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The *sps* Genes Encode an Original Legionaminic Acid Pathway Required for Crust Assembly in *Bacillus subtilis*

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ABSTRACT The crust is the outermost spore layer of most *Bacillus* strains devoid of an exosporium. This outermost layer, composed of both proteins and carbohydrates, plays a major role in the adhesion and spreading of spores into the environment. Recent studies have identified several crust proteins and have provided insights about their organization at the spore surface. However, although carbohydrates are known to participate in adhesion, little is known about their composition, structure, and localization. In this study, we showed that the spore surface of Bacillus subtilis is covered with legionaminic acid (Leg), a nine-carbon backbone nonulosonic acid known to decorate the flagellin of the human pathogens Helicobacter pylori and Campylobacter jejuni. We demonstrated that the spsC, spsD, spsE, spsG, and spsM genes of Bacillus subtilis are required for Leg biosynthesis during sporulation, while the spsF gene is required for Leg transfer from the mother cell to the surface of the forespore. We also characterized the activity of SpsM and highlighted an original Leg biosynthesis pathway in B. subtilis. Finally, we demonstrated that Leg is required for the assembly of the crust around the spores, and we showed that in the absence of Leg, spores were more adherent to stainless steel probably because of their reduced hydrophilicity and charge.

IMPORTANCE *Bacillus* species are a major economic and food safety concern of the food industry because of their food spoilage-causing capability and persistence. Their persistence is mainly due to their ability to form highly resistant spores adhering to the surfaces of industrial equipment. Spores of the *Bacillus subtilis* group are surrounded by the crust, a superficial layer which plays a key role in their adhesion properties. However, knowledge of the composition and structure of this layer remains incomplete. Here, for the first time, we identified a nonulosonic acid (Leg) at the surfaces of bacterial spores (*B. subtilis*). We uncovered a novel Leg biosynthesis pathway, and we demonstrated that Leg is required for proper crust assembly. This work contributes to the description of the structure and composition of *Bacillus* spores which has been under way for decades, and it provides keys to understanding the importance of carbohydrates in *Bacillus* adhesion and persistence in the food industry.

KEYWORDS *Bacillus subtilis*, bacterial adhesion, crust, legionaminic acid, nonulosonic acid, spores

The *Bacillaceae* are able to engage in a process of cellular differentiation called sporulation, which results in the formation of a mature spore capable of surviving adverse environmental conditions and of being dispersed into the environment. Because of these properties, *Bacillus* spores are ubiquitous in the environment and

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Accepted 6 July 2020 Published 18 August 2020 contaminate many raw materials of the food industry, e.g., in the dairy industry, causing heavy economic losses and a health risk to consumers (1).

The Bacillus spore consists of a series of concentric layers. Its core contains highly condensed DNA wrapped around proteins called "small soluble acid proteins," which protect the DNA from physicochemical stresses. The core is surrounded by an inner membrane and a germ cell wall that are themselves covered by a spore-specific peptidoglycan called cortex (2, 3). The cortex is surrounded by an outer membrane, which is itself surrounded by the three-layered coat, composed of at least 80 different proteins (4). The coat proteins are recruited and assembled by morphogenetic proteins, with at least one morphogenetic protein per coat layer: SpoIVA for the basement layer, SafA for the inner coat, and CotE for the outer coat (5-8). In Bacillus subtilis and most of the Bacillus strains producing spores devoid of an exosporium, the outermost layer of spores is the crust (9). It is made of proteins and carbohydrates, and it plays a major role in the surface and adhesion properties of spores (9). Thus, the mechanical removal of the crust renders the spore less hydrophilic and more adherent to stainless steel (9). Therefore, the presence or absence of this outermost layer could determine the spreading properties of *B. subtilis* spores and their resistance to the cleaning-in-place procedures used in food industries (1, 10).

At least six proteins have been identified as constituents of the crust: CotV, CotW, CotX, CotY, CotZ, and CgeA (11–15). The crust structure is mainly provided by the CotV, CotX, and CotY proteins, among which, CotY would be the major structural component (15). Interestingly, CotV and CotX share homologue domains and putative N-glycosylation motifs, which make these proteins candidates for glycosylation (15). McKenney and colleagues suggested that CotX, CotY, and CotZ are the crust morphogenetic proteins (12), and recent studies have confirmed that these proteins play a major role in crust assembly (14, 15). CotZ could be the main morphogenetic protein, since it is involved in the proper localization of most of the crust proteins (14). CotV could also be involved in the crust assembly by propagating the crust structure from the polar cap-like structure to the middle part of the spore (15). CotZ, which required CotE and CotO but none of the crust proteins for localization, seems to have an important anchoring function for the crust in addition to its assembly function (15). CotW plays a role in maintaining the structural integrity of the crust and might play an anchoring role at the interface of crust and coat (14, 15). Finally, transmission electron microscopy (TEM) experiments showed that the crust of the cotX and cgeA mutants is assembled, but only loosely attached to spores, thus suggesting that CgeA and CotX play roles in the interaction of the crust with the coat (14).

While the crust protein interaction network begins to be better described, the localization, nature, and structure of the glycans on the spore surface remain mostly unknown. It was shown that the spore surface of B. subtilis contains at least rhamnose, galactose, guinovose, glucosamine, and muramic lactam (9, 16). Although the structure of the glycans composed of these monosaccharides is unknown, it was suggested that the spore surface contains at least two different glycans: one associated with the outer coat and at least one other linked to the crust (15, 17). The spsM, spsABCDEFGIJKL, yfnHGFED, ytdA-ytcABC, and cgeAB-cgeCDE genes were identified as participating in the morphology and the surface properties of the crust (15–18). Mutations in the sps genes result in the production of less hydrophilic spores, devoid of perceivable halo after negative staining with India ink, thus suggesting these genes play a role in spore surface glycosylation (16–18). Contrastingly, mutations in each of the yfnHGFED genes do not affect the spore hydrophilicity but modify the crust structure and expand the glycan layer (15, 17). Finally, spores of a *cgeD* mutant strain are less hydrophilic and the superficial saccharide layer is expanded, just as it is with the mutants of the yfnHGFED genes (17). These data resulted in the identification of some of the enzyme-encoding genes that synthesize the superficial spore glycans. However, to date, both these genes' functions and the biosynthetic pathway in which they are involved remain unknown, with the notable exception of the spsIJKL genes, known to encode the biosynthesis pathway converting D-glucose-1-phosphate to dTDP-L-rhamnose (19-23).



FIG 1 The *sps* genes participate in the surface and adhesion properties of *B. subtilis* spores. (A) Schematic representation of the *spsABCDEFGUKL* and *spsM* genes in *B. subtilis* PY79. Transcriptional activators and promoters are shown with arrows and broken arrows, respectively. Potential stem-loop structure is indicated with a lollipop. The annotation of the *sps* genes is presented in Table S1 in the supplemental material. (B) Surface hydrophilicity of spores of the PY79 and PY79 $\Delta spsABCDEF$ ($\Delta spsABCDEF$) strains evaluated by MATH assays. (C) Surface hydrophilicity of spores of the PY79, PY79 $\Delta spsM$ ($\Delta spsM$), and PY79 $\Delta spsM$ amy*E::spsM* ($\Delta spsM$ amy*E::spsM*

Here, we sought to define the functions of the *sps* genes and their roles in crust biosynthesis. Through an approach that combines genetic and biochemical methods, we demonstrated that six of the *sps* genes encode an original CMP-legionaminic acid (CMP-Leg) pathway, and we showed that legionaminic acid (Leg) is required for proper crust assembly. This work contributes to a better understanding of the outermost spore layer composition and structure that influence the adhesion and spreading of *B. subtilis* in the environment.

RESULTS

The sps genes contribute to the surface and adhesion properties of Bacillus subtilis spores. The sps genes are distributed in two loci on the *B. subtilis* chromosome (Fig. 1A). The first locus contains the spsABCDEFGIJKL genes and the second contains the spsM gene. The two loci are transcribed in the mother cell from σ^{K} -dependent promoters located upstream from the spsA and spsM genes (16, 18, 24–26). This transcriptional regulation indicates that the products of the sps genes are produced concomitantly in the mother cell during the late stage of sporulation. To evaluate the role of the sps genes, mutant strains of the spsABCDEF and spsM genes were con-

structed in the B. subtilis PY79 strain (subsequently named PY79). Spores were produced, and their surface properties were characterized. The hydrophilicity of the spores was assessed through microbial-adhesion-to-hydrocarbons (MATH) (Fig. 1B and C). Most PY79 spores remained in the aqueous phase throughout the MATH experiment, which reflected a marked hydrophilic character. In contrast, the percentage of spores in the aqueous layer for the $\Delta spsABCDEF$ and $\Delta spsM$ mutant strains decreased over time and tended toward zero after 240 s of agitation. These results indicate that the ΔspsABCDEF and ΔspsM mutant strains are less hydrophilic than the PY79 strain, as is consistent with previous studies (16-18). In addition, the complementation of the ΔspsM mutant strain restored the hydrophilic properties of the PY79 strain (Fig. 1C). This indicates that the phenotype observed with the $\Delta spsM$ mutant strain is not due to a polar effect of the mutation on flanking genes expression. The overall spore charge at neutral pH was then evaluated using zetametry (Fig. 1D). The zeta potentials of the $\Delta spsABCDEF$ and $\Delta spsM$ spores were around -20 mV, indicating that they are less charged than the PY79 spores (-46.4 mV). These data demonstrate that the spsABCDEF and spsM gene products contribute to the hydrophilicity and the negative charge of spores that may lead to changes in spore adhesion properties. The role of the sps genes in the adhesion of spores to stainless steel coupons was therefore examined (Fig. 1E). With the PY79 strain, an average of 1.8×10^3 spores adhered to the stainless-steel coupons, while 12 and 5 times more spores adhered with the $\Delta spsABCDEF$ and $\Delta spsM$ mutant strains, respectively, thus indicating that the sps gene products influence spore adhesion properties.

The sps genes are required for crust assembly. The surface and adhesion properties of spores of the $\Delta spsABCDEF$ and $\Delta spsM$ mutant strains were very similar to those of spores lacking the crust (see Fig. S1 in the supplemental material), which suggests that the *sps* genes are important for proper crust assembly. To test this assumption, the spore morphology of the PY79, $\Delta spsABCDEF$, and $\Delta spsM$ strains was observed by TEM using spore sections contrasted with ruthenium red. This staining enables the observation of the carbohydrate-rich layer on the spore surface (27). The crust forms an exosporium-like structure that is clearly visible for PY79 spores (black arrows) (Fig. 2). This exosporium-like structure was no longer observed on the spore surfaces of the $\Delta spsABCDEF$ and $\Delta spsM$ mutant strains, thereby confirming that the *sps* genes are required for proper crust assembly. However, a loose low-contrast material was still visible, mostly at the spore poles (blue arrows). This could be reminiscent of a polar cap-like structure from which the crust propagates to the middle part of the spore. These observations suggest that the products of the *sps* genes participate in the propagation of the crust or in anchoring the crust to the outer coat.

The sps genes participate in spore surface glycosylation. The function of the enzymes encoded by the sps genes and the biosynthetic pathways in which these enzymes are involved have not been described. To better define the role of the sps genes, the amount of neutral sugars on the PY79 spore surface was measured. For this purpose, spores were mechanically treated with a French press to remove their surface fraction without damaging the spore coat (9). This surface fraction contains the crust proteins and the potentially associated carbohydrates as well as the constituents located between the outer coat proteins and the crust. The amount of neutral sugars in the surface fraction was then measured by the phenol-sulfuric method (28) (see Fig. S2). It was significantly lower in the surface fraction of the $\Delta spsABCDEF$ and $\Delta spsM$ mutant strains than in the PY79 strain. Knowing the putative function of the sps genes (Table S1), this might indicate that the *spsABCDEF* and *spsM* genes are involved in the production or the attachment of one or several neutral sugars on the spore surface. However, given that the $\Delta spsABCDEF$ and $\Delta spsM$ mutant strains lack most of the crust (Fig. 2), it is also possible that the neutral sugar decrease was an indirect consequence of the crust assembly defect, such as the loss of one or several crust-linked glycan(s).

The surface fraction of spores contains legionaminic acid. Similar phenotypes being obtained with the $\Delta spsABCDEF$ and $\Delta spsM$ mutant strains, these genes probably



FIG 2 The *sps* genes are required for crust assembly. TEM images of spore sections after ruthenium red staining. The experiments were performed with the spores of the PY79, PY79 $\Delta spsABCDEF$ ($\Delta spsABCDEF$), and PY79 $\Delta spsM$ ($\Delta spsM$) strains. White arrows, coat layers; black arrows, crust; blue arrows, cap-like structures.

participate in the same carbohydrate biosynthetic pathway. To better understand the role of the Sps proteins in the spore surface glycosylation, in silico analysis was performed using RaptorX software with the primary sequences of the SpsA-G and SpsM proteins (29). The RaptorX software predicts a tridimensional structure of a query sequence and compares this predicted structure to known protein structures from the Protein Data Bank. The outputs of this analysis are presented in Table S1 and they are further described in the discussion section. By using gene annotations and RaptorX predictions, a predictive biosynthetic pathway was constructed (Fig. 3A). This pathway is reminiscent of the pseudaminic acid (Pse) biosynthetic pathway of Helicobacter pylori and the Leg biosynthetic pathway of Campylobacter jejuni (30, 31). Pse and Leg are nine-carbon-backbone sialic acid analogs known to be crucial for flagellin glycosylation and virulence of C. jejuni and H. pylori (32-36). To ensure that the spore surface contains sialic acid-related compounds, the potential nonulosonic acids (NulOs) from the surface fraction of the PY79 strain were derivatized by 1,2-diamino-4,5-methylenedioxybenzene (DMB), a fluorogenic reagent that shows high specificity for NulOs, and analyzed by reverse-phase high-performance liquid chromatography coupled to fluorescence detector (RP-HPLC-FL) (Fig. 3B). RP-HPLC-FL analysis showed three peaks with retention times of \sim 8.5 min (peak 1), 14.9 min (peak 2), and 20.5 min (peak 3), which were then analyzed in liquid chromatography mass spectrometry (LC-MS). Unlike in peaks 2 and 3, an MS signal at m/z 451 was identified in peak 1, corresponding to the [M+H]⁺ adduct of a compound with a molecular mass of 450.25 Da (data not shown).



FIG 3 The *sps* genes encode a pathway required for legionaminic acid addition on the spore surface of *B. subtilis*. (A) Predicted Leg pathway in *B. subtilis*: I, UDP-*N*-acetyl- α -D-glucosamine (UDP-GlcNAc); II, UDP-2-acetamido-2,6-dideoxy- α -*D*-xylo-hexose-4-ulose (UDP-4-keto-6-deoxy-GlcNAc); III, UDP-4-amino-4,6-dideoxy-*N*-acetyl- α -D-glucosamine (UDP-4-amino-6-deoxy-GlcNAc); IV, UDP-2,4-diacetamido-2,4,6-trideoxy- α -D-glucose (UDP-2,4-diNAc-6-deoxy-Glc); V, 2,4-diacetamido-2,4,6-trideoxy- α -D-glucose (UDP-2,4-diNAc-6-deoxy-Glc); V, 2,4-diacetamido-2,4,6-trideoxy- α -D-glaco-nonulosonic acid (legionaminic acid or Leg); VII, CMP-5,7-diacetamido-3,5,7,9-tetradeoxy-D-glycero- β -D-galacto-nonulosonic acid (CMP-legionaminic acid or CMP-Leg). (B) Chromatograms of the RP-HPLC-FL experiments performed on the surface fractions of spores of the PY79, PY79 Δ *spsABCDEF* (Δ *spsABCDEF*), and PY79 Δ *spsM* (Δ *spsM*) (Δ *spsM*) strains. The RP-HPLC-FL experiments were carried out on at least three independent surface fractions for each strain. One representative chromatogram is presented. (C) Identification by LC/ESI-MS³ of DMB-Leg5Ac7Ac. Representative MS³ spectra of DMB-Leg5Ac7Ac, [M+H]⁺ at *m/z* 451. The ion [M+H-18]⁺ at *m/z* 433 is indicated by the black diamond. (D) Structure of DMB ([M+H]⁺ = 229), DMB-Leg ([M+H]⁺ = 541), and of the major structure fragments ([M+H]⁺ = 433, [M+H]⁺ = 415.1, [M+H]⁺ = 376).

This signal corresponds to the expected mass of DMB-Leg, DMB-Pse, or one of their isomers that exhibits a $[M+H]^+$ value at m/z 451 (37, 38). The identification was then confirmed by MS³ (Fig. 3C and D). MS³ analysis showed recurrent losses of H₂O (m/z 433, m/z 415, and m/z 356) and *N*-acetyl groups (m/z 374), which agrees with the fragmentation pattern previously obtained with *N*-acetylneuraminic acid (Neu5Ac) and other NulOs (36, 37). Furthermore, the retention time of peak 1 perfectly matched that of the Leg standard (see Fig. S3), thus indicating that the surface fraction of PY79 spores contains Leg.

The *sps* genes expressed during the late stage of sporulation are required for legionaminic acid addition on the spore surface. To determine whether the *sps* genes are involved in Leg biosynthesis, the surface fractions of the PY79, $\Delta spsABCDEF$, and $\Delta spsM$ spores were analyzed by RP-HPLC-FL (Fig. 3B). Interestingly, Leg (peak 1) was undetectable in the surface fraction of the $\Delta spsABCDEF$ and $\Delta spsM$ mutant strains.

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These results demonstrate that spsM and at least one of the spsABCDEF genes are required for Leg biosynthesis or transfer to the spore surface. To study the kinetics of the sps genes transcription, a chromosomal transcriptional fusion was constructed by inserting the promoter region of the spsA gene fused to a promoterless mCherry gene into the *amyE* gene, and the resulting strain was designated PY79 *amyE*::P_{spsA}-mCherry. The kinetics of sps genes expression was then determined in the Spo8 sporulation medium (see Fig. S4A). The PY79 amyE::P_{spsA}-mCherry strain displayed mCherry fluorescence 6 h after the transition to the stationary phase (t6) and peaked at t8, which is in agreement with a previous study (18). Transcription of the sps genes is under the control of σ^{K} , indicating that the Sps enzymes and their products are localized in the mother cell (16, 18, 24-26). To define the timing of Leg production in the mother cell, a kinetics analysis of Leg production was performed under the same growth conditions (Fig. S4B). Consistently with the transcription kinetics, the Leg was detectable in the PY79 strain at t8 and peaked at t10, approximately 2 h following the transcription peak. Together, these results indicate that Leg is produced in the mother cell of sporulating cells during the late stage of sporulation.

SpsM is the first enzyme of the legionaminic acid biosynthesis pathway. The predictive three-dimensional structure of SpsM is close to that of the enzymes CapE, WbjB, and PseB (Table S1). These enzymes catalyze the C-4/C-6 dehydration and C-5 epimerization of the UDP-GlcNAc to produce UDP-4-keto-6-deoxy-AltNAc, and they share a conserved catalytic domain with a distinctive $T-(x)_{\alpha}-M/Y-(x)_{3}-K$ catalytic triad (32, 39-41). In Staphylococcus aureus, the replacement of the methionine by an alanine in the catalytic triad of CapE causes a partial activity loss for this enzyme (42). To determine whether SpsM also shares this catalytic domain, a protein sequence alignment was performed (Fig. 4A). This analysis identified, in the SpsM protein sequence, a $T-(x)_9$ -M- $(x)_3$ -K catalytic triad in a well-conserved putative catalytic domain. Mutations were introduced into the coding sequence of the *spsM* gene by directed mutagenesis to replace methionine 146 or lysine 150 with an alanine (Fig. 4A). Both mutations caused a decrease in the hydrophilicity and global spore charge (Fig. 4B and C). In addition, M146A and K150A mutations led to a partial and a total loss of Leg in the surface fraction of spores (Fig. 4D). These results indicate that the M146 and K150 amino acids are in the SpsM catalytic site and are required for full SpsM activity. CapE, WbjB, PgIF, PseB, and Pen-Pal share C-4/C-6 dehydratase activity and use a nucleotideactivated GlcNAc as the substrate (32, 39-41, 43), thus suggesting that SpsM shares these properties. However, the activity of these enzymes slightly varies from one to the other. For example, PgIF only has a C-4/C-6 dehydratase activity, while PseB has an additional C-5 epimerase activity (40) (Fig. 4E). To better define SpsM activity, the pen or pal genes of Bacillus thuringiensis ATCC 35646 and the pseB, legB, and pglF genes of Campylobacter jejuni NCTC11168 were introduced downstream of the spsM promoter into the *amyE* locus of the PY79 Δ*spsM* mutant strain. The enzymatic reactions catalyzed by Pen-Pal, PgIF, LegB, and PseB are summarized in Fig. 4E (30, 32, 40, 43). Then, the relative amounts of Leg in the mother cells of the *AspsM amyE::PspsM-pen*, *AspsM* amyE::P_{spsM}-pal, ΔspsM amyE::P_{spsM}-pen-pal, ΔspsM amyE::P_{spsM}-pseB, ΔspsM amyE:: P_{spsM} -pgIF, and $\Delta spsM$ amyE:: P_{spsM} -legB strains were evaluated at t10 by RP-HPLC-FL (Fig. 4F). The addition of the pen, pal, legB, or pseB gene to the ΔspsM mutant strain did not restore, or only partially restored, Leg production in the mother cells. Contrastingly, the addition of the *pen-pal* or *pglF* gene fully restored Leg production, thus indicating that SpsM catalyzes the C-4/C-6 dehydration of the UDP-GlcNAc to produce UDP-4keto-6-deoxy-GlcNAc. Together, these results also demonstrate that SpsM is the first enzyme of the CMP-Leg pathway in B. subtilis (Fig. 3A).

spsC, *spsD*, *spsE*, and *spsG* are required for Leg biosynthesis, while *spsF* is required for Leg transfer to the forespore surface. In the CMP-Leg biosynthesis pathway of *B. subtilis* proposed in Fig. 3A, the SpsM product is successively processed by SpsC, SpsD, SpsG, SpsE, and SpsF. It was shown above that one or more of the *spsABCDEF* genes are involved in Leg biosynthesis or transfer to the forespore surface



FIG 4 SpsM is a C-4/C-6 dehydratase using UDP-GlcNAc as a substrate. (A) Alignment of the protein sequence of SpsM with the protein sequences of CapE (Staphylococcus aureus), WbjB (Acinetobacter baumannii), PgIF (Campylobacter jejuni), PseB (Helicobacter pylori), and Pen (Bacillus thuringiensis). The PDB number corresponding to each sequence is indicated in parentheses. The three conserved residues of the catalytic triad are black boxed. The arrows indicate residues modified by directed mutagenesis. The experiments presented in panels B. C. and D. were carried out with PY79 AspsM amvF::spsM (spsM). PY79 AspsM amvF::spsM M146A (M146A), and PY79 ΔspsM amyE::spsM K150A (K150A) strains. (B) Surface hydrophilicity of spores evaluated by MATH assays. (C) Surface charge of spores evaluated by zetametry assays. ****, $P \le 0.0001$ for M146A or K150A versus *spsM* by Mann-Whitney test. (D) Relative amounts of Leg in the surface fractions of spores measured by RP-HPLC-FL. The results are relative to the amount of Leg of the PY79 $\Delta spsM$ amyE::spsM strain. nd, not detectable; ***, $P \leq 0.001$ for M146A or K150A versus spsM by Welch's t test. (E) Enzymatic reactions catalyzed by Pen-Pal, PgIF, LegB, and PseB in B. thuringiensis and C. jejuni. The solid arrows represent the enzymatic reactions catalyzed by the enzymes. The dotted arrows represent the remaining parts of the pathways and point to the final products. (F) Relative amounts of Leg in the mother cells of the PY79 ΔspsM amyE::spsM (spsM), PY79 ΔspsM amyE::pen (pen), PY79 AspsM amyE::pal (pal), PY79 AspsM amyE::pen-pal (pen-pal), PY79 AspsM amyE::pglF (pglF), PY79 AspsM amyE::legB (legB), and PY79 ΔspsM amyE::pseB (pseB) strains measured at t10 by RP-HPLC-FL. The results are relative to the amount of Leg in the PY79 $\Delta spsM$ amyE::spsM strain. ns, not significant; ***, $P \le 0.001$; ****, $P \le 0.0001$ for pen, pal, pen-pal, pg/F, legB, or pseB versus spsM by Welch's t test. The error bars represent the SDs of the means.

(Fig. 3B). However, the involvement of each of these genes in the CMP-Leg pathway remained to be verified. To check the validity of the CMP-Leg pathway proposed in Fig. 3A, mutants of the *spsD* and *spsF* genes were constructed in the PY79 strain. The *spsD* and *spsF* genes encode a putative acetyltransferase and a putative cytidylyltransferase, respectively (Table S1). The spores of both mutants were less hydrophilic and negatively charged than those of the PY79 strain (see Fig. S5A, B, and E). The complementation of the $\Delta spsD$ mutant strain restored the hydrophilic properties of spores, while the complementation of the $\Delta spsF$ mutant strain is discussed in Text S1. The surface fraction of spore was extracted and the Leg amount quantified (Fig. 5A). In parallel, the amounts of Leg in the mother cells of the PY79, $\Delta spsD$, and $\Delta spsF$ strains were evaluated at t10 (Fig. 5B). Leg was absent from the surface fractions of the $\Delta spsF$ mutant

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FIG 5 The *spsC*, *spsD*, *spsE*, and *spsG* genes are required for Leg biosynthesis, while *spsF* is required for Leg transfer to the forespore surface. The amounts of Leg were measured in the surface fractions of spores (A and C) and the mother cells of sporulating *B. subtilis* cells at t10 (B and D) by RP-HPLC-FL. The experiments were performed with PY79, PY79 *ΔspsD* (*ΔspsD*), and PY79 *ΔspsF* (*ΔspsF*) strains (A and B) or 168, 168 *ΔspsC* (*ΔspsC*), 168 *ΔspsE* (*ΔspsE*), and 168 *ΔspsG* (*ΔspsG*) strains (C and D). The results were standardized by the OD₆₀₀ of the spore preparations (A and C) or the OD₆₀₀ of the cultures at t10 (B and D). ns, not significant; nd, not detected; nt, not statistically tested because the data set did not pass the Shapiro-Wilk normality test (two values of three were equal to zero). The error bars represent the SDs of the means. ****, $P \leq 0.0001$ for Δ versus PY79 or 168 by Welch's *t* test.

strains and from the mother cell of the $\Delta spsD$ mutant strain during sporulation. In contrast, the amount of Leg measured in the mother cell of the $\Delta spsF$ mutant strain was comparable to that of the PY79 strain. These results demonstrate that the spsD gene is required for Leg production, while the spsF gene is required for Leg transfer from the mother cell to the forespore. To test whether the spsC, spsE, and spsG genes are also required for Leg biosynthesis, we used the $\Delta spsC$, $\Delta spsE$, and $\Delta spsG$ mutants constructed in B. subtilis 168 strain (44). The SpsC, SpsE, and SpsG proteins of the 168 strain shared 100% of identity with the SpsC, SpsE, and SpsG proteins of the PY79 strain. In addition, the amount of Leg in the spore surface fraction of the 168 strain was similar to that of the PY79 strain (Fig. 5A and C and Fig. S3). Leg was absent from the spore surface fraction of the $\Delta spsC$ and $\Delta spsE$ mutant strains, and the Leg amount was strongly reduced in the $\Delta spsG$ mutant strain (Fig. 5C). Moreover, the quantification of Leg in the mother cells of the $\Delta spsC$, $\Delta spsE$, and $\Delta spsG$ mutant strains did not detect significant amounts of Leg (Fig. 5D). These data demonstrate that spsC, spsE, and spsG genes are required for Leg biosynthesis and are in accordance with the CMP-Leg pathway proposed in Fig. 3A.

SpsA and SpsB are not the transferase of the CMP-Leg. The final acceptor(s) of the Leg on the forespore surface as well as the enzyme(s) involved in the transfer of the Leg to the final acceptor(s) remained to be identified. The *spsA* and *spsB* genes encode putative glycosyltransferases (Table S1). Glycosyltransferases represent a subclass of enzymes that catalyze glycoside linkage synthesis by the transfer of a monosaccharide from an activated donor substrate to an acceptor. Therefore, the SpsA and SpsB proteins were candidates for transferring the Leg from the CMP-Leg to one or more acceptor(s) on the forespore surface. To test this hypothesis, spores of the PY79 $\Delta spsA$ and PY79 $\Delta spsB$ mutant strains were produced. Spores of both mutant strains were less



FIG 6 Legionaminic acid is linked to the crust. The amounts of Leg were measured in the surface fractions of spores by RP-HPLC-FL. The experiments were performed with PY79, PY79 $\Delta spsA$ ($\Delta spsA$), and PY79 $\Delta spsB$ ($\Delta spsB$) strains (A) or PY79, PY79 *cotE::erm* (*cotE::erm*), PY79 *cotX::erm* (*cotX::erm*), and PY79 *cotZ::erm* (*cotZ::erm*) strains (B). The results were standardized by the OD₆₀₀ of the spore preparations. ns, not significant.; ****, $P \leq 0.0001$ for Δ versus PY79 by Welch's *t* test. The error bars represent the SDs of the means.

hydrophilic and negatively charged than spores of the PY79 strain, and complementations restored the hydrophilic properties of spores of the PY79 strain (Fig. S5C, D, and E). This indicates that the *spsA* and *spsB* genes are involved in spore surface maturation. To determine whether the modifications of the surface properties of spores of the $\Delta spsA$ and $\Delta spsB$ mutant strains were due to a defect in Leg transfer to the forespore surface, the surface fraction of both mutants was analyzed in RP-HPLC-FL (Fig. 6A). Surprisingly, the amounts of Leg in the surface fractions of the $\Delta spsA$ and $\Delta spsB$ mutant strains were similar to that of the PY79 stain, indicating that *spsA* and *spsB* genes are not required for Leg transfer to the forespore surface.

Legionaminic acid is linked to the crust. To determine whether the Leg is linked to the crust or the outer coat, the spore surface fractions of the PY79 *cotE::erm*, PY79 *cotX::erm*, and PY79 *cotZ::erm* mutant strains were analyzed in RP-HPLC-FL (Fig. 6B). It was shown previously that spores of the *cotE* mutant strain lack both the outer coat and the crust, while spores of the *cotZ* mutant strains only lack the crust (14, 45). The *cotX* mutant stain has an assembled crust, but it is loosely attached to spores (14). However, in our hands, we observed that the hydrophilicity of spores of the *cotX* mutant strain decreased during the washing of the spore preparations, which would indicate that the crust is lost under our experimental conditions (data not shown). Therefore, if the Leg was linked to the crust, a decrease of the Leg amount was expected in the surface fractions of the *cotE*, *cotX*, and *cotZ* mutant strains by comparison with that of the PY79 strain. In contrast, if the Leg was linked to the outer coat, a decrease of the Leg amount was only expected in the surface fraction of the *cotE* mutant strain. The inactivation of the *cotE*, *cotX*, and *cotZ* genes strongly reduced the amount of Leg in the surface fraction of spores, indicating that Leg is linked to the crust.

DISCUSSION

We previously showed in *B. subtilis* that the glycans of the crust are composed of rhamnose, quinovose, glucosamine and muramic lactam, this last being known to be part of the bacterial peptidoglycan (9). In this study, we showed that the surface of *B. subtilis* spores also contained Leg, a NulO identified for the first time on the surface of bacterial spores. NulOs are composed of nine-carbon-backbone monosaccharides differing by structural variations at the C-5, C-7, and C-9 positions (46). NulOs notably encompass sialic acids, such as Neu5Ac and the ketodeoxynonulosonic acid, as well as nonsialic acids, such as Leg and Pse. Biosynthesis of NulOs proceeds through a conserved enzymatic mechanism starting by the condensation of a 6-carbon monosaccharide with a 3-carbon pyruvate. The 9-carbon intermediate is then activated to a CMP-sugar, before being transferred to an acceptor. The CMP-Pse biosynthesis pathway of *H. pylori* is an example of a NulO pathway that has been extensively studied (31, 47,

48). In *H. pylori*, the nucleotide activated Pse is synthesized from UDP-GlcNAc by six consecutive enzymes. The CMP-Leg pathway in *C. jejuni* is quite similar, except that the sugar precursors are GDP linked, rather than UDP linked. It also differs in the epimerization performed at C-2, C-4, and C-5 of the 6-deoxy-hexoses intermediates, resulting in stereochemical differences at C-5, C-7, and C-8 of the final nonulosonates (30). In contrast to that of *H. pylori*, the Pse pathway of *B. thuringiensis* involves seven enzymes (43). The two pathways differ by the enzymes catalyzing the first reaction of the pathway, i.e., the synthesis of the UDP-4-keto-6-deoxy-AltNAc from the UDP-GlcNAc. In *H. pylori* this reaction is catalyzed by PseB, while it is catalyzed by Pen and Pal in *B. thuringiensis*. Interestingly, it was shown that the coincubation of Pen and Pal with UDP-GlcNAc *in vitro* leads to the formation of UDP-4-keto-6-deoxy-AltNAc and UDP-4-keto-6-deoxy-GlcNAc (43).

By using gene annotation and RaptorX software, we identified SpsM as the putative enzyme catalyzing the first reaction of the CMP-Leg pathway in B. subtilis, and we demonstrated that the *spsM* gene is required for Leg biosynthesis. We also showed that the insertion of the *pen-pal* and *pgIF* genes into the *amyE* locus of the $\Delta spsM$ mutant strain restored Leg production. Pen-Pal and PgIF were previously shown to convert UDP-GlcNAc to UDP-4-keto-6-deoxy-GlcNAc (43, 49). More surprisingly, the addition of the *legB* gene in the $\Delta spsM$ mutant strain did not fully restore Leg production. LegB is the first enzyme of the CMP-Leg pathway in C. jejuni, and it catalyzes the C-4/C-6 dehydration of the GDP-GlcNAc (30). These data indicate that SpsM is a C-4/C-6 dehydratase, but unlike LegB, it preferentially uses UDP-GlcNAc as a substrate rather than GDP-GlcNAc. These results also imply that the following enzymes of the CMP-Leg pathway in B. subtilis use UDP-linked precursors, indicating that the CMP-Leg pathway of B. subtilis differs from the only complete CMP-Leg pathway described to date in C. jejuni (30). By a similar approach to that carried out with SpsM, we identified SpsC, SpsD, SpsG, SpsE and SpsF as the following enzymes of the pathway, we predicted their functions (see Text S2 and Fig. S6 in the supplemental material), and we reconstructed the CMP-Leg pathway of B. subtilis (Fig. 3A). These in silico data are consistent with our experimental results, which showed that the spsC, spsD, spsG, spsE and spsF genes are required for CMP-Leg biosynthesis. To summarize, this study identified an original CMP-Leg pathway in B. subtilis consisting of six enzymes and using UDP-linked sugars as precursors.

Once activated, the Leg must be transferred to the forespore surface by a specific transferase. There are currently no reports on the bacterial enzymes transferring Leg to cell surfaces. However, crystal structures of four bacterial sialyltransferases have been reported (50-54). These bacterial sialyltransferases have been classified into four carbohydrate-active enzyme database (CAZy) families, including GT38, GT42, GT52, and GT80, based on their amino acid sequence similarities (55, 56). However, in B. subtilis, no glycosyltransferase belongs to any of these four CAZy families. Due to their predicted glycosyltransferase function and the vicinity of the spsA and spsB genes to the genes of the CMP-Leg pathway, the SpsA and SpsB proteins were the most likely transferases of the CMP-Leg in B. subtilis. An analysis performed with the NCBI conserved domain database identified a putative TagB domain in the primary sequence of SpsB. The TagB protein is involved in the priming step of polyglycerol phosphate wall teichoic acid synthesis in B. subtilis (57). Interestingly, two enzymes containing a TagB domain have recently been identified as plausible candidates for a Pse and a Leg derivative transferase in the bacteria Tannerella forsythia (58). In addition, the protein sequence of SpsB contains one DEG and three HP domains. Both D/E-D/E-G and HP motifs are well conserved among different sialyltransferase families with otherwise little or no sequence identity, and these motifs were shown to be important for enzyme catalysis and CMP-Neu5Ac binding (59). Surprisingly, we showed that SpsB is not required for Leg transfer to the forespore surface. Nevertheless, at this stage, it is not possible to exclude that SpsB transfers another still unidentified NulO or that spsB deletion is compensated by another Leg transferase with another acceptor on the forespore surface. The SpsA protein is a transferase of the GT-2 glycosyltransferase

family whose three-dimensional structure has been resolved in complex with Mn-dTDP and Mg-DTP (60, 61). The protein sequence of SpsA contains an asparagine doublet, acting as the DxD motif frequently found in the active site of glycosyltransferases. The asparagine interacts both with the ribose moiety of the nucleotide sugar donor and the divalent cation, thus stabilizing the charged phosphate groups in the nucleotide sugar donor (61). Sialyltransferases usually lack a DxD motif and do not require divalent metal cations for enzymatic activity (62). Accordingly, we showed that SpsA is not required for the transfer of Leg to the forespore surface. These findings suggest that SpsA is not a CMP-Leg transferase. It is more likely that SpsA participates in the transfer of a monosaccharide to a glycan which is being synthesized.

In this study, we showed that the Leg on the spore surface is linked to the crust. Furthermore, we demonstrated that the abolition of Leg production leads to a crust assembly defect. It was shown elsewhere that Pse is essential for flagellar assembly in Campylobacter spp. and H. pylori (32, 34). In H. pylori, single Pse moieties are attached by O-linkages to the structural flagellar proteins FlaA and FlaB, at up to 7 and 10 Ser/Thr residues, respectively. In C. jejuni, the flagellum is also decorated, but with more complex pseudaminic acids, including one with acetamido groups (34, 35, 63). More recently, it was also shown that Leg is present on the flagellins of C. jejuni NCTC 11168 and that flagellin glycosylation is highly heterogeneous, with up to 6 different sugars singly present at a given site (64). Therefore, it is likely that several proteins of the crust of B. subtilis spores are O-glycosylated with Leg on one or several glycosylation sites, and we hypothesize that these O-linked Leg participate in the assembly or the stabilization of the interactions between the crust proteins and/or in the anchoring of the crust to the outer coat. The way by which Leg stabilizes the interactions between the spore surface proteins remains to be defined. However, NulOs are well known to participate in carbohydrate-protein interactions, e.g., mediation of cell-cell adhesion via lectins or cell-cell communication (63). NulOs are hydrophilic and negatively charged at neutral pH and they typically occur as the terminating units of *N*-glycans, *O*-glycans, and glycolipids, which could explain the multiplicity of biological functions they fulfill. In this study, we showed that the abolition of Leg production resulted in a decrease in the negative charge and hydrophilicity of B. subtilis spores, making them more adherent to stainless steel. However, similar surface modifications were obtained with spores of the $\Delta spsA$ and $\Delta spsB$ mutant strains that still produce Leg, suggesting that the hydrophilicity and the negative charge of spores are provided by a crust glycan, rather than by Leg.

The total amounts of neutral sugars in the surface fractions of the $\Delta spsABCDEF$ and $\Delta spsM$ mutant strains were halved compared to that of the wild-type strain. Given that most of the crust is lacking on the spore surfaces of both mutants, this result suggests that approximately one-half of the monosaccharides are located on the outer coat surface and the other half are linked to the crust, which agrees with previous studies (15, 17). The precise localization, structure, and composition of the glycans containing these monosaccharides remain to be defined. However, it has been suggested that one of these glycans contains rhamnose and another one contains galactose (15). It was shown previously that *spsM* deletion leads to a loss of rhamnose on the spore surface, and we confirmed this by gas chromatography-mass spectrometry (GC-MS) experiments (data not shown) (16). Knowing the activity of SpsM, the loss of rhamnose in the $\Delta spsM$ mutant strain is obviously an indirect consequence of Leg production abolition. Since Leg is required for crust assembly, it is therefore likely that rhamnose moieties are constitutive of a crust-linked glycan. Using the data from this study, we propose a speculative model for the organization of the *B. subtilis* spore surface (Fig. 7).

In most of the *B. subtilis* strains, the *spsM* gene is interrupted by the sp β prophage. The prophage is excised from the chromosome both during sporulation and in response to stress (DNA damage), thus reconstituting a functional *spsM* gene (16). Another study indicates that the SigY sigma factor promotes the maintenance of sp β in the chromosome of *B. subtilis* (65). As the other ECF-type sigma factors, SigY activity is regulated by stresses and nitrogen deficiency has been shown to increase SigY



FIG 7 Hypothetical model of the *B. subtilis* spore surface organization. In this model, one or several crust proteins are glycosylated by Leg or by a crust-linked glycan containing Leg. Leg could participate in the assembly and/or the stabilization of the interactions between the crust proteins possibly through a carbohydrate-protein interaction. The crust proteins and the crust-linked glycan give the spores their hydrophilicity and negative charge that define the adhesion properties of *B. subtilis* spores. Leg might also participate in the anchoring of the crust proteins to the outer coat by interacting through a carbohydrate-protein interaction with the outer coat proteins or by a carbohydrate-carbohydrate interaction with a coat-linked glycan. The interactions presented in this model are speculative. They are extrapolated from current knowledge about NulOs in bacteria (see Discussion). In the absence of data about the structure of the coat-linked and crust-linked glycans, the number of monosaccharides that make up the glycans and the position of the bonds between monosaccharides are just indicative of a possible structure.

activity (66). Therefore, it is likely that stresses encountered by *B. subtilis* during sporulation modulate the excision of $sp\beta$ and consequently the production of Leg. Knowing that Leg is required for crust formation and that presence or absence of crust determines the surface and adhesion properties of spores, it is likely that the environmental conditions encountered by *B. subtilis* during sporulation influence the adhesion properties of spores. In the context of food industries, this could mean that production, cleaning and disinfection processes encountered by the sporulating *Bacillus* cells contaminating the production lines might influence the adhesion properties of the resulting spores. This adaptive strategy would explain the high persistence of these microorganisms on food industry production lines.

MATERIALS AND METHODS

Bacterial strains and growth conditions. The strains used in this study are described in Table 1. *Escherichia coli* K-12 strain TG1 was used as host for the construction of plasmids and cloning experiments. *B. subtilis* cells were transformed as described previously (67). *E. coli* strains were grown at 37°C in Luria Broth (LB). *Bacillus* strains were grown at 30°C or 37°C in LB, Nutrient Broth (NB), or Spo8, a sporulation medium (68). The following concentrations of antibiotic were used for bacterial selection: 100 μ g/ml ampicillin and 20 μ g/ml chloramphenicol for *E. coli*, and 1 μ g/ml erythromycin and 5 μ g/ml chloramphenicol for *Bacillus*. Spores were produced at 30°C on Spo8-agar as previously described (68). After a 10-day incubation period and when over 95% of spores were obtained, spores were harvested by scraping the surfaces of plates, washing them five times in chilled sterile water (1,500 × g for 15 min), and storing them in sterile water at 4°C until use. The titer of spore preparations was evaluated by plating on NB-agar and by measuring the optical density at 600 nm (OD₆₀₀).

DNA manipulations. Total DNA was extracted from *B. subtilis* cells using the Puregene Yeast/Bact kit (Qiagen, France). Plasmid DNA was extracted from *E. coli* using a SmartPure plasmid kit (Eurogentec, Belgium). Restriction enzymes and T4 DNA ligase (New England BioLabs, USA) were used in accordance with the manufacturer's recommendations. Oligonucleotide primers were synthesized by Eurofins

TABLE 1 Strains used in this study

Strain	Trait or relevant genotype	Reference or source
B. subtilis PY79		BGSC, 1A747
B. subtilis PY79 ∆spsABCDEF	spsABCDEF-deficient strain	This study
B. subtilis PY79 ∆spsA	spsA-deficient strain; markerless	This study
B. subtilis PY79 ∆spsB	spsB-deficient strain; markerless	This study
B. subtilis PY79 ∆spsD	spsD-deficient strain; markerless	This study
B. subtilis PY79 ∆spsF	spsF-deficient strain; markerless	This study
B. subtilis PY79 ∆spsM	spsM-deficient strain; markerless	This study
B. subtilis PY79 cotE::erm	<i>cotE</i> -inactivated strain; Ery ^r	This study
B. subtilis PY79 cotX::erm	<i>cotX</i> -inactivated strain; Ery ^r	This study
B. subtilis PY79 cotZ::erm	<i>cotZ</i> -inactivated strain; Ery ^r	This study
B. subtilis PY79 ∆spsA amyE::spsA	Complementation of the spsA-deficient strain; Cm ^r	This study
B. subtilis PY79 ∆spsB amyE::spsB	Complementation of the <i>spsB</i> -deficient strain; Cm ^r	This study
B. subtilis PY79 ∆spsD amyE::spsD	Complementation of the <i>spsD</i> -deficient strain; Cm ^r	This study
B. subtilis PY79 ∆spsF amyE::spsF	Complementation of the <i>spsF</i> -deficient strain; Cm ^r	This study
B. subtilis PY79 ∆spsM amyE::spsM	Complementation of the <i>spsM</i> -deficient strain; Cm ^r	This study
B. subtilis PY79 ∆spsM amyE::spsM M146A	M146A mutated allele of <i>spsM</i> in the amyE locus of the <i>spsM</i> -deficient strain; Cm ^r	This study
B. subtilis PY79 ∆spsM amyE::spsM K150A	K150A mutated allele of <i>spsM</i> in the <i>amyE</i> locus of the <i>spsM</i> -deficient strain; Cm ^r	This study
B. subtilis PY79 ΔspsM amyE::pen	pen gene in the amyE locus of the spsM-deficient strain; Cmr	This study
B. subtilis PY79 ΔspsM amyE::pal	pal gene in the amyE locus of the spsM-deficient strain; Cm ^r	This study
B. subtilis PY79 ΔspsM amyE::pen-pal	pen and pal genes in the amyE locus of the spsM-deficient strain; Cm ^r	This study
B. subtilis PY79 ∆spsM amyE::legB	legB gene in the amyE locus of the spsM-deficient strain; Cmr	This study
B. subtilis PY79 ∆spsM amyE::pglF	pgIF gene in the amyE locus of the spsM-deficient strain; Cmr	This study
B. subtilis PY79 ∆spsM amyE::pseB	pseB gene in the amyE locus of the spsM-deficient strain; Cm ^r	This study
B. subtilis PY79 amyE::P _{spsA} -mCherry	P _{spsA} - <i>mCherry</i> transcriptional fusion in the <i>amyE</i> locus of the PY79 strain: Cm ^r	This study
B. subtilis 168		BGSC, 1A1
B. subtilis 168 ∆spsC	spsC-deficient strain; Km ^r	BGSC, BKK37890
B. subtilis 168 ΔspsE	spsE-deficient strain; Km ^r	BGSC, BKK37870
B. subtilis 168 ΔspsG	spsG-deficient strain; Km ^r	BGSC, BKK37850
B. thuringiensis serovar israelensis ATCC 35646	· · ·	BGSC, 4Q12

Genomics (Germany). PCRs were performed in a SimpliAmp thermal cycler (Applied Biosystems, USA). Amplified fragments were purified using the SmartPure PCR kit (Eurogentec, Belgium). Digested DNA fragments were separated on 1% (w/v) agarose gels after digestion and extracted from gels using the SmartPure gel kit (Eurogentec, Belgium). Nucleotide sequences of plasmid inserts were determined by Sanger sequencing (Eurofins Genomics, Germany).

Construction of the *B. subtilis* **recombinant strains.** Gene deletion and gene interruption were performed by homologous recombination with pMAD and pMUTIN4, respectively, as previously described (69, 70). The deletion mutants were constructed by deleting the targeted genes from the ATG to the stop codon without introducing exogenous DNA to avoid disrupting downstream genes transcription. Complementation was introduced in the *amyE* gene using the plasmid pBS1C (67). The recombinant strains, oligonucleotides, and plasmids used in this study are described in Table 1 and Tables S2A and B in the supplemental material. Nucleotide sequences of the introduced chromosomal modifications were verified by Sanger sequencing (Eurofins Genomics, Germany).

Surface and adhesion properties of spores. Microbial-adhesion-to-hydrocarbons (MATH), zetametry, and adhesion assays were performed as described previously (9).

Electron microscopy. To observe spore surface features, spores were fixed and stained as previously described (9). Ultrathin sections were cut on a Leica UC7 ultramicrotome and collected on 150-mesh copper grids. The sections were examined under a Jeol JEM-2100 transmission electron microscope at an accelerated voltage of 200 kV.

Preparation of the intracellular extracts and spore surface fractions. A cell pellet obtained by centrifugation (12,000 × *g*, 15 min, 4°C) and washed twice with cold sterile ultrapure water was resuspended in 3 ml of sodium citrate buffer (0.01 M, pH 5.5) and then transferred to a screw tube containing 400 mg of glass beads (acid-washed, G1277-500G; Sigma). The screw tube was shaken in a MiniBeadBeater-16 (Biospec Products) for 2 min and then cooled on ice. This treatment was repeated 4 times. The lysate was then centrifuged (6,000 × *g*, 15 min, 4°C), and 1 ml of the supernatant was ethanol precipitated. The precipitate was recovered in 500 μ l of sterile ultrapure water and kept at -20° C for further analyses. The surface fraction of spores was removed by three successive passages of spore suspensions through a French press (SLM Instruments, Urbana, IL) at 20,000 lb/in². Spores were separated from the surface fraction by two successive centrifugations (4,500 × *g*, 15 min, 4°C), and the supernatant was kept at -20° C until use.

Dosage of neutral sugars. The dosage of the neutral sugars in the surface fraction of spores was performed as described previously (28). The amounts of neutral sugars were standardized by the OD_{600} of the spore preparations.

Release and derivation of NulOs. 500 μ l of spore surface fraction or intracellular extract was lyophilized. DMB-coupled NulOs were obtained as described previously (36). Shortly, dried glycoconjugates were hydrolyzed at 80°C for 2 h in 0.1 M trifluoroacetic acid to release the NulOs. NulOs were subsequently coupled to DMB by heating the samples at 50°C for 2 h in the dark in 7 mM DMB, 1 M β -mercaptoethanol, 18 mM sodium hydrosulfite in 0.02 mM trifluoroacetic acid.

Analysis of DMB derivatives on HPLC. The NulO derivatives were separated isocratically on a C₁₈ reverse-phase HPLC column (Waters, 4.6 mm by 150 mm, 3.5 μ m. or Wakopak Handy ODS, 4.6 mm by 250 mm, 6 μ m) using a solvent mixture of acetonitrile/methanol/water (7:9:84 [vol/vol/vol]), and they were detected by a fluorimeter (Waters 474, excitation wavelength [λ_{exc}] = 373 nm, emission wavelength [λ_{em}] = 448 nm). The Leg amount was estimated by measuring the area of peak 1 and by reporting this area to a standard range of Neu5Ac.

Analysis of DMB derivatives on micro-LC/ESI-MS³. Analyses were performed in positive ion mode on an amaZon speed ETD ion trap mass spectrometer equipped with the standard electrospray ionization (ESI) ion source and controlled by Hystar 3.2 software (Bruker Daltonics). DMB-coupled NulO separation was achieved on a micro-LC system (Prominence LC-20AB; Shimadzu, Kyoto, Japan). Samples were applied to a reverse-phase Luna C₁₈-2 column (150 mm by 1.00 mm, 3- μ m particles; Phenomenex), and they were separated using an isocratic elution of acetonitrile/methanol/water (6:4:90 [vol/vol/vol)] at a flow rate of 70 μ l/min. The targeted MS³ scans for DMB-coupled NulO were performed using ultrascan mode (26,000 atomic mass units [amu]/s). NulO species were identified by referring to elution positions and MS³ fragmentation of Leg and Pse standards.

SUPPLEMENTAL MATERIAL

Supplemental material is available online only. **TEXT S1**, DOCX file, 0.1 MB. **TEXT S2**, DOCX file, 0.1 MB. **FIG S1**, TIF file, 1.4 MB. **FIG S2**, TIF file, 0.6 MB. **FIG S3**, TIF file, 0.8 MB. **FIG S4**, TIF file, 1.2 MB. **FIG S6**, PDF file, 1.2 MB. **FIG S6**, PDF file, 1.5 MB. **TABLE S1**, DOCX file, 0.1 MB. **TABLE S2**, DOCX file, 0.1 MB.

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