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The Key Role of Peptidoglycan in the Opsonization of Staphylococcus Aureus

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ABSTRACT In an effort to determine the staphylococcal cell surface component(s) of importance in opsonization, cell walls (peptidoglycan and teichoic acid) and peptidoglycan were isolated from Staphylococcus aureus strain H grown in [3H]glycine-containing broth. After incubation of the cell walls and peptidoglycan with various opsonic sources, uptake by human polymorphonuclear leukocytes was measured. The opsonic requirements for phagocytosis of cell walls and peptidoglycan were found to be similar to those of intact bacteria. Removal of teichoic acid from the cell wall did not affect opsonization. Likewise, a teichoic acid-deficient mutant strain of S. aureus H was opsonized in a manner similar to that of the parent strain. Immunoglobulin G functioned as the major heat-stable opsonic factor and both the classical and alternative pathways participated in opsonization. Kinetic studies revealed that opsonization of peptidoglycan, as well as C3–C9 consumption by peptidoglycan, proceeded at a slower rate via the alternative pathway (C2-deficient serum) than when the classical pathway was present (normal serum). The ability of peptidoglycan to activate C3–C9 was significantly reduced when normal and C2-deficient sera were preabsorbed with peptidoglycan at 2°C suggesting that antibodies to peptidoglycan may be involved in activation of both the classical and alternative complement pathways. Thus, peptidoglycan appears to be the key cell wall component involved in staphylococcal opsonization, and it is suggested that host response to peptidoglycan, a major cell wall component of most gram-positive bacteria, may be related to the development of “natural immunity” to this group of microorganisms.

INTRODUCTION

In recent years increased attention has focused on the biological significance of the cell walls of gram-positive bacteria. In particular, peptidoglycan, a major component of the cell wall of most gram-positive bacteria, has been shown to play a significant role in determining the nature of host response to this group of bacteria (1, 2). In the case of Staphylococcus aureus, this cell wall component has been demonstrated to possess endotoxin-like properties (3), to activate the serum complement (C) system (4), and to elicit a cell-mediated immune response (5, 6). Few studies, however, have examined the role of the cell wall components of S. aureus in the process of opsonization, and results have been conflicting regarding the relative importance of these components in this process (7–12).

In this investigation the role of the two major components of the S. aureus cell wall—the peptidoglycan and teichoic acid—in the process of opsonization was studied by measuring the uptake of radioactively labeled bacteria and purified cell wall components by human polymorphonuclear (PMN) leukocytes. The nature of the opsonic factors involved in this process and the kinetics of C activation were examined. Data is presented which demonstrates that opsonization by both heat-stable (IgG) and heat-labile serum factors is mediated by cell wall peptidoglycan. Peptidoglycan was found capable of activating both the classical and alternative pathways of the C system. It is proposed that C1 may play a role in activation of the alternative complement pathway by peptidoglycan and that activation of both the classical and alternative pathways is initiated by antigen-antibody complex formation.

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1 Abbreviations used in this paper: CH₅₀, total hemolytic complement; HBSS, Hanks' balanced salt solution with 1% gelatin; PBS, phosphate-buffered saline; PMN, polymorphonuclear.
METHODS

Bacterial strains

*S. aureus* strain H, chosen because of its well-established cell wall structure (13), was kindly provided by Sir James Baddiley's laboratory, University of Newcastle-upon-Tyne, England. *S. aureus* HSm* (a spontaneous streptomycin-resistant isolate of strain H) and a teichoic acid-deficient mutant, strain 52A5 (14), were generously provided by J. T. Park, Tufts University, Boston, Mass. The cell wall of strain 52A5 appears to be composed almost entirely of peptidoglycan (15). The organisms were maintained on Trypticase Soy Agar (Difco Laboratories, Detroit, Mich.) slants.

Cultivation and radioactive labeling of bacteria

Bacteria were grown in peptone yeast extract broth (16) with aeration on an orbital shaker at 37°C. For phagocytosis studies of intact bacteria, the organisms were labeled by inoculating several colonies of bacteria into 20 ml peptone yeast extract broth containing 40 μCi [2-3H]glucose (sp act 5–15 Ci/mmole, New England Nuclear, Boston, Mass.). After an 18-h incubation at 37°C, the bacteria were washed three times in phosphate-buffered saline (PBS), pH 7.4, and resuspended in PBS to a final concentration of 1 × 10⁹ colony-forming units/ml using a spectrophotometric method confirmed by a plate colony count. For preparing radioactively labeled cell walls, organisms were grown in peptone yeast extract broth, 2 liters containing [2-3H]glucose, 0.4 μCi/ml, and were harvested in late exponential growth phase (OD₆₅₀ nm 1.5–2.0, Spectronic 20, Bausch & Lomb Scientific Optical Products Div., Rochester, N.Y.). Large scale growth of unlabeled bacteria (26 liters) was achieved in a New Brunswick Fermentor (New Brunswick Scientific Co., Inc., New Brunswick, N.J.) at 37°C with forced aeration (5 liters/min) using antifoam (Anti-foam aerosol spray, Sigma Chemical Co., St. Louis, Mo.). 2 liters of a 15-h culture was used as an inoculum, and the bacteria were grown for 6–7 h and harvested at an OD₆₅₀ of 1.0–1.5. The cultures were harvested in a continuous flow centrifuge (Beckman J-21) using a JCF-2 rotor (Beckman Instruments Inc., Spinco Div., Palo Alto, Calif.) operating at 2,500 g with a flow rate of 250 ml/min. The organisms were washed once in cold distilled water and then frozen (−15°C) overnight before disruption. The yield was 80 g wet weight of cells from 26 liters of culture.

Isolation of cell walls, peptidoglycan, and teichoic acid

Unless otherwise indicated, all operations were carried out at 4°C and all centrifugations were at 16,319 g for 10 min at 4°C. Washed organisms in distilled water were mixed with glass beads (0.10–0.11 mm, VWR Scientific Div., UNIVAR, San Francisco, Calif.), and dispersed by agitation in a Vibrogen Cell Mill (R.H.O. Scientific Inc., Conmack, N. Y.) for 20 min at maximum speed. The glass beads were removed by filtration on a coarse glass sinter and the filtrate was centrifuged to deposit the white cell walls overlaying yellower, unbroken organisms. Care was taken to avoid resuspending unbroken organisms. The crude cell walls were washed once in distilled water, resuspended in 200 ml 2% sodium dodecyl sulphate, and then stirred overnight at room temperature to inactive autolytic enzymes and to aid in stripping of membrane components. The cell walls were then washed four times in water at room temperature before being resuspended in 200 ml 0.05 M Tris-HCl containing MgCl₂ (5 mM) plus 2.5 μg/ml each of RNase (EC 3.1.4.5, 2,588 U/mg) and DNase (EC 3.1.4.22, 2,500 U/mg) (each type 1 from Worthington Biochemical Corp., Freehold, N.J.) and incubated at 37°C for 1 h with gentle shaking (100 rpm). To remove proteinaceous material including protein A (17), 200 μg/ml trypsin was added (EC 3.4.14.4, 189 U/ml, Worthington Biochemical Corp.), and the incubation continued for 4 h. The walls were washed once in water and then stirred with an equal volume of 80% phenol for 30 min at room temperature to remove lipoteichoic acid (18). After extraction, the walls were washed six times in water and then lyophilized. 1,680 mg of cell wall was recovered from 26 liters of culture.

Cell walls (1,260 mg) suspended in 75 ml water were mixed with 75 ml of 20% (wt/vol) TCA and were incubated at 60°C for 90 min to remove teichoic acid (19). The suspension was centrifuged to yield peptidoglycan which was washed four times with water before being lyophilized (yield 1,067 mg, 35.5% weight of wall removed). Teichoic acid was recovered from the supemate by adding 750 ml cold EtOH and allowing to stand at 4°C overnight. The precipitate recovered by centrifugation was resuspended in 14 ml 10% TCA and allowed to precipitate again overnight at 4°C with 70 ml EtOH. The product was washed twice with acetone and allowed to air dry.

Typically, cell walls were labeled to a specific activity of ≈ 5 × 10⁶ cpm/mg dry weight. The hot TCA treatment resulted in about 10% loss of counts indicating little solubilization of glycine bridges occurred (20) and the specific activity was roughly double that of the cell walls. Preparations were stored frozen at −15°C at a concentration of 5–10 mg dry weight/ml, and brief treatment in a sonic bath (Branson Sonic Power Co., Danbury, Conn.) was used to help attain an even suspension before each opsonization procedure.

Chemical characterization of cell walls, peptidoglycan, and teichoic acid

Quantitative examination. Carefully weighed cell walls, peptidoglycan, and teichoic acid were hydrolyzed in sealed tubes with 6 M HCl for 18 h at 105°C. Acid was removed under vacuum over anhydrous CaCl₂ in the presence of NaOH pellets; the dried residues were twice moistened with 0.5 ml water and redried. Amino acids and amino sugars were estimated on an amino acid analyzer (Beckman model 120B Amino Acid Analyzer, Beckman Instruments, Fullerton, Calif. Inc.). Phosphorus was estimated by the method of Fiske and Subba Row as described by Leloir and Gardini (21). Total fatty acids in 10-mg amounts of preparation were determined as described by Wilkinson (22).

Qualitative examination. Teichoic acid (5 mg) was hydrolyzed in 2 M HCl for 3 h at 105°C in sealed tubes; acid was removed as described above. Part of the hydrolysate was treated with alkaline phosphatase (EC 3.1.3.1, 1 mg, type 1, calf intestinal mucosa, Sigma Chemical Co.) in 0.025 M NH₄HCO₃ containing 0.005 M NH₄OH, pH 8.3, overnight at 37°C. Alkaline phosphatase-treated and untreated hydrolysates were examined by descending chromatography on Whatman no. 1 paper in the solvent systems 6 propan-1-OL:3.088 NH₃:1 H₂O (by volume) and 6 butan-1-OL:4 pyridine: 3 H₂O (by volume) (23). Reducing compounds were detected with the alkaline AgNO₃ reagent (24) and compounds with amino groups by ninhydrin treatment (0.25% wt/vol in acetone plus heating at 100°C for 5 min).
**Opsonic sources**

Serum was pooled from five healthy donors who denied previous staphylococcal infection (normal serum). To block C1, normal serum was chelated with ethylene glycoltetraacetic acid in the presence of MgCl2 (MgEGTA) in a 10-M to 10-mM concentration with respect to serum (25). C2-deficient serum was obtained from a patient with genetically determined complete absence of C2 and normal levels of all other C components. Serum was stored in 0.5-ml aliquots at −70°C. Heat-inactivated serum was prepared by heating thawed aliquots of serum at 56°C for 30 min. Purified human IgG was provided by Dr. D. Roos, Central Laboratory of the Blood Transfusion Service, Amsterdam, Netherlands. Serum and IgG were diluted to specified concentrations with Hank's balanced salt solution containing 1% gelatin (HBSS) just before use. Calcium- and magnesium-free HBSS was used to dilute MgEGTA-chelated serum.

**Opsonization procedure**

According to Chatterjee (26), $1 \times 10^8$ whole organisms yield 25 μg of cell walls. Consequently, 0.1-ml aliquots of intact bacteria (1 $\times 10^8$ colony-forming units) and 25 μg of cell walls were used in all opsonization experiments. Studies using 25 and 12.5 μg of peptidoglycan yielded similar findings; results were reported using 25 μg of peptidoglycan. Each aliquot was incubated at 37°C in a 12 $\times$ 75-mm plastic tube, BioQuest, BBL & Falcon Products, Becton, Dickinson & Co., Cockeysville, Md., containing 1.0 ml of the indicated opsonic source for specified times followed by centrifuging at 4,000 g for 15 min (4°C). The supernate was discarded and the pellet resuspended in 0.5 ml HBSS.

**PMN leukocytes**

Blood from healthy donors was drawn into heparinized syringes (10 U heparin/ml blood). PMN leukocytes were prepared using a method modified from Büyum (27) in which the blood was centrifuged for 15 min at 200 g, and the plasma-platelet layer was removed with a Pasteur pipette. The leukocyte layer was then withdrawn, placed in plastic tubes, and then diluted with two parts Spinner’s Minimal Essential Medium (IX, Grand Island Biological Co., Grand Island, N.Y.) 8 ml of this suspension was carefully layered onto 3 ml Ficoll-Isoaque (lymphocyte separation medium, Litton Bionetics, Kensington, Md.). After centrifuging at 200 g for 30 min the mononuclear cell layer was removed with a Pasteur pipette. To obtain pure PMN leukocytes, the remaining medium was withdrawn and discarded, and 0.87% NH4Cl in sterile water was added to the tube containing the erythrocyte-PMN leukocyte pellet which was then rotated for 10 min at room temperature. After centrifuging for 10 min at 160 g, the PMN leukocyte pellet was washed an additional two times with 0.87% NH4Cl to lyse the remaining erythrocytes. The PMN leukocytes were resuspended finally in HBSS and total and differential leukocyte counts were performed. The final leukocyte pellets were adjusted to a concentration of $10^7$ PMN leukocytes/ml HBSS. Contamination by mononuclear cells was evaluated by Wright’s stained smears and generally did not exceed 1%.

**Phagocytosis mixtures and assays**

To each 0.5 ml suspension of opsonized radioactively labeled intact bacteria, cell walls, and peptidoglycan, 0.5 ml of PMN leukocyte suspension was added. These mixtures were incubated at 37°C in an incubator shaker (New Brunswick Scientific Co., Inc.) at 250 rpm. PMN leukocyte uptake of bacteria, cell walls, and peptidoglycan was quantitated using a modification of a previously described method (28). Briefly, leukocyte-associated radioactivity was determined by taking duplicate 100-μl samples from each phagocytosis mixture at specified time intervals, placing the samples in 3 ml PBS, pH 7.4, in polypropylene vials (Bio-vials, Beckman Instruments Inc.) and washing the leukocytes three times in PBS by differential centrifugation at 160 g. Final pellets were resuspended in 3 ml scintillation liquid (toluene containing fluorarloy [TLA, Beckman Instruments, Inc.] and 20% Biosolv-3 [Beckman Instruments, Inc.]) and counted in a liquid scintillation counter (Beckman LS-250). To determine total radioactivity (representing both leukocyte-associated and nonleukocyte-associated bacteria, cell walls, or peptidoglycan), duplicate 100-μl samples were placed in 3 ml PBS at the end of the assay period and centrifuged at 4,000 g for 15 min. Supernates were discarded, pellets resuspended in scintillation liquid, and samples counted in the scintillation counter. An average of duplicate values was used in all calculations. PMN leukocyte uptake of radioactively labeled bacteria, cell walls, and peptidoglycan at given sampling times was calculated using the formula: % uptake = (cpm in leukocyte pellet)/(total cpm) $\times$ 100.

To study opsonization of soluble peptidoglycan fragments, purified peptidoglycan was digested with the mureinolytic enzyme, lysostaphin. 0.1 mg (0.1 ml) peptidoglycan was incubated in 0.9 ml PBS containing 10 μg/ml lysostaphin (Schwarz-Mann Div., Becton, Dickinson & Co., Orangeburg, N.Y.) for 60 min at 37°C followed by centrifuging at 10,000 g for 30 min. Supernates were then heated in a boiling water bath for 15 min to inactivate lysostaphin. 50-μl aliquots of the supernate (peptidoglycan fragments) were then incubated with opsonic sources for 15 min. PMN leukocytes were added, and uptake was determined as described above. Total radioactivity in these mixtures was determined by suspending 100-μl samples directly into scintillation liquid. A control mixture of 0.5 mg peptidoglycan was incubated in PBS for 60 min; after centrifuging, the supernate was discarded and the pellet resuspended in 1.0 ml PBS containing 10 μg/ml lysostaphin which had been previously heat-inactivated. This mixture was then heated in a boiling water bath after which 50-μl aliquots were removed for opsonization and phagocytosis studies.

**Measurement of C activation by purified peptidoglycan**

0.1-ml aliquots of peptidoglycan, at specified concentrations, and of water (control) were added to 12 $\times$ 75-mm plastic tubes (BioQuest, BBL & Falcon Products) containing 1.0 ml normal serum. After a 30-min incubation at 37°C, the tubes were centrifuged at 4,000 g for 15 min (4°C) and supernates were stored at −70°C before measurement of total hemolytic complement (CH50).

To study the kinetics of C activation in normal serum and in C2-deficient serum, 0.1-ml (1.0 mg) aliquots of peptidoglycan were added to 1.0-ml samples of serum followed by incubating at 37°C for indicated time intervals. Control mixtures containing 0.1 ml distilled water were incubated simultaneously and removed at the same time intervals. After centrifuging at 4,000 g for 15 min (4°C), the supernates were stored at −70°C before measurement of C3–C9.

The effect of preabsorption of normal and C2-deficient sera with peptidoglycan on the subsequent ability of peptidoglycan...
to activate the serum C system was studied by incubating 1.0-ml samples of serum three times for 60 min at 2°C with either 0.1-ml (1.0 ng) aliquots of peptidoglycan or 0.1-ml aliquots of distilled water (nonabsorbed serum). After each 60-min absorption period, samples were centrifuged for 15 min at 4,000 g (2°C) and the supernatant was added to the next precooled tube containing either peptidoglycan or water. After the final centrifugation, the serum samples were added to tubes containing 0.1-ml aliquots of peptidoglycan, at indicated concentrations, and were then incubated at 37°C for specified time intervals. Control tubes containing absorbed and nonabsorbed serum and 0.1 ml distilled water were likewise incubated at 37°C. After incubation, samples were centrifuged at 4,000 g for 15 min (2°C) and the supernates stored at −70°C before C3–C9 determination.

Hemolytic titrations of C were performed using sheep erythrocytes in acid citrate dextrose (Willer Laboratories, Stillwater, Minn.). Intermediate cell preparation (CT4) and the assays of CH50 and C3–C9 were performed as previously described (29–32). For CH50, the optical density was read at 541 nm and for C3–C9 titration at 412 nm. Results were expressed as 50% lysis, and titrations were corrected for day-to-day variations by including a known internal standard human serum.

**Electron microscopy**

Cell walls and peptidoglycan were examined by a Siemens 102 electron microscope (Siemens Medical/Industrial Groups, Istlin, N. J.) using previously described methods (33). In addition, 0.1-ml aliquots of peptidoglycan (1 mg) were incubated in plastic tubes (BioQuest, BBL & Falcon Products) containing 1.0 ml 10% normal serum, 10% heat-inactivated serum, and HBSS for 60 min. After centrifuging for 15 min at 4,000 g, supernates were discarded and 0.5 ml HBSS was added to each tube. Leukocyte suspensions (0.5 ml) were then added and the mixtures incubated for 15 min at 37°C. The leukocytes were washed three times with cold PBS, fixed in 1.5% glutaraldehyde in PBS, and then thin sections were examined.

**RESULTS**

**Chemistry of the cell walls, peptidoglycan, and teichoic acid**

Table I shows the quantitative analysis for the cell wall preparations used in this study. It can be seen that amino acids other than those normally considered to be part of the staphylococcal peptidoglycan were either not detected or present in only small amounts. The lack of the full range of protein amino acids indicates little contamination of the cell walls with proteinaceous materials including protein A. On the basis of the analyses presented here, and knowledge of the S. aureus peptidoglycan structure (19, 34), it can be seen that the average subunit of the peptidoglycan contains isoglutamine, 1.0; lysine, 1.0; alanine, 2.0; glycine, 4.0; serine, 0.1; N-acetylmuramic acid, 1.0; N-acetylglicosamine, 1.0; O-acetyl, 0.6; and the formula weight is 1,138. Using the lysine value it can be seen that 50.5% of the dry weight of the wall is accounted for by the peptidoglycan. For isolated peptidoglycan, 92.4% of the dry weight of the preparation is accountable for as peptidoglycan components. Assuming that ribitol phosphate and N-acetylglucosamine are equimolar then one can take ribitol, 1.0; phosphate, 1.0; N-acetylglucosamine, 1.0; alanine, 0.16 as the average teichoic acid sub-unit (see below) and this gives a formula weight of 410. Thus, 41% of the dry weight of the cell wall is present as teichoic acid plus 50.5% as peptidoglycan thereby accounting for 91.5% of the dry weight of the cell wall. It is likely that metal counterions to negatively charged groups in the teichoic acid and peptidoglycan make up a considerable portion of the unaccounted weight of these preparations (35). 82% of the teichoic acid preparation is accountable for as teichoic acid components.

<table>
<thead>
<tr>
<th>Component</th>
<th>Cell walls</th>
<th>Peptidoglycan</th>
<th>Teichoic acid</th>
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</thead>
<tbody>
<tr>
<td>Muramic acid</td>
<td>180.4</td>
<td>280</td>
<td>Trace</td>
</tr>
<tr>
<td>Glucosamine</td>
<td>682</td>
<td>412</td>
<td>1,560</td>
</tr>
<tr>
<td>Glutamic acid</td>
<td>277</td>
<td>668</td>
<td>24</td>
</tr>
<tr>
<td>Lysine</td>
<td>444</td>
<td>812</td>
<td>48</td>
</tr>
<tr>
<td>Alanine</td>
<td>849</td>
<td>1,588</td>
<td>310</td>
</tr>
<tr>
<td>Glycine</td>
<td>1,826</td>
<td>3,460</td>
<td>120</td>
</tr>
<tr>
<td>Serine</td>
<td>48</td>
<td>68</td>
<td>—</td>
</tr>
<tr>
<td>Phosphate</td>
<td>1,000</td>
<td>84</td>
<td>2,000</td>
</tr>
<tr>
<td>Aspartic acid</td>
<td>26.4</td>
<td>24</td>
<td>—</td>
</tr>
<tr>
<td>Threonine</td>
<td>8.8</td>
<td>8</td>
<td>—</td>
</tr>
<tr>
<td>Valine</td>
<td>8.8</td>
<td>8</td>
<td>—</td>
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<tr>
<td>Isoleucine</td>
<td>8.8</td>
<td>8</td>
<td>—</td>
</tr>
<tr>
<td>Leucine</td>
<td>8.8</td>
<td>8</td>
<td>—</td>
</tr>
<tr>
<td>Histidine</td>
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<tr>
<td>Arginine</td>
<td>8.8</td>
<td>16</td>
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</tr>
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</table>

*Cell walls, peptidoglycan, and teichoic acid were isolated from S. aureus strain H.*

Additional evidence that the peptidoglycan is free of teichoic acid is the finding that 95% of the cell wall phosphorus was removed by the hot TCA extraction and the ratio of the glucosamine to muramic acid changed from an excess to being almost equimolar.

Qualitative examination of the teichoic acid hydrolysate revealed components with chromatographic mobilities characteristic of glucosamine, ribitol, alanine, glucosaminol phosphate, ribitol di- and monophosphate, and glucosaminol ribitol. Amino acid analyses of the teichoic acid revealed glucosamine and alanine to be the major components. The absence of muramic acid and the low amounts of other amino acids show that the teichoic acid fraction does not contain major amounts of peptidoglycan. In an effort to determine whether the cell wall and peptidoglycan preparations were contaminated with lipoteichoic acid, the total fatty acid composition of the preparations was determined. Fatty acids are markers of cell membrane

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lipids and/or lipoteichoic acid. This was compared with the fatty acid composition of "crude cell walls" that had not been treated with sodium dodecyl sulfate, phenol, or enzymes but had simply been washed six times in water. Five major fatty acid methyl ester peaks were obtained from the crude cell walls by gas-liquid

Figure 1: Electron micrographs of cell walls, (a) \( \times 20,265 \), (b) \( \times 100,000 \) and peptidoglycan, (c) \( \times 20,265 \), (d) \( \times 100,000 \) isolated from S. aureus H.
chromatography. Chromatography of the fatty acid methyl ester fraction from an equal weight of the purified cell walls used in this study revealed only one very small “peak” on the trace that was difficult to estimate. Similar examination of the peptidoglycan revealed two extremely small peaks. Summation of the total detector response revealed that the cell walls and peptidoglycan had at least 97.4 and 98.7%, respectively, less fatty acids than the crude cell walls. The cell wall and peptidoglycan preparations were contaminated with fatty acids at levels < 1 nmol/mg. These results, plus the fact that we can account for most of the weight of the preparations in known components as well as the proven value of sodium dodecyl sulfate and phenol extractions in removing lipoteichoic acid (18, 36), indicate the preparations are not significantly contaminated with lipoteichoic acid.

Thus, the qualitative and quantitative analyses indicate that the cell walls (composed of peptidoglycan and teichoic acid) and peptidoglycan used in this investigation are highly purified and chemically consistent with the observations of other investigators (15, 34).

Electron micrographs of glutaraldehyde–osmium tetroxide-fixed, embedded sections of cell walls and peptidoglycan are shown in Fig. 1. The preparations are quite homogenous and devoid of whole organisms and debris. Both cell walls and peptidoglycan reveal a characteristic trilamellar appearance and septa (37). The morphologic appearance of the peptidoglycan is somewhat collapsed and folded in comparison to the cell walls. In general, the peptidoglycan is thinner in sectioned material than the cell walls, a finding in agreement with previous studies (37, 38); detailed morphometric measurements, however, were not performed.

Phagocytosis of S. aureus H and of isolated cell walls and peptidoglycan

Kinetics of uptake by PMN leukocytes. Intact S. aureus H, isolated cell walls (peptidoglycan with covalently linked teichoic acid) and purified peptidoglycan were incubated with 1% normal serum, 1% heat-inactivated serum, and HBSS for 15 min before adding PMN leukocytes. As shown in Fig. 2, uptake of intact bacteria, cell walls, and peptidoglycan was similar at 3, 10, and 15 min. At 15 min, ≈70% of each of the three types of “particles” opsonized with normal serum were leukocyte-associated compared with ≈25% and virtually no uptake of particles which had been incubated with heat-inactivated serum and HBSS, respectively. These results suggested that removal of teichoic acid from the cell wall preparations did not significantly alter opsonization and phagocytosis by PMN leukocytes.

To determine whether leukocyte-associated radioactivity represented attached and/or ingested peptidoglycan, electron microscope examination was performed (Fig. 3). After opsonization with normal serum, peptidoglycan was present within phagocytic vacuoles in the leukocytes and significant degranulation has occurred. When preincubated with HBSS, essentially no peptidoglycan was found within the leukocytes. Opsonization with heat-inactivated serum resulted in less ingestion than when normal serum was used as an opsonic source but in greater ingestion than when no opsonin was present (HBSS).

Opsonic titer of normal and heat-inactivated sera. When 1% serum was used as an opsonic source PMN leukocyte uptake of cell walls (peptidoglycan with teichoic acid) and peptidoglycan was found to be similar. This result, as stated above, suggested that removal of teichoic acid from the cell wall did not significantly affect opsonization. To test the possibility that removal of teichoic acid might change the titer of serum capable of effective opsonization, intact bacteria, cell walls, and peptidoglycan were incubated for 15 min with varying concentrations of normal serum and heat-inactivated serum before adding leukocytes and determining uptake at 15 min (Fig. 4). Phagocytosis of all three particles was significantly reduced and to the same extent when 0.1% normal serum and when 1.0% heat-inactivated serum were used as opsonic sources thereby demonstrating that the opsonic titer was not influenced by the removal of teichoic acid from the cell wall. When purified human IgG, in concentrations of 2.5, 1.0, 0.1, and 0.01 mg/ml, was used as an opsonic source, uptake was found to be similar to that of 5.0, 1.0, 0.1, and 0.01% heat-inactivated serum, respectively (data not shown). This suggested that IgG was the major opsonic factor in heat-inactivated serum and implies
that purified human IgG contains antibodies against peptidoglycan.

*Phagocytosis of S. aureus HSm* and 52A5. Although the results of the above experiments suggested that peptidoglycan was the major component involved in opsonization by normal and heat-inactivated sera,
and that cell wall teichoic acid played little or no role in this regard, electron microscope studies of thin sections of cell walls and peptidoglycan (Fig. 1) had indicated that these preparations differed morphologically as well as in their previously defined chemical compositions. Therefore, it seemed possible that removal of teichoic acid from the cell wall might have affected opsonization independently of the change brought about in chemical makeup. To investigate this possibility, opsonization of a teichoic acid-deficient mutant (52A5) was compared to that of its parent strain (HSm^6) (Fig. 5). As shown, phagocytosis of the teichoic acid-deficient mutant did not differ significantly from that of the parent strain when varying concentrations of normal and heat-inactivated sera were used as opsonic sources. This result provided further evidence that the teichoic acid component of the cell wall did not have a major influence on bacterial opsonization. After each experiment the typeability of the strains was checked with phage 52A (39). Teichoic acid forms part of the receptor site for this phage (14). The parent organism remained susceptible to this phage and strain 52A5 was not susceptible, thereby indicating that reversion of the 52A5 strain to a teichoic acid-containing form had not occurred.

The role of C1 and C2 in opsonization. To study the role of the classical C pathway in opsonization, C1 was blocked by chelating normal serum with MgEGTA, and opsonization in the absence of C2 was studied by using serum from a patient with complete C2 deficiency. Intact bacteria, cell walls, and peptidoglycan were incubated for 60 min with 1% concentrations of normal, heat-inactivated, MgEGTA-chelated, C2-deficient, and heat-inactivated C2-deficient sera before adding PMN leukocytes and measuring uptake at 15 min. Normal serum was the most effective opsonic source for intact bacteria, cell walls, and peptidoglycan, uptake being significantly greater than when either MgEGTA-

![Figure 4](image4.png)  
**FIGURE 4** PMN leukocyte uptake at 15 min of intact *S. aureus* H, isolated cell walls and purified peptidoglycan opsonized with varying concentrations of normal serum (A) and heat-inactivated serum (B). Results represent means of three experiments.

![Figure 5](image5.png)  
**FIGURE 5** PMN leukocyte uptake of *S. aureus* HSm^6 (parent strain) and 52A5 (teichoic acid-deficient mutant) preincubated with varying concentrations of (A) normal serum and (B) heat-inactivated serum before adding leukocytes and determining uptake at 15 min. Results represent means of three experiments.

![Figure 6](image6.png)  
**FIGURE 6** PMN leukocyte uptake of intact *S. aureus* H, isolated cell walls, and purified peptidoglycan preincubated for 60 min with normal serum (NS), MgEGTA-chelated serum (X), heat-inactivated serum (ΔS), C2-deficient serum (C2 def) and heat-inactivated C2-deficient serum (ΔC2 def) before adding leukocytes and determining uptake at 15 min. Results represent means of five experiments. Significance of differences in uptake comparing NS and X, P < 0.001 for intact bacteria, cell walls and peptidoglycan; comparing NS and C2 def, P < 0.025, 0.001, 0.001 for intact bacteria, cell walls and peptidoglycan respectively; comparing C2 def and ΔC2 def, P < 0.001 for intact bacteria, cell walls and peptidoglycan (t test for paired data).
Chelated or C2-deficient sera were used as opsonic sources (Fig. 6). Although MgEGTA-chelated serum was not significantly more opsonic than heat-inactivated serum, C2-deficient serum had a greater opsonic capacity than heat-inactivated C2-deficient serum.

These results suggested that opsonization could proceed via a heat-labile pathway other than the classical pathway when C2 was blocked but not when C1 was blocked. When the period of opsonization was limited to 15 min rather than 60 min, however, C2-deficient serum was no more effective than heat-inactivated C2-deficient serum as an opsonic source, opsonization with the other serum sources not being affected by the shorter opsonization period (data not presented). This finding indicated that opsonization via the second heat-labile pathway proceeded more slowly than when opsonization was carried out in the presence of the classical pathway.

Effect of lysostaphin treatment of peptidoglycan on opsonization. To determine whether relatively small, soluble fragments of peptidoglycan could be opsonized by normal and heat-inactivated sera, peptidoglycan was treated with lysostaphin and the fragments incubated with 10% normal serum, 10% heat-inactivated serum, and HBSS before adding PMN leukocytes and measuring leukocyte-associated radioactivity at 3, 10, and 15 min. Whereas there was uptake of intact peptidoglycan in the control mixtures similar to that observed in earlier experiments (Fig. 2), no leukocyte-associated radioactivity was recorded when peptidoglycan fragments were studied. These results suggested that for effective opsonization a certain particle size of the peptidoglycan was required.

C activation by peptidoglycan

Effect of peptidoglycan concentration on C activation in normal serum. As results from the above experiments indicated that the peptidoglycan of S. aureus H was the major C-activating component of the cell wall, additional studies were performed in which consumption of C by peptidoglycan was measured. When purified peptidoglycan was incubated for 30 min with normal serum in concentrations of from 5.0 to 0.01 mg/ml serum, a clear dose-response relationship was established (Table II). When concentrations of 1.0–5.0 mg were used, CH50 activity was not detectable.

Kinetics of C activation in normal and C2-deficient sera. The ability of peptidoglycan to activate C in the presence and in the absence of C2 was studied by incubating normal and C2-deficient sera with peptidoglycan, at a concentration of 1.0 mg/ml, at 37°C. Samples were removed at 1, 5, 15, 60, and 120-min intervals and centrifuged at 2°C. The hemolytic titer of C3–C9 was measured in the supernatant serum, and values were compared with C3–C9 levels measured in simultaneous controls (normal and C2-deficient sera incubated with water for the same time intervals). Consumption of C3–C9 in the peptidoglycan-treated samples was expressed as a percentage relative to the C3–C9 levels in the controls (Fig. 7). As shown, activation of complement occurred both in the presence (normal serum) and in the absence (C2-deficient serum) of an intact classical pathway; however, the rate of activation in C2-deficient serum was significantly slower than in normal serum (Fig. 7).

TABLE II

<table>
<thead>
<tr>
<th>Peptidoglycan mg</th>
<th>CH50</th>
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</thead>
<tbody>
<tr>
<td>5.0</td>
<td>&lt;101</td>
</tr>
<tr>
<td>2.5</td>
<td>&lt;10</td>
</tr>
<tr>
<td>1.0</td>
<td>&lt;10</td>
</tr>
<tr>
<td>0.5</td>
<td>13</td>
</tr>
<tr>
<td>0.1</td>
<td>25</td>
</tr>
<tr>
<td>0.01</td>
<td>33</td>
</tr>
<tr>
<td>Control</td>
<td>31</td>
</tr>
</tbody>
</table>

* 1-ml aliquots of normal serum were incubated with equal volumes of purified peptidoglycan, in concentrations from 5.0 to 0.01 mg, and of water (control) for 30 min at 37°C. After centrifuging, CH50 was measured in the supernatant serum. † <10 = not detectable.

Figure 7. Kinetics of C activation by purified peptidoglycan in normal and C2-deficient sera. 1-ml aliquots of normal and C2-deficient sera were incubated at 37°C with equal volumes of water (controls) and peptidoglycan (1.0 mg) for 1, 5, 15, 60, and 120 min. Consumption of C3–C9 in the peptidoglycan-treated serum samples is expressed as a percent of the simultaneous control samples.

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Effect of preabsorption of serum with peptidoglycan on subsequent C activation. The ability of peptidoglycan to activate the C system in normal serum was found to be significantly impaired when serum was preabsorbed with peptidoglycan at 2°C. Although there was 92, 89, and 37% consumption of C3–C9 in control serum after a 30-min incubation with 5.0, 1.0, and 0.1 mg peptidoglycan, respectively, there was <25% C3–C9 consumption in serum that had been preabsorbed with peptidoglycan. Likewise, preabsorption of C2-deficient serum with peptidoglycan at 2°C had an inhibitory effect on the subsequent capacity of peptidoglycan to activate the C system. After 30-min and 60-min incubations with 1.0 mg peptidoglycan, there was 9 and 44% C3–C9 consumption in nonabsorbed C2-deficient serum compared with 0 and 27% activation in C2-deficient serum which had been preabsorbed with peptidoglycan. The absorption procedure itself did not significantly influence C3–C9 levels (after absorption, C3–C9 levels were 422 and 540 U/ml in normal and C2-deficient sera compared with control values of 437 and 500 U/ml, respectively). These data suggested that peptidoglycan activation of the C system, via both the classical and alternative pathways, depended upon the presence of an absorbable, noncomplement-related serum factor.

DISCUSSION

The cell wall of most S. aureus strains is composed of three major constituents—peptidoglycan, teichoic acid, and protein A. The peptidoglycan component is a linear polymer of repeating β-1–4-linked N-acetylmuramyl-N-acetylglucosamine and N-acetylmuramic acid residues. Attached to the carboxyl group of each N-acetylmuramic acid is a tetrapeptide cross-linked by a pentaglycine bridge to a neighboring peptide unit. Approximately 50% of the weight of the cell wall is contributed by peptidoglycan, and it is this component which provides the wall its basic rigidity (40). Cell wall teichoic acids, comprising ≈40% of cell wall weight, are charged polymers of ribitol–phosphate linked to muramic acid residues of the peptidoglycan (41). Protein A, contributing ≈5% to cell wall weight, is a protein with unique properties which, in general, interfere with the process of opsonization (42, 43). In this investigation, trypsin digestion was used to remove protein A and other cell wall proteins. Also lipoteichoic acid, which may be another component of the staphylococcal cell surface (18), was extracted, and the two major constituents of the cell wall—peptidoglycan and teichoic acid—were studied. The staphylococcal cell surface may be viewed as a mosaic of different domains where all of the major cell wall components, including peptidoglycan and teichoic acid, can be sufficiently exposed to interact with external factors.

The relative importance of the peptidoglycan and teichoic acid components of the S. aureus cell wall in the process of opsonization has been a matter of controversy. Evidence has previously been presented supporting the role of both teichoic acid (7–10) and peptidoglycan (11, 12) in this regard. These studies, however, have used indirect methods to assess opsonization, either by immunizing animals with cell wall components and measuring the protective effect of such immunization (8, 9) or by absorbing serum with cell wall components and determining the effect of such absorption on serum opsonic activity (7, 11, 12).

In this study, PMN leukocyte uptake of intact S. aureus H and of highly purified cell walls—composed of peptidoglycan and teichoic acid—and of isolated peptidoglycan was measured quantitatively using [3H]glycine-labeled bacteria, cell walls, and peptidoglycan. When normal human serum and heat-inactivated serum were used as opsonic sources, intact bacteria, cell walls, and peptidoglycan were taken up by PMN leukocytes in a similar manner. All three particles were most efficiently opsonized by normal serum, and uptake did not occur in the absence of serum factors. Electron microscope studies revealed that opsonized peptidoglycan was readily ingested by PMN leukocytes. Studies with purified IgG indicated that IgG was the major heat-stable serum opsonic factor, a finding reported by other investigators (44–46) for different staphylococcal strains.

As isolated cell walls (peptidoglycan and teichoic acid) and purified peptidoglycan were phagocytized in a comparable fashion, it appeared that the removal of teichoic acid from the cell wall did not significantly affect opsonization, thereby implying that the process of opsonization was primarily mediated by peptidoglycan. Electron microscope studies, however, had revealed that the cell wall and peptidoglycan preparations were not morphologically identical; in thin sections the peptidoglycan was, in general, more collapsed. Therefore, the possibility existed that this structural difference might influence opsonization in a manner independently of the well-defined chemical difference of these particles. To help clarify this possibility, opsonization of a teichoic acid-deficient mutant strain of S. aureus H was compared with that of the parent strain. The absence of teichoic acid in the cell wall did not alter opsonization by normal and heat-inactivated sera which added support to the hypothesis that teichoic acid played an insignificant role in bacterial opsonization.

The finding that there was no PMN leukocyte uptake of lysostaphin-degraded peptidoglycan which had been preincubated with normal and heat-inactivated sera suggested that a certain particle size is required for effective opsonization. Shayeegani and colleagues (11) reported similar findings using different methods and
enzymes. Grov and co-workers (47) have recently demonstrated that the inhibition of leukocyte migration by staphylococcal peptidoglycan is also dependent upon the size of the peptidoglycan fragment.

Other investigators have demonstrated that the classical (46, 48) and the alternative (46, 49, 50) C pathways are involved in staphylococcal opsonization. In an attempt to delineate which heat-labile serum factors participated in the opsonization of *S. aureus* H and of the peptidoglycan component, phagocytosis of intact bacteria, cell walls, and peptidoglycan was measured after opsonization with normal, MgEGTA-chelated, heat-inactivated, C2-deficient, and heat-inactivated C2-deficient sera. Opsonization of all three particles was optimal when normal serum was used as an opsonic source. In the absence of C2, opsonization was less effective, both the opsonic capacity and the rate of opsonization being less with C2-deficient serum than with normal serum; however, opsonization did proceed via a heat-labile system (the opsonic capacity of C2-deficient serum being significantly reduced by heat-inactivation). On the other hand, heat-labile opsonic activity was abolished in the absence of C1 (MgEGTA-chelated serum).

These results suggest that opsonization of intact *S. aureus* H and of peptidoglycan is most effective in the presence of an intact classical pathway. However, it appears that opsonization can proceed via another heat-labile pathway when C2 is blocked but not when C1 is blocked. Other investigators (51, 52) have demonstrated that C1 can play a role in the activation of the alternative complement pathway (the "C1-bypass" mechanism). From the results in this study, it appears that the C1-bypass may be operative in the opsonization of *S. aureus* H peptidoglycan by C2-deficient serum, and that blocking C1 thereby interferes with alternative pathway activation.

Activation of the serum C system by peptidoglycan was further studied by incubating peptidoglycan with normal serum and measuring C consumption. A dose-response relationship was established in which CH50 was not detectable after a 30-min incubation with peptidoglycan in concentrations of 1.0–5.0 mg/ml serum. The kinetics of C activation were then investigated using normal and C2-deficient sera. Although the rate of C3–C9 consumption was significantly faster and proceeded to a greater degree in normal serum, C3–C9 was also consumed in C2-deficient serum. These findings were consistent with observations made in the functional phagocytosis assay wherein C2-deficient serum was not as effective as normal serum as an opsonic source and opsonization proceeded at a slower rate in C2-deficient serum than in normal serum. Similar findings have been reported in studies of opsonization of other staphylococcal strains (28, 42). Recently, Bokisch (53) has reported that peptidoglycans from *Streptococcus* group A, *Staphylococcus epidermidis*, and *Micrococcus luteus* were capable of activating both the classical and the alternative C pathways in human serum. Pryjma and colleagues (4) have demonstrated the same capacity for *S. aureus* peptidoglycan using serum from guinea pigs and mice.

In an attempt to determine whether *S. aureus* peptidoglycan was capable of directly activating the classical and alternative C pathways or whether C activation was initiated by antigen-antibody complex formation, normal and C2-deficient sera were preabsorbed with peptidoglycan at 2°C before adding peptidoglycan at 37°C and measuring C3–C9 consumption. Preabsorption of both serum sources with peptidoglycan was found to limit the ability of peptidoglycan to subsequently activate the classical and alternative pathways of C. These preliminary findings suggest that C activation by peptidoglycan depends upon the presence of an absorbable, noncomplement-related factor, presumably immunoglobulin. Wheat and co-workers (54) have presented data to support this hypothesis. Activation of the classical C pathway, in general, is known to be initiated by the combining of IgG or IgM with antigen (55). There is a growing body of evidence which suggests that, in certain systems, activation of the alternative complement pathway may likewise depend upon antigen-antibody formation (46, 56, 57). Antibodies of the IgG class, to *S. aureus* peptidoglycan have been shown to be present in normal human serum (58). However, there is a need for studies utilizing purified C components to evaluate the possibility that peptidoglycan may directly interact with C1 and/or directly activate the alternative C pathway as described for certain gram-negative bacterial cell wall components (59).

Over the past decade it has been demonstrated that the recognition of staphylococci by PMN leukocytes is mediated by receptors within the phagocytic membrane with specificity for the Fc fragment of IgG and for an activated form of the third component of C, C3b (60, 61). Staphylococcal peptidoglycan would appear to play a central role in the initiation of this recognition process. Other investigators (1, 62) have proposed that the peptidoglycan, a major cell wall component of most gram-positive bacteria, may play a significant role in the development of "natural immunity" to this group of microorganisms. Recognition of all gram-positive bacteria may, therefore, be a manifestation of host response to this common cell wall component.

It should be emphasized that although cell wall teichoic acid was found not to be important in promoting opsonization in the serum sources used in this study, the possibility remains that teichoic acid is of importance in fostering opsonization in serum from patients with *S. aureus* infections in which increased levels of teichoic acid antibodies have been demon-
strated (63). Future studies will be directed at investigating this possibility.

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