**Bacillus subtilis** YngB is a UTP-glucose-1-phosphate uridylyltransferase contributing to wall teichoic acid modification and glycolipid formation during anaerobic growth

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Abstract
UTP-glucose-1-phosphate uridylyltransferases (UGPases) are enzymes that produce UDP-glucose from UTP and glucose-1-phosphate. In *Bacillus subtilis* 168, UDP-glucose is required for the decoration of wall teichoic acid (WTA) with glucose residues and the formation of glucolipids. The *B. subtilis* UGPase GtaB is essential for UDP-glucose production under standard aerobic growth conditions, and *gtaB* mutants display severe growth and morphological defects. However, bioinformatics predictions indicate that two other UGPases, are present in *B. subtilis*. Here, we investigated the function of one of them named YngB. The crystal structure of YngB revealed that the protein has the typical fold and all necessary active site features of a functional UGPase. Furthermore, UGPase activity could be demonstrated in vitro using UTP and glucose-1-phosphate as substrates. Expression of YngB from a synthetic promoter in a *B. subtilis* gtaB mutant resulted in the reintroduction of glucose residues on WTA and production of glucolipids, demonstrating that the enzyme can function as UGPase in vivo. When wild-type and mutant *B. subtilis* strains were grown under anaerobic conditions, YngB-dependent glycolipid production and glucose decorations on WTA could be detected, revealing that YngB is expressed from its native promoter under anaerobic condition. Based on these findings, along with the structure of the operon containing yngB and the transcription factor thought to be required for its expression, we propose that besides WTA, potentially other cell wall components might be decorated with glucose residues during oxygen limited growth condition.

Introduction
The cell envelope of bacteria is composed of several sugar-containing polymers, including peptidoglycan, capsular polysaccharides, lipopolysaccharide (LPS) in Gram-negative bacteria and secondary cell-wall polymers such as teichoic acids in Gram-positive bacteria (1-5). Secondary cell-wall polymers in Gram-positive bacteria can either be complex and made up of different repeating sugar units, or more simple glycerol- or ribitol-phosphate polymers that are further decorated with sugar residues (6-8). Under standard aerobic growth conditions, the model Gram-positive organism *Bacillus subtilis* strain 168 produces two different types of teichoic acid (TA). Lipoteichoic acid (LTA) is a polyglycerolphosphate polymer that is linked by the glycolipid anchor diglucosyl-diacylglycerol (Glc2-DAG) to the outside of the bacterial membrane and further decorated with D-alanine and N-acetylg glucosamine (GlcNAc) residues, and wall teichoic acid (WTA) is a polyglycerol phosphate polymer covalently linked to peptidoglycan and decorated with D-alanine and glucose residues (9,10). Both polymers are made up of glycerol phosphate repeating units but are produced by separate pathways. Whereas LTA is polymerized on the outside of the cell, WTA is polymerized within the cell (7,8). Under phosphate-limiting growth conditions, at least part of the WTA is replaced with teichuronic acid, a non-phosphate containing anionic cell-wall polymer (11,12).

For the synthesis as well as decoration of bacterial cell-wall polymers, several important enzymes producing nucleotide-activated sugars are required (13,14). For the decoration of LTA with GlcNAc residues, UDP-GlcNAc is utilized, which is also an essential precursor for peptidoglycan and WTA synthesis (7,15). UDP-GlcNAc is produced from glucosamine-1-phosphate by the bifunctional enzyme GlmU, which has acyltransferase and uridylyltransferase activity (16). The nucleotide-activated sugar is subsequently used by a multi-enzyme glycosylation machinery for the modification of LTA with GlcNAc residues on the outside of the cell. For this process, it is thought that the membrane-linked glycosyltransferase CsbB transfers the GlcNAc residue from UDP-GlcNAc onto the lipid carrier undecaprenyl phosphate (C55-P) to generate the lipid-linked sugar intermediate C55-P-GlcNAc (17-19). Next, this intermediate is transferred across the membrane with the aid of the small membrane protein and proposed flippase enzyme GtcA, which belongs to the GtrA protein family (20,21). The GlcNAc residues are finally added to the LTA polymer by the multi-membrane spanning GT-C-type glycosyltransferase YfhO (19).

The glycosylation process of WTA in *B. subtilis* is much simpler. The glucose residues are attached to the polymer within the cytoplasm of the cell by the glycosyltransferase TagE using UDP-glucose as substrate (22,23). The nucleotide activated sugar precursor UDP-glucose is produced from UTP and glucose-1-phosphate by UTP-
glucose-1-phosphate uridylytransferase (UGPase) enzymes (14). UGPases are widespread in bacteria and are often named GalU. For several Gram-positive as well as Gram-negative bacterial pathogens, GalU has been shown to be required for full virulence due to its involvement in biofilm formation, capsule and/or LPS biosynthesis (24-27). A well-characterized GalU equivalent in B. subtilis is GtaB, and the first gtaB mutants were isolated as part of studies investigating phage-resistant B. subtilis strains (10). These studies also revealed that the phage resistance is due to the lack of glucose decorations on WTA (28,29). In B. subtilis 168, UDP-glucose is not only required for the decoration of WTA with glucose residues but also for the production of glycolipids. An abundant glycolipid found in the membrane is Glc2-DAG, which serves as the lipid anchor for LTA and is produced by the transfer of two glucose molecules from UDP-glucose onto the membrane lipid diacylglycerol (DAG) by the glycosyltransferase UgpP (or sometimes also named YpfP) (30-33). Several independently obtained B. subtilis gtaB mutants have now been characterized. These mutants are resistant to certain types of phages, display morphological and growth defects, lack glucose decorations on WTA and are unable to produce glycolipids under standard laboratory growth conditions (31,34). Furthermore, no UGPase activity could be detected in lysates prepared from gtaB mutant strains, indicating that GtaB is the sole functional UGPase in B. subtilis under these conditions (28). However, apart from GtaB, two other predicted UGPases, YtdA and YngB are encoded in the B. subtilis 168 genome: as part of this study we investigated the function of YngB (35,36).

The yngB gene is thought to be part of the yngABC operon (35), but an additional internal promoter appears to present in yngB driving yngC expression. Furthermore, expression of yngC has been reported to be under control of a sigma M-dependent promoter (37). YngA belongs to the GtrA protein family and could therefore, similar as proposed for the GtcA enzyme involved in the LTA glycosylation process, function as flippase enzyme and be required for the transfer of a sugar-linked C55-P lipid intermediate across the membrane. YngC is a membrane protein belonging to the DedA family of proteins. The function of the protein is not clear, but proteins belonging to this family have been shown to be important for multiple processes in other bacteria, including cell division, membrane composition and antibiotic resistance (38-40). The transcription of the yngABC operon is activated by the transcription factor YcJ, which forms a two-component system (TCS) with the histidine sensor kinase YclK (35,41,42). The biological role of the YcJ-YclK TCS remains unknown but it has been shown that the yclJK regulon is upregulated during low oxygen conditions (43,44). Hence the proposed UGPase YngB could potentially function under oxygen-limiting conditions. However, the expression might be more complex as the expression of the yclJK genes in turn is controlled by the ResDE two-component system, a signal transduction system known to play a key role in the expression of both aerobic and anaerobic respiration-related genes in B. subtilis (45,46).

In this study, we aimed to determine if YngB is a functional UGPase enzyme and to provide insight into its biological function. Based on its structure as well as its in vitro enzymatic activity, we showed that YngB is a functional UGPase. YngB can also function in vivo as UGPase leading to the decoration of WTA with glucose residues and glycolipid production in the absence of GtaB, when expressed from a synthetic promoter. YngB-dependent glycosylation of WTA and glycolipid production was also observed when B. subtilis was grown under anaerobic fermentative growth condition. This revealed that besides GtaB, YngB is expressed from its native promoter under anaerobic growth conditions and functions as UGPase. Based on these findings and previous reports on the transcription control of the yngABC and other operons in B. subtilis, the potential decoration of other cell wall structures with glucose residues during oxygen limited growth condition will be discussed.

Results

The crystal structure of the B. subtilis YngB protein reveals a putative UDP-glucose binding site

Three paralogous UGPases, GtaB (BSU_35670), YtdA (BSU_30850) and YngB (BSU_18180) are encoded in the genome of B. subtilis strain 168 (Fig. S1). GtaB has been characterized as a UGPase and in its absence B. subtilis lacks glucose decorations on WTA and is unable to produce glycolipids during vegetative growth (28,31). Here we set out
to determine whether YngB is a bona fide UGPase and to investigate its biological function. An amino acid sequence alignment of the B. subtilis proteins GtaB, YngB and YtdA with the UGPase enzymes A4JT02 from Burkholderia vietnamiensis (PDB code: 5I1f), GalUHp from Helicobacter pylori (PDB code: 3juk) and GalUCg from Corynebacterium glutamicum (PDB code: 2pa4), for which structures with bound UDP-glucose are available (47,48), revealed that most of the UDP-glucose binding residues (Fig. S1, colored in yellow) as well as the metal-chelating residue (Fig. S1, colored in cyan) are conserved in the three B. subtilis proteins (14,47,48). To determine the crystal structure of YngB, selenomethionine-substituted protein was produced in E. coli and purified as a C-terminal His-tag fusion protein. The protein crystallized as a dimer in the asymmetric unit and the structure was solved at 2.73 Å by experimental phasing (Fig.1A and Table S1). YngB displayed a Rossmann fold with alternating α-helices and β-strands, which is commonly found in nucleotide-binding proteins (49). More specifically, the YngB monomer contains a central β-sheet surrounded by α-helices (47,50). Similar to the dimer interactions described for homologous UGPases, hydrogen bonds are formed between Tyr102 residues on the two β-strands located at the interface of the monomers, producing an extended central β-sheet spanning across the two subunits (50,51). The overall structure of YngB is very similar to homologous UGPases including A4JT02, GalUHp and GalUCg as shown by superimposition (Fig. S2). The outer most C-terminal α-helix is one of the most variable region and either absent in GalUHp or in a different conformation in GalUCg (Fig. S2). To locate the substrate-binding site, the YngB and UDP-glucose bound H. pylori GalUHp structures (PDB code: 3juk) (47) were overlayed. The structures superimposed with an r.m.s.d. of 1.47 Å and the putative UDP-glucose binding pocket could easily be identified in YngB (Fig.1B & 1C). Based on this alignment it can be predicted that residues Gly110, Gln105, Ala13, Gly14, Glu29, and Lys28 in YngB interact with the uridine moiety; Asp133 is involved in the chelation of a Mg²⁺ ion; Asp134, Val204, Gly172, and Glu191 interact with the glucose moiety; and Lys192 interacts with the diphosphate moiety (Fig. 1C). Based on this structural analysis, YngB has all the required features expected of a bona fide UGPase enzyme.

YngB shows UGPase activity in vitro
To determine if YngB has UGPase activity in vitro, enzyme assays were performed using a method previously described for assessing the UGPase activity of the GalU enzyme from Erwinia amylovor (52). For this assay, purified proteins are incubated with α-glucose-1-phosphate (G-1-P) and UTP and active UGPases will convert these substrates into UDP-glucose and pyrophosphate. The generated pyrophosphate is then hydrolyzed by a pyrophosphatase to two molecules of inorganic phosphate, which is quantified calorimetrically. To assess the enzymatic activity of YngB, assays were performed with recombinant YngB protein using recombinant GtaB protein as control, and increasing concentrations of G-1-P and a fixed concentration of UTP or increasing concentrations of UTP and fixed concentration of G-1-P. These experiments revealed that YngB possesses UGPase activity, and Michaelis-Menten curves could be produced for both enzymes acting on either substrate (Fig. 2). From these data, apparent Kₘ values of 45.6 ± 3.24 μM (GtaB) and 42.1 ± 20.2 μM (YngB) were calculated for G-1-P and 49.5 ± 10.2 μM (GtaB) and 62.9 ± 13.8 μM (YngB) for UTP, respectively. k_cat values of 1.06 s⁻¹ (GtaB) and 0.264 s⁻¹ (YngB) were calculated for G-1-P and 1.04 s⁻¹ (GtaB) and 0.293 s⁻¹ (YngB) for UTP, respectively. These enzyme assays confirmed that YngB has UGPase activity in vitro and revealed that under the assay conditions used, YngB had a lower turnover number compared to GtaB.

Expression of YngB from an inducible promoter leads to glucosylation of WTA
In B. subtilis strain 168, glucose is transferred onto WTA by the glycosyltransferase TagE using UDP-glucose as substrate (23). Under standard aerobic growth conditions, GtaB appears to be the only enzyme that produces UDP-glucose, as glucose is absent from WTA in gtaB mutant strains (28,34,53). This is somewhat at odds with our data showing that B. subtilis YngB protein is a functional UGPase enzyme. Possible explanations could be that YngB uses a different sugar or nucleotide as substrate in vivo or that it is not expressed under standard aerobic growth conditions. To address these issues, the yngB gene or, as a control, the gtaB gene, were placed under control of the synthetic IPTG-inducible P_hyperspank
promoter (short \text{P}_{\text{hyper}} promoter) and introduced into the chromosome of the \text{A}gt\text{a}B single mutant and the \text{A}gt\text{a}B\text{A}yng\text{B} double mutant strain. The presence of glucose on WTA in the different \textit{B. subtilis} strains was initially assessed by fluorescence microscopy after staining the bacteria with fluorescently labeled concanavalin A, a lectin that specifically binds to terminal glucose residues (34). The different \textit{B. subtilis} strains were grown aerobically in medium supplemented with IPTG and culture samples were taken at mid-log growth phase for microscopy analysis. A fluorescence signal was observed for the wild-type and \text{A}yng\text{B} single mutant strain, but as expected was absent from the \text{A}gt\text{a}B single and the \text{A}gt\text{a}B\text{A}yng\text{B} double mutant strains (Fig. 3). Consistent with previous observations, \text{A}gt\text{a}B mutant bacteria displayed morphological defects and the cells were curled and showed some bulges, which was also seen for \text{A}gt\text{a}B\text{A}yng\text{B} double mutant cells (Fig 3. Expression of either \text{yng}B or gtaB from the inducible promoter (\text{P}_{\text{hyper}}) in the \text{A}gt\text{a}B single or \text{A}gt\text{a}B\text{A}yng\text{B} double mutant strains fully or at least partially complemented both phenotypes; the cells showed again binding to the fluorescent lectin and bacteria complemented with GtaB had a normal rod-shaped morphology and the bacteria complemented with YngB showed an improved cell morphology and the cell were less curved (Fig. 3). These results indicate that YngB is not expressed from its native promoter under standard aerobic growth conditions; however, when expressed from an inducible promoter, YngB can function as UGPase \it{in vivo}. To further confirm that YngB expression leads to the decoration of WTA with glucose resides, WTA was isolated from wild-type, the \text{A}gt\text{a}B mutant and strains \text{A}gt\text{a}B \text{P}_{\text{hyper-gta}B} and \text{A}gt\text{a}B \text{P}_{\text{hyper-yng}B} and analyzed by NMR and LC-MS. The $^1$H NMR spectrum obtained for WTA isolated from the wild-type strain revealed a peak at 5.2 ppm likely derived from the hydrogen atom at the anomeric carbon of the glucose residue on WTA (23) (Fig. 4A). This peak was absent in the sample derived from the \text{A}gt\text{a}B mutant strain but could again be detected in samples isolated from strains \text{A}gt\text{a}B \text{P}_{\text{hyper-gta}B} and \text{A}gt\text{a}B \text{P}_{\text{hyper-yng}B} (Fig. 4A), indicating that WTA is indeed decorated with glucose residues upon expression of YngB from an inducible promoter. For the LC-MS analysis, the purified WTA was hydrolyzed with hydrogen fluoride and the depolymerized species were characterized. A species with a mass of 253.09 \textit{m/z} corresponding to a glycerol-glucose repeating unit was detected for samples derived from the wild-type as well as strains \text{A}gt\text{a}B \text{P}_{\text{hyper-gta}B} and \text{A}gt\text{a}B \text{P}_{\text{hyper-yng}B} (Fig. 4B) but was absent from the \text{A}gt\text{a}B sample (Fig. 4B). These data highlight that YngB can function as UGPase \it{in vivo} and produce UDP-glucose, which can subsequently be used for the decoration of WTA with glucose moieties.

**Expression of YngB from an inducible promoter leads to the formation of glycolipids**

UDP-glucose is also used for the production of glycolipids in \textit{B. subtilis} and the glycosyltransferase UgpP transfers one or more glucose moieties onto the membrane lipid diacylglycerol (DAG) (30,32). It has been reported that glycolipids are absent in \textit{B. subtilis gta}B mutant strains (31,54), however, our data suggest that expression of YngB in gtaB mutant strains should restore glycolipid production. To experimentally verify this, total membrane lipids were isolated from wild-type, \text{A}gt\text{a}B single and \text{A}gt\text{a}B\text{A}yng\text{B} double mutant strains as well as complementation strains expressing either gtaB (\text{P}_{\text{hyper-gta}B}) or yngB (\text{P}_{\text{hyper-yng}B}) from the IPTG-inducible promoter. The isolated membrane lipids were separated by thin layer chromatography and glycolipids were visualized by staining with α-naphthol and sulfuric acid (Fig. 5). Several major bands were observed for the wild-type strain, two of which (Fig. 5 top and middle band) could be further characterized by MALDI-TOF MS (Fig.6) and MALDI-TOF MS/MS (Fig. S3). Based on the mass and fragmentation pattern, they are most consistent with Glc$_2$-DAG (top band) and Glc$_3$-DAG (middle band) sodium adducts with different fatty acid chain lengths (Table 1 and Table S2). These bands were absent in the \text{A}gt\text{a}B single and \text{A}gt\text{a}B\text{A}yng\text{B} double mutant strains, but again present upon expression of either gtaB or yngB from the IPTG-inducible promoter (Fig. 5A and 5B). The identity of these lipids was again consistent with sodium adducts of the glycolipids Glc$_2$-DAG (top band) and Glc$_3$-DAG (middle band) as assessed by MALDI-TOF MS and MALDI-TOF MS/MS (Table. 1 and Table S2). The glycolipids produced by a \text{A}yng\text{B} mutant and complementation strain \text{A}yng\text{B} \text{P}_{\text{hyper-yng}B} were also analyzed, however, no clear differences in the glycolipid profile were observed for these strains as compared to the wild-type strain (Fig. 5C). These data confirm that YngB is not required...
for glycolipid production under standard aerobic growth, presumably because the enzyme is not produced under these conditions and that GtaB is the main enzyme responsible for UDP-glucose production. However, when YngB is expressed from a synthetic promoter, it can take over the function of GtaB.

**YngB is produced under anaerobic growth condition, leading to decoration of WTA with glucose residues and glycolipid production**

In previous work, it has been shown that expression of genes in the *yngABC* operon is upregulated by the transcription activator YclJ (35). Expression of *yclJ* itself is under control of the transcription factor ResD, which is produced under oxygen-limitation conditions (42-44). It is therefore possible that YngB is expressed from its native promoter under anaerobic growth conditions through a pathway involving ResD and YclJ. To determine if YngB is expressed from its native promoter under anaerobic growth condition and contributes to the decoration of WTA with glucose molecules and glycolipid production, the wild-type *B. subtilis* strain 168 and the isogenic Δ*gtab* and Δ*yngB* single and Δ*gtabΔyngB* double mutant strains were grown in an anaerobic chamber under fermentative growth condition. The presence of glucose on WTA and the production of glycolipids was assessed by fluorescence microscopy and TLC analysis as described above. Clear fluorescence signals were observed for the wild-type, Δ*gtab* mutant and Δ*yngB* mutant strains under anaerobic fermentative growth conditions, indicating that WTA is also decorated with glucose residues during anaerobic growth (Fig. 6A). Only cells of the Δ*gtabΔyngB* double mutant were no longer stained (Fig. 6A). These data suggest that under these growth conditions YngB is produced from its native promoter and functions as a second UGPase enzyme next to GtaB. Consistent with the fluorescence microscopy data, glycolipids could be detected by thin layer chromatography in wild-type, Δ*gtab* and Δ*yngB* single mutant strains, but not in the Δ*gtabΔyngB* double mutant following growth under anaerobic, fermentative growth condition (Fig. 6B). Some differences in the glycolipid profiles were observed for lipid samples isolated from the wild-type following growth under aerobic or anaerobic conditions (Fig. 6B). A number of the slower migrating glycolipids were absent in samples derived from the anaerobically grown cultures, however bands likely corresponding to Glc2-DAG and Glc3-DAG were present in both samples (Fig. 6B). Furthermore, a reduced glycolipid signal was observed for the Δ*gtaB* mutant compared to the wild-type and Δ*yngB* mutant strain (Fig. 6B). This is consistent with the microscopy results and the observation that only some but not all of the Δ*gtaB* mutant cells seemed to contain glucose decorations on their WTA. Taken together, the data show that GtaB is the main UGPase enzyme producing UDP-glucose under both aerobic and anaerobic fermentative growth conditions. Furthermore, our data show that YngB is a functional UGPase that augments UDP-glucose production under oxygen limiting conditions. Under these conditions, it functions together with GtaB to produce UDP-glucose, for the production of glycolipids and the decoration of WTA with glucose residues and as discussed below potentially also other cell wall structures.

**Discussion**

In bacteria, nucleotide-activated sugars are key sugar donors for glycosylation processes (52,55). In the Gram-positive, spore-forming bacterium *B. subtilis*, one of the best characterized nucleotide-activated sugar-synthesizing enzymes is GtaB. Up to now all UDP-glucose produced in *B. subtilis* has been attributed to the activity of GtaB, despite the presence of two orthologous proteins, YngB and YtdA. Here we show that, based on its crystal structure and in vitro and in vivo biochemical activity, the *B. subtilis* YngB protein is a functional UGPase (Figs. 1 and 2). The necessary glucose-1-phosphate and UTP substrate-binding residues could be identified in the YngB structure (Fig. 1) and it is likely that YngB synthesizes UDP-glucose via the same catalytic mechanism proposed for other members of this family (47,48,52).

The main reason why all UDP-glucose production in *B. subtilis* has been attributed to GtaB is likely due to the specific growth conditions or developmental stages in which YngB and YtdA are produced. The findings presented in this study are consistent with previous reports that GtaB is likely the sole functional UGPase during vegetative aerobic growth (Fig. 3 and 4). *ytdA* is under the control of the sporulation-specific transcription factor sigma K and hence only expressed during late stages of the sporulation process (56). It has
been suggested that YtdA is a UGPase involved in the production of the polysaccharide layer on spores, however no clear phenotype could be identified for a ytdA mutant strain (36). We present here experimental evidence that YngB is produced and can synthesize UDP-glucose when bacteria are grown under anaerobic conditions (Fig. 7). However, GtaB appears to be also the main UGPase in cells grown under anaerobic conditions, and YngB activity could only be revealed in a gtaB mutant strain (Fig. 7). Therefore, similar as done here, to reveal a function of YtdA as UGPase, its activity and contribution to the production of the polysaccharide layer on spores might need to be assessed in a gtaB mutant strain.

Under aerobic growth condition, an aberrant morphology was observed for the ΔgtaB mutant, and in place of rod-shaped bacteria, the cells were curved (Fig. 3). This is consistent with previous observations, and it thought that the aberrant morphology is due to the lack of glycolipids rather than the absence of glucose residues on WTA, since a ugpP (ypfP) mutant but not a tagE mutant showed similar morphological defects (31,53,57). Consistent with these findings, the ΔgtaBΔyngB double mutant strain, which is unable to produce glycolipids, showed similar morphological defects (Figs. 3 and 5). Furthermore, cellular UDP-glucose levels are thought to impact the cell size of B. subtilis via a second moonlighting function of UgtP (58). Under nutrient rich conditions, UDP-glucose, which is present under these conditions at high levels, will bind to UgtP. The substrate-bound UgtP protein will not only produce glycolipids but will also interact with the key cell division protein FtsZ and in this manner inhibit the maturation of the cytokinetic ring leading to an increase in cell size before cell division is completed (58). On the other hand, under nutrient limiting conditions, a reduction in the availability of UDP-glucose in the cell is thought to favor UgtP oligomerization, preventing it from inhibiting FtsZ and allowing cells to divide at a smaller cell mass leading to cells of reduced size (58,59). Under anaerobic growth conditions, B. subtilis grows a lot slower as compared to aerobic conditions and produced also shorter rod-shaped cells compared to aerobic growth conditions (Figs 3 and 7). Interestingly, no aberrant morphology was observed for ΔgtaB or ΔgtaBΔyngB mutant bacteria, and both strains produced short rod-shaped cells (Fig. 7). While the gtaB single mutant is able to produce some glycolipids even under anaerobic conditions, this observation is in particular interesting for the ΔgtaBΔyngB double mutant strain, which is unable to synthesize glycolipids under these conditions (Fig. 7). These data indicate that under anaerobic growth conditions, the ability of bacteria to produce glycolipids is not essential for cells to maintain their normal rod-shape. Furthermore, similar to nutrient poor growth conditions, the growth rate of B. subtilis is drastically reduced under anaerobic growth conditions. This will likely also lead to reduced cellular UDP-glucose levels in wild-type cells, which in turn would favor UgtP oligomerization preventing it from interacting with FtsZ. As a consequence, cell division can take place when bacteria have a smaller cell mass and cell size, and this is indeed observed for bacteria grown under anaerobic conditions. Shorter cells might be less susceptible to the disruption of other cellular pathways, such as glycolipid synthesis, in order to maintain their cell shape.

The finding that YngB contributes to the production of UDP-glucose under anaerobic growth conditions is consistent with previous reports on its expression control. Transcription of the yngABC operon has been reported to be activated by YcIJ, a transcription factor and part of the YcJK two-component system, whose expression itself is upregulated during oxygen limitation by the ResDE two-component system (35,42,44). Previous studies have revealed that genes regulated by YcIJ include the three genes of the yngABC and the two genes of the ykcBC operon (35,42) (Fig. 8A). Here, we show that the UDP-glucose produced by YngB can be utilized for the glucosylation of WTA and glycolipid production. However, given the predicted function of the proteins encoded by the yngABC operon and the ykcBC operon, we speculate that the UDP-glucose produced by YngB under anaerobic growth conditions could potentially be utilized for the glycosylation of other cell wall polymers or extracellular cell-wall components (Fig. 8). Genes forming part of the YcIJ regulon, encode putative glycosyltransferases and a transporter that may allow glucosylation to take place extracellularly, in addition to the utilization of the UDP-glucose intracellularly. For example, WTA is glycosylated intracellularly in B. subtilis by TagE, and this
enzyme directly adds the sugar residue onto the WTA backbone using a nucleotide-activated sugar as precursor. In contrast, LTA is glycosylated extracellularly using a multi-component transmembrane glycosylation system (21). A nucleotide-activated sugar, in the case of *B. subtilis* UDP-GlcNAc, is first linked by the glycosyltransferase CsbB to the C₅₅-P lipid carrier in the cytosol of the cell, the C₅₅-P-sugar intermediate is subsequently flipped across the membrane, likely by the GtrA-type membrane protein GtcA, and the sugar is finally added onto the LTA polymer on the outside of the cell by the multi-membrane spanning GT-C-type glycosyltransferase YfhO (19-21) (Fig. 8B). Proteins with homology to CsbB, GtcA and YfhO are encoded in the YelJ-controlled yngABC and ykcBC operons. YkcC shows homology to the glycosyltransferase CsbB. YngA is similar to GtcA, a GtrA protein family member and hence predicted to mediate the transport of a C₅₅-P-sugar intermediate across the membrane (35,60). Finally, YkbB is predicted to be a multi-membrane spanning GT-C-fold glycosyltransferase similar to YfhO (Fig. 8C). By analogy to the function of CsbB, GtcA and YfhO, which utilize UDP-GlcNAc for the transfer of GlcNAc onto the LTA polymer on the outside of the cell, we speculate that YkcC, YngA and YkbB constitute a multi-component transmembrane glycosylation system (Fig. 8C). Since the transcription of yngABC and ykcBC operons is predicted to be activated during anaerobic growth conditions, the YkcC-YngA-YkbB system might function only during specific growth conditions. As shown here, YngB is a functional UGPase, and hence we speculate that UDP-glucose is the likely substrate for the YkcC-YngA-YkbB multi-component transmembrane glycosylation system (Fig. 8C). While the glycosylation target in the bacterial cell envelope is currently unknown, it will be interesting to investigate this further in future studies. Furthermore, while UDP-glucose produced by YngB is utilized by TagE and UgtP, it will also be interesting to determine if, through specific protein/protein interactions, the UDP-glucose produced by YngB can be more efficiently fed towards the YkcC-YngA-YkbB multi-component glycosylation system, as compared to TagE and UgtP, enzymes which are supplied with UDP-glucose by GtaB.

In conclusion, we provide the first experimental evidence that the *B. subtilis* YngB protein is a functional UGPase, which is produced under anaerobic growth conditions. The UDP-glucose synthesized by YngB is utilized for the glycosylation of WTA as well as glycolipid production. However, we speculate that the UDP-glucose produced by YngB might also enter other glycosylation pathways, leading to the decorating of other cell envelope components with glucose residues under anaerobic or other growth conditions, which result in the activation of genes forming part of the YclJ regulon.

**Experimental procedures**

**Bacterial strains and growth condition**

All strains used in this study are listed in Table S3. *Escherichia coli* and *B. subtilis* strains were grown in Luria-Bertani (LB) medium at 37°C unless otherwise stated. LB medium for *B. subtilis* was supplemented with 0.2 % D-glucose for aerobic growth condition. For anaerobic fermentative growth of *B. subtilis*, a single colony was first inoculated in LB medium and grown at 37°C aerobically until reaching an OD₆₀₀ of 1.0. Then 100 µl of the bacteria culture was spread on 2 × yeast extract tryptone (YT) agar plates (16g/L tryptone, 10g/L yeast extract, 5g/L NaCl, 1% glucose, 20 mM K₃PO₄ pH 7.0, 15g/L Bacto agar). The plates were incubated for 18 hr at 37°C in an anaerobic cabinet (Don Whitley Scientific) with an atmosphere of 10% CO₂, 10% H₂, and 80% N₂. Bacterial cultures were supplemented with appropriate antibiotics at the following final concentrations: *E. coli* cultures, ampicillin (Amp) at 100 µg/ml and kanamycin (Kan) at 30 µg/ml; for *B. subtilis* cultures, Kan at 10 µg/ml, erythromycin (Erm) at 5 µg/ml, spectinomycin (Spec) at 100 µg/ml.

**Strain and plasmid construction**

All primers used in this study are listed in Table S4. For the construction of plasmids for the expression and purification of the C-terminally His-tagged *B. subtilis* GtaB and YngB proteins, the gtaB (BSU_35670) and yngB (BSU_18180) genes were amplified by PCR from *B. subtilis* 168 genomic DNA using primer pairs ANG3161/ANG3162 and ANG3163/ANG3164, respectively. The PCR products were digested with NcoI and XhoI and ligated with plasmid pET28b cut with the same
restriction enzymes. The resulting plasmids pET28b-gtaB-cHis and pET28b-yngB-cHis were recovered in E. coli strain XL1-Blue, yielding strains ANG5206 and ANG5207, respectively. Sequence of the inserts for pET28b plasmids was confirmed by sequencing using primers ANG111 and ANG112. For protein expression, the plasmids pET28b-gtaB-cHis and pET28b-yngB-cHis were introduced into E. coli strain BL21(DE3), yielding strains ANG5208 and ANG5209.

For the construction of B. subtilis expressing gtaB or yngB from the IPTG-inducible hyperspang promoter (P\text{hyper}), the gtaB (BSU_35670) and yngB (BSU_18180) genes were amplified by PCR using B. subtilis 168 genomic DNA as template and primer sets ANG3203/ANG3204 and ANG3205/ANG3206, respectively. The PCR products were digested with HindIII and NheI and ligated with plasmid pDR111 cut with the same restriction enzymes. The resulting plasmids pDR111-gtaB and pDR111-yngB were recovered in E. coli strain XL1-Blue, yielding strains XL1-Blue-pDR111-gtaB and XL1-Blue-pDR111-yngB. The sequences of the inserts in pDR111 plasmids were confirmed by sequencing using primers ANG1671 and ANG1672. Plasmids pDR111, pDR111-gtaB, and pDR111-yngB were linearized with ScaI and introduced into the wild-type B. subtilis strain 168 yielding 168 amy::spec P\text{hyper} (ANG5675), 168 amy::spec P\text{hyper-gtaB} and 168 amy::spec P\text{hyper-yngB} respectively. Next, the chromosomal DNA of strain 168ΔgtaB::kan (ANG5277) was introduced to strains 168 amy::spec P\text{hyper}, 168 amy::spec P\text{hyper-gtaB} and 168 amy::spec P\text{hyper-yngB} yielding strains 168ΔgtaB::kan amy::spec P\text{hyper} (ANG5676), 168ΔgtaB::kan amy::spec P\text{hyper-gtaB} (ANG5677) and 168ΔgtaB::kan amy::spec P\text{hyper-yngB} (ANG5678), respectively. The chromosomal DNA of strain 168ΔyngB::kan (ANG5263) was introduced into 168 amy::spec P\text{hyper} and 168 amy::spec P\text{hyper-yngB} yielding strains 168ΔyngB::kan amy::spec P\text{hyper} (ANG5679) and 168ΔyngB::kan amy::spec P\text{hyper-yngB} (ANG5680). The chromosomal DNA of strain 168ΔyngB::erm (ANG5659) was introduced into strain ANG5677, yielding strain 168ΔgtaB::kan ΔyngB::erm amy::spec P\text{hyper-gtaB}(ANG5682). The chromosomal DNA of strain 168ΔgtaB::erm (ANG5658) was introduced into strains ANG5679 and ANG5680, yielding strains 168ΔgtaB::erm ΔyngB::kan amy::spec P\text{hyper} (ANG5681) and 168ΔgtaB::erm ΔyngB::kan amy::spec P\text{hyper-yngB} (ANG5683). The deletion of the gtaB and yngB genes was confirmed by PCR using primer sets ANG3197/ANG3198 and ANG3199/ANG3200, respectively. The integration of the gtaB gene at the \text{amyE} site was confirmed by PCR using primer sets ANG1663/ANG3204 and ANG1664/ANG3203. The integration of the yngB gene at the \text{amyE} site was confirmed by PCR using primer sets ANG1663/ANG3206 and ANG1664/ANG3205. The integration of plasmid pDR111 at the \text{amyE} site for strain 168 amy::spec P\text{hyper} was confirmed by PCR using primers sets ANG1664/ANG1671 and ANG1663/ANG1672.

Expression and purification of GtaB and YngB

E. coli strain BL21(DE3) pET28b-gtaB-cHis (ANG5208) was grown in LB medium at 30°C with shaking until reaching an OD\text{600} of 0.6. Protein expression was induced by the addition of IPTG to a final concentration of 0.5 mM and the cultures were incubated overnight at 16 °C with agitation. Bacterial cells were harvested by centrifugation and washed once with cold 500mM NaCl, 50mM Tris pH 7.5 buffer and the bacterial pellets stored at -20°C for future use. For the protein purification, the bacterial cells were suspended in 20 ml cold buffer A (500mM NaCl, 50mM Tris pH 7.5, 5% glycerol, 10 mM imidazole) supplemented with cOmplete™ protease inhibitor cocktail (Roche), 100 ug/ml lysozyme, 10 ug/ml DNase, followed by passing the cell suspension twice through a French press cell at 1100 psi. For the purification of GtaB-cHis, the cell lysate was loaded onto a 5ml HisTrap column equilibrated with buffer A. The column was washed with 5 column volumes of buffer A followed by elution using a linear gradient of 10 column volumes from buffer A to buffer B (500mM NaCl, 50mM Tris pH 7.5, 5% glycerol, 500 mM imidazole). Elution fractions containing GtaB-cHis were pooled and thrombin was added to cleave off the C-terminal His-tag. The protein was dialyzed at room temperature against 1L buffer C (500mM NaCl, 50mM Tris pH 7.5, 5% glycerol) for 1 hr, followed by overnight dialysis against 1L of fresh buffer C. The protein solution was then loaded onto a Superdex 10/60 Hilo size exclusion column (GE Healthcare) equilibrated with buffer C. Following size-exclusion chromatography, the purified protein was concentrated using a PES 10-kDa cut off Pierce™ protein concentrator.
For the expression of selenomethionine substituted YngB-cHis, strain BL21(DE3) pET28b-"ngB-cHis (ANG 5209) was grown in LB medium at 37°C for 12 hr. Bacterial cells were then washed once with minimal medium (0.5 g/L NaCl, 1.0 g/L (NH₄)₂SO₄, 7.5 g/L KH₂PO₄, 23.25 g/L K₂HPO₄, 0.246 g/L MgSO₄, 7.2 g/L glucose, 100 mg/L lysine, 100 mg/L phenylalanine, 100 mg/L threonine, 50 mg/L isoleucine, 50 mg/L leucine, 50 mg/L valine, 0.1 mM CaCl₂, 10 ml/L 100 × Kao and Michayluk vitamin solution) and grown overnight at 37°C in the minimal medium containing 42 mg/L methionine. The next day, bacterial cells were washed with the minimal medium and grown in the minimal medium containing 42 mg/L selenomethionine at 37°C until reaching an OD₆₀₀ of 0.5 - 0.6. At this point 2.9 g/L additional glucose and IPTG to give a final concentration of 0.5 mM, 16°C overnight with agitation. Cells were harvested, washed and stored as described above.

For the productive of native YngB-cHis used in the kinetic assay, methionine was added to the medium in place of selenomethionine. For the purification of SeMet YngB-cHis or native YngB-cHis, the cell lysate was loaded by gravity flow onto a column of Ni-NTA resin (Qiagen) equilibrated with buffer A. The column was washed with 30 ml of buffer A and 30 ml of buffer D (500 mM NaCl, 50 mM Tris pH 7.5, 5% glycerol, 50 mM imidazole). The proteins were eluted in 5 × 1 ml fractions using buffer B. The elution fractions were pooled and subjected to size-exclusion chromatography and the purified protein concentrated as described above.

**Protein crystallization, structural solution and analysis**

SeMet YngB-cHis crystals were obtained by the sitting drop-method in 0.2M potassium citrate tribasic monohydrate, 0.05M lithium citrate tribasic tetrahydrate, 0.1M sodium phosphate monobasic monohydrate, 25% PEG6000, using a protein concentration of 6 mg/ml. SeMet YngB-cHis crystals were cryo-protected with 30% ethylene glycol and flash frozen in liquid nitrogen. Datasets were collected at the I03 Beamline at the Diamond Light Source (Harwell Campus, Didcot, UK). Data indexing, integration, scaling, and merging was done using the xia2 3dii pipeline (61). The selenium sites and initial phases were solved using CRANK2 (62). Structure refinement was performed with Refmac (63) and model building with Coot (64). Data collection and refinement statistics are summarized in Table S1. The figures with the structure were generated using PyMol.

**Enzyme kinetic analysis**

The steady state kinetics assays with purified GtaB and YngB-cHis proteins were performed using a previously described method with some modifications (52). Briefly, enzyme assays were performed in 96-well plates in 100 μl reaction volumes. The reactions contained 50 mM Tris pH 7.5, 500 mM NaCl, 5 % glycerol, 10 mg/ml MgCl₂, 0.05U of *E. coli* pyrophosphatase (NEB), 100 nM GtaBs or 100 nM YngB₉₃-cHis. For measuring the Kₘ value for α-glucose-1-phosphate (G-1-P), the reactions contained 200 μM UTP and α-glucose-1-phosphate at a concentration of 200 μM, 150 μM, 100 μM, 50 μM, 12.5 μM, 6.3 μM, or 3.1 μM. For measuring the Kₘ value of UTP, the reactions contained 200 μM α-glucose-1-phosphate and UTP at a concentration of 200 μM, 150 μM, 100 μM, 50 μM, 12.5 μM, 6.3 μM, or 3.1 μM. Reactions were performed in triplicate and reactions without GtaBs and YngB₉₃-cHis were used as negative controls. Reactions were incubated at room temperature for 1 min for GtaBs and 4 min for YngB₉₃-cHis so that less than 20% of the substrates were converted in the reactions. The reactions were terminated by adding 100 μl of Biomol® Green (Enzo® Life Sciences) and after a 20 min incubation at room temperature, the absorbance was measured at 620 nm. Michaelis-Menten non-linear fitting was performed using Prism.

**Detection of glucose residues on WTA using fluorescently labelled lectin Concanavalin A**

Glucose modifications on WTA were detected by fluorescence microscopy using a previously described method with minor modifications (34). Single colonies of the different *B. subtilis* strains were used to inoculate 5 ml LB medium and the cultures grown overnight at 37°C. The overnight cultures were back diluted 1:100 into 25 ml of fresh LB medium supplemented with 1 mM IPTG and the cultures grown at 37°C until reaching mid-log growth phase (OD₆₀₀ between 0.4 and 0.6). Cells equivalent to 100 μl of a culture with an OD₆₀₀ of 0.5 were pelleted by centrifugation at 17,000 × g for 1 min and washed once with PBS pH 7.4. Bacterial
Isolation of WTA and its analysis by NMR and UPLC-MS

*B. subtilis* strains were grown in 2 L LB medium supplemented with 1 mM IPTG at 37 °C. Once the cultures reached an OD<sub>600</sub> of 0.6, the cells were harvested by centrifugation and WTA isolated using a previously described method (19). The NMR analysis of WTA was performed as previously described (19,20). Briefly, 2 mg of WTA from each strain was suspended and lyophilized twice in 500 μl D<sub>2</sub>O of 99.96% purity. Lyophilized WTA at the final step was suspended in 500 μl D<sub>2</sub>O of 99.96% purity and NMR spectra were recorded on a 600-MHz Bruker Advance III spectrometer equipped with a TCI cryoprobes. NMR spectra were recorded at 303 K with a total recycling time of 5 s and a 1H flip angle of approximately 30°. Two independent experiments were performed, and very similar spectra were obtained and one spectrum for each strain is shown. For the UPLC-MS analysis of the purified WTA, the method used was adapted from previously described protocols (20,65). Briefly, 2 mg of the purified WTA was lyophilized in deionized distilled H<sub>2</sub>O. The lyophilized WTA was then depolymerized into monomeric units by hydrolysis of the phosphodiester bonds using 48% hydrofluoric acid for 20 hr at 0°C. The depolymerized WTA material was subjected to UPLC-MS analysis as described previously (65). All data were collected and processed using the MassLynx software, version 4.1 (Waters Corp., USA).

Isolation of membrane lipids and TLC analysis

For the isolation of total membrane lipids, the different* B. subtilis* strains were grown overnight at 37°C in 100 ml LB medium supplemented with 1 mM IPTG. Cells were harvested by centrifugation, and total membrane lipids isolated as described previously (66). TLC analysis and detection of glycolipids was performed as described (66). Briefly, isolated lipids were suspended in chloroform and 0.5 mg were spotted on Å60 silica gel plates (Macherey-Nagel). Lipids were separated using a developing solvent of chloroform : methanol : H<sub>2</sub>O (65:25:4). Plates were sprayed with 0.5% α-naphthol in 50% methanol and then with 95% sulfuric acid. Glycolipids were visualized as purple bands by a final heating step. For the glycolipid analysis of bacteria grown under anaerobic fermentative growth conditions, colonies obtained on agar plates following incubated at 37°C in an anaerobic cabinet were scraped off the plates and suspended in 0.1M sodium citrate pH 4.7 for subsequent membrane lipid isolation and TLC analysis as described above. Representative data from three independent experiments are shown.

MALDI-TOF MS and MALDI-TOF MS/MS analysis of glycolipids

MALDI-TOF analysis of glycolipids was performed using a previously described method with some minor modifications (66). A total of 5 × 0.5 mg of lipids were spotted on silica plates and separated by TLC as described above. The silica matrix with the lipids from appropriate areas of the TLC plates were scraped off into glass tubes and extracted overnight at room temperature with 6 ml of a 1:1 methanol : chloroform mix for each sample. Next day, the silica matrix was removed by filtering the solutions through classic Sep-pak silica cartridges (Waters) pre-equilibrated with 6 ml methanol followed by 6 ml chloroform. The filtered samples with the extracted lipids were dried under
a stream of nitrogen. Dried lipids were suspended in 50 µl chloroform and aliquots mixed 1:1 with matrix. The matrix consisted of a 9:1 mixture of 2,5-dihydroxybenzoic acid and 2-hydroxy-5-methoxybenzoic acid (super-DHB, Sigma-Aldrich) at a final concentration of 10 mg/ml dissolved in chloroform : methanol at a ratio of 9:1. One µl sample was spotted onto disposable MSP 96 polished steel plate. As calibration standard, the peptide calibration standard II (Bruker) in 0.1% TFA was mixed 1:1 with IVD Matrix α-Cyano-4-hydroxycinnamic acid. The samples were analyzed on a MALDI Bi Typer Sirius system (Bruker Daltonik, Germany). The mass profiles were acquired using the FlexControl 3.4 software (Bruker Daltonik, Germany) with mass spectra scanned in the m/z range of 600 to 2,000. Spectra were recorded in the reflector positive ion mode (laser intensity 95%, ion source 1 = 10.00 kV, ion source 2 = 8.98 kV, lens = 3.00 kV, detector voltage = 2652 V, pulsed ion extraction = 150 ns). Each spectrum corresponded to an ion accumulation of 5,000 laser shots randomly distributed on the spot. Representative spectra from two independent experiments are shown. The obtained spectra were processed with default parameters using the FlexAnalysis v.3.4 software (Bruker Daltonik, Germany). For the MALDI-TOF MS/MS analysis, MS/MS fragmentation profiles were acquired on a 4800 Proteomics Analyzer (with TOF-TOF Optics, Applied Biosystems, plate: 384 Opti-TOF 123 mm × 84 mm AB Sciex NC0318050, 1016629) using the reflectron mode. Samples were analyzed operating at 20 kV in the positive ion mode. MS/MS mass spectrometry data were analyzed using the Data Explorer software version 4.9 from Applied Biosystems.

**Data availability**
The atomic coordinates and structure factors have been deposited in the Protein Data Bank under accession code 7B1R.
https://doi.org/10.2210/pdb7B1R/pdb

**Author contribution statement**
Chih-Hung Wu: Conceptualization, Investigation, Data analysis, Visualization, Writing – original draft preparation. Jeanine Rismondo: Investigation, Supervision, Data analysis, Writing – review & editing. Rhodri M. L. Morgan: Data analysis, Supervision, Writing – review & editing. Yang Shen: Investigation, Data analysis, Writing – review & editing. Martin J. Loessner: Data analysis, Writing – review & editing. Gerald Larrouy-Maumus: Investigation, Data analysis, Writing – review & editing Paul S. Freemont: Conceptualization, Funding acquisition, Supervision, Data analysis, Writing – review & editing. Angelika Gründling: Conceptualization, Funding acquisition, Data analysis, Supervision, Writing – original draft preparation.

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**Conflict of Interest**
The authors declare no conflicts of interest in regards to this manuscript.
| Possible fatty acid chain length | Chemical formula | Predicted molecular mass | Wild-type | ΔgtaB P<sub>hyper-gtaB</sub> | ΔgtaB P<sub>hyper-yngB</sub> |
|---------------------------------|------------------|--------------------------|-----------|-----------------------------|-----------------------------|
| **Top band: Glc<sub>2</sub>-DAG** |                  |                          |           |                             |                             |
| (30:0)                          | C<sub>45</sub>H<sub>84</sub>Na<sub>1</sub>O<sub>15</sub> | 887.57                   | 887.51    | 887.94                      | 887.90                      |
| (31:0)                          | C<sub>46</sub>H<sub>86</sub>Na<sub>1</sub>O<sub>15</sub> | 901.59                   | 901.95    | 901.97                      | 901.81                      |
| (32:0)                          | C<sub>47</sub>H<sub>88</sub>Na<sub>1</sub>O<sub>15</sub> | 915.60                   | 915.95    | 915.87                      | 915.91                      |
| **Mid band: Glc<sub>3</sub>-DAG** |                  |                          |           |                             |                             |
| (30:0)                          | C<sub>51</sub>H<sub>94</sub>Na<sub>1</sub>O<sub>20</sub> | 1049.62                  | 1049.85   | 1049.90                     | 1049.77                     |
| (31:0)                          | C<sub>52</sub>H<sub>96</sub>Na<sub>1</sub>O<sub>20</sub> | 1063.64                  | 1063.85   | 1063.90                     | 1063.93                     |
| (32:0)                          | C<sub>53</sub>H<sub>98</sub>Na<sub>1</sub>O<sub>20</sub> | 1077.65                  | 1077.85   | 1077.38                     | 1077.90                     |
Figure 1. YngB crystal structure and proposed UDP-glucose binding site residues. A, crystal structure of YngB shown in cartoon representation. The protein crystalized as a dimer in the asymmetric unit, and individual monomers are shown in orange and cyan respectively. B, overlay of the *B. subtilis* YngB (cyan) and the UDP-glucose bound *H. pylori* GalU*HP* (3JUK) (grey) structures. The structures are shown in ribbon representation, with UDP-glucose and UDP-glucose binding residues shown in stick representation. C, enlarged view of the substrate binding site from the alignment shown in panel B. The magenta sphere represents a magnesium ion; ionic interactions with the magnesium ion and hydrogen bonds are shown as dashed lines. Proposed UDP-glucose binding residues in YngB are Gly110, Gln105, Ala13, Gly14, Glu29, and Lys28 interacting with the uridine moiety; Asp133 chelating the magnesium ion; Asp134, Val 204, Gly172, and Glu191 interacting with the glucose moiety; and Lys 192 interacting with the diphosphate moiety.
Figure 2. Enzyme activity of GtaB and YngB. (A-B), Michaelis-Menten kinetics using purified GtaB or YngB proteins and A, increasing concentrations of the substrate glucose-1-phosphate (G-1-P) or B, increasing concentrations of the substrate UTP. The measured apparent $K_m$ values for glucose-1-phosphate were $45.6 \pm 3.24 \ \mu$M (GtaB) and $42.1 \pm 20.2 \ \mu$M (YngB) and for UTP, $49.5 \pm 10.2 \ \mu$M (GtaB) and $62.9 \pm 13.8 \ \mu$M (YngB), respectively. The experiment was performed three times with technical replicates. A representative graph from one experiment and plotting the mean and SD from the technical replicates is shown. The SD values for some data points were too small to be displayed on the graph. Michaelis-Menten curves were produced with Prism and $K_m$ values given are the mean ± SD from the three independent experiments.
Figure 3. Microscopy analysis and detection of glucose modifications on WTA produced by aerobically grown wild-type *B. subtilis*, mutant and complementation strains. The indicated *B. subtilis* strains were grown under aerobic growth conditions to mid-log phase in LB medium supplemented with IPTG and glucose. Bacteria were prepared for microscopy analysis and stained with the fluorescently labelled Alexa Fluor™ 594 Concanavalin A lectin to detect glucose modifications on WTA as described in the experimental procedure section. The experiment was performed 3 times and representative phase contrast, fluorescence and merged images are shown for each strain. Scale bars represent 10 µm.
Figure 4. NMR and UPLC-MS analysis of WTA isolated from wild-type *B. subtilis*, mutant and complementation strains. *A*, $^1$H NMR spectra of WTA isolated from wild-type *B. subtilis* (WT $P_{\text{hyper}}$), mutant $\Delta gtaB P_{\text{hyper}}$, and complementation strains $\Delta gtaB P_{\text{hyper}}-gtaB$ and $\Delta gtaB P_{\text{hyper}}-yngB$. The peak at 5.2 ppm (highlighted in orange) is likely derived from the hydrogen atom at the anomeric carbon of glucose residues on WTA (23). The experiment was performed twice, and similar results were obtained with the peak at 5.2 ppm absent in the $gtaB$ mutant but present in all other strains. The spectra from one experiment are shown. *B*, UPLC-MS spectra of hydrolyzed WTA samples derived from the same strains as described in *A*. The fragment with a mass signal of 253.09 (m/z) corresponding to glucose-glycerol species is absent in the $gtaB$ mutant but present in all other strains.
Figure 5. TLC analysis and detection of glycolipids produced by aerobically grown wild-type *B. subtilis*, mutant and complementation strains. Total membrane lipids were isolated from different *B. subtilis* strains following overnight growth under aerobic conditions in LB medium supplemented with IPTG and glucose. Lipids were separated by TLC and glycolipids visualized by spraying the plates with α-naphthol and 95% sulfuric acid and heating. Strains used were *A*, wild-type *B. subtilis* (WT *P* hyper), the gtaB mutant ∆gtaB *P* hyper, and complementation strains ∆gtaB *P* hyper-gtaB and ∆gtaB *P* hyper-yngB; *B*, wild-type *B. subtilis* (WT *P* hyper), the gtaB/yngB double mutant ∆gtaBΔyngB *P* hyper and complementation strains ∆gtaBΔyngB *P* hyper-gtaB and ∆gtaBΔyngB *P* hyper-yngB; and *C*, wild-type *B. subtilis* (WT *P* hyper), the yngB mutant ∆yngB *P* hyper, and complementation strain ∆yngB *P* hyper-yngB. Three independent experiments were performed, and a representative TLC plate image is shown. (*) marks likely Glc2-DAG and (**) likely Glc1-DAG glycolipid bands as assessed by MALDI-TOF mass spectrometry.
Figure 6. MALDI-TOF analysis of glycolipids isolated from wild-type *B. subtilis*, mutant and complementation strains. The glycolipid bands marked with * (top band) and ** (middle band) in Figure 5 were extracted from TLC plates for samples derived from wild-type *B. subtilis* (WT \(P_{\text{hyper}}\)), the \(\Delta gtaB\) mutant \(\Delta gtaB P_{\text{hyper}}\), and complementation strains \(\Delta gtaB P_{\text{hyper}}-gtaB\) and \(\Delta gtaB P_{\text{hyper}}-yngB\) and analyzed by MALDI-TOF M/S in the positive ion mode. Lipids were analyzed from 3 independent experiments and the spectra for one experiment is shown. The observed and expected masses of sodium adducts of specific glycolipids are summarized in Table 1. The data indicate that the top bands (*) corresponds to Glc2-DAG and the middle bands (**) to Glc3-DAG.
Figure 7. Detection of glucose modification on WTA and glycolipid analysis of wild-type and mutant B. subtilis strains following growth under anaerobic conditions. Wild-type B. subtilis (WT Phyper) and the gtaB and yngB single and double mutant strains ΔgtaB Phyper, ΔyngB Phyper, and ΔgtaBΔyngB Phyper were grown in an anaerobic chamber under fermentative growth condition on 2 x YT plates. A, bacteria were prepared for microscopy analysis and stained with the fluorescently labelled Alexa Fluor™ 594 Concanavalin A lectin to detect glucose modifications on WTA as described in the experimental procedure section. The experiment was performed 3 times and representative phase contrast, fluorescence and merged images are shown for each strain. Scale bars represent 10 µm. B, total membrane lipids were isolated from the different B. subtilis strains following growth in an anaerobic chamber, separated by TLC and glycolipids visualized by spraying the plates with α-naphthol and 95% sulfuric acid and heating. As control, a lipid sample isolated from the wild-type strain grown aerobically was run alongside the other samples. The experiment was performed 3 times and a representative TLC plate is shown.
Figure 8. Proposed glycosylation pathway involving the UGPase YngB and the YkcC-YngA-YkcB multi-component transmembrane glycosylation system. A, the transcription factor YclJ activates the transcription of genes belonging to the ykcBC and yngABC operons. B, the B. subtilis LTA glycosylation model requires CsbB, GtcA and YfhO for the transfer of GlcNAc onto the LTA. The glycosyltransferase CsbB produces C$_{55}$-P-GlcNAc, which is subsequently transported across the membrane by GtcA and the GlcNAc is finally attached to the LTA polymer on the outside of the cell by the multimembrane spanning GT-C-fold glycosyltransferase YfhO. C, proposed glycosylation pathway leading under anaerobic growth conditions to the transfer of glucose onto an uncharacterized target in the cell envelope. As shown in this study, YngB is a functional UGPase that can produce UDP-glucose and we hypothesize, that the predicted glycosyltransferase YkcC produces C$_{55}$-P-glucose, which is transported across the membrane by YngA. The glucose residue is then transferred by the predicted multimembrane spanning GT-C-fold glycosyltransferase YkB to an unknown target within the bacterial cell envelope. Panel B was adapted from model figure presented in Rismondo et al., 2018 (19) and Rismondo et al., 2020 (20).
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