Subcellular Localization of Monogalactosyldiacylglycerol Synthase in *Synechocystis* sp. PCC6803 and Its Unique Regulation by Lipid Environment

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Abstract

Synthesis of monogalactosyldiacylglycerol (GalDAG) and digalactosyldiacylglycerol (GalGalDAG), the major membrane lipids in cyanobacteria, begins with production of the intermediate precursor monoglucosyldiacylglycerol (GlcDAG), by monoglucosyldiacylglycerol synthase (MGS). In *Synechocystis* sp. PCC6803 (*Synechocystis*) this activity is catalyzed by an integral membrane protein, Sll1377 or MgdA. In *silo* sequence analysis revealed that cyanobacterial homologues of MgdA are highly conserved and comprise a distinct group of lipid glycosyltransferases. Global regulation of lipid synthesis in *Synechocystis* and, more specifically, the influence of the lipid environment on MgdA activity have not yet been fully elucidated. Therefore, we purified membrane subfractions from this organism and assayed MGS activity in *vivo*, with and without different lipids and other potential effectors. Sulfoquinovosyldiacylglycerol (SQDAG) potently stimulates MgdA activity, in contrast to other enzymes of a similar nature, which are activated by phosphatidylglycerol instead. Moreover, the final products of galactolipid synthesis, GalDAG and GalGalDAG, inhibited this activity. Western blotting revealed the presence of MgdA both in plasma and thylakoid membranes, with a high specific level of the MgdA protein in the plasma membrane but highest MGS activity in the thylakoid membrane. This discrepancy in the subcellular localization of enzyme activity and protein may indicate the presence of either an unknown regulator and/or an as yet unidentified MGS-type enzyme. Furthermore, the stimulation of MgdA activity by SQDAG observed here provides a new insight into regulation of the biogenesis of both sulfolipids and galactolipids in cyanobacteria.

Introduction

Proposed by the endosymbiotic theory to be the ancestors of plant chloroplasts, cyanobacteria are commonly used to investigate the biogenesis of the thylakoid membrane (TM). Two major ideas currently prevail, one considering cyanobacterial TM to be a physically separate entity and the other stating that this membrane has either permanent or transient contact with the plasma membrane (PM) [1]. Although much work on the synthesis of both proteins and pigments of the photosystems and their regulation has been performed with the unicellular cyanobacterium *Synechocystis* sp. PCC6803 (hereafter referred to as *Synechocystis*) [2–4], much less is known about the lipid components. If the TM and PM have no connection, both membranes might synthesize their own lipids. Alternatively, vesicular lipid transport, such as in chloroplasts [5], could occur between these membranes, although there is at present no direct evidence for this theory. On the other hand, if the membranes are connected, lipids synthesized by either or both could flow between them. Accordingly, a better understanding of lipid synthesis and its regulation in cyanobacteria is central to understanding the process of thylakoid biogenesis.

Although the major lipid constituents of membranes in most non-photosynthetic organisms are phospholipids, the major membrane lipids of chloroplasts and cyanobacteria do not contain a phosphorus (P) atom, possibly because both plants and many bacterial species have limited access to inorganic phosphorus (P) [6–9]. Monoglucosyldiacylglycerol (GalDAG: 1,2-diacyl-3-O-[β-D-galactopyranosyl)-(sn-)glycerol), one of the most abundant membrane lipids in the biosphere, often constitutes 50 mol% or even more of the total lipid content of cellular (or chloroplast) membranes in photosynthetic organisms [10,11]. Not only does GalDAG play an important structural role in membranes but it is also vital for the correct functioning of the photosynthetic complexes and pathways [12]. The pathways by which GalDAG is synthesized in plant chloroplasts and cyanobacteria are quite distinct. In chloroplasts, MGD1 catalyses the reaction of sn-1,2-diacylglycerol with UDP-activated galactose (UDP-1-α-galactose) [13] whereas in cyanobacteria diacylglycerol is first combined with UDP-glucose by monoglucosyldiacylglycerol synthase (or MGS) to yield the precursor monoglucosyldiacylglycerol (GlcDAG: 1,2-diacyl-3-O-[β-D-glucopyranosyl)-(sn-)glycerol) [14]. In *Synechocystis* MGS is encoded by the gene *sll1377* or *mgdA* [15,16] and deletion of this
gene renders cells non-viable [15]. GlcDAG does not accumulate in *Synechocystis* cells under normal growth conditions, since its glucose moiety is rapidly converted into galactose by an as yet uncharacterized epimerase, producing GalGalDAG [14]. GalDAG is the acceptor substrate for formation of the other major galacto-lipid, digalactosyldiacylglycerol (GalGalDG: 1,2-diacyl-3-[α-D-galactopyranosyl-(1→6)-O-[β-D-galactopyranosyl]-]in-glycerol) through a pathway thought to be conserved from cyanobacteria to plants [17].

The various homologues of MGS can potentially be regulated both transcriptionally and posttranslationally, but the lipid environment is a major regulatory factor of this enzymatic activity. In organisms such as *Acholeplasma laidlawii*, *Streptococcus pneumoniae*, *Mycoplasma pneumoniae* and *M. genitalium* GlcDAG synthesis is dependent on the presence of an activator lipid, the anionic phosphatidylglycerol (PG) [18–21] Enzymatic activity is stimulated further by the presence of small amounts (up to 10 mol%) of phosphatidylethanolamine (PE), possibly due to anionic phosphatidylglycerol (PG) [18–21] Enzymatic activity is stimulated further by the presence of small amounts (up to 10 mol%) of phosphatidylethanolamine (PE), possibly due to the activation of an activator lipid, the anionic phosphatidylglycerol (PG) [18–21].

**Materials and Methods**

**Media and Cell Growth**

*Synechocystis* sp. 6803 was grown in glass bottles with 1 L BG11 medium [23] at 30°C, bubbling with sterile air and at a light intensity of 50 μE sec⁻¹ m⁻². Cell cultures with an OD₇₃₀ 1–1.5 were harvested by centrifugation (8,000 g, 4°C, 15 minutes), washed once in cold 20 mM potassium phosphate buffer (pH 7.5) and cell pellets were stored at −80°C. *E. coli* C41 (DE3) was routinely grown in LB medium (liquid or solid) for cloning and in 2× LB medium for enzyme overexpression, both supplemented with 50 μg mL⁻¹ kanamycin.

**Fractionation of *Synechocystis* Membranes**

The pellets obtained as described above were washed once again with chilled lysis buffer (20 mM potassium phosphate buffer, pH 7.5), pelleted and resuspended in 1 mL lysis buffer containing protease inhibitors (Complete, EDTA-free, Roche). This suspension was then lysed by vortexing vigorously in the presence of 0.17–0.18 mm glass beads (Sartorius). Sucrose gradients were performed and aqueous two-phase partitioning carried out according to previously established protocols [24], with these modifications. Two-phase systems were incubated on ice for 15 minutes between centrifugations and thoroughly mixed by inversion in closed ice box. This procedure yields “light” (low-density) plasma membrane (PM1, less than 200 μg protein/L culture with an OD₇₃₀ = 1) and “heavy” (high-density) plasma membrane (PM2, roughly 2–3 mg protein/L culture), as well as purified TM (approximately 20 mg protein/L culture). PM1 separates in an orange band between 10 and 30% (w/w) borders in the sucrose gradient, while PM2 and TM are isolated from the 38–42% (w/w) region of this gradient using aqueous two-phase partitioning [24]. Protein was quantified by the Peterson procedure [25] and samples stored at −80°C for further use.

**Protein Electrophoresis and Western Blotting**

To assess purity, 5 μg protein from each fraction was solubilized in SDS-PAGE sample buffer supplemented with 7M urea, incubated for 30 minutes at room temperature and electrophoresed in 12% TGE precast SDS-PAGE gels (BioRad). The gels were then transferred to PVDF membranes, which were subsequently probed with antibodies for characteristic TM (PsAA and CP43, Agrisera) and PM proteins (KtrE, obtained from Nobuyuki Uozumi and PixJ1, peptide antibody [4]), as well as for MgdA (a kind gift from Mie Shimojima). The blots were developed using an anti-rabbit antibody conjugated with horseradish peroxidase (GE Healthcare), enhanced chemiluminescence (ECL Prime, GE Healthcare) and a CCD camera (LAS-4000 Fujifilm), in the “Precision” mode and with automatic exposure time. For comparison of the migration of the bands detected to that of the prestained markers (PageRuler Plus, Thermo Scientific), blot images were merged with the MultiGauge software (v. 3.2, Fujifilm). Each individual blot was performed in triplicate.

**Cloning MgdA Variants**

Truncated MgdA constructs were generated in order to examine the function of each individual transmembrane region. All primers consisted of a gene-specific annealing sequence of approximately 20 bp and common overhangs containing the recombination sites (see Table S1). The major vector employed was pNIC28-Bsa4 (GenBank accession EF198106), developed by Posher Gileadi at SGC Oxford. All targets were fused with an N- or C-terminal hexahistidine tag to simplify purification, as well as a cleavage site for TEV protease to facilitate removal of this tag.

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**Figure 1. Western blotting analysis of membrane subfractions isolated from *Synechocystis*.** Each gel lane was loaded with 5 μg membrane protein and probed with the antibodies indicated. PM1– low-density plasma membrane; PM2– high-density plasma membrane; TM – thylakoid membrane. All individual blots were performed in triplicate, using independent samples. For further details, please see the Materials and Methods. doi:10.1371/journal.pone.0088153.g001
Expression of MgdA Variants in *E. coli*

For expression of MgdA variants, single colonies from LB-agar plates were grown overnight in 5 mL 2× LB medium supplemented with 50 μg/mL kanamycin at 37°C, shaking at 200 RPM. The following morning, 2 mL of the starter culture was inoculated into 100 mL fresh 2× LB medium containing antibiotic and grown for another 2 hours, after which the temperature was reduced to 22°C and the cultures incubated for an additional 60 minutes. Expression was induced by addition of 1 mM isopropyl β-D-1-thiogalactopyranoside [IPTG] and the cells allowed to grow for 22 hours at 22°C with shaking. Cultures were harvested by centrifugation (3,000 g, 4°C, 15 minutes), washed in wash buffer (100 mM HEPES with 20 mM MgCl₂, pH 8.0), resuspended to an OD₆₀₀ = 40 in the assay buffer (100 mM HEPES, 20 mM MgCl₂ and 20 mM CHAPS, pH 8.0) and sonicated three times for 5 minutes each in a water-bath sonicator (Fisherbrand) to obtain whole cell lysates, which were stored in aliquots at −80°C until further use.

For large-scale expression of full-length MgdA, 500-mL cultures were induced as described above and total membrane isolated as follows: after washing in wash buffer supplemented with Complete EDTA-Free protease inhibitors (Roche Applied Science), the cells were homogenized (four passes at 12,000 psi), debris removed by slow-speed centrifugation (3,000 g, 4°C, 15 minutes) and the total membrane fraction then obtained by high-speed centrifugation (200,000 g, 4°C, 45 minutes). These fractions were resuspended in storage buffer (wash buffer supplemented with 10% glycerol) and kept in aliquots at −80°C. Protein content were quantified by the Peterson method [25].

**Assay of MGS Activity and Thin-layer Chromatography (TLC)**

MGS activity was assayed as described previously [18], with certain modifications. In brief, either 600 μg total membrane protein or 200 μg protein of the different *Synechocystis* membrane fractions was washed once with wash buffer, centrifuged (250,000 g, 4°C, 20 minutes) and resuspended in 100 μL of assay buffer. These suspensions were subsequently incubated on ice for 30 min, sonicated in a water-bath sonicator (1 minute, 3 times) and combined with 80 μL of mixed lipid micelles (Avanti Polar Lipids) [18], followed by additional 30 minutes of incubation on ice. 10 μL of UDP-[14C]glucose solution (Perkin-Elmer, 0.02 μCi, μL⁻¹, 302 μCi,μmol⁻¹, diluted 1:5) was then added and incubation continued for another 30 minutes at 30°C, after which the reaction was stopped by adding 390 μL chloroform-methanol (1:2) and lipid phases extracted according to Bligh and Dyer [28,29]. Thereafter, these lipid phases were vacuum-dried, redissolved in 40 μL chloroform-methanol (2:1) and loaded onto an aluminum-backed TLC plate (Merck), which was developed with a chloroform-methanol:28% NH₄OH system (78:2:6). After TLC run, the plate was dried and exposed at least 22 hours on a Kodak Imaging Screen K (Kodak) and analyzed with a Typhoon scanner (Typhoon Trio, GE Healthcare) using standard settings for autoradiography. Activity was calculated based on the intensity of the radioactive GlcDAG band in comparison to serial dilutions standard of UDP-[14C] glucose, employing the ImageQuant TL software (v. 7.0, GE Healthcare).

**Pulse-labeling of Newly Synthesized Lipids in *Synechocystis***

*Synechocystis* cultures (at OD₇₃₀ = 1.5) were centrifuged, cell pellets were washed three times with fresh BG11 and then divided into 4 individual cultures in 100 mL E-flasks (final OD₇₃₀ of 15 in 30 mL of BG11). Following 2 hours of incubation at 30°C with air bubbling, acyl chains were labeled by addition of 12 μCi [14C]-acetate (Perkin-Elmer) to each culture (corresponding to a final concentration of 7.6 μM), which can be taken up rapidly by the cells. Then, these mixtures were incubated at 30°C with air bubbling and at the time intervals indicated, labeling was stopped by addition of 1 mL of 1 M sodium acetate (pH = 7.5) to each culture. Cells were transferred to a 50-mL Falcon tube, cooled rapidly in liquid nitrogen to slow down metabolism and centrifuged (15 minutes at 8,000 g, 4°C). The resulting pellets were flash-frozen in liquid nitrogen and stored at −80°C, until preparation of total membrane, PM1, PM2 and TM fractions as described above. 5,000 CPM from the total membrane fraction at each time point and 2,000 CPM from each isolated fraction were subjected to TLC and analyzed as described above.

**In silico Analysis**

Transmembrane helices were predicted using TMHMM 2.0 (www.cbs.dtu.dk/services/TMHMM), TOPCONS (http://octopus.cbr.su.se), ΔG Prediction server (http://dgpred.cbr.su.
Subcellular Localization of MgdA Protein and Activity in Synechocystis

In plant chloroplasts, lipid synthesis occurs exclusively in the two envelope membranes, rather than in the intraorganellar TM. Therefore, formation of the TM should be entirely dependent on lipid transport from these envelopes, most likely in the form of lipid vesicles [30]. To localize synthesis of the galactolipid precursor GlcDAG in Synechocystis, we investigated the distribution of both MgdA protein and activity in the different membranes.

The purity of the Synechocystis membrane fractions isolated by sucrose gradient and two-phase partition [24] was confirmed by western blots (Figure 1). The marker proteins for the TM (CP43 and PsaA) only exist in TM fraction, while PM marker proteins (PixJ1 [4] and KtrE (also known as DgdA [31])) are only present in PM1 and PM2 fractions. In comparison to the unambiguous distribution of these markers, MgdA was detected in both PM1 and PM2, as well as in TM, although the amount present in TM was significantly lower (Fig. 1).

To our surprise, the TM fraction exhibited the highest MGS activity, followed by the PM1 and the PM2 fractions (Fig. 2). Previously, Omata and Murata [32] reported that “lighter”, “cytoplasmic” membranes isolated from Synechococcus on a sucrose gradient demonstrated higher MGS activity than “heavier”

membranes, which is consistent with our present findings, since the PM1 fraction should correspond to their lighter fraction. Moreover, using our aqueous two-phase system we were able to further separate their “heavy”, “thylakoid” fraction into the TM and PM2 fractions (as shown in [24]) and found that the highest specific activity was present in TM (Fig. 2). Accordingly, the lower specific activity observed by Omata and Murata for the “thylakoid membrane” fractions [32] may be due to the simultaneous presence of PM2 (with low specific activity) in the same fraction as TM.

Localization of de novo Lipid Synthesis in Synechocystis Membranes

When examining de novo lipid synthesis in Synechocystis membranes by labeling acyl chains with $^{14}$C-acetate, it was necessary to label for at least 10 minutes (Figure S1). Within 10 minutes, selectively labeled GlcDAG had already accumulated in the total membrane fraction; during the following 3 hours, over 90% of this labeled precursor was converted into GalDAG, while, surprisingly, labeling of GalGalDAG required more than 24 hours (Fig. 3A). The relatively rapid biosynthesis of GlcDAG and its conversion to GalDAG highlight the indispensability of these lipids in cyanobacteria. It was shown previously that deletion of the enzyme responsible for synthesis of GalGalDAG (DgdA) exerts only minor effects on cell growth and photosynthesis reactions [17] and its slow rate of synthesis here also supports a less critical role for this lipid in Synechocystis. The non-sugar lipids PG and SQDAG were also labeled, even at the earliest time-points, and the level of this labeling appeared to remain rather constant.
In experiments designed to further clarify the localization of \textit{GlcDAG} synthesis, purified PM (PM1 and PM2) and TM fractions from the pulse-labeled cells were all found to contain radio-labeled \textit{GlcDAG} even at the first time-point (10 minutes) tested after addition of \([^{14}\text{C}]\)-acetate (Fig. 3B). This observation is consistent with the presence of MGS activity in all these membranes (Fig. 2). Furthermore, the PM1, PM2 and TM fractions all epimerized \textit{GlcDAG} to GalDAG. Therefore, it can be concluded that all of these membranes harbor the machinery required to synthesize \textit{GlcDAG} and convert it to GalDAG.

Influence of the Lipid Environment on MGS Activity

TM is the most abundant membrane fraction in most cyanobacterial cells and large amounts of sugar lipids are required as building blocks for its biogenesis. This explains the necessity for higher glucosyltransferase activity in the TM rather than PM2. However, the lower level of MgdA protein in the TM (Fig. 1) raises the question as to how sugar lipid synthesis is regulated in \textit{Synechocystis}. Since the lipid composition of the membrane influences lipid glycosyltransferase (GT) activities in other organisms [18–21,33], we tested this possibility here as well.

Under normal conditions GalDAG and GalGalDAG can account for approximately 50 and 26 mol\% of the total lipid in \textit{Synechocystis} membranes [17]. However, the presence of either of these lipids in our mixed-micelle assay system at any of the concentrations tested lowered MgdA activity (Fig. 4A), with GalDAG exerting a more pronounced effect. We propose that this represents feedback inhibition. In \textit{Synechocystis}, the pathways for sulfolipid and galactolipid synthesis share the same initial substrate (DAG) and must therefore be delicately balanced. Koichiro Awai and co-workers [17] have shown that when GalGalDAG synthesis is prevented (by a \textit{dgdA} knock-out mutation), the amount of SQDAG in the membranes increases by 60\% (from 14.3 to 22.5 mol\%), while the level of GalDAG increases by only 20\%.

Figure 4. MGS activity in the presence of mixed micelles containing different amounts of various lipids and with different concentrations of K\textsubscript{2}SO\textsubscript{4}. The values shown are averages of four independent experiments, using 600 mg total membrane protein from two independent cultures per assay. 100\% activity corresponds to 3.8±0.17 pmol UDP-glucose incorporated h\(^{-1}\) mg total protein\(^{-1}\). A. The influence of GalDAG (◯) and GalGalDAG (□) on MgdA activity. B. The influence of SQDAG (•), PE (■) and PG (○) on MgdA activity. C. The influence of sulfate ions (●) on MGS activity in the presence of 30 mol\% PG. D. Influence of SQDG (△) on recombinant full-length MgdA activity. 100\% activity corresponds to 4.60±0.20 pmol UDP-glucose incorporated h\(^{-1}\) mg total protein\(^{-1}\). doi:10.1371/journal.pone.0088153.g004
Synechocystis but is not found in the inner membranes of this organism, by a mechanism involving surface charge and appears to be the major regulator of lipid synthesis, also been reported. The more detailed discussion will be elucidated in time such pronounced stimulation of a lipid GT by SQDAG has fold at the highest concentration tested (40 mol%). This is the first time such pronounced stimulation of a lipid GT by SQDAG has fold at the highest concentration tested (40 mol%). This is the first such pronounced stimulation of a lipid GT by SQDAG has been reported. The more detailed discussion will be elucidated in the later paragraphs.

Membrane Curvature has Little Effect on MgdA Activity

The ratio between lipids that are prone to form bilayers or not influences membrane curvature and lipid phase equilibria, which in turn affect functional features of membrane proteins [18,35,36]. For example, the activity of lipid GTs is sensitive to membrane curvature and surface charge [18,19,21,33,37]. In addition, in the presence of anionic lipids MGS and DGS in A. laidlawii (alMGS and alDGS) are stimulated by the zwitterionic (neutral) PE and other nonbilayer-prone lipids [18,35,36].

Both GalDAG and GalGalDAG have neutral headgroups, but only the latter forms bilayers [36]. However, addition of either of these two major sugar lipids with different effects on membrane curvature impaired MgdA activity (Fig. 4A). Moreover, although GalDAG and PE both have neutral headgroups and are not prone to form bilayers, PE stimulated MgdA activity somewhat (Fig. 4A and B). Similarly, GalGalDAG and SQDAG, both of which form bilayers, exert opposite effects (Fig. 4A and B). It appears likely that in Synechocystis MgdA activity is not influenced by the bilayer curvature, in contrast to alMGS and alDGS in A. laidlawii [18,35]. We suggest that, as final products of the pathway, GalDAG and GalGalDAG interact with and allosterically regulate MgdA (or a regulatory interacting partner). This would allow the cells to maintain the proper lipid composition of their membranes by reducing production of the precursor GlcDAG when sufficient galactolipids are already present (Fig. 5).

SQDAG is a Potent and Specific Activator of Synechocystis MgdA

Synechocystis MgdA activity was specifically and very potently stimulated by SQDAG, a negatively charged lipid component of cyanobacterial membranes (Fig. 4B). In an attempt to understand the underlying mechanism, we performed an experiment similar to the one previously conducted with the A. laidlawii enzymes [40]. Since the SQDAG headgroup contains a sulfonic acid rather than phosphate moiety, MgdA activity was assayed in the presence of increasing concentrations of sulfate ions and unexpectedly, as shown in Fig. 4C, as little as 20 mM sulfate, in the presence of 30 mol% PG, decreased this activity by 52% and this suppression was even greater at higher concentrations of sulfate. Thus, the stimulation by SQDAG cannot be explained either by its negative surface or sulfate/sulfonic acid moiety.

When mgdA gene was expressed in E. coli, only a small amount of the recombinant enzyme was produced but the bacterial membranes exhibited potent MGS activity that was stimulated by SQDAG in the same manner as above (Fig. 4D). This observation provides strong support for the unique and probably specific interaction of SQDAG with MgdA. SQDAG contains a large polar head with a negative charge located some distance from the hydrophobic portion of the membrane. As sulfate alone did not activate MgdA, activation by SQDAG must be somehow dependent on the nature of the head group and/or of the entire molecule. It is noteworthy that lipids with a similar (sugar) head group, but no negative charge (such as GalDAG and GalGalDAG), were inhibitory (Fig. 4A).

Curiously, synthesis of SQDAG is not an absolute requirement for all cyanobacteria. Synechocystis strains lacking the sqdB gene require exogenous SQDAG for survival, whereas in Synechococcus elongatus sp. PCC7942 (Synechococcus) such a knockout produced no obvious effect on cell growth [39]. This difference might be explained by the recent observation that deletion of the sqdB gene affects DNA replication and cell division in Synechocystis, but not in Synechococcus [40].

We propose that this unique regulation of MgdA activity by SQDAG might be involved in maintaining lipid homeostasis in cyanobacteria, especially under stress conditions. For example, in the absence of inorganic phosphate, both Synechocystis and Synechococcus accumulate SQDAG and GalGalDAG at the expense of PG [41]. Such an exchange of phospholipids for sulfolipids has been suggested to play an important evolutionary role for picocyanobacteria, which must compete with heterotrophic
bacteria for a limited supply of phosphate [42]. The situation is similar in freshwater environments [43–45] and Synechocystis can also take up and incorporate exogenous lipids such as SQDAG [43].

Control of lipid-synthesizing enzymes in both Synechococcus and Synechocystis must be at the post-translational, rather than the transcriptional level, since in neither organism is MgdA among the proteins whose expression is regulated by the SphS-SphR two-component system in response to inorganic phosphate [46,47]. In fact, MgdA can be regulated post-translationally by temperature, both \textit{in vivo} and \textit{in vitro} [16], and, as our present results demonstrate (Fig. 4D), by SQDAG, GalDAG and GalGalDAG. As a freshwater cyanobacterium, Synechocystis is often subjected to restricted and/or fluctuating supplies of phosphate [7] and in such circumstances it may be more efficient to utilize a lipid species that does not contain phosphate, i.e., SQDAG, to regulate lipid biosynthesis and/or as part of an effective rescue mechanism when little phosphate is available. A higher SQDAG concentration would then respond to inadequate levels of galactolipids by activating MgdA and vice-versa (Fig. 5). The level of SQDAG is also related to the availability of inorganic phosphate and, since the amount of galactolipid (namely GalGalDAG) is elevated during phosphate starvation [17], the synthesis of phospho-, sulfo- and galactolipids is probably coordinated in cyanobacteria by regulating the activity of appropriate enzymes.

However, the mechanism underlying this regulation on MgdA by SQDAG remains unknown. Several laboratories have observed direct interaction with or binding of SQDAG to different proteins \textit{in vitro}, e.g., a protein that initiates replication in humans [48] and \textit{E. coli} DNA polymerase I, which is inhibited by SQDAG [49]. In addition, this lipid stimulates protein import through the chloroplast outer envelope mediated by the precursor protein translocon [50]. To determine whether or not SQDAG regulates MgdA through direct interaction will require further experimental investigations.

Figure 6. Conservation of transmembrane helices (TMH) among MGS homologues in cyanobacteria. ClustalW sequence alignments for several MGS homologues, with TMH (as predicted by the OCTOPUS server) highlighted by line boxes. These sequences were selected from a BlastP similarity search utilizing Sll1377 as the template and refer to enzymes from (top to bottom) \textit{Paulinella chromatophora}, Nostoc sp. PCC7120, \textit{Anaabaena variabilis} ATCC29413, \textit{Nostoc punctiforme} PCC73102, \textit{Cyanothece} sp. PCC7424, \textit{Synechococcus} sp. PCC7002 and \textit{Synechocystis} sp. PCC6803.

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Conservation of the Transmembrane Helixes of MgdA among Cyanobacteria

Organisms related evolutionarily often retain specific features in the sequences and structures of their proteins and common regulatory mechanisms. Transmembrane domains are not only an important structural feature of proteins, but can also serve important regulatory roles. We chose typical model cyanobacteria, from single-cell to filamentous and both nitrogen-fixing and non-nitrogen-fixing species, to investigate conservation of sequence and secondary structure. When we analyzed the amino acid sequence of MgdA in silico, the 4 prediction programs employed all identified 5 putative transmembrane helices (TMH) that appear to be a conserved feature of cyanobacterial homologs (Fig. 6). In MgdA these are localized essentially to the N- and C-terminal regions (at residues TMH1 46–66, TMH2 68–88, TMH3 361–381, TMH4 389–409, and TMH5 435–455), flanking the putative catalytic domain and active site that face the cytoplasmic surface, with pronounced sequence similarity and remarkable conservation of these five transmembrane domains. Even in the case of the MGS homologue in the "chromatophore" inclusions of the amoeba Paulinella chromatophora, which are thought to be of cyanobacterial origin and the product of a recent endosymbiotic event [51], there is pronounced sequence similarity, with two TMH in the N-terminal and three in the C-terminal region (Fig. 6).

To evaluate the importance of different TMH segments in Synechocystis experimentally, several deletion variants were expressed in E. coli and assayed for activity. Even though many of these were expressed at a level higher than the full-length protein, none of the truncated constructs exhibited any glucosyltransferase activity, even in the presence of SQDAG or any other lipid activator (Fig. S5). In comparison, the full-length construct produced GlcDAG under the same conditions as those used for MgdA in Synechocystis (Fig. 4D). Thus, the transmembrane helices appear to be important for correct folding and/or enzymatic activity and may be one reason why MgdA responds preferentially to SQDAG, rather than PG, as do lipid GT enzymes without transmembrane domains.

Regulation of the Synthesis of Sugar Lipids in Synechocystis – One Step Closer to the Answer?

Our findings that the MgdA protein and activity are not distributed in the same manner among Synechocystis membranes could have two explanations (Fig. 7). First, an as yet unknown MGS-type enzyme located exclusively in the TM, may be responsible for the high activity observed there. As previously mentioned, deletion of the sll1377 gene is lethal to Synechocystis [15], so such an enzyme, if it exists, is unable to compensate for the loss of MgdA under these circumstances.

A second alternative could be the presence of a stimulator of MgdA activity in PM1 and TM. Such a regulator has already been suggested to enhance MgdA activity and thereby maintain membrane integrity under conditions of heat shock [32]. We attempted to isolate complexes containing MgdA, both by twodimensional clear-native/SDS-PAGE electrophoresis, as well as by co-immunoprecipitation with the antibody against MgdA (Figs. S3 and S4). Regrettably, MgdA appeared to migrate alone in the gels and the antibody failed to pull-down putative MgdA-containing complexes, which may indicate that such a protein regulator does not exist or that its interaction with MgdA is not preserved during use of these approaches.

To examine whether reversible phosphorylation is involved in the regulation of MgdA, prior to addition of the enzymatic...
substrate we treated both PM2 and TM preparations with alkaline phosphatase in the presence of SQDAG. Since both treated and untreated preparations showed similar levels of activity (Fig. S2) this does not seem to be the case. Another possible explanation is that MgdA is regulated by a small molecule, as it-dGMP regulates the GT enzymes PgaC and PgaD that synthesizes the exopolysaccharides in E. coli [53].

The Synechocystis enzyme Slr1508 (DgdA) has GalGalDAG synthase activity in vivo [17] and deletion of the dgdA gene results in cells containing none of this lipid [17]. Strangely, this enzyme has also been reported to be part of a sodium-dependent potassium membrane transporter complex (Ktr) in Synechocystis [31,54] and details concerning its reaction mechanism and the galactose donor are still unclear, since no assay for its in vitro activity is presently available. We detected the DgdA protein exclusively in PM (Fig. 1), implying that GalGalDAG synthase should occur only at this location. This raises the question as to whether GalGalDAG is transported by vesicular traffic or by intermembrane diffusion through connections between the PM and TM, previously postulated to be PM1 [4]. While no consensus has been reached, accumulating evidence for a connection between the PM and TM [1,3,4,55] supports lateral membrane diffusion (as hypothesized in Fig. 7).

Concluding Remarks

Here, our initial observation of the discrepancies between the subcellular localization of protein and enzymatic activity involved in the crucial first step in galactolipid synthesis by Synechocystis motivated us to investigate the regulation of MgdA in greater detail. We have demonstrated that this MgdA has unique regulatory features, in particular the potent activation by sulfolipid SQDAG, which distinguishes it from other GlcDAG or GalDAG synthases. Pulse-labelling experiments with [14C]-acetate, as well as measurements of enzymatic activity revealed that GlcDAG and GalDAG are synthesized in both PM and TM, with highest specific activity in the latter. Whether MgdA is regulated post-translationally, by interaction with a small effector molecule or by a protein partner remains to be discovered. The possible existence of a second MGS in Synechocystis and the interaction of SQDAG with MgdA are still open questions that will require further investigation.

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