to liberate electrons from water, a very stable compound. As a by-product, breathable oxygen is released into the atmosphere (Barber and Kühlbrandt 2000; Karukstis et al. 1990). This oxidation of water is achieved by four consecutive univalent oxidation steps at the Mn-cluster where the intermediary electron transfer component is a redox active tyrosine (see (Debus 1992; Renger 1993) for review).

In order to withstand such a harsh environment, nature has afforded the PSII reaction centre with the means to prevent and deal with the oxidation of neighbouring proteins and pigments (Anderson 2001).
Figure 7 - The energy of light absorbed by pigment molecules (P₁-Pₙ) in the antenna system hops rapidly from molecule to molecule by resonance energy transfer until it is trapped (kᵣ) in an electron-transfer reaction in a reaction centre (shaded molecule) or released back (kᵣ⁻¹) to the antenna pigment bed where it may be dissipated as either heat or fluorescence (Renger 1992; Zubay 1993).
Table 5 - Polypeptides of the PSII core complex including the inner antenna and the reaction center (RC). All are encoded by the chloroplast genome except PsbW (Erickson 1998).

<table>
<thead>
<tr>
<th>Gene</th>
<th>Subunit</th>
<th>Apparent MW (kDa)</th>
<th>Function</th>
<th>Associated Pigments</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>PSII RC</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>psbA</td>
<td>D1</td>
<td>32</td>
<td>$Y_z$ and binds $Q_B$, Pheo, P680, Mn</td>
<td>chl a, $\beta$-carotene</td>
</tr>
<tr>
<td>psbD</td>
<td>D2</td>
<td>34</td>
<td>$Y_D$ and binds $Q_A$, Pheo, P680</td>
<td>chl a, $\beta$-carotene</td>
</tr>
<tr>
<td>psbE</td>
<td>cyt $b_{559}$-$\alpha$</td>
<td>6</td>
<td>Binds heme, photoprotection</td>
<td>chl a, $\beta$-carotene</td>
</tr>
<tr>
<td>psbF</td>
<td>cyt $b_{559}$-$\beta$</td>
<td>4</td>
<td>Binds heme, photoprotection</td>
<td>chl a, $\beta$-carotene</td>
</tr>
<tr>
<td>psbI</td>
<td>I</td>
<td>4</td>
<td>PSII stability</td>
<td>chl a, $\beta$-carotene</td>
</tr>
<tr>
<td><strong>Inner Antenna</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>psbB</td>
<td>CP47</td>
<td>50</td>
<td>Excitation energy transfer, binds OEE1</td>
<td>chl a, lutein, $\beta$-carotene</td>
</tr>
<tr>
<td>psbC</td>
<td>CP43</td>
<td>47</td>
<td>Excitation energy transfer, binds OEE1</td>
<td>chl a, lutein, $\beta$-carotene</td>
</tr>
<tr>
<td><strong>Core Complex</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>psbH</td>
<td>H</td>
<td>9</td>
<td>Photoprotection</td>
<td></td>
</tr>
<tr>
<td>psbJ</td>
<td>J</td>
<td>5</td>
<td>PSII assembly?</td>
<td></td>
</tr>
<tr>
<td>psbK</td>
<td>K</td>
<td>4</td>
<td>PSII stability, assembly</td>
<td></td>
</tr>
<tr>
<td>psbL</td>
<td>L</td>
<td>5</td>
<td>$Q_A$ function</td>
<td></td>
</tr>
<tr>
<td>psbM</td>
<td>M</td>
<td>5</td>
<td>?</td>
<td></td>
</tr>
<tr>
<td>psbN</td>
<td>N</td>
<td>4</td>
<td>PSII stability</td>
<td></td>
</tr>
<tr>
<td>psbS</td>
<td>S</td>
<td>22</td>
<td>chl chaperonin, antenna, qE</td>
<td>chl a</td>
</tr>
<tr>
<td>psbT</td>
<td>T</td>
<td>3</td>
<td>PSII stability</td>
<td></td>
</tr>
<tr>
<td>PsbW</td>
<td>W</td>
<td>6</td>
<td>?</td>
<td></td>
</tr>
<tr>
<td>psbX</td>
<td>X</td>
<td>4</td>
<td>$Q_A$ function</td>
<td></td>
</tr>
</tbody>
</table>
Figure 8 - Schematic diagram of PSII in the thylakoid membrane (Erickson 1998).
Figure 9 - Model of the pigment-protein distribution of the PSII complex including the light-harvesting complex II (LHCII) and the core complex (CCII). Molecular weights are shown within the structures, labelled by subunit and sum of the chlorophylls they contain is described beneath (adapted from (Thornber et al. 1991)).
One other group of very important proteins occur in the lumen of the thylakoid membrane. The oxygen evolving complex is composed primarily of 3 extrinsic polypeptides with molecular weights of 33, 23 and 16 kDa, referred to as oxygen-evolution enhancer 1 (OEE1) OEE2 and OEE3 respectively (Erickson and Rochaix 1992). A brief summary of these components of the water oxidizing complex is listed in table 4. Their function is reviewed in (Nugent et al. 2001). Figure 10 illustrates the S cycle, also referred to as the Kok cycle (Kok et al. 1970), as water is oxidized and a four step linear mechanism operates to release molecular oxygen on the donor side of PSII.
Figure 10 - The S cycle. Possible Mn oxidation state changes for two of the Mn (of four in the cluster) within the water oxidizing complex. Two others are suggested to remain in the same oxidation state. At each S state (Sn) one proton and one electron are released (Nugent et al. 2001).
2.2.3. Heterogeneity

Heterogeneity among photosystem II is not a new concept. More than twenty-five years ago the notion that two distinct PSII species may exist surfaced for the first time (Melis and Homann 1975). This phenomenon was expanded to further explain the biphasic nature of the kinetics of PSII activity. It may be useful to describe these early measurements with the following equation:

\[
\begin{align*}
Z \cdot P680 \cdot Q & \leftrightarrow \frac{k_1}{k_2} Z \cdot P680^+ \cdot Q^- \leftrightarrow \frac{k_3}{k_4} Z^+ \cdot P680 \cdot Q^-
\end{align*}
\]

where \( Z \) is the secondary electron donor on the oxidizing side of the PSII reaction centre, \( P680 \), and \( Q \) is the primary electron acceptor. The rate constants \( k_1, k_2, k_3, \) and \( k_4 \) represent the photochemical rate constant, the thermochemical rate constants including the luminescent back reaction and the removal of primary oxidizing charges from the reaction centre, and the slow back reaction leading to the dark restoration of the fluorescence induction in DCMU poisoned chloroplasts. \( k_4 \), moderated by \( k_3 \), does not significantly alter the fluorescence kinetics (Melis and Homann 1975). Biphasic kinetic data suggested that two distinct populations of PSII were present in the chloroplast labelled PSII\( \alpha \) and PSII\( \beta \) (Black et al. 1986; Melis and Duysens 1979; Melis and Homann 1976). A slow linear phase upon examination of the kinetics of the photoreduction of \( Q \) is attributed to PSII\( \beta \). It is described as a monophasic, first-order function of time. Its slope is described as the rate constant \( k_\beta \), measured at 4.8 s\(^{-1} \). Alternatively, the photoactivity of PSII\( \alpha \) reveals a fast phase described as non-first-order. The kinetics are calculated by subtracting the slow first-order phase from the global event. The rate constant, \( k_\alpha \), estimated from the slope of the semilogarithmic plot at the start of the kinetic, \( T_{\text{init}} \), to be 11.2 s\(^{-1} \) is associated with PSII\( \alpha \) (Melis and Anderson 1983). The authors suggest that the nonlinearity of PSII\( \alpha \) photoreduction of \( Q \) is based on the its ability to aggregate into clusters of PSII\( \alpha \) centres, concomitantly, increasing its absorption cross section. As the slope (or the rate constant) of PSII\( \alpha \) increases progressively with time, the closure of the reaction centres becomes clear. Fewer and fewer reaction centres in the
PSIIα-aggregated clusters are open to receive absorbed excitation energy from their shared antenna. The terminal value of the slope of, $k_\alpha$, $T_{\text{final}}$, approached 40-50 s$^{-1}$, four to five times higher than the value at $T_{\text{init}}$. The relative ratio of $T_{\text{final}}$ to $T_{\text{init}}$ is suggested to be useful in calculating the size the PSIIα cluster. This may suggest that up to five individual PSIIα centres aggregate to construct the cluster. Furthermore, many PSIIα clusters many occupy the appressed membrane regions of thylakoid membranes (Melis and Anderson 1983).

Different light-harvesting antenna sizes for PSIIα and PSIIβ also become apparent upon analysis of the biphasic kinetic data of photoreduction of Q by Melis and Anderson, 1983. The portion of PSII centre classified as PSIIβ is 25%. Therefore the remaining portion, 75% are PSIIα centres. The ratio of PSII to PSI is 1.9 so, the individual reaction centre ratios are; PSIIα/PSI = 1.43 and PSIIβ/PSI = 0.48. These ratios, coupled with the values of $k_\alpha$ and $k_\beta$, allowed the authors estimate the absolute number of chlorophylls associated with PSIIα ($N_\alpha$), PSIIβ ($N_\beta$) and PSI ($N_{P700}$) that are involved in excitation energy transfer to their respective reaction centres. These parameters were determined this way:

\[
\frac{\text{Chl}}{\text{PSI}} = \frac{\text{PSII} \alpha}{\text{PSI}} N_\alpha + \frac{\text{PSII} \beta}{\text{PSI}} N_\beta + N_{P700}
\]  

(10)

\[ k_\alpha = cIN_\alpha \]  

(11)

\[ k_\beta = cIN_\beta \]  

(12)

\[ k_{P700} = cIN_{P700} \]  

(13)

where the overall ratio of the total Chl per PSI reaction centre is represented by $\text{Chl/PSI}$. The actinic light intensity, $I$, and a proportionality constant dependent upon the quantum yield of photochemistry, $c$, couple with the rate constants associated with each of the three reaction centre types allows the calculation of the absolute number of Chl associated with each reaction centre type in equations 11, 12 and 13. In each case $c$ is kept constant based on the assumption that the quantum yields for photochemistry for PSIIα, PSIIβ and PSI are similar (> 0.8). Based on these
calculations the authors report values of $N_\alpha = 234$, $N_\beta = 100$ and $N_{F700} = 209$ (Melis and Anderson 1983). These values are in agreement with (Thielen and Van Gorkom 1981b). In summary the authors suggest that the relative amounts of chlorophyll associated with $\text{PSII}_\alpha$, $\text{PSII}_\beta$ and PSI are 57%, 8% and 35% respectively. Hence, large PSII$\alpha$ harbours seven times more chlorophyll than smaller PSII$\beta$ and 1.8 times more than PSI. (Melis and Anderson 1983)

The identification of PSII heterogeneity and its characterization remains somewhat enigmatic, relative to the great deal of attention it has received. But even more mysterious is the physiological significance behind this evolutionary puzzle. Since PSII heterogeneity has been found in chloroplasts from virtually all higher plants which employ stacked and unstacked regions of thylakoid membranes, some advantage must be conferred to the organism which orchestrates this paradigm (Armond et al. 1976).

Melis has suggested a physiological significance for PSII heterogeneity that has rarely been debated. He reported that PSII heterogeneity exists in two modes (figure 11), both of which are dependant upon the location of PSII$\alpha$ (in the grana partition regions) and PSII$\beta$ (in stroma-exposed membranes). In the first mode, PSII antenna heterogeneity may be a product of the development of this multi-subunit pigment-protein complex (Guenther et al. 1988; Guenther and Melis 1990; Melis 1985; Melis 1991). PSII$\beta$ lacks the LHCII-peripheral antenna.
Figure 11 - The two modes of PSII heterogeneity are dependent upon the location of PSII in either grana or stroma lamellae regions of the thylakoid membrane. (Melis 1985; Melis 1991).
Figure 12 - The PSII repair cycle (Guenther and Melis 1990; Melis 1991).
The second mode, reducing side heterogeneity, is best described by the PSII repair cycle (figure 12). Photoinhibition is a regulatory response that plants may undergo under high light intensities. Photodamage may originate from an over-reduced PQ pool or from the oxidizing side of PSII. Both theories may be true and operate under certain conditions. Some researchers suggest that there is a threshold photon flux density for photodamage whereas others suggest that there is a constant, albeit low, probability for photodamage to occur with every absorbed photon (Anderson et al. 1998; Melis 1999). Irreversible photodamage to the reaction centre proteins brings the onset of photoinactivation of electron transport (Aro et al. 1993; Melis 1991). PSII-Q\(_{\beta}\) nonreducing units establish a repair state of the photochemical donor side of PSII. Here, the damaged 32 kDa reaction centre named D1 (Q\(_{\beta}\)-binding) protein is replaced. PSII repair is initiated by an uncoupling of the LHCII peripheral antenna from the damaged unit within stacked grana. The new PSII\(\alpha\)-like centre migrates towards the stroma-exposed region where the damaged D1 protein is replaced, rendering it a photochemically competent centre, still incapable of Q\(_{\alpha}\)-oxidation via Q\(_{\beta}\). PSII\(\beta\) is then activated upon its conversion to the Q\(_{\alpha}\)-reducing form prior to its ultimate migration back to stacked grana regions and reassociation with the LHCII-peripheral antenna. At this stage the newly functional PSII\(\alpha\) unit has been formed (Guenther and Melis 1990). The distinctive lipid composition of thylakoid membranes easily permits the migration of these large complexes. Only 10% of the lipid content is derived from phospholipids. The remaining 90% is made up of four-fifths mono- and digalactosyl diacylglycerols and one-tenth sulfoquinivisyl diacylglycerols. A high degree of unsaturation within the acyl chains of these lipids give the thylakoid membrane its high fluidity (Barber 1990; Voet and Voet 1995). An observed correlation with growth conditions may also support this hypothesis. However the proportion of PSII\(\beta\) centres occurring in non-photoinhibited material has been reported to be as high as one-third, a unduly large share (Hemelrijk and Van Gorkom 1996).
Figure 13 - A hypothetical schematic representation of the association of different phases in the PSII repair cycle (see figure 12). High light intensities cause D1 photodamage and phosphorylation of most PSII proteins within the appressed grana regions of the thylakoid membrane (1). Cleavage is triggered by conformational changes in the damaged D1 polypeptide (2). Dephosphorylation of D1 offers a superior substrate for protease (3) and is required before proteolytic cleavage (4). The 23 kDa D1 fragment is digested further as a newly synthesized D1 is inserted into the thylakoid membrane (4). Cofactors are reassociated as the new D1 is processed and palmitoylated. The repaired PSII complex migrates back to the appressed grana regions (6) and where final assembly of the LHC and OEC occur. Phases following steps (2) and (3) occur in unknown locations. For clarity, only the heterodimeric reaction centre polypeptides are shown (Aro et al. 1993).
A semi-related physiological significance of PSII heterogeneity has been suggested by Weis, (Timmerhaus and Weis 1990). The state transitions component (qT) of non-photochemical quenching (qN) (see (Allen 1992) for review) is a physiological adaptation in plants where excess absorbed excitation energy by PSII is alternatively channelled towards PSI via phosphorylation of the major light-harvesting antenna (LHCII). Traditionally, many authors suggest that a portion of the LHCII, bound to PSII, becomes phosphorylated (p-LHCII) and migrates into nonappressed PSI rich regions. Weis and coworkers suggest that p-LHCII may not form a complex with PSI since P700-photooxidation is not (or barely) stimulated when assayed under this condition. The supercomplex PSIIβ·LHCII-P·PSI is formed (Timmerhaus and Weis 1990; Yakovlev et al., 2002). This model is not supported by the relatively low Chl a/b (Chl b being constitutive to LHCII) ratios in stroma reported by others (Guenther et al. 1988).

PSII heterogeneity does not end at that relatively simple separation of α- and β-centres. Many authors suggest that some sub-heterogeneity may exist amongst PSIIα within grana regions of thylakoid membranes. It has been suggested that PSIIα is regionally divided into three separate species as they occur in concentric circles with the stacked grana core as illustrated in figure 14 (Albertsson et al. 1990b).
Figure 14 - Heterogeneity may also exist among PSIIα centres (Albertsson et al. 1990b).
The significance of PSII heterogeneity discussed previously does not adequately address the conundrum of linear electron transport in series. How can an excited electron from PSII be transported to a position where it is able to reduce NADP⁺ if functional PSII centres are located in thylakoid domains far removed from PSI? Lateral flow of electrons throughout thylakoid membranes is thought to be coordinated by two mobile carriers, the water-soluble plastocyanin and the pool of plastoquinone (five to ten per PSII). The former, migrating within the inner thylakoid space, escorts electrons from specific sites at cyt b₆/f to PSI, and the latter, associated with the acyl lipids in the thylakoid lipid bilayer, acts as a Qₒ-cyt b₆/f oxidoreductase (Kirchhoff et al. 2000).

Shortly after the non-detergent method of thylakoid membrane fractionation was established this question was addressed. In grana core fractions (8S) it was found that the ratio or PQ/Q was 6.4. This implies that the PQ pool can accept 12-14 electrons. Surprisingly, this value is near that of whole thylakoid. Therefore, PQ present in the stroma remains unavailable to PSIIα in the grana, and is not an active participant in linear electron transport (Yu and Albertsson 1993). So, if the PQ pool of the grana minds only PSIIα, then PQ does not act as an electron shuttle within the phospholipid thylakoid bilayer between grana and stroma. It is suggested that a 1:1:1 relationship among the three electron transport complexes in higher plants: PSII, cyt b₆/f and PSI may exist. Since the PQ pool may occur as high as six molecules per PSI complex, the cyt b₆/f complex stores reducing equivalents at a value slightly greater than 2 molecules per P700, and, more than 2 molecules of PC occur for each PSI complex (illustrated in figure 15). This is consistent with (Albertsson et al. 1990b) where the authors report that the operations of the grana and stroma act independently of each other. The α-centres of each photosystem perform oxygenic non-cyclic electron transport photo-phosphorylation in the grana, while PSIβ performs cyclic electron transport photo-phosphorylation in the stroma (Albertsson et al. 1990b; Graan and Ort 1984).
Figure 15 - The stochiometric relationship among the three large electron transport complexes (PSII, Cyt b₆/f and PSI) in thylakoid membranes and their mobile electron carriers (PQ and PC) pools from original source, H₂O, to final target, NADP⁺. Electron transport inhibitors (DCMU, DBMIB and KCN) are indicated at their specific effector sites. Electron capacity relative to PSI is shown in brackets (Graan and Ort 1984).
Table 6 - Some of the early work done to characterize PSIIα and PSIIβ centres (Hodges and Barber 1986).

<table>
<thead>
<tr>
<th>Characteristic</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>PSIIα in grana and PSIIβ in stroma</td>
<td>(Anderson and Melis 1983)</td>
</tr>
<tr>
<td>PSIIα antenna &gt; PSIIβ antenna (red shifted)</td>
<td>(Thielen and Van Gorkom 1981a)</td>
</tr>
<tr>
<td>PSIIβ not associated with the 2 electron gate</td>
<td>(Thielen and van Grokum 1981)</td>
</tr>
<tr>
<td>Different primary stable electron acceptors</td>
<td>(Horton 1981)</td>
</tr>
<tr>
<td>PSIIβ can be preferentially altered by high herbicide conc.</td>
<td>(Horváth et al. 1984)</td>
</tr>
<tr>
<td>PSIIα fluorescence affected by high [Mg^{2+}], not PSIIβ</td>
<td>(Melis and Ow 1982)</td>
</tr>
<tr>
<td>Different fluorescence emission characteristics</td>
<td>(Brearley and Horton 1984)</td>
</tr>
</tbody>
</table>
Contrary to the characteristics of PSII heterogeneity listed in table 6, the biphasic data may be interpreted as being linked to different degrees of PSII connectivity, regardless of membrane differentiation or differential antenna sizes. Or, it may be described as differences among the degree of chloroplast integrity and its link to DCMU affinity (Dekker et al. 1999).

Spatial separation of protein complexes within the thylakoid membranes of higher plants offers challenges other than those considering mobile electron carriers. The diffusion of protons plays an important role in energetic coupling within chloroplasts since ATP production is directly dependent upon the translocation of protons by the ATP-synthase, far removed from the source. The non-uniform distribution of transmembrane pH differences laterally among thylakoid membranes may be a product of spatial heterogeneity (Dubinskii and Tikhonov 1997).

PSII heterogeneity is not unique. Heterogeneity has also been suggested to exist among PSI centres as well. Similar nomenclature relative to PSII heterogeneity is maintained. PSI centres with large antennae are classified as PSIα centres, whereas PSI centres associated with smaller antennae are classified as PSIβ centres consisting of 65% and 35% of all PSI centres respectively (Andreasson et al. 1988; Svensson et al. 1991). Andreasson et al. 1988 reported that P700 photooxidation kinetics clearly indicate two distinct populations of PSI, distinguishable by their relative antenna sizes, however, it remained difficult to quantify the difference. Some authors suggest that PSIα employs an antenna 30% larger than PSIβ (Andreasson et al. 1988; Svensson et al. 1991).
Figure 16 - The model of the thylakoid membrane system of higher plant chloroplasts. Spatial separation of α- and β-centres of PSII and PSI is clearly illustrated. PSIα is located in stroma exposed end membrane regions of the grana partition whereas PSIβ is found primarily in the stroma lamellae. PSIIα is found exclusively within the appressed regions of the grana partition, whereas PSIIβ is found primarily in the stroma lamellae. The cyt b6-f and ATP synthase are ubiquitous throughout the structure (Albertsson 2001; Allen and Forsberg 2001; Svensson et al. 1991).
2.3. Chlorophyll Fluorescence

The key light absorbing pigments of photosynthesis are chlorophylls. In higher plants and algae this pigment is Chl a. In primitive photosynthetic bacteria it is bacteriochlorophyll a (BChl a) or b. Without one or another of these pigments, no organism is known to be able to carry out photosynthesis (Sauer 1975). The experimental portion of this study is largely based on capturing chlorophyll fluorescence. The fact that chlorophyll is green is hardly an accident. The probability that chlorophyll will absorb a green photon in its vicinity is relatively low, nearly all are transmitted through or reflect from leaves. Blue and red photons are much more readily absorbed by chlorophyll. Upon absorption, their energies are used to raise an electron from the ground state to the second excited state or first excited state respectively. Electrons in the second excited state rapidly and radiationlessly decay to the first excited state. This loss occurs so rapidly that it cannot be captured during photosynthesis. The Pauli exclusion principle states that no two electrons can be in the same detailed quantum state. In the same orbital, two electrons must maintain opposing spin orientations. In the lowest energy states, electrons fill the orbitals in pairs. The absorbance of a photon promotes an electron from the electronic ground state (HOMO, highest occupied molecular orbital) to an excited state, (LUMO, lowest unoccupied molecular orbital). An electron in an excited state with a spin opposite its former neighbour, the excited singlet state, is likely to result from excitation energy absorption since spin reversal requires a magnetic interaction between that promoted electron and its surroundings (figure 17). Triplet states (upon a spin reversal event) are more probable from an excited singlet because the energy distribution is broad, giving rise to a decrease in the magnetic couple between the electron pair, which is inversely proportional to the interactions with its surroundings. Chlorophyll in the excited triplet state, 3Chl*, is dangerously reactive since its lifetime in a more widespread distribution exceeds 10⁻³ s, whereas the lifetime of an excited singlet (1Chl*) is closer to 10⁻⁶ s. This difference is a product of the spin reversal. A decay to the ground state would require a re-reversal of the spin polarization. It is easier to transfer the energy to O₂, which would ultimately give rise to dangerous ¹O₂ following formation of the ³O₂. Interestingly, chlorophylls in photosynthesis employ singlet excited states.
This is the primary reason why energy transfer is so fast.
Figure 17 - Schematic representation of the electronic coupling between two identical pigment molecules, P₁ and P₂. P₁ in its lowest excited state (P₁*) can either transfer its energy to P₂, initially in its ground state, giving rise to P₂* (top), or, transfer the electron to P₂, reducing it, P₂ −, and itself becoming oxidized P₁ +. Exciton transfers are common in antenna complexes whereas electron transfers are common in reaction centres (Renger 1992).
Once electrons in the second excited state (or any higher excited state) rapidly and radiationlessly decay to the first excited state, bioenergetics and photochemistry begin here (see figure 18) (Seaton and Walker 1992). This is very typical of many large aromatic molecules. Also typical of these types of molecules are a) the occurrence of the Stokes’s shift following the thermal relaxation of the higher excited state (figure 19), b) the short (5 ns) fluorescence lifetime in organic solvents, such as acetone, ether, and alcohols, c) a fluorescence yield of merely 30% in organic solvents due to competing processes for the deactivation of the excited state (see equations 14, 15, 16, and 17) and d) fluorescence emissions and absorption are governed by the same kind of relationships among wavefunctions. Therefore the fluorescence emission maintains essentially the same polarization as the long-wavelength absorption (Sauer 1975)

\[
\text{Chl}^* \xrightarrow{k_F} \text{Chl} + h\nu \quad \text{Fluorescence} \tag{14}
\]

\[
\text{Chl}^* \xrightarrow{k_i} \text{Chl} + \text{heat} \quad \text{Internal conversion, radiationless} \tag{15}
\]

\[
\text{Chl}^* \xrightarrow{k_T} \text{Chl}^T + \text{heat} \quad \text{Intersystem crossing to triplet state} \tag{16}
\]

\[
\text{Chl}^* + Q \xrightarrow{k_Q} \text{Chl} + Q^* \quad \text{Quenching} \tag{17}
\]

The reactions listed in equations 14-17 are kinetically first-order in Chl* concentration. The fluorescence lifetime of Chl* is dependent upon the rate constants, \(k_{\text{alpha}}\), for these equations. This is commonly referred to as the quantum yield, described by equation 18:

\[
\Phi_F = \frac{k_F}{k_F + k_i + k_T + k_Q[Q] + \ldots} \tag{18}
\]

The quantum yield of fluorescence \(\Phi_F\) is can be calculated by dividing the fluorescence rate by the sum of all the rates of deactivation described in equations 14-17. The rate constant \(k_F\) is an intrinsic property of the molecule, governed by both its ground and excited state quantum-mechanical properties. Its reciprocal, \(\tau_0\), the natural lifetime of the excited state, describes the
lifetime that the excited state would maintain in the absence of any competing deactivation processes. Therefore, since the denominator of equation 5 is the reciprocal of the actual (observed) lifetime, \( \tau \), equation 18 is rewritten in equation 19 (Sauer 1975)

\[
\Phi_F = \frac{\tau}{\tau_0}
\]

The fluorescence yield is dependent upon various factors, first and foremost, the redox state of intersystem electron carriers, namely the plastoquinone pool. This is most noticeable in the presence of DCMU, an electron transport inhibitor the blocks the transfer to PQ as the electron attempts to leave PSII (see figure 15). Samples inhibited with DCMU offer the highest level of fluorescence since the reaction centres of PSII remain closed (fully reduced). Secondly, the S-state sequence (figure 10) mediates the fluorescence yield. Saturating flashes effectively reduced downstream electron acceptors. The maximum initial fluorescence follows the second flash in a series (giving rise to \( S_3 \)), when steady light is applied. Furthermore, increasing electric field across the membrane, a component of the proton motive force (PMF), decreases the level of fluorescence from state P to state S witnessed in the fluorescence induction curve (Weis et al. 1987). And finally, LHCII, the accessory antenna of PSII, by undergoing reversible phosphorylation, and inducing \( q_T \), may alter the level of fluorescence following the state transition (Allen 1992; Allen and Holmes 1986; Gregory 1989; Wollman 2001).
Figure 18 - Light absorbed by chlorophyll (if R = CH₃ then Chl a, if R = CHO then Chl b) in the Qᵥ transition or the Qₓ transition excites an electron to excited states S₁ or S₂ respectively. The absorption of shorter wavelengths (Bₓ) excites an electron to S₃ however the energetic difference is lost rapidly by internal conversion (IC) as heat, similar to the loss from S₂ to S₁. If in intersystem crossing (ISC) to the triplet state (T₁), followed by phosphorescence (E'), or a radiationless decay (not shown) to the ground state (S₀) does not occur, the remaining absorbed excitation energy may be given off as fluorescence (F). Each possibility is associated with its respective rate constant (kα) (Renger 1992; Seaton and Walker 1992).
Figure 19 - Stoke's loss. The energy emitted as fluorescence (F) upon relaxation to the ground state is significantly less than the energy absorbed (A) by the molecule to promote it to the excited state (O'Connor and Phillip 1984).
2.4. Thylakoid Membrane Preparations

Evidence that PSIIα centres originate from appressed grana and PSIIβ centres originate from stroma exposed membranes is abundant. Decades ago, long before PSIIα/β heterogeneity was described, the German pioneers of chlorophyll fluorescence microscopy observed bright PSII fluorescence from grana using conventional bright-field optics (Heitz 1936), which was significantly different from the relatively non-fluorescent stroma lamellae (Metzner 1954). These results were extended by studies with confocal scanning laser microscopy which revealed brightly fluorescent grana over a non-fluorescent stroma background (van Spronsen et al. 1989). Isolation of thylakoid membranes enriched in particular pigment-protein complexes are of great interest in order to study their specific activities. The ability to isolate pigment-protein complexes in a state that closely resembles their native environment would be the ideal way to study them. Commonly, harsh detergents are employed to tear thylakoid membranes apart, and separate the pigment-protein complexes by centrifugation. BBY particles, named after Berthold, D.A., Babcock, G.T. and Yocum, C.F., who were the first to isolate them (Berthold et al. 1981), have been employed in countless studies on PSII. Triton X-100 is a nonionic detergent (octylphenol ethylene oxide condensate) which is often used in biochemical applications to solubilize proteins. It is considered a comparatively mild, non-denaturing detergent. It does absorb in the ultraviolet region of the spectrum, however, so can interfere with protein quantification. It is suggested however, that BBY particles do harbour PSIIα centres (Lam et al. 1983).

Another detergent method describes a procedure that involves n-dodecyl-α-D-maltoside (α-DM), is suggested to be relatively mild in comparison to Triton X-100. A reduced amount of chemical detergent, coupled with a minimal exposure time limits the fragmentation of the fragile macromolecular complexes to very low levels (van Roon et al. 2000). Various types of PSII supercomplexes, megacomplexes as well as heptameric associations of trimeric LHCII, the icosianamer, can be harvested with this relatively mild detergent (Dekker et al. 1999). α-DM effectively solubilizes the stroma exposed portion of the thylakoid membrane, leaving the appressed grana regions relatively intact. Appressed membrane fragments, joined in pairs,
originating from grana closely resemble PSII membranes isolated with Triton X-100, the BBY particles (Berthold et al. 1981), however stochiometry of PSII supercomplexes and their higher order of regularity is better preserved. The authors report that the most important advantage of the use of α-DM is the intactness of the solubilized fractions, which allow researchers to closely examine stromal proteins including native PSI and the ATP synthase complex as they occur in their intrinsic state. The authors also note that the α-DM treatment offers a relatively low yield of the PSII membranes and is not nearly as cost effective as the Triton X-100 procedure (van Roon et al. 2000).

The preparation of BBY particles using Triton X-100 (Berthold et al. 1981) and the α-DM isolation method (van Roon et al. 2000) differ from the isolation method developed by the Photosynthesis Group at the University of Lund, Lund, Sweden in several ways. Researchers at the University of Lund have developed a method of thylakoid membrane isolation based on native domains that is detergent free. For this study, thylakoid membranes were fractionated into five domains. The fractions were obtained via the non-detergent method illustrated in figure 20. The stacked grana were represented in three of the fractions. The grana core fraction (BS), the grana lamellae (B3) and the grana margins (Ma). The grana margin fraction is different from the other two since it is composed primarily of stroma exposed membrane. The unstacked stroma lamellae were represented in the remaining fractions, T3 and purified stroma lamellae fraction labelled Y100 (figure 21).

This novel method of isolation combines fragmentation via sonication with centrifugation in an aqueous two-phase partition system to separate different parts of the thylakoid membrane. All of the fragments are in vesicular form to maintain the highest level of photochemical activity (Albertsson et al. 1994). The details of the isolation are described in the appendix, section 8.1. Below, each of the different fractions are described.
Figure 20 - Summary of the method of non-detergent domain specific thylakoid membrane isolation employed by the Photosynthesis Group, Lund Universitet, Lund Sweden, used in this study (Albertsson et al. 1994).
Figure 21 - Thylakoid membranes are fractionated primarily into the grana fraction (B3) and the stroma fraction (T3) along the red dashed line. The B3 fraction can then be divided into the grana core (BS) and margin fraction. The Y100 fraction is a purified stroma lamellae fraction of which its origin is not precisely known (Mamedov et al. 2000).
The Grana (B3)

The appressed grana regions of the thylakoid membrane are isolated in the B3 fraction during aqueous two-phase partitioning. Following the fragmentation by sonication the B3 fraction migrates to the bottom phase (Dextran T500) of the aqueous two phase partitioning system, opposite the migration of the stroma lamellae (T3) fraction towards the upper phase (PEG 4000). Evidence supporting the notion that PSIIα centres are enriched in appressed grana membranes is well founded (Anderson and Melis 1983). Of all the chlorophyll, 63% originates form the grana, whereas the remaining 37% originates from stroma lamellae. Inside-out B3 vesicles have a diameter of 0.3 μm (Albertsson et al. 1990d).

The Stroma (T3)

The stroma, fractionated by sonication, migrates toward the upper phase of the separation vesicle, with PEG 4000. PSII in stroma occurs as PSIIβ (Anderson and Melis 1983; Guenther and Melis 1990; Henrysson and Sundby 1990; Melis and Homann 1976). Of all the reaction centres in the T3 fraction, only 20% are PSII. Another characteristic of the stroma fraction is the occurrence of the ATPsynthase within the membrane vesicles. The large CF₁ subunit that rests on top of the CF₂ subunit simply can not fit within tightly stacked thylakoid membranes that occur in grana (Allen and Forsberg 2001; Jansson et al. 1997; Miller and Staehelin 1976).

The Grana Core (B5)

The procedure mentioned in section 3.1 describes how the B3 fraction can be further fractionated into two different membrane vesicles. Inside-out B3 vesicles subjected to sonication at a relatively very low ionic strength (1 mM MgCl₂) PSI-enriched domains are sheared off and turned right-side-out. Within an aqueous two-phase system, where inside-out vesicles prefer the lower phase and right-side-out vesicles prefer the upper phase, fragmentation and separation are achieved simultaneously (Svensson and Albertsson 1989). This second round of aqueous two-phase partitioning renders a sub-fraction of inside out vesicles containing the grana core (labelled
BS). The yield, 6% from isolated whole thylakoid membranes, is considerably low. These PSII-enriched preparations are not pure PSII. About 20-30% of all reaction centres is PSI on a per chlorophyll basis. They have a diameter of 0.2 μm and they maintain their high oxygen evolving capacity. Some promising PSII preparations obtained using detergent methods (BBY particles for example) may also maintain high oxygen evolving activity but probably do not preserve the native structure of the thylakoid membrane (Berthold et al. 1981; Pålsson et al. 1990). The BS preparation in vesicle form has not been treated with detergents, and is an interesting candidate for the study of PSII in the native form. Comparisons between these two PSII preps have illustrated the differences between them clearly. Relative light-saturation curves have revealed that the slope of the plots of BBY are much less steep that those of PSIIα. The slope of the BBY actually falls between that of PSIIα and PSIIβ. The authors have concluded that the antenna size of the BBY is smaller than that of PSIIα since the detergent treatment likely disconnected some of the LHCII. Of course this is not the case upon consideration of the slope of the light-saturation curves of BS fractions rendered by the non-detergent method. The slopes are very near those of PSIIα, therefore, no evidence of LHCII disconnection was reported (Yu and Albertsson 1993).

More evidence indicating that the BS fraction maintains antenna post-preparation is described by the absorption spectra at room temperature. A stronger absorption peak attributed to Chl b compliments the low Chl a/b reported elsewhere (Albertsson and Yu 1988; Svensson and Albertsson 1989). The BS fraction reports a lower Chl a/b ratio than the whole thylakoid preparations (see table 7) which is in agreement with the relatively high Chl b content of LHCII. At lower temperatures, resolution improves and the Chl b contributions are illustrate more clearly (Yu and Albertsson 1993).

The Grana Margins (Ma)

The top phase of a second round of aqueous two-phase partitioning contains right-side out vesicles which consist of the margins, or nonappressed regions of the grana (Anderson et al. 1999). At 1 mM MgCl₂, sonication shears the PSI-enriched parts from the inside-out B₃ vesicles
and become right-side-out (Svensson and Albertsson 1989). Unlike the appressed membrane regions of the grana stack, these stroma exposed domains on the grana exterior harbour the ATP synthase consisting of $\text{CF}_0$ and $\text{CF}_1$ (Allen and Forsberg 2001; Jansson et al. 1997; Miller and Staehelin 1976; Webber et al. 1987).

**Purified Stroma (Y100)**

The T3 fraction can be fractionated further using a mechanical press to disintegrate the right-side-out vesicles. The resultant membrane fractions from the Yeda press operating at 10 MPa of nitrogen pressure are a very pure suspension of stroma lamellae labelled the Y100 fraction. The Y100 fraction is suggested to be rich in PSIIβ. Fluorescence induction kinetics reveal that the typical biphasicity, the basis for PSII heterogeneity, is absent, and only a slow rise phase is maintained. Also noted is the tremendously low photochemical yield, $F_v/F_M$, relative to not only whole thylakoid, but especially other fractions derived from supposed PSIIα-rich regions of higher plant thylakoid membranes. PSIIα:PSIIβ stoichiometry, which are typically reported as approximately 3:1 in whole thylakoid, are as low as nearly 1:9 in the Y100 fraction (Henrysson and Sundby 1990).

The thylakoid membrane vesicles described above were a generous gift F. Mamedov and R. Danielsson under the direction of S. Styring from the Photosynthesis Group, Lunds Universitet, Lund, Sweden

A protein profile of these thylakoid membrane fractions is displayed in figure 22. Table 7, describes a summary of the data obtained in a recent paper which employed these fractions. Their relative activities with respect to their spatial distribution are illustrated in figure 23. Juhler et al. 1993 published their findings on the composition of photosynthetic pigments in the thylakoid membrane vesicles. Most interestingly, β-carotene is enriched in PSI-rich fractions. Lutein is present in a 5:2 ratio relative to Chl b in all fractions. Stroma fractions (T3 and Y100) contain about nearly twice as much neoxanthin per Chl b than grana fractions (B3 and BS). Violaxanthin is evenly distributed among all fractions at a ratio of approximately 5 molecules per 100 total
chlorophyll. Generally, about 1 carotenoid is present for every 4 chlorophylls throughout the thylakoid membrane (Juhler et al. 1993). Zeaxanthin content is less than 1% of all carotenoids in all fractions. These results are best summarized in table 8. Table 9 offers a summary of the PSIIα/β stoichiometry found throughout the literature to aid in the understanding of the material employed in this study.
- SDS-PAGE in the presence of urea showing the polypeptide pattern in thylakoid membranes (lane 3) and subthylakoid membrane vesicles in lanes 1 - purified stroma, 2 - stroma lamellae, 4 - grana margins, 5 - grana, and 6 - grana core (Jansson et al. 1997).
Table 7 - PSII electron transfer activity in different domains of the thylakoid membranes from spinach (summarized from (Mamedov et al. 2000)).

<table>
<thead>
<tr>
<th>Domain</th>
<th>O₂ evolution [μmol of O₂ (mg of Chl)⁻¹ h⁻¹]</th>
<th>O₂-centres (% of total centres, (V_{H₂O}/V_{DPC}))</th>
<th>(F_\text{/}F_₀)</th>
<th>Chl (a/b)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thy</td>
<td>120</td>
<td>80</td>
<td>0.7</td>
<td>2.9</td>
</tr>
<tr>
<td>BS</td>
<td>250-300</td>
<td>91</td>
<td>1.1</td>
<td>1.8-2.0</td>
</tr>
<tr>
<td>B3</td>
<td>200-250</td>
<td>84</td>
<td>0.96</td>
<td>2.2-2.4</td>
</tr>
<tr>
<td>Ma</td>
<td>102</td>
<td>66</td>
<td>0.48</td>
<td>3.0-3.3</td>
</tr>
<tr>
<td>T3</td>
<td>80</td>
<td>43</td>
<td>0.27</td>
<td>4.5-5.0</td>
</tr>
<tr>
<td>Y100</td>
<td>0</td>
<td>0</td>
<td>0.2</td>
<td>6.0-6.7</td>
</tr>
</tbody>
</table>
Figure 23 - PSII activity is highest in the PSIIα-rich grana core region of the thylakoid membrane, relative to PSIIβ-rich stroma lamellae regions, a phenomenon that occurs opposite its degradation (Mamedov et al. 2000).
Table 8 - Summary of the pigment composition of the thylakoid membrane fractions based on a % of the total molar pigment (Juhler et al. 1993).

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Carotenoids*</th>
<th>Chlorophylls</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Neo.</td>
<td>Viola.</td>
</tr>
<tr>
<td>Thylakoid</td>
<td>2.5</td>
<td>3.7</td>
</tr>
<tr>
<td>BS</td>
<td>3.6</td>
<td>3.9</td>
</tr>
<tr>
<td>B3</td>
<td>3.4</td>
<td>3.9</td>
</tr>
<tr>
<td>Ma</td>
<td>2.6</td>
<td>4.2</td>
</tr>
<tr>
<td>T3</td>
<td>1.4</td>
<td>3.9</td>
</tr>
<tr>
<td>Y100</td>
<td>1</td>
<td>3.5</td>
</tr>
</tbody>
</table>

* Carotenoids = Neo. - neoxanthin; Viola. - violaxanthin; Lut. - lutein; β-car. - β-carotene
Table 9 - PSIIα/β stoichiometry gathered from various sources and authors.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>PSIIα</th>
<th>PSIIβ</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>% PSII</td>
<td>64-75</td>
<td>25-36</td>
<td>(Andreasson et al. 1988)</td>
</tr>
<tr>
<td>% in grana</td>
<td>66</td>
<td>33</td>
<td>(Albertsson et al. 1990b)</td>
</tr>
<tr>
<td>% in stroma</td>
<td>15-20</td>
<td>80-85</td>
<td>(Albertsson et al. 1990b)</td>
</tr>
<tr>
<td>% in BS</td>
<td>0/0</td>
<td>0/0</td>
<td></td>
</tr>
<tr>
<td>% in Y100</td>
<td>12</td>
<td>88</td>
<td>(Henrysson and Sundby 1990)</td>
</tr>
<tr>
<td>Rel. conc.</td>
<td>1.43</td>
<td>0.48</td>
<td>(Melis and Anderson 1983)</td>
</tr>
<tr>
<td>Chl a/b</td>
<td>2.4*</td>
<td>4.6*</td>
<td>(Albertsson et al. 1990b)</td>
</tr>
<tr>
<td>% total Chl†</td>
<td>38</td>
<td>3</td>
<td>(Albertsson et al. 1990b)</td>
</tr>
<tr>
<td>% PSII Chl</td>
<td>92.7</td>
<td>8.3</td>
<td>(Albertsson et al. 1990b)</td>
</tr>
<tr>
<td></td>
<td>88</td>
<td>12</td>
<td>(Melis and Anderson 1983)</td>
</tr>
<tr>
<td>Chl a/core RC complex</td>
<td>40</td>
<td>40</td>
<td>(Melis and Anderson 1983)</td>
</tr>
<tr>
<td>Chl antenna</td>
<td>240</td>
<td>130</td>
<td>(Thielen and Van Gorkom 1981b)</td>
</tr>
<tr>
<td></td>
<td>230</td>
<td>100</td>
<td>(Melis and Anderson 1983)</td>
</tr>
<tr>
<td></td>
<td>250±40</td>
<td>120±20</td>
<td>(Melis 1991)</td>
</tr>
<tr>
<td>Chl a/b proteins</td>
<td>106 Chl a</td>
<td>45 Chl a</td>
<td>(Melis and Anderson 1983)</td>
</tr>
<tr>
<td></td>
<td>84 Chl b</td>
<td>15 Chl b</td>
<td>(Melis and Anderson 1983)</td>
</tr>
<tr>
<td>Rel e⁻ transport</td>
<td>1.6</td>
<td>0.23</td>
<td>(Melis and Anderson 1983)</td>
</tr>
<tr>
<td>Q₈ reduction</td>
<td>Yes</td>
<td>No</td>
<td>(Graan and Ort 1986; Melis 1985; Thielen and van Gorkum 1981)</td>
</tr>
</tbody>
</table>

* 1.8-2.0 in BS
† 6.0-6.7 in Y100
‡ PSII accounts for 41%, PSI accounts for 59% of total Chl
2.5. Gaussian Decomposition of Absorption Spectra

Proper analysis of the absorption spectra and its Gaussian decomposition of any photosynthetic apparatus can, with significant accuracy, estimate a subset of bands that may directly correspond to the relative absence and/or presence of any particular chlorophyll species in a set of samples (Ikegami and Itoh 1988; Zucchelli et al. 1994).

Absorption changes of pigments may be influenced by several mechanisms: (a) charge-induced energy shift, (b) pigment-pigment interaction, (c) mutual polarisability interaction and (d) distortion of the conjugated double bond system via conformation induction or nearby changes. The $Q\gamma$ transition of Chl $a$ in ethanol peaks at 663 nm. The same pigment, when bound to Lhc proteins, reveals multiple absorptions in the 660 to 730 range, primarily at 677 and 670 nm with relative dipole strengths of 1:2 (Croce et al. 2000). Primary $Q\gamma$ transitions of chl $b$ occur at 652 and 644 nm with relative dipole strengths of 2:1 (Hemelrijk et al. 1992). More specifically five Gaussian bands can be resolved from isolated PSII antenna. Unlike the $Q\gamma$ absorption spectra of bacterial reaction centres which consist of four bacteriochlorophylls (Bchl) over a range of as much as 3000 cm$^{-1}$, the PSII reaction centre employs six Chl $a$ over a range of merely 500 cm$^{-1}$ (Gall et al. 1998). The absorption maxima ($Q\gamma$ transitions) are present at 660, 670, 678, 684, and 695 nm. Chl $b$ usually renders a single spectral band and usually absorbs maximally near 648 nm (Jennings et al. 1993). It is clear that the absorption spectrum of an light-harvesting complex is the product of individual chromophore absorption at different protein sites, mediated also by coupling between pigments. Attempts have been made to modify the native chlorophylls within the PSII reaction centre, but to no great achievement. Ideally, chromophore substitution would expand the $Q\gamma$ absorption range to one similar to that of the bacterial reaction centre, thereby, expediting spectroscopic discrimination of individual chromophore contributions. Nevertheless, regardless of the method of chromophore exchange, it remains unclear exactly which of the six reaction centre chlorophylls was exchanged (Gall et al. 1998). Longer wavelength absorption bands are typically associated with PSI (Andreeva and Velitchkova 1998; Cramer and Butler 1968; Kochubey and Samokhval 2000).
Linear dichroism studies have shown that Qy transition dipoles of Chl b within the PSII antenna are oriented greater than 35° out of the membrane plane. Similarly shorter wavelength Chl a molecule Qy transitions are oriented right about that same angle. Long wavelength forms of Chl a (λ > 676 nm) employ Qy transition dipoles very near to the plane of the membrane (Breton and Vermeglio 1982; Zucchelli et al. 1994).

2.6. PSII Absorbsance Cross-section

Energy transfer among Chl within the antenna of PSII is governed by the distance between them. It can occur in two ways. Resonance energy transfer (intermolecular transfer, also known as Förster transfer) depend on the degree of overlap between the fluorescence emission spectrum of the donor and the absorption band of the acceptor. This excitation transfer is not a series of fluorescence emissions followed by rapid reabsorptions, but rather a intimate coupling among the closely spaced chlorophyll molecules in the antenna. The relative orientations of the donor and the acceptor within the pigment-protein matrix affects the efficiency of the transfer. This transfer is inversely proportional to the sixth power of the distance between donor and acceptor. A distance of 2 nm between pigments takes approximately 1 ps. Förster R₀ values (critical distance) have been known for some time. Typically, a Chl a-Chl a transfers requires an R₀ of 100 Å, whereas a Chl b-Chl a transfer requires only 80-90 Å (van Grondelle 1985). The beauty of the Lhc proteins that make up the bulk of the PSI I antenna is their ability to hold chlorophylls at precise orientations and proper distances for optimal energy transfers. This distance is typically less than 5 nm, however chlorophylls held too close to one another may quench each other. Ultimately, resonance excitation energy transfer is a means to funnel exciton towards and into the reaction centre, P680.

The second method, delocalized exciton coupling (intramolecular transfers) occurs when pigment molecules are within 1.5 nm of each other. Direct interaction among molecular orbitals of two molecules exists such that excitation is shared between them. Delocalized exciton coupling occurs at much faster rates than resonance energy transfers, such that they are virtually instantaneous since the distance travelled is so short. In PSII antenna, which harbour carotenoids
(Car) as well as Chl a and Chl b, intramolecular energy transfers are required to move excitation from the short lived Car* within van der Waals distances for efficient energy transfer.

The measurement of PSII absorbance cross sections is dependent on these inter- and intramolecular interactions and quantifies the functional size of the chlorophyll network employed by the photosynthetic unit to drive primary photochemistry. Single turnovers of PSII to generate Q₅ are accomplished by pump flashes of varying intensities. The pump flash is short enough (6 ns in this study) to close PSII centres shortly before a weak probe flash used to assay the fluorescence yield. A typical pump-probe experiment measures the fluorescence yield at various pump flash intensities from zero, F₀, to saturating, Fₛₐₜ, the maximal level of fluorescence achieved with a single turnover saturating flash. Fₛₐₜ is not to be confused with Fₘ, since the fluorescence yield following a short high light single turnover flash can not reach the maximal level of fluorescence, severely underestimating Fᵥ/Fₘ. Not until the flash lasts for longer than 200 ms can Fₘ be accurately measured (Lazár 1999).

2.7. Fluorescence Induction Kinetics

Chl a fluorescence induction is a widely used technique to probe primary photochemistry non-invasively, but with high sensitivity. The data collected with this technique, employed for more than six decades, is often difficult to interpret. About ten years ago researchers began questioning whether or not to continue pursuing its elusive benefits (Holzwarth 1993; Lazár 1999). More recently however, these intriguing measurements are proving to be useful in determining biophysical parameters such as antenna size with respect to PSII heterogeneity in addition to rates of quinone reduction (Lazár et al. 2001).

The purpose of structural heterogeneity of PSII has been assayed by fluorescence induction. The area over the fluorescence induction curve is a measure of the number of photons (quanta) not released as fluorescence while open PSII centres are reduced and converted to their closed states. Conceivably, this yield may be used to perform photochemistry. Time resolution offers insight into the progress of said photochemistry. Furthermore, the size of the acceptor pool
may be estimated by measuring the size of the area if the light-induced charge separation is
maintained in these pools (Melis and Homann 1975). Semilogarithmic plots have illustrated two
kinetically different phases, one fast and nonexponential, and the other slow and exponential.
These two phases have been attributed to the two distinct forms of PSII discussed previously,
PSIIα and PSIIβ respectively (Hodges and Barber 1986).

2.8. Picosecond Fluorescence Decay Kinetics

Light energy absorbed by a chlorophyll molecule establishes an exciton with some finite
equilibration time in the antenna chlorophyll. Both thermal equilibration within protein matrices and
spatial equilibration among the entire chlorophyll antennae are considered here. Rapid
equilibration times, relative to the fluorescence lifetimes, are indicative of a trap-limited system,
since the fluorescence emission spectra are maintained constant, regardless of the excitation
wavelength. If this were not so, in a diffusion-limited system, the equilibration time is comparable
to the overall fluorescence lifetime, therefore, the excitation at far-red wavelengths would shorten
substantially the fluorescence lifetimes since molecules neighbouring the reaction centre would be
preferentially excited.

The origins of the various variable fluorescence parameters are resolved by picosecond
fluorescence decay kinetics at any given fluorescence yield, be it $F_0$, $F_M$ or $F_{SAT}$. Fluorescence
decay offers much more information than other “slower” fluorescence measurements by revealing
more intricate interpretations of fluorescence yield changes. Coupled with a suitable kinetic model
of the origins of chlorophyll fluorescence decay components, fluorescence decay measurements
are useful in the development of simulations of experimental results. Unfortunately, the time-
resolved spectroscopic techniques that attempt to measure these kinetics remain problematic since
excited state transfer processes that precede electron transfer obstruct their detection. The rate
constants derived from this measurement, however, remain useful in illustrating the primary
photochemical events in photosynthesis and add to the accurate description of a comprehensive
kinetic model of the energy transfer processes in PSII (Vasil'ev et al. 2001).
Since the primary events in photosynthetic energy transfer are multicomponent in nature, the observed kinetics depends upon not only $\lambda_{\text{exc}}$, but also the detection wavelength, $\lambda_i$. In a series of $\lambda_i$, within a certain range, at a particular $\lambda_{\text{exc}}$, the contribution of each component is very likely to change as one scans the data set. These kinetic components, or fluorescence lifetimes, can be analysed in a couple of different ways. Traditionally, these types of data were analysed separately using single-decay analysis. A biexponential decay:

$$I(t, \lambda_{\text{exc}}, \lambda_i) = \sum_{j=1}^{n} (A_j, \tau_j) \cdot \exp(-t / \tau_j)$$

(20)

with $n=2$ would render four unique parameters with two amplitudes, $A_j$, coupled to two fluorescence lifetimes. This analysis offers only these two lifetimes with the same amplitude at all $\lambda_{\text{exc}}$ and $\lambda_i$. Single-decay analysis does not account for the fact that certain fluorescence lifetimes are connected, or even constant, across all individual decay data collections. Therefore, more fitting parameters are determined from the data set than required (Holzwarth 1996).

The alternative to single-decay analysis is global analysis. A single parameter set for all experiments from a combined analysis performed in a single analysis run may be generated by taking explicit account of connected parameters within a systems across an entire experiment. In the example illustrated above with a biexponential decay where $n=2$, a relationship exists among the fluorescence lifetimes across the experiments that can be describe by the simple identity;

$$\tau_1(\lambda_1) = \tau_1(\lambda_2) = \ldots = \tau_1(\lambda_i)$$

(21)

and;

$$\tau_2(\lambda_1) = \tau_2(\lambda_2) = \ldots = \tau_2(\lambda_i)$$

(22)

This simple relationship, where one of two fluorescence lifetimes, $\tau_1$ and $\tau_2$, is associated with each detection wavelength, $\lambda_i$, in a series, illustrates how these lifetimes may be commonly connected across experiments. With global analysis of this simplified example the total number of parameters, $P$ generated from a complete data set is with $M$ experiments,
\[ P_{\text{global}} = 2 \times M + 2 \]  
\[ (23) \]

whereas, with single-decay analysis, \( P \) is:

\[ P_{\text{single}} = 4 \times M \]
\[ (24) \]

\( P_{\text{single}} \) is larger than \( P_{\text{global}} \) in all cases where \( M > 1 \). Therefore, from identical data sets, a global analysis provides the same information with less parameters. Global analysis allows increased accuracy in the values of the extracted parameters so, more complex systems, with fluorescence lifetimes very close to one another can ultimately be achieved (Holzwarth 1996).

Global analysis was first applied by Knorr and Harris, 1981. Within the realm of photosynthesis research, global analysis was first applied by Holzwarth et al., 1987, Wendler et al., 1986 and, Wendler and Holzwarth 1987. Global target fitting, and its employment within PSII heterogeneity is a decade old exercise (Roelofs et al. 1992). The authors have suggested from this pioneer paper, that PSII\( \alpha \) and PSII\( \beta \) exhibit different molecular functioning with respect to the primary processes. These differences may emerge from a different molecular structure of the reaction centres and/or a different local environment of these centres. Spatial heterogeneity was suggested as well.

Several studies have suggested that separate lifetime components can be observed from PSII\( \alpha \) and PSII\( \beta \). Lifetimes of 250 ps and 500 ps have been observed for PSII\( \alpha \) centres at \( F_o \). Meanwhile lifetimes of 100 ps and 500 ps have been reported for PSII\( \beta \) (Beauregard et al. 1991; McCauley et al. 1990; Schatz et al. 1988). At \( F_o \) it is reported that one lifetime is expected to reflect the Pheo re-oxidation kinetics of several hundred picoseconds. The other, much shorter lifetime, is believed to be proportional to the number of pigment molecules in the antenna. Accordingly, a PSII\( \beta \) centre, with 100 chlorophyll molecules associated with its antenna should have a fluorescence lifetime slightly longer than that which is observed for reaction centres isolated from \textit{Synechococcus} PCC 7942 (80 ps) which contain 80 chlorophyll molecules per antenna. Furthermore, one may also predict that the 250 chlorophyll antenna of PSII\( \alpha \) centres should emit a fluorescence lifetime three times longer than that of the cyanobacteria. Common to both \( \alpha \)- and
β- centres, is a third component of about 500 ps, originating from reaction centre primary photochemistry (McCauley et al. 1990).

The current study is a comprehensive effort with one goal. To characterize PSIIα and PSIIβ comparatively and quantitatively. Comparisons to some of the top photosynthesis research within the realm of PSII heterogeneity will aid in this endeavour. The five biophysical techniques employed in this study to describe the significance of these differences are:

- Low temperature absorption spectroscopy
- PSII absorbance cross section
- Fluorescence induction kinetics analysis
- 77K fluorescence emission spectroscopy
- Time-correlated single photon counting (TC-SPC) of chlorophyll fluorescence decay kinetics

These techniques, some of which based on tested models of the primary events in photosynthesis, have been chosen for this study for three reasons. First, three of the five have never been performed on the domain specific thylakoid membrane fractions, only on the whole thylakoids of higher plants. Secondly, they offer superior resolution to probe deeply into the structure and function of photosynthetic complexes. Lastly, together, these techniques are best suited to analyse the two modes of PSII heterogeneity discussed earlier. Antenna heterogeneity with respect to pigment composition and size will be probed by the low temperature absorption spectra and the PSII absorbance cross sections, respectively. PSII antenna size will also be described with respect to the fluorescence induction kinetics. The heterogeneity of PSIIα (Albertsson et al. 1990c; Albertsson et al. 1990d) will also be addressed. TC-SPC of chlorophyll fluorescence decay kinetics will illustrate the reducing side heterogeneity. The data presented here, representative of PSII in different domains of the thylakoid membrane, may be compared to proposed properties and distributions of PSIIα and PSIIβ centres.
3. Materials and Methods

3.1. Preparation of Thylakoid Membrane Fractions

All thylakoid membrane fractions were prepared by the photosynthesis group at the University of Lund, Lund, Sweden, under the direction of S. Styring. The nondetergent method (Albertsson et al. 1994) of isolation is described in detail in appendix 8.1.

3.2. Gaussian Decomposition of Absorption Spectra

Measurements were obtained at 10K using a cryostat (Advanced Research Systems, Inc., model DE-202, Allentown, PA) coupled with a helium compressor (APD Cryogenics, Inc., model HC-2D-1, Allentown, PA) controlled by a microprocessor-based digital temperature indicator/controller (Scientific Instruments, Inc. Series 9600, West Palm Beach, CA). Thylakoid fractions were resuspended in resuspension buffer (see appendix 1) with 60% glycerol as a cryoprotectant and sealed between two, 2.5 cm polycarbonate lexan discs, separated by a 2 mm thick PVC ring and sealed in the cryostat chamber. A beam of white light was cast through a pinhole towards the chamber. Transmitted light was then sent through frosted glass towards an intensified diode array detector interface (model 1461 EG&G Princeton Applied Research, Princeton, NJ). The resultant signal was plotted as the log of transmittance (minus the background) against wavelength.

Nonlinear curve fitting was done with the commercial software package Origin 4.1. Ten Gaussian sub-bands where strategically placed according to major negative peaks present in the second-order derivative of the absorption spectra similar to (Schmid et al. 2001). The shape of each sub-band was determined by the equation:

\[ y = y_0 + \frac{A}{w \sqrt{\pi/2}} e^{-\frac{(x-x_c)^2}{w^2}} \]  

where \( y \) is the area contained beneath the curve, \( w \) is its half-width in nanometres and \( x_c \) is the centre wavelength of its peak. All thylakoid membrane fractions were fitted globally by sharing nine
of the ten peak centres (all but x<sub>c</sub>7 at ~676 nm) and letting their half-maximum widths and contained areas free using the Levenberg-Marquardt non-linear curve fitting algorithm to a reduced Chi-square of 6.68 x 10<sup>-6</sup>.

3.3. PSII Absorbance Cross-section

The pump-probe method of fluorimetry was used to determine the absorption cross-section of PSII. A pulsed Neodymium:Yttrium-Aluminum-Garnet laser, Nd:YAG (Spectron Laser Systems, Rugby, U.K. model SL456G-10) coupled with a Beta-Barium Borate Optical Parametric Oscillator (GWU-Lasertechnik VirI R 2, Erftstadt, Germany) was used to fire a 6 ns pump flash at 435 nm via a fibre optic cable to a 250 μL flow through cuvette (~1 mL/sec) containing a thylakoid preparation (5 μg·Chl·ml<sup>-1</sup>) to obtain F<sub>SAT</sub>, the saturating level of fluorescence induced by a single-turnover pulse of light. Non-actinic 60 μs probe pulses of light were supplied by four low intensity light-emitting diodes, at 450 nm excitation, focussed at 90° to the direction of the pump flash 100 μs after the pump flash. Chlorophyll fluorescence was measured at 90° to the direction of the probe flash from the cuvette through a fibre optic light guide towards a photomultiplier tube (Hamamatsu RG967) behind the monochromator. In the absence of the pump flash, the level of chlorophyll fluorescence obtained from probe flashes alone represented F<sub>0</sub>.

A motorized light polarizer was used to adjust the flash energy before the beam was cast on the flow through cuvette. Relative intensities of each pump flash were gathered from reflected actinic light from a glass slide positioned between the light polarizer and the fibre optic, to a photodiode connected to a 9-volt biasing battery. The pump-probe triggers, photodiode and the photomultiplier tube are connected to channels 1, 2 and 3, respectively of a digital storage oscilloscope (Tektronics, model TDS 540) and uploaded to a computer data collection program. Each data point was the average of 20 signals at a pulse frequency of 2.5 Hz. Flash saturation curves were acquired using a set program of positions on the motorized light polarizer in the path of the pump pulse (Figure 24).
Figure 24 - PSII absorbance cross-sections were collected using this experimental setup. Increasing relative fluorescence was measured as flash intensity increased and managed by the DOS-based software program GLIB.
Curve-fitting was performed with Origin 4.01. The cumulative Poisson single-hit probability distribution (Mauzerall and Greenbaum 1989) was used to fit the data:

\[ \Phi(I) = \Phi_{\text{MAX}} \left(1 - e^{-I \sigma}\right) \]  

(26)

For variable fluorescence from \( F_0 \) to \( F_M \), equation 26 is written as:

\[ F(I) = F_0 + F_M \left(1 - e^{-I \sigma}\right) \]  

(27)

where \( I \) is the flash intensity, \( \Phi(I) \) is the yield of photoproduction measured (fluorescence), \( \Phi_{\text{MAX}} \) is the maximum yield (\( F_M \)) and \( \sigma \) is the effective absorption cross-section, representing a homogeneous population of photosystems.

3.4. Fluorescence Induction Kinetics

The photochemical events mediated by PSII can be measured with room temperature fluorescence induction kinetics as an intrinsic probe. At room temperature fluorescence is emitted primarily from PSII. The contribution from PSI is merely 20% of the fluorescence signal in whole thylakoid. The first fluorescence induction experiments were conducted by Kautsky and Hirsh in 1931, hence it is often referred to as the Kautsky effect. Basically, fluorescence induction represents a plot of the measured fluorescence intensity as a function of time of continuous illumination (Lazár 1999).

The theoretical model of chl \( a \) fluorescence induction with consideration of a heterogenous population of PSII is described in Strasser and Stirbet, 1998. This model is based solely on the acceptor side of PSII (referred to as the complex model (Stirbet and Strasser 1996)). The portion of electron transport chain which involves Pheo, \( Q_A \) and the PQ pool are included in their simulations (Strasser and Stirbet 1998).
The theoretical relative variable fluorescence, \( V(t) \), of PSII centres of type \( i \) can be calculated by equation 31, reported in (Strasser and Stirbet 1998; Strasser and Stirbet 2001):

\[
V_i(t) = \frac{B_i(t)}{1 + C[1 - B_i(t)]}
\]

(28)

where \( B_i(t) \) is the fraction of the closed reaction centres (reduced QA) of the unit type \( i \) with:

\[
C = \frac{p(F_M - F_0)}{F_0}
\]

(29)

where \( C \) is the parameter for the curvature of the hyperbola, and \( p \) is the overall probability of connectivity between the PSII units.

According to Strasser's model, PSII\( \beta \) centres are considered isolated units (therefore \( C_\beta = 0 \)). So, if \( C=1 \) for a mature plant, on average, then a heterogenous system with 60% of the total PSII centres being \( \alpha \)-centres, \( C_\alpha \), is predicted to be equal to 1.67. Subsequently, the total relative variable fluorescence, \( V_{sum}(t) \) can be calculated as:

\[
V_{sum}(t) = \sum_i f_i V_i(t)
\]

(30)

where \( f_i \) is the fraction of PSII centres of type \( i \) relative to the total number of PSII centres in the sample (Strasser and Stirbet 1998).

Limited sample quantities allowed for the collection of fluorescence induction kinetics from the whole thylakoid membrane vesicles, B3, grana partition vesicles and T3, stroma lamellae vesicles only. For each sample a small volume (3 mL in a quartz cuvette) of membrane vesicles at a chlorophyll concentration of 10 \( \mu \)g/mL was dark adapted for 5 min. A 650 nm beam triggered by a pulse generator (Hewlett Packard 8011A), pulse width 200 ms at 500 \( \mu \)mol·m\(^{-2}\)·s\(^{-1}\), was focussed on a randomizing fibre optic cast on the cuvette 90° to the photomultiplier (Hamamatsu
RG967) screened by a monochromator with a 680 nm longpass filter. The data were visualized on a digital storage oscilloscope (Tektronics, model TDS 540) and uploaded to a computer data collection program (GLIB). The averages of four kinetic traces was plotted semi-logarithmically as relative fluorescence (a.u.) versus time (s). During the fitting procedure, performed with a user-defined fit function in Microcal Origin 4.1, a model for homogeneous populations was employed. The more complex heterogenous model was unnecessary since fractionated thylakoid membranes offer nearly pure PSIIα and PSIIβ from the B3 fraction and the T3 fraction respectively. Also, the connectivity constant in equation 31 was left free during the fit to solve for $B(t)$. Values of $B(t)$ were calculated at the slow initial rise of fluorescence, $B(O)$, the inflection point of the exponential rise, $B(M)$, and the final plateau, $B(P)$.

### 3.5. 77K Fluorescence Emission Spectra

A small volume (30-40 µL) of thylakoid membrane fractions at a chlorophyll concentration of 10 µg/ml in resuspension media were frozen at 77K in a heat-sealed Pasture pipette. At this temperature, the resolution of the multiple pigment-protein complexes is greatly enhanced. A monochromatic beam (435 nm) from an arc lamp screened with a monochromator (ScienceTech, model 9055) struck the sample 90° to the angle of emission detection by an intensified diode array detector interface (model 1461 EG&G Princeton Applied Research, Princeton, NJ). The resultant signal was plotted as relative fluorescence (a.u.) versus wavelength.

### 3.6. Picosecond Fluorescence Decay Kinetics

Measurement of time dependent fluorescence intensity may offer tremendous insight into the primary events in photosynthesis. More explicitly, this measurement allows researchers to predict, with significant accuracy, the relative fates of P680⁺. Time-resolution is obtained when the excitation beam is pulsed (or modulated), rather than continuous in intensity. The signal intercepted is commonly the response to an optical excitation this brief (within the picosecond range in this study) excitation pulse at a particular wavelength $\lambda_{exc}$ and is time-resolved by the
detector (Sauer and Debreczeny 1996). The arrival of a photon released by fluorescence at the
detector is compared to the time of the excitation pulse. These photons arriving at particular time
delays relative to the excitation pulse are gated into channels of a predetermined temporal width.
A multichannel analyzer collects the fluorescence decay kinetic over a time scale determined by
the time per channel, multiplied by the number of channels. In this study the time resolution is
approximately 0.012 ns per channel on about 1000 channels. High signal-to-noise data in a
relatively short time is permitted since fluorescence photons at multiple time delays are collected
simultaneously. Time resolution is limited only by the pulse width of the excitation beam and the
time response of the emission detector (Sauer and Debreczeny 1996).

A single photon timing apparatus driven by a picosecond pulsed diode laser was used to
determine the kinetics of chlorophyll fluorescence decay on a sub-nanosecond time scale. (Bruce
and Miners 1993). Excitation pulses were delivered at 406 nm from a pulsed diode laser
(PicoQuant GmbH, Berlin, Germany, PDL 800-B, model LDH-C-2-042) with a pulse width of 54
ps (figure 25). Chlorophyll fluorescence was measured from the sample (5 μg·Chl·ml⁻¹) by a
Hamamatsu R-3809 microchannel plate photomultiplier behind a double monochromator (model
DH-10, Instruments SA, Inc., Metuchen, NJ), 90° to the angle of incident light yielding and
instrument response function (IRF) of approximately ~58 ps. This IRF was collected as a
background decay at 406 nm under experimental conditions. Data are collected as voltage when
a photon emitted as chlorophyll fluorescence. Upon its capture, the time difference from excitation
is used to plot its amplitude with respect to decay time. It is done this way since the entire
fluorescence yield is produce by only about 10% of the photons from the pulsed diode laser. If the
data collection started upon trigger of the excitation beam, the chlorophyll fluorescence signal
would almost never be detectable. The electronic design of SPC apparati discussed in (Ainbund
Figure 25 - The PicoQuant, PDL 800-B with the LDH-C 400 laser head.
Figure 26 - Single-photon counting apparatus employed in picosecond chlorophyll fluorescence decay kinetics studies.
Measurements when PSII centres are open ($F_o$) and when PSII centres are closed ($F_m$) were performed at an excitation frequency of 5 MHz and 10 MHz and cold flow rates of $\sim$1 mL/s and $\sim$4 mL/s respectively.

Fluorescence decay data were collected for nine detection wavelengths between 675 nm and 730 nm at $F_m$, with 20,000 counts in the peak channel, each of which representing approximately 0.012 ns.

Data analyses were performed using a global target analysis and fitting package on a Linux-based system written by S. Vasil’ev similar to (Vasil’ev et al. 1998). Global analysis is a procedure in which all decays from the various emission wavelengths collected are analysed simultaneously in order to minimize the number of free parameters to fit. The fluorescence emission decay kinetic traces collected at each of the nine wavelengths where compared to the room temperature fluorescence emission spectra and fit, simultaneously for the desired number of components. As a general rule, more components are easier to fit to the data than fewer components. The resultant plot is a decay associated spectra (DAS). The DAS illustrate the relative contributions of each of the components in the fitting parameters to the spectra of emission wavelengths collected. Each decay component is measured with respect to the fluorescence lifetime and may be assigned to an early event in photosynthesis.

In each DAS fit with five unique exponentials a 120 ps to 260 ps component at $F_m$ and a 100 ps to 140 ps component at $F_o$ can be subsequently deconvoluted in to two separate components. A PSII-type component with an amplitude most dominant in the short wavelength region of the DAS and a PSI-type component, most dominant in the long wavelength region of the DAS are easily resolved from the mixed component by calculating the sum of a known PSII decay (multiplied by a factor) and a known PSI decay. Therefore, the result is two distinct components with identical lifetimes representative of PSII and PSI from a single mixed component with that lifetime.
4. Results

4.1. Gaussian Decomposition of Absorption Spectra

It is reasonable to suspect that each specialized thylakoid membrane fraction may offer its own unique pigment-protein profile. Low temperature absorption spectroscopy, and its very fine resolution, offers the best opportunity to probe this uniqueness. Each absorption spectrum (figures 27-32) is plotted as absorption (a.u.) versus wavelength (nm). Ten Gaussian bands have been resolved in each spectrum. The analyses of these spectra is described in table 10. Table 22 (see appendix 8.2) offers insight into their relative assignments.

The ratio of Chl $a/b$ was calculated by the sum of the Chl $a$ absorption bands (multiplied by the molar extinction coefficient) divided by the Chl $b$ band (multiplied by the molar extinction coefficient). Appendix 8.3 shows a plot of these molar extinction coefficients in cm$^{-1}$/M versus wavelength (Du et al. 1998).
Figure 27 - Ten Gaussian sub-bands were fit globally to all five thylakoid membrane fractions simultaneously. Each plot represents the sub-bands (green), the data (blue) and the fit (red).
Figure 28 - Global Gaussian decomposition of the 10K absorption spectrum (black) of the grana fraction (B3) and its fit (red). Green bands (centred at nm, nm and nm) represent red shifted PSI components, whereas the orange band represent chl b Q transition (centred at nm), primarily associated with PSII.
Figure 29 - Global Gaussian decomposition of the 10K absorption spectrum (black) of the stroma fraction (T3) and its fit (red). Green bands (centred at nm, nm and nm) represent red shifted PSI components, whereas the orange band represent chl b Qy transition (centred at nm), primarily associated with PSII.
Figure 30 - Global Gaussian decomposition of the 10K absorption spectrum (black) of the grana core fraction (BS) and its fit (red). Green bands (centred at nm, nm and nm) represent red shifted PSI components, whereas the orange band represent chl $b$ $Q_{y}$ transition (centred at nm), primarily associated with PSII.
Figure 31 - Global Gaussian decomposition of the 10K absorption spectrum (black) of the grana margins fraction (Ma) and its fit (red). Green bands (centred at nm, nm and nm) represent red shifted PSI components, whereas the orange band represent chl b Q\textsubscript{y} transition (centred at nm), primarily associated with PSII.
Figure 32 - Global Gaussian decomposition of the 10K absorption spectrum (black) of the purified stroma fraction (Y100) and its fit (red). Green bands (centred at nm, nm and nm) represent red shifted PSI components, whereas the orange band represent chl $b$ $Q_{y}$ transition (centred at nm), primarily associated with PSII.
Table 10 - Ratio of the Chl $a/b$ (Q$_y$-transition) for thylakoid membrane fractions from Gaussian bands in figures 27 - 32.

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Chl $a/b$ (Q$_y$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BS</td>
<td>3.2</td>
</tr>
<tr>
<td>B3</td>
<td>3.0</td>
</tr>
<tr>
<td>Ma</td>
<td>4.0</td>
</tr>
<tr>
<td>T3</td>
<td>5.3</td>
</tr>
<tr>
<td>Y100</td>
<td>4.6</td>
</tr>
</tbody>
</table>
4.2. PSII Absorbance Cross-section

Antenna side heterogeneity is illustrated further upon examination of the absorption cross-sections of PSII as it occurs in different thylakoid membrane fractions. Each flash saturation curve is plotted as relative fluorescence (a.u.) versus the log of light intensity, each step was kept constant among all membrane fractions. PSII absorbance cross-sections are easily interpreted by the coordinates of the inflection point of the sigmoidal curve. Relatively large PSII centres (with respect to their antennae) will demonstrate curves that are shifted to the left, or towards lower photon densities. This is very apparent upon examination of the PSII absorbance cross-section of the grana core (figure 34).

The opposite is true for smaller PSII centres. The probability of striking a small PSII antenna at lower photon densities is quite low. Therefore, the PSII absorbance cross-section of thylakoid fractions with small chlorophyll antennae will be shifted to the right, or towards higher photon densities. The purified stroma fraction illustrates this phenomenon very well (figure 34).

PSII absorbance cross-sections of the grana, stroma lamellae, and grana margins are presented in figures 33 and 34 respectively. Table 11 offers a summary of the data demonstrated by these measurements. The $\delta$-value is numerical measure of the effective absorbance cross section. They have been normalized to a grana core relative absorbance cross section of 1. The grana core employs the largest relative absorbance cross section, $\delta = 1$, whereas the purified stroma employs the smallest, $\delta = 0.20$. 
Figure 33 - Flash saturation curves of the grana fraction (B3) and the stroma lamellae fraction (T3). Each point represents the average of twenty individual single-turnover flashes. Lines are best fits to Poisson distribution with a computer-generated error of ~3%.
Figure 34 - Flash saturation curves of the grana core fraction (BS) and the purified stroma lamellae fraction (Y100). Each point represents the average of twenty individual single-turnover flashes. Lines are best fits to Poisson distribution with a computer-generated error of ~3%.
Table 11 - PSII absorbance cross section ($\delta$) obtained from flash saturation curves, figures 33 and 34. The fit error is no more than $\pm 5\%$. 

<table>
<thead>
<tr>
<th>Fraction ($\chi^2$)</th>
<th>$\delta$ (Relative Cross section*)</th>
</tr>
</thead>
<tbody>
<tr>
<td>BS</td>
<td>1</td>
</tr>
<tr>
<td>B3</td>
<td>0.90</td>
</tr>
<tr>
<td>Ma</td>
<td>0.31</td>
</tr>
<tr>
<td>T3</td>
<td>0.27</td>
</tr>
<tr>
<td>Y100</td>
<td>0.20</td>
</tr>
</tbody>
</table>

* Relative to the grana core fraction with a fixed cross section of 1.
4.3. Fluorescence Induction Kinetics

The rise of chlorophyll fluorescence has been an intrinsic probe of both antenna heterogeneity and reducing-side heterogeneity. Here the rate of PSII closure may be measured by applying model fitting functions to experimentally obtained data. Figures 35, 36 and 37 are semilogarithmic plot of the fluorescence rise curves for whole thylakoid membrane vesicles, stacked grana membrane vesicles and unstacked stroma lamellae vesicles respectively. With each curve its fit, generated by Strasser's model for homogenous populations of PSII, is plotted as well (Strasser and Stirbet 1998; Strasser and Stirbet 2001).
Figure 35 - Fluorescence induction kinetic from whole thylakoid and its fit (bold) using Strasser's model for connectivity within homogeneous PSII populations (Strasser and Stirbet 1998; Strasser and Stirbet 2001).
Figure 36 - Fluorescence induction kinetic from B3, grana partition, and its fit (bold) using Strasser’s model for connectivity within homogeneous PSII populations (Strasser and Stirbet 1998; Strasser and Stirbet 2001).
Figure 37 - Fluorescence induction kinetic from the T3, stroma lamellae, fraction thylakoid membranes and its fit (bold) using Strasser's model for connectivity within homogeneous PSII populations (Strasser and Stirbet 1998; Strasser and Stirbet 2001).
Table 12- Connectivity and rate constants obtained from the fitting fluorescence induction kinetic traces with Strasser’s model for homogenous populations (Strasser and Stirbet 1998; Strasser and Stirbet 2001).

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Connectivity</th>
<th>Rate constant</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thylakoids</td>
<td>1.89</td>
<td>941.48</td>
</tr>
<tr>
<td>(3.6 x 10^{-4})</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Grana</td>
<td>2.08</td>
<td>1350.19</td>
</tr>
<tr>
<td>(2.2 x 10^{-4})</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Stroma Lamellae</td>
<td>0.00</td>
<td>956.68</td>
</tr>
<tr>
<td>(2.45 x 10^{-3})</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
4.4. 77K Fluorescence Emission Spectra

The 77K fluorescence emission spectra in figure 38 have been clearly divided into PSII-rich thylakoid membrane fractions and PSI-rich thylakoid membrane fractions. These high resolution spectra were assayed for their relative ratios of typical PSII emissions ($F_{695}$) to typical PSI emissions ($F_{730}$). The $F_{695}/F_{730}$ is useful in determining PSII:PSI ratios, employed in the kinetic modelling of the primary events in PSII performed in section 4.6. Also determined from these spectra are the $F_{665}/F_{695}$ ratios. This parameter offers insight into the degree of LHCII association among the thylakoid membrane fractions. Both of these measurements are presented in table 13.
Figure 38 - 77K fluorescence emission spectra of PSII-rich thylakoid membrane fractions (left) and PSI-rich thylakoid membrane fractions (right) and whole thylakoid membranes (inset). See table 13.
Table 13 - Ratios of $F_{695}/F_{730}$ and $F_{685}/F_{695}$ derived from the 77K fluorescence emission spectra in figure 38. The error of these spectra is less than 3%.

<table>
<thead>
<tr>
<th>Fraction</th>
<th>$F_{695}/F_{730}$ (a.u.)</th>
<th>$F_{685}/F_{695}$ (a.u.)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thylakoid (Thy)</td>
<td>0.88</td>
<td>0.60</td>
</tr>
<tr>
<td>Grana Core (BS)</td>
<td>2.20</td>
<td>0.88</td>
</tr>
<tr>
<td>Grana (B3)</td>
<td>1.62</td>
<td>0.81</td>
</tr>
<tr>
<td>Margin (Ma)</td>
<td>0.29</td>
<td>0.73</td>
</tr>
<tr>
<td>Stroma Lamellae (T3)</td>
<td>0.34</td>
<td>0.65</td>
</tr>
<tr>
<td>Purified Stroma (Y100)</td>
<td>0.08</td>
<td>--</td>
</tr>
</tbody>
</table>
4.5. Picosecond Fluorescence Decay Kinetics

The primary events in photosynthetic energy transfer and electron transport towards the production of stored chemical energy are under investigation here. These analyses are useful in predicting the earliest events in photosynthesis, up to the reduction of $Q_A$. Picosecond chlorophyll fluorescence decay kinetics are collected in two states. The first of these is $F_M$. Simply by slowing the sample pump speed to the measurement cuvette to 1 mL/s, increasing the frequency of the pulsed diode laser (407 nm) to 10 MHz and applying a saturating halogen lamp upon excitation, PSII centres are essentially closed ($Q_A$ reduced) and the chlorophyll fluorescence level is at a maximum. The herbicide DCMU is used to enhance the certainty that $F_M$ is achieved. The addition of DCMU also permits the use of a reduced intensity saturating light intensity (relative to DCMU-absent measurements), since higher intensities damage PSII centres (Aro et al. 1993; Krause 1988; van Wijk et al. 1993).

Each emission wavelength is associated with a weighted residual demonstrating how well the decay curve fits along the ~12 ns time window (see figure 39 for example). Weighted residual plots in the appendix for whole thylakoid and each of the five membrane fractions at $F_M$ reveal a good distribution about the median in every wavelength with the exception of some noise at the fit start.

Figures 40, 41, 42, 43, 44 and 45 are the globally fit DAS at $F_M$ for all six thylakoid membrane preparations including whole thylakoid, the grana fraction, the stroma lamellae fraction, the grana core fraction, the grana margin fraction and the purified stroma lamellae fraction respectively. Each DAS plots the amplitude of four components of a five component fit versus the fluorescence emission wavelength.
Figure 39 - Picosecond decay kinetics from dark adapted whole thylakoid membranes (685 nm fluorescence emission) representative of the difference between $F_M$ and $F_0$, including the instrument response function (IRF).
Figure 40 - Decay associated spectra of globally fitted picosecond chlorophyll fluorescence decay kinetics from whole thylakoid at F<sub>M</sub>. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Figure 41 - Decay associated spectra of globally fitted picosecond chlorophyll fluorescence decay kinetics from grana core at $F_M$. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Figure 42 - Decay associated spectra of globally fitted picosecond chlorophyll fluorescence decay kinetics from whole grana at $F_M$. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Figure 43 - DAS of globally fitted picosecond chlorophyll fluorescence decay kinetics from grana margins at $F_M$. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Figure 44 - DAS of globally fitted picosecond chlorophyll fluorescence decay kinetics from whole stroma lamellae at $F_v$. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Figure 45 - DAS of globally fitted picosecond chlorophyll fluorescence decay kinetics from purified stroma lamellae at $F_M$. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Table 14 - Fluorescence lifetimes, and their relative yield (%), from global fitting analysis for thylakoid fractions at $F_M$. $T_1$ of 11 ps (exciton equilibrium (Roelofs et al. 1992)) has been omitted. Uncertainty of lifetimes and yields is approximately 10%.

<table>
<thead>
<tr>
<th>Fraction ($\chi^2$)</th>
<th>Fluorescence Lifetime (ns)</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$T_2$</td>
<td>$T_3$</td>
</tr>
<tr>
<td>Thy (1.216)</td>
<td>0.12</td>
<td>0.58</td>
</tr>
<tr>
<td></td>
<td>(15.7*, 22.5*)</td>
<td></td>
</tr>
<tr>
<td>BS (1.162)</td>
<td>0.16</td>
<td>0.65</td>
</tr>
<tr>
<td></td>
<td>(20.1*, 4.4*)</td>
<td></td>
</tr>
<tr>
<td>B3 (1.1)</td>
<td>0.14</td>
<td>0.59</td>
</tr>
<tr>
<td></td>
<td>(18.3*, 7.7*)</td>
<td></td>
</tr>
<tr>
<td>Ma (1.135)</td>
<td>0.13</td>
<td>0.54</td>
</tr>
<tr>
<td></td>
<td>(20.5*, 23.2*)</td>
<td></td>
</tr>
<tr>
<td>T3 (1.25)</td>
<td>0.09</td>
<td>0.24</td>
</tr>
<tr>
<td></td>
<td>(61.5)</td>
<td>(7.0*, 16.4*)</td>
</tr>
<tr>
<td>Y100 (1.191)</td>
<td>0.09</td>
<td>0.26</td>
</tr>
<tr>
<td></td>
<td>(80.8)</td>
<td>(3.5*, 8.2*)</td>
</tr>
</tbody>
</table>

* - deconvoluted PSII component
+ - deconvoluted PSI component
At $F_o$, the measurement is very similar to that at $F_M$. Experimentally, these kinetics are determined with a few minor differences in the equipment set up relative to the $F_M$ collection set up. The flow through the pump is increased to $\sim 4 \text{ mL/s}$ from $\sim 1.5 \text{ mL/s}$, the frequency of the pulsed laser diode is decreased to 5 MHz from 10 MHz and the saturating halogen lamp is set to off. These technical alterations, coupled with the absence of the herbicide, DCMU, permit PSII centres to remain open ($Q_A$ oxidized). At $F_o$, the chlorophyll fluorescence decays much faster than those occurring at $F_M$. This is demonstrated in figure 39. Weighted residuals, similar to those obtained at $F_M$ are displayed in figure 56 for all six thylakoid membrane preparations. The globally fit DAS generated from chlorophyll fluorescence decay curves at $F_o$ are shown in figures 46, 47, 48, 49, 50 and 51 for whole thylakoid, the grana fraction, the stroma lamellae fraction, the grana core fraction, the grana margin fraction and the purified stroma lamellae fraction respectively. Each DAS plots the amplitude of four components of a five component fit versus the fluorescence emission wavelength.
Figure 46 - Decay associated spectra of globally fitted picosecond chlorophyll fluorescence decay kinetics of whole thylakoid at $F_0$. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Figure 47 - Decay associated spectra of globally fitted picosecond chlorophyll fluorescence decay kinetics from the grana core at $F_0$. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Figure 48 - Decay associated spectra of globally fitted picosecond chlorophyll fluorescence decay kinetics of whole grana at F₀. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Figure 49 - Decay associated spectra of globally fitted picosecond chlorophyll fluorescence decay kinetics grana margins at $F_0$. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Figure 50 - Decay associated spectra of globally fitted picosecond chlorophyll fluorescence decay kinetics whole stroma lamellae at $F_0$. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Figure 51 - Decay associated spectra of globally fitted picosecond chlorophyll fluorescence decay kinetics of purified stroma lamellae at $F_0$. Four components of a five component fit are plotted. An 11 ps component has been omitted (Durrant et al., 1992).
Table 15 - Fluorescence lifetimes, and their relative yield (%), from global fitting analysis for thylakoid fractions at F₀. T₁ of 11 ps (exciton equilibrium (Roelofs et al. 1992)) has been omitted. Uncertainty of lifetimes and yields is approximately 10%.

<table>
<thead>
<tr>
<th>Fraction (χ²)</th>
<th>Fluorescence Lifetime (ns)</th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>T₂</td>
<td>T₃</td>
<td>T₄</td>
<td>T₅</td>
<td></td>
</tr>
<tr>
<td>Thy (1.18)</td>
<td>0.12</td>
<td>0.34</td>
<td>0.67</td>
<td>4.6 (&lt; 1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(33.6*, 12.7*)</td>
<td>(39.8)</td>
<td>(13.4)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>BS (1.331)</td>
<td>0.11</td>
<td>0.30</td>
<td>0.67</td>
<td>7.98 ( &lt; 1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(57.5*, 2.4*)</td>
<td>(38.2)</td>
<td>(1.9)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>B3 (1.293)</td>
<td>0.10</td>
<td>0.30</td>
<td>0.54</td>
<td>5.04 ( &lt; 1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(48.2*, 5.8*)</td>
<td>(38.6)</td>
<td>(7.3)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ma (1.281)</td>
<td>0.10</td>
<td>0.31</td>
<td>0.84</td>
<td>4.35 ( &lt; 1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(48.8*, 24.4*)</td>
<td>(24.1)</td>
<td>(2.6)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>T3 (1.25)</td>
<td>0.04</td>
<td>0.14</td>
<td>0.43</td>
<td>2.72 ( &lt; 1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(37.1)</td>
<td>(38.4*, 12.7*)</td>
<td>(16.8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Y100 (1.27)</td>
<td>0.04</td>
<td>0.14</td>
<td>0.43</td>
<td>2.38 ( &lt; 1)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>(71.7)</td>
<td>(6.5*, 18.3*)</td>
<td>(3.3)</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* - deconvoluted PSII component
+ - deconvoluted PSI component
A visual analysis of the DAS clearly indicate the 100 to 200 ps component (T_2 in grana derived fractions, and T_3 in stroma lamellae derived fractions) consists of a mixture of PSII and PSI fluorescence decays. Representative PSII and PSI components were used to deconvolute these mixed spectra into separate PSII and PSI components maintaining their respective lifetimes. This process offered two advantages. First, it is now clear what amount of these mixed spectra is associated with PSII fluorescence decay and PSI fluorescence decays. The DAS at F_M, figures 40-45, at F_o, figures 46-51, illustrate this deconvolution. The newly calculated PSII and PSI components are represented by broken lines. They are plotted in the same colour as the original mixed component that they were deconvoluted from. The addition of these novel spectra renders a near perfect match their respective two component origins. Secondly, the normalized yield of the PSII component derived from the mixed component, coupled to the remaining original PSII components in the DAS permit a more precise calculation of F_v (calculated by F_M - F_o) associated with each thylakoid fraction. The parameter F_v/F_o was utilized by Mamedov et al., 2000, to describe the robustness of the same thylakoid membrane fractions as those employed in the current study. The values that they obtained are relatively low due to the method used to measure these fluorescence levels. These data are presented in tables 16, 17 and 18.
Table 16 - Ratios of the relative contributions of deconvoluted fluorescence decay components from the DAS at $F_M$ (figures 40-45).

<table>
<thead>
<tr>
<th>Fraction</th>
<th>PSII/PSI ($T_2$)</th>
<th>PSII/PSI ($T_3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thylakoid</td>
<td>0.7</td>
<td>N.D.</td>
</tr>
<tr>
<td>Grana Core</td>
<td>4.57</td>
<td>N.D.</td>
</tr>
<tr>
<td>Grana</td>
<td>2.38</td>
<td>N.D.</td>
</tr>
<tr>
<td>Grana Margins</td>
<td>0.88</td>
<td>N.D.</td>
</tr>
<tr>
<td>Stroma Lamellae</td>
<td>N.D.</td>
<td>0.43</td>
</tr>
<tr>
<td>Purified Stroma Lamellae</td>
<td>N.D.</td>
<td>0.43</td>
</tr>
</tbody>
</table>

Table 17 - Ratios of the relative contributions of deconvoluted fluorescence decay components from the DAS at $F_O$ (figures 46-51).

<table>
<thead>
<tr>
<th>Fraction</th>
<th>PSII/PSI ($T_2$)</th>
<th>PSII/PSI ($T_3$)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Thylakoid</td>
<td>2.65</td>
<td>N.D.</td>
</tr>
<tr>
<td>Grana Core</td>
<td>23.96</td>
<td>N.D.</td>
</tr>
<tr>
<td>Grana</td>
<td>8.3</td>
<td>N.D.</td>
</tr>
<tr>
<td>Grana Margins</td>
<td>2.02</td>
<td>N.D.</td>
</tr>
<tr>
<td>Stroma Lamellae</td>
<td>N.D.</td>
<td>3.02</td>
</tr>
<tr>
<td>Purified Stroma Lamellae</td>
<td>N.D.</td>
<td>0.36</td>
</tr>
</tbody>
</table>
Table 18 - Comparison of variable fluorescence between (Mamedov et al. 2000) employing flash-induced fluorescence measurements and the current study using novel DAS.

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Mamedov et al., 2000</th>
<th>This study</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$F_v/F_o$</td>
<td>$F_v/F_o$</td>
</tr>
<tr>
<td>Thylakoid</td>
<td>0.7</td>
<td>2.94</td>
</tr>
<tr>
<td>BS</td>
<td>1.1</td>
<td>4.72</td>
</tr>
<tr>
<td>B3</td>
<td>0.96</td>
<td>4.32</td>
</tr>
<tr>
<td>Ma</td>
<td>0.48</td>
<td>3.18</td>
</tr>
<tr>
<td>T3</td>
<td>0.27</td>
<td>2.25</td>
</tr>
<tr>
<td>Y100</td>
<td>0.2</td>
<td>10.2</td>
</tr>
</tbody>
</table>
5. Discussion

The non-detergent method of thylakoid membrane fractionation employed in this study is clearly a superior preparation tool relative to harsh detergent methods of isolation. Oxygen evolution in the BBY particles (Berthold et al. 1981) is high, but, since detergent concentrations as low as 0.01-0.02 % are known to interrupt the energy transfer among LCHII and the reaction centre of PSII, they make a very unreliable candidate for studies on exciton migration in PSII-LHCII complexes (Yu and Albertsson 1993).

The collective literature describing PSII heterogeneity since Melis and Homann, 1975, has rendered a concrete and broad data set. Consistency with the wealth of research that has been undertaken since the first notion of PSII heterogeneity is crucial to the current study. First, many trends that illustrate the widespread idea of spatial separation (Albertsson et al. 1992; Albertsson et al. 1994; Anderson and Melis 1983) become apparent upon examination of the Gaussian decomposition of the absorption spectra illustrated in figures 27 - 32. The absorption band assigned to Chl b at ~651 nm is most dominant in the grana fraction and more so in the grana core fraction. Longer wavelength bands, typical of PSI absorptions add more weight to the spectra in stroma derived fractions. The Chl a/b ratios have been calculated based on these Gaussian bands, and are listed in table 10. A firm trend exists in these data that suggest that grana and grana core are enriched in PSII relative to the stroma derived fractions. This trend is also apparent in the 77K fluorescence emission spectra in section 4.4. A high F695/F730 is indicative of a relatively large PSII population, and subsequently smaller PSI population in a heterogeneous sample of thylakoid membranes. The spatial distribution of PSII and PSI centres (Albertsson et al. 1990b; Albertsson et al. 1992; Anderson and Melis 1983) is well illustrated by the Gaussian decomposition of the absorption spectra and the 77K fluorescence emission spectra in this study since these data were obtained from domain specific regions of the thylakoid membrane.

These data are consistent with the proposed distribution in this way. The grana core, suggested to be PSII-rich (Svensson and Albertsson 1989; Yu and Albertsson 1993), maintains the highest F695/F730 of all thylakoid membrane vesicles. The grana fraction, consisting of whole grana
membrane vesicles, maintains an $F_{695}/F_{730}$ between that of the grana core and the grana margins. This is not surprising since the grana margins, unlike the grana core are suggested to be a PSI-rich region of thylakoid membrane (Anderson et al. 1999; Webber et al. 1987; Webber et al. 1988; Wollenberger et al. 1994; Wollenberger et al. 1995). The PSI-rich (and PSII-poor) characteristics are also true of the stroma lamellae and purified stroma lamellae fractions. These data are consistent with the Chl $a/b$ ratios that were calculated from the Gaussian deconvolution of the absorption spectra in section 4.1, table 10.

One further line of evidence that is consistent with these notions of spatial separation of the photosystems lies in the deconvolution of the mixed component from the DAS. Tables 16 and 17 list the novel PSII/PSI ratios from within the mixed component at $F_M$ and $F_0$ respectively. From these DAS, the mixed component was determined by a visual test and was apparently $T_2$ in grana derived fractions and whole thylakoid and $T_3$ in stroma derived fractions. In any case, the lifetime of the mixed component was between 120 ps and 260 ps at $F_M$, and between 100 ps and 140 ps at $F_0$. $T_2$ in stroma derived fractions was clearly not mixed since its lifetime was too short (below 100 ps at $F_M$ and $F_0$) and its amplitude was most dominant in the low energy region of the DAS, a typical characteristic of PSI fluorescence decays. The highest novel PSII/PSI, not surprisingly was measured from the deconvolution of $T_2$ of the grana core. The relative yields of the novel components are $20.1\%$ and $4.4\%$ from the overall sum of all of the exponentials. The novel PSII/PSI ratio decreases steadily as the measured parameter moves away from the grana core, towards the stroma lamellae regions of the thylakoid, where this measurement plummets to 0.43, more than ten times less than grana core. Furthermore, the grana margin region of the thylakoid membrane renders a novel PSII/PSI below one.

Secondly, the notion of spatial separation of PSII$\alpha$ and PSII$\beta$ centres needs to be addressed with respect to these novel thylakoid membrane fractions. The first piece of evidence that suggests that this innovative distribution may exist appears in the 77K fluorescence emission spectra. Previously the measurement of the $F_{695}/F_{730}$ parameter proved useful upon description of the spatial separation of PSII and PSI. A similar measurement may be employed to gain insight
into the relative amounts of PSIIα and PSIIβ in domain specific regions of the thylakoid membrane. The relative association of the LHCII complex with PSII increases as the $F_{685}/F_{695}$ ratio at 77K increases. This degree of association between the accessory light-harvesting antenna and the core complex is highest in grana core (2.20) and decreases as one moves towards the stroma lamellae (0.65), as described in table 13. This parameter was not measurable in the purified stroma lamellae fraction since these peaks were present at the same level as noise in the spectrum. Visually, one notices that this region of the spectrum, figure 38, is far below measurable values relative to the other fractions. This may give an indication of its shear lack of LHCII (and PSI-richness) in the membrane vesicles. Using the model of the pigment protein distribution of the PSII complex figure 9, the $F_{685}/F_{695}$ measurement may be used to estimate the number of chlorophylls maintained by PSII centres in these thylakoid membrane vesicles. A number of assumptions are required for this analysis. First, one must assume that equal coupling among the chromophores of each thylakoid membrane fraction exists, as well as equal rates of primary charge separation within the reaction centre. Assuming that PSII in grana core, reported to employ the largest antenna (Mamedov et al. 2000; Svensson and Albertsson 1989; Yu and Albertsson 1993), harbours the full 244 chlorophylls as stated in this model, it may be possible to estimate what portions of the whole complex may be retained by other PSII centre species in these isolated fractions. This analysis suggests that the smallest $F_{685}/F_{695}$, which is indicative of a poor association between PSII and the accessory LHCII, exists in the stroma lamellae fraction. A value of only 0.65 translates to 180 chlorophylls. The most parsimonious resolution states that only two or three, of four, LHCIIb pigment-protein complexes may be associated with PSII centres in this region of the thylakoid membrane.

Of course the LHCIIb subunit may not be the only possible target for dissociation. LHCIIa, -c, -d and -e could be likely candidates as well. A large number of combinations of associations among these proteins could suit this analysis. Table 19 illustrates this analysis. Since the values of $F_{685}/F_{695}$ listed in table 13 are all within approximately 20% of each other, regardless of the trend that they follow, these results may remain difficult to quantify in the current study. A nonlinear
relationship between the $F_{685}/F_{695}$ measurement of LHCII association, and the excited state pigment dynamics may also be a factor in the poor quantification of these data. Nevertheless, consistent with the early research on spatial separation of two PSII species based on antenna size (Albertsson et al. 1990a; Albertsson et al. 1992; Anderson and Melis 1983; Thielen and Van Gorkom 1981a), among many other more recent papers, these results agree with a domain based separation of PSIIα and PSIIβ.
Table 19 - Hypothetical association of LHCIIb with each thylakoid membrane fraction based on the model for the pigment-protein distribution within the PSII complex in figure 9 (Thornber et al. 1991).

<table>
<thead>
<tr>
<th>Fraction</th>
<th>Chlorophylls*</th>
<th>LHCIIb subunits†</th>
</tr>
</thead>
<tbody>
<tr>
<td>Grana core (BS)</td>
<td>244</td>
<td>4</td>
</tr>
<tr>
<td>Grana (B3)</td>
<td>224</td>
<td>3-4</td>
</tr>
<tr>
<td>Grana Margins</td>
<td>202</td>
<td>3</td>
</tr>
<tr>
<td>Stroma lamellae (T3)</td>
<td>180</td>
<td>2-3</td>
</tr>
</tbody>
</table>

* calculated from the F685/F695 considering the grana core as the entire complex
† assuming that the loss of each subunit corresponds to 42 chlorophylls, consistent with the model suggesting a total of 4 subunits harbouring 168 chlorophylls.
The relative sizes of the light-harvesting antennae associated with PSII in different regions of the thylakoid membrane in relation to the spatial separation of PSIIα and PSIIβ is further explored upon examination of the PSII absorbance cross sections (figures 33 - 34). Larger chlorophyll antennae result in a large fluorescence yield at lower light intensities. The chlorophyll antenna associated with PSII centres in the grana core fraction render larger cross-sections relative to others, consistent with the idea that PSIIα centres with larger antennae dominate this fraction. The smallest PSII antenna, are found in purified stroma lamellae (the Y100 fraction), consistent with the proposed location PSIIβ centres harbouring small antennae. These findings are in agreement with Anderson and Melis, 1983, where differential antenna size among domain specific PSII centres was first described, and also agree with the findings based on the 77K fluorescence emission spectra analyses. These findings are illustrated by the δ values listed in table 11. The general trend of decreasing δ is followed by these values. However the degree to which this value decreases is much more than those suggested in the literature and what is reported in table 19. These data suggest that the PSII absorbance cross section measured from purified stroma lamellae is merely 20% of that measured from the grana core. This difference in antenna size is much larger than the popular 50% difference reported across the field. Fluorescence induction kinetics collected from grana partition membrane vesicles and stroma lamellae membrane vesicles, fit with Strasser's model (Strasser and Stirbet 1998; Strasser and Stirbet 2001), and agree with the wealth of spatial distribution data beginning as early as Jahns and Schweig 1995. Since the intensity and duration of the excitation flash remained constant from sample to sample, the differential rate constants undoubtedly due to the different antenna sizes (Thielen and Van Gorkom 1981a), are illustrated clearly as well in the PSII absorbance cross sections for the grana vesicles and stroma lamellae vesicles (figure 33).

It is reasonable now to suggest that the spatial separation of PSII and PSI between the stacked grana partition and the unstacked stroma lamellae (Albertsson et al. 1990b; Albertsson et al. 1992; Anderson and Melis 1983) and between the appressed grana core and the stroma exposed grana margins (Anderson et al. 1999; Webber et al. 1987; Webber et al. 1988;
Wollenberger et al. 1994; Wollenberger et al. 1995) is clearly demonstrated by the domain specific, nondetergent isolated thylakoid membrane fractions prepared by F. Mamedov and R. Danielsson under the direction of S. Styring of the Photosynthesis Group at Lunds Universitet, Lund, Sweden. It is also reasonable to accept the notion of PSIIα/β separation based on these data. For the purpose of analogous comparison the BS grana core fraction may be referred to as the α-fraction. Furthermore, the Y100 purified stroma lamellae may be referred to as the β-fraction.

In the last decade global target analysis of picosecond chlorophyll kinetics including an analysis of PSII heterogeneity has remained virtually untouched. Roelofs et al., 1992 was the first study to examine the characteristics of the primary processes in PSIIα and PSIIβ centres to any appreciable degree. The results of their paper suggested the following. 1) PSIIα and PSIIβ centres exhibit a different molecular functioning. 2) Differential molecular structure of the reaction centres and/or relative local environment may give rise to these differences. 3) Spatial separation may have a role as well (Roelofs et al. 1992).

Data from Roelofs et al., 1992, is summarized in table 20 and has been corrected to allow a valid comparison to the DAS rendered from this study. This was accomplished by considering only the plotted components of the DAS, leaving out the 10 ps decay. The uniqueness of the non-detergent thylakoid membrane preparations employed in this study, relative to Holzwarth's whole thylakoid analysis (Roelofs et al. 1992), is most likely the key determinant responsible for the differences that become apparent upon comparison of these kinetic data. Of course, variation among samples does exist to some degree, however, general trends among the DAS and their associated lifetimes and respective yields are usually maintained. Unlike the present study, Roelofs et al., 1992, were not afforded the luxury of performing kinetic analyses on isolated, but intact, PSIIα and PSIIβ centres. DAS from Roelofs et al., 1992 were scanned and redigitized in figures 52, Fw and figure 53, Fo, to aid in the comparison. For the purpose of this comparison table 21 describes the assignment of the novel DAS components using the same nomenclature as Roelofs et al., 1992. In this case, DAS obtained from the grana core and purified stroma lamellae fractions represented the PSIIα population PSIIβ population respectively. This assignment is in
agreement with the overwhelming wealth of literature that suggests the existence of this spatial separation (Albertsson et al. 1990b; Albertsson et al. 1992; Anderson and Melis 1983).
Figure 52 - DAS at $F_M$ for whole thylakoid membranes from Roelofs et al., 1992. The PSI component is plotted in black. Supposed PSII$\alpha$ components are plotted in red. Supposed PSII$\beta$ components are plotted in green. The 11 ps exciton equilibrium component has been omitted, as in the original document (Roelofs et al. 1992).
Figure 53 - DAS at F₀ for whole thylakoid membranes from Roelofs et al., 1992. The PSI component is plotted in black. Supposed PSIIα components are plotted in red. Supposed PSIIβ components are plotted in green. The PSII closed component is plotted in blue. The 11 ps exciton equilibrium component has been omitted, as in the original document (Roelofs et al. 1992).
Table 20 - Results from the global target analyses of the fluorescence kinetics at $F_0$ and $F_M$ from (Roelofs et al. 1992) employing whole thylakoid membranes. The reduced $\chi^2$-value for the combined ($F_M$ and $F_0$) global target analysis is 1.0662.

<table>
<thead>
<tr>
<th>Case</th>
<th>Lifetimes (yield, %)*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PSI (ns)</td>
</tr>
<tr>
<td>$F_0$</td>
<td>0.104 (30.5)</td>
</tr>
<tr>
<td>$F_M$</td>
<td>0.104 (30.3)</td>
</tr>
</tbody>
</table>

* The lifetime component reflecting the exciton equilibrium process (34%-37%) appearing in the original paper is not reported to allow comparison to the present study. The yields reported here have been recalculated to allow this subtraction.
Table 21 - Comparison of the present study to Roelofs, et al., 1992, at F_M and F_0 using assignments from Roelofs et al., 1992. Fluorescence lifetimes, $T_m$ (ns) are obtained from the target global fitting procedure. The associated (%) yields are in parentheses. Fluorescent yields of deconvoluted components in the current study are assigned to either PSI or PSII.

<table>
<thead>
<tr>
<th></th>
<th>T_2, PSI/PSII_{fast}</th>
<th>T_3, PSI_{slow}/PSII_1</th>
<th>T_4, PSI_2</th>
<th>T_5, PSI_3</th>
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<tr>
<td><strong>F_M</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PSI\alpha</td>
<td>0.654 (9.5)</td>
<td>1.785 (35.8)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>N.D.</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PSI\beta</td>
<td>0.375 (6.5)</td>
<td>2.921 (18)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>F_0</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PSI\alpha</td>
<td>0.253 (29.6)</td>
<td>0.519 (14.2)</td>
<td></td>
<td>1.855 (1.3)</td>
</tr>
<tr>
<td>PSI\beta</td>
<td>0.322 (14.2)</td>
<td>0.686 (10.2)</td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>Present study</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td><strong>F_M</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PSI\alpha†</td>
<td>0.16 (4.4, 20.1)</td>
<td>0.65 (26.6)</td>
<td>1.74 (44)</td>
<td>3.76 (4.9)</td>
</tr>
<tr>
<td>PSI\beta‡</td>
<td>0.09 (80.8)</td>
<td>0.26 (8.2, 3.5)</td>
<td>0.88 (5.4)</td>
<td>2.28 (2.1)</td>
</tr>
<tr>
<td><strong>F_0</strong></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>PSI\alpha†</td>
<td>0.11 (2.4, 57.5)</td>
<td>0.3 (38.2)</td>
<td>0.67 (1.9)</td>
<td>7.98 (&lt;1)</td>
</tr>
<tr>
<td>PSI\beta‡</td>
<td>0.04 (71.4)</td>
<td>0.14 (18.3, 6.5)</td>
<td>0.43 (3.3)</td>
<td>2.38 (&lt;1)</td>
</tr>
</tbody>
</table>

† Other data in the current study suggest that the BS fraction is essentially a PSI\alpha preparation.
‡ Other data in the current study suggest that the Y100 fraction is essentially a PSI\beta preparation.
The results from global analysis reported by Roelofs et al., 1992 in table 21 suggest alternative fluorescence decay lifetimes relative to those reported in the present study. Unlike the data presented in this paper at $F_0$, a low amplitude, long-lived fluorescence lifetime, usually about 2 ns or greater, is not reported for measurements taken at $F_M$. However, they do report a similar fluorescence decay component as the longer-lived of two PSII components associated with both $\alpha$- and $\beta$-centres at this level of chlorophyll fluorescence.

The current study describes fluorescent lifetimes for PSII$\beta$ centres that are shorter than those of PSII$\alpha$ centres at $F_M$ and $F_0$. This may be because of the structure and functionality of these PSII centres. Large, shared antenna PSII$\alpha$ centres which are electron transport competent (capable of $Q_b$ reduction) offer alternative fates of the excited state relative to the small, isolated antenna PSII$\beta$ centres which cannot. Of the small absorption that the PSII$\beta$ centres are capable of, more often than not, the result is probably some sort of quenching mechanism coupled with a fast release of low energy as heat, rather than moving towards charge stabilization. This explanation is supported by the PSII repair cycle (figures 12 and 13) where PSII$\beta$ centres in stroma exposed regions of thylakoid membranes harbour a damaged, semi-repaired nonfunctional or no D1 reaction centre protein at all. In this state, photochemistry and the further reduction of PQ is not an option (Guenther and Melis 1990; Melis 1991). It may be advantageous for plants to protect damaged reaction centres during repair via quenching in order to reduce the production of dangerously reactive oxygen species that lead to proteolysis.

Holzwarth's group describe longer fluorescence lifetimes of PSII$\alpha$ relative to those of PSII$\beta$ in only one case. Table 21 lists the fluorescence lifetimes for Roelofs et al., 1992, at $F_M$ and $F_0$. Only at $F_M$, where the fastest of two PSII components is that of the $\beta$-centre, does this study agree with the current one. It is difficult to access the reasons for these differences among the relative lifetimes of PSII$\alpha$ and PSII$\beta$ since not all components have been resolved in Roelofs et al., 1992.

The relative fluorescence yield of the two PSII components derived from the DAS at $F_M$ of each fraction reports an interesting phenomenon. In cases where PSII$\alpha$ is enriched, the faster decay component offers a lower yield relative to the slower decay component. This occurs in the
grana ($T_3/T_4 = 0.78$) and more-so in grana core ($T_3/T_4 = 0.60$). In fractions derived from stroma-exposed regions of thylakoid, the grana margins, the stroma lamellae fraction and the purified stroma fraction the $T_3/T_4$ ratios are 1.16, 1.9, 2.17 respectively.
6. Conclusions

Heterogeneity of PSII, related to its spatial distribution within the thylakoid membranes of higher plants is not a new concept (Albertsson et al. 1990b; Albertsson et al. 1992; Anderson and Melis 1983). However its understanding and significance have remained elusive over two decades of research. The present study aimed to offer insight on the biophysical properties of PSIIα and PSIIβ.

This study sought to address the notion of spatial separation within thylakoid membranes. The domain specific, nondetergent, thylakoid membrane fractions prepared by F. Mamedov and R. Danielsson under the direction of S. Styring from the Photosynthesis Group at Lunds Universitet, Lund, Sweden, offer an ideal candidate for this study. The first element of spatial heterogeneity is a product of the PSII/PSI distribution. This distribution was demonstrated clearly by the Gaussian decomposition of the absorption spectra. The red-shifted PSI absorption bands (Cramer and Butler 1968) were more dominant in the stroma exposed fractions of the thylakoid membrane. In addition, higher Chl a/b ratios are obtained from grana and more so from grana core. Since PSII is relatively rich in Chl b, compared to PSI, one can assume that these fractions agree with this trend. More evidence supporting the spatial separation of PSII and PSI was illustrated by the 77K fluorescence emission spectra. F_{690}/F_{730} was lowest in stroma exposed membrane fractions, known to be PSI-enriched. These include the stroma lamellae fraction (T3), purified stroma lamellae fraction (Y100), and the grana margin fraction.

The second element of spatial heterogeneity lies in the domain specific distribution of PSIIα and PSIIβ. The data presented here clearly demonstrate this distribution. First, further examination of the 77K fluorescence emission spectra via evaluation of the F_{685}/F_{695}, which reveals the relative association of LCHII, shows that this parameter is highest in grana (B3) and grana core (BS). These data coupled with the PSII absorption cross section agree with the notion of a separation of large PSIIα in grana and small PSIIβ in stroma (Anderson and Melis 1983; Thielen and Van Gorkom 1981a) and the PSII repair cycle where damaged PSII centres are stripped of the LHCII prior to migration towards the unstacked regions of the membrane (Aro et al. 1993; Guenther
and Melis 1990; Melis 1991). This study probed the differential antenna sizes of PSIIα and PSIIβ one other way. Proper analysis of fluorescence induction kinetics offers information about both antenna and reducing side heterogeneity of PSII centres; however, these enigmatic data are often difficult to describe (Holzwarth 1993). Arguably, PSII centres employing larger chlorophyll antennae may close their reaction centres (fully reduce Q_A) at greater rates than those employing smaller chlorophyll antennae under the same experimental conditions. Strasser’s models for the theoretical yield of F_v based on PSII connectivity are useful in some respects upon measurement of these rates (Strasser and Stirbet 1998; Strasser and Stirbet 2001). PSIIα-rich grana vesicles and PSIIβ-rich stroma vesicles were valuable to this question. It was found that indeed, the rate of PSII closure was greater in the B3 fraction, relative to the T3 fraction. Another interesting finding was noted when the connectivity constants (equation 31) for domain specific PSIIα and PSIIβ were left free rather than fixing them at Strasser’s theoretical values of 1.67 and 0 respectively for whole thylakoid membrane. For PSIIβ centres this value remained nil whereas, in PSIIα-rich grana membrane vesicles it remained relatively close to the model at 2.08. This may be interpreted as an indication of the vast exciton sharing that is reported upon description of the lake (or matrix) model of PSII units (Beauregard and Trissl 1999). It seems that in a relatively homogenous population the measurement of large shared PSII antennae is not clouded by the small isolated antenna of PSIIβ producing the typical biphasic kinetics see in many early papers on PSII heterogeneity (Black et al. 1986; Melis and Duysens 1979; Melis and Homann 1975; Melis and Homann 1976).

Spatial separation between photosystems I and II, and also between the two species of PSII, α- and β-centres have been established with these novel thylakoid membrane vesicles. Accordingly, an analysis of the relative picosecond fluorescence decay kinetics with respect to domain specific PSII heterogeneity was possible with the novel membrane fractions. Work on PSII heterogeneity with respect to TC-SPC on chlorophyll fluorescence decay kinetics on the submicrosecond time scale has remained dormant since Roelofs et al., 1992. The DAS from Holzwarth’s paper were recreated in figure 52 at F_M and figure 53 at F_0 (Roelofs et al. 1992).
Unlike the current study, in only one case does the decay of the PSIIα component have a longer lifetime than the corresponding PSIIβ component reported by Roelofs et al., 1992. At Fₘ, Roelofs et al., 1992, describe fluorescence lifetimes of 0.654 ns and 0.375 ns for PSIIα and PSIIβ respectively. The current study describes a data set where all fluorescence lifetimes of PSIIα centres are longer lived than the respective PSIIβ centres. This fast chlorophyll fluorescence decay of PSIIβ centres may be a constituent of a rapid quenching mechanism that protects damaged PSII centres during their repair in unstacked stroma lamellae. Since long fluorescence lifetimes of \( ^3 \text{Chl}^* \) enhance the generation of dangerously reactive \( ^1 \text{O}_2 \) when photochemistry is not an option, its rapid qN is the most advantageous way for a plant to recover the damaged PSII (Aro et al. 1993; Guenther and Melis 1990; Melis 1991).
7. Literature Cited


Beauregard K, Trissl H-W. 1999. Theories for kinetic and yields of fluorescence and photochemistry: How, if at all, can different models of antenna organization be distinguished experimentally? Biochimica et Biophysica Acta. 1409, 125-142.


Lyon MK .1998. Multiple crystal types reveal PSII to be a dimer. Biochimica et Biophysica Acta. 1364, 403-419.


Staehelin LA (1986) Chloroplast structure and supramolecular organization of photosynthetic


Raton, FL)


8. Appendices
Nondetergent Fractionation of Thylakoid Membranes

Reagents

Preparation Medium
- 50 mM sodium phosphate buffer (pH 7.4)
- 5 mM MgCl₂
- 300 mM sucrose

Washing Medium I
- 10 mM tricine (pH 7.4)
- 5 mM MgCl₂
- 300 mM sucrose

Washing Medium II
- 10 mM sodium phosphate buffer (pH 7.4)
- 5 mM NaCl
- 1 mM MgCl₂
- 100 mM sucrose

Polymer Mixture
- 3.32 g of 20% (w/w) dextran T500
- 1.66 g of 40% (w/w) PEG 3350 (4000)
- 0.53 g of 0.2 M sodium phosphate buffer (pH 7.4)
- 0.30 g of 0.1 M NaCl
- 1.07 g of 10 mM MgCl₂
- 1.33 g of 0.1 M sucrose
- 2.45 g ddH₂O

Fractionation I
- 2 g of thylakoid suspension to 9.66 g of Polymer Mixture to give the following final concentrations:
  - 5.7% dextran T500
  - 5.7% PEG 3350 (4000)
  - 10 mM sodium phosphate buffer (pH 7.4)
  - 20 mM sucrose
  - 3 mM NaCl
  - 1 mM MgCl₂
- Sonicate with ½-inch horn in six 30s bursts with 60s between bursts in a cylindrical aluminum tube immersed in ice and water.
  - The ultrasonic intensity output setting is 7, with 20% duty pulses
- 6.43 g of pure lower phase and 5 g of pure top phase are added to:
  - 5.7% dextran T500
  - 5.7% PEG 3350 (4000)
  - 10 mM sodium phosphate buffer (pH 7.4)
  - 20 mM sucrose
  - 5 mM NaCl
- mixed at 4 °C
- centrifuged at 2000 g for 3 min to achieve separation
- upper and lower phases are separated and washed twice in 10 ml of fresh upper and lower phase respectively
- the final upper phase following wash with fresh lower phase constitutes the T3 fraction
- the final lower phase following wash with fresh upper phase constitutes the B3 fraction
Fractionation II
- grana cores and margins were separated from within the B3 fraction
  - Sonicate with ½-inch horn in twelve 30s bursts with 60s between bursts in a cylindrical aluminum tube immersed in ice and water.
  - The ultrasonic intensity output setting is 7,
  - with 20% duty pulses
  - upper and lower phases are diluted three times in:
    - 10 mM sodium phosphate buffer (pH 7.4)
    - 5 mM NaCl
    - 100 mM sucrose
  - centrifuged at 100000 g for 90 min.

Fractionation III
- Y100 in a buffer similar to Washing Medium II (except 5 mM MgCl₂) are disintegrated in a Yeda press at 10 MPa nitrogen pressure
- homogenate is diluted 5x in MgCl₂-free buffer and centrifuged at 40000x g for 30 min.
- membrane vesicles in supernatant are sedimented at 100000x g for 45 min.

Resuspension Buffer
- 0.33 M Sorbitol
- 10 mM HEPES (pH 7.8)
- 15 mM MgCl₂
- 10 mM NaCl
- 3 mM Ascorbic Acid
- 0.5 g/L B.S.A. (only when noted)
<table>
<thead>
<tr>
<th>Fraction</th>
<th>Par. 1</th>
<th>Par. 2</th>
<th>Par. 3</th>
<th>Par. 4</th>
<th>Par. 5</th>
<th>Par. 6</th>
<th>Par. 7</th>
<th>Par. 8</th>
<th>Par. 9</th>
<th>Par. 10</th>
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<tbody>
<tr>
<td>BS xc</td>
<td>620</td>
<td>633.74797</td>
<td>640.69468</td>
<td>649.74823</td>
<td>664.08294</td>
<td>670.69474</td>
<td>676.4956</td>
<td>681.77687</td>
<td>695</td>
<td>710</td>
</tr>
<tr>
<td>A</td>
<td>1.06719 ±0.9616</td>
<td>1.47673 ±0.46119</td>
<td>2.24253 ±0.22396</td>
<td>4.50411 ±0.15683</td>
<td>±0.258 ±0.29244</td>
<td>±0.31217 ±0.06423</td>
<td>±0.02122</td>
<td></td>
<td></td>
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</tr>
<tr>
<td>B3 xc</td>
<td>620</td>
<td>633.74797</td>
<td>640.69468</td>
<td>649.74823</td>
<td>664.08294</td>
<td>670.69474</td>
<td>676.56555</td>
<td>681.77687</td>
<td>695</td>
<td>710</td>
</tr>
<tr>
<td>A</td>
<td>0.92585 ±0.77134</td>
<td>1.24887 ±0.33232</td>
<td>2.09841 ±0.56275</td>
<td>3.76746 ±0.48172</td>
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<td>±0.02249</td>
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<tr>
<td>Ma xc</td>
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<td>633.74797</td>
<td>640.69468</td>
<td>649.74823</td>
<td>664.08294</td>
<td>670.69474</td>
<td>676.68729</td>
<td>681.77687</td>
<td>695</td>
<td>710</td>
</tr>
<tr>
<td>A</td>
<td>1.44194 ±0.80348</td>
<td>1.9599 ±0.20918</td>
<td>1.8578 ±0.33599</td>
<td>3.39397 ±0.39109</td>
<td>±0.34936 ±0.73399</td>
<td>±0.29764 ±0.10855</td>
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<td></td>
<td></td>
</tr>
<tr>
<td>T3 xc</td>
<td>620</td>
<td>633.74797</td>
<td>640.69468</td>
<td>649.74823</td>
<td>664.08294</td>
<td>670.69474</td>
<td>676.96404</td>
<td>681.77687</td>
<td>695</td>
<td>710</td>
</tr>
<tr>
<td>A</td>
<td>1.26882 ±0.64429</td>
<td>1.11457 ±0.27426</td>
<td>1.71672 ±0.46297</td>
<td>4.14717 ±0.35224</td>
<td>±0.24028 ±0.36141</td>
<td>±0.71532 ±0.54864</td>
<td>±0.24881</td>
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</tr>
<tr>
<td>Y100 xc</td>
<td>620</td>
<td>633.74797</td>
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<td>664.08294</td>
<td>670.69474</td>
<td>677.50234</td>
<td>681.77687</td>
<td>695</td>
<td>710</td>
</tr>
<tr>
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<td>3.90308 ±0.45729</td>
<td>±0.1404 ±0.45729</td>
<td>±0.61402 ±0.36141</td>
<td>±0.12321 ±0.79836</td>
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</tr>
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</table>

Table 22: Parameters (Par.) of the global Gaussian decomposition of the low-temperature absorption spectra. Peak centers (xc) half-widths (w) and amplitudes are fixed with the exception of xc.
Figure - 54 - Molar extinction coefficients of Chls a and b (Du et al. 1998).
Figure 55- Residuals generated from global analysis of decay kinetics of thylakoid membrane fractions at $F_M$. $\chi^2$ values for each plot are A) whole thylakoid - 1.216, B) B3 fraction - 1.10, C) T3 fraction - 1.25, D) BS fraction - 1.162, E) Ma fraction - 1.135 and F) Y100 - 1.191. Each residual represents the distribution about the median for each of nine emission wavelengths collected from 675 nm (top) to 730 nm (bottom).
Figure 56- Residuals generated from global analysis of decay kinetics of thylakoid membrane fractions at F₀. χ² values for each plot are A) whole thylakoid - 1.18, B) B3 fraction - 1.293, C) T3 fraction - 1.259, D) BS fraction - 1.446, E) Ma fraction - 1.281 and F) Y100 - 1.316. Each residual represents the distribution about the median for each of nine emission wavelengths collected from 675 nm (top) to 730 nm (bottom).