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Novel mixed-linkage β-glucan activated by c-di-GMP in *Sinorhizobium meliloti*

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An artificial increase of cyclic diguanylate (c-di-GMP) levels in Sinorhizobium meliloti 8530, a bacterium that does not carry known cellulose synthesis genes, leads to overproduction of a substance that binds the dyes Congo red and calcofluor. Sugar composition and methylation analyses and NMR studies identified this compound as a linear mixed-linkage $(1 \rightarrow 3)(1 \rightarrow 4)$ - β -D-glucan (ML β -glucan), not previously described in bacteria but resembling ML β-glucans found in plants and lichens. This unique polymer is hydrolyzed by the specific endoglucanase lichenase, but, unlike lichenan and barley glucan, it generates a disaccharidic \rightarrow 4)- β -D-Glcp-(1 \rightarrow 3)- β -D-Glcp-(1 \rightarrow repeating unit. A two-gene operon bgsBA required for production of this ML β -glucan is conserved among several genera within the order Rhizobiales, where bgsA encodes a glycosyl transferase with domain resemblance and phylogenetic relationship to curdlan synthases and to bacterial cellulose synthases. ML β-glucan synthesis is subjected to both transcriptional and posttranslational regulation. bgsBA transcription is dependent on the exopolysaccharide/guorum sensing ExpR/SinI regulatory system, and posttranslational regulation seems to involve allosteric activation of the ML β-glucan synthase BgsA by c-di-GMP binding to its C-terminal domain. To our knowledge, this is the first report on a linear mixed-linkage $(1\rightarrow 3)(1\rightarrow 4)$ - β -glucan produced by a bacterium. The S. meliloti ML β-glucan participates in bacterial aggregation and biofilm formation and is required for efficient attachment to the roots of a host plant, resembling the biological role of cellulose in other bacteria.

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n bacteria the second messenger cyclic diguanylate (c-di-GMP) regulates the transition from a motile to a sedentary lifestyle, so that high concentrations enhance the production of extracellular matrix components, exopolysaccharides (EPSs), and proteins (1–5). The c-di-GMP is synthesized from two molecules of GTP by the action of diguanylate cyclases (DGCs) and is hydrolyzed to 5'-phosphoguanylyl-(3'-5')-guanosine by specific phosphodiesterases. It has been determined that artificial increments of the intracellular levels of c-di-GMP by the over-expression of DGCs often trigger phenotypes related to the synthesis of EPSs and biofilm formation (5, 6).

Rhizobia produce a complex array of glucidic molecules including LPSs, capsular polysaccharides (CPSs), cyclic glucans, high-molecular-weight neutral polysaccharide (glucomannan), gel-forming polysaccharide, and EPSs (7–9). In addition, these symbiotic bacteria secrete other compounds, such as lipo-chitin oligosaccharides, which specifically mediate the interaction with their legume hosts (10).

In contrast to structural polysaccharides (i.e., CPSs or LPSs), EPSs are weakly associated with the cell outer surface and often are released into the environment (8). EPSs are either branched or linear homo- or heteropolymers of a wide variety of D-monosaccharides and form a highly hydrated layer on the cell surface, providing protection against environmental biotic and abiotic stresses. EPSs also play a major role in cell aggregation, flocculation, and biofilm formation, critical processes in the initial stages of bacterial interaction with eukaryotic hosts (11, 12). Finally, many rhizobia also use specific EPSs as signaling molecules that are indispensable for the invasion of host plants (13, 14).

Sinorhizobium meliloti, an endosymbiont of alfalfa (Medicago sativa), produces two structurally distinct EPSs: EPS-I, or succinoglycan, and EPS-II, or galactoglucan; both are regulated by the ExpR/Sin quorum-sensing system and are important for nodulation of legume hosts (13, 15, 16). However, in contrast to many members of the Rhizobiaceae family, the model strain S. meliloti 8530 (Sme) lacks the gene clusters required for the biosynthesis of other common D-glucose homopolysaccharides such as cellulose or curdlan. Two types of cellulose synthase (CS) systems have been described in bacteria. Group I is found in the α , β , and γ subdivisions and is encoded by a three-gene operon *bcsABC*/ acsABC, plus a glycosylhydrolase gene which can be linked to the main operon or located elsewhere in the genome (17, 18). Group II typically is found in α -Proteobacteria, such as Agrobacterium tumefaciens, and is encoded by two adjacent operons: celABC, homologs to group I bcsA, bcsB and the glycosylhydrolase, respectively, and celDE(FG) with no significant sequence similarity to group I genes (19). A. tumefaciens and other Rhizobiales also carry genes for the biosynthesis of curdlan (crdASC, crdR, and *pssAG*), another high-molecular-weight linear β -glucan composed of D-glucose residues linked by $\beta(1\rightarrow 3)$ bonds instead of the $\beta(1\rightarrow 4)$ bond found in cellulose (20). Production of both curdlan and cellulose seem to be activated by the second messenger c-di-GMP, although only the allosteric activation of CSs through c-di-GMP binding to a PilZ domain, located in the C-terminal region

Significance

We report a novel linear mixed-linkage $(1\rightarrow 3)(1\rightarrow 4)$ - β -glucan produced by the bacterium *Sinorhizobium meliloti* upon raising cyclic diguanylate (c-di-GMP) intracellular levels. This unique bacterial polysaccharide resembles $(1\rightarrow 3)(1\rightarrow 4)$ - β -glucans found in cereals and certain lichens but has a distinctive primary structure. Genes, proteins, and regulatory pathways for producing this new polymer are described. Our findings open the possibility of using bacteria to produce $(1\rightarrow 3)(1\rightarrow 4)$ - β -glucans, which are receiving increasing interest as bioactive compounds, and provide new elements to disclose molecular mechanisms of c-di-GMP regulation as well as for investigating the evolution, activity, and specificity of glycosyl transferases.

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The authors declare no conflict of interest.

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of BcsA/CelA proteins, has been demonstrated (21–23). A PilZ domain apparently is absent in curdlan synthases (24); nonetheless c-di-GMP activation could occur by a different mechanism, because this second messenger is able to regulate the production of other EPSs by interacting with proteins involved in EPS extrusion (25, 26). Here we have exploited c-di-GMP activation of bacterial EPSs to unveil an undisclosed linear mixed-linkage (ML) $(1\rightarrow3)(1\rightarrow4)$ - β -glucan in a cellulose-nonproducing bacterium, *S. meliloti* 8530.

Results

Effects of Increased c-di-GMP Levels on the Production of EPSs by *S. meliloti* 8530. In cellulose-producing (Cel⁺) bacteria, raising c-di-GMP intracellular levels (i.e., by overexpression of a DGC) triggers production of cellulose and other extracellular molecules (5, 6, 27). *S. meliloti* 8530 does not bear any of the known CS operons in its genome; however, overexpression of the DGC PleD* generated wrinkled and red-stained colonies in medium with Congo red (CR), in deep contrast to the wild-type colonies (*SI Appendix*, Fig. S1*A*). Furthermore, the overexpression of *pleD** (pJBpleD*) also increased the calcofluor (CF)-derived fluorescence in both solid and liquid cultures (Fig. 1 and *SI Appendix*, Fig. S1*B*).

In liquid media and particularly in minimal medium (MM), Sme overexpressing pleD* (Sme pJBpleD*) displayed a strong aggregative behavior with most bacteria forming flocs (SI Ap*pendix*, Fig. S24). Scanning electron microscopy of flocs showed that bacteria were encased in a dense matrix of filaments (SI Appendix, Fig. S2B). Adherence to and biofilm formation on test tube walls also were enhanced (SI Appendix, Fig. S3A); moreover, those biofilms fluoresced in the presence of CF under UV light (SI Appendix, Fig. S3B), indicating that the CF⁺ substance was part of the biofilm matrix. However, biofilm formation could not be quantified because it detached easily from the abiotic surfaces and was removed during the Cristal Violet staining process (Materials and Methods). The PleD*-dependent EPS matrix produced by Sme also enhanced its adhesion to alfalfa roots and the formation of a thick CF⁺ biofilm coating the root surface (SI Appendix, Fig. S3C). All these phenotypes correlated with a strong increase in the c-di-GMP intracellular levels provoked by PleD*. Cell extracts from Sme harboring the plasmid pJBpleD* contained 596.86 pmol of c-di-GMP/mg total protein, whereas the amounts of c-di-GMP from Sme carrying the pJB3Tc19 empty vector were below the HPLC-MS limit of detection (SI Appendix, Supplementary Methods).

Characterization of a Novel β -Glucan Produced by S. meliloti. CR binds basic or neutral polysaccharides as well as some proteins,



Fig. 1. Quantification of CF-derived fluorescence of *S. meliloti* 8530 and its derivative *Smb20391::Nm* mutant harboring different plasmid constructions. Results are expressed in arbitrary units standardized by the $OD_{600} \pm SE$ from three independent cultures.



Fig. 2. CF-derived fluorescence of *S. meliloti* 8530 (Wt) and mutant derivatives carrying the plasmid pJBpleD* or the empty vector pJB3Tc19. (*Right*) Bacterial cultures were imaged under UV light after growth at 28 °C in solid TY medium supplemented with CF (200 μ g/mL). (*Left*) The diagram depicts the positions of the strains.

whereas CF is more specific for polymers containing β (1-3)- or β (1-4)-D-glucopyranosyl units (28, 29). *Sme* produces at least one EPS, known as "EPS-I" or "succinoglycan," which binds CF but not CR (30, 31). Indeed, *Sme* shows a greenish CF-derived fluorescence; however pJBpleD* conferred a blue CF fluorescence (Fig. 2) which was retained in an *exoY* mutant lacking EPS-I (32). This finding suggested that high c-di-GMP levels generated by PleD* stimulated the production of a CF-binding polymer different from EPS-I.

A purification approach was designed to isolate and identify this substance. The Sme pJBpleD* strain was grown in flasks with liquid MM, and the flocs generated were recovered after 3 d using a sieve (Materials and Methods). Like curdlan and cellulose, the Sme substance was water insoluble, but, unlike cellulose, it could be dissolved in diluted NaOH solutions (0.5 M). Curdlan is soluble in dilute alkali (0.25 M NaOH) and, after aqueous suspensions are heated above 80 °C and subsequently cooled, produces high-set thermo-irreversible gels (33). This gelling behavior was not observed with the Sme substance, so several rounds of boiling/cooling/centrifugation cycles were followed by a lyophilization step to obtain a substance with a cottony appearance. Monosaccharide analysis showed that D-glucose was the only component, and the methylation analysis indicated that these glucopyranoses were substituted at positions $\rightarrow 3$ and $\rightarrow 4$) in a 1:1 ratio. The polysaccharide was insoluble in water or in DMSO; therefore a few milligrams were treated with a mixture of acetic anhydride and pyridine to acetylate its hydroxyl groups and improve its solubility in organic solvents. After purification by size-exclusion chromatography (SEC), a fraction of the polysaccharide was solubilized in CDCl₃ and was studied by NMR. Assignments were made using bidimensional experiments [DQF-COSY, total correlated spectroscopy (TOCSY), rotating-frame Overhauser spectroscopy (ROESY), and heteronuclear single-quantum coherence (HSQC)] and are listed in Table 1. NMR spectra showed two different units of glucose, labeled as "unit A" and "unit B," with a ratio close to 1:1. Chemical shifts (¹H and ¹³C) indicated their β configuration and that they were present in the pyranose form (Fig. 3A). Moreover, unit A was substituted at position $\rightarrow 4$), whereas unit B was substituted at position \rightarrow 3). The results were in good agreement with the chemical shifts of acetylated cellulose (34) and $(1\rightarrow 3)$ - β -D-glucopentaose peracetate (35). Finally, the ROESY spectrum (SI Appendix, Fig. S4) allowed the sequence of residues in the polysaccharide to be deduced, because intense cross-peaks were found between signals H-1 in unit A and H-3 in unit B and between H-1 in unit B and H-4 in unit A. Considering all these data, we propose that the acetylated polysaccharide has the repeating unit $\rightarrow 3$)- β -D-Glcp-(1 $\rightarrow 4$)- β -D-Glcp-(1 \rightarrow (Fig. 3B).

This β -glucan structure resembles ML β -glucans such as lichenan and barley β -glucan, which can be hydrolyzed specifically by lichenase, an endohydrolase whose target site is widely assumed to be all $(1\rightarrow 4)$ bonds immediately following $(1\rightarrow 3)$

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Table 1. Chemical shifts and assignments of acetylated ML β-glucan purified from 5. meliloti 8530 overexpressing pleD* (Sme pJBpleD*)

Unit	Nucleus	Position						
		1	2	3	4	5	6a	6b
A: →4)-β-D-Glc <i>p</i> -(1→	¹ H	4.49	4.78	5.05	3.69	3.56	4.3	4.1
	¹³ C	100.9	71.0	72.7	75.7	72.4	62.2	
B: →3)-β-D-Glc <i>p</i> -(1→	¹ H	4.26	4.9	3.76	4.88	3.56	4.3	4.1
	¹³ C	100.7	72.8	78.8	67.8	72.4	62.2	

bonds (36). It does not hydrolyze pure $(1\rightarrow 3)$ - β -D-glucans (i.e., curdlan) nor $(1\rightarrow 4)$ - β -D-glucans (cellulose). Incubation of lichenan and barley β -glucan with lichenase produced distinctive mixtures of tri-, tetra-, and larger cello-oligosaccharides which could be separated by TLC (Fig. 4), in agreement with previous reports (37). Lichenase also was able to hydrolyze the S. meliloti glucan, although in this case the only product of hydrolysis was a disaccharide (Fig. 4). This result agreed with the NMR data and further supported the notion that the repeating unit of the Sme β-glucan contains alternating $\beta(1\rightarrow 3)$ and $\beta(1\rightarrow 4)$ bonds (Fig. 3B).

The bgsBA Operon Is Required for ML β-Glucan Production. The S. meliloti 8530 genome encodes at least two glycosyl transferases



Fig. 3. Structure of the ML β -glucan. (A) ¹H-NMR (500 MHz) and ¹H-¹³C (125 MHz) HSQC spectra of the acetylated ML β-glucan purified from S. meliloti 8530 overexpressing pleD* (Sme pJBpleD*). (B) Proposed structure of the ML β-glucan. The repeating unit is indicated by brackets.

(GTs) of unknown function, SMb20391 and SMb202460, annotated as hypothetical CSs (38). The PleD*-dependent phenotypes were tested with transposon-induced mutants in both genes (SI Appendix, Table S1). The SMb20460 mutant behaved like the wild type; however, the SMb20391 mutant had lost the PleD*dependent CR⁺ and CF⁺ phenotypes (SI Appendix, Fig. S5) and the aggregative behavior observed in the wild type.

The SMb20391 ORF overlaps by 19 bp the upstream ORF SMb20390, which encodes a putative membrane fusion protein (MFP) belonging to the AcrA/EmrA/HlyD family of transporters. RT-PCR experiments indicated that SMb20390 and SMb20391 are cotranscribed (SI Appendix, Fig. S6A).

An in-frame deletion of SMb20390 (Δ SMb20390) was constructed, and, like the SMb20391 mutant, the Δ SMb20390 pJBpleD* mutant showed a CR⁻ and CF⁻ phenotype and did not aggregate or form biofilm at high c-di-GMP levels. This result indicated that both the SMb20390 and SMb20391 genes are responsible for the c-di-GMP-induced phenotypes observed in Sme. This notion was confirmed further after genetic complementation. To avoid possible functional problems resulting from an imbalance between SMb20390 and SMb20391 protein levels, two new plasmids were constructed, one harboring the complete operon SMb20390-91 cloned in pJB3Tc19 (pJB9091) and a second one carrying SMb20390-91 and pleD* (pJBpleD*9091) (SI Appendix, Fig. S7). Both plasmid constructions were able to restore the CR⁺ and CF⁺ phenotypes in the SMb20391::Nm mutant (Fig. 1). Although the CF-derived fluorescence was much higher when *pleD** was present, the two genes in multicopy were able to provide CR⁺ and CF⁺ phenotypes even under physiological c-di-GMP conditions. We also measured the relative yield of ML β -glucan in cultures, which represented 47.8% of the dry weight of the wild-type Sme (pJBpleD*) cultures. As expected, no ML β-glucan was produced by the SMb20391::Nm mutant overexpressing PleD*. The SMb20391::Nm mutant complemented with pJB9091 displayed a low (2.1% of culture dry weight) but nevertheless measurable yield of ML β -glucan. The ML β -glucan yield was highest in cultures of the mutant carrying both *pleD** and SMb20390-20391 genes in multicopy (pJBpleD*9091), accounting for 77.5% of the culture dry weight. Therefore, two conditions seem to be required for high production of the ML β -glucan by Sme: (i) the presence of a functional copy of the SMb20390-91 operon and (ii) high intracellular levels of c-di-GMP. Given their requirement for ML β -glucan production, we propose that *SMb20391* be renamed *bgsA* (with "*bgs*" standing for " β -glucan synthesis") and that SMb20390 be renamed "bgsB."

c-di-GMP Binding to BgsA. The bgsB gene product contains an HlyD-like domain found in diverse proteins involved in the export of a broad variety of compounds (39) and shows homology to MFPs, accessory proteins that cooperate with a variety of transporters. MFPs span the periplasm, linking the inner and outer membranes and facilitating the formation of a channel to expel the substrate from the cell (40).

Hydropathy analysis of the 664-aa BgsA protein sequence using the TMHMM 2.0 prediction program (41) revealed seven potential transmembrane (TM) helices (Fig. 5), in contrast to the



Fig. 4. TLC of β -glucans hydrolyzed with lichenase. Lane 1: untreated *S. meliloti* ML β -glucan; lane 2: *S. meliloti* ML β -glucan incubated with lichenase; lane 3: untreated lichenan; lane 4: lichenan + lichenase; lane 5: untreated barley β -glucan; lane 6: barley β -glucan + lichenase; lane 7: mixture of glucose, maltose, maltotetraose, maltopentaose, and maltohexaose.

eight TMs present in CSs (42, 43). The large hydrophilic region between TM III and IV (amino acid residues 90-375) contains residues and motifs conserved in the GT-2 glycosyl transferases family (D,D,D₃₅QxxRW), including the DXD signature involved in the addition of the UDP-sugar to the nascent polysaccharide (44) and the QXXRW sequence motif likely required for holding the growing glycan chain in the active site (Fig. 5) (45). After TM VII a C-terminal cytoplasmic segment was predicted (residues 532-664) which contains no known functional domains. This protein organization strongly resembles the structure predicted for the A. tumefaciens curdlan synthase (CrdS) (24) and CS; the latter contains a PilZ signature in its C-terminal cytoplasmic domain involved in c-di-GMP binding (22). However, neither CrdS nor BgsA seems to bear a PilZ domain nor the recently described c-di-GMP-binding GIL domain that is present in BcsE-like proteins (46). We nevertheless tested the ability of BgsA to bind c-di-GMP, using a previously reported fluorescence-based dot blot assay with 2'-Fluo-AHC-c-di-GMP (47). Crude extracts of Escherichia coli cells expressing full-length BgsA gave weak positive results, although BgsA expression probably was poor, because these cells had strongly delayed growth, and the protein could not be detected in SDS/PAGE gels. However, extracts of E. coli cells expressing the cytoplasmic C-terminal segment of BgsA (BgsA-C; the last 139 residues beginning with Ala526) (Fig. 5) showed significant binding to 2'-Fluo-AHC-c-di-GMP compared with cell extracts expressing the empty vector (Fig. 6A). The signal from the BgsA-C extract was lower than with the PilZ domain protein PA3353 (46), but this difference could be caused by different protein-expression levels and/or affinities for the ligand. Moreover, the binding of the BgsA-C extract to 2'-Fluo-AHC-cdi-GMP could be outcompeted by growing concentrations of pure, unlabeled c-di-GMP (Fig. 6B). These results provide strong evidence that BgsA can bind c-di-GMP through its C-terminal cytoplasmic domain, suggesting allosteric activation of this GT.

ExpR Activates Transcription of the *bgsBA* **Operon**. The impact of c-di-GMP on the transcription of the *bgsBA* operon was assayed by quantitative RT-PCR (RT-qPCR) in the absence and presence of PleD*. Although *bgsA* transcription was slightly higher with PleD*, this variation was not statistically significant and suggested that c-di-GMP does not affect the transcription of *bgsBA* (*SI Appendix*, Fig. S6B). This suggestion was verified in a strain carrying a transcriptional *bgsA*:*gus* fusion. β -Glucuronidase activity in strain *SMb20391::Nm-gus* carrying pJBpleD* was 158.4 ± 13.85, similar to 171.7 ± 22.04 in the strain with the empty vector pJB3Tc19.

bgsA (*SMb20391*) was found previously among *Sme* genes whose expression was dependent on *expR* and *sinI* genes (48) and recently has been confirmed to be a member of the Sin/ExpR regulon (49). We confirmed the ExpR dependence of *bgsA* transcription (*SI Appendix*, Fig. S6B) and extended it to *bgsB*, which showed a 50-fold transcript reduction in the *expR* mutant. Moreover, *expR* and *sinI* mutants overexpressing PleD* were impaired in the emission of the c-di-GMP–induced blue CF fluorescence (Fig. 2), showed a CR⁻ phenotype, and did not exhibit the PleD*-dependent flocculation in liquid media. Furthermore, the ExpR dependence of ML β-glucan production was confirmed in the *expR* mutant Rm1021 (38), which did not display any of the c-di-GMP–associated phenotypes described for *Sme*.

ML β-Glucan Is Involved in the Attachment of S. meliloti to Alfalfa Roots. We evaluated the impact of the new ML β-glucan on bacterial colonization and attachment to alfalfa roots. For microscopy studies, variants of wild-type Sme and bgsA (SMb20391::Nm) mutant strains expressing two different fluorescent proteins, GFP and DsRed, were used. As a control for plasmid stability and to rule out any possible interference of the fluorescent proteins with bacterial physiology or ML β-glucan production, a flocculation assay was carried out first with mixtures of the Sme (pJBpleD*) strain tagged with either of the fluorescent proteins, and flocs formed after 3 d were observed under the epifluorescence microscope. These flocs were constituted by similar numbers of each tagged variant (SI Appendix, Fig. S84). In contrast, when bacterial mixtures contained the wild-type (pJBpleD*, pBBR-gfp) and the bgsA mutant (pJBpleD*, pBBR-dsRed) cells, the wild-type cells were found almost exclusively in the flocs. Most mutant cells were in the surrounding medium, and few were located within the flocs (SI Appendix, Fig. S8B), suggesting that cells not producing the ML β -glucan were excluded from the cell network.

Confocal laser scanning microscopy (CLSM) was used to study competitive root attachment and colonization under high c-di-GMP conditions. Twenty-four hours after inoculation Bgs⁺ cells firmly attached to roots exceeded the *bgsA* mutant cells by more than 200fold and formed a strong biofilm with large cell aggregates covering most of the root surface, whereas Bgs⁻ cells were dispersed (Fig. 7).



Fig. 5. Topology model of BgsA (SMb20391) in the inner cell membrane (IM) predicted by TMHMM 2.0 (41) and presented by TOPO2 (www.sacs.ucsf. edu/cgi-bin/open-topo2.py). The putative catalytic residues of the D,D, $D_{35}QxxRW$ motif are shown in yellow, and the predicted DXD motif is enclosed in a box. The 139-aa C-BgsA peptide used in c-di-GMP-binding experiments is shown as gray diamonds.

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Fig. 6. Dot blot c-di-GMP-binding assays. (A) Dot fluorescence images of extracts of *E. coli* cells expressing the C-terminal fragment of BgsA (pQE80L:: C-BgsA) (lane 3), the PilZ domain protein PA3353 (pET21::PA3353) (lane 2), or the empty vector (pQE80L) (lane 1) immobilized onto nitrocellulose membranes and incubated with 2'-Fluo-AHC-c-di-GMP. (*B*) Dot fluorescence of C-BgsA lysates incubated with a mix of 0.1 μ M 2'-Fluo-AHC-c-di-GMP and increasing amounts of unlabeled c-di-GMP.

Bacterial attachment to roots also was studied at physiological levels of c-di-GMP. Plant roots were inoculated with 1:1 mixtures of the wild-type 8530 and the *bgsA SMb20391::Nm* strain at two different inoculum densities, 10^5 or 10^3 total cells applied per root. The BgsA⁻ mutant cells firmly attached to roots were clearly outnumbered by the wild-type cells (Fig. 8), suggesting that Bgs⁺ bacteria attach to roots more effectively than those lacking the ML β -glucan. Taken together, the results support the notion that production of the ML β -glucan is required for efficient attachment of *S. meliloti* to the alfalfa root surface and consequently for root colonization.

Influence of ML β-Glucan on Nodulation. Alfalfa nodulation with bacteria at high c-di-GMP levels could not be tested properly because the PleD* plasmid was lost rapidly under nonselective conditions. At physiological c-di-GMP levels the BgsA⁻ mutant formed pink wild-type–like nodules indicative of nitrogen fixation. Nodulation kinetics by individual strains were determined at two different inoculum cell densities, 10^5 or 10^3 total cells applied per root; no significant differences between the wild-type and the *bgsA SMb20391::Nm* strain could be observed (*SI Appendix*, Fig. S9). Competitive nodulation was tested by inoculating roots with 1:1 mixtures of the wild type and the *bgsA* mutant, also applied at cell densities of 10^5 or 10^3 total cells per root. In both instances each strain occupied nearly 50% of the nodules, suggesting that under our experimental conditions the *bgsA* mutant is as competitive for nodulation as the wild type.

The bgsBA Operon Is Widespread Among Rhizobiales. The bgsBA operon is required for the synthesis and/or secretion of the novel ML β -glucan uncovered in Sme. Given the features predicted for the encoded proteins, it is most likely that BgsA is the synthetase and that BgsB may participate in the export of the glucan chain. Similar to what occurs in cellulose or curdlan production (18, 20), it is likely that additional yet unknown components are necessary for full activity of the ML β-glucan synthesis system. We could identify bgsBA-like operons in genomes of the genera Sinorhizobium, Rhizobium, and Agrobacterium in the Rhizobiaceae and Methylobacterium in the Methylobacteriaceae, all within order Rhizobiales (Fig. 9). For instance, all available sinorhizobial (S. meliloti and S. fredii) and most Methylobacterium genomes bear a bgsBA-like operon, although only some strains of Rhizobium carry these genes. Among agrobacteria, only A. radiobacter K84 (a nonpathogenic biocontrol strain) carries a bgsBA-like operon. A phylogenetic analysis of BgsA (β-glucan), CrdS (curdlan), and CelA/ BcsA/AcsA (cellulose) synthases showed two monophyletic clades separating Cel synthases from Crd and Bgs synthases (Fig. 9), with the two latter forming clearly diverging clusters. Unlike S. meliloti 8530, which carries no CS genes, a Cel system coexists with either a Bgs or a Crd system in most available Rhizobiales genomes. Coexistence of Crd and Bgs systems is far less frequent, although distinctive Cel, Crd, and Bgs synthases are borne by some strains of methylobacteria (Fig. 9). Taken together, the data suggest that CrdS and BgsA share a common ancestor and that divergent evolution may have shaped their specific activities toward synthesis of either (1,3)- β -glucan or (1,3) (1,4)- β -glucan, respectively.

Discussion

c-di-GMP is a common activator of several bacterial EPSs, such as cellulose in many bacteria, poly-β-1,6-N-acetilglucosamine in enterobacteria, or alginate, Psl, and Pel in Pseudomonads (50, 51). In this work, we have exploited this feature to uncover an otherwise cryptic EPS produced by the α -proteobacterium S. meliloti 8530 when its intracellular c-di-GMP levels are artificially increased. This new bacterial polymer is a linear ML $(1\rightarrow 3)(1\rightarrow 4)$ - β -D-glucan. It resembles the ML β -glucans found in cereals and lichens, but the bacterial ML β-glucan presents a unique primary structure. Whereas the plant ML β -glucans contain distinctive ratios of cello-oligosaccharides of 3, 4, or higher degrees of polymerization, linked by $(1\rightarrow 3)$ bonds (37), the bacterial ML β -glucan contains a \rightarrow 3)- β -D-Glcp-(1 \rightarrow 4)- β -D-Glcp-(1 \rightarrow repeating unit. ML β -glucans are attracting increasing industrial interest as bioactive ingredients and food additives (52). The structural features of ML β -glucans, including the ratio of β -(1 \rightarrow 4)/(1 \rightarrow 3) linkages, the abundance of long cellulosic fragments, and molecular size, determine their physical and physiological properties (53). The S. meliloti ML β -glucan shares some properties with other bacterial β -glucans such as its insolubility in water (i.e., cellulose and curdlan) and solubility in dilute alkali (curdlan); however, the precise description of its



Fig. 7. Competitive root attachment and colonization with fluorescence-tagged strains. CLSM images of *M. sativa* roots inoculated with 1:1 mixtures of *S. meliloti* 8530-GFP and mutant *SMb20391*-dsRed, both overexpressing PleD*. (Magnification, 4×.) The wild type (green) forms dense biofilms coating the root, whereas *SMb20391* cells (red) are found in low numbers and are dispersed.



Fig. 8. Attachment of *S. meliloti* to alfalfa roots. Alfalfa roots were inoculated with 1:1 mixtures of wild-type strain 8530 and the *bgsA* mutant *SMb20391::Nm* (inoculum*) at two cell densities, 10^5 (*Left Bars*) or 10^3 (*Right Bars*). Cells firmly attached to roots were counted 1 and 3 d after inoculation. Bars show the proportion (%) of wild-type (gray and hatched portions of bars) and mutant (white portions of bars) cells attached to roots. Error bars indicate SDs.

physical features and rheological properties shall be the subject of future specific studies.

At least two genes, bgsA and bgsB, likely encoding a GT-2 and a membrane fusion protein, respectively, are required for the production of this extracellular ML β -glucan. BgsA is a strong candidate to be the ML β -glucan synthase, because it shares sequence, domain conservation, and predicted secondary structure with other bacterial GT-2 proteins, particularly curdlan (CrdS) and Cel synthases. In Cel synthases the C-terminal region encloses the PilZ domain involved in c-di-GMP binding and required for enzyme activation (22). Although BgsA does not seem to contain a PilZ signature, we provide strong evidence that it binds c-di-GMP through its C-terminal region, which could represent a new c-di-GMP-binding domain that awaits characterization. Therefore, it is likely that induction of ML β-glucan by c-di-GMP takes place through allosteric activation of the BgsA GT. It also will be of interest to identify the DGC(s) involved in the provision of c-di-GMP to BgsA and the physiological conditions that promote its activation. Synthesis of the ML β-glucan also is dependent on the ExpR/SinI system which regulates bgsBA transcription, thereby linking ML β-glucan to the production of other S. meliloti EPSs (7) Thus, ML β -glucan biosynthesis seems to be regulated at multiple levels, a situation that also has been proposed for cellulose biosynthesis in A. tumefaciens (54).

We observed that high production of ML β -glucan facilitates aggregation and biofilm formation by *S. meliloti* 8530 under freeliving conditions and also on the alfalfa root surface. ML β -glucan overproduction greatly enhances bacterial attachment to the host plant root and the formation of a dense biofilm on the root surface, from which mutants unable to produce ML β -glucan seem to be excluded. Moreover, under physiological c-di-GMP levels *bgsA* mutants show reduced attachment to alfalfa roots. The involvement of ML β -glucan in *S. meliloti*'s firm attachment to roots and its aggregation and biofilm formation on abiotic and biotic surfaces suggest that it can be of particular importance for bacterial persistence and multiplication in the soil and rhizosphere under field conditions. This evokes the biological role of related glucans such as cellulose, an important adherence factor in very diverse bacteria (55), including rhizobia. Like the *S. meliloti* ML β -glucan, cellulose also provides Rhizobium leguminosarum with a root attachment advantage; however, both cellulose-deficient and -overproducing strains show wild-type nodulation properties (56, 57) similar to those of the S. meliloti bgsA mutants. With resemblance, A. tumefaciens cellulose-deficient mutants also show reduced attachment to plant cells but remain fully virulent (54) except under very particular experimental conditions [e.g., when the infection sites are washed shortly after inoculation], suggesting that cellulose may be important for bacterial attachment to plant wound sites under adverse field conditions (58). Phylogenetic analyses suggest that, like S. meliloti, other Rhizobiales, including nonnodulating plant-interacting bacteria such as Agrobacterium and Methylobacterium, also are able to produce ML $(1\rightarrow 3)(1\rightarrow 4)$ - β -D-glucan. This novel polymer adds to the plethora of surface polysaccharides that help rhizobia and other plant-associated bacteria thrive in the changing soil and rhizosphere environments.

Materials and Methods

Bacteria and Culture Conditions. Bacterial strains and plasmids used in this work are listed in *SI Appendix*, Table S1. Starting cultures of *S. meliloti* strains were grown overnight at 28 °C on tryptone-yeast extract-CaCl₂ (TY) broth (59) or MM medium (60). When required, antibiotics and other compounds were added at the following final concentrations: tetracycline, 10 µg/mL; CF, 50 µg/mL (in solid medium) or 100 µg/mL (in liquid medium). The plasmids pJBpleD*, pJB3Tc19, pJB9091, and pJBpleD9091 were introduced in rhizobia by conjugation using the *E. coli* β 2163 donor strain (61) in matings as previously described (62).

Construction of S. meliloti Mutants and Complementing Plasmids. Molecular biology techniques were performed according to standard protocols and manufacturers' instructions. For the construction of Sme SMb20391::Nm and SMb20460::Nm mutants, the phage Φ M12 (63) was used to transduce mutations from the donors S. meliloti 2011mTn5STM.1.08.B12, 2011mTn5STM.4.13. E03, and 2011mTn5STM.4.03.E12 to Sme, as previously described (64). Mutants were verified by PCR and Southern hybridization with specific probes. For the construction of Sme \triangle SMb20390 with an unmarked in-frame deletion, two fragments flanking the ORF were amplified separately, using primer pairs op90-F/20390L-R (496 bp) and 20390R-F/SM_b20391-R (408 bp) (SI Appendix, Table S2). These two fragments were purified and used as template DNA for an overlapping PCR with primers op90-F and SM_b20391-R. The final fragment was cloned into pCR-XL-TOPO (Invitrogen) and sequenced. A correct Sphl/BamHI DNA insert was isolated and subcloned into the suicide plasmid pK18mobSacB (65), generating the pK18△20390. The pK18△20390 plasmid was introduced into Sme by biparental conjugation with an E. coli S17.1 donor, and the SMb20390 deletion of 1,035 bp (402490-403524) was generated by homologous recombination following the procedures described in ref. 65. The *\DeltaSMb20390* mutant was corroborated by PCR and Southern hybridization with specific probes. Diagrams of the plasmid constructions used in the complementation experiments are depicted in SI Appendix, Fig. S7. For pJB9091 and pJBpleD*9091 constructions, two primers flanking the SMb2039-91 operon were used (op90-F and op91-R) in a PCR with Sme genomic DNA as a template. The 3,720-bp fragment containing both genes was cloned in pCR-XL-TOPO (Invitrogen) and sequenced. A correct EcoRI DNA insert was isolated and subcloned into pJB3Tc19 and pJBpleD* to generate pJB9091 and pJBpleD*9091, respectively. The correct orientation of both constructions was verified with appropriate restrictions enzymes.

Gene-Expression Assays. For RT-PCR experiments, RNA extractions were carried out using the Qiagen RNeasy RNA purification kit (Qiagen). Total RNA (1 µq) treated with RNase-free DNase I Set (Roche) was reverse-transcribed using SuperScript III reverse transcriptase (Invitrogen) and random hexamers (Roche) as primers. RT-qPCR was performed on an iCycler iQ5 (Bio-Rad). Each 25-µL reaction contained 1 µL of cDNA, 200 nM of each primer, and iQ SyBrGreen Supermix (Bio-Rad). Control PCRs of the RNA samples also were performed to confirm the absence of contaminating DNA. Samples were denatured initially by heating at 95 °C for 3 min, followed by a 35-cycle amplification and quantification program (95 °C for 30 s, 55 °C for 45 s and 72 °C for 45 s). Melting-curve analysis was conducted to ensure amplification of a single product. The efficiency of each primer pair (E) was determined by running 10-fold serial dilutions (four dilution series) of Sme genomic DNA as template and generating a standard curve by plotting the log of the dilution factor against the cycle threshold (C_T) value during amplification of each dilution. Amplification efficiency was calculated using the formula



Fig. 9. Neighbor-joining phylogenetic analysis of BgsA proteins and cellulose and curdlan synthases. Bootstrap values (1,000 replicates) are indicated at branching points. Schematic diagrams of ORFs in representative operons are shown by black arrows for glycosyltransferases, gray arrows for membrane fusion proteins, and white arrows for all others.

 $[E = (10^{(1/a)} - 1) \times 100]$, where *a* is the slope of the standard curve. The relative expression of the *SMb20390* and *SMb20391* genes was normalized to that of 16S rRNA gene.

A transcriptional reporter gene fusion also was used to study the effect PleD* on the expression *bgsBA* operon. A strain with an oriented miniTn5gus inserted in *bgsA* was used (*SMb20391::Nm-gus*) (*SI Appendix*, Table S1), and β -glucuronidase was determined as previously described (62).

CF-Binding Assays. Starting cultures of rhizobial strains were prepared as detailed above. After washing with MM, cultures were diluted 1:100 into 10-mL flasks containing MM supplemented with CF (100 μ M final concentration) at 28 °C under agitation for 48 h. Then cultures of the different strains were centrifuged for 10 min at 3,220 × g. Supernatant containing broth with unbound CF of each biological replicate was removed. The pellet was suspended in 2 mL of distilled water and placed in the wells of 24-well plates. Similar growth of all strains was confirmed, and CF-binding measurements for three biological replicates of each strain were performed in a PTI fluorimeter (Photon Technology International); the results were expressed in arbitrary units \pm SE.

Bacterial Attachment to Plant Roots. Alfalfa (*Medicago sativa* L.) seeds were surface-sterilized and germinated as described by Olivares et al. (66). Seed-lings were placed on 12×12 cm plates (12 seedlings per plate) containing nitrogen-free medium (66) solidified with 1.15% (wt/vol) purified agar and were incubated for 3 d in a plant-growth chamber (16-h light/8-h dark photoperiod, 24 °C/16 °C day/night temperature, 75% relative humidity) before inoculation. For root attachment of bacteria at high c-di-GMP levels, seedlings were transferred into a flask containing 25 mL of nitrogen-free plant nutrient solution with a mixture of the wild type and the *SMb20391:: Nm* mutant (10⁹ cells each), both expressing PleD* and tagged with GFP or

dsRed, and were agitated gently at 28 °C for 24 h. After this time, roots were separated from shoots, and four groups of three roots each were prepared and washed vigorously four times with 1 mL of sterile deionized water in Eppendorf tubes to remove bacteria unbound or weakly bound to roots. After these washings, one group of three roots was used for CLSM with a Nikon Eclipse TE2000-U microscope at the technical service of Estación Experimental del Zaidín Consejo Superior de Investigaciones Cientificas. One milliliter of MM with 2 mM EDTA was added to each of the remaining three groups of roots, and bacteria firmly bound to roots were released by two cycles of vortexing for 1 min and sonication for 1 min. Serial dilutions were spread onto TY with and without Nm and were incubated at 28 °C for 3-4 d; then cfus of each strain were counted. For root attachment of bacteria at physiological c-di-GMP levels, a modification of the protocol described by Anollés and Favelukes was followed (67). Seedlings prepared as above were kept in plates, and each root was inoculated with 30 µL of nitrogen-free plant nutrient solution containing mixtures of Sme and SMb20391::Nm mutant in a 1:1 ratio. Two different inoculum densities were tested, so 10⁵ or 10³ cells were applied to each root. Plants were kept in the plant-growth chamber, and bacteria firmly attached to roots were recovered and counted as above, at 1 and 3 d after inoculation.

Nodulation Experiments. Alfalfa seedlings were prepared as above, except that nine plants per plate were used. For nodulation kinetics each strain was inoculated onto the roots of 45 plants by spreading 30 μ L of a bacterial suspension (prepared in a nitrogen-free plant nutrient solution) along each root surface. Two inoculum densities were tested, so 10⁵ or 10³ cells were loaded on each root. Plants were maintained in the growth chamber, and the number of visible nodules was recorded daily. For competitive nodulation, plants were prepared as above, but bacterial suspensions contained

mixtures of the wild type and the *SMb20391::Nm* mutant in a 1:1 ratio. Two inoculum densities were tested, so 10^5 or 10^3 total cells were loaded on each root. Nodule occupancy was determined 15 d after inoculation. At least 100 nodules were collected, surface sterilized, and individually crushed. The contents of each nodule were spread on TY-agar plates with and without Nm and were incubated at 28 °C for 3–4 d.

ML β-Glucan Isolation and Chemical Characterization. Starting cultures of Sme pJBpleD* were diluted 1:100 in 5-L flasks containing 0.5 L of liquid MM supplemented with tetracycline and were grown with slow shaking (80 rpm) for 3 d at 28 °C. The flocs formed were recovered using a sieve with a 500- μ m cutoff and then were placed in a 50-mL Falcon tube (Fisher Scientific) and washed with 30 mL of boiling Milli-Q water (Millipore Corporation). After boiling for 5 min, the material was cooled at room temperature and centrifuged for 20 min at $3,220 \times g$. The supernatant was discarded, and the washing process was repeated four times. The β -glucan obtained was frozen and lyophilized overnight. For quantification of β -glucan yield, the different strains were grown in 35 mL of MM in 250-mL flasks and were incubated and processed as described above, except that the sieve runoffs were recovered and centrifuged, and bacterial pellets were lyophilized and weighed to calculate the amount of the β -glucan dry weight relative to the total dry weight of the culture. Monosaccharides of the β -glucan were identified upon gas liquid chromatography (GLC)-MS separation of their per-O-trimethylsilylated methyl glycosides (68). The absolute configuration of monosaccharides was assigned following GLC-MS analysis of their per-O-trimethylsilylated (S)- and (R,S)-2-butyl glycosides as described by Gerwig et al. (69). For methylation analysis, a further purification step on the β -glucan purification was applied to increase its final purity. The β -glucan was obtained as described above, and the flocs trapped on the sieve were dissolved in 15 mL of 1% NaOH for 1 h with shaking and were dialyzed in water using molecular porous-membrane tubing (Spectrum Labs) until neutral pH was obtain in the sample, which then was frozen and lyophilized overnight. Lyophilized samples were methylated using the method of Ciucanu and Costello (70). The methylated polysaccharide was purified by SEC on Sephadex LH-20, using CH₂Cl₂:MeOH (1:1) as the eluent. Hydrolysis was carried out first with 88% HCOOH (100 °C, 1 h) and then, after evaporation of the acid, with 2 M trifluoroacetic acid (120 °C, 4 h). Reduction with NaBD₄ and acetylation were performed according to the methods of Kim and coworkers (71), yielding the corresponding partially methylated and acetylated alditols, which were solubilized in CH₂Cl₂ and analyzed by GLC-MS. For acetylation of polysaccharide, 6 mg of polysaccharide were resuspended in 1 mL of pyridine:acetic anhydride (1:1 ratio) for 24 h at room temperature The solution was poured into 5 mL of ice water, and the precipitate was recovered by centrifugation, rinsed with cold water, and freeze dried. This procedure was repeated once. The sample was resuspended in CH₂Cl₂:MeOH (1:1 ratio, vol/vol) and purified by SEC on Sephadex LH-20 with the same eluent. Fractions containing carbohydrates were vacuum-evaporated, dissolved in 300 µL of CDCl₃, and studied by NMR. Spectra were recorded at 300 K on a Bruker Avance III 500 MHz spectrometer operating at 500.40 MHz (¹H) and 125.8 MHz (^{13}C) . Chemical shifts are given in parts per million, using the residual CHCl₃ signal (δ_{H} 7.26 ppm, δ_{C} 77.16 ppm) as reference. Standard Bruker sequences were used for 2D experiments (72). DQF-COSY, TOCSY, and ROESY were performed by using a data matrix of $256 \times 2,048$ points to digitize a spectral width of 5,515 × 5,515 Hz. TOCSY and ROESY experiments were acquired with mixing times of 80 and 100 ms, respectively. The 2D heteronuclear one-bond proton-carbon correlation experiment was registered in the ¹H-detection mode via HSQC. A data matrix of 128 \times 1,024 points was used to digitize a spectral width of 20,843 (F1) \times 5,515 (F2) Hz.

Five milligrams of polysaccharides were digested with 10 U of a commercial lichenase (*endo*-1,3 (4)- β -glucanase; Megazyme) in 1 mL of water overnight at room temperature, including respective controls without the enzyme. Samples were centrifuged at 16,000 × g for 10 min; then supernatants were frozen and lyophilized, dissolved in water, and studied by TLC on silica gel 60 Alugram Sil G/UV (Macherey-Nagel) using butanol:acetic acid:water (2:1:1) as eluent. Carbo-

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hydrates were detected with orcinol:sulfuric acid (73). A mixture of glucose, maltose, maltotriose, maltotetraose, maltopentaose, and maltohexaose (Sigma-Aldrich) oligosaccharides were used as size markers.

Phylogenetic Analyses. Protein sequence similarity searches were carried out with the BLASTP program from the National Center for Biotechnology Information (74). Alignment was performed with CLUSTALW (75). Phylogenetic and molecular evolutionary analyses were conducted using MEGA, version 5.2 (76), using the following phylogenetic parameters: analysis, phylogeny reconstruction; statistical method, neighbor-joining; number of bootstrap replications, 1,000; substitutions type, amino acid; model/method, Poisson model; rates among sites, uniform rates; pattern among lineages, same (homogeneous); gaps/missing data treatment; and pairwise deletion.

Dot Blot c-di-GMP-Binding Assays. Plasmids pET28b::BgsA and pQE80L::C-BgsA were constructed to express the entire or the last 139 amino acids of BgsA, respectively. A 2,068-bp DNA fragment containing full-length bgsA was amplified with primers op91-F and op91-R, cloned into pCR-XL-TOPO, and subsequently subcloned as an EcoRI fragment into pET28b⁺. The correct orientation of the construction was checked using Xhol. Primers C-BgsA-F and C-BgsA-R were used to amplify a 488-bp DNA fragment encoding the last 139 amino acids of BgsA. The amplicon was cloned in pCR-XL-TOPO and subsequently was subcloned as a HindIII/BamHI fragment into pQE80L. E. coli strains carrying pET28b::BgsA, pQE80L::C-BgsA (SI Appendix, Table S1), respective empty vectors (as negative controls), and pET21:PA3353 overexpressing the Pseudomonas aeruginosa PilZ domain protein PA3353 [used as a positive control (47)] were grown to OD₆₀₀ 0.5–0.7 and induced with 1 mM of isopropyl 1-thio-β-D-galactopyranoside for 3 h at 37 °C. After centrifugation, pellets from 8 mL of culture were resuspended in 500 μL of lysis buffer [50 mM Tris (pH 7.5), 10 mM NaCl, 1 mM DTT; and 5% glycerol] and were lysated by sonication (six cycles of 15 s on ice; Branson Sonifier 250). Protein expression was evaluated by SDS/PAGE, and all lysates were standardized with the total protein content determined by Bradford assay (77). For the c-di-GMP dot blotbinding assays, we modified a previously described protocol (47). Briefly, 3 μ g of each lysate sample were spotted on 0.45-µm nitrocellulose membrane discs (Millipore) and left to dry. The membranes were blocked in Tris buffer saline with 0.2% Tween 20 (TBS-T) and 5% skim milk for 1 h at room temperature. Blocked membranes were incubated with 1 µM of 2'-O-(6-[Fluoresceinyl]aminohexylcarbamoyl)-c-di-GMP (2'-Fluo-AHC-c-di-GMP; Biolog) in TBS-T containing 5% skim milk for 1 h at room temperature in a dark room. After washing with TBS-T for 10 min, the dot blots were scanned with a Pharos FX Plus fluorescence imager (Bio-Rad) using the SYBR Green I settings and were quantified by densitometry with the Quantity One software (Bio-Rad). Competitive assavs were performed as indicated above, but membranes were incubated with mixtures of 0.1 µM of 2'-Fluo-AHC-c-di-GMP and increasing quantities (0.1-30 µM) of unlabeled c-di-GMP.

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