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Review

Photosystem II and the unique role of bicarbonate: A historical perspective [☆]Dmitriy Shevela ^{a,*}, Julian J. Eaton-Rye ^b, Jian-Ren Shen ^c, Govindjee ^{d,e,f,g,**}^a Centre for Organelle Research, University of Stavanger, Kristine Bonnevis vei 22, N-4036 Stavanger, Norway^b Department of Biochemistry, University of Otago, PO Box 56, Dunedin 9054, New Zealand^c Division of Bioscience, Graduate School of Natural Science and Technology/Faculty of Science, Okayama University, Okayama 700-8530, Japan^d Department of Plant Biology, University of Illinois at Urbana-Champaign, 265 Morrill Hall, 505 South Goodwin Avenue, Urbana, IL 61801, USA^e School of Life Sciences, Jawaharlal Nehru University, New Delhi, 110067, India^f Department of Biochemistry, University of Illinois at Urbana-Champaign, 265 Morrill Hall, 505 South Goodwin Avenue, Urbana, IL 61801, USA^g Center of Biophysics & Computational Biology, University of Illinois at Urbana-Champaign, 265 Morrill Hall, 505 South Goodwin Avenue, Urbana, IL 61801, USA

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ABSTRACT

In photosynthesis, cyanobacteria, algae and plants fix carbon dioxide (CO₂) into carbohydrates; this is necessary to support life on Earth. Over 50 years ago, Otto Heinrich Warburg discovered a unique stimulatory role of CO₂ in the Hill reaction (i.e., O₂ evolution accompanied by reduction of an artificial electron acceptor), which, obviously, does not include any carbon fixation pathway; Warburg used this discovery to support his idea that O₂ in photosynthesis originates in CO₂. During the 1960s, a large number of researchers attempted to decipher this unique phenomenon, with limited success. In the 1970s, Alan Stemler, in Govindjee's lab, perfected methods to get highly reproducible results, and observed, among other things, that the turnover of Photosystem II (PSII) was stimulated by bicarbonate ions (hydrogen carbonate): the effect would be on the donor or the acceptor, or both sides of PSII. In 1975, Thomas Wydrzynski, also in Govindjee's lab, discovered that there was a definite bicarbonate effect on the electron acceptor (the plastoquinone) side of PSII. The most recent 1.9 Å crystal structure of PSII, unequivocally shows HCO₃⁻ bound to the non-heme iron that sits in-between the bound primary quinone electron acceptor, Q_A, and the secondary quinone electron acceptor Q_B. In this review, we focus on the historical development of our understanding of this unique bicarbonate effect on the electron acceptor side of PSII, and its mechanism as obtained by biochemical, biophysical and molecular biological approaches in many laboratories around the World. We suggest an atomic level model in which HCO₃⁻/CO₃²⁻ plays a key role in the protonation of the reduced Q_B. In addition, we make comments on the role of bicarbonate on the donor side of PSII, as has been extensively studied in the labs of Alan Stemler (United States) and Vyacheslav Klimov (Russia). We end this review by discussing the uniqueness of bicarbonate's role in oxygenic photosynthesis and its role in the evolutionary development of O₂-evolving PSII. This article is part of a Special Issue entitled: Photosynthesis Research for Sustainability: from Natural to Artificial Photosynthesis.

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1. Introduction

1.1. Role of inorganic carbon and its interconversion in living organisms

Carbon dioxide (CO₂) is not only a greenhouse gas in the Earth's atmosphere, but also a key metabolite in living organisms, where it plays an essential role in such fundamental biological processes as respiration and

photosynthesis. Due to its ability to exist in equilibrium with carbonic acid (H₂CO₃) and bicarbonate (HCO₃⁻, IUPAC's recommended term is *hydrogen carbonate*, but in this review we use its traditional and well-known term '*bicarbonate*') (see Fig. 1); CO₂ produced by cells during aerobic metabolism of glucose and fats provides the acid (H⁺ and CO₂) and base (HCO₃⁻) components for the so-called *bicarbonate buffering system*. This buffering system maintains both intracellular and extracellular pH.

Abbreviations: CA, carbonic anhydrase; Chl, chlorophyll; DCMU, 3-(3,4-dichlorophenyl)-1,1-dimethylurea (diuron); DPC, 1,5-diphenylcarbazine; HCO₃⁻, hydrogen carbonate (bicarbonate) ion; MS, mass spectrometry; NHI, non-heme iron; OEC, oxygen-evolving complex; P680, primary electron donor molecule (Chl) in Photosystem II; P700, primary electron donor molecule (Chl) in Photosystem I; Pheo, pheophytin; PQ, plastoquinone; PQH₂, plastoquinol; PSI, Photosystem I; PSII, Photosystem II; Q_A, primary quinone electron acceptor of PSII; Q_B, secondary quinone electron acceptor of PSII; RC, reaction center; S_i, redox state of the OEC, where *i* is the number of stored oxidizing equivalents; TL, thermoluminescence

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* Corresponding author at: Centre for Organelle Research, University of Stavanger, Kristine Bonnevis vei 22, N-4036 Stavanger, Norway. Tel.: +47 518 31810; fax: +47 518 31860.

** Corresponding author at: Department of Plant Biology, University of Illinois at Urbana-Champaign, 265 Morrill Hall, 505 South Goodwin Avenue, Urbana, IL 61801, USA. Tel.: +1 217 337 0627; fax: +1 217 244 7246.

E-mail addresses: dmitry.shevela@uis.no (D. Shevela), gov@illinois.edu (Govindjee).

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The interconversion of inorganic carbon, on the other hand, allows rapid transport of its species ($\text{CO}_2/\text{HCO}_3^-/\text{CO}_3^{2-}$) in all cells. While HCO_3^- is poorly soluble in biological membranes, CO_2 can freely diffuse in and out of the cell. Therefore, $\text{HCO}_3^- \rightarrow (\text{H}_2\text{CO}_3) \rightarrow \text{CO}_2$ interconversion facilitates the transport of inorganic carbon in the form of CO_2 into intracellular space, while the reversed conversion ($\text{CO}_2 \rightarrow (\text{H}_2\text{CO}_3) \rightarrow \text{HCO}_3^-$) provides trapping of the CO_2 within the cell in the form of HCO_3^- . Although the reversible hydration of CO_2 and dehydration of HCO_3^- occurs spontaneously, even in the absence of catalysts, most—if not all—organisms have Zn-containing *carbonic anhydrases* (CAs) that catalyze this ubiquitous conversion. By speeding up these reactions (k_{cat} can reach $\sim 1 \times 10^6 \text{ s}^{-1}$), CAs play an essential role in a wide range of biochemical and physiological processes [1].

All photosynthetic organisms need atmospheric CO_2 to store harvested energy from sunlight in the form of energy-rich carbohydrates. However, in the oxygenic photosynthesizers (cyanobacteria, algae and higher plants), CO_2 is not only required as the terminal electron acceptor to synthesize carbohydrates, but also for the regulation of photosynthetic electron transport in Photosystem II (PSII), the enzyme responsible for light-induced primary charge separation and subsequent water oxidation [2]. The latter is known as the 'bicarbonate effect'. This review summarizes historical discoveries related to the 'bicarbonate effect' and outlines our current state of knowledge about the location and role of HCO_3^- in PSII.

1.2. The 'bicarbonate effect' and Otto Heinrich Warburg

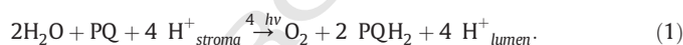
Despite the fact that the reduction of CO_2 to carbohydrates within the Calvin–Benson cycle is driven by the products of the light reactions of photosynthesis, ATP and NADPH, it does not directly require light, and thus, belongs to the photosynthetic light-independent ('dark') reactions [3]. Until the discovery (1958) of the 'bicarbonate effect' on the light-dependent electron flow by Otto Warburg and Günter Krippahl, CO_2 was assumed to be involved only in the 'dark' and not in the 'light' reactions [4]. Warburg and Krippahl found that the high rates of the Hill reaction (the reaction which allows the study of electron flow in isolated broken chloroplasts illuminated in the presence of an artificial electron acceptor *via* the measurements of O_2 production) required the presence of CO_2 in the gas phase above the sample suspension. Specifically, it was shown that the O_2 evolution rate measured in grana isolated from kohlrabi leaves in the presence of quinone (as electron acceptor) was significantly higher when argon atmosphere above the sample contained 1.4% CO_2 (v/v) (see Fig. 2). In spite of the earlier difficulties with its reproducibility, many research groups confirmed this phenomenon (outlined in Section 3.1). Later on, Alan Stemler and Govindjee [5] significantly improved reproducibility of the bicarbonate effect by developing a reliable method of $\text{CO}_2/\text{HCO}_3^-$ depletion.

Otto Warburg believed that the observed phenomenon provides evidence for his 'photolyte theory', in which O_2 originates from the splitting of 'activated CO_2 ', not from water. In 1964, he noted "As was expected, no proof of water photolysis survived the discovery of 'active CO_2 '" [6]. Despite this mistaken interpretation, the finding made by Warburg and Krippahl was fundamental to subsequent research of the 'bicarbonate effect' on light-induced electron transport

during photosynthesis. Their discovery initiated long-term debates about possible action site(s) and role(s) of inorganic carbon on photosynthetic O_2 production. Thus, intensive studies by many laboratories explored the possibility that HCO_3^- (CO_3^{2-}) (and not CO_2) is required for both PSII electron transport efficiency and for the photo-assembly of the inorganic core (the Mn_4CaO_5 cluster) of the O_2 -evolving complex (OEC) of PSII (see Sections 2 and 3; for previous historical overviews, see [7–14]).

1.3. Photosystem II and the sites of the 'bicarbonate effect'

PSII is a large multi-component pigment-protein complex, which is incorporated into the thylakoid membrane of all oxygenic photosynthetic organisms (for reviews on PSII, see [2,15]). Fig. 3 shows a schematic view of PSII in higher plants and green algae and its important redox cofactors, which are thought to be the same as in cyanobacteria (for further details on the cyanobacterial PSII structure, see [16] and [17]). PSII acts as a water:plastoquinone oxidoreductase, catalyzing the following reaction:



Thus, the light-induced charge separation between the reaction center (RC) chlorophyll (Chl) molecules in the D1 protein (Chl_{D1} and P680) and pheophytin (Pheo_{D1}), and the formation of the stabilized radical ion pair $\text{P680}^{+\bullet}\text{Pheo}^{-\bullet}$ (for reviews, see [18,19]), lead to two reactions: (1) water splitting (oxidation) to O_2 , protons and electrons on the luminal side of PSII, the so-called *electron donor side* of PSII with $\text{P680}^{+\bullet}$ as the driving force and, (2) the reduction of plastoquinone (PQ) to plastoquinol (PQH_2) on the stromal, the *electron acceptor side*, of PSII with proton uptake and $\text{Q}_\text{A}^{-\bullet}$ acting as the reductant.

Extensive data show that HCO_3^- ions, under appropriate experimental conditions, have effects on both the acceptor side and the donor side reactions of PSII (Fig. 3). The focus of this review is the bicarbonate effects related to the electron acceptor side, where HCO_3^- is known to bind (see Fig. 3 and Section 3.2 for the current model) and to play an important role in facilitating the reduction of Q_B , and in protonation reactions near the Q_B -site. We, however, briefly discuss here, for completeness, the possible roles of HCO_3^- on the 'donor side' reactions of PSII. For historical surveys on the discoveries of the HCO_3^- effect on the PSII donor side, see several references [9,10,14,20–22].

2. Bicarbonate and the donor side of Photosystem II

The role of HCO_3^- on the donor side reactions of PSII has been extensively studied by many researchers, but mainly in the laboratories of Alan Stemler (University of California, USA) and Vyacheslav Klimov (Institute of Basic Biological Problems, Russia), as mentioned earlier.

In the early 1970s, Stemler with co-workers [5,23] were the first to propose the water-oxidizing side of PSII as a possible site for the HCO_3^- effect (Fig. 4). However, in 1975, Thomas Wydrzynski and Govindjee obtained evidence for the participation of HCO_3^- ions in the electron transfer kinetics on the acceptor side of PSII [24] (Fig. 5). This discovery was supported by numerous subsequent experiments (see

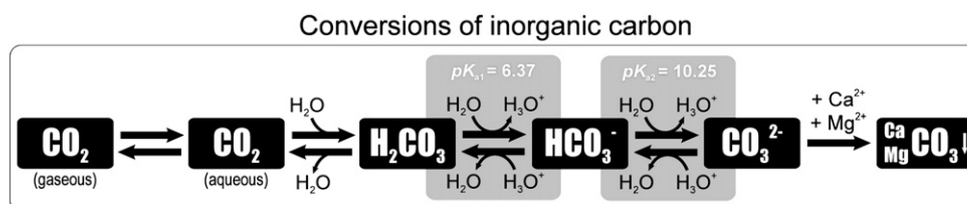


Fig. 1. Conversion of inorganic carbon species including acid–base ionization/dissociation constant (pK_a) values for hydrogen carbonate (bicarbonate) anion. See text for further details.

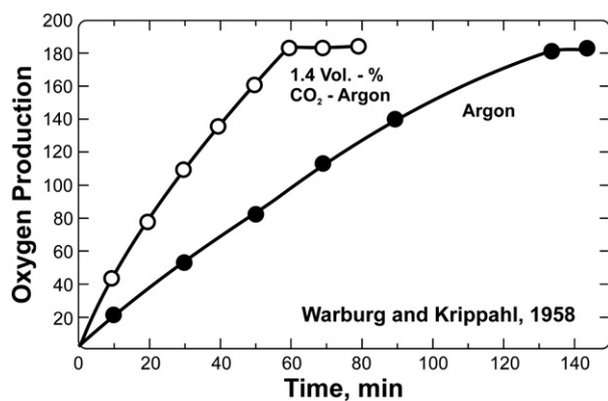


Fig. 2. CO₂ (HCO₃⁻) effect on the rate of the Hill reaction as was found by Otto Warburg and Günter Krippahl [4] in isolated kohlrabi grana suspended in 0.1% KCl. The measurements were performed in the presence of 2.1 mg quinone as electron acceptor under argon (closed symbols) or argon + 1.4% CO₂ (v/v) (open symbols) in the gas phase.

165 Section 3), and later on, the non-heme iron (NHI) between Q_A and Q_B
 166 was shown to play an essential role in HCO₃⁻ binding [12,25]. On the
 167 other hand, some (among them Helmut Metzner, Werner Kreuzt, and
 168 Alan Stemler) believed that HCO₃⁻ may act as a substrate or a chemical
 169 intermediate in photosynthetic O₂ evolution, possibly coupled with CA
 170 activity [26–28]. Thus, Stemler and collaborators continued to investigate
 171 the possible involvement of HCO₃⁻ ions in the mechanism of O₂ evolution
 172 on the oxidizing side of PSII (reviewed in [9,22]). Stemler's reports, as well
 173 as reports of some others [29,30] indicated that HCO₃⁻ may affect both the
 174 electron acceptor and donor sides of PSII. Undoubtedly, however, the dis-

covery of the 'acceptor-side' effect inadvertently affected the search for
 specific effects of HCO₃⁻ on the donor side, and inevitably led to a contro-
 versy on the interpretation of the 'bicarbonate effects'.

Since the mid 1990s, the idea for an additional role of HCO₃⁻ on the
 electron donor side of PSII was revived by a series of experiments per-
 formed in the laboratory of Vyacheslav Klimov. The studies by Klimov
 and collaborators indicated that HCO₃⁻ ions are required for (1) the
 efficient photo-induced assembly of the Mn₄CaO₅ cluster capable of
 water splitting, (2) the stability of the OEC, and (3) the protection
 of the donor side of PSII against photoinhibition and thermoinactiva-
 tion (reviewed in [10,14]). Other groups (see, for instance, [31–35])
 also obtained indication for the requirement of HCO₃⁻ on the water-
 splitting side of PSII. However, the binding site(s) and the role(s) of
 HCO₃⁻ ions in the water-splitting reaction of PSII remain unclear
 (and, therefore, appear 'questionable' to the authors; see Fig. 3). The
 following main proposals for the involvement of HCO₃⁻ in the events
 on the water-oxidizing side of PSII have been considered:

- (i) Exchangeable HCO₃⁻ is an intermediate substrate for photosynthetic
 water oxidation; water is delivered to the Mn₄CaO₅ cluster in the
 form of HCO₃⁻ (or peroxidicarbonic acid; H₂C₂O₆). Initially proposed
 by Helmut Metzner [27] as an alternative to Warburg's 'photolyte
 theory' (mentioned above) and later elaborated by Alan Stemler,
 [26] and by Paul Castelfranco with co-authors [36], this hypothe-
 sis has become obsolete, in our opinion, because of various studies
 using isotope ratio mass spectrometry (MS) in combination with
¹⁸O-labeling of H₂O and HCO₃⁻ [37–41], UV spectrophotometry
 under high backpressure of CO₂ [38], and light-induced FT-IR dif-
 ference spectroscopy [42].

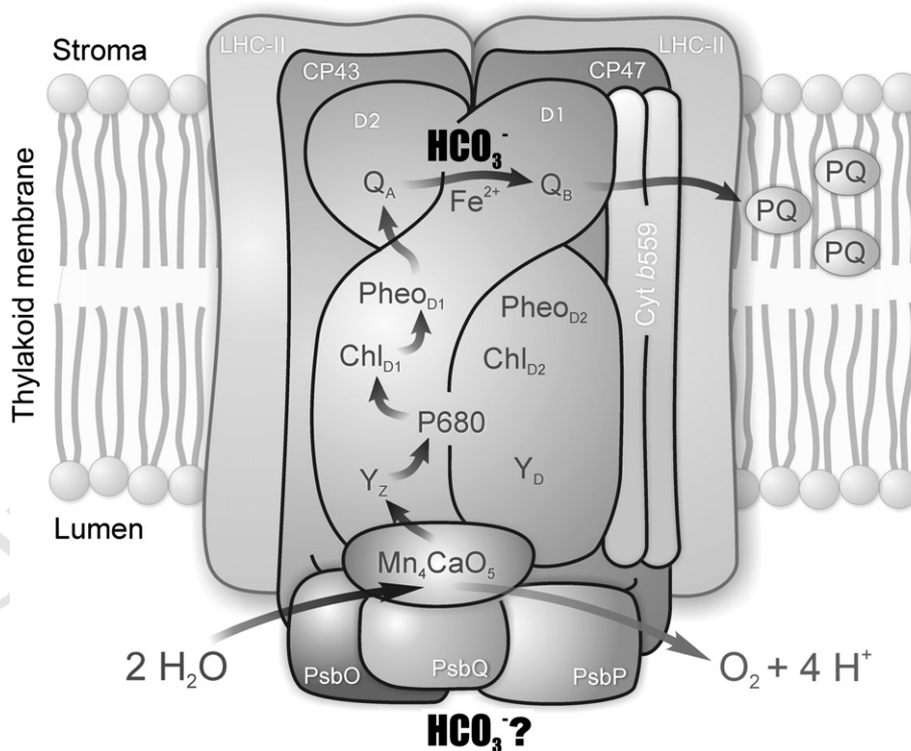


Fig. 3. Schematic representation of PSII in higher plants and green algae (only core proteins are shown) and two sites (acceptor and donor) where bicarbonate (HCO₃⁻; hydrogen carbonate) has effects. While the acceptor side bicarbonate is known to bind to the NHI (Fe²⁺) between Q_A and Q_B, the exact location of the donor side bicarbonate is unknown. The acceptor side bicarbonate may also be bound to the NHI in the form of carbonate (CO₃²⁻). In cyanobacteria the sites of HCO₃⁻ effects are the same, but some components of PSII are different (for further details see [15]). The pathway of the electron flow through PSII is shown by black arrows. Other abbreviations: D1 and D2, the reaction center proteins; P680, the reaction center Chl molecule; Chl_{D1}, the primary electron donor on D1; Pheo_{D1}, the primary electron acceptor on D1 (pheophytin); Chl_{D2} and Pheo_{D2}, symmetrically related cofactors on D2 (inactive branch; do not participate in linear electron transfer through PSII); Mn₄CaO₅, inorganic core of the OEC; Y_Z (on D1) and Y_D (on D2), the redox active tyrosine residues; PQ, mobile plastoquinone molecule; CP43 and CP47, Chl-protein complexes of 43 and 47 kDa; LHC-II, light-harvesting complex II; PsaO (33 kDa), PsaP (23 kDa) and PsaQ (17 kDa), extrinsic proteins of PSII; Cyt b559, redox active cytochrome b559.

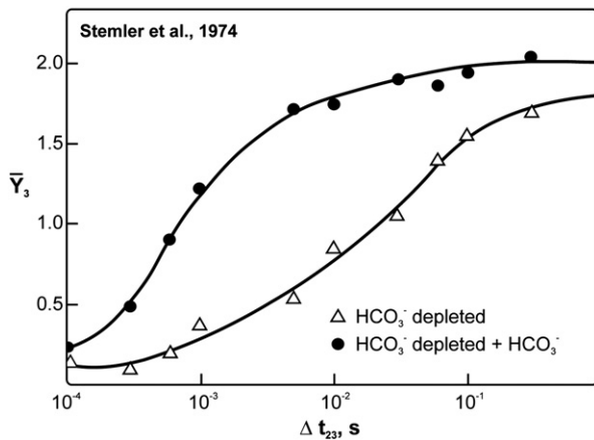


Fig. 4. O₂ yield obtained on the third flash (Y_3) as a function of the dark-time between the second and the third flash (Δt_{23}) as measured by Stemler et al. [23] in dark-adapted HCO₃⁻-depleted chloroplast suspensions in the presence and the absence of bicarbonate. The frequency of the main flash train was 1 Hz. Y_3 values were normalized with respect to the steady-state O₂ yield. Open triangles: HCO₃⁻-depleted chloroplast suspensions were injected onto the Pt electrode to final Chl concentration of 0.3 mg ml⁻¹. The measurements were performed in buffered medium, which contained 0.25 M NaCl, 0.04 M Na acetate, 0.05 M Na phosphate buffer (pH 6.8), 20 μg ml⁻¹ of ferredoxin, and 0.5 mM NADP⁺. Closed circles: the same as above but after re-addition of 10 mM NaHCO₃. Adapted and modified from [23].

(ii) *Non-exchangeable, tightly bound HCO₃⁻ being a structural part of the Mn₄CaO₅ cluster may alter the redox properties of the Mn cations, and thus, is required for the functionality and stability of the assembled OEC.* The studies carried out by Klimov and collaborators showed stabilizing and protective effects of HCO₃⁻ on the donor (water-oxidizing) side of PSII [43–50]. One of the interpretations of the

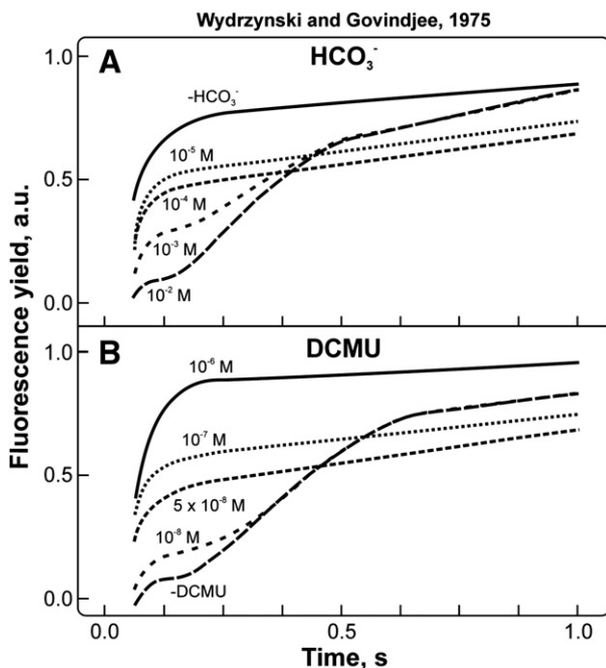


Fig. 5. First experimental evidence for the action of bicarbonate on the electron acceptor side of PSII reported by Wydrzynski and Govindjee in 1975 [24]. A comparison of variable Chl *a* fluorescence on concentration of HCO₃⁻ and an inhibitor of PSII DCMU. (A) HCO₃⁻-depleted chloroplasts at various concentrations of NaHCO₃. (B) Non-HCO₃⁻-depleted chloroplasts at various concentrations of DCMU. Before the measurements, the samples were incubated in the dark for 5 min. Note that the fluorescence induction curves obtained under HCO₃⁻-free conditions look like the one obtained after addition of 10⁻⁶ M DCMU. Fluorescence was measured at 685 nm upon excitation with a broad blue light at a Chl concentration of 12.5 μg ml⁻¹. Modified and adapted from [24].

observed effects was the idea that HCO₃⁻ may function as a ligand to the Mn₄CaO₅ cluster or an integral cofactor of the OEC [10]. In addition, in the PSII crystal structure by Ferreira et al. [51] at a resolution of 3.5 Å, HCO₃⁻ (or CO₃²⁻) anion was tentatively included as a ligand bridging Mn and Ca ions within the OEC. However, the latter could not be supported by the most recent X-ray crystallography studies of PSII at higher resolutions [16,17,52]. At the same time, all these crystallographic studies clearly displayed HCO₃⁻ as a ligand of the NHI between Q_A and Q_B (for further details, see Section 3.2). Earlier sensitive differential infrared gas analyzer and MS measurements [7,53] also clearly showed only ~1 HCO₃⁻/CO₂ molecule bound per PSII RC (see Section 3.1.2.3). Further, a recent re-examination of the structural coupling of HCO₃⁻ to the OEC by FT-IR spectroscopy provided no indication for any HCO₃⁻-bands from the OEC during the S-state transitions [42]. This is also consistent with the results obtained by flash-induced O₂ evolution pattern (FIOP) studies, where the redox potentials of the S states of the OEC were found to be unaffected by HCO₃⁻ depletion via washing with CO₂/HCO₃⁻-free buffer [54]. Moreover, evidence for the absence of tightly bound HCO₃⁻ in the first coordination sphere of the Mn₄CaO₅ cluster was obtained by isotope ratio MS [41] (for details, see Section 3.1.3.3) and GC-MS [55] studies. In addition, HCO₃⁻, as a structural part of the OEC has not been supported by the computational models based on density functional theory (DFT) and quantum mechanics/molecular mechanics (QM/MM) studies [56,57]. It is, therefore, very unlikely that HCO₃⁻ is a ligand or strongly coupled cofactor to the Mn₄CaO₅ cluster in its assembled state.

(iii) *Acting as a transient ligand to Mn ions, HCO₃⁻ is a native cofactor in the photo-assembly (photo-activation) process of the Mn₄CaO₅ cluster that assembles in the OEC-depleted PSII centers that are free of inorganic cofactors, but HCO₃⁻ is not part of the assembled cluster.* This suggestion is based on the results obtained by Klimov and co-workers [10] demonstrating a pronounced stimulating effect of HCO₃⁻ ions on the electron donation from exogenous Mn²⁺ ions to Mn-depleted PSII and the photo-induced reconstitution of the functional OEC [43–45,48,58]. Further experiments in collaboration with the group of Charles Dismukes provided evidence for the requirement of HCO₃⁻ (CO₃²⁻) for *in vitro* light-driven assembly of the Mn₄CaO₅ cluster (for details, see [31,59]; for reviews on the photo-assembly of the OEC, see [60,61]). Although electrochemical characterizations of Mn-HCO₃⁻ complexes [62–64] as well as electron paramagnetic resonance/electron spin echo envelope modulation (EPR/ESEEM) spectroscopy studies of assembly intermediates [65,66] strongly support this idea, there is no experimental data demonstrating a HCO₃⁻ requirement for the assembly process *in vivo*.

(iv) *HCO₃⁻ indirectly stabilizes the OEC by binding to extrinsic proteins or some other protein components of PSII in the vicinity of the Mn₄CaO₅ cluster.* Pobeguts et al. [67] demonstrated a protective effect of HCO₃⁻ against extraction of the extrinsic proteins (especially PsbO—the Mn-stabilizing protein) of the OEC after treatment of pea PSII membrane fragments with urea. Moreover, the specific high-affinity binding of HCO₃⁻ (or CO₂) to the PsbO protein has been proposed based on the recent observations of HCO₃⁻-dependent re-arrangements in the PsbO protein [68]. However, as mentioned above, no HCO₃⁻ was detected bound on the protein components belonging to the donor side of PSII of thermophilic cyanobacteria by Umena et al. [17] in their recent crystal structure at 1.9 Å resolution. Nevertheless, since significant differences are known to exist between proteins of cyanobacteria and plants (reviewed in [69,70]), the possibility of HCO₃⁻ binding to protein components in higher plants needs to be addressed by future experiments.

(v) *Mobile, exchangeable HCO₃⁻ is involved in proton removal during photosynthetic water oxidation; it may work coupled with the*

PSII-donor-side-associated CA. Deprotonation reactions and removal of protons away from the OEC are thought to have significant impact on the thermodynamics of water splitting [71]. Ananyev et al. [34] proposed that HCO_3^- may play an indirect role in water splitting as a proton transfer mediator and recent results support this proposal [32,35]. In fact, such an interpretation of the bicarbonate effect on the water-oxidizing side of PSII may explain a large body of existing data, especially in connection with the PSII-associated CA activity, which was experimentally shown in many studies [72–78]. Stemler, based on circumstantial evidence, was the first to suggest that a thylakoid CA might be involved in the ‘donor-side’ effects of HCO_3^- [74,79–81]. Experiments by Shutova et al. [32] show that in *Chlamydomonas* (*C.*) *reinhardtii*, both HCO_3^- and Cah3 (the CA protein in *C. reinhardtii* associated with the PSII donor side) have specific ‘donor-side’ effects on proton release steps, but not on electron transfer. Moreover, there are also some indications for a similar role of CA and HCO_3^- in higher plants, although both CA and HCO_3^- requirements were found to be lower than that observed in *C. reinhardtii* [82]. Shutova et al. [32] suggested that a CA/ HCO_3^- system in *C. reinhardtii* may facilitate proton removal away from the OEC during water splitting by accelerating interconversion between HCO_3^- and CO_2 (see Fig. 1). Indeed, if the luminal “working” pH under illumination is 5.4–5.7, as shown recently [83,84], one can assume that due to the strong deficit of HCO_3^- species at this pH range, the presence of CA activity is ‘naturally’ required for the fast production of these species from CO_2 .

We note here, that to our knowledge, most of the above bicarbonate-related investigations of cyanobacteria, algae, and higher plants, except for a few with intact alga *Chlamydomonas stellata* [29] and the hypercarbonate-requiring cyanobacterium *Arthrospira maxima* [85,86] have been limited to *in vitro* studies of isolated thylakoids, PSII membrane fragments and PSII particles. Therefore, we emphasize that the effect and the function of HCO_3^- on the donor side of assembled PSII may be different (if any) when the protein environment is intact as to when it is disrupted, e.g., as a result of sample preparation. Thus, further research is needed to study this option and to elucidate the role of HCO_3^- on the water-oxidizing side of PSII.

3. Bicarbonate and the acceptor side of Photosystem II

In contrast to what little is known regarding the effect of bicarbonate on the donor side of PSII, we know a great deal about the role of HCO_3^- on the acceptor-side of PSII—and its binding is obvious in the high resolution structures of cyanobacterial PSII RCs. Therefore, the remaining historical perspective in this review will focus on the research efforts related to the discoveries of the bicarbonate effect on the electron flow within PSII, and the HCO_3^- -dependent regulation of electron transport on the acceptor side. Hence, the current state of our knowledge about the location and the function of HCO_3^- is also discussed in detail.

3.1. Time-line of discoveries on HCO_3^- in the electron flow of Photosystem II

3.1.1. The early work: from Otto Warburg to Norman Good

In 1948, Boyle [87] had observed that O_2 evolution by ground-up spinach leaves, when *p*-benzoquinone was added, was absent when KOH was included (to absorb CO_2) in the center well of a manometer vessel; thus, Boyle concluded that CO_2 was necessary for the benzoquinone Hill reaction. Although the conclusion was confirmed by Warburg and Krippahl in 1960 [88], Boyle’s results were artifacts as suggested by Warburg and Krippahl (1958) [4] and, as shown, in 1961, by Abeles et al. [89]: benzoquinone in the main

vessel distilled into the KOH-soaked filter paper in the center well and the mixture consumed O_2 balancing O_2 evolution from the broken leaves. The discovery of the ‘bicarbonate effect’ by Warburg and Krippahl has already been mentioned in Section 1.2.

- In 1961, Abeles et al. [89] confirmed Warburg and Krippahl’s results in kohlrabi chloroplasts, i.e., requirement of CO_2 for the Hill reaction. Warburg’s idea that O_2 arose from CO_2 (see Section 1.2) had to be tested. For this purpose, Abeles et al. used MS that distinguishes O_2 evolution and metabolism of CO_2 . They observed changes only in O_2 release and none in CO_2 metabolism. Thus, Warburg’s idea was not supported. However, Abeles and co-workers could not reproduce this effect in sugar beet chloroplasts leading them to conclude that the effect was not universal.
- During 1960–1962, Stern and Vennesland [90,91] observed that the ferricyanide-supported Hill reaction, in spinach and kohlrabi chloroplasts suspended in buffered media, declined much faster, with time, with CO_2 present. Addition of CO_2 restored Hill activity. Further, in 1963, Vennesland, who was still supporting Warburg’s point of view, reported stimulation of the Hill reaction with different electron acceptors, using thylakoids from various plant sources [92].
- In 1962, Izawa [93] introduced the use of CA to the reaction medium, while CO_2 was being removed; this hastened the time of CO_2 -depletion and gave much more reliable results; Izawa found larger effects in broken than in intact chloroplasts.
- In 1963, Heise and Gaffron [94] reported decreases in O_2 evolution during the Hill reaction with *p*-benzoquinone in the cyanobacterium *Anacystis nidulans* (*Synechococcus elongatus* strain PCC 7942) and in the green alga *Scenedesmus obliquus* (strain D3) in the absence of CO_2 . However, these authors suggested that this effect is not an important one since many different metabolic reactions have been shown to be dependent on traces of CO_2 .
- During 1963–1965, Good [95,96] discovered that CO_2 dependence of the Hill reaction, in pea chloroplasts, was highly influenced by the addition of anions, particularly of formate and acetate; none of the anions used could act as bicarbonate; thus, bicarbonate was considered to have a specific stimulatory effect in electron transport during the Hill reaction; uncouplers of phosphorylation had no effect on electron transport in CO_2 -depleted chloroplasts.
- During 1964 and 1965, Punnett and Iyer [97], Punnett [98] and Batra and Jagendorf [99] discovered that in addition to the effects of $\text{CO}_2/\text{HCO}_3^-$ on electron transport, an additional, although a different effect, exists on photophosphorylation. In their 1978 review on the bicarbonate effect, Govindjee and Van Rensen [11] have called this separate effect, the “Punnett Effect”; however, it will not be discussed further in this review since CO_2 was not a requirement for phosphorylation, whereas it is a requirement for electron transport.
- In 1967, West and Hill [100] confirmed the existence of the stimulatory role of CO_2 in both dichlorophenol indophenol (DCPIP) and ferricyanide Hill reactions in pea chloroplasts, and as Izawa had stated, the effect was larger in broken, than in intact, chloroplasts.

3.1.2. Work at the University of Illinois at Urbana-Champaign (UIUC): from Alan Stemler to Jin Xiong and collaborations with other groups

Most of the research up to this point was aimed to see if there was an effect of CO_2 on the Hill reaction, i.e., electron flow from water to NADP^+ (see a review [11]). There were, in general, considerable variations in the magnitude of the effects; further, the conditions producing them showed considerable differences. Govindjee, one of the authors of this review, presented a lecture to a graduate level course in late 1960s or early 1970s, where he talked about this effect emphasizing the *out-of-this-world* ideas of Otto Warburg—that this effect implies that O_2 comes from CO_2 . To the surprise of Govindjee, one of his own doctoral students in the class wanted to pursue this as his PhD thesis project. Govindjee attempted to discourage such an undertaking as it was very risky, but then the student Alan Stemler persisted. The rest is history. We present below a time line of research from 1973 to 1998 in Govindjee’s laboratory in this area of research (also see a different perspective in Stemler’s reviews

[9,21]). We define the ‘bicarbonate effect’ as follows: addition of bicarbonate to $\text{CO}_2/\text{HCO}_3^-$ -depleted samples restores fully the electron transport to that without the depletion process.

3.1.2.1. 1970s: research at the UIUC and collaboration with labs in Berkeley, Leiden and Berlin.

In 1973, Stemler and Govindjee [5] worked out a procedure (by flushing isolated broken chloroplasts with nitrogen in a medium containing a high anion concentration at low pH) to remove bicarbonate; they obtained a large (5-fold) and reproducible effect of bicarbonate on the DCPIP Hill reaction in these samples; they suggested that HCO_3^- was bound in darkness and released in light. In view of their experiments with diphenylcarbazide (DPC), they had suggested that the effect was only on the O_2 -evolving side of PSII. This suggestion was challenged in 1975 by Wydrzynski and Govindjee [24] (see below).

In early 1974, Stemler and Govindjee [101] proceeded to perfect the methods of HCO_3^- -depletion further including the effects of light intensity and differences between the rates of O_2 evolution and ferricyanide reduction, again in broken maize chloroplasts, suggesting the possible existence of non- O_2 -evolving centers, and even an effect of bicarbonate on the rate of photoinactivation. These concepts still remain to be further investigated.

In 1974, Stemler and Govindjee [102] reported, working still with broken maize chloroplasts, complex effects of bicarbonate on Chl *a* variable fluorescence induction and delayed light emission, including an initial faster rise of Chl *a* fluorescence (from the minimum “O” level to the intermediate “I” level) in HCO_3^- -depleted conditions (in hindsight, a hint of an effect on the electron acceptor side); they suggested that HCO_3^- may stabilize the S_1 state in the dark, and, simply, bicarbonate is of critical importance in the initial photochemical process.

In collaboration with Gerald Babcock, then at the University of California, Berkeley, Stemler and Govindjee, in 1974 [23] presented the following findings: (1) bicarbonate decreases the probability of so-called “misses” in the system; (2) the turnover time of PSII is increased by $\text{CO}_2/\text{HCO}_3^-$ -depletion since the rate of dark relaxation of the S-states ($\text{S}_1' \rightarrow \text{S}_2$; $\text{S}_2' \rightarrow \text{S}_3$) is severely retarded in bicarbonate depleted broken maize chloroplasts (Fig. 4); in our current understanding, this may be either due to effects on the PSII acceptor or the donor or both sides; (3) the final O_2 -evolving reaction, after accumulation of four positive charges, is independent of bicarbonate; and (4) HCO_3^- has no effect on the dark deactivation of the higher oxidation states, S_2 and S_3 .

In 1975, Wydrzynski and Govindjee [24], as mentioned above, provided the first evidence that there was a clear effect of bicarbonate on the electron acceptor side of PSII: (1) absence of HCO_3^- led to a faster rise of Chl *a* fluorescence (reflecting reduction of Q_A to Q_A^-) in systems where the O_2 -evolving system was blocked (e.g., by Tris-washing) and artificial electron donors (e.g., NH_2OH , MnCl_2 , hydroquinone and even DPC) were added to replace water; (2) effect of increasing concentrations of the herbicide 3-(3,4-dichlorophenyl)-1,1-dimethylurea (DCMU; diuron), which blocks electron flow from Q_A^- to Q_B , mimics increasing $\text{CO}_2/\text{HCO}_3^-$ -depletion conditions (Fig. 5). These results leave no doubt about an effect of bicarbonate on the electron acceptor side of PSII. This, of course, does not mean that there is no bicarbonate effect on the donor side of PSII.

In 1976, Jursinic, also in Govindjee's lab, and in collaboration with Warden [103] demonstrated a major effect of bicarbonate on the electron acceptor side of PSII by using three separate and independent methods: EPR signal II “very fast”, corresponding to tyrosine Z radical, fast Chl *a* fluorescence yield changes; and delayed light emission; although no effects were observed on the electron donor side of PSII, a reversible inactivation of PSII RC activity was observed.

In 1976, Govindjee, in collaboration with Pulles, R. Govindjee, Van Gorkom and Duysens [104], discovered, using spinach chloroplasts, that HCO_3^- -depletion inhibits the re-oxidation of the reduced form of the secondary electron acceptor Q_B (Q_B^{2-}) by the PQ pool. Results on the effects of DCMU supported this conclusion. Flash-number dependent measurements on Chl *a* fluorescence yield established that in HCO_3^- -depleted samples, the “two-electron gate”, on the electron acceptor side of PSII, was non-functional (Fig. 6).

In 1977, Khanna et al. [105] performed the first, so-to-say, biochemical surgery of the bicarbonate effect, using artificial electron acceptors (silicomolybdate (SM), oxidized diaminodurene (DAD), and methyl viologen (MV)) and donors (DPC, reduced DAD), acting at specific sites, and the inhibitors (2,5-dibromo-3-methyl-6-isopropyl-*p*-benzoquinone (DBMIB) and DCMU) of electron flow, also at specific sites (see Fig. 7). Results were clear: (1) there was no bicarbonate effect on Photosystem I (PSI); (2) there was no bicarbonate effect on the water-oxidizing side of PSII; and (3) there was a definite inhibition of electron flow on the PSII acceptor side, in agreement with the work of Wydrzynski and Govindjee [24].

In 1977, in collaboration with Siggel and Renger, in Berlin, Khanna and Govindjee performed absorption spectroscopy to decipher the nature of electron carriers on the PSII acceptor side that were affected by $\text{CO}_2/\text{HCO}_3^-$ -depletion and the reversal after HCO_3^- re-addition [106]. The formation of Q_B^{2-} was reversibly slowed down, by a factor of 10–20 fold, from ~500 μs to ~8 ms. However, a much larger effect of $\text{CO}_2/\text{HCO}_3^-$ -depletion was in the slowing down of the reduction of PQ pool to ~100 ms; this was consistent with the measurements of Govindjee et al. [104], where the fluorescence decay after the 3rd and subsequent flashes was in the range of 150 ms; this was also reflected in the slowed reduction of oxidized P700. This is clearly the major bottleneck produced by bicarbonate depletion.

3.1.2.2. 1980s: further research at UIUC, and collaboration with other labs in Berlin and in Wako Shi (Japan).

In view of the fact that reduction of Q_B to PQH_2 requires protonation, it became obvious that bicarbonate must be playing a role through protonation, and, thus, in 1980, in collaboration with Junge's research group, in Berlin, Khanna et al. [107] measured the effect of HCO_3^- -depletion on the proton uptake and release, using pH indicator dyes neutral red (internal space) and bromo-cresol purple (external space); the results of HCO_3^- -depletion on protons were remarkable: not only was the release of protons into the internal space dramatically reduced, but there was no proton uptake by the PQ pool at the outer side of the membrane (Fig. 8). Whatever was the detailed mechanism, effects on protonation by bicarbonate depletion on PSII were firmly established.

In 1981, and in collaboration with the research group of Arntzen, and with Van Rensen, Khanna et al. [108] provided information suggesting that the binding of bicarbonate is on the same protein that binds the herbicide atrazine; further results suggested complete inactivation of a part of the total number of electron transport chains. These conclusions were based on: (1) a shift in the binding constant of atrazine in bicarbonate-depleted thylakoid membranes indicating decreased affinity of atrazine; (2) trypsin treatment, which modifies PSII at the level of Q_B , strongly diminished stimulation by bicarbonate addition to HCO_3^- -depleted thylakoids. These conclusions were confirmed by measurements on atrazine-resistant plants (Fig. 9).

In 1982, in collaboration with Van Rensen, Vermaas et al. [109] used the herbicide ioxynil, which is different from atrazine used earlier by Khanna et al. [108]; inhibition of electron transport by ioxynil increased at decreasing bicarbonate levels (Fig. 10). An interesting

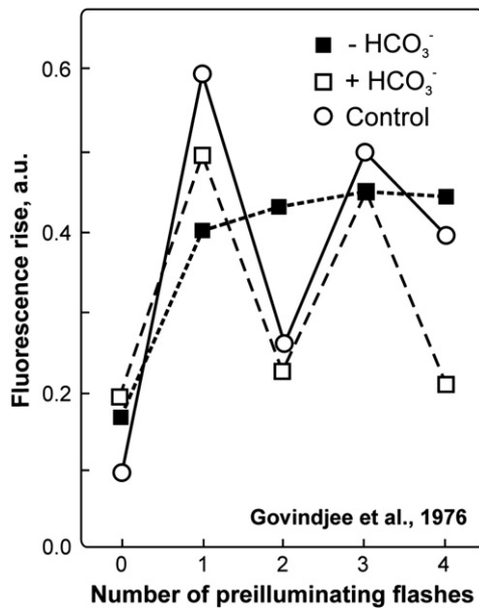


Fig. 6. Increase of DCMU-induced Chl *a* fluorescence as a function of flash number in HCO₃⁻-depleted (closed squares), HCO₃⁻-depleted plus 20 mM NaHCO₃ (open squares), and control (open circles) spinach chloroplast suspensions as measured by Govindjee et al. [104]. Other conditions: [Chl] = 20 μg ml⁻¹; [DCMU] = 5 μM. The measurements were done in the presence of 1 mM NH₂OH used as an artificial electron donor. Modified and adapted from [104].

528 conclusion of this study was that the binding sites of herbicide and
 529 bicarbonate, although similar, are not identical.
 530 • In 1984, Blubaugh and Govindjee [30] came to the conclusion that
 531 bicarbonate has 2 binding sites: (1) a high affinity binding site
 532 close to where DCMU binds; this binding is inhibited by light; and
 533 (2) a low affinity binding site, which requires light, and is where
 534 bathocuproine may bind, and, thus, this could be the one effect on
 535 the donor side of PSII. These results and conclusions need further
 536 investigations. Considering the high resolution structure of PSII
 537 [17] where only one bicarbonate site has been seen, all experiments

dealing with two bicarbonate binding sites need to be re-examined
 and proven by additional experiments using newer methodologies.
 • In 1984, Eaton-Rye and Govindjee [110] extended the conclusions
 of Khanna et al. [105], using MV as electron acceptor, and provided
 additional evidence that there was no effect of bicarbonate in PSI,
 and that there was a specific effect on the PSII electron acceptor
 side (reduction of PQ) that was not dependent on the use of formate
 to remove bicarbonate. They suggested “the observed large slow
 component in HCO₃⁻-depleted samples results from an altered
 equilibrium of Q_A⁻ with PQ and/or PQH₂ at the Q_B binding site”.
 This conclusion was consistent with their hypothesis that removal
 of HCO₃⁻ results in a retardation of the PQ/PQH₂ exchange reactions
 of the two-electron gate. Further, it has been suggested that this
 may be due to changes in the association constants for one or
 more of the PQ/PQH₂ species and/or by affecting the protonation
 reactions of the partially reduced plastoquinone anion or the
 doubly reduced plastoquinol; it was only after the first full turnover
 of the two-electron gate that the full effect of HCO₃⁻ depletion could
 be observed. This also explained the observation of Govindjee et al.
 [104] and Robinson et al. [111] that the decay of Chl *a* fluorescence
 after the 1st flash is less inhibited than after the 3rd and subsequent
 actinic flashes, but intermediate after the 2nd flash.
 • In 1984, Govindjee et al. [112], working in the laboratory of Inoue
 (Wako Shi) in Japan, confirmed, through thermoluminescence (TL)
 measurements that the bicarbonate depletion affected PSII on the
 electron acceptor side, in the Q_AQ_B region. They discovered (1) a 6–10 °C
 shift, to a higher temperature, in the S₂Q_B⁻ TL band; (2) a reduction
 in TL intensity upon prolonged depletion of bicarbonate; and (3) elimi-
 nation, after the first few flashes, of the characteristic period four oscil-
 lations in TL intensity as a function of the flash number. On the other
 hand, addition of DCMU produced the same S₂Q_A⁻ TL band, at about
 +20 °C in both depleted and reconstituted samples. These results sug-
 gest (1) the initial effect of CO₂/HCO₃⁻ depletion is to increase the acti-
 vation energy for S₂(S₃)Q_B⁻ recombination; (2) with further depletion,
 the incidence of this recombination decreases and the cycling of the
 S₂Q_B⁻ and S₃Q_B⁻ recombination is inhibited through effects at the Q_B
 apo-protein. These bicarbonate depletion effects were fully reversible
 if HCO₃⁻ was added to HCO₃⁻-depleted samples (i.e., reconstituted sam-

Some inhibitors, artificial electron donors and
 electron acceptors of photosynthetic electron transport

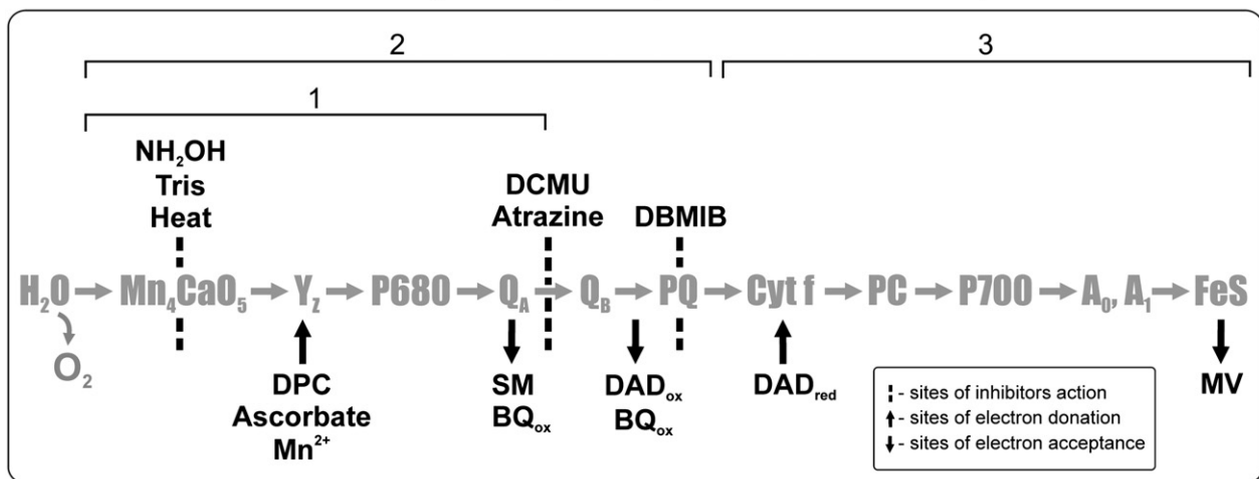


Fig. 7. Isolation of the photosynthetic electron transfer chain into several segments (1, 2, and 3) by using artificial electron donors and acceptors in combination with specific inhibitors of electron carriers. Abbreviations: DPC, diphenylcarbazine; SM, silicomolybdate; BQ, benzoquinone; DAD, diaminodurene; DBMIB, 2,5-dibromo-3-methyl-6-isopropyl-p-benzoquinone; DCMU, diuron (3-(3,4-dichlorophenyl)-1,1-dimethylurea); MV, methyl viologen; Cyt *f*, cytochrome *f*; PC, plastocyanin; P700, RC Chl of PSI; A₀, and A₁, primary electron acceptors of PSI; FeS, iron sulfur centers of PSI. Other abbreviations are as in Fig. 3. Modified from [105].

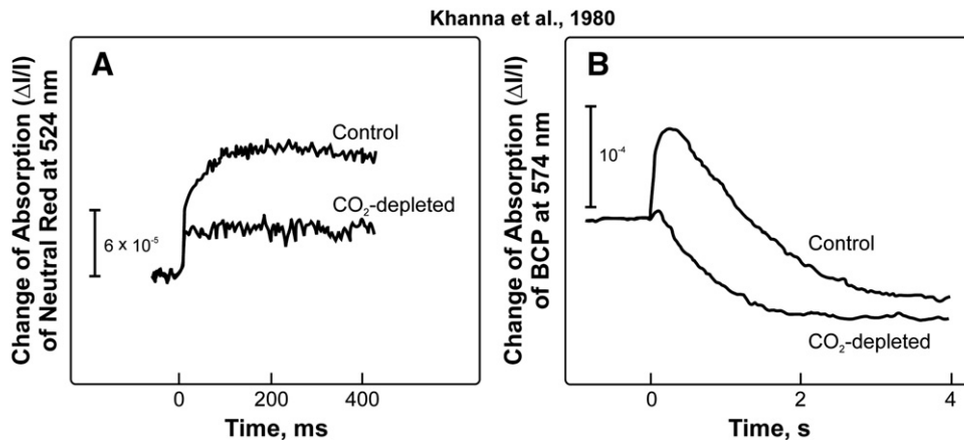


Fig. 8. Flash-induced kinetics of proton release as measured by Khanna et al. [107] in control and CO_2 -depleted spinach thylakoids. (A) Proton release kinetics monitored by absorption changes of neutral red (NR) at 524 nm. Signals represent a difference between two transient signals (signal obtained in the absence of imidazole minus signal obtained in the presence of imidazole). The assays were performed in a medium containing 20 mM KCl, 2 mM MgCl_2 , 0.5 mM $\text{K}_3[\text{Fe}(\text{CN})_6]$, 0.3 μM nonactin, 10 μM NR, and 1.3 mg ml^{-1} bovine serum albumin (BSA) at pH 7.0. (B) Proton release kinetics as indicated by absorption changes of bromocresol purple (BCP) at 574 nm. Reaction medium (pH 6.4) contained 20 mM KCl, 2 mM MgCl_2 , 0.5 mM $\text{K}_3[\text{Fe}(\text{CN})_6]$, and 20 μM BCP. In both cases the final concentration of Chl in reaction mixture was 10 $\mu\text{g ml}^{-1}$. For illumination, saturating single-turnover flashes ($\tau_{1/2} = 15 \mu\text{s}$) were used, and the obtained signals were averaged over 10 flashes. Dark time between flashes was 10 s. Modified and adapted from [107].

576 ples). A conformational change of the PSII complex in the region of the
577 Q_B apo-protein was suggested to be responsible for these effects.
578 • The CO_2 concentration in water solutions ($[\text{CO}_{2(\text{aq})}]$) is a function of
579 Henry's Law of solubility and the partial pressure of CO_2 (g) in the
580 air above the water (see Fig. 1). Concentrations of other inorganic
581 carbon species, i.e., HCO_3^- and CO_3^{2-} , vary with pH, and therefore,
582 the ratio $[\text{HCO}_3^-]/[\text{CO}_{2(\text{aq})}]$ is pH dependent (for details, see [113]).
583 The total concentration of dissolved inorganic carbon increases at
584 the pH range between 6 and 9 due to an increase in HCO_3^- species.
585 In 1986, Blubaugh and Govindjee [114], taking advantage of the pH
586 dependence of the ratio $[\text{HCO}_3^-]/[\text{CO}_2]$ at equilibrium to vary effectively
587 the concentration of one species while holding the other species
588 constant, discovered that the Hill reaction was stimulated in
589 direct proportion with the equilibrium $[\text{HCO}_3^-]$, but was independent
590 of the equilibrium $[\text{CO}_2]$ (Fig. 11). Thus, they suggested that
591 HCO_3^- is the species, which binds to the effector site, while CO_2 is
592 the diffusing species [115].

593 • In 1988, Blubaugh and Govindjee [116], using kinetic analysis of rates
594 of electron flow versus $[\text{HCO}_3^-]$, came to the conclusion that there are
595 two high affinity bicarbonate binding sites, apparently with cooperative
596 binding. We now ask where is the second bicarbonate binding
597 site, if it really exists? On the PSII electron donor side? Or at another
598 site on the electron acceptor side? As mentioned above, since in a re-
599 cent high-resolution PSII structure there is no indication for two
600 HCO_3^- molecules [17] (also see Section 3.2) the two binding site
601 concept needs to be re-examined with new experimental approaches.
602 Another conclusion was that bicarbonate is an essential activator for
603 PSII and that complete removal of HCO_3^- would result in zero electron
604 transport activity [116].
605 • In 1988, Eaton-Rye and Govindjee [117,118] provided a detailed
606 study of flash number dependent analysis of Chl *a* fluorescence
607 decay in spinach thylakoids at different pH values. The concept

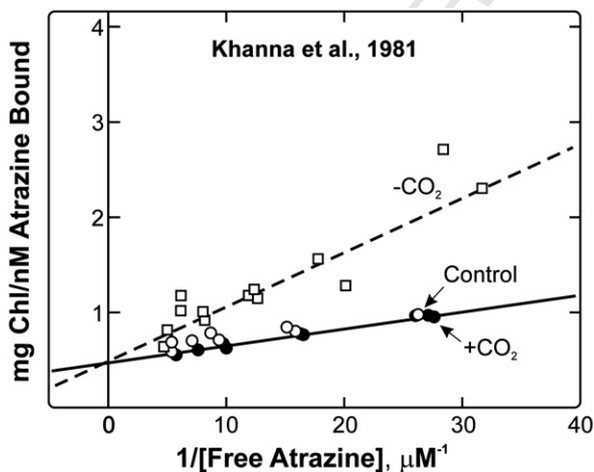


Fig. 9. ^{14}C -labeled atrazine binding to CO_2 -depleted (open squares), CO_2 -depleted plus 20 mM NaHCO_3 (closed circles), and control (open circles) pea membrane thylakoids as reported by Khanna et al. [108]. The data were represented as plots of double reciprocal (mg Chl/nM bound atrazine) vs. $1/[\text{free atrazine}]$. Thylakoids were incubated at 23 $^\circ\text{C}$ with various concentrations of ^{14}C -labeled atrazine. The amount of bound atrazine was calculated from the difference between the total radioactivity added to the thylakoids and the amount of free atrazine found in the supernatant after centrifugation. Modified and adapted from [108].

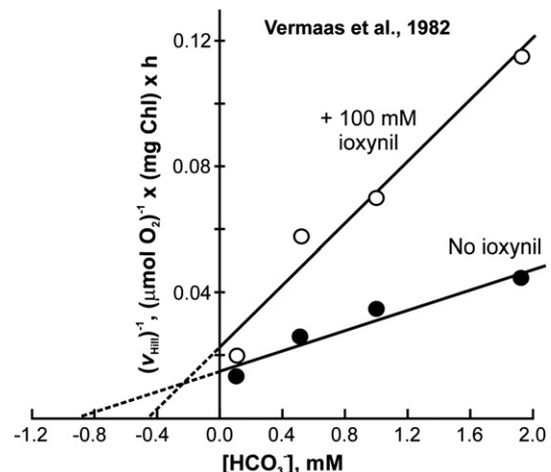


Fig. 10. Double reciprocal plot of the ferricyanide Hill reaction rate (v_{Hill}) as a function of the bicarbonate concentration in the absence (closed circles) and presence (open circles) of 100 nM ioxynil in pea thylakoids as reported by Vermaas et al. [109]. The samples were incubated with bicarbonate for 2 min. The measurements of O_2 evolution rates were done in the presence of 0.5 mM $\text{K}_3[\text{Fe}(\text{CN})_6]$. Ioxynil was added 3.5 min prior to the measurements. Adapted and modified from [109].

608 that bicarbonate was involved in protonation was fully supported: a
609 model of bicarbonate acting as a proton donor to the protein disso-
610 ciable group believed to participate in the protonation of reduced
611 Q_B was discussed, as well as the possibility of HCO_3^- being a ligand
612 to the NHI in the Q_A-Fe-Q_B complex of the PSII RC. In addition, in
613 **1988**, (1) Cao and Govindjee [119] reported a bicarbonate effect in
614 a cyanobacterium *Synechocystis* sp. PCC 6803; and (2) Garab and
615 collaborators [120] provided evidence through TL measurements
616 that CO_2 does affect charge accumulation in intact leaves.

617 • In **1989**, (1) in collaboration with the Crofts' lab, Govindjee et al.
618 [121], using fast fluorescence changes, failed to observe any signifi-
619 cant effect of bicarbonate on electron donation from tyrosine Z (Y_Z)
620 to P680 or in the formation of $P680Q_A^-$; and (2) in collaboration
621 with the lab of Colin Wraight [122], a total absence of CO_2/HCO_3^- -
622 depletion effect was observed between the quinones both in chro-
623 matophores and RCs in the purple bacterium *Rhodobacter* (*R.*)
624 *sphaeroides*. This was followed, in **1992**, by the work of Wang (in
625 Wraight's Lab at Urbana, IL) and Cao (in Govindjee's lab) who, in collab-
626 oration with Oesterhelts' lab in Munich [123] asked if bicarbonate in
627 PSII is equivalent of Glu (M234 in *R. sphaeroides*) in bacterial RCs in
628 binding to the NHI. Michel and Deisenhofer [124] had earlier suggested
629 this notion. None of the mutants of M-234, where Glu was changed to
630 Val, Gln or Gly, showed any difference in the HCO_3^- -reversible formate
631 effect, confirming the absence of bicarbonate effect in these anoxygenic
632 photosynthetic bacteria.

633 3.1.2.3. 1990s: continued research at UIUC, and collaboration with other
634 labs in the USA and in labs around the World (Canada, China, Israel, Finland,
635 Switzerland, France, Germany, and The Netherlands).

636 • Following the lead of Khanna et al. [108] and Vermaas et al. [109] that
637 had suggested an overlap of binding sites of bicarbonate and herbicides
638 in higher plants, Govindjee, working in collaboration with Vernotte,
639 Peteri, Astier and Etienne, found, in **1990** [125] that the herbicide-
640 resistant mutants of the cyanobacterium *Synechocystis* sp. PCC 6714,
641 that are altered in specific amino acids in their D1 protein, show differ-
642 ential sensitivity to formate treatment. Yield of O_2 in a sequence of
643 flashes, Chl *a* fluorescence transients and Chl *a* fluorescence yield
644 decay after a flash revealed that the resistance of cells to formate treat-
645 ment was in the following (highest to lowest) order: [double
646 D1-mutant] A251V/F211S > [single D1-mutant] F211S > wild type >

[single D1-mutant] S264A. These results established the involvement
647 of the D1 protein in bicarbonate/formate binding, but gave no further
648 clue to the precise site of binding. From the PSII crystal structure [17],
649 these residues are rather close to Q_B ; changes in these residues may per-
650 turb the proper binding of Q_B , giving rise to indirect effects on the bind-
651 ing of bicarbonate/formate. 652

653 • In **1991**, using membrane-inlet mass spectrometry (MIMS) and a in-
654 frared gas analyzer, Govindjee in collaboration with Weger, Turpin,
655 Van Rensen, Devos and Snel, [53] showed that formate replaces
656 HCO_3^- from its binding site in PSII (see Fig. 12 and legend for experi-
657 mental details). Addition of 100 mM formate to spinach thylakoids re-
658 leased from $\sim 0.4 HCO_3^-/CO_2$ to $1.3 HCO_3^-/CO_2$, confirms the earlier idea
659 [12,125] that the bicarbonate effect occurs through the binding of
660 HCO_3^- to PSII, and that the addition of formate removes HCO_3^-/CO_2
661 from its binding site, leading to inhibition of electron flow. This did
662 not support the experiments and conclusions of Alan Stemler [126]. It
663 appears that about 1 HCO_3^- (at pH 6.5) is released by formate addition.
664 Further, in 1995, Oscar et al. [127] established the "bound-bicarbonate"
665 rather than the "inhibitory anion or the empty site" hypothesis of
666 Jursinic and Stemler [128] by showing CO_2 release under their experi-
667 mental conditions. 667

668 • Further evidence that the D1 protein was involved in the HCO_3^- effect
669 on PSII was obtained, in **1991**, by Govindjee et al. [129], using a
670 D1-L275F strain and several other mutants of *C. reinhardtii*, in collab-
671 oration with labs at University of Geneva, Switzerland. The L275F mu-
672 tant failed to show the HCO_3^- -reversible formate effect suggesting to
673 the authors that a significant change in formate (bicarbonate) binding
674 had occurred in helix V of the D1 protein near His involved in NHI
675 binding. Further, with the exception of the S264A mutant, which is
676 considerably more sensitive to formate than the wild type, five
677 other different [V219I, A251V, F255Y, G256D and cell-wall deficient
678 CW-15] mutants displayed a relatively similar response to formate
679 as wild type. Absence of a formate effect on a PSII-lacking mutant
680 seemed to confirm the sole involvement of PSII in the 'bicarbonate ef-
681 fect'. These results suggested that specific areas of the D1-protein are
682 more important than the others in formate/bicarbonate binding, but
683 they did not give precise clues. Lack of effect may not only be due to
684 the geometric organization of the structure, but may also be due to a
685 replacement with similar residues. The search continued. 685

686 • In **1991**, Xu and Govindjee [130], in collaboration with the laboratory
687 of Tony Crofts, presented a detailed kinetic investigation on spinach
688 thylakoids, as well as a model of HCO_3^- -reversible formate/formic 688

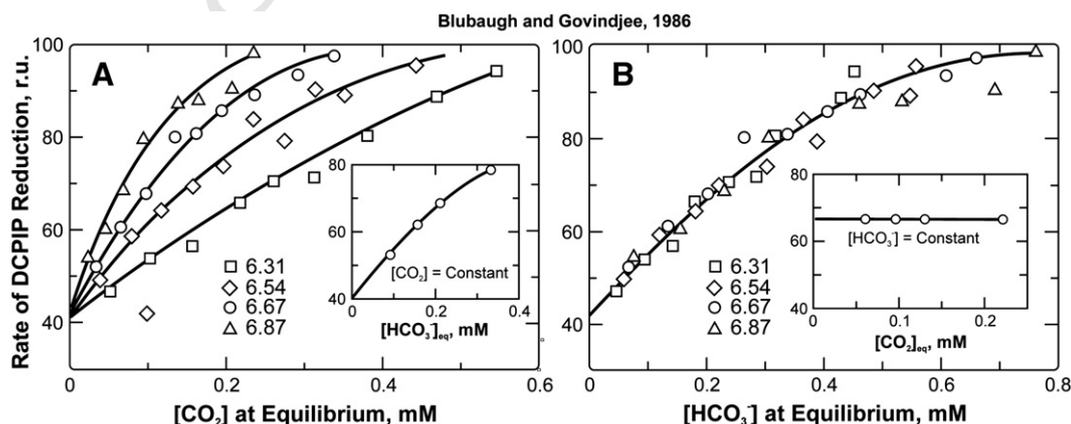


Fig. 11. The rate of 2,6-dichlorophenolindophenol (DCPIP) reduction measured by Blubaugh and Govindjee [114] in CO_2 -depleted thylakoids as a function of the equilibrium CO_2 (A) and HCO_3^- (B) concentrations. The reduction rate of DCPIP was calculated from the decrease in absorbance at 600 nm and normalized to the control rate. The control rates (in μmol (DCPIP^{red}) mg (Chl)⁻¹ h^{-1}), estimated separately for each curve (pH value) by adding 2.5 mM HCO_3^- to the CO_2 -depleted samples, were the following: 209 at pH 6.31 (open squares); 212 at pH 6.54 (open diamonds); 191 at pH 6.67 (open circles); and 192 at pH 6.87 (open triangles). $NaHCO_3$ was added 3 min prior to illumination. Inset in (A): the effect of the equilibrium $[HCO_3^-]$ on the Hill reaction, with the $[CO_2]$ held constant at 0.1 mM. Inset in (B): the effect of the equilibrium $[CO_2]$ on the Hill reaction, with the $[HCO_3^-]$ held constant at 0.2 mM. Modified and adapted from [114].

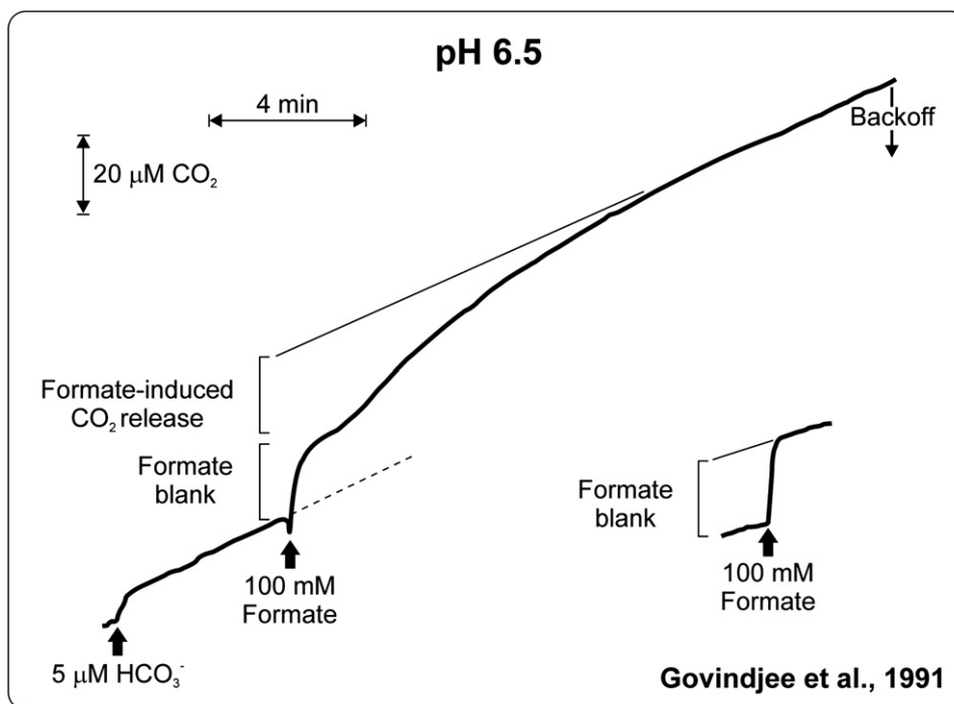


Fig. 12. First detection of formate-induced release of CO₂ from spinach thylakoids as measured by MIMS by Govindjee et al. [53]. The addition of formate (to 100 mM) induced a rapid increase in the CO₂ signal (which corresponded to a formate injection artifact, i.e. formate blank) followed by a slow release of CO₂ from thylakoids (left). Repetitive addition of formate to the same sample induced only the initial rapid CO₂ release related to artifact of formate injection (right). CO₂ was continuously monitored at $m/z = 44$. The measurements were performed at 20 °C and pH 6.5 in the presence of external carbonic anhydrase (final concentration of 0.5 μg/ml) in order to facilitate the equilibration between inorganic carbon species.

Modified and adapted from [53].

acid effect. In agreement with earlier reports [104,118], electron flow from Q_A^- to Q_B^- was shown to be slowed down, and the notion that CO₂/HCO₃⁻-depletion leads to a loss of protonation necessary for stabilization of Q_B^- became the dominant mechanism. However, their conclusion that it is formic acid, not formate, which binds to the acceptor side of PSII has not been pursued since then, and it remains to be further investigated and deserves additional studies.

- In their detailed review in 1988, Blubaugh and Govindjee [12] had presented models for HCO₃⁻ action in PSII, and had also suggested that positively charged Arg residues may be involved in bicarbonate binding. In order to test this idea, Cao et al. [131], in 1991, made D2-R233Q and D2-R251S mutants in the cyanobacterium *Synechocystis* sp. PCC 6803, and, based on both O₂ evolution and Chl *a* fluorescence measurements, suggested that these two Arg residues must be somehow involved in bicarbonate binding in PSII. In the current high-resolution PSII structure [17], these two residues are located on the stromal surface 15–16 Å away from bicarbonate, and thus their effect may be indirect, through effects on the hydrogen-bonding network linking the bicarbonate to the stromal surface.
- In 1992, in collaboration with Pfister and Strasser's research group, Govindjee et al. [132] extended the earlier work on several herbicide resistant D1-C. *reinhardtii* mutants [129] and concluded that D1-S264, but not D1-L275, D1-F255 and D1-V219, plays an important role in the functioning of HCO₃⁻ and PQ in PSII; the role of D1-G256 could not be determined in this study. (See also [133] for the role of D1-S264 and the absence of the role of D1-L275.) The high-resolution structure of PSII [17] now shows that D1-S264 is hydrogen-bonded to Q_B^- (see Section 3.2).
- The role of various D1 amino acids in the HCO₃⁻ effect, this time, by using herbicide-resistant mutants of the cyanobacterium *Synechococcus*

- sp. PCC 7942, was examined by Cao and Xiong, in 1992, in collaboration with Hirschberg and Ohad, in Israel [134]. Interestingly, the hierarchy of the equilibrium dissociation constant for bicarbonate (highest to lowest) was D1-F255L/S264A > D1-F255Y/S264A ~ D1-S264 ~ D1-F255Y > Wild type, establishing the importance of D1-S264 and D-F255 in the HCO₃⁻ binding niche directly or indirectly. Again, a role of bicarbonate in protonation and stabilization of Q_B^- was emphasized, a recurring concept since the earlier observations [107].
- By 1993, the following conclusions were made [135]: (1) formate, azide, nitrite and nitric oxide inhibited electron flow in thylakoids and cells, and these effects were significantly and uniquely reversed by bicarbonate; (2) with formate treatment, a remarkably strong HCO₃⁻-reversible slowing down of Q_A^- reoxidation after the second and subsequent flashes, but not after the first flash, was observed; (3) a hypothesis was in place suggesting that bicarbonate functions as a proton shuttle stabilizing the binding niche of Q_B^- and stimulating PQH₂ formation (and, perhaps, even its oxidation) in some manner; (4) this effect somehow involves both D1 and D2, directly, or indirectly, particularly the region where herbicides bind, and part of this was based on several mutant studies (e.g., D1-S264A, D1-L275F, D2-R251S, D2-R233Q, D2-R139H, among others); (5) possible involvement of "Fe" in the " Q_A -Fe- Q_B " complex was also implicated; (6) this effect was unique to PSII since electron transport in the " Q_A -Fe- Q_B " complex of both green and purple bacteria (including M-E234G, Q and V mutants) was insensitive to HCO₃⁻-reversible inhibitors.
- In 1995, Mäenpää et al. [136] made an interesting observation in *Synechocystis* sp. PCC 6803: HCO₃⁻-reversible formate effect on $Q_A Q_B$ was several fold less in the CA1 mutant (that had Glu 242, Glu 243, and Glu 244 deleted, and where Gln 241 was changed to His; these changes being in the de-loop of the D1 protein). These

results may be related to differences in the accessibility of the anions and/or due to changes in the redox properties of Q_A/Q_A^- in the mutant—perhaps, an indirect effect.

- In **1996**, in collaboration with the research group of Sayre, attempts were made to test the importance of D1-R269 in *C. reinhardtii* [137]; it was difficult to obtain firm conclusions since the used D1-R269G mutant was unable to grow photosynthetically and to evolve O_2 : it had many defects. In the current high-resolution PSII structure [17], D1-R269 is hydrogen-bonded to D2-T243, which is probably needed to maintain the proper orientation of D2-Y244 in order for it to be able to hydrogen-bond to the bicarbonate (see Section 3.2).
- In **1996**, Xiong et al. [138] presented a 3-dimensional model of the D1/D2 protein and the cofactors, using the bacterial RCs, and predicted the HCO_3^- binding niche in PSII; it was modeled in the NHI site, providing a bidentate ligand to the iron. In their model, a bicarbonate ion that was suggested to be stabilized by D1-R257, was said to donate a proton to Q_B^2- through the D1-H252 residue, whereas a water molecule was proposed to donate another proton to Q_B^2- ; Xiong et al. also proposed a positively charged water channel, near Q_B and the NHI, for transporting water and HCO_3^- . It is now indeed known [17] that D1-H252 is hydrogen-bonded to Q_B through D1-S264, and there are water molecules close to D1-H252 that could serve as proton donor to Q_B (see Section 3.2). However, D1-R257 is at a distance of 8.6 Å from Q_B and further away from the bicarbonate; thus, it cannot be directly involved in this hydrogen-bond network.
- In **1997**, Govindjee et al. [7] presented data on Chl *a* fluorescence yield changes after light flashes 1–6 in spinach thylakoids at pH 6.0; they showed a bicarbonate effect on both the electron donor and electron acceptor sides in the same samples. The donor side effect was shown by a decrease in maximum fluorescence, and the acceptor side effect by a slowing down of the fluorescence decay due to Q_A^- oxidation. Using a sensitive differential infra-red gas analyzer they showed the presence of 0.8–1.25 bicarbonate ions bound per PSII RC in maize and pea thylakoids. These results were in agreement with earlier published data obtained by time-resolved MIMS on spinach thylakoids [53] (Fig. 12). Govindjee et al. [7] suggested that bicarbonate bound to the acceptor side is required for PSII activity, both on the acceptor and the donor sides in the same experiment and in the same sample; in this hypothesis, conformational changes may need to be invoked.

3.1.2.4. 2000s: new conclusions, collaboration with research group of Tony Crofts.

- In **2008**, Rose et al. [139], using both Chl *a* fluorescence, and TL measurements, provided the following conclusions on the D1-R257 mutation (D1-R257E, D1-R257M, and D1-257K): although the forward rate of electron transfer from Q_A to Q_B was little affected, the two-electron gate on the acceptor side of PSII was thermodynamically perturbed in the R257 mutants; this led to a decrease in the overall electron transfer rate from water to PQ. The effects on equilibrium constants of the two-electron gate are likely due to changes in coulombic fields on changing the net charge in the neighborhood of the Q_B site, suggesting that the electrostatic environment plays an important role in the mechanism of PSII. The bicarbonate-reversible formate effect on the Q_B site had been shown to be on the protonation events at this site [117,118]. Dramatic differences of the bicarbonate effect on the D1-R257 mutants, observed earlier [140], might thus have a basis in changes in the redox potential and the stability of the Q_B site, observed in this research. It, thus, seems that although D1-R257 is not close to the binding site of HCO_3^- on the NHI, it has a significant effect on the PSII reactions in the Q_B -region.
- As the model for the role of HCO_3^- had been evolving, it was generally thought that the first proton for the stabilization of Q_B^- came from D1-H252, and, thus, removal of HCO_3^- did not exhibit its major effect on the electron transport from the reduced Q_A to Q_B ,

but it had a large effect on the electron flow from the reduced Q_A to Q_B^- , and the succeeding reactions; the idea that bicarbonate provides this second proton, becoming carbonate, is the current picture. Carbonate, in turn, picks up a proton from D1-E244, finishing the cycle. The HCO_3^- ions (or water protons) outside the PSII complex provide the missing protons to the Glu (see current model in Section 3.2). The idea of involvement of D1-H252 in the first protonation was discussed by Petrouleas and Crofts [141], based on the experiments of Padden (see [142]; and paper in preparation). We note that depending upon the severity of HCO_3^- depletion procedure, an inhibition after the 1st flash is also observed explaining effects on TL band due to $S_2Q_B^-$ recombination (see earlier discussion).

3.1.3. Work around the World related to the site of bicarbonate binding

Research summarized below focuses on the studies related to key observations of bicarbonate binding to the electron acceptor side of PSII, mainly on the Q_A -NHI- Q_B niche and the PQ pool.

3.1.3.1. The 1980s.

- In **1984**, Vermaas and Rutherford [143] were among the first ones to focus on the relationship of bicarbonate to the Q_A -NHI- Q_B niche of PSII. They discovered that removal of HCO_3^-/CO_2 , in PSII membrane fragments from *Brassica napus*, led to a very large increase in the EPR signal at $g = 1.82$ that is due to the $Q_A^-Fe^{2+}$ complex, and, that this effect was fully reversible when bicarbonate was added back. This result identified bicarbonate to be either located near this complex, or, to play a crucial role in affecting the conformation of the Q_AFe complex.
- In **1987**, Diner and Petrouleas [144] showed reversible decrease in the quadrupole splitting of the NHI Mossbauer spectra, upon bicarbonate depletion. This confirmed the concept of bicarbonate acting on the electron acceptor side of PSII.
- In **1988**, Nugent et al. [145], using EPR measurements on both NHI ($g = 6$) and $Q_A^-Fe^{3+}$ ($g = 1.82$) in PSII particles, from both the thermophilic cyanobacterium *Phormidium laminosum* (Fig. 13A) and *Spinacea oleracea* (spinach) (Fig. 13B), suggested that bicarbonate binds close to the NHI and affects Q_A , Q_B as well as the NHI. Further, they found that the NHI was oxidized only when bicarbonate was present (also see [146]). These results supported the conclusions of Govindjee and coworkers (see Sections 3.1.2.1 and 3.1.2.2) that bicarbonate plays a central role in providing conditions for efficient electron flow on the acceptor side of PSII [11,147].
- In **1988**, Michel and Deisenhofer [124] in their perspective in the journal *Biochemistry* wrote "Having in mind the well-known effects of bicarbonate at the electron-accepting site of PSII, we consider bicarbonate as a likely candidate to be the fifth iron ligand in D1 and D2". They suggested that bicarbonate occupies the place of M-E232 of anoxygenic bacterial RC.

3.1.3.2. The 1990s.

- In **1990**, Diner and Petrouleas [25], using NO, instead of formate, to remove CO_2/HCO_3^- , showed that $g = 4$ EPR signal of Fe^{2+} -NO was diminished when bicarbonate was added, favoring the concept that HCO_3^- is a ligand to the NHI.
- In **1991**, Diner et al. [148] presented a detailed overview on the iron-quinone electron acceptor complex of PSII. Here, they reviewed the literature on the bicarbonate effect in PSII and discussed various models for the binding and functioning of bicarbonate at the Q_A -NHI- Q_B complex.
- In **1995**, Hienerwadel and Berthomieu [149] provided the first IR spectroscopy evidence for bicarbonate binding on the acceptor side of PSII, using FT-IR difference spectroscopy, and ^{13}C -labeled HCO_3^- . Binding of bicarbonate to the NHI was strongly supported

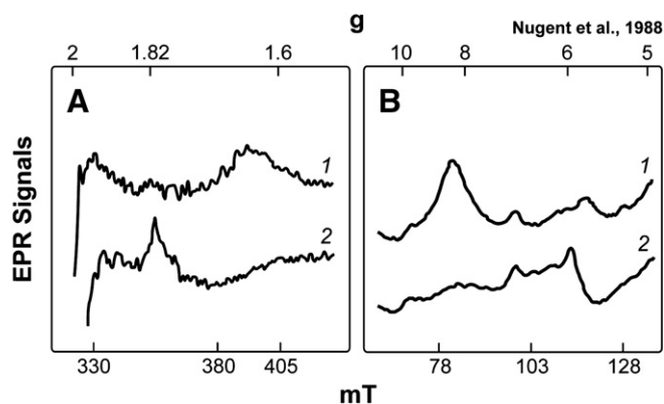


Fig. 13. EPR data showing the effect of HCO_3^- removal on the acceptor side of PSII as reported by Nugent et al. [145]. (A) EPR spectra of the Q_A -NHI region ($g=1.82$) in dark-adapted PSII particles from the cyanobacterium *Phormidium laminosum* upon 5 min illumination at 77 K in the absence (spectrum 1) and the presence of 100 mM formate (spectrum 2). (B) EPR spectra of the NHI (Fe^{3+}) region ($g=6$) of the dark-adapted PSII particles from spinach in the absence (spectrum 1) and the presence (spectrum 2) of 100 mM formate. For further details see [145]. Modified and reproduced from [145].

875 by this study; it was suggested that bicarbonate is a monodentate
876 ligand of the oxidized iron, but a bidentate ligand of the reduced
877 form of iron, and exhibits hydrogen bonds with the protein.

878 3.1.3.3. The 2000s.

879

- 880 • In **2001**, Berthomieu and Hienerwadel [150] looked for the specific
881 interactions of bicarbonate with the protein; here, they used lactate,
882 glycolate and glyoxylate, instead of formate or NO, to remove inorganic
883 carbon. Further, these authors concluded, from their studies,
884 that one proton is released upon iron oxidation, and suggested
885 that pH dependence of the iron couple may reflect deprotonation
886 of D1-H215, a “putative” iron ligand located at the “ Q_B ” pocket.
887 (This proton release was suggested to have a different mechanism
888 from that involved in the functioning of bicarbonate.) They con-
889 cluded that a ‘hydrogen network’ exists from the NHI towards the
890 “ Q_B ” pocket involving bicarbonate and D1-H215 (see current
891 model in Section 3.2).
- 892 • In **2008**, in search for proof (or absence of proof) for the binding of
893 HCO_3^- to the electron donor side of PSII, Shevela et al. [41,151] re-
894 examined and extended the MIMS experiments reported earlier
895 by Govindjee et al. [53] and Stemler [126]. Govindjee et al. [53]
896 had presented clear evidence for the release of $\text{CO}_2/\text{HCO}_3^-$ induced
897 by formate addition (Fig. 12); however, the binding site for this
898 anion was not specified in this study. Based on the previous experi-
899 mental data, indicating the binding of HCO_3^- to the NHI at the ac-
900 ceptor side, it was assumed that formate removes HCO_3^- from this
901 binding site. Formate, however, was reported to bind both at the ac-
902 ceptor and donor sides of PSII [152]. It was, therefore, unclear, from
903 which binding side(s) in PSII the released CO_2 had originated in the
904 previous study [53]. In the MIMS study of Shevela et al. some experi-
905 ments were performed with an H_2^{18}O enrichment, which allowed the
906 detection of CO_2 isotopologues at $m/z=46$ ($\text{C}^{16}\text{O}^{18}\text{O}$), and $m/z=48$
907 (C^{18}O_2). Since the Faraday cups used for the detection of $\text{C}^{16}\text{O}^{18}\text{O}$ and
908 C^{18}O_2 were amplified by 10 and 100, respectively, than the one used
909 for the detection of non-labeled CO_2 ($m/z=44$), the ^{18}O -enrichment
910 greatly increased the sensitivity of the MS instrument (compare signal
911 amplitudes in Fig. 14A, B, and C). The results obtained not only fully con-
912 firmed the formate-induced release of $\text{CO}_2/\text{HCO}_3^-$ reported earlier by
913 Govindjee et al. [53] (Fig. 12), but also clearly demonstrated that the re-
914 leased $\text{HCO}_3^-/\text{CO}_2$ originates *only* from the acceptor side, and *not* from
915 the donor side of PSII (for experimental details see Fig. 14 and its

legend). We also note here that, in the same year (**2008**), evidence for
916 the binding of HCO_3^- on the electron acceptor side of PSII and the ab-
917 sence of bicarbonate bound to the donor side was presented in a FT-IR
918 spectroscopy study by Aoyama et al. [42] and in a GC-MS study by
919 Ulas et al. [55]. Thus, the focus of action on the mechanism remained
920 on the Q_A -NHI- Q_B complex.

- 921 • In **2009**, Cox et al. [153] continued EPR studies on the Q_A -NHI- Q_B
922 complex of PSII, initiated in the 1980s and 1990s (see above), but
923 they added DFT calculations. They looked at the native $g\sim 1.9$ form
924 as well as the $g\sim 1.84$ form, which is the well known signal in purple
925 bacterial RCs (where bicarbonate does not bind, see Section 3.1.2.2)
926 and that is occurring in PSII when they are treated with formate
927 that removes $\text{CO}_2/\text{HCO}_3^-$. The calculations led Cox et al. to conclude
928 that the doubly charged carbonate ion (CO_3^{2-}) is responsible for the
929 $g\sim 1.9$ form of the semiquinone-iron signal; and carbonate, rather
930 than bicarbonate (HCO_3^-), is the ligand to the NHI; the latter is in
931 apparent contradiction to what we believe was the conclusion of
932 Berthomieu and Hienerwadel (see above). It is highly likely that
933 both bicarbonate and carbonate can bind to the NHI depending
934 upon the precise physical and chemical status of the system since
935 carbonate is formed from bicarbonate when the latter would be do-
936 nating a proton to stabilize $\text{Q}_\text{B}^{\cdot -}$ (see Section 3.2)
- 937 • In **2009**, Takahashi et al. [154] dug deeply into the question of HCO_3^-
938 binding at the NHI in PSII using FT-IR, as Berthomieu and Hienerwa-
939 del [150] had done, and included DFT calculations as well. Their
940 study included specific ^{13}C -Tyr labeling together with a deuteration
941 effect to provide evidence from Tyr IR modes to indicate Tyr involve-
942 ment in hydrogen bonding to bicarbonate. The results obtained indi-
943 cated that a Tyr (either D1-Y246 or D2-Y244; see Section 3.2) side
944 chain in “a hydrogen bond donor-acceptor form” is strongly coupled
945 to the NHI; this was suggested to provide a hydrogen bond to the ox-
946 ygen of the bicarbonate ligand. Thus, Takahashi et al. were the first to
947 propose that a key “Tyr residue coupled to the NHI may play a key role
948 in the regulatory function of the iron-bicarbonate center by stabilizing
949 the bicarbonate ligand and forming a rigid hydrogen bond network
950 around the NHI.”
- 951 • In **2011**, Sedoud et al. [155] provided a thorough study on the ef-
952 fects of formate binding on the EPR of the quinone-NHI electron ac-
953 ceptor complex using light flash experiments and reached the
954 conclusion that the effect was maximum after the 3rd flash indicat-
955 ing that the major effect of formate treatment ($\text{HCO}_3^-/\text{CO}_2$ removal)
956 is on the $\text{Q}_\text{B}\text{H}_2$ exchange. This conclusion is in agreement with the
957 earlier results of flash number dependence on Chl *a* fluorescence
958 observed by Govindjee et al. [104] and on absorption changes by
959 PQ, as measured by Siggel et al. [106]. However, this does not pre-
960 clude, at all, the participation of bicarbonate in the protonation of
961 $\text{Q}_\text{B}^{\cdot -}$. An integrated model would include both effects although the
962 bottleneck reaction that would control the net electron flow may
963 very well be this exchange reaction that would lead to slower oxida-
964 tion of PQH_2 .
- 965 • In **2011**, Chernev et al. [156] investigated the NHI-(bi)carbonate com-
966 plex using μs -resolution X-ray absorption spectroscopy (XAS) after
967 laser flash excitation of PSII membrane particles. An interpretation
968 of the observed spectral changes revealed that the coordination of bi-
969 carbonate at the Fe^{2+} may change from a bidentate to a monodentate
970 ligation (carboxylate shift) after the formation of Q_A^- . Based on the
971 obtained data and DFT calculations as well as on previous XAS experi-
972 ments showing that no $\text{Fe}^{2+} \rightarrow \text{Fe}^{3+}$ transition occurs during the
973 electron transfer from Q_A to Q_B in the type II photosynthetic RCs
974 [157], Chernev et al. proposed that a coordination flexibility of the li-
975 gand (bicarbonate in PSII and glutamate in bacterial RCs) is essential
976 for the functioning of the NHI-carboxyl complex in the interquinone
977 electron transfer.
- 978 • In **2011**, Müh et al. [158] have beautifully reviewed PQ reduction in
979 PSII. They suggest that one water molecule is there in the PSII struc-
980 ture that interacts with D1-H252, and two water molecules bridge
981

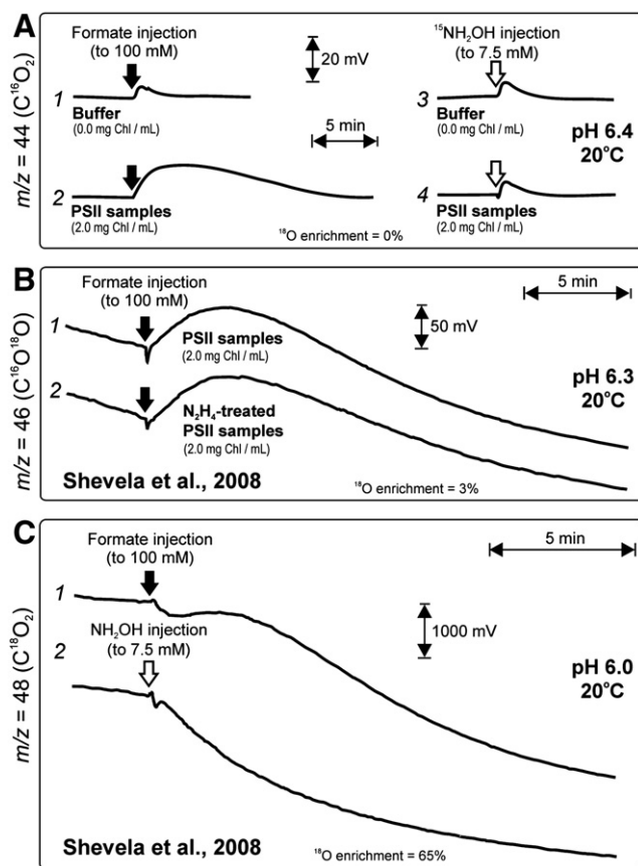


Fig. 14. Probing for binding sites of $\text{HCO}_3^-/\text{CO}_3^{2-}$ in PSII by isotope ratio MIMS in spinach PSII membrane fragments. These data confirmed the formate-induced release of CO_2 reported earlier by Govindjee et al. (see Fig. 12) and demonstrated that all released $\text{CO}_2/\text{HCO}_3^-$ originates from the electron acceptor and none from the donor side of PSII. (A) The addition of formate (to final concentration of 100 mM; black arrows) to PSII membranes at pH 6.3 and 20 °C induced a slow release of CO_2 (detected at $m/z = 44$) which was much above the artifact caused by injection of formate into the buffer with no samples (compare traces 1 and 2). Destruction of the possible binding site (the Mn_4CaO_5 cluster) via the addition of strong reductant NH_2OH (to final concentration of 7.5 mM; open arrows) does not lead to a release of $\text{CO}_2/\text{HCO}_3^-$ (compare traces 3 and 4). During the reduction NH_2OH is known to produce N_2O . In order to shift the signal of N_2O from $m/z = 44$ to $m/z = 46$, and thus to avoid possible overlay of the CO_2 and N_2O signals the ^{15}N -labeled NH_2OH was used for these experiments. (B) CO_2 release upon formate addition (to 100 mM) to 'control' PSII membranes (trace 1) is the same as in the case of PSII membranes without the Mn_4CaO_5 cluster (due to pre-incubation with 80 mM NH_2NH_2 for 75 min) (trace 2). CO_2 was detected at pH 6.3 and 20 °C as $^{16}\text{O}^{18}\text{O}$ at $m/z = 46$ due to ^{18}O -enrichment with H_2^{18}O (3%). (C) Formate-induced release of CO_2 (trace 1) compared with the absence of CO_2 release upon injection of NH_2OH (trace 2) as detected at $m/z = 48$ at pH 6.0 and 20 °C. To get the highest possible sensitivity the experiments were performed with high ^{18}O -enrichment level (~65%). All measurements were done in the presence of externally added carbonic anhydrase (to a final concentration of 3 $\mu\text{g}/\text{ml}$) to facilitate equilibration between CO_2 and HCO_3^- and by this to allow the detection of all dissolved inorganic carbon as CO_2 along. Modified and adapted from [41].

Shen reported, in 2003 [161], the PSII structure from *T. vulcanus* at a 3.7 Å resolution. These structures did not allow the assignment of bicarbonate in PSII, either on the donor, or the acceptor side. The first assignment of bicarbonate was reported by Ferreira et al., in 2004 [51], in their PSII structure from *T. elongatus* at 3.5 Å resolution, in which they assigned two HCO_3^- ions, one at the donor side and the other at the acceptor side. The bicarbonate at the donor side was assigned to be a direct ligand to the Mn_4CaO_5 cluster. The density that was assigned to a putative bicarbonate, however, was not found in the subsequent structures at higher resolutions of 2.9–3.0 Å [52,162]. In the most recent structure of PSII determined at a resolution of 1.9 Å [17], bicarbonate was also not found at the donor side. Since in this high resolution structure, all of the ligands for the 4 Mn ions and the Ca ion were determined, which showed that each of the Mn has 6 ligands and the Ca ion has 7 ligands, there is no room for the presence of a bicarbonate in the immediate ligand sphere of the Mn_4CaO_5 cluster, at least in the assembled, active PSII complex. It is also highly unlikely that a well-defined HCO_3^- could be missed in an electron density map with a resolution beyond 2.0 Å, as the electron density for the bicarbonate at the acceptor was clearly defined and visible [17]. One can assume that bicarbonate strongly bound to the Mn_4CaO_5 cluster might be lost due to reduction of high-valence Mn ions ($\text{Mn}^{\text{III}}\text{Mn}^{\text{IV}}_2$) to Mn(II), which is known to take place under X-ray doses used for structure determination by X-ray crystallography [163]. However, in view of recent MS and FT-IR data [41,42,55,151] showing the absence of tightly bound bicarbonate to the Mn_4CaO_5 cluster, this option can be excluded.

Fig. 15A shows the position of the bicarbonate on the acceptor side in a PSII monomer determined at 1.9 Å resolution [17]. While the global position of the bicarbonate could be assigned in the structures with a resolution in the range of 3.0–3.5 Å, its detailed environment including the presence and the positions of water molecules surrounding it has to be determined at a much higher resolution, which is now achieved at 1.9 Å. Based on this structure, the bicarbonate serves as a bidentate ligand to the NHI, which is located just under the surface of the stromal side of the membrane region. This bicarbonate is surrounded by hydrophilic residues and water molecules, indicating that it is in a highly hydrophilic environment.

As we can see from Fig. 15B, there is a very small proteinaceous region from the HCO_3^- toward the stromal solution; thus, protons from the stromal side are expected to have easy access to the site of bicarbonate. In order for an efficient and uni-directional transfer of protons to be able to occur, however, hydrogen-bond networks are expected to be present. In fact, well-defined hydrogen-bond networks have been found linking the bicarbonate to the stromal bulk solution. As shown in Fig. 16, the 3rd oxygen in the bicarbonate that was not ligated to the NHI is hydrogen-bonded to a water molecule (W1138A in the 1.9 Å structure, PDB ID: 3ARC). This water molecule has four hydrogen bonds with its neighboring groups, among which, two are Tyr and Ser residues of the D1 protein (D1-Y246 and D1-S268), and the 3rd one is another water molecule (W675A). This 2nd water molecule (W675A) extends the hydrogen-bond network to the stromal surface through another water molecule W2195D. A plausible hypothesis is: after reduction of Q_B by the reduced Q_A , protons could be easily taken in from the stromal bulk solution through this hydrogen-bond network, and transferred to the site of bicarbonate, which may be further transferred to the reduced Q_B through D1-H272 and D1-H215.

Both the 1st (W1138A) and the 2nd (W675A) water molecules, which are hydrogen-bonded to the bicarbonate, have a tetragonal configuration, bearing 4 hydrogen bonds with their neighboring molecules. The amino acid residues surrounding them thus seem to be important for holding these two water molecules in a proper position, in order to form the proper hydrogen-bond network connecting the bicarbonate to the stromal side. These residues include D1-S268, D1-Y246 for the 1st water molecule, and D1-E244, D2-T243 for the 2nd water molecule. Changes in one of these residues may therefore disturb the positions of the water molecules, and thereby disrupt the proper hydrogen-bond

D1-E244 and D1-Y246, and these could very well be involved in proton pathways (see Figs. 4A and 6A in [158]). They independently propose, as Govindjee, in collaboration with Crofts and Padden [142,159] has suggested that the first proton enters via D1-H252 and the second via D1-Y246, and that it may involve D1-E244.

3.2. The crystal structure at a resolution 1.9 Å and the current model for bicarbonate function

The crystal structure of PSII was first reported by Zouni et al. in 2001 [160] at a resolution of 3.8 Å from a thermophilic cyanobacterium *Thermosynechococcus* (*T.*) *elongatus*. Subsequently, Kamiya and

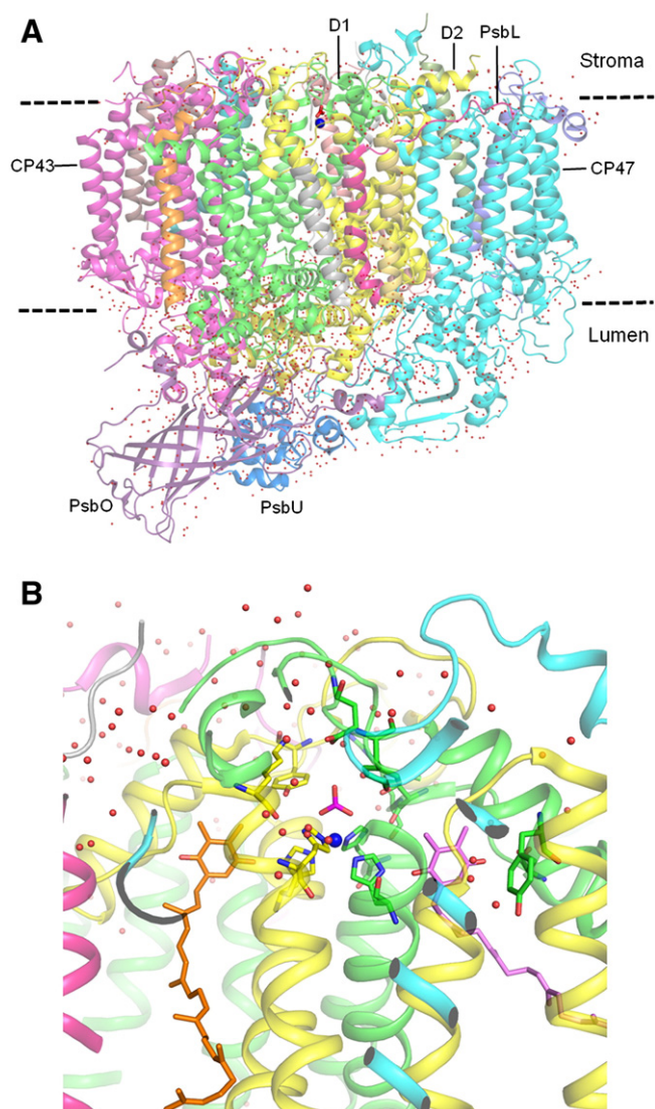


Fig. 15. (A) Structure of a PSII monomer determined at a 1.9 Å resolution [17]. View from the direction perpendicular to the membrane normal. Dashed lines represent the cytoplasmic (stromal) and lumenal surface of the membrane, respectively. Color codes: green, D1; yellow, D2; cyan, CP47; dark pink, CP43; red, PsbL; light pink, PsbO; light blue, PsbU. The blue ball in the middle represents the NHI, and a magenta molecule above the NHI represents bicarbonate. (B) An enlarged view of the NHI and bicarbonate region shown in (A). The color codes are the same as for (A).

networks. D2-K264 is not hydrogen-bonded to any of the two water molecules, but is hydrogen-bonded to D2-E242, and also close to D1-E244, one of the residues hydrogen-bonded to the 2nd water molecule. Alteration of D2-K264 may therefore perturb the orientation of D2-E242, resulting in an effect on the position of the 2nd water molecule.

We know that there is another short H-bond network that connects Q_B to the stromal surface, which is composed of D1-S264 and D1-H252. D1-H252 is located in a small dent in the stromal surface and is hydrogen-bonded to a water molecule directly, which is further hydrogen-bonded to another water molecule. A number of additional water molecules are found in the vicinity of these water molecules, indicating that D1-H252 is located in a highly hydrophilic area. Thus, protons may also be easily taken from this area of the stromal surface and transferred to Q_B through D1-H252 and D1-S264. In view of the previous functional studies [142], it is plausible to suggest that the first proton to protonate Q_B^- is taken up through D1-H252 and D1-S264, and the second proton is transferred via bicarbonate to D1-H272 and D1-H215, and, finally to Q_B through the H-bond network (see Fig. 16, and its legend). HCO_3^- that must become CO_3^{2-} ,

after giving up its proton to Q_B^{2-} , may get its proton back from the stroma via D1-E244 [158,159]. However, further functional studies are required to prove or disprove the order of these protonation events. In addition, since there are indications for a change of the bicarbonate coordination to the NHI from bidentate to monodentate upon electron transport from Q_A towards Q_B [150,156], there might be alternative proton paths newly created by possible accompanying conformational changes.

In view of the above picture of the environment and plausible function of bicarbonate, we recommend comparative biochemical and biophysical studies on appropriate site-directed mutants of D1-E244; D1-Y246; D1-S268; D2-T243; D2-E242; and D2-K264.

4. Uniqueness of role of bicarbonate in oxygenic photosynthesis

The requirement of PSII for bicarbonate (carbonate) has been observed at the level of intact leaves, isolated thylakoids and PSII-enriched membrane fragments from plants, algae, and cyanobacteria, but never in the RCs of anoxygenic photosynthetic bacteria (see sections above and references therein). It appears, therefore, that by being a ligand to the NHI between Q_A and Q_B , and binding to amino acids of the D1 and D2 proteins of PSII in these organisms, bicarbonate/carbonate plays a unique role only in oxygenic photosynthesis: it stabilizes the Q_A -NHI- Q_B structure of the PSII RC, and, thus, allows efficient electron transport and protonation of Q_B^- via certain amino acids around Q_B (Fig. 16). We ask: why does the PSII RC have, unlike its bacterial cousin, a bicarbonate ion liganded to its NHI? The simple answer is that it may have a regulatory function here in PSII electron flow. Under normal conditions, bicarbonate may be bound and function in protonation events, as discussed above; however, when the plant is exposed to drought, high light and high temperature, the stomata may close, leading to a decrease of the internal $[CO_2]$. Similar decreases in $[CO_2]$ are expected in algae and cyanobacteria that do not have stomata [164]. This would lead to a decrease in $[HCO_3^-]$ limiting PSII activity.

The effect of bicarbonate depletion within PSII is not only on the electron acceptor side, but also on the donor side, although the exact location (or binding site) responsible for this effect of HCO_3^- is not known [9,14,20]. The effect of HCO_3^- on the water-oxidizing side of PSII has also been seen *in vitro* in all oxygenic organisms (higher plants, algae, and cyanobacteria) (for details see Section 2). There is, however, lack of observations of this effect in intact organisms. Many experimental data obtained on isolated PSII membrane fragments and PSII core preparations are consistent with a unique role of HCO_3^- in initiating and/or facilitating assembly of the inorganic core of the OEC from OEC-depleted PSII RCs (e.g., arising as a result of disassembly of the OEC under stress conditions or when newly synthesized) and Mn^{2+} ions (reviewed in [61]). There are also indications for the functioning of HCO_3^- in the assembled OEC [14]. Thus, for instance, newer data suggest that mobile (loosely bound or even non-bound) bicarbonate may facilitate deprotonation of the Mn_4CaO_5 cluster (opposite to the protonation reactions assigned for the ‘acceptor-side’ HCO_3^-) [32,34]. By “picking up” the protons that are produced during water splitting, HCO_3^- *per se* or in concert with CA may play a regulatory function against over-acidification of the lumen in the proximity of the water-oxidizing site, and by this, protect the OEC against destabilization and predisposition to photoinhibition.

5. Bicarbonate and evolutionary development of the O_2 -evolving Photosystem II

All O_2 -producing photosynthetic organisms (cyanobacteria, green algae, and plants) have the same Mn_4CaO_5 inorganic core and very similar RC core proteins forming the basis for PSII capable of catalyzing oxidation of water. The available geological and geochemical data indicate that nature created this single type of enzyme as early as 3.2 Ga or as late as 2.4 Ga ago [165–168]. The role of bicarbonate (CO_2) in the evolutionary

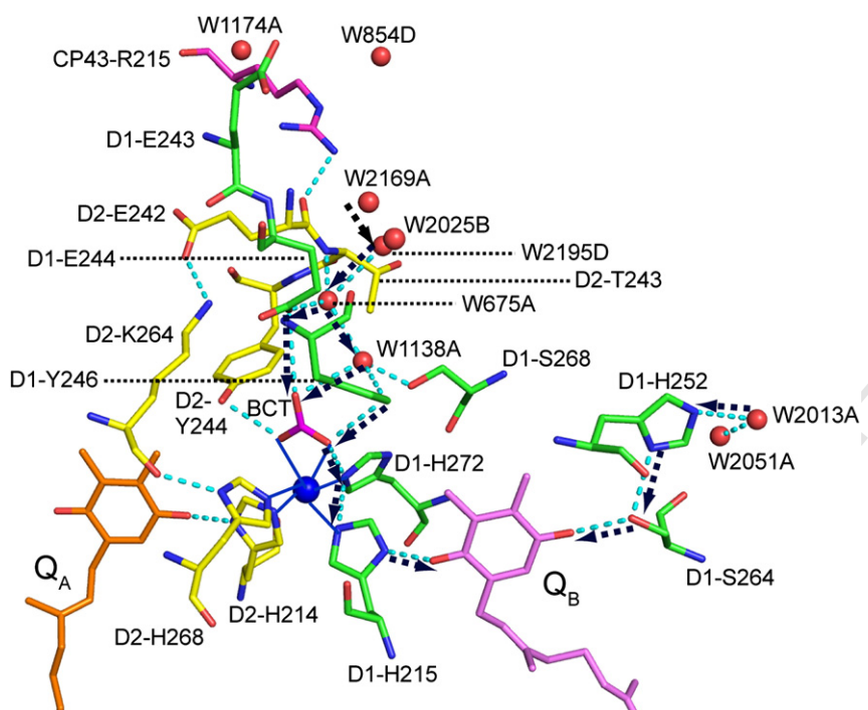


Fig. 16. Hydrogen-bond networks around Q_B , the NHI and bicarbonate. Blue lines represent coordination bonds, and dashed lines in cyan indicate hydrogen-bonds. Arrows in dashed, black lines indicate possible flow of protons towards the Q_B molecule. One of the protons (say the first one) may be picked up through D1-H252 and D1-S264 to protonate Q_B^- , while the second one may be transferred through bicarbonate, D1-H272, and D1-H215 to Q_B (see Section 3.2 for more details). Figure based on data of Umena et al. [17].

1139 development of the first O_2 -evolving cyanobacteria-like organisms is ob-
 1140 vious, since the presence of HCO_3^-/CO_3^{2-} bound between Q_A and Q_B in the
 1141 RC is unique, as it exists only in oxygenic photoautotrophs, whereas it is
 1142 absent in all anoxygenic photosynthesizers [122,123]. The coupling of bi-
 1143 carbonate as a ligand to facilitate Q_B^- protonation and, thus, the electron
 1144 transfer in the first O_2 -producing organisms *via* replacement of the Glu li-
 1145 gand in anoxygenic bacterial RCs (as first suggested in [124]; for further
 1146 details, see Section 3.1.3.1) could be simply an additional evolutionary
 1147 step from anoxygenic towards oxygenic photosynthesis [169].

1148 There are also indications for a key role of Mn–bicarbonate complexes
 1149 in the evolutionary origin of the water-oxidizing inorganic core of the
 1150 OEC of PSII [59,170]. The unique capability of bicarbonate to form easily
 1151 oxidizable complexes with Mn ions has been demonstrated in numerous
 1152 electrochemical and EPR studies (see, for instance, [62,64,65]). Since the
 1153 oxidation potentials of the $Mn^{2+}-HCO_3^-$ complex (520–680 mV) were
 1154 found to have close values to the midpoint redox potentials of the primary
 1155 electron donor (P) in the RCs of non-oxygenic bacteria, Dismukes et al.
 1156 [170] suggested, that these complexes (which could be formed under
 1157 much higher concentrations of dissolved CO_2 (HCO_3^-) in the ancient
 1158 ocean than at present) were probably used as a source of electrons by
 1159 some Archean anoxygenic bacteria—ancestors to the first oxygenic cyano-
 1160 bacteria. Results obtained both on wild type contemporary purple
 1161 bacteria [171] and on mutants of *R. sphaeroides* with modified midpoint
 1162 redox potentials of the P/P⁺ RC couple [33] support this idea by showing
 1163 that the formation of the Mn^{2+} –bicarbonate complexes stimulate
 1164 electron donation from Mn^{2+} to type II RCs of these anoxygenic
 1165 bacteria.

1166 6. Concluding remarks

1167 As proved by recent X-ray crystallography studies of PSII [16,17],
 1168 in the cyanobacterial RC, there is only evidence for a single bound bi-
 1169 carbonate at the NHI. The evidence for “bicarbonate” as ligand to the
 1170 quinone–iron complex derived from a large body of data makes it
 1171 clear that there is a role for this ligand *in vivo*. The presence of bicar-
 1172 bonate as a bidentate ligand to the NHI bridging Q_A and Q_B is now

1173 firmly established [16,17]. This set the key stone to a huge body of
 1174 studies that have established a role of bicarbonate in facilitating proton
 1175 transfer and, thereby, accelerating electron transfer between Q_A ,
 1176 Q_B and from Q_B into the PQ-pool; the absence of bicarbonate might
 1177 down-regulate this electron transfer step. Since this action prevails
 1178 in all oxygenic organisms, the structural and functional role of bicar-
 1179 bonate has arisen very early in evolution. There is also an effect on the
 1180 reoxidation of PQH₂. Comparative biochemical and biophysical stud-
 1181 ies on site-directed mutants of tyrosines near the HCO_3^- binding site
 1182 is expected to provide key information on the mechanistic role of bi-
 1183 carbonate in these reactions.

1184 There is another, though less well defined role of bicarbonate on the
 1185 donor side of PSII. A particular binding site close to the Mn_4CaO_5 cluster
 1186 is absent in the high-resolution structure [17]. Since the roles of HCO_3^-
 1187 as a mobile substrate of PSII or as a direct tightly bound ligand to the
 1188 Mn_4CaO_5 cluster are excluded by numerous studies, a direct involvement
 1189 of HCO_3^- in the water-oxidizing process can now be ruled out. There is,
 1190 however, undeniable evidence for an essential role of HCO_3^- in the pro-
 1191 cess of photoactivation. Further experiments are required to evaluate
 1192 the possibility of HCO_3^- involvement in the deprotonation reactions of
 1193 the OEC. The indirect effects of HCO_3^- on water oxidation (such as, protec-
 1194 tion against thermoinactivation, photoinhibition, protein extraction, and
 1195 treatments with some reductants) need to be studied and characterized
 1196 further. Moreover, one should clarify whether HCO_3^- ions have the
 1197 same function on the donor side of PSII in intact photosynthetic systems.

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