Chlorophyll Biosynthesis Gene Evolution Indicates Photosystem Gene Duplication, Not Photosystem Merger, at the Origin of Oxygenic Photosynthesis

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Abstract

An open question regarding the evolution of photosynthesis is how cyanobacteria came to possess the two reaction center (RC) types, Type I reaction center (RCI) and Type II reaction center (RCII). The two main competing theories in the foreground of current thinking on this issue are that either 1) RCI and RCII are related via lineage divergence among anoxygenic photosynthetic bacteria and became merged in cyanobacteria via an event of large-scale lateral gene transfer (also called "fusion" theories) or 2) the two RC types are related via gene duplication in an ancestral, anoxygenic but protocyanobacterial phototroph that possessed both RC types before making the transition to using water as an electron donor. To distinguish between these possibilities, we studied the evolution of the core (bacterio)chlorophyll biosynthetic pathway from protoporphyrin IX (Proto IX) up to (bacterio)chlorophyllide a. The results show no dichotomy of chlorophyll biosynthesis genes into RCI- and RCII-specific chlorophyll biosynthetic clades, thereby excluding models of fusion at the origin of cyanobacteria and supporting the selective-loss hypothesis. By considering the cofactor demands of the pathway and the source genes from which several steps in chlorophyll biosynthesis are derived, we infer that the cell that first synthesized chlorophyll was a cobalamin-dependent, heme-synthesizing, diazotrophic anaerobe.

Key words: redox-switch, selective-loss, manganese, cyanobacteria, fusion.

Introduction

The origin of oxygenic photosynthesis introduced a new high-potential electron acceptor into microbial ecosystems (Holland 2006) and enzymatic reaction mechanisms (Raymond et al. 2002), marking the onset of pivotal changes in geochemical cycles and biochemical pathways. Oxygen first began to accumulate in the atmosphere approximately 2.4 billion years ago, and its subsequent accumulation in the oceans was slower, such that full oxic conditions were only reached approximately 635-580 Ma (Arnold et al. 2004; Lyons 2007; Canfield et al. 2008; Scott et al. 2008; Lyons et al. 2009; Sahoo et al. 2012). This "new chemistry" had far-reaching impact on the evolutionary process.

Chlorophyll-based photosynthesis arose among eubacteria, where it is currently found among six phyla. Chlorobia (green sulfur bacteria [GSB]), firmicutes (heliobacteria), and acidobacteria have anoxygenic Type I reaction centers (RCIs), whereas chloroflexi (green nonsulfur bacteria, GNSB of filamentous anoxygenic phototrophs, FAPs) and some proteobacterial organisms (purple nonsulfur bacteria and purple sulfur oxidizing bacteria [Dahl et al. 2005]) perform anoxygenic photosynthesis with Type II reaction centers (RCIIs) (Xiong and Bauer 2002a; Bryant et al. 2007; Magueo Chew and Bryant 2007; Blankenship 2010). Only cyanobacteria and, via endosymbiosis, photosynthetic eukaryotes (Margulis and Bermudes 1985) perform oxygenic photosynthesis, having both the Photosystem I (PSI) and the Photosystem II (PSII) complexes that are homologous to RCI and RCII, respectively (Michel and Deisenhofer 1988; Hauska et al. 2001; Neerken and Amesz 2001). Although at the sequence level there is

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almost no detectable similarity between RCI and RCII, their structural and cofactor arrangements are unquestionably homologous and clearly indicate common ancestry (Schubert et al. 1998; Barber et al. 2000; Sadekar et al. 2006).

Chlorophyll (Chl) and bacteriochlorophyll (Bch) are required for oxygenic and anoxygenic photosynthesis, respectively, where they serve two essential functions. As light-harvesting antennae, they act as photon funnels, absorbing light and channeling its energy to reaction centers (RCs), where chlorophylls perform their second function: photochemical charge separation to create strong oxidants and reductants, sending low-potential electrons through the electron transport chain (ETC). In anoxygenic phototrophs, bacteriochlorophylls are the main pigments. In contrast, chloroplasts and cyanobacteria possess only chlorophylls, which can, however, also be residually present in the RC of some anoxygenic photosynthetic bacteria (Kobayashi et al. 2000; Hohmann-Marriott and Blankenship 2011; Sarrou et al. 2012; Tsukatani et al. 2012). With exception of chlorophyll c, chemically, both chlorophylls and bacteriochlorophylls are chlorins; reduced magnesium-containing cyclic tetrapyrroles with an additional fifth isocyclic ring. Their main differences concern the level of unsaturation affecting the system of conjugated double bonds. Chlorophyll has a single bond between carbons C17 and C18 (IUPAC numbering), and bacteriochlorophyll has an additional C7-C8 single bond (Maqueo Chew and Bryant 2007; Niedzwiedzki and Blankenship 2010). There are 11 major types of bacteriochlorophyll and chlorophylls (Hohmann-Marriott and Blankenship 2011) differing with respect to the substituents of the tetrapyrrole ring, hence their differing absorption properties, which allow organisms to specialize toward different spectral niches, most notably as a function of depth in the water column (Glaeser et al. 2002; Manske et al. 2005; Gomez Maqueo Chew et al. 2007; Kiang, Siefert, et al. 2007; Kiang et al. 2007; Stomp et al. 2007; Chen et al. 2010).

The enzymes involved in chlorophyll metabolism are the only set of photosynthesis-related proteins common to all phototrophs (Mulkidianian et al. 2006), and thoughts on chlorophyll evolution have a long history. Central to the topic is the Granick hypothesis (Granick 1965), which posits that the evolution of the chlorophyll biosynthetic pathway followed the sequential inventions of new enzymes to generate more stable products. This premise has been widely used to study the evolution of the chlorophyll pathway, in particular as a proxy for the evolution of photosynthesis itself (Olson and Pierson 1987a, 1987b; Xiong and Bauer 2002a, 2002b; Gupta 2012). However, other hypotheses are still discussed, and the issue is debated (Lockhart et al. 1996; Blankenship 2001). In 2000, Xiong et al. studied phylogenies for 9 of the 17 enzymes then known to be involved in the (bacterio)chlorophyll pathway. They suggested that (bacterio)chlorophylls first arose within purple bacteria (proteobacteria) and that the pathway's emergence involved the recruitment and duplication of

homologous enzymes such as nitrogenase subunits and cobalt chelatase from cobalamin biosynthesis (Xiong et al. 1998; Xiong and Bauer 2002a, 2002b). More recently, and based on phylogenies and sequence signatures of the (B)ChlNBL complex, which is responsible for catalyzing the last step of the (bacterio)chlorophyll core pathway, an origin of chlorophyll in the Gram-positive heliobacteria lineage has been suggested (Gupta 2012).

Here, we address the evolution of the chlorophyll biosynthesis pathway to distinguish between competing theories for the origin of two photosystems in cyanobacteria. The RCs of photosystems I and II are clearly related at the level of three-dimensional structure (Nitschke and Rutherford 1991: Schubert et al. 1998; Baymann et al. 2001; Allen and Puthiyaveetil 2005; Allen et al. 2011; Hohmann-Marriott and Blankenship 2011), and the issue is how they came to reside within the ancestral cyanobacterial genome. The "fusion" (or "merger") hypothesis asserts that the two photosystems diverged during the evolution of anoxygenic photosynthetic lineages and became merged in the founding cyanobacterium via lateral gene transfer (LGT) (Mathis 1990; Meyer 1994; Xiong and Bauer 2002a; Hohmann-Marriott and Blankenship 2011). In contrast, the "duplication" hypothesis asserts that the photosystems diverged within a protocyanobacterial ancestor and subsequently underwent vertical inheritance and export, via LGT, to diverse anoxygenic photosynthetic lineages (Olson and Pierson 1987a, 1987b; Baymann et al. 2001; Olson 2001; Allen 2005; Mulkidjanian et al. 2006). Were the merger hypothesis correct, the genes of chlorophyll biosynthesis in anoxygenic photosynthetic lineages should reflect early lineage splittings and hence the same deep divergence as the photosynthetic RCs. In the case that the two photosystems arose via duplication in a protocyanobacterium, the chlorophyll biosynthetic pathway should not reflect ancient lineage splittings, that is, there should be no deep dichotomy into RCI and RCII-specific chlorophyll biosynthetic pathways types. To distinguish between these possibilities, we have studied the evolution of the core (bacterio)chlorophyll biosynthetic pathway from Proto IX up to (bacterio)chlorophyllide a, which is the last chemical intermediate common to all chlorophyll types.

Materials and Methods

Data

The list of organisms was extracted from the GOLD table as accessed on the 16 June 2010 (Bernal et al. 2001). Genomes of photosynthetic prokaryotes were downloaded from RefSeq database release 41 of 9 May 2010 (Pruitt et al. 2005). In addition, the recently sequenced genome of *Candidatus Chloracidobacterium thermophilum* was also included (Garcia Costas et al. 2012). Homologous proteins involved in (bacterio)chlorophyll biosynthesis were identified by Basic

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Local Alignment Search Tool (BLAST) (Altschul et al. 1997) within the data set of downloaded genomes (17 eukaryotes and 87 prokaryotes, supplementary table S1, Supplementary Material online) using the proteins shown in supplementary table S2, Supplementary Material online, as queries. The BLAST lists were filtered for *E* values better than 10⁻¹⁰ and amino acid identities of ≥25%. Paralogs from the cobalamin pathway or later steps in the bacteriochlorophyll *c* pathway were removed. Because some enzymes of chlorophyll biosynthesis (BchE/ChIE, BciA, and light-dependent NADPH:protochlorophyllide oxidoreductase [LPOR]) belong to protein families with members catalyzing reactions in other pathways, possible paralogs were identified by reBLASTing against the entire National Center for Biotechnology Information (NCBI) refseq database.

Sequence Alignments and Phylogenetic Analysis

Proteins were aligned using MUSCLE (Edgar 2004) using its default parameters. Trees were reconstructed using PhyML 3.0 (Guindon and Gascuel 2003) with the WAG+I+G model and four rate categories. Alignments are available upon request and nonconcatenated phylogenetic trees are given as supplementary figures S1–S12, Supplementary Material online.

Topology Comparisons

Topology testing was performed using the approximately unbiased (AU) test and the Shimodaira–Hasegawa (SH) test (Shimodaira and Hasegawa 1999; Shimodaira 2002). Where justified, initial constrained trees with an imposed separation of (*B*)*Chls* genes from RCI- and RCII-containing organisms were created with an in-house perl script and optimized in PhyML 3.0 (Guindon and Gascuel 2003). The site likelihoods of the constrained and the nonconstrained trees were calculated using PhyML 3.0 and the AU and the SH tests performed using CONSEL v0.1k (Shimodaira and Hasegawa 2001).

Results

Taxonomic Distribution of Photosynthesis and Bacteriochlorophyll Biosynthesis Genes

The taxonomic distribution of the photosynthetic apparatus in a sample of sequenced prokaryotic genomes is presented in figure 1A. Photosynthesis is absent among archaebacteria and present in only 6 of the 24 bacterial phyla currently recognized by NCBI taxonomy. Within those six phyla, photosynthesis is, in general, sparsely distributed. All cyanobacterial genomes and, with one exception—the recently sequenced *Ignavibacterium album* bacterium (an early diverging member of the chlorobia phylum) (lino et al. 2010; Liu et al. 2012), all chlorobial genomes sequenced so far have genes coding for the photosynthetic apparatus. *Ignavibacterium* is a facultative aerobe, lacks photosynthesis genes, and is currently the only

nonphototrophic chlorobium to have been cultured (Liu et al. 2012). Nevertheless, the dense distribution of photosynthetic genes in cyanobacteria and chlorobia suggests their presence in the ancestors of these phyla. The same can also be assumed for chloroflexi, at least at the class level. The presence of 1 to 3 chlorophyll biosynthesis genes in Dehalococcoides sp. BAV1, Dehalococcoides sp. CBDB1, and Herpetosiphon aurantiacus probably indicates either LGT or past presence of the pathway and its subsequent loss. In contrast, within proteobacteria and the firmicutes, even at the class level, a sparse distribution of photosynthetic representatives is found, indicating either late acquisition(s) of photosynthesis or many independent losses of it. Because photosynthetic genes are often encoded in superoperons (Alberti et al. 1995; Xiong et al. 1998) or lineagespecific small photosynthetic gene clusters (Xiong et al. 2000), LGT scenarios can, in principle, account for the patchy distribution of photosynthesis among eubacteria. This would also be consistent with the presence of photosynthesis genes, including both photosystems I and II, in the marine phage metagenome (Lindell et al. 2004; Sharon et al. 2009). However, scenarios of vertical inherence have also been discussed, at least for the evolution of RCI-containing organisms (Mix et al. 2005).

Genes involved in chlorophyll biosynthesis were identified in our data set of 104 complete sequenced genomes from photosynthetic organisms (see Materials and Methods). Their taxonomic distribution is presented in figure 1B. The enzymes of the core chlorophyll biosynthesis pathway are present in all photosynthetic organisms, regardless of which RC type the organism uses. The few differences that are observed with respect to gene presence or absence concern O_2 -dependent and O_2 -independent alternative enzymes for some steps (see later). Clearly, a single pathway of chlorophyll biosynthesis was present in the ancestral photosynthetic bacterium.

Mg Chelatase

The first unique intermediate of chlorophyll biosynthetic pathway, Mg-protoporphyrin IX (fig. 2A, step 1), is generated by the insertion of Mg²⁺ into protoporphyrin IX. Biochemical and genetic analysis identified a Class I ATP-dependent magnesium chelatase, composed of three subunits BchH, BchI, and BchD, that catalyzes a reaction (Walker and Willows 1997; Willows and Beale 1998; Bollivar 2006) consisting of an activation and a chelation step. In the presence of both ATP and Mg²⁺, an AAA+ motor complex is assembled from a hexameric (Willows et al. 2004) or heptameric (Reid et al. 2003) BchI ring connected to a hexameric BchD ring (Lundqvist et al. 2010) (the activation step). Proto IX binds to the BchH catalytic subunit, and its transient interaction with the formed AAA+ motor complex leads to the insertion of Mg²⁺ in the tetrapyrrole macrocycle (the chelation step).



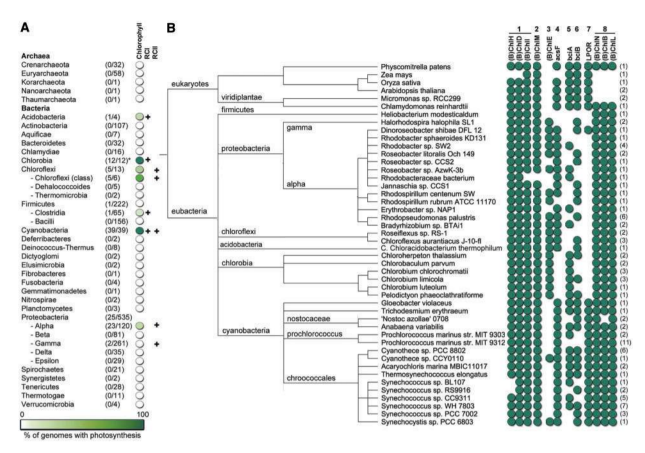


Fig. 1.—Distribution of photosynthesis in prokaryotes. (A) Distribution of photosynthesis in completely sequenced genomes of prokaryotic organisms organized according to their phyla and class. The color gradient of each circle reflects the normalization of species within a certain taxon that have genes coding for photosynthetic apparatus versus the total number of organisms from that taxa whose genome has been completely sequenced. Numbers in brackets represent the photosynthetic species within that taxon and the total number of species with completely sequenced genomes. *Ignavibacterium album, the only known nonphotosynthetic member of chlorobia phylum was not included in our data set. (B) Taxonomic distribution (presence or absence) of the different genes involved in the biosynthesis of (bacterio)chlorophyll from Mg-protoporphyrin IX to chlorophyllide a. Columns correspond to genes involved in the different steps of chlorophyll photosynthesis. A modified taxonomic tree is represented on the left where leafs are grouped by taxonomy and same pattern of presence and absence of genes. Numbers between brackets represent number of organisms with the same pattern belonging to same taxonomic group: Physcomitrella patens; Zea mays; Oryza sativa; Arabidopsis thaliana (2): A. thaliana and Vitis vinifera; Micromonas sp. RCC299 (2): Micromonas sp. RCC299 and Ostreococcus lucimarinus CCE9901; Chlamydomonas reinhardtii; Heliobacterium modesticaldum Ice1; Halorhodospira halophile SL1 (2): H. halophile SL1 and Allochromatium vinosum DSM180; Dinoroseobacter shibae DFL 12; Rhodobacter sphaeroides KD131; Rhodobacter sp. SW2 (4): Rhodobacter sp. SW2, R. sphaeroides ATCC17025, R. sphaeroides ATCC17029, and R. sphaeroides 241; Roseobacter litoralis Och149 (2): Ros. litoralis Och149 and Ros. denitrificans OCh114; Roseobacter sp. CCS2; Roseobacter sp. AzwK-3b; Rhodobacterales bacterium HTCC2083; Jannaschia sp. CCS1; Rhodospirillum centenum SW; Rho. rubrum ATCC11170; Erythrobacter sp. NAP1; Rhodopseudomonas palustris (6): BisB18, HaA2, CGA009, BisA53, BisB5, and TIE-1; Bradyrhizobium sp. BTAi1 (2): sp. BTAi1 and sp. ORS278; Roseiflexus sp. RS-1 (2): Roseiflexus sp. and Roseiflexus castenholzii DSM13941; Chloroflexus aurantiacus J-10-fl (3): Chloroflexus aurantiacus J-10-fl, Chloroflexus aggregans DSM9485, and Chloroflexus sp. Y-400-fl; C. Chloracidobacterium thermophilum: Candidatus Chloracidobacterium thermophilum; Chloroherpeton thalassium (2): Chloroherpeton thalassium ATCC35110 and Prosthecochloris aestuarii DSM271; Chlorobaculum parvum (2): NCIB 8327 and Chlorobaculum tepidum TLS; Chlorobium chlorochromatii (3): Chlorobium chlorochromatii CaD3, Chlorobium phaeobacteroides BS1, and P. vibrioformis DSM 265; Chlorobium limicola (3): Chlorobium limicola DSM245, Chlorobium phaeobacteroides DSM266, and Chlorobium ferrooxidans DSM13031; Chlorobium luteolum DSM273; Pelodictyon phaeoclathratiforme BU-1; Gloeobacter violaceus PCC7421; Trichodesmium erythraeum IMS101; "Nostoc azollae" 0708 (2): "Nostoc azollae" 0708 and Nostoc punctiforme PCC73102; Anabaena variabilis (2): ATCC29413 and Nostoc sp. PCC7120; Prochlorococcus marinus str. MIT9303 (2): str. MIT9303 and MIT 9313; Prochlorococcus marinus str. MIT9312 (11): str. MIT9312, subsp. pastoris. str. CCMP1986, subsp. pastoris. str. CCMP1375, str. NATL1A, str. NATL2A str. MIT9202, str. MIT9215, str. MIT9211, str. MIT9515, str. AS9601, and str. MIT9301; Cyanothece sp. PCC8802 (6): sp. PCC8802, sp. PCC7424, sp. PCC7425, sp. ATCC51142, sp. PCC7822, and sp. PCC8801, Cyanothece sp. CCY0110; Acaryochloris marina MBIC11017 (2): Aca. marina MBIC11017 and Microcystis aeruginosa NIES843; Thermosynechococcus elongatus BP-1; Synechococcus sp. BL107; Synechococcus sp. RS9916 (2): sp. RS9916 and sp. WH7805; Synechococcus sp. CC9311 (5): sp. CC9311, sp. WH8109, sp. WH8102, sp. CC9902, and sp. CC9605; Synechococcus sp. WH7803(7): sp. WH7803, sp. JA-3-3Ab, elongatus PCC7942, elongatus PCC6301, sp. 2-3Ba2-13, sp. PCC7335, and sp. WH5701; Synechococcus sp. PCC7002 (3): sp. PCC7002, sp. RS9917, and sp. RCC307; and Synechocystis sp. PCC6803.

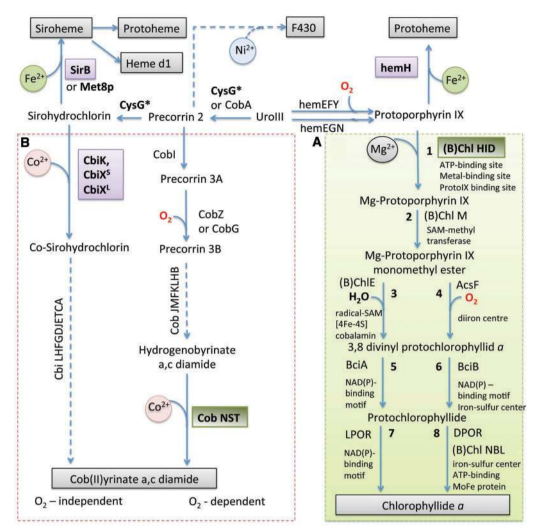


Fig. 2.—Schematic representation of the tetrapyrrole biosynthesis pathway. Based on information from Martens et al. (2002), McGoldrick et al. (2005), Heinemann et al. (2008), Storbeck et al. (2010) and Bali et al. (2011). The different metal chelatases are represented in bold and colored according to their sequence and/or structural similarity. The numbers correspond to the different steps of the common core of (bacterio)chlorophyll pathway. UrollI, the last common intermediate of all tetrapyrroles, can either be methylated by CysG (or CobA) to form precorrin-2 or suffer decarboxylation and oxidation to form Protoporphyrin IX via the O2-dependent route (HemE, HemF, HemY) or the O2-independent route (HemE, HemG, HemN). Precorrin-2 is the branching point for F430 (a Ni²⁺-containing tetrapyrrole), the cobalamin O₂-dependent route, and Sirohydrochlorin. Iron can be inserted into Sirohydrochlorin by SirB or by multifunctional oxidases such as Met8p or CysG to form siroheme. Siroheme can be further modified to form heme d1 or heme, according to the alternative archaeal heme synthesis pathway. Protoporphyrin IX is the branch point to hemes and chlorophylls. Iron is inserted into protoporphyrin IX (classical pathway). (A) Specific (bacterio)chlorophyll steps: the incorporation of Mg²⁺ into protoporphyrin IX occurs by the action of (B)ChlHID complex (step 1), resulting in the formation of Mg-protoporphyrin. A methyl transferase (B)ChIM subsequently converts Mg-protoporphyrin into Mg-protoporphyrin IX monomethyl ester (step 2), which is the substrate for a cyclase reaction, involving water (B)ChIE (step 3) or molecular oxygen (AcsF, step 4). The cyclase forms divinyl protochlorophyllide a, an intermediate that contains a fifth ring (ring E), characteristic of all (bacterio)chlorophylls. The next reaction consists in the reduction of the 8-vinyl group by an 8-vinyl reductase, BciA (step 5) or BciB (step 6). Finally, the reduction of the IV pyrrole ring of protochlorophyllide to form chlorophyllide is performed by the action of the light-dependent LPOR (step 7) or the light-independent (DPOR) complex formed by (B)Chl LNB. (B) Alternative pathways for cobalamin synthesis. In the O2-independent cobalamin pathway, cobalt is inserted into Sirohydrochlorin to form the Co-Sirohydrochlorin intermediate, via one of the three homologous chelatases CibK, CibX^S, or CibX^L homologs to the SirB chelatase and the HemH chelatase from the classical heme pathway. Co-Sirohydrochlorin is then sequential converted into Cob(II)yrinate a,c diamide by the action of 10 enzymes (Cbi LHFGDJETCA). In the O2dependent pathway, Cobl performs the methylation of precorrin-2 originating precorrin-3A that will be oxidized by one of the two O2-dependente enzymes (CobZ or CobG) originating precorrin-3B. A series of reactions involving cob JMFKLHB genes occur until hydrogenobyrinate a,c diamide is formed. At this stage, Co²⁺ ion is inserted into the ring by the CobNST (homologous to BChl HID chelatase from the (bacterio)chlorophyll pathway) forming Cob(II)yrinate a,c diamide, the merger point between the two cobalamin pathways.



The Mg-chelatase complex has both sequence and structural homology with the Class I cobalt chelatase of the O₂-dependent cobalamin biosynthetic pathway (Lundqvist et al. 2009). The CobN, CobS, and CobT subunits of the trimeric cobalt chelatase are homologous with the BchH/ChlH, Bchl/ChlI, and BchD/ChlD subunits of magnesium chelatase, respectively (Fodje et al. 2001; Lundqvist et al. 2009). This homology has been interpreted as reflecting ancient duplication and divergence (Lundqvist et al. 2009). Although there is a broad taxonomic distribution of the *CobN* gene among cobalamin-dependent organisms, a much narrower distribution is observed for the genes that compose the AAA+ motor complex (*CobS* and *CobT*) (Rodionov et al. 2003).

The Catalytic Subunit BchH/ChlH

Chlorobaculum tepidum, similar to most members of GSB, has three paralogous genes for this subunit, named BchS, BchH, and BchT. Single- and double-mutant experiments (Gomez Maqueo Chew et al. 2009) showed that strains with only the BchH or the BchS gene retained sufficient Mg-chelatase activity to be viable, but its activity was maximal if the BchS gene was present together with BchH. This was also confirmed by biochemical characterization of the recombinant enzymes (Johnson and Schmidt-Dannert 2008). However, in all mutagenesis experiments, a decrease in the production of bacteriochlorophyll c, the main photosynthetic pigment of GSB, was detected. This suggests that the different isoforms function in end-product regulation and/or substrate channeling of the Bch c intermediates (Johnson and Schmidt-Dannert 2008; Gomez Maqueo Chew et al. 2009).

The existence of *BchH/ChlH* paralogs is not unique to the GSB. All chloroflexi have at least one additional gene coding this subunit (Gomez Maqueo Chew et al. 2009), and the same is also true for some cyanobacterial species (Lohr et al. 2005). Isoforms of the gene also occur in some eukaryotes (Lohr et al. 2005) although, in this case, different duplication events are possibly at their origin.

The schematic diagram summarizing the BchH/ChlH phylogenetic tree is presented in figure 3A. A clear dichotomy between the different subunit isoforms can be observed. The upper clade contain sequences similar to the BchH gene of Chlorobaculum tepidum TLS. With the exception of Chlorobium chlorochromatii, all organisms from the seven phyla considered contain at least one copy the BchH/ChlH gene. In this group, three distinct groups are retrieved: 1) chloroflexales and chlorobia, 2) acidobacteria and proteobacteria, and 3) heliobacteria and cyanobacteria with photosynthetic eukaryotes branching as basal members. Recent events of gene duplication can also be observed in the chloroflexales ancestor and at the species level of three land plants.

Notably, there is no relationship between the type of RC and the organization of the sequences within the tree, indicating a single origin of this pathway. Moreover, a close

relationship between chlorobia and chloroflexi sequences is observed, a relationship found in most of our phylogenetic analysis. Based inter alia on the observation that fundamental aspects of core carbon and energy metabolism in chlorobia and chloroflexi are different, it has been suggested that lateral transfer of genes underlying chlorosome structure and chlorophyll biosynthesis might have occurred between these groups (Raymond et al. 2002). However, the close phylogenetic proximity of chlorophyll biosynthesis genes among organisms that contain different RC types is not confined to these two groups.

The lower clade of the tree contains sequences similar to the *Chlorobaculum tepidum* TLS *BchS* and *BchT* genes, belonging to chlorobia, chloroflexi, cyanobacteria, and green algae. In this clade, two distinct groups are retrieved, one comprising the cyanobacterial and plastid sequences and the other with chloroflexi BchH2 and chlorobial BchS and BchT sequences. A duplication event in the ancestor of the chlorobia appears to be the likely origin of the two isoforms.

The role of these subunits has been proposed to be either end-product regulation and/or substrate channeling of the Bch c intermediates (Johnson and Schmidt-Dannert 2008; Gomez Maqueo Chew et al. 2009). Both chlorobia and chloroflexi have Bch c pigments, but the cyanobacteria synthesize only chlorophyll a, raising the question of what the role of these cyanobacterial homologs might be. The *Gloeobacter violaceus* ChlH2 sequence alone falls outside these two main clades; furthermore, it is more similar to homologs in nonphotosynthetic lineages (euryarchaeotes and firmicutes) than to its photosynthetic homologous (data not shown). This suggests that *Gloeobacter* ChlH2 is not involved in chlorophyll biosynthesis.

The presence of BchH/ChlH isoforms in three out of the six considered bacterial phyla and their monophyly within the two clades may suggest their presence in the ancestral chlorophyll pathway, in which case the isoforms would have been lost (or not LGT acquired) in proteobacteria, acidobacteria, and heliobacteria.

The AAA+ Motor: Bchl/Chll and BchD/ChlD

The Bchl/Chll subunit is a member of the AAA⁺-ATPase family of proteins, which includes proteins of diverse function (Fodje et al. 2001). With the exception of some land plants (Kobayashi et al. 2008; Huang and Li 2009) and green algae (Lohr et al. 2005), there is a consensus in the literature that photosynthetic organisms only have one Bchl/Chll subunit. However, a study of two recombinant isoforms of Bchl from *Prosthecochloris vibrioformis* showed that both had Mg chelatase activity in vitro (Petersen et al. 1998), and in the recently sequenced genome of *Chloroflexus aurantiacus*, three genes were annotated as *Bchl* (Tang et al. 2011). In our search, several isoforms belonging to both chlorobia and chloroflexi were retrieved. These enzymes are distinct from the aerobic

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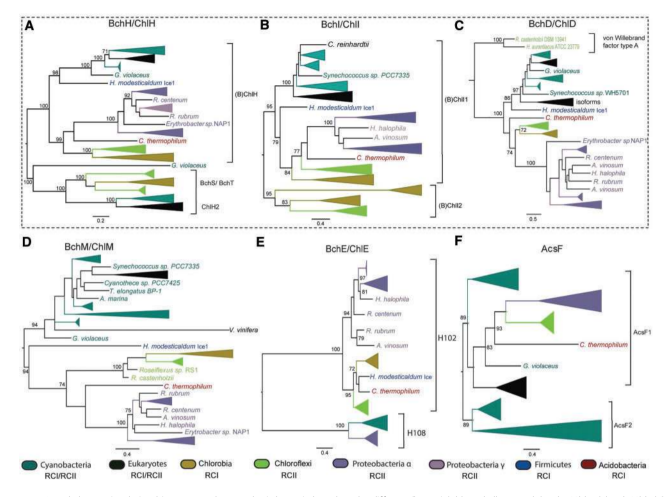


Fig. 3.—Phylogenetic relationships among photosynthetic bacteria based on the different (bacterio)chlorophyll genes. (A) BchH/ChIH, (B) BchI/ChII, (C) BchD/ChID, (D) BchM/ChIM, (E) BchE/ChIE, and (F) AcsF.

cobalamin chelatase *CobS* gene and were included in the analysis (fig. 3*B*).

The Bchl/Chll tree is divided into two clades, one comprising Bchl/Chll genes among all photosynthesizers and one containing the different isoforms from GSB and GNSB, which we call the GSB/GNSB clade. In the former, two major groups can be observed, one with heliobacteria, chlorobia, chloroflexi, acidobacteria, and proteobacteria (branching in that order) and a second one, where the cyanobacteria and eukaryotic sequences cluster together. The isoforms present in the GSB/ GNSB clade, as in the case of G. violaceus ChIH2 sequence, have high similarity with sequences similar to hypothetical magnesium chelatases from nonphotosynthetic organisms. Although the results from P. vibrioformis indicate that both isoforms have identical activities in vitro, their sequences show higher similarity with sequences from organisms that do not synthetize bacteriochlorophyll, questioning their role in chlorophyll biosynthesis. Specifically, within delta-proteobacteria, proprionobacterales, firmicutes, and euryarchaeota, some organisms possess isoforms of the Bchl/Chll gene in addition to CobS. Other cobalamin producers possess only the Bchl/Chll gene and lack CobS. In these organisms, the Bchl/Chll genes probably substitute the missing CobNST genes (Xiong et al. 1998; Lohr et al. 2005). The distributions of Bchl/Chll and CobS homologs suggest a way in which pre-existing building blocks could have been recruited into the assembly of the ancestral chlorophyll and O₂-dependent cobalamin pathway.

The N-terminus of *BchD/ChID* gene, which is also a member of the AAA+-ATPase family of proteins (Fodje et al. 2001), has a segment of approximately 260 amino acid residues with sequence homology to Bchl/ChII (Masuda et al. 1999). BchD/ChID has a proposed structural role in magnesium chelatase, functioning as a platform for the convergence of the other two subunits (Axelsson et al. 2006). The BchD/ChID phylogenetic tree in figure 3C was rooted by two sequences of von Willebrand factor Type A, which is considered to be an ancient protein domain (Ponting et al. 1999). The overall BchD/ChID tree topology is very similar to the upper part of the BchH/ChIH tree with the exception of the position of *C. Chloracidobacterium thermophilum*. There are



two distinct BchD/ChlD clades. One contains acidobacteria, the GSB/GNSB clade, and proteobacteria. In the second, heliobacteria, cyanobacteria, and eukaryotic sequences cluster. There are two eukaryotic clades, one composed of green algae isoforms that branch between heliobacteria and cyanobacteria, and a second in which at least one copy from every eukaryotic organism is represented in a sister group of the *G. violaceus* sequence. However, this branch is poorly supported. As in the case of *Bchl/Chll*, copies of this gene are also present in nonphotosynthetic organisms.

Mg Protoporphyrin IX Methyltransferase

The next step in the pathway of synthesis is the insertion of a methyl group by the enzyme Mg-protoporphyrin IX methyl transferase (BchM/ChlM) to form the Mg-protoporphyrin IX monomethyl ester (MPE) (fig. 2A, step 2). This enzyme belongs to the Class I S-adenosylmethionine-(SAM-)dependent methyltransferase family (Gibson and Hunter 1994) and transiently interacts with BchH/ChlH (Johnson and Schmidt-Dannert 2008). With the exception of Arabidopsis thaliana, all photosynthetic organisms have only one BchM/ChlM gene. There is strong support for monophyly of the major photosynthetic taxa, and in this case, because cyanobacteria and eukaryotes cluster together, while heliobacteria branch closer to the chlorobia/chloroflexi sister groups and acidobacteria closer to proteobacteria (fig. 3D).

Mg-Protoporphyrin IX Monomethyl Ester Oxidative Cyclases

The next step in the pathway is the formation of the isocyclic ring by the conversion of MPE to 3,8-divinyl protochlorophyllide a. This complex reaction entails the cyclization of the ring and the introduction of a keto group at carbon 13¹. This step is catalyzed by either of two different and unrelated enzymes: the oxygen-dependent enzyme AcsF (fig. 2A, step 4) and the oxygen-independent enzyme (BchE), both of which catalyze the reaction (fig. 2A, step 3). The arguably more ancient reaction of the oxygen-independent cyclase, BchE, uses water as electron donor and is widely distributed among heliobacteria, proteobacteria, GNSB, and GSB. Although all cyanobacteria have the AcsF enzyme, some, including Synechococcus sp. PCC 6803, also harbor a functional ChIE gene (Minamizaki et al. 2008) that can complement Rhodobacter capsulatus BchE-deficient strains (Bollivar 2006). BchE/ChlE is a radical-SAM enzyme with homology to adenosylcobalamin-binding domain. In R. capsulatus and Chlorobium limicola strain 1230, the enzyme appears to be vitamin B12 (cobalamin) dependent (Fuhrmann et al. 1993; Gough et al. 2000). However, these two sequences lack the canonical signature motif (DXHXXG) that typically contains the cobalt coordinating histidine residue (Drennan et al. 1994). The BchE/ChlE alignment reveals, however, a highly conserved histidine (H102 Chlorobaculum tepidum TLS numbering) that might function as the missing cobalt ligand, and, in sequences where H102 is missing, another histidine residue (H108 *Synechocystis* sp. *PCC6803* numbering) is present close by, whereas no functional role of either in B12 binding has been shown. The two clades of the phylogenetic BchE tree (fig. 3*E*) can be divided by the position of this residue

Chlorobaculum tepidum has seven BchE homologous genes, whereby only three (BchE, BchR, and BchQ) are involved in bacteriochlorophyll biosynthesis (Gomez Maqueo Chew et al. 2007). BchR and BchQ act as methyl transferases performing specific methylations in the later steps of Bacteriochlorophyll c pathway (Gomez Maqueo Chew et al. 2007) and probably arose from later lineage-specific duplication events. These paralogs and all BchE-related genes with higher sequence similar to homologs from nonphotosynthetic organisms were removed from the phylogenetic analysis.

The BchE tree reveals two well-supported major clades, one containing the *Chlorobaculum tepidum* and *R. capsulatus* sequences and another where cyanobacterial homologs and proteobacterial BchE isoforms are sister groups (fig. *3E*). Within the first clade, there is a clear separation between proteobacteria sequences from the other taxa. In this tree, the sequences belonging to heliobacteria and acidobacteria cluster between chloroflexi and chlorobia.

AcsF is the enzyme from the O₂-dependent pathway. It is present in cyanobacteria, photosynthetic eukaryotes, and chloroflexi, as well as in some photosynthetic proteobacteria. This oxygen-dependent enzyme most likely had its origin within the cyanobacteria and was vertically inherited and horizontally transferred to other photosynthetic organisms. Until the discovery of photosynthesis in Acidobacteria, this enzyme was only present in RCII-containing organisms. The presence of an O₂-dependent enzyme in an RCI phototroph is interesting due to the usual oxygen sensitivity of the iron-sulfur centers of RCI (Tsukatani et al. 2012). However, the C. Chloracidobacterium thermophilum RCI center is not oxygen sensitive, probably because of adaptation of the organism to the aerobic lifestyle (Tsukatani et al. 2012). The AcsF phylogenetic tree can be divided into two major clades, both containing cyanobacterial sequences (fig. 3F). Nostocales and Cyanothece taxa have two AcsF isoforms named here AcsF1 and AcsF2. At first sight, the dichotomy of the tree might seem to be a reflection of this duplication event. Nevertheless, none of the clades contain sequences from all represented cyanobacteria organisms. Specifically, the major branch contains AcsF1 from all photosynthetic phyla chosen, but no Prochlorococcus sequences are present. These cluster in the lower branch of the tree, with the other cyanobacterial AcsF2 isoforms. This could suggest that the oxygenic photosynthetic ancestor (or at least the extinct ancestor of nostocales and Cyanothece and possibly Prochlorococcus) had two AcsF isoforms, which were selectively lost in some organisms, with the AcsF1 isoform having been laterally transferred to

GBE

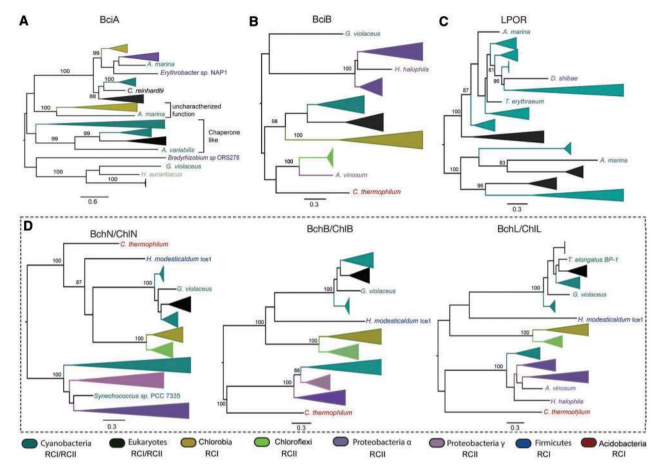


Fig. 4.—Phylogenetic relationships among photosynthetic bacteria based on the different (bacterio)chlorophyll genes. (A) BciA, (B) BciB, (C) LPOR, and (D) DPOR with BchN/ChlN, BchB/ChlB, and BchL/ChlL gene trees.

other photosynthetic lineages. The *C. Chloracidobacterium* thermophilum sequence is basal to the group of noncyanobacterial AcsF sequences and has higher sequence similarity with the chloroflexi clade.

(Di)Vinyl Reductases

of Prochlorococcus With exception some strains (Chisholm et al. 1992) and heliobacteria (bacteriochlorophyll q), (bacterio)chlorophylls have an ethyl group at position 8 of ring II. The reduction of the 8-vinyl group by a 8-vinyl reductase enzyme may occur at various steps of the chlorophyll biogenesis pathway (Tripathy and Rebeiz 1988; Kim and Rebeiz 1996; Adra and Rebeiz 1998; Kolossov and Rebeiz 2001) either in a sequential or in a parallel route (Parham and Rebeiz 1995; Nagata et al. 2005), thereby generating some of the observed heterogeneity of chlorophylls and their intermediates. For clarity, here we show a sequential representation, keeping the other possible alternatives in mind.

There are at least two unrelated enzymes that catalyze this reaction: BciA and BciB (or slr1923) (Islam et al. 2008; Ito et al.

2008; Bryant et al. 2012). BciA (fig. 2A, step 5) belongs to the short-chain dehydrogenases/reductases family having a ligand-binding site and a NAD(P)+/NAD(P)H-binding site. It is present in some organisms belonging to chlorobia, cyanobacteria, eukaryotes, and proteobacteria. Mutagenesis and biochemical experiments identified the *BciA* gene in *Chlorobaculum tepidum*, *A. thaliana*, and *Oryza sativa* (Nagata et al. 2005; Nakanishi et al. 2005; Chew and Bryant 2007; Wang et al. 2010). As seen in figure 4A, these orthologo genes form a well-supported clade that can be subdivided into two groups, one comprising GSB and proteobacteria sequences and another with cyanobacteria and eukaryotic sequences. The only exception is the *Acaryochloris marina* sequence that clusters within the proteobacterial clade.

Because of the broad distribution of NAD(P)H-binding sites (Rossman folds) in proteins, database searching retrieved many additional hits (lower part of the tree). BciA shares sequence similarity with a proposed chaperone for quinone binding in PSII (Ermakova-Gerdes and Vermaas 1999) and also with some sequences of unknown function among



chlorobia. Interestingly, another *Aca. marina* isoform is positioned with the functionally undetermined chlorobial sequences. To root the tree, a group of five sequences belonging to photosynthetic and nonphotosynthetic organisms was used.

The *BciB* gene (fig. 2*B*, step 6) is present in all phyla considered here, and, with exception of gamma-proteobacteria, the groups are all monophyletic (fig. 4*B*). BciB has high homology with coenzyme F420-reducing hydrogenase only found in methanogenic archaea (Islam et al. 2008). The *G. violaceus* sequence, basal to the proteobacterial clade, represents a second copy (an isoform) of the *BciB* gene in the *Gloeobacter* genome and was probably a later acquisition via lateral transfer from proteobacteria. The BciB tree is one of the few cases where chloroflexi and chlorobia are not sister groups, with chloroflexi closer to *Allochromatium vinosum* and *C. Chloracidobacterium thermophilum* sequences. Also, BciB shows separation between the major proteobacterial clade and the other photosynthetic taxa.

Protochlorophyllide Reductases

The last step of the common metabolic pathway of (bacterio)chlorophyll consists of the reduction of the IV pyrrole ring of protochlorophyllide to form chlorophyllide. Two distinct enzymes have evolved to catalyze this reaction, the LPOR and the light-independent protochlorophyllide reductase (DPOR) composed of three different subunits, BchL/ChlL, BchN/ChlN. and BchB/ChlB.

LPOR (fig. 2A, step 7) belongs to the short chain alcohol dehydrogenases family and is present in oxygenic photosynthetic organisms (Dahlin et al. 1999). This enzyme most likely originated in cyanobacteria and was acquired by eukaryotes at the origin of plastids (Yang and Cheng 2004). Because of its involvement in the development of etiolated and greening tissues (Fujita 1996), LPOR from different organisms has been studied extensively in terms of mechanism, function, localization, and expression (summarized in Reinbothe et al. [2010]). Several isoforms of this nuclear encoded enzyme are present in eukaryotic phototrophs (named PorA, PorB, and PorC in A. thaliana) with differential expression and function along A. thaliana development (Masuda et al. 2003). Besides the isoforms present in eukaryotes, additional putative LPORs belonging to cyanobacteria and Dinoroseobacter shibae DFL 12 were retrieved via database searching. The Dinoroseobacter gene clusters within the Synechococcus clade and is probably a recent gene acquisition. The tree in figure 4C can be divided into two major clades, each one containing cyanobacterial and eukaryotic homologs. The upper part of the tree contains biochemically characterized LPORs. The several PorA, PorB, and PorC enzymes found in angiosperms cluster together, indicating duplication and functional or regulatory specialization during plant evolution.

The lower part of the LPOR tree contains cyanobacterial isoforms and eukaryotic isoforms, the latter, so far, uncharacterized. Curiously, in this subtree, angiosperm (*Vitis vinifera* and *Oryza sativa*) and algal sequences (*Micromonas* sp. and *Ostreococcus lucimarinus*) form two statistical significant groups, the former clustering with *Aca. marina* and *Cyanothece* sequences and the later with marine cyanobacteria (*Synechococcus* and *Prochlorococcus*). It is possible that these two major LPOR isoforms existed before the origin of plastids and were acquired then, with subsequent differential loss. This suggestion is in agreement with the presence of multiple LPOR isoforms in modern cyanobacteria.

The DPOR (fig. 2A, step 8) is present in all photosynthetic phyla, having been lost in most of the angiosperm species (Fong and Archibald 2008). Its three subunits, (B)ChlLNB, have homology with a second chlorin reductase complex BchXYZ present only in bacteriochlorophyll-containing organisms. This complex is responsible for the reduction of the C7–C8 bond, converting chlorophyllide into bacterochlorophyllide. (B)ChlLNB also has homology with the nitrogenase complex (nifHDK) responsible for nitrogen fixation in some bacterial and archaeal taxa (Leigh 2000; Dedysh et al. 2004; Raymond et al. 2004). The nifHDK complex is proposed to have an ancient origin and to be at the base of the (B)ChlLNB complex (Dorr et al. 2003; Martin 2012).

Two groups of cyanobacteria are observed in (B)ChlLNB trees (fig. 4D). Sequences from marine cyanobacteria (Synechococcus and Prochlorococcus) cluster with proteobacterial homologs, whereas the remaining cyanobacterial sequences and the two eukaryotes known to have (B)ChlLNB (Physcomitrella patens and Chlamydomonas reinhardtii) cluster next to the chlorobia/chloroflexi clades. Events of LGT between cyanobacteria and proteobacteria are well documented (Badger and Price 2003; Beiko et al. 2005; Zhaxybayeva et al. 2006), especially for the closely related Synechococcus and Prochlorococcus genera. However, the polarity of this LGT (who was donor, who was recipient) is still not established (Bryant et al. 2012; Gupta 2012). In the different DPOR trees, the heliobacterial sequences align either with nonmarine cyanobacteria or with the chlorobia/cloroflexi group, whereas the acidobacterial sequence is basal either to the proteobacterial clade (BchB/ChlB and BchL/ChlL) or to the Heliobacterium-containing clade (BchN/ChIN).

Discussion

Individually, the trees for proteins underlying chlorophyll biosynthesis are complex. Although they do not all tend strongly to reflect a single underlying topology, they do have aspects in common. Their main implication in the context of this article is as follows: taken as a whole, the trees for chlorophyll biosynthesis appear to distinguish between competing hypotheses for the presence of two serially linked photosystems at the



origin of water-splitting photosynthesis. This behavior is statistically supported by the AU and SH tests, with rejection of trees where separation of RCI and RCII organisms was imposed.

Two Cyanobacterial RCs: Gene Duplication, Not Lineage Merger

The most important observation from this study is that there is no coevolutionary pattern linking chlorophyll biosynthesis gene phylogeny with either RCIs or RCIIs; in other words, we observe neither Type I- or Type II-specific chlorophyll biosynthesis genes nor Type I- or Type II-specific chlorophyll biosynthesis gene phylogenies. Because RCs cannot undergo evolution in the absence of chlorophyll, this lack of coevolutionary pattern linking chlorophyll biosynthesis to the divergence of RCI and RCII allows us to exclude the widely discussed possibility that RCI and RCII diverged via lineage splitting and became reunited in cyanobacteria via a largescale gene transfer event (Mathis 1990; Blankenship 1992; Meyer 1994; Blankenship and Hartman 1998; Xiong et al. 1998, 2000; Blankenship 2001; Xiong and Bauer 2002a, 2002b; Blankenship 2010), of the kind that Hohmann-Marriott and Blankenship (2011) call "fusion theories." Indeed, the consistently close proximity of the GSB (chlorobia, with RCI) and green nonsulfur bacteria (chloroflexi, with RCII) in chlorophyll biosynthesis trees argues strongly against the view that there was a deep evolutionary split in chlorophyll biosynthesis corresponding to a lineage split between RCI and RCII, one that would be expected to have pulled (B)Chl genes in tow. This is all the more true given the tendency for the chlorophyll biosynthesis genes from proteobacteria (with RCII) to branch in close proximity to their cyanobacterial homologs, rather than with homologs from green nonsulfur bacteria. In addition, the chlorophyll biosynthesis genes of C. Chloracidobacterium thermophylum (with RCI) usually branch in close proximity to RCII-containing taxa (proteobacteria or chloroflexi).

If we thus exclude ancient lineage splitting and later (re)union in cyanobacteria at the evolutionary origin of two RC types coexisting in the same cell, the simplest competing alternative, and one widely discussed in the literature (Olson and Pierson 1987a; Vermaas 1994; Olson 2001; Allen 2005; Mulkidjanian et al. 2006), is that gene duplication giving rise to RCI and RCII within the same genome gave rise to the photosystem configuration in oxygenic photosynthesis. This possibility is easily reconciled with chlorophyll biosynthesis phylogenies. An immediately ensuing question is: in which lineage did the gene duplication take place? Occam's razor clearly favors the premise that the photosystem genes underwent duplication in an ancestral cyanobacterium—a protocyanobacterium—because cyanobacteria are the only group where genes for both RC types have remained present and

expressed. It is possible to assume that the duplication took place elsewhere, but there is no obvious alternative location.

What Use Might Two Photosystems Be?

Olson and Pierson (1987a, 1987b) have suggested an ancient gene duplication event in a cyanobacterium for the origin of the two photosystems. However, that formulation accounted for the distribution of chlorophyll-based photosynthesis and RC types among prokaryotic groups exclusively by vertical inheritance and differential loss. Today, with the number of bacterial lineages having grown, the distribution of photosynthesis and RCs (fig. 1) has become much more sparse than in 1987, such that differential loss alone is unlikely to account in full for these distributions, especially because photosynthesis genes are observed to be mobile in the marine phage metagenome, and because substantial amounts of LGT have indeed been shown for photosynthetic lineages (Raymond et al. 2002; Huang and Gogarten 2007; Shi and Falkowski 2008). Hence, a mechanistic mixture of some vertical inheritance, some differential loss, and some lateral transfer, relative amounts of which might differ from gene to gene across (B)Chl synthesis, needs to be invoked, because the trees for (B)Chl biosynthesis genes are insufficiently similar for artifacts of phylogenetic reconstruction to account for their differences.

However, how much and what kind of LGT might be required to explain the distribution of photosynthesis among prokaryotes? Various possibilities have recently been discussed, for example, by Bryant et al. (2012) and Gupta (2012). However, that question is not the focus of this article. Rather, the question at our focus is how two photosystems came to reside within a single genome so as to give rise to oxygenic photosynthesis. Gene duplication in a protocyanobacterium is the alternative most compatible with present data. If there were two photosystems, what were they doing?

This question has to do with the transition from anoxygenic photosynthesis to water splitting. Blankenship and Hartman (1998) suggested that hydrogen peroxide (H_2O_2) might have been an initial electron donor far more chemically accessible than water, for a linear two-photosystem ETC. Mulkidjanian et al. (2006) suggested that H_2 might have been the initial electron donor for a two-photosystem ETC. Nisbet and coworkers (Nisbet et al. 1995; Nisbet and Sleep 2001) suggest that the manganese complex evolved either to handle excess of peroxide or as a toxic weapon against competitors.

An alternative suggestion (Mathis 1990; Allen and Puthiyaveetil 2005) differs from the foregoing models with respect to the presumed function of the two photosystems. It posits that the two photosystems in the protocyanobacterium operated in a temporally regulated manner (Allen 2005), for example, as an H₂S-oxidizing and NAD⁺-reducing RCI when H₂S was available as it occurs in modern *Chlorobium* or as a light-driven proton pump (cyclic electron flow



thorough RCII as in *Rhodobacter*) when H₂S was not available. Before the water-splitting complex had evolved, what would a bacterium with two different and specialized, nonoxygenic photosystems have done with them? Probably just what modern bacteria do: express them when needed, with the help of a regulatory switch. This model implies relatively strict regulation of the RC genes, because in the event that regulation failed, for example, through mutation of the regulatory protein constituting the redox switch, both RCIs and RCIIs became expressed in the absence of H₂S, the protocyanobacterium would be exposed to a lethal level of oxidative stress (there is no way to turn off assembled photosystems), unless it could extract electrons from an environmentally available donor

In principle, such a donor could have been H₂O₂, or H₂, or an organic compound such as succinate. However, it also could have been aqueous Mn^{IVIII} cations, which have the interesting property of giving off low potential electrons under ultraviolet (UV) radiation (photooxidation) (Anbar and Holland 1992; Hakala et al. 2006; Allen and Martin 2007; Russell et al. 2008). Before the accumulation of atmospheric oxygen (and hence ozone), UV was a larger component of the solar radiation that reaches the Earth's surface than it is today. Thus, the presence of light for photosynthesis would have also meant the presence of UV radiation for Mn^{II/III} photooxidation. Photooxidation of aqueous, environmental Mn^{II/III} could have thus literally "pushed" electrons into photooxidized chlorophyll of RCII. That would have led to a lethal log-jam of electrons, or over-reduction, in the electron-transport cycle unless the protocyanobacterium was simultaneously expressing RCI. This complex would redirect the surplus membrane-bound electrons stemming from photosystem II into CO₂-reduction and thereby create precisely the flow of electrons seen in cyanobacteria today: a linear flow through two photosystems.

Obviously, such an ETC would hardly have been perfect from the beginning. And clearly there is a difference between tapping environmental Mn atoms photooxidatively, one at a time, and the establishment of the resident, catalytic Mn₄Ca center in cyanobacterial RCII water-splitting complex. However, the overall contours of the circuit would have been right, from manganese at RCII and going on through RCI to NADH and ultimately to CO₂. The key to water splitting would then have entailed the transition from exploiting an environmental supply of soluble manganese, where each electron-donating Mn ion would reach photosystem II by simple diffusion, to holding four manganese atoms (and a calcium of as-yet-unknown function) in place. Dissolved Mn²⁺ has long been known to donate electrons to photosystem II, and thus to reconstitute noncyclic electron transport, in isolated chloroplasts biochemically depleted of the capacity to oxidize water to oxygen (Cheniae 1970). Following adsorption of Mn to its donor side, a fine tuning of photosystem II by natural selection to optimize its reduction/oxidation potential could have allowed it to oxidize a now biologically portable manganese reservoir four times in a row.

It is notable that the Mn atoms of the water-splitting complex are bound directly by the proteins of the photosystem II RC, without an intervening protein or electron carrier. This suggests that no major evolutionary invention was required for photosystem II to tap environmental Mn as an electron source. In line with that, it has recently been shown (Allen et al. 2012) that an engineered, Mn-binding RCII of *R. sphaeroides* will produce O_2 from O_2^- in the presence of Mn in a light-dependent reaction in which photodamage is impeded in comparison with that in a wild-type, Mn-free RC. Allen et al. (2012) interpret this observation as an important clue to the origin of oxygenic photosynthesis.

In what sort of setting could suitably high concentrations of Mn^{I/III} have accumulated for such series of events to transpire? Mn is not a good candidate for high enough concentrations to make the model viable in an open ocean setting. However, in a locally circumscribed freshwater setting, sufficiently high Mn^{I/III} concentrations could, in principle, have accumulated.

In this context, it is of particular interest that Blank and Sáchez-Baracaldo (2010) recently provided evidence that oxygenic photosynthesis arose in a freshwater environment, based on the basal phylogenetic position of freshwater cyanobacteria and the derived phylogenetic position of marine cyanobacteria.

Related Pathways and the Cell That Invented Chlorophyll

Chlorophylls coordinate Mg²⁺ and belong to the tetrapyrrole family, which includes cobalamin (Co²⁺), heme (Fe²⁺), sirohem (Fe²⁺), heme d1 (Fe²⁺), and F_{430} (Ni²⁺) (Heinemann et al. 2008; Warren and Smith 2009; Zappa et al. 2010). All tetrapyrrole biosynthetic pathways are related in the sense that they start from the universal precursor, δ -aminolevulinic acid, and share three enzymatic steps that generate the tetrapyrrole macrocycle uroporphyrinogen III (UroIII) (fig. 5). Proto IX is the common precursor for chlorophyll synthesis and heme synthesis via the classical pathway that occurs in eubacteria and eukaryotes (Heinemann et al. 2008). Precorrin-2 is the common precursor for cobalamin, heme synthesis via the alternative pathway recently discovered in archaebacteria (Storbeck et al. 2010; Bali et al. 2011), siroheme (a cofactor in some nitrite and all sulfite reductases (Tripathy et al. 2010), heme d1 (a cofactor only present in the bacterial cd_1 nitrite reductases (Allen et al. 2005), and F₄₃₀ pathways. F₄₃₀ is a cofactor that is critical to methanogenesis and has only been found in methanogens so far (Thauer 1998). In addition to two routes for heme synthesis, there are also two routes for cobalamin synthesis, one O₂ dependent (often called the late pathway due to the late insertion of Mg²⁺) and one O₂ independent (early insertion of Mg²⁺) (Martens et al. 2002; Warren et al. 2002). The O2-dependent cobalamin pathway involves the Co²⁺ chelatase CobNST, the three subunits of

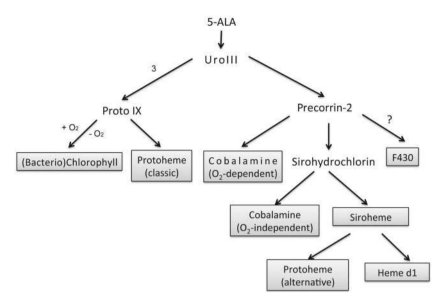


Fig. 5.—Schematic representation of the bifurcations present in the global tetrapyrrole pathway. The biosynthetic pathway of all tetrapyrroles begins with the condensation of eight molecules of 5-amino-levulinate to uroporphyrinogen-II (UroIII), the first cyclic tetrapyrrole. This compound can be sequentially converted to protoporphyrin IX in three enzymatic steps. This is the last common precursor for (bacterio)chlorophyll synthesis and heme synthesis via the classical pathway (present in most eubacteria and eukaryotes). Alternatively, Urolll can be converted to precorrin-2, the branching point of F430 synthesis, of the O₂-dependent cobalamin pathway, and of the formation of Sirohydrochlorin. At the level of Sirohydrochlorin, the O₂-independent cobalamin pathways diverge from the formation of siroheme. Siroheme may be used as a cofactor, further transformed into heme d1 or transformed into heme according to the alternative heme pathway present in archaea and some eubacteria.

which are related to (B)ChlHID. It has been suggested that (B)ChlHID arose from CobNST (Xiong et al. 2000), but the converse might be more likely given the presence of one O₂-dependent step in the late cobalamin pathway (Martens et al. 2002; McGoldrick et al. 2005). In the early (O2 independent) cobalamin pathway, the CbiK/CbiX^L/CbiX^S chelatases are related to the SirB and HemH chelatases of the siroheme and classical heme pathways, respectively (Schubert et al. 1999; Brindley et al. 2003; Romao et al. 2011). The chelatase for the F₄₃₀ pathway is unknown, but genes related to (B)ChIHID are present in methanogens and encode candidates for the Ni²⁺chelatase. In some organisms, the insertion of iron and cobalt into sirohydrochlorin is performed by multifunctional chelatases (CysG and Met8p) (Spencer et al. 1993; Fazzio and Roth 1996; Schubert et al. 2002) (Class III).

What did the cell that invented chlorophyll biosynthesis have in terms of tetrapyrrole pathways? It definitely contained cobalamin and might have had a cobalamin biosynthesis pathway (the O₂-independent type, obviously), because the O₂independent route to 3,8-divinyl protochlorophyllide a via (B)ChlE, a cobalamin-dependent reaction (Fuhrmann et al. 1993; Gough et al. 2000). Of course, it could also have just been cobalamin dependent; some cyanobacteria synthesize cobalamin, and many, however, acquire it from the environment via high affinity importers (Tang et al. 2012). The cell that invented chlorophyll also had a classical heme pathway, because barring chlorophyll, the three steps from Urolll to Proto IX are specific to classical heme synthesis (Heinemann et al. 2008) and hence preceded the (B)Chl pathway. The first cell with chlorophyll probably also fixed nitrogen, because the subunit of DPOR, (B)ChlNBL, is related to nitrogenase subunits (Xiong et al. 1998). We propose that the cell that invented chlorophyll was a cobalamin-dependent, heme-synthesizing, diazotrophic anaerobe.

Conclusions

The phylogeny and evolution of the chlorophyll biosynthesis core pathway were analyzed both at the gene and pathway level. The lack of coevolution of chlorophyll biosynthesis genes with RCIs or RCIIs permits us to exclude the widely discussed possibility that RCI and RCII diverged via lineage splitting and became reunited in cyanobacteria via a largescale gene transfer (fusion) event. Moreover, it can be concluded that the primordial photosynthetic organism performed nitrogen fixation, synthesized heme, and was cobalamin dependent.

Note Added in Proof

In support of the genetic mobility of anoxygenic photosysnthesis, photosynthesis is encoded on a plasmid in Roseobacter littoralis (Kalhöfer et al. 2012).



Supplementary Material

Supplementary figures S1–S12 and tables S1 and S2 are available at *Genome Biology and Evolution* online (http://www.gbe.oxfordjournals.org/).

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