Structure/Function/Dynamics of Photosystem II Plastoquinone Binding Sites

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Abstract: Photosystem II (PSII) continuously attracts the attention of researchers aiming to unravel the riddle of its functioning and efficiency fundamental for all life on Earth. Besides, an increasing number of biotechnological applications have been envisaged exploiting and mimicking the unique properties of this macromolecular pigment-protein complex. The PSII organization and working principles have inspired the design of electrochemical water splitting schemes and charge separating triads in energy storage systems as well as biochips and sensors for environmental, agricultural and industrial screening of toxic compounds. An intriguing opportunity is the development of sensor devices, exploiting native or manipulated PSII complexes or ad hoc synthesized polypeptides mimicking the PSII reaction centre proteins as biosensing elements. This review offers a concise overview of the recent improvements in the understanding of structure and function of PSII donor side, with focus on the interactions of the plastoquinone cofactors with the surrounding environment and operational features. Furthermore, studies focused on photosynthetic proteins structure/function/dynamics and computational analyses aimed at rational design of high-quality bio-recognition elements in biosensor devices are discussed.

Keywords: Molecular docking, molecular dynamics simulations, Photosystem II, plastoquinone, plastoquinone binding site, protein dynamics.

1. STRUCTURAL OVERVIEW OF PSII CORE COMPLEX

Photosynthetic organisms have the unique ability of entrapping and using solar energy to oxidize water to molecular oxygen through the highly structured multi-subunit protein-chlorophyll complexes of photosystem I (PSI) and photosystem II (PSII), arranged in supra-molecular assemblies embedded in thylakoid membranes. These structured assemblies are organized in a way to ensure supreme energy conversion efficiency and effective diffusion of small molecules (plastoquinone) interconnecting the two photosystems functions in a fine-tuned electron transport chain and leading to the synthesis of ATP and reducing equivalents. PSII stands out for its remarkable ability to act as a water:plastoquinone oxidoreductase, extracting electrons from water in physiological conditions. The outstanding features of PSII have motivated numerous research efforts aimed to clarify its structural and functional relationships, which until recently have been hindered by the lack of a high resolution X-ray crystal structure. Our current understanding of the PSII structure/function/dynamics rationale has been largely acquired by comparative studies and assumptions with the simpler and better-known purple bacterial reaction centre (bRC) [for review on bRC see 1, 2]. In the last decade, the crystal structure resolution at 2.9-3.5 Å has exponentially improved the knowledge of atomic details of dimeric and monomeric forms of PSII core complex (PSIIcc) of the thermophilic cyanobacterium Thermosynechococcus elongatus [3-7]. More recently, Umena and coworkers provided the highly resolved 1.9 Å structure from Thermosynechococcus vulcanus [8]. This structure features a rich collection of atomic details among which a cofactors arrangement made up by 35 chlorophylls, 2 phaeophytins, 11 β-carotenes, 2 plastoquinones, 2 heme irons, 1 non-heme iron, 4 Mn atoms, 3-4 Ca atoms (among which, 1 is in the Mn,Ca-cluster), 3 Cl ions (among which, 2 are in the vicinity of the Mn,Ca-cluster), and 1 (bi)carbonate ion. From the electron density map it was possible for the first time to locate all the atoms building up the Mn,CaO-cluster and to define the precise position of all the metal ligands and of the substrate water molecules. In this high-resolution structure the primary plastoquinones (PQs) QA and QB occupy positions very similar to those observed in previous crystallographic studies. However, the third plastoquinone molecule observed in the structure solved by Guskov and coworkers [5], the so called QC, has not been observed in this last structure casting some doubts on the real physiological function of QC. These structural data together with the numerous biochemical and spectroscopic methods, and computational analyses contribute to solve the enigma of PSII structure/function/dynamics specific correlation.
PSIIcc consists about 20 different polypeptide chains, with the majority in the form of trans-membrane and α-helical subunits. Among these, PsbA (D1) and PsbD (D2) subunits, with five trans-membrane helices (TMH) each, form the architecture of PSII RC and host the major redox-active cofactors organized in two quasi symmetrical branches: two main chlorophyll a (Chla), PD1 and PD2, two accessory Chla, ChlD1 and ChlD2, two peripheric Chla, ChlEZ1 and ChlEZ2, two Pheophytin a, PheoD1 and PheoD2, and two PQs, QA and QB (Fig. 1). On the stromal (cytosol in prokaryotes) side of the complex between the primary and secondary PQs, a non-heme iron interacting with a bicarbonate anion has been also revealed. It is electrostatically associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to the two PQs, optimizing charge distribution and H-bonding of the QA-Fe-QB bridge and concomitantly associated to

Fig. (1). Schematic representation of PSIIcc main protein components and electron transfer cofactors. The reaction centre (D1/D2) heterodimer is reported in light (D2) and dark (D1) yellow, respectively; core antenna proteins CP47 and CP43 are in green; the Oxygen Evolving Complex (OEC) – in blue. PSIIcc associated cofactors: the primary (PD1/PD2) and accessory (ChlD1/ChlD2) Chla molecules are in cyan, pheophytin a (PheoD1/PheoD2) are in gold brown, plastoquinones (QA, QB and QC) are in yellow, the heme of cytochrome b\textsubscript{559} (Cyt. b\textsubscript{559}) is colored green and the non-heme iron in grey.

2. PSII PLastoquinones

The two chemically identical PQ molecules, hosted into the QA and QB binding sites of the RC heterodimer, have the distinctive role of stabilization of primary charge separation and conduction of electrons towards the photosynthetic electron transport chain, respectively. The existence of a third plastoquinone, QC, in the PSII core complex has been supported by Cyt.b\textsubscript{559} photoreduction and redox titration experiments and lately identified by the 2.9 Å resolution crys-
The role of this quinone molecule in the PSII function is still under debate and will be discussed below. The binding energies for QA, QB and QC obtained by fragment molecular orbital (FMO) calculations based on the 2.9 Å X-ray structure are -56.1 kcal/mol, -37.9 kcal/mol and -30.1 kcal/mol, respectively [23]. The FMO calculations indicated a decrease of the relative contribution of the PQ/protein matrix interactions to the total PQ interaction energy in the order QA>QB>QC, with concomitant increase of the relative contribution of surrounding lipids and cofactors (Table 1). Thus, the FMO results drew a correlation between the increasing of PQ mobility and the growing relative contribution of lipids and cofactors interaction.

### 2.1. Plastoquinone at QA Binding Site

The plastoquinone at QA site tightly interacts with the RC protein matrix, does not undergo protonation, and is not exchangeable. It functions as a single electron recipient and mediates the double reduction of the second quinone molecule at the QB binding pocket. In contrast, the double reduced and protonated QB moves away from its binding site and is promptly replaced by a new PQ molecule (as well as by artificial quinones or herbicides), leading to consider this molecule as a substrate rather than a tightly bound cofactor. Although chemically identical, QA and QB demonstrate distinguishable redox potentials, necessary for the potentialisation of PSII forward electron transfer and the fine tuning of the backward recombination reactions [24, 25]. The different functional and redox-properties of the PSII plastoquinone molecules are largely determined by the interactions with the surrounding environment (H-bonding, hydrophobic interfaces, π-stacking), and the conformation of the individual binding sites and phytyl chains [4-6]. FMO quantum chemical calculations based on the 2.9 Å X-ray structure of *T. elongatus* pointed out that the QA phytyl tail is responsible for more than 60% of the total molecule interactions and QA stabilization is mainly due to interactions with amino acid residues [23], Table 1 and Table 2. The position of the isoprenoid tail is restrained mostly by van der Waals contacts with TMH-d of D2, TMH-a of D1, PsbL and PsbT and the accessory ChlD1 (Fig. 2). The 2.9 Å resolution structure revealed an additional stabilization of the QA molecule due to the interaction of the terminal unit of the isoprenoid tail with π-electron clouds of the phenyl rings of D1-Phe52 and PsbT-Phe10 [5]. QA head is fixed by two H-bonds, formed by the keto-oxygen atoms with the D2-Phe261 backbone amide group and the side chain N of D2-His214, and by π-stacking with D2-Trp253 indole ring, occurring in an offset-stacked manner [4, 5]. The static picture of QA interactions within its binding niche obtained by X-ray analyses has been further complemented by spectroscopic studies. Simultaneous 1H and 14N Two-Dimensional Hyperfine Sublevel Correlation spectroscopy shed new light on the strength and orientation asymmetry of H-bonding of QA anion radical, establishing that when QA is in its reduced form, the H-bond involving D2-Phe261 amide group is significantly stronger than the one attributed to D2-His214 [26, 27]. Similarly, computational studies provided stronger interaction energies of QA with D2-Phe261, D2-Thr217 and D2-Trp253 than with QA-D2-His214 [23], (Fig. 2, left bottom panel).

Important properties of the primary PQ molecule, proven in bacteria, cyanobacteria and higher plants, is the existence of the one-electron redox couple QA/QA· in two conformations with highly different redox potential [28]. The well-recognized ability of herbicides binding at QB site to increase or decrease the susceptibility of PSII to photodamage was related to herbicide-induced changes at midpoint potential of the QA [29-31]. Recent studies have strengthened a previ-

| Factors Contributing to PQs Interactions | Interaction Energy Distribution, % |
|----------------------------------------|----------------------------------|
|                                       | PQ at QA Binding Niche | PQ at QB Binding Niche | PQ at QC Binding Niche |
| Protein matrix PQs interactions         | 100.0                | 82.0                | 56.0                |
| PQs head contribution                   | 38.0                 | 44.9                 | 8.8                 |
| PQs tail contribution                   | 62.0                 | 56.1                 | 91.2                |
| Lipids and cofactors PQs interactions   | None                 | 18.0                 | 44.0                |
| PQs head contribution                   | -                     | 0.0a                | 10.0                |
| PQs tail contribution                   | -                     | 100.0               | 90.0                |
| Relative head contribution to total PQs interaction energy | 38.0 | 35.6 | 44.6 |
| Relative tail contribution to total PQs interaction energy | 62.0 | 64.4 | 55.4 |

*The repulsive QB interactions with lipids and cofactors counteract to the attractive one.*
ously found correlation [32, 33] between the $Q_A/Q_{A^{\cdot -}}$ midpoint potential changes and PSII donor side integrity level [28, 34]. The $Q_A$ low potential form is present in functional PSII reaction centres while donor-side $Ca^{2+}$ extraction or Mn cluster disassembly is associated with an increase of (approx. 150 mV) in the $Q_A$ redox potential. Müh and coworkers [35] and Cardona and coworkers [36] have recently addressed in comprehensive reviews the physiological significance of the $Q_A$ redox shift in guiding a “safe” PSII charge recombination reactions and protecting PSII from photodamage during the assembly of the RC. In addition, an up-shift of the $Q_A$ midpoint potential by ca. +38 mV was observed in T. elongatus in photo-oxidative stress conditions, when in the PSII RC the PsbA3 (D1) isofrom preferentially replaces the PsbA1 (D1) protein, expressed in optimal conditions. This finding evidenced the capability of the D1 amino acid composition to influence the $Q_A$ redox potential, most probably affecting the $Q_A$-His214(D2)-Fe-His215(D1)-$Q_B$ interaction chain [37].

The rationale of the $Q_A$ redox potential switching in photo-protection should be viewed in light of the very recent study conducted by Boussac and coworkers [38], demonstrating faster recombination reaction in PsbA3 than in PsbA1 PSII-RC, which could thwart $^{1}O_2$ generation initiated by triplet state of the RC primary chlorophyll molecules.

2.2. Plastoquinone at $Q_B$ Binding Site

The site hosting the secondary quinone, $Q_B$, is situated in a symmetry-related manner to $Q_A$ binding spot around the non-heme iron (Fig. 2, top panel). Because of its substrate-like behaviour it was not always possible to assign the electron density found in the presumed $Q_B$ binding site to a PQ molecule [6]. The problem of the $Q_B$ binding site electron density inhomogeneity was surpassed by crystallization of the PSII core complex paired with a herbicide molecule [7]. However, in the 2.9 Å structure it was possible to model seven out of the nine $Q_B$ isoprenoid units [5, 11]. In contrast to the secondary quinone acceptor in bRC, found in two different positions called “proximal” and “distal” [39], PSII $Q_B$ was located in only one position, corresponding to the “proximal” one in bRC. The $Q_B$ binding site is formed mainly by the D1 RC protein residues (Table 2). The keto-oxygen atoms of the $Q_B$ head group form H-bonds with D1-Ser264 hydroxyl group and the side chain N of D1-His215. In contrast to $Q_A$, $Q_B$ has a possibility to form a third H-bond with the backbone amide group of D1-Phe265 (2.84 Å) [5, 8], (Fig. 2, centre panel). Most probably, the latter is involved in the stabilization of the SQ state of the $Q_B$ molecule during the reduction/protonation process [36]. The $Q_B$ isoprenoid tail, located into the recently described PQ/PQH$_2$ exchange channel II, is inclined in relation to the one of $Q_A$ [5]. Similarly to $Q_A$, the contribution of the phytol tail to the total $Q_B$ interaction energy is calculated to be more than 60%, but differently from the $Q_A$ the surrounding protein matrix contributes for a smaller fraction of the binding energy [23], Table 1. Based on the interaction strength of the $Q_A$ and $Q_B$ with the respective amino acid residues obtained by FMO calculations, Hasegawa and Noguchi [23] have questioned the role of the H-bonds established by the PQ keto-oxygen groups in the $Q_A$ and $Q_B$ binding. The authors observed more substantial contribution of the non-covalent interactions of $Q_A$ head group with D2-Phe261 and D2-Trp253 and $Q_B$ head group with D1-Phe255 and D1-Phe265 to the total PQ binding energies than the above mentioned H-bonding. Instead, it has been unambiguously demonstrated the importance of the H-bonding patterns of PSII PQs in determining their redox potential [40, 41]. The relative contribution of PSII integral lipids and cofactors to the total $Q_B$ interaction energy is less than 20% and is mainly limited to the PQ tail (Table 1 and Table 2). The $Q_B$ phytol chain is accommodated in channel II of the PSII internal cavity, connecting the $Q_B$ binding site with the membrane PQ pool, and is in close interactions with the phytol chain ChlD2 and the acyl chains of MGDG18, and to a lesser extent to SQDG4, DGDG6 and MGDG7.

The reduction/protonation mechanism of the secondary quinone has been a long-term challenge for researchers in photosynthesis. The $Q_B$ reduction occurs in two successive photoreactions including the formation of SQ intermediates. The first reaction is fast, occurring within 0.2-0.4 ms, and generates a SQ tightly bound into the $Q_B$ pocket, which can accept a second electron in a somewhat slower process – 0.6-0.8 ms [42]. Experimental data suggested that $Q_B$ becomes protonated before the second electron transfer occurs, thus avoiding the formation of highly charged and unstable $Q_AQ_B^{2-}$ species [35, 36] and the protonation occurs more promptly if $Q_A$ is in its reduced state [10]. The resulting plastochinol is characterized by low affinity for the $Q_B$ binding niche and is promptly released into the membrane.

In bRC the protonation process has been well established: the first $H^+$ necessary to neutralize the charge on the distal (in relation to the non-heme Fe position) keto-oxygen atom of the SQ is provided by the Ser L223 amino acid trough Asp L213, while the second one is released by the Glu L212 residue. In plant and cyanobacterial PSII RCs the details of this phenomenon are still under investigation. Similarly to bRC, it has been supposed that the first protonation is accomplished by D1-Ser264 via D1-His252, while different possibilities have been explored for the mechanism of the second protonation [35, 36, 43]. Recently, by large-scale quantum mechanical/molecular mechanical approach based on the 1.9 Å resolution crystal structure, the D1-His215 was suggested as the sought $H^+$ donor for the keto-oxygen of $Q_BH^+$ proximal to the non-heme Fe [10]. The authors speculated that the resulting D1-His215 anion, in turn, is protonated in a redox-process, which most probably includes bicarbonate, D1-Tyr246 (Fig. 2, centre panel) and water as well as rearrangement of H-bonds network within the $Q_B$ binding niche.

2.3. Additional PSII Plastoquinone Binding Sites

A third plastoquinone molecule ($Q_C$) associated to PSIIcc was identified in the 2.9 Å resolution crystal structure of T. elongatus, and five of its isoprenoid units were successfully modelled into the PQ/PQH$_2$ exchange channel I, between the TMH of PsbJ and Cyt$b$$_{559}$ α and β-subunits [5]. In a subsequent 3.6 Å resolution crystallographic study of monomeric PSII core complex from T. elongatus, a trace of electron density was revealed in the same position but it was not possible to assign it to a specific molecule [6]. No electron density was detected in the putative $Q_C$ binding side in the most detailed 3D structure at 1.9 Å resolution from
Table 2. Factors involved in PSII plastoquinone interactions within QA, QB and QC binding niche, summarized from Hasagawa and Noguchi [23]. Amino acid residues providing attractive or repulsive effects with |ΔE'|≥2.0 kcal/mol and lipids and cofactors providing attractive or repulsive effects with |ΔE'|≥0.1 kcal/mol are presented. Amino acids, lipids and cofactors are ordered following the decrease of the interaction energy with the corresponding PQ molecule.

| Factors Involved in PQs Interactions | PQ at QA Binding Niche | PQ at QB Binding Niche | PQ at QC Binding Niche a |
|-------------------------------------|------------------------|------------------------|--------------------------|
| Protein Matrix                      |                        |                        |                          |
| Head attractive interactions        | D2-Phe261, -Thr217, D2-Tmp253, -His214, D2-Met246, -Ser262 | D2-Phe38, -Tyr42, D1-Phe211, PsbF-Leu26, -Thr25, D2-Arg26, -Arg128 | PsbJ-Gly20, -Val21, PsbJ-Gly18, -Ala17, Psb-Met19, -Ile24, PsbF-Thr30, -Ile31, PsbE-Phe31 |
| Tail attractive interactions        | D2-Gly200, -Gly203, D1-Phe52, PsbL-Leu30, PsbT-Phe10, PsbL-Leu27, D2-Phe270, D1-Ile77 | D1-Phe255, -Phe265, D1-Val219, -Leu218, D1-His215 | None |
| Repulsive interactions              | Head: D2-Gly215, D2-Arg251, Tail: D2-Leu279, D2-Val204, -Val274 | Head: D1-Gly253, D1-Asn266, Tail: D2-Leu43, PsbF-Val23, -Ala22, D1-Leu271, D2-Asp25, PsbE-Glu7 | Head: PsbE, Tail: PsbT |
| Lipids and Cofactors b               |                        |                        |                          |
| Head attractive interactions        | -                      | -                      | MGDG18, ChlD2, MGDG7, CarD2, DGDG6, SQDG4 |
| Tail attractive interactions        | -                      | ChlD2, MGDG18, SQDG4, DGDG6, MGDG7 | CarD2, MGDG18, DGDG6 |
| Repulsive interactions              | -                      | Head: MGDG18, ChlD2, Tail: MGDG7, CarD2 |

a For QC, all PSII amino acid residues providing attractive effects with |ΔE'|≥1.0 kcal/mol are presented.

b The abbreviations for lipids and molecular cofactors are same as define in Section 1 and the List of Abbreviations.

T. vulcanus [8], and this binding spot was neither occupied by PQ nor by PSII herbicide terbutryn in T. elongatus PSII/terbutryn crystal complex resolved at 3.2 Å resolution [7]. The uncertainty of QC presence in the different structures has been ascribed to differences in adopted purification and crystallization procedures, and photosynthetic sample pretreatment, to the low binding affinity of QC, or to the lack of a well-defined binding pocket [5, 7]. In fact, neither H-bonds nor π–π interactions were observed for QC stabilization and, most probably; the molecule binding is a result of the possible van der Waals interactions with the nearby lipid molecules, ChlD2, CarD2 and phytlyl tail of QB [5, 23], Table 2. PQ Head attractive interactions and (Fig. 2). FMO calculations based on the 2.9 Å resolution structure pointed out that more than 90% of the QC interaction energy with the protein matrix and with PSII integral lipids and cofactors is due to its isoprenoid chain (Table 1). In addition, the relative contribution of the interactions with the surrounding protein matrix to QC total binding energy is about 36%, that is much lower compared to that calculated for QA and QB [23] – about 100% and 82%, respectively (Table 1). The high mobility of the QC molecule has been attributed to its peculiar interaction with the environment, which is characterized by the head group rimming predominantly by a belt of lipids and cofactors, and the tail fencing to proteins (Table 2). The X-ray localization of a third PQ molecule in the 2.9 Å PSII structure supported earlier findings on the occurrence of additional quinones (different from QA and QB), located in the PQ hydrophobic cavity close to the Cytb559 [44, 45]. The physiological role of the QC is still under investigation. Guskov and coworkers [5] proposed three hypothetical mechanisms for PQ/PQH2 exchange, in which QC is alternating with QB or is the PQ molecule waiting to enter into the QB-binding site, thus contributing to the rapid turnover of the PQH2 with a new PQ molecule from the membrane pool. Experimental evidences have suggested a strong involvement of QC in a PSII photoprotective electron pathway mediated by Cytb559 [20-22]. It was hypothesized that Cytb559 could act as a plastoquinol oxidase (oxygen reductase) or a superoxide oxidase/reductase enzymes; these functions have been extensively documented in other reviews [15, 16, 36]. A distinctive feature of this heme-containing protein is its redox heterogeneity. In PSII membrane fragments 75% of Cytb559 is in its high potential form, 16% is in its intermediate potential form and 9% is in its low potential form; its redox properties and reactivity have been correlated to the occupancy of the QC site [22]. A biphasic pattern of Cytb559 reduction by PQH2 has been observed in PSII membrane fragments containing predominantly oxidized Cytb559, and experimental results suggested that a PQ/PQH2 molecule...
bound to the Q_C site is responsible for the fast phase (with rate constant of 100 ms$^{-1}$) of the Cyt.b$_{559}$ reduction [46]. The slow Cyt.b$_{559}$ reduction phase (with rate constant of 2.4 min$^{-1}$) was attributed to formation of redox equilibrium between a set of Cyt.b$_{559}$ redox forms and “free” PQ/PQH$_2$ from the membrane pool. The most likely mechanisms suggested by the authors for the Cyt.b$_{559}$ fast reduction requires: i) a location preferably encouraging one-electron redox-reaction; ii) a polar environment allowing accommodation of the PQ/SQ/PQH$_2$ and occurrence of the so called L substances in anionic form (such as ADRY reagents, dinoseb, high concentration of DCMU, tetraphenylboron); and iii) a fine-tuned Cyt.b$_{559}$/quinone interaction requiring well-defined quinone-protein interactions. However, the structure of the Q_C binding site described in the 2.9 Å resolution crystallographic study does not fulfil any of these requirements. The discrepancy between the structurally identified and experimentally suggested additional PSII PQ binding sites was found also in previous studies [20-22]. In order to distinguish between the Q_C site and the presumed polar quinone site of PSII that binds PQ/SQ/PQH$_2$/L, the authors proposed to assign the latter as Q_D [46]. These experiments led to hypothesize the existence of two different PQ binding sites, Q_C and Q_D, which host two different PQ molecules or one bound quinone that can occupy two different positions.

3. DYNAMICS STUDIES IN PHOTOSYNTHESIS RESEARCH

3.1. Neutron Scattering Techniques

The determination of protein structure and the analysis of conformational changes are the key necessary to understand the biological function and the activity of a protein. The structural characterization of proteins, however, constitutes only part of the puzzle because their function requires a specific dynamic. The protein motions arise on a wide range of timescales, from femtoseconds for bonds vibrations to milliseconds or seconds, for more complex processes that require a structural rearrangement. The dynamics of proteins contributes to the thermodynamic stability of their functional states and plays an important role in various processes, such as allosteric processes, where the correlations between structural fluctuations are able to transmit information between distant sites of the protein.

The study of dynamic processes at the microscopic level has been developed using a wide range of high resolution spectroscopies (Mössbauer spectroscopy, nuclear magnetic resonance (NMR), time-resolved fluorescence). We should add elastic, quasi-elastic and inelastic neutron scattering (ENS, QENS and INS, respectively) technique, which offers...
the advantage of covering length/distance (0.5-10 Å) and energy (0-200 meV) scales that correspond to thermal fluctuations and provide information on the geometry of the movements [47, 48]. The neutrons as a dynamics probe are highly sensitive to movements of hydrogen atoms, which are uniformly distributed within biological structures and represents nearly half of the atoms in them. On temporal and spatial scales accessible with neutron scattering, the hydrogen atoms trace the movements of the chemical groups, which they are linked to, such as amino acids side chains. In addition, it is also possible to mask part of the signal using specific deuterium, since deuterium has a weak scattering signal with respect to that of hydrogen [48]. The neutron scattering experiments, with their peculiar chance to exploit the isotopic substitution technique, permit to integrate and give complement to data obtained by numerical simulation and through specific molecular biology techniques. Moreover, they provide unique information useful for the characterization of the dynamical heterogeneity of biological macromolecules, from globular protein up to membrane complexes, in the temporal range $10^{-12} - 10^{-9}$ s.

The neutron scattering is an invaluable technique to tackle a plethora of important open questions in Biophysics such as the definition of correlations between structure, dynamics and functionality, the correlations between the interaction of water molecules with proteins and its role in the folding process and enzymatic activity, the observation of how the hydrogen bond network between solvent and biological macromolecules is affected and modified by selective genetic mutation, and hence the local flexibility and biological efficiency of the molecules [49-51].

### 3.2. Neutron Scattering in Photosynthetic Research

Photosynthetic research exploring the relationship between the function and the dynamics is a new field of interest. However, neutron scattering has been demonstrated to be a suitable tool to investigate protein dynamics in both complex multi-subunit photosynthetic systems and whole cells [52, 53].

A pioneer quasi-elastic neutron scattering study on the purple bacterium *Rhodobacter sphaeroides* RC proteins has demonstrated the effect of genetic point mutations for the overall protein dynamics [52]. The RC dynamics over a temperature range of 280 K was probed in a wild type and two non-functional mutants, carrying alanine residues on the place of Glu212 and Asp213 in the L-subunit. The Glu212 residue is highly conserved in bRC and is involved in the secondary ubiquinol protonation (see section 2.2). The crystallographic structure of the mutant proteins showed a local configuration of the active site more open than that of the wild type. One of the parameters indicative for the internal protein motions, the mean square displacement ($\langle u^2 \rangle$), calculated for the two mutants resulted to be higher than for the wild type protein suggesting a higher flexibility [52]. These results correlated the enhanced flexibility of mutated RC proteins with a lack of functionality, thus suggesting the requirement of a rigid core for the accomplishment of protein function in a multi-protein complex, such as RC involved in highly tuned photosynthetic reactions. This ground-breaking experiment opened the way to the urgency to understand if the core rigidity is essential for the charge transfer process, and if this property is shared by all the photosynthetic systems, and how this information can be applied to the design of bio-sensors and organic semiconductors.

An original green microalga species from the *Chlorococcal* order, which was never studied for its high tolerance to radiations, has been isolated in the Institute Laue Langevin research reactor storage pool. The microorganism able to live and reproduce in extreme environments was investigated by Farhi and coworkers combining neutron scattering experiments of internal dynamics with results of radiation effects on the microalgae physiology, metabolism and genomics [53]. The microalgae were stressed with radiation doses up to 20 kGy (2 Mrad), and studied by NMR, looking for modification in the metabolism, and by ENS, looking for both dynamical flexibility and structural macromolecular changes in the cells. While the sugar metabolism was slightly reduced and only at high radiation dose, it was probed that the effect of a high irradiation on micro-algae does not lead to protein denaturation, but rather to reduction of macromolecules flexibility – as showed by the reduction of mean square displacement ($\langle u^2 \rangle$) in (Fig. 2) of reference [53].

Pieper and coworkers have performed exhaustive neutron scattering studies on protein dynamics of PSII membrane fragments and antenna complex LHCII as a function of the temperature and sample hydration level [54]. They showed that PSII membrane fragments undergo a hydration dependent "dynamical transition" at about 240 K, underling the crucial role of hydration water for PSII flexibility. Above this temperature, PSII membrane fragments exhibit structural flexibility due to the presence of fast (picosecond) internal protein motions, while there is only harmonic vibrational dynamics at lower temperatures. It was then speculated that diffusive protein dynamics are indispensable for enabling QA reoxidation by QB at temperatures above 240 K, which explains the strong dependence of this electron transfer step on temperature and hydration level of the sample.

A newly developed technique of time-resolved QENS offers the advantage of simultaneous monitoring of protein internal dynamics and functional activity in specific sample functional states. This method was firstly applied for dynamics-function correlation in bacteriorhodopsin, a membrane embedded protein that functions as a light-driven proton pump in *Halobacterium salinarum*, and lately for the more complex PSII membrane fragments with inhibited QA to QB electron transfer [54, 55]. At physiological temperatures the calculated mean-square displacement of the hydrogen atoms, characterizing the flexibility of the system, revealed a significantly increased flexibility between 310 and 320 K [56]. The authors claimed that this transition is evidently correlated with the detachment of OEC from the membranes, as suggested by its absence in samples in which OEC has been removed prior to the QENS experiments. The transition is not accompanied by significant changes in the molecular organization of the pigment protein complexes, but the role of monomerization of PSII dimers cannot be ruled out.

Lately, preliminary results on intracellular water collective dynamics investigation on *Chlamydomonas* cells, carrying both native and mutated D1 RC proteins, have been also addressed. The authors proposed a comparison between fully
hydrated and partially de-hydrated deuterated whole cells. It was observed a distinct sound propagation speed between the two hydration levels suggesting a more rigid structure of hydration water than intracellular water [57].

4. COMPUTATION ANALYSES FOR MODIFICATION OF Qb BINDING POCKET AMINO ACID COMPOSITION: FUNDAMENTAL RESEARCH AND APPLICATIONS

In the last decades, several mutational studies on D1 protein were conducted to determine the effects induced by aminoacidic substitutions located in the stromal loop of the D1 protein. Mutations resulted in an impairment of the electron transfer between Q₃ and Q₈ leading to the identification of residues often involved in modulation of PSII photoinhibition susceptibility and/or herbicide specificity [58-62]. Computational studies on PSII D1 protein aimed at rational modification of the Q₈ binding pocket have been scarce partly because a high resolution three-dimensional structure of PSII has been available only relatively recently and partly due to the difficulties in generating site-specific mutants of this protein.

The first structure of PSII from _T. elongatus_ has been published only in 2004 and at a resolution of only 3.5 Å [PDB code 1S5L, 3]. The same year a 3.2 Å resolution structure has been published independently [PDB code 1W5C, 63] and refined the following year at 3.0 Å resolution [PDB code 2AXT, 4]. However, this last structure had still an insufficient resolution to uncover all the atomic details of this macromolecular proteins-cofactor complex such as the precise architecture of the OEC. Nonetheless, the Loll _et al._ crystal structure was sufficiently detailed to allow the first computational studies on the Q₈ binding pocket as well as homology modelling of PSIIs from other organisms. The first of such studies was probably that of Loll and coworkers [64] who modelled the three different copies of _T. elongatus_ D1 protein (PsbA1-A3) within the 3.0 Å resolution crystal structure of PSII, revealing that most of the D1 amino acid variants were located in the direct vicinity of redox-active cofactors of the electron transfer chain. However, proper computational studies aimed at selecting amino acid variants within the Q₈ binding pocket able to modify the recognition properties of the pocket itself appeared only the following year [65]. In this study, homology modelling of the _Chlamydomonas reinhardtii_ D1 protein was carried out based on the _T. elongatus_ crystal structure, followed by in _silo_ mutagenesis and energy minimization analyses to identify functional amino acids in the D1 protein interacting with herbicides. Following the results of theoretical calculations, three site directed mutants were produced by site-directed mutagenesis and characterized by fluorescence analysis. This study demonstrated that D1 mutants S268C and S264K could be used in the development of biocatalysts suitable for the selective detection of triazine and urea classes of herbicides, respectively. The study was further refined the same year undertaking an in _silo_ study of the _C. reinhardtii_ D1-D2 proteins with the aim of designing mutants with increased affinity for atrazine [66]. The final outcome of this work was the design and selection of D1 mutants hypothetically able to increase atrazine binding affinity by orders of magnitude, representing a useful theoretical guide for the development of high affinity atrazine biosensors.

With the increase in computing power, it has become feasible to adopt more sophisticated structural bioinformatics approaches, such as molecular dynamics (MD) simulations, for the study of the recognition properties of the Q₈ binding pocket toward natural and exogenous ligands. The first all atom MD simulations of the complete _T. elongatus_ PSII structure (including all cofactors) embedded in a lipid membrane was published in 2011 [67]. The aim of this work was that of uncovering the fine details of atrazine molecular interactions within the Q₈ binding pocket using a full atomistic (and thus realistic) description of PSIIcc. Atrazine was first docked in the vicinity of the Q₈ binding pocket and then let free to diffuse during a 10 ns MD simulations trajectory. During the simulations atrazine bound deeper into the Q₈ pocket establishing hydrophobic interactions with PSII D1-Phe255 phenyl ring and D1-Met214 side chain, and a single strong hydrogen bond with D1-His215 (Fig. 3). On the other hand, no binding partner appeared to stabilize other groups of the atrazine molecule such as the chloride atom (bound in an energetically unfavourable position in the vicinity of the aliphatic residue D1-Val219) and the N4 and N5 protons for which no hydrogen bonding partner was observed. The conclusions of this study were that rational design of site-directed mutants was likely to increase atrazine affinity for the Q₈ binding pocket. In particular, it was suggested that, among others, mutation of D1-Phe265 residue with a polar hydrogen bond acceptor residue could stabilize the atrazine N5 proton and atrazine binding within the pocket.

![Fig. (3). Schematic view of the interactions of atrazine within the PSII Q₈ binding pocket predicted through a 10 ns MD simulations run [66]. For clarity, only the D1 backbone (orange ribbon) and residues mentioned in the text are shown. The dashed green line highlights the hydrogen bond between atrazine and His215. Atrazine carbon atoms are in green, nitrogen atoms are in blue, sulphur atom is in yellow and the chloride atom is in magenta.](image)

CONCLUSIONS AND PERSPECTIVES

The advances in the crystallographic characterization of PSIIcc have broaden the understanding of PSII function and organization allowing to identify the atomic interactions tak-
ing place between the RC protein matrix, electron transport cofactors and thylakoid membrane lipids. Unravelling finer molecular details and mechanisms of the PSII structure, function and dynamics is mandatory to support novel biotechnological applications exploiting its unique properties of charge separation, electron transfer and water splitting. Specifically, it is possible to exploit PSII enriched particles or their active subcomponent to develop receptor elements suitable to be exploited in photosynthesis-based biosensors for the detection of environmental pollutants. The latter, in fact, can interact with the PSII donor side interrupting the light-induced electron transfer, generating a measurable optoelectronically detectable signal. In this context, insights into atomic interactions and dynamics of polypeptides and cofactors provided by both molecular dynamics or docking studies will help to design novel bio-recognition elements having improved selectivity and sensitivity towards different classes of compounds. Studies exploring the relationship between the function and the dynamics of the photosynthetic complex systems are still in their nascent stages, along with MD simulations. However, these techniques could offer a real possibility of investigating the dynamics associated with a molecule’s biological function and complement the progress of this technology. Besides, the extraordinary properties of water-splitting could be used to design bio-inspired device for bio-energy production, while the charge separation features could be exploited to design molecular photovoltaics.

**CONFLICT OF INTEREST**

The authors confirm that this article content has no conflicts of interest.

**ACKNOWLEDGEMENTS**

This publication is supported by Grant of COST Action TD1102. COST (European Cooperation in Science and Technology) is Europe’s longest-running intergovernmental framework for cooperation in science and technology funding cooperative scientific projects called ‘COST Actions’. With a successful history of implementing scientific networking projects for over 40 years, COST offers scientists the opportunity to embark upon bottom-up, multidisciplinary and collaborative networks across all science and technology domains. For more information about COST, please visit www.cost.eu.

**LIST OF ABBREVIATIONS**

| Abbreviation | Description |
|--------------|-------------|
| ADRY         | Acceleration of the deactivation reactions of system Y |
| bRC          | Bacterial reaction centre |
| Car          | Carotenoids |
| Chl          | Chlorophyll |
| CP43         | PSIIcc chlorophyll apoprotein encoded by \( psbC \) gene |
| CP47         | PSIIcc chlorophyll apoprotein encoded by \( psbB \) gene |
| Cyt.\( b_{559} \) | Cytochrome \( b_{559} \) |
| DGDG         | Digalactosyldiacylglycerol |
| ENS          | Elastic Neutron Scattering |
| FMO          | Fragment Molecular Orbital |
| INS          | Inelastic neutron scattering |
| L substances | Chemical compounds that induce negative shift in the midpoint potential of high potential form of Cyt.\( b_{559} \) |
| LHCII        | Light Harvesting Complex II |
| MD           | Molecular Dynamics |
| MGDG         | Monogalactosyldiacylglycerol |
| NMR          | Nuclear Magnetic Resonance |
| OEC          | Oxygen Evolving Complex |
| PDB          | Protein Data Base |
| PG           | Phosphatidylglycerol |
| RC           | Reaction Centre |
| Pheo         | Pheophytin |
| PQQ\( H_2 \) | Plastoquinol |
| PQs          | Plastoquinones |
| PsbE and PsbF PSIIcc proteins | Corresponding to \( \alpha \) - and \( \beta \)-subunit of cytochrome \( b_{559} \) |
| PSI          | Photosystem I |
| PSII         | Photosystem II |
| PSIIcc       | PSII core complex |
| Q\( A \)     | PSII primary quinone electron acceptor |
| Q\( B \)     | PSII secondary quinone electron acceptor |
| Q\( C \) and Q\( D \) | PQ molecules supposed to be bound to PSIIcc |
| QENS         | Quasi-elastic neutron scattering |
| SQ           | Semiquinones |
| SQDG         | Sulfocinnamoyl-diacylglycerol |
| terbutryn    | 2-tert-Butylamino-4-ethylamino-6-methylthio-1,3,5-triazin |
| TMH          | Trans-membrane helices |

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