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Wall teichoic acids govern cationic gold nanoparticle interaction with Gram-positive bacterial cell walls†

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Molecular-level understanding of nanomaterial interactions with bacterial cell surfaces can facilitate design of antimicrobial and antifouling surfaces and inform assessment of potential consequences of nanomaterial release into the environment. Here, we investigate the interaction of cationic nanoparticles with the main surface components of Gram-positive bacteria: peptidoglycan and teichoic acids. We employed intact cells and isolated cell walls from wild type *Bacillus subtilis* and two mutant strains differing in wall teichoic acid composition to investigate interaction with gold nanoparticles functionalized with cationic, branched polyethylenimine. We quantified nanoparticle association with intact cells by flow cytometry and determined sites of interaction by solid-state ³¹P- and ¹³C-NMR spectroscopy. We find that wall teichoic acid structure and composition were important determinants for the extent of interaction with cationic gold nanoparticles. The nanoparticles interacted more with wall teichoic acids from the wild type and mutant lacking glucose in its wall teichoic acids than those from the mutant having wall teichoic acids lacking alanine and exhibiting more restricted molecular motion. Our experimental evidence supports the interpretation that electrostatic forces contributed to nanoparticle–cell interactions and that the accessibility of negatively charged moieties in teichoic acid chains influences the degree of interaction. The approaches employed in this study can be applied to engineered nanomaterials differing in core composition, shape, or surface functional groups as well as to other types of bacteria to elucidate the influence of nanoparticle and cell surface properties on interactions with Gram-positive bacteria.

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Introduction

A molecular-level understanding of nanomaterial interactions with bacterial cell surfaces is needed to effectively design nanomaterial-enabled products intended to modulate bacterial populations (e.g., antimicrobials, antifouling surfaces) and for assessing the potential effects of engineered nanomaterials released into the environment. The increase in engineered nanomaterial production and their incorporation into commercial products makes their introduction into the environment inevitable during some portion of their life cycle.¹

Bacteria are vital to many biogeochemical processes and may play a role in the introduction of nanomaterials into some food webs. Molecular-level insight into the interactions of bacteria with nanomaterials may facilitate their design to either target harmful bacteria more effectively or reduce their negative impacts on beneficial microbiota.

The distinct cell envelope architectures of Gram-negative and Gram-positive bacteria² lead to the expectation that their interactions with charged nanoparticles differ, as indeed has been shown in earlier studies.^{3–7} The Gram-negative cell envelope is composed of two phospholipid membranes with a thin

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layer of peptidoglycan sandwiched between them; the outer leaflet of the outer membrane is composed primarily of lipopolysaccharides. In contrast, Gram-positive bacterial cells are bounded by a single cell membrane, which includes lipoteichoic acids anchored into it, and a thick cell wall composed of peptidoglycan with covalently attached wall teichoic acids (WTAs). Previous investigation of electrostatically driven association of nanoparticles with the Gram-negative bacterial outer membrane employed intact and lipopolysaccharide-depleted bacteria and supported lipid bilayers incorporating lipopolysaccharides to demonstrate that lipopolysaccharide structure influenced the extent and distance from the outer membrane lipid bilayer of nanoparticle binding.⁸ Experimental modeling Gram-positive cell surfaces to probe the nanomaterial-bacterium interface requires a different approach be taken. The prominent Gram-positive bacterial cell wall surface structures lack lipid-like character, precluding the use of lipid bilayers to study interactions with nanoparticles. Also, given that teichoic acids can account for up to 50% of the mass of the cell wall⁹ and assist in maintaining cation homeostasis for the cell, understanding their role in interacting with nanoparticles is vital, particularly for nanoparticles functionalized with cationic groups.¹⁰

For *Bacillus subtilis* SB491, a Gram-positive bacterium found in soil and in human and ruminant gastrointestinal tracts, the hydroxyl groups of the glycerol units of the wall teichoic acid poly(glycerolphosphate) backbone can be substituted with glucose (Glc) or alanine (Ala) residues (Fig. 1). Wall teichoic acids are covalently bound to cell wall peptidoglycan *via* a phosphodiester bond formed between the phosphate group bound to the *N*-acetylglucosamine (GlcNAc) residue in the WTA disaccharide linkage unit and the C6 hydroxyl of *N*-acetylmuramic acid (MurNAc) residue in peptidoglycan. Each WTA molecule contains between 45 to 60 glycerolphosphate units.¹¹ Peptidoglycan is composed of alternating *N*-acetylglucosamine (GlcNAc) and MurNAc residues, connected *via* a MurNAc *D*-lactyl group to a tetrapeptide (*L*-alanyl-*D*- γ -glutamyl-*meso*-diaminopimetyl-*D*-alanine). The *D*-Ala of the tetrapeptide is bound to the *meso*-diaminopimetyl of a pentapeptide (*L*-alanyl-*D*- γ -glutamyl-*meso*-diaminopimetyl-*D*-alanine-*D*-alanine), which in turn is bound to another MurNAc residue. Approximately every ninth MurNAc unit contains an attached WTA polymer. For *Bacillus subtilis*, the crosslinking of peptidoglycan strands results in a meshwork having a reported effective pore size of 4.24 or 5 nm.^{12,13}

Bacteria synthesize WTAs within the cytoplasm and translocate them through the cell membrane. Prior to translocation, Glc is added to the C2 hydroxyl of the WTA glycerol *via* the wall teichoic acid glycosyltransferase Tag E (encoded by the teichoic acid glycerol (*tagE*) gene).¹⁴ After translocation, a proposed *D*-alanyl carrier protein ligase (encoded by the *D*-alanyl-lipoteichoic acid A (*dltA*) gene) attaches *D*-Ala to the WTAs also to the C2 hydroxyl of the WTA glycerol.^{10,15} In the lipoteichoic acids of *B. subtilis*, approximately 9% of the glycerolphosphate moieties become substituted with *D*-Ala and 64% with Glc.¹⁵ The degree of substitutions of wall teichoic acids have not been reported to our knowledge. Deletion of the *tagE* gene results in

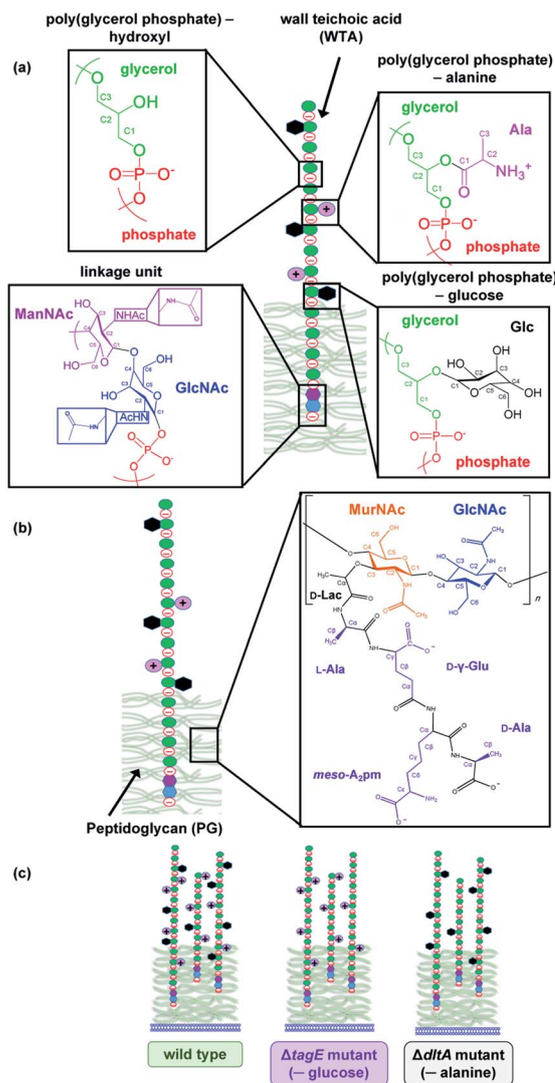


Fig. 1 Gram-positive bacterial cell wall, including cytoplasmic membrane (blue), peptidoglycan (light green), and teichoic acids (red and green). Structures with labelled carbons of (a) the poly(glycerolphosphate) backbone, alanine and glucose side chains, disaccharide linkage unit between the peptidoglycan and wall teichoic acids, and (b) the sugar and amino acid residues in peptidoglycan are shown. (c) Pictorial representations of the cells prior to sacculi isolation of the three species. The wall teichoic acid of wild type contains both alanine and glucose, that of $\Delta tagE$ contains only alanine, and that of $\Delta dltA$ contains only glucose. Abbreviations: ManNAc, *N*-acetylmannosamine; Glc, glucose; GlcNAc, *N*-acetylglucosamine; MurNAc, *N*-acetylmuramic acid; *D*-Lac, *D*-lactyl; *L*-Ala, *L*-alanyl; *D*- γ -Glu, *D*- γ -glutamyl; *meso*-A₂pm, *meso*-diaminopimetyl; *D*-Ala, *D*-alanine.

elaboration of WTA lacking Glc attached to the poly(glycerolphosphate) backbone.¹⁴ Deletion of genes in the *dlt* operon of *B. subtilis* results in the production of teichoic acids lacking *D*-Ala and a concomitant increase in methicillin susceptibility.¹⁶ Similarly, the absence of *D*-alanylation in Gram-positive *Lactococcus lactis* resulted in decreased resistance to cationic antimicrobials nisin and lysozyme.¹⁷ These results reinforce the notion that the lack of *D*-Ala in teichoic acids can



alter interaction of the cell surface with external factors, such as antimicrobial agents and potentially nanomaterials.

In the present study, we investigated the interactions of WTA from the Gram-positive bacterium *Bacillus subtilis* with gold nanoparticles functionalized with cationic branched polyethylenimine (bPEI-AuNPs). The bPEI-AuNPs were chosen for the colloidal stability conferred by their strong positive charge (*vide infra*) as well as for the availability of bPEI for use as an experimental control. We focused on the cell surface components responsible for the interaction of cationic nanoparticles with bacterial cell surfaces because prior investigation demonstrated negligible interaction of wild type *B. subtilis* cells with anionic gold nanoparticles.³ We employed three strains of *B. subtilis* possessing the same peptidoglycan structure but differing in WTA structure due to genetic modifications: wild type (SB 491), the *tagE* knockout strain $\Delta tagE$ with non-glycosylated WTA, and the *dltA* knockout strain $\Delta dltA$ with WTA lacking D-Ala. Experiments employed intact cells or bacterial exoskeletons (sacculi) composed of peptidoglycan and covalently bound WTAs. We note that the sacculi did not include lipoteichoic acids which are anchored in the cell membrane or can be noncovalently associated with the peptidoglycan matrix.¹¹ Stable isotope-labelled sacculi were used to facilitate investigation of nanoparticle interaction with WTA and peptidoglycan *via* solid-state nuclear magnetic resonance (NMR) spectroscopy. Solid-state NMR has been previously applied to characterize WTA of different bacterial species, propose divalent cation binding sites on bacterial sacculi, investigate the interaction of cationic polymers with sacculi, and determine the influence of antibiotics on cell wall composition.^{18–20} To our knowledge, the present work represents the first solid-state NMR investigation of nanomaterial interaction with bacterial cell wall components. Flow cytometry performed on intact bacterial cells showed that WTA composition impacts cationic gold nanoparticle association with bacterial cells. Solid-state ³¹P- and ¹³C-NMR experiments performed on hydrated sacculi allowed identification of the chemical groups in peptidoglycan and WTA involved in binding.

Results and discussion

Bacterial cell surface charge

We investigated the interaction of cationic bPEI-AuNPs with the surfaces of wild type and WTA-mutant Gram-positive *Bacillus subtilis* cells. We hypothesized that bPEI-AuNP association with the cell surfaces is governed predominantly by electrostatic interactions. We evaluated the impact of the genetic mutations on overall cell surface charge by two approaches: whole-cell electrophoretic mobility (u_e) measurement and the binding of the cationic protein cytochrome *c*. The permeability of bacterial surface structures (*e.g.*, WTA) to ions and water (*i.e.*, the “softness” of the surface) precludes determination of cell surface ζ -potential from electrophoretic mobility measurements.²¹ We therefore compared electrophoretic mobilities of the bacterial cells directly^{22,23} recognizing that in addition to the potential at the polyelectrolyte–solution interface, u_e depends on the thickness, charge density, permeability, and homogeneity of the

soft layer.²⁴ All three strains exhibited negative electrophoretic mobility (Fig. 2a), largely attributable to the phosphate groups in the WTA on the cell surfaces. The electrophoretic mobilities of the three strains were statistically indistinguishable.

We measured the binding capacity of the cationic protein cytochrome *c* (+8 at pH 7) for *B. subtilis* cells of each strain as a proxy for their relative amounts of anionic surface charge. The absorbance of cytochrome *c* solutions at 530 nm was used to determine the amount of the protein removed from solution upon exposure to bacterial suspensions to provide a measure of the amount of negative charge on the cell surfaces.¹⁶ Fig. 2b shows that the reduction in A_{530} was larger for $\Delta tagE$ strain than for the $\Delta dltA$ strain indicating that the former possessed a larger number of negatively charged surface sites. The extent of cytochrome *c* binding to the wild type and $\Delta dltA$ strains was statistically indistinguishable, suggesting comparable anionic surface site densities. This and the equivalent u_e for these strains contrasts with earlier reports of increased negative surface charge in *dltA*-gene deleted strains.^{11,16} Our results are, however, consistent with those reported for a different Gram-positive bacterium, *Lactococcus lactis*, for which changes in the extent of D-alanization did not impact electrophoretic mobility.¹⁷

WTA composition

To verify that the *B. subtilis* mutants expressed the expected phenotypes, we examined the Ala and Glc contents of WTA isolated from the three strains. We hydrolyzed Ala from WTA and derivatized it with Marfey's reagent to allow detection *via* absorbance at 820 nm.^{25,26} D-Ala was clearly present in WTA from both wild type and $\Delta tagE$ strains, but was significantly diminished in the $\Delta dltA$ sample (Fig. S1†). The clear differences in the chromatograms indicate that knockout of the D-alanyl-lipoteichoic acid A ($\Delta dltA$) gene resulted in the expected phenotype.

We conducted solution ³¹P-NMR experiments to obtain evidence for the Glc content of isolated WTA from each *B. subtilis* strain. Fig. 3 compares the ³¹P-NMR spectra from the strains collected using a pulse sequence to directly observe P nuclei. The ³¹P-NMR spectra of isolated WTA exhibit three main peaks centered at approximately 0.73 (peak A), 0.38 (peak B),

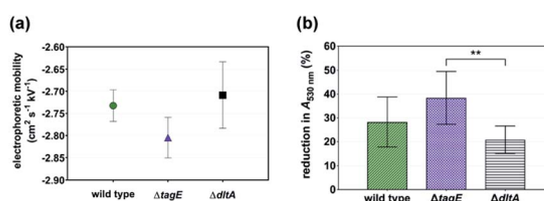


Fig. 2 Assessment of the surface charge of intact wild type and WTA mutant *Bacillus subtilis* cells: (a) electrophoretic mobility of cells and (b) reduction in absorption at 530 nm to assess the binding of cationic cytochrome *c* to intact cells; a larger reduction in A_{530} nm corresponds to a more negative cell surface charge (**, $p < 0.01$). The $\Delta tagE$ and $\Delta dltA$ mutants lack glucose and alanine residues, respectively, in their wall teichoic acids.



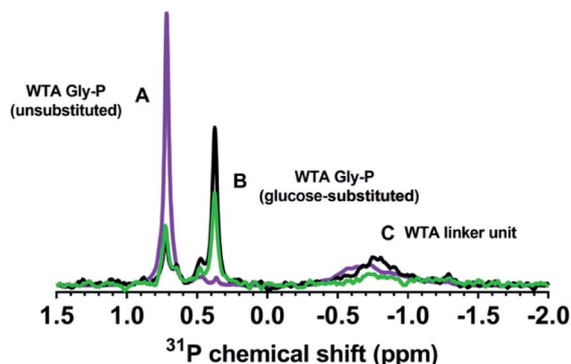


Fig. 3 Solution ^{31}P -NMR spectra of isolated WTAs confirm WTA phenotypes of the wild type (green), ΔtagE mutant (purple), and ΔdltA mutant (black) of *Bacillus subtilis*. Peak A reflects unsubstituted glycerolphosphate groups, peak B is due to glucose-substituted glycerolphosphate, and peak C is assigned to phosphorus in the linker unit. Intensity from ΔtagE mutant spectrum was reduced by 20-fold to compare relative peak positions. The amounts of WTA analyzed for each strain were from cell suspensions normalized to equivalent optical density. Sodium phosphate was used as an internal standard (chemical shift = 2.5 ppm at pH \sim 8) and was added at the same concentration in each sample to allow quantitative comparison of samples.

and -0.80 (peak C) ppm. Peaks A and B are both assigned to the phosphorus in the WTA poly(glycerolphosphate) backbone (Fig. 1) based on previous reports for wall teichoic¹⁸ and lipoteichoic acids.²⁷ Peak C is assigned to phosphorus in the linker unit as previously reported.^{28–30} A distinctive feature of the ΔtagE mutant (WTA glycosyltransferase Tag E knockout) WTA spectrum is the absence of peak B. Because peak A arises from the unsubstituted glycerol,⁵¹ we deduce that peak B corresponds to glycerolphosphate substituted with Glc. The proximity in space of an electron-rich Glc residue to the phosphorus nuclei likely causes the observed upfield position of peak B relative to peak A.³¹ The absence of peak B in the ΔtagE spectrum provides clear evidence that the *tagE* gene deletion resulted in the intended phenotypic changes.

Peak positions in ^{31}P -NMR spectra of WTA from the ΔdltA strain closely resemble those in the spectrum from the wild type. We attribute the lack of a unique ^{31}P feature corresponding to the presence of D-Ala on WTA to hydrolysis of the ester bond and loss of this substituent during the WTA isolation process.^{32,33} Quantitative comparison of the wild type and ΔdltA WTA spectra leads to three important conclusions. First, the ΔdltA mutant likely has more linker units and therefore a larger number of WTA chains than does the wild type. Peak C is more prominent in the spectrum from the ΔdltA mutant than in that from the wild type strain. The ratio of the integrated areas of peaks A + B (WTA backbone phosphates) to that of peak C (phosphorus in the linker unit) was higher for the wild type than for the ΔdltA strain ($p < 0.05$, $n = 4$; Table 1). (The peak ratio for the ΔtagE mutant did not differ from the other two strains ($p > 0.05$.) The broadness of peak C likely reflects a range of chemical environments for the phosphorus in the linker units. Second, the ΔdltA mutant has shorter WTA chains based on the

Table 1 Solution ^{31}P -NMR peak ratios of isolated wall teichoic acids

	Wild type	ΔtagE	ΔdltA
(A + B) : C ^a	2.10 \pm 0.40	1.66 \pm 0.10	1.39 \pm 0.36
A + B	2.33 \pm 0.14	—	3.04 \pm 0.68
A : B ^a	0.76 \pm 0.08	—	0.54 \pm 0.20

^a Wild type and ΔdltA differ ($p < 0.05$).

first conclusion and the observation that the integrated peak areas of A + B, which correspond to the total glycerolphosphates in WTA chains do not differ significantly in wild type and ΔdltA mutant. Third, the extent of Glc substitution (which gives rise to peak B) is higher for the ΔdltA strain than the wild type as indicated by comparison of the integrated areas of peak A and B (A : B in Table 1). In summary, the ^{31}P -NMR data are consistent with ΔdltA mutant having more numerous and shorter WTA chains with more Glc substitution than wild type.

The properties of the bacterial cells and their WTA described above confirmed that the mutations led to the expected changes in WTA structure. However, the trends in net cell surface charge ran counter to the straightforward explanation that ΔtagE differed only in lacking Glc and ΔdltA differed only in lacking D-Ala. Rather, the results suggest that cell surface characteristics of the ΔtagE and ΔdltA mutants differ beyond the mere absence of Glc and D-Ala, respectively. Comparison of signal intensities in the ^{31}P -NMR spectra suggests that WTA from the ΔtagE strain contained a larger amount of phosphate relative to that from the other two strains. This was corroborated by colorimetric phosphorous analysis (Fig. S2†).³⁴ Therefore, we hypothesized that the small variations in surface charge among *B. subtilis* mutants arose from differences in the length or number of WTA chains. Wall teichoic acid contributes substantially to the surface charge of Gram-positive bacteria. Phosphorus analysis alone cannot discriminate whether the higher P content of the WTA from the ΔtagE strain was due to a larger number of or longer WTA chains. Nonetheless, more negative surface charge of the ΔtagE relative to the ΔdltA strain (Fig. 2b) may be related to the larger amount of glycerolphosphate in the former. Comparison of the phosphate content of wild type and ΔtagE strains suggests that factors in addition to WTA glycerolphosphate (e.g., lipoteichoic acids, not included in the total phosphate determination here) may contribute to overall negative surface charge of the cells.

Solid-state ^{31}P -NMR analysis of sacculi

We acquired solid-state ^{31}P -NMR spectra of isolated sacculi, which contain only peptidoglycan and WTA, from each strain and compared them with the solution ^{31}P -NMR spectra for the corresponding isolated WTAs (Fig. 4a–c). The solution and solid-state NMR spectra resembled one another for each strain, although the peaks in the solid-state spectra exhibited the broadening expected from the restricted molecular motion of WTA covalently bound to and embedded within the peptidoglycan meshwork of the sacculi. Larger molecules have shorter transverse, T_2 , relaxation times, resulting in broader peaks.³⁵



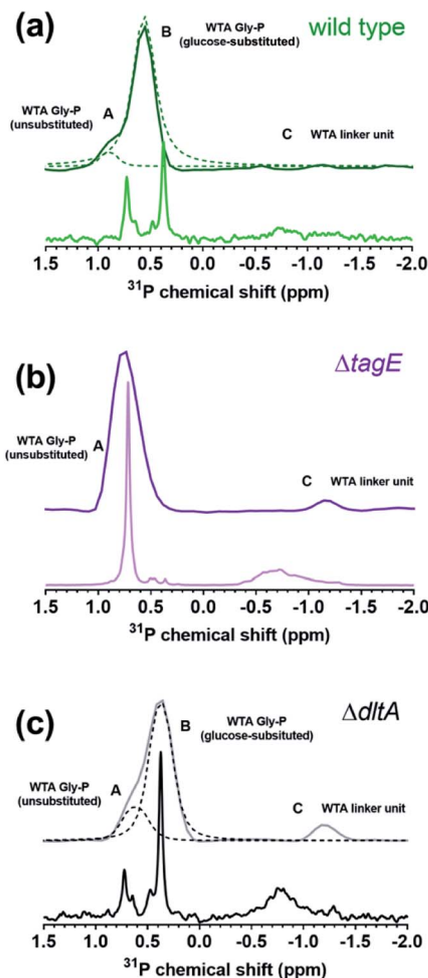


Fig. 4 Solution and solid-state ^{31}P -NMR spectra for (a) wild type, (b) ΔtagE , and (c) ΔdltA strains of *Bacillus subtilis*. Dashed lines in the solid-state spectra indicate deconvoluted peaks. Peak A reflects unsubstituted glycerolphosphate groups, peak B is due to glucose-substituted glycerolphosphate, and peak C is assigned to phosphorus in the linker unit. Sodium phosphate (at 2.5 ppm) and hexamethylphosphoramide (at 29.97 ppm) were used as internal standards for the solution and solid-state spectra, respectively.

Peak shifts are presented in Tables S1 and S2,[†] and additional observations are provided in the ESI.[†]

The positions of peaks A and B differ slightly in the deconvoluted solid-state spectra for wild type (Fig. 4a) and ΔdltA (Fig. 4c) relative to the solution NMR spectra; this is attributable to either the presence of shoulders on the peaks in the solution spectra or differences in chemical environment of the phosphorus nuclei (*i.e.*, the presence of peptidoglycan in the sacculi samples). Overall, the close resemblance of ^{31}P -NMR spectra of isolated WTA and cell wall sacculi suggests that the phosphorus signals in the sacculi samples arise exclusively from WTA, confirming the purity of the isolated sacculi with respect to other P-containing species. Though unlikely given the thorough washing steps, any phospholipids remaining in the sacculi or WTA samples would produce signals in the same spectral region (between 1 and -1 ppm).^{36,37} Specifically, the ^{31}P

chemical shifts produced by phosphatidylethanolamine and phosphatidylglycerol lipids (the dominant phospholipids in Gram-positive bacterial membranes) occur at approximately -0.25 ppm and 0.24 ppm, respectively.³⁸ The absence of such peaks confirms the purity of both sacculi and isolated WTA.

Solid-state ^{13}C -NMR analysis of sacculi

We further probed the sacculi by solid-state ^{13}C -NMR (Fig. 5). Labeled structures are in Fig. 1 and enlarged in Fig. S3.[†] Peaks in these spectra reflect the C atoms in the peptidoglycan amino acid and amino sugar residues and in the WTA (amino) sugar residues and glycerol. The four large peaks dominating the spectra reflect the most abundant cell wall components, which are primarily associated with the GlcNAc and MurNAc amino sugar residues in peptidoglycan. Peak assignments were based on previous reports^{39,40} and are provided in Table S3.[†] The carbons of the WTA glycerol and Glc (when present) produces smaller peaks. The Glc carbon peaks were often difficult to discern; we consider the assignments of these peaks tentative. The other smaller peaks reflect the remaining C atoms in the amino sugar residues in peptidoglycan and the WTA linker unit, and the peptide and D-lactyl group in peptidoglycan.

The ^{13}C spectra for the three strains closely resemble one another with respect to peak positions. The peaks attributed to Glc residues were quite small and are indicated in Fig. 5 by arrows in the spectra of the sacculi from the wild type and ΔdltA strains; these peaks are displayed more clearly in Fig. S4.[†] Wall teichoic acid from the wild type and ΔdltA strains contains Glc, while the WTA of ΔtagE is unsubstituted (see also Fig. 4). Given the nature of cross polarization experiments, rigorous,

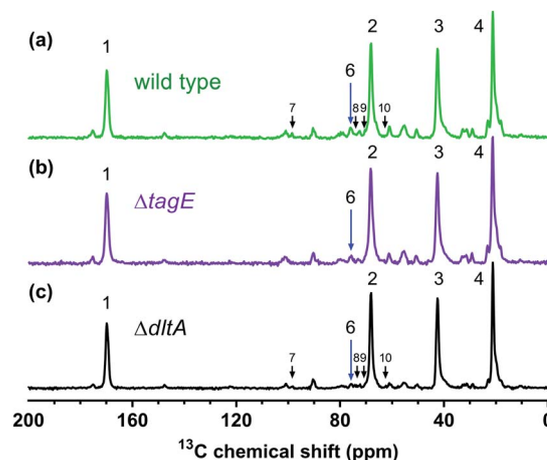


Fig. 5 Solid-state ^{13}C -NMR spectra of sacculi from (a) wild type, (b) ΔtagE , and (c) ΔdltA strains of *Bacillus subtilis*. Arrows in black indicate peaks tentatively assigned to Glc carbons. The peaks are assigned as follows: (1) carbonyl carbons in peptidoglycan amino acid and amino sugar residues and in WTA linker unit amino sugar residues; (2) C3 in the peptidoglycan amino sugar residues; (3) C2 in the peptidoglycan and WTA linker unit amino sugar residues; (4) methyl groups in the peptidoglycan and WTA linker unit amino sugar residues; (5) glycerol C1 and C3; (6) glycerol C2 (blue arrow); (7) C1 from Glc (tentative); (8) C2, C3, and C5 from Glc (tentative); (9) C4 from Glc (tentative); and (10) C6 from Glc (tentative).



quantitative comparison of peak areas among the three strains is not possible. Cross polarization transfers magnetization from a nearby network of abundant $\frac{1}{2}$ -spin nuclei (^1H in our case) to the observed nucleus (^{13}C) during the contact period. Thus, peak intensity and area depend on both the abundance of the observed nuclei and the number of protons on nearby carbon nuclei (within 3–4 Å). The solution ^{31}P -NMR results indicated that the $\Delta dltA$ strain contains more Glc than wild type. We further note that the $\Delta tagE$ strain exhibited the largest ^{13}C chemical shift at C1 and C3 and the smallest at C2 of the glycerolphosphate (Table S3†).

To provide a measure of the rigidity of the peptidoglycan meshwork and the covalently bound WTA molecules, we quantified longitudinal relaxation time-constants, T_1 , for ^{13}C in sacculi from the three *B. subtilis* strains using inversion-recovery experiments (Fig. S6 and S7†). Our analysis focused on nuclei for which we can assume dipole–dipole interaction dominates relaxation rather than chemical shift anisotropy. We therefore omitted data from carbonyl carbons from the analysis. Relaxation times were measured between -8 and 30 °C to determine the molecular regime (small or large) based on the increase in molecular motion with temperature. With the exception of C γ in the Glu residues of $\Delta tagE$ sacculi, T_1 decreased as temperature increased for all carbon nuclei analyzed (fitted slopes showed non-zero linear dependence; Fig. S5 and Table S4†). For residues exhibiting a decrease in T_1 with increasing temperature, larger T_1 values indicate hampered motion and higher structural rigidity and smaller T_1 values reflect more rapid relaxation due to higher molecular motion.^{18,41}

We determined T_1 values for the ^{13}C nuclei in peptidoglycan and WTA indicated in Fig. 6, as well as for C γ of Glu and C β of *meso*-A₂pm in peptidoglycan (Fig. S5†). We focused on peaks attributable solely to peptidoglycan or to WTA. At 30 °C, the temperature closest to the sample temperature in the solid-state NMR experiments, T_1 values for peptidoglycan amino sugar residues and the WTA glycerolphosphate of $\Delta dltA$ were larger than those of the wild type and $\Delta tagE$ strains (Fig. 6). In

particular, T_1 of C2 (the location of Glc substitution in the wild type and $\Delta dltA$ strains) was larger in $\Delta dltA$ than for wild type and $\Delta tagE$, reflecting more restricted molecular motion for the $\Delta dltA$ strain. Differences among T_1 values for the three strains were also apparent at 18 and 12 °C, but not at lower temperatures ($p > 0.05$; Fig. S6†). The T_1 values indicate that the molecular motion of peptidoglycan strands from the $\Delta dltA$ strain are more restricted than those from the wild type and $\Delta tagE$ strains, which exhibit similar molecular motion. Likewise, the WTA from the $\Delta dltA$ strain exhibits more restricted molecular motion than does that from the other two strains. The glycerolphosphate carbons bearing $-\text{OH}$ groups (*viz.* C1 and C3) exhibit somewhat higher molecular motion in the $\Delta tagE$ strain than in the other two strains. For resonances associated with amino acid residues in peptidoglycan, T_1 values did not differ among the three strains at any temperature ($p > 0.05$).

The lower molecular motion of WTA in $\Delta dltA$ sacculi relative to those of the other two strains is consistent with the more abundant Glc substitution of its glycerolphosphate (Table 1), the larger number of WTA molecules bound to peptidoglycan (Fig. 3), and less protrusion into solution due to fewer glycerolphosphate repeat units per WTA molecule (Fig. S7†). As noted above, the $\Delta dltA$ strain appears to produce a larger number of shorter WTA molecules than the other two strains. Shorter WTA strands dissipate energy less efficiently, leading to longer relaxation times. Both solution ^{31}P -NMR and phosphorus analysis indicate that the $\Delta tagE$ strain contains the most glycerolphosphate groups, although this strain lacks Glc side chains, and the distance that WTA molecules protrude beyond the peptidoglycan meshwork is unknown. Overall, we hypothesize that differences in molecular motion result from the difference in type and number of substituents on the WTA glycerol, and the extent to which the WTA molecules protrude from the peptidoglycan layer. Specifically, the glycerol substituent Glc is larger than $-\text{OH}$; high amounts of Glc substitution and close proximity of Glc residues to other Glc residues would slow the molecular motion of WTA. Furthermore, more hindered molecular motion is expected for shorter WTA as they extend into solution from the peptidoglycan matrix to a smaller degree.

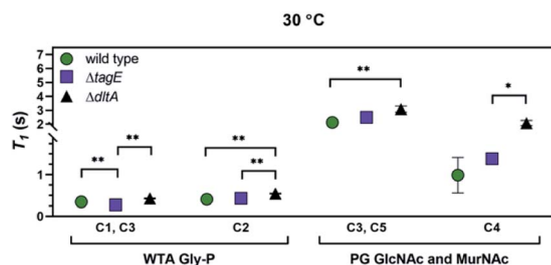


Fig. 6 T_1 relaxation time constants at 30 °C for the indicated ^{13}C resonances of sacculi from wild type, $\Delta tagE$, and $\Delta dltA$ strains of *B. subtilis*: GlcNac C3 and MurNac C5 of peptidoglycan, GlcNac and MurNac C4 of peptidoglycan, glycerolphosphate C1 and C3 of wall teichoic acids, and glycerolphosphate C2 of wall teichoic acids. Non-linear fit statistical significance: *, $p < 0.05$; **, $p < 0.01$. Error bars represent standard error in non-linear fitting from T_1 relaxation time equation. See Fig S6† for T_1 relaxation times for other temperatures. Abbreviations: GlcNac, *N*-acetylglucosamine; MurNac, *N*-acetylmuramic acid.

Nanoparticle properties

To investigate the differential interactions between the *B. subtilis* strains and cationic nanoparticles, we exposed the bacteria to bPEI-AuNPs with core diameters of 10.9 ± 1.8 nm as indicated by their localized surface plasmon resonance peak,⁴² in good agreement with the diameter reported by the manufacturer based on analysis of transmission electron microscopy images (12.1 ± 0.8 nm). We characterized the hydrodynamic and electrokinetic properties of the particles in 0.025 M NaCl buffered to pH 7.4 with 0.002 M HEPES, the solution we used to study the interaction of the bPEI-AuNPs with intact cells. Under these solution conditions, the number mean hydrodynamic diameter was 37.7 ± 0.3 nm, and the ζ -potential was $+23 \pm 5.9$ mV. The bPEI ligands are covalently attached to the nanoparticle surfaces. Nevertheless, we determined that a 0.93 nM



solution of bPEI-AuNPs contained 792 ± 72 nM bPEI, using the reported average molecular mass for bPEI of 25 000 Da.

Nanoparticle–cell surface association

We employed flow cytometry to quantify the number of *B. subtilis* cells of each strain that had bPEI-AuNPs associated with them.^{3,8} We previously demonstrated that flow cytometry can be used to quantify the number of bacterial cells having AuNPs associated with cell surfaces and that the measurements correlate well with qualitative TEM observations.^{3,43} Cells were stained with a membrane-permeant nucleic acid stain to allow intact cells to be sorted from cellular debris. The strong localized surface plasmon resonance scattering signal from the AuNPs allowed cells with surface-associated AuNPs to be discriminated from those lacking associated AuNPs. The detected AuNP signals in flow cytometry did not necessarily originate from individual AuNPs; previous TEM analyses have revealed the likelihood of NP aggregation on bacterial surfaces.^{3,6,7} Fig. 7 shows the proportion of cells from each strain with bPEI-AuNPs associated with their surfaces (of 10^4 bacterial cells). Fewer $\Delta dltA$ cells had AuNPs associated with them than did the wild type and $\Delta tagE$ strains ($p < 0.01$). No difference was observed between the wild type and $\Delta tagE$ strains.

To gain further insight into the chemical basis for the difference in bPEI-AuNP association with cells from the three strains, we employed solid-state ^{31}P - and ^{13}C -NMR. Solid-state ^{31}P -NMR spectra were collected from sacculi from the three *B. subtilis* strains before and after exposure to 34 nM bPEI-AuNPs and to 29.1 μM free bPEI polymer, the estimated amount of free polymer in a 34 nM AuNP solution (Fig. 8). For the wild type and $\Delta tagE$ strains, the peaks associated with backbone phosphates shifted upfield upon nanoparticle interaction (Table S2[†]). For the $\Delta dltA$ strain, the dominant peak in the spectrum (peak B) appears to shift downfield upon nanoparticle addition. For all three strains, the peaks associated with backbone phosphates shifted substantially downfield upon exposure to free bPEI polymer.

Upfield peak shifts typically result from shielding due to addition of electron density to the observed nuclei. Downfield

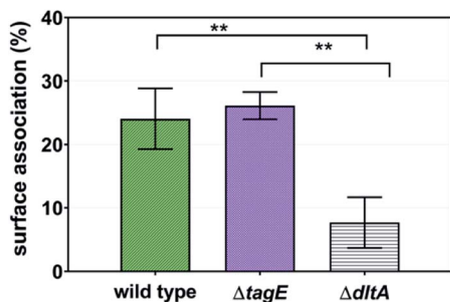


Fig. 7 Percentage of cells from each *B. subtilis* strain with surface-associated bPEI-AuNPs from flow cytometry measurements of 10^4 bacterial cells. Bacterial cultures were exposed for 10 min to 0.93 nM bPEI-AuNPs.

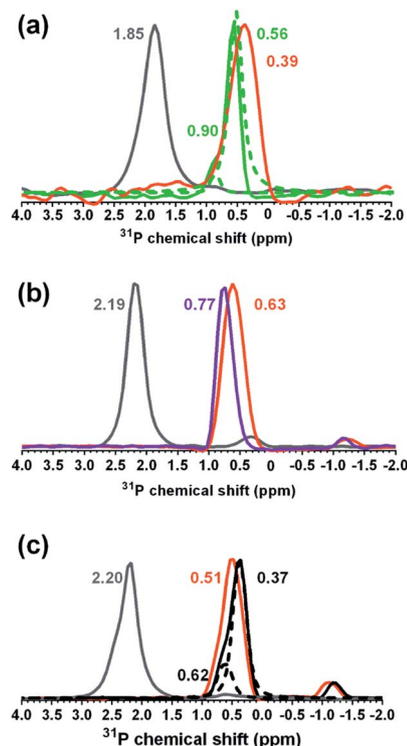


Fig. 8 Solid-state ^{31}P -NMR spectra for the (a) wild type (green), (b) $\Delta tagE$ (purple), and (c) $\Delta dltA$ (black) strains before and after exposure to 34 nM bPEI-AuNPs and 29.12 μM free bPEI polymer. In (a) and (c), deconvolved spectra are shown with dashed lines. The orange traces reflect phosphorus signals upon interaction with 34 nM AuNPs. Grey traces correspond to sacculi exposed to 29.12 μM free bPEI polymer, the estimated amount of free polymer in a 34 nM AuNP solution. Numerical values in green, purple, or black (sacculi for each strain), orange (with AuNPs), and grey (with free bPEI polymer) are the corresponding chemical shifts for predominant peaks and deconvolved shoulders.

peak shifts arise from deshielding, due to removal of electron density. The downfield peak shifts observed for all strains upon exposure to bPEI polymer (grey traces in Fig. 8) reflect removal of electron density from the P nuclei, likely due to formation of hydrogen bonds between primary amines of bPEI and WTA phosphate oxygen.⁴⁴ In contrast, addition of bPEI-AuNPs produced peak shifts in the opposite direction for the wild type and $\Delta tagE$ strains. This change is clearly not the result of free bPEI polymer. These results imply that wild type and $\Delta tagE$ phosphorus nuclei experienced an increase in electron density in the presence of bPEI-AuNPs, which we attribute to the proximity of the AuNPs which contain delocalized electrons.⁴⁵

Exposure to either the bPEI-AuNPs or the free bPEI caused broadening of the ^{31}P peaks in spectra acquired from all three strains (Tables S5–S7[†]). This broadening may be due to heterogeneity in the interaction of the WTA phosphorus atoms with both bPEI-AuNPs and free bPEI polymer.⁴¹ Peak broadening may also arise from reduced mobility and slower relaxation times due to the presence of bPEI-AuNPs. Restricted WTA motion could result from the interactions with bPEI-AuNPs or free bPEI polymer. In a similar manner, peak broadening in ^1H -



NMR spectra of a potential cancer therapeutic molecule was attributed to its reduced mobility in proximity to the metal core of poly(ethylene glycol)-AuNPs.⁴⁶

We further investigated changes in the solid-state ¹³C-NMR spectra induced by interaction of sacculi from each strain with bPEI-AuNPs (Fig. 9) and free bPEI polymer (Fig. S8†). Carbon nuclei in wild type and $\Delta tagE$ strains clearly experienced more changes in their chemical environments when exposed to bPEI-AuNPs and free bPEI than did the $\Delta dltA$ strain as indicated by changes in chemical shift, peak width, and normalized intensity. Changes to these parameters indicate an alteration in the chemical environment of observed nuclei and suggest possible interaction sites. Previous studies have suggested that nanoparticles can affect the signal intensities of interacting biomolecules to a much larger extent than chemical shifts.⁴⁷

Spectral changes upon introduction of bPEI-AuNPs were similar for the wild type and $\Delta tagE$ strains, with the exception of those associated with the WTA Glc in the wild type. For both strains, exposure to bPEI-AuNPs resulted in substantial attenuation of the intensities of peaks associated primarily with carbons in the amino sugar residues, glycerolphosphate, and Glc (wild type only) of WTA (Tables S5 and S6†). The glycerolphosphate peaks decrease substantially in intensity, and many of the peaks corresponding to the linker unit GlcNAc and ManNAc residues broaden to the point of disappearing into the baseline (Fig. S9†). Other peaks exhibiting decreased intensity, albeit less pronounced than those of the WTA groups, are associated with carbons in the amino acid residues of peptidoglycan (*meso*-A₂pm, L- and D-Ala, Glu) and in the D-Lac

connecting MurNAc to L-Ala (Fig. S9†). The negative charges of D-γ-Glu (and *meso*-A₂pm if not crosslinked) carboxylate groups likely enhanced interaction with bPEI polymer relative to carbons in the glycan sugar residues. Peaks associated with amino acid and amino sugar residues in peptidoglycan decreased somewhat in intensity and shifted slightly downfield (<0.35 ppm), but otherwise remain largely unaltered. Exposure to free bPEI polymer produced spectral changes that resembled those observed in the presence of bPEI-AuNPs, although peak intensity also decreased for some carbon nuclei in the GlcNAc and MurNAc residues of peptidoglycan (Fig. S10†). The spectral changes indicate that WTAs are the primary binding sites for wild type and $\Delta tagE$ sacculi for both AuNPs and free bPEI polymer, and the chemical environment of some of the carbon residues in the peptide of peptidoglycan are also affected. More peaks decrease in the presence of free bPEI polymer relative to bPEI-AuNPs, suggesting that the peptidoglycan meshwork is less accessible to bPEI bound to the NPs relative to free bPEI.

Peak intensities in the spectra from $\Delta dltA$ sacculi remained largely unaltered upon exposure to bPEI-AuNPs or free bPEI polymer (Table S8†). Specifically, peaks corresponding to WTA glycerolphosphate, GlcNAc, and ManNAc were minimally impacted by the presence of nanoparticles. Small decreases in peak intensity and narrowing of peak shoulders were apparent for a few resonances, including the D-Lac group between MurNAc and L-Ala, the WTA linker GlcNAc and ManNAc C5 and C3, and glycerolphosphate C1, 2, and 3 (Fig. S11†). Similar, rather minor changes are observed in the presence of free bPEI.

Both flow cytometry analysis of whole bacteria cells and solid-state NMR analysis of bacterial sacculi revealed low association of bPEI-AuNPs with the surface of the $\Delta dltA$ mutant relative to the other two strains. The flow cytometry results for the $\Delta dltA$ and $\Delta tagE$ strains are consistent with expectations based on cell surface charge as reflected by cytochrome *c* binding (Fig. 2b). The less negatively charged strain, $\Delta dltA$, had much lower proportion of cells associated with AuNPs, suggesting that electrostatic forces contributed to the interactions between whole cells and bPEI-AuNPs. Foxley *et al.* previously reported on the importance of Coulombic interactions between the cationic primary amines of bPEI polymer and the anionic phosphate groups in WTA in methicillin-resistant *Staphylococcus aureus* with slightly different WTA structure from *B. subtilis*.⁴⁴ Similar interactions between the bPEI coating the AuNPs and the WTA glycerolphosphate in the three *B. subtilis* strains is expected. Upon addition of free bPEI polymer, we observed downfield shifts of the dominant peaks in ³¹P-NMR spectra for each strain which corresponded to the most abundant form of glycerol substitution (glucose-substituted (peak B) for wild type and $\Delta dltA$, and unsubstituted (peak A) for $\Delta tagE$). Similar downfield peak shifts were previously reported for a strain of *B. subtilis* with poly(ribitol phosphate) WTA molecules upon exposure to bPEI in solution ³¹P-NMR experiments.⁴⁴ In contrast, we observe small upfield peak shifts in ³¹P-NMR for wild type and $\Delta tagE$ strains upon exposure to bPEI-AuNPs. Comparison of flow cytometry results from the wild type and $\Delta dltA$ strains suggests the electrostatic charge at the cell surface is not the sole factor influencing bPEI-AuNP association with

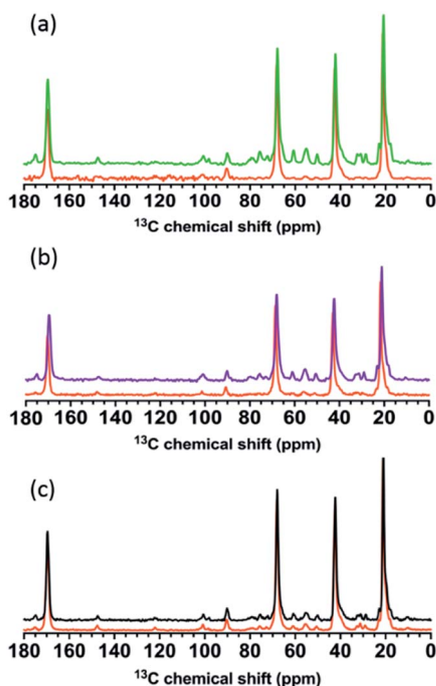


Fig. 9 Solid-state ¹³C-NMR spectra of (a) wild type, (b) $\Delta tagE$, and (c) $\Delta dltA$ before (green, purple, and black, respectively) and after (orange) exposure to 34 nM bPEI-AuNPs.



the bacteria. The surface potentials/charges of wild type and $\Delta dltA$ were not distinguishable by the methods employed; nonetheless, a significantly higher proportion of wild type than $\Delta dltA$ cells were associated with bPEI. We hypothesize that the accessibility of WTA at the cell surface affects association between the bPEI-AuNPs and the bacterial cells.

The T_1 measurements show that specific carbons in $\Delta dltA$ sacculi exhibit more restricted molecular motion than in those of the wild type, $\Delta tagE$, or both. This more constrained molecular motion may be due to a larger proportion of the WTA being embedded within the peptidoglycan meshwork in the $\Delta dltA$ mutant than in the wild type (supported by the evidence for a larger number of shorter WTA strands in the $\Delta dltA$ mutant than in the wild type), making the WTA physically unable to interact with the bPEI-coated AuNPs. The lower flexibility of the $\Delta dltA$ WTA may also reflect the presence of more Glc substituents than are attached to the WTA of the wild type. The $\Delta dltA$ WTA that is available to interact with AuNPs may be less flexible and unable to form as many contacts with the nanoparticles as the WTA from the wild type. Interestingly, although both wild type and $\Delta dltA$ WTA contain Glc, T_1 values are higher for a larger number of $\Delta dltA$ WTA carbons than in the wild type. This further suggests wild type has longer WTA than $\Delta dltA$, supporting the interpretation that the accessibility of the WTA to the AuNPs influences the extent of NP association. Overall, the T_1 results are consistent with observations that wild type and $\Delta tagE$ strains are similar in their interaction with bPEI-AuNPs, while the $\Delta dltA$ strain interacts with the nanoparticles to a much smaller extent as shown by flow cytometry. Peaks in the ^{31}P -NMR spectra from wild type and $\Delta tagE$ sacculi exhibit small upfield peak shifts in the presence of AuNPs, indicating enhanced electron density around the phosphorus nuclei, while a small downfield shift in the main peak is apparent for $\Delta dltA$. Many changes are apparent in the ^{13}C -NMR spectra for sacculi from wild type and $\Delta tagE$ strains upon interaction with bPEI-AuNPs, while minimal changes are observed for sacculi from the $\Delta dltA$ strain. Although the $\Delta dltA$ WTA may be shorter or more buried within the peptidoglycan matrix relative to that in wild type *B. subtilis*, free polymer can more readily penetrate the cell wall than bPEI-AuNPs. This hypothesis is consistent with the pronounced downfield peak shift seen in ^{31}P -NMR for $\Delta dltA$ exposed to free bPEI, while the downfield shift upon exposure to bPEI-AuNPs is quite small.

Conclusions

We have examined the interaction of cationic bPEI-AuNPs with intact bacteria and their exoskeletons (sacculi) using wild type and two mutant strains of *Bacillus subtilis* that differ in the composition of their wall teichoic acids. Through a combination of flow cytometry and solution and solid-state NMR, we have established a relationship between bacterial cell wall composition and the association of cationic AuNPs. Flow cytometry data reveal significantly more bPEI-AuNP associated to the wild type and $\Delta tagE$ strains, the latter of which lacks Glc in its WTA, than to the $\Delta dltA$ strain, the WTA of which lacks Ala and exhibits the most restricted molecular motion. Solid-state ^{31}P -NMR spectra suggest

AuNPs come into closer proximity to the wall teichoic acid molecules of wild type and $\Delta tagE$, compared to $\Delta dltA$, as evidenced by an upfield shift when nanoparticles are present. Solid-state ^{13}C -NMR spectra reveal that bPEI-AuNPs induced spectral changes primarily to the resonances corresponding to the linker unit between WTA and peptidoglycan, to glycerol in the WTA backbone for wild type and $\Delta tagE$, and to the Glc carbons in wild type. For the $\Delta dltA$ strain, exposure to bPEI-AuNPs impacted the chemical environment of WTA and peptidoglycan carbon nuclei to only a small extent. The $\Delta dltA$ strain GlcNAc and MurNAc of peptidoglycan and the WTA glycerol generally exhibited more restricted molecular motion relative to wild type and $\Delta tagE$ as ascertained from temperature-dependent T_1 relaxation experiments. The reduced molecular motion of the WTA in the $\Delta dltA$ strain likely reflected WTA extending to a lesser extent into solution relative to the other two strains and resulted in corresponding diminished interaction with the cationic AuNPs. Similar effects may be seen in cell wall interaction with cationic antimicrobial peptides. The nanoparticle cores used here are inherently larger than the effective pore size of peptidoglycan meshwork and are not expected to penetrate into the cell wall.⁴⁸

Our experimental evidence supports the interpretation that electrostatic forces contribute importantly to interactions of Gram-positive bacterial surfaces with cationic bPEI-AuNPs and that the accessibility of negatively charged moieties in teichoic acid chains influence the degree of interaction. We note that wall teichoic acid structural variations did not significantly impact cell viability (Fig. S12†) indicating that cell wall mutations may alter NP binding, but these binding differences do not necessarily result in changes to bacterial viability.

We found that major components of *B. subtilis* walls that interacted with nanoparticles include the poly(glycerolphosphate) backbone in WTA, the amino sugars in the disaccharide linker unit, and the peptides in peptidoglycan. Although our findings were based on one type of cationic nanoparticle and one Gram-negative bacterial species, the results demonstrate the importance of the structure and composition of wall teichoic acid in modulating interaction with nanoparticles. The approaches taken in the present study can be extended to additional systems, including engineered nanomaterials differing in core composition, shape, or surface functional groups to elucidate the influence of these properties on interactions with Gram-positive bacteria. Knowledge of such relationships can aid the design of nanomaterials to intentionally target cells to limit microbial growth or to minimize undesired impacts on bacteria.

Materials and methods

Chemicals and nanoparticles

The chemicals and nanoparticles used, suppliers, and purities are described in the ESI.†

Nanoparticle characterization

To verify the nanoparticle properties reported by the manufacturer, the stock particle solution was diluted 100× in 0.025 M



NaCl buffered to pH 7.4 with 0.002 M HEPES, and UV-Vis extinction spectra were acquired (Beckman DU 7500) to determine AuNP core diameter and concentration.⁴² Particle hydrodynamic diameter and ζ -potential were derived from dynamic and electrophoretic light scattering measurements (Brookhaven ZetaPALS). All measurements were made in triplicate. The particle suspension was stored at 4 °C, and the colloidal stability of the suspension was determined prior to all experiments. The structure of cationic branched polyethylenimine (bPEI) is shown in Fig. S13.†

Bacterial culture and characterization

Bacterial culture. *Bacillus subtilis* wild-type SB 491, $\Delta tagE$, and $\Delta dltA$ strains were purchased from the Bacillus Genetic Stock Center (Columbus, OH). Bacterial colonies were inoculated into sterile lysogeny broth (LB) from LB agar plates and allowed to grow at 37 °C under shaking (275 rpm, ~24 h) until the culture reached an optical density at 600 nm (OD_{600}) of 0.8. Bacterial cells were harvested by centrifugation (750g, 10 min), washed in equal volume Dulbecco's phosphate-buffered saline (D-PBS) buffer, and resuspended in 0.025 M NaCl buffered to pH 7.4 with 0.002 M HEPES.

Bacterial cell electrophoretic mobility. The electrophoretic mobility of bacterial cells was measured using a Brookhaven Instruments Zeta Potential Analyzer (Holtville, NY) following published protocols.^{49,50} Bacterial motility was inhibited with 0.001 M sodium azide to prevent such motion from confounding the interpretation of electrophoretic mobility measurements.^{50,51} The bacterial cells were removed from the sodium azide solution by centrifugation (750g, 10 min), resuspended in 0.025 M NaCl buffered to pH 7.4 with 0.002 M HEPES (OD_{600} adjusted to 0.2), and placed in sample cuvette with the electrode. Electrodes were cleaned between measurements in 10% ethanol for 30 min to reduce electroflocculation, wherein adhesion of cells to the electrodes leads to reductions in the measured absolute electrophoretic mobility over time.⁵²

Bacterial cell cytochrome *c* binding capacity. The cytochrome *c* assay was adapted from published protocols.^{16,53,54} Briefly, the OD_{578} of the bacterial suspension (in 0.025 M NaCl buffered to pH 7.4 with 0.002 M HEPES) was adjusted to 0.94. A 75 μ L aliquot of 10 mg mL⁻¹ cytochrome *c* solution was added to 1.425 mL of each bacterial suspension, as well as to a solution lacking bacteria as a negative control. Samples were lightly agitated for 10 min at ambient temperature to increase mass transport of cytochrome *c* to cell surfaces before centrifugation to sediment cells (12 000g, 15 min). The supernatants containing unbound cytochrome *c* were then removed and oxidized with 20 μ L of 30% H₂O₂. The difference between absorbance at 530 nm of supernatants after exposure to cells and of the negative control provide an indication of the relative surface charge of the cell surfaces.

Flow cytometry to investigate bacteria association with nanoparticles. The fraction of the bacterial cells that had AuNPs attached on their surfaces was assessed by flow cytometry.^{3,8} Briefly, bacterial cultures were diluted in 0.025 M NaCl buffered to pH 7.4 with 0.002 M HEPES to an OD_{600} of 0.2 and exposed to

0.9 nM bPEI-AuNP for 10 min. The fluorescent dye, SYTO9 (Thermo-Fisher, Waltham, MA) was added to stain the nucleic acid for 15 min. Ten thousand cells were sorted from each sample in triplicate using a flow cytometer equipped with a 20 mW, 488 nm laser (Becton Dickinson LSRII SORP). Side scattering intensity based on the plasmonic extinction of AuNPs was monitored to identify the population of cells with nanoparticles bound on surfaces. Exposure of the three *B. subtilis* strains to either 0.93 nM bPEI-AuNPs or to the amount of free bPEI polymer in such a suspension (Fig. S12†) produced no discernable impact on bacterial viability (assay described in the ESI†).

Wall teichoic acid isolation and characterization

Wall teichoic acid isolation. Wall teichoic acid was extracted and isolated from each bacterial strain according to a published protocol.³² Briefly, a bacterial culture (20 μ L) was sedimented by centrifugation (2000g, 10 min), washed with 25 mL of 0.05 M MES at pH 6.5, and resuspended in 25 mL of 4% [wt/vol] SDS in 0.05 M MES at pH 6.5. Each suspension was adjusted to the same OD_{600} value to ensure approximately the same number of cells were extracted. The bacterial samples were placed in boiling water for 1 h, centrifuged (4000g, 10 min), and resuspended with 1 mL of 4% [wt/vol] SDS in 0.05 M MES at pH 6.5. The samples were sedimented by centrifugation (14 000g, 10 min), and washed with 2% NaCl [wt/vol] buffered to pH 6.5 with 0.05 M MES, followed by 0.05 M MES at pH 6.5. The samples were incubated with proteinase K (0.02 M Tris-HCl, 0.5% [wt/vol] SDS, 20 μ L mL⁻¹ of proteinase K, pH 8.0) at 50 °C for ~4 h and washed again with 2% NaCl [wt/vol] buffered to pH 6.5 with 0.05 M MES and at least twice with ultrapure water. Finally, the samples were suspended in 1 mL of 0.1 M NaOH and gently shaken at room temperature for ~16 h. Cell wall debris was removed by centrifugation (14 000g, 10 min), and the supernatant containing isolated WTA was collected for further analysis.

Solution ³¹P-NMR of isolated WTA. Isolated WTA were analyzed by solution ³¹P-NMR spectroscopy using a Bruker Avance III HD nanobay 400 MHz spectrometer with BBO smartprobe (³¹P-NMR frequency at 161.87 MHz) with a sample temperature of 298 K. The extracted WTA from each bacterial sample was placed in 5 mm NMR tubes with 20% D₂O. Sodium phosphate was used as an internal standard (chemical shift = 2.5 ppm at pH ~8).

Phosphate quantification of isolated WTA. To investigate the differences in phosphate content on WTA from each mutant, phosphate was quantified using a colorimetric method based on the color change resulting from formation of a phosphomolybdate complex.^{55,56} The isolated WTA from each strain was washed by placing 200 μ L of WTA sample in a test tube with 1 mL of 10% Mg(NO₃)₂·6H₂O in ethanol. The mixture was evaporated to dryness over a strong flame with constant shaking until brown fumes disappeared and a white powder was left. Once cooled to room temperature, 1 mL of 1 M HCl was added to the test tubes, which were then placed in a boiling water bath for 15 min to hydrolyze inorganic phosphate. A 1 : 1 mixture of 10% ascorbic acid and 0.42% ammonium molybdate in 1 M



H₂SO₄ was used to complex inorganic phosphate from WTA during incubation for 20 min at 45 °C. The absorbances of reaction mixtures were measured at 820 nm to quantify the amount of phosphate against a calibration curve constructed from potassium phosphate standards.

Isolation and characterization of bacterial sacculi

Isolation of bacterial sacculi. Bacterial cultures were grown in either LB media for solid-state ³¹P-NMR experiments or ¹³C- and ¹⁵N-labeled minimal (M9L) media (see ESI† for composition, Tables S8 and S9†) for solid-state ¹³C-NMR experiments. Details of the bacterial culture for isolation of bacterial sacculi are provided in the ESI.†

Solid-state NMR of bacterial sacculi. Isolated WTA were analyzed by solid-state ³¹P and ¹³C-NMR spectroscopy. Lyophilized sacculi samples (1 mg for ³¹P and 4 mg for ¹³C-NMR experiments) were packed into inserts by centrifugation. A solution of 0.05 M HEPES buffered to pH 7.4 was prepared. Before collecting spectra, 1 mL of this solution alone, with nanoparticles, or with bPEI polymer was added to the lyophilized sample. For samples containing both sacculi and nanoparticles, a total volume of 1 mL contained both 0.05 M HEPES, pH 7.4 and 34 nM bPEI-AuNPs. For samples containing both sacculi and free bPEI, a total volume of 1 mL contained both 0.05 M HEPES, pH 7.4 and free bPEI equivalent to amount of free polymer in 34 nM of nanoparticles. This was done at least 24 h prior to sample collection. The method to quantify the free polymer is described in the ESI.†

Following thorough sonication and vortexing, samples were sedimented for 30 min at 25 000g in a tabletop centrifuge at room temperature (Eppendorf, 5417R). After removing the supernatant, samples were packed into solid-state NMR inserts (Bruker, Kel-F). Inserts were placed into 4 mm diameter rotors (Bruker, Kel-F) and spun at 10 000 Hz during data collection. We collected 17 920 scans for each ³¹P spectrum and 512 scans for each ¹³C spectrum. The maximum distance between ¹³C nuclei sensed through space was roughly 20 Å.⁵⁷ As a control experiment, 34 nM bPEI-AuNPs were analyzed without sacculi by ¹³C-NMR. No signal was produced from these samples over the spectral acquisition time. We referenced ¹³C-NMR spectra to an external adamantane sample (chemical shifts of 38.5 and 29.5 ppm).⁵⁸ For ³¹P-NMR spectra, hexamethylphosphoramide was used as an internal reference (chemical shift at 29.97 ppm). All spectra were processed in MestReNova, including baseline correction and curve smoothing. ¹³C-NMR spectra were normalized by scaling the peak at ~20 ppm to 100 with respect to intensity.

T₁ relaxation experiments. Temperature-dependent ¹³C longitudinal T₁ relaxation time constants were measured by inversion-recovery after direct excitation of ¹³C nuclei using a Bruker Avance III 500 spectrometer equipped with a Doty 4 mm MAS probe with variable temperature capabilities. Nine interscan delay times were used and ranged from 0.0001 to 8 s. Sample temperatures were set to 30, 18, 12, 6 and -8 °C following temperature calibration described below. Acquisition time (aq) was 30 ms, and 64 scans per experiment were taken.

Sample temperatures were calibrated following a published method.⁵⁹ The chemical shifts of K⁷⁹Br were measured over a range of temperature settings, and the actual sample temperatures (T_{actual}) were calculated based on the expected relationship between T_{actual} and the measured ⁷⁹Br chemical shift: (K). The reference chemical shift was obtained at ambient temperature of 22 °C at a magic angle spinning frequency of 3 kHz, decoupler off (PLW2 = PLW12 = 0 W), repetition rate d1 = 5 s. Parameters were then changed to the experimental values of a spinning frequency of 8 kHz, ¹H decoupler strength = 104 kHz (pw90 = 2.4 μs at 200 W), aq = 30 ms, d1 = 15 s. Additional temperature points were then set, and the actual sample temperatures were obtained from the chemical shift difference from the reference.⁵⁹ One measurement was taken at each temperature and interscan delay time.

Conflicts of interest

There are no conflicts to declare.

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