Characterisation of *Eubacterium* cell wall: peptidoglycan structure determines arthritogenicity

X Zhang, M Rimpiläinen, E Šimelyte, P Toivanen

Abstract

**Objective**—To elucidate factors involved in the arthritogenicity of bacterial cell walls.

**Methods**—For characterisation of an arthritogenic *Eubacterium aerofaciens* cell wall, peptidoglycan-polysaccharide (PG-PS) polymers were isolated by removing cell wall associated proteins (CWPs), PG and PS moieties were separated, and an attempt was made to de-O-acetylate PG-PS. The cell wall of *E limosum* was used as a non-arthritogenic control. The chemical composition of these cell wall preparations was analysed by gas chromatography-mass spectrometry. Also, their ability to resist lysozyme degradation and to sustain experimental chronic arthritis was tested.

**Results**—The observations made with the cell wall of *E aerofaciens*, an anaerobic inhabitant of the human intestine, were compared with those reported from a pathogenic *Streptococcus*, showing that in both strains a complex consisting of PG-PS is required for the induction of chronic arthritis. The PS moiety most probably protects PG from enzyme degradation, allowing prolonged tissue persistence and leading to the chronic synovial inflammation. CWPs attached to PG-PS are not necessary for this function. O-Acetylation of PG, which is required for arthritogenicity of the streptococcal cell wall, seems not to be present in the arthritogenic *E aerofaciens* PG or only occurs to a small degree; attempts to de-O-acetylate the *E aerofaciens* cell wall did not affect its arthritogenicity or lysozyme resistance.

**Conclusion**—The results obtained indicate that the source of bacterial cell wall plays no part in the chemical or structural requirements for PG to induce chronic cell wall arthritis in the rats; the chemical structure of the PG moiety is decisive.

The intestinal flora of patients with newly developed rheumatoid arthritis (RA) is significantly different from that of non-RA controls. This difference is primarily due to anaerobic bacterial species—namely, to *Eubacterium aerofaciens*. On the other hand, a single intraperitoneal (IP) injection of cell walls isolated from this Gram positive bacterium, a normal inhabitant of the human gut, induces a chronic arthritis in Lewis rats, whereas cell walls from a related *E limosum* are non-arthritogenic.

The Gram positive bacterial cell wall skeleton mainly contains peptidoglycan (PG), polysaccharide (PS), and the cell wall associated proteins (CWPs) (fig 1). The CWPs in the Gram positive cell wall extend from the plasma membrane through the PG and PS layers to the cell surface and are covalently attached to the PG. They interact with the extracellular environment and contribute to the development of bacterial infections. CWPs of *Staphylococcus aureus* interact with specific host molecules either by promoting attachment to the host tissues or by preventing the phagocytic clearance. CWPs of *Mycobacterium leprae* are the major stimulants of the cell mediated immunity, and CWPs of *M tuberculosis* are known to play a part in establishing protective immunity. The M protein of the group A streptococcal CWPs is a major virulence factor with superantigenic activity. Group A streptococcal M protein has shared epitopes with the articular cartilage and synovium, and systemic administration of streptococcal cell wall induces a chronic, destructive arthritis in susceptible rat strains.

The bacterial cell wall PG-PS complex can initiate and sustain a chronic destructive inflammation. PG-PS isolated from human intestinal flora bacteria can induce chronic arthritis in rats. The arthritogenic ability of cell wall constituents is dependent on resistance to biodegradation and consequent persistence in the tissues. The PS moiety...
covalently bound to PG is essential for the PG-PS complexes to persist and induce chronic inflammation. Streptococcal PG-PS did not sustain chronic arthritis when the PS moiety was removed, and PS alone induced only mild oedema in the joints. Another structural basis for streptococcal arthropathogenicity is the presence of an acetyl moiety at the C-6 hydroxyl group of N-acetylmuramyl residues of PG, producing the corresponding 2,6-diacetylmuramyl PG (O-acetylated PG).

Gonococcal O-acetylated PG is more resistant to enzyme degradation than non-O-acetylated PG, and subsequently persists in the tissues longer, leading to chronic arthritis. De-O-acetylated group A streptococcal PG-PS induced less severe chronic arthritis than native PG-PS. In this study we aimed at defining the arthritogenic component(s) of the E. aerofaciens cell wall to determine whether the same requirements are valid for a representative of the normal human gut flora and a pathogenic Streptococcus.

**Methods**

**BACTERIA**

*Eubacterium aerofaciens* ATCC 25986 was obtained from the Culture Collection, University of Gothenburg, Sweden, with a nomination CCUG 28087. *E. limosum* ATCC 8486 was purchased from the American Type Culture Collection, Rockville, MD, USA. Both strains were grown overnight under strictly anaerobic conditions at 37°C in BBL Schaedler Broth (Becton Dickinson Microbiology Systems, Cockeysville, MD, USA) to the late logarithmic phase. They were characterised by 16S rDNA sequence analysis as described by Jalava et al.

**CRUDE CELL WALL PREPARATION**

Bacterial crude cell walls were isolated as described previously. Briefly, the cells were broken with glass beads (diameter 0.45–0.50 mm) in an MSK cell homogeniser (B Braun Melsungen AG, Melsungen, Germany). The cell walls were collected by centrifugation, washed three times with distilled water, and sonicated in an ice bath for 75 minutes (Branston Sonifier, Smith Kline Co, Danbury, Conn., USA). The sonicated cell wall suspension was centrifuged at 10 000 × g, 4°C, for 20 minutes. The supernatant was centrifuged by ultracentrifugation (Sorvall Ultracentrifuge OTD65B, rotor 60 Ti, Du Pont Co, Wilmington, USA) at 100 000 × g, 4°C, for 60 minutes. The pellet containing the cell wall was suspended in water. For chemical analysis the cell walls were dialysed in membrane (MWCO 3500, Spectrum, CA, USA) against distilled water. For IP injection into rats the cell wall suspension was sterilised by heating at 90°C for 30 minutes. The sterility was checked by cultures on agar plates at 37°C and at room temperature under aerobic and anaerobic conditions; no bacterial growth was detected after two days of culture. The endotoxin tests by E-TOXATE (Sigma Chemical Co, MO, USA) were also found to be negative. These procedures were applied for all cell wall preparations used.

**PG-PS POLYMERS**

To obtain PG-PS polymers, the CWPs were removed; the crude cell walls were treated with proteolytic enzymes, protease (Subtilisin Carlsberg, 5 mg/g wet weight), proteinase K (2 mg/g wet weight), pepsin (20 mg/g wet weight), and papain (5 mg/g wet weight). All enzymes were purchased from Sigma Chemical Co, MO, USA. After lyophilisation the cell walls were extracted with chloroform-methanol (2:1 vol/vol, 40 mg/ml) at room temperature for four hours to remove lipoteichoic acid or glycosyl glycerides. After washing once with water and dialysis against distilled water, the cell walls were treated by 8 M lithium chloride at room temperature for one hour to remove any non-covalently bound CWPs. The cell walls were collected by ultracentrifugation and washed twice with distilled water. They were extracted with 0.1 M EDTA at room temperature for one hour to remove any divalent cation stabilised oligomeric proteins. After ultracentrifugation and washing twice with distilled water the cell walls were suspended in acetone at room temperature for one hour and further mixed at 4°C for 30 hours to remove possible contaminating proteins and endotoxin. The cell walls were collected by ultracentrifugation, washed three times with distilled water, and dialysed against distilled water. The preparations thus obtained are PG-PS complexes as shown by gas chromatography-mass spectrometry (GC-MS) analysis.

**PS AND SOLUBLE PG POLYMERS**

PG and PS polymers were isolated from *E. aerofaciens* ATCC 25986 as described previously for *Lactobacillus*. Briefly, the lyophilised cell walls were extracted with 10% trichloroacetic acid at 60°C for four hours. The suspension was centrifuged at 38 800 × g at 4°C for 30 minutes and washed once with distilled water. The supernatant contained the PS fraction, and the pellet the PG fraction. The soluble PG was obtained as described previously. The PG fraction was lyophilised and suspended in water at 10 mg/ml and sonicated (Branson Sonifier) for 180 minutes (18 × 10-minute cycles with breaks for cooling between the cycles) in an ice bath. The remaining insoluble PG fraction was removed by centrifugation at 10 000 × g at 4°C for 30 minutes. The supernatant (soluble PG fraction) was collected and dialysed against distilled water for 48 hours.

**DE-O-ACYLATION OF CELL WALL**

The treatment of the cell walls for de-O-acylation was done as described previously. Briefly, the cell walls were lyophilised, suspended in 0.05 M sodium hydroxide at 37°C for four hours. After washing three times with distilled water, the cell walls were fragmented by sonication and subsequently separated by differential centrifugation as described above for the cell wall preparation.
The chemical composition of the cell wall preparations was analysed by GC-MS as described previously. The derivatised molecules were ionised by the electron impact method and analysed in the selected ion monitoring mode using single positive ions at a mass to charge ratio (m/z). Sugars were analysed as alditol acetate derivatives with the same ions as monitored by Gilbart et al, except m/z 289 for rhamnose. Fucose was used as internal standard for rhamnose, allose for glucose/galactose, and N-methyl-D-glucamine for muramic acid. Amino acids were analysed as butyl heptafluorobutyl derivatives with the same ions as used by Gilbart et al. L-Norleucine was used as internal standard for glycine, serine, alanine, leucine, cysteine, threonine, isoleucine, and valine; L-methionine for phenylalanine, glutamic acid/glutamine, proline, ornithine, lysine, aspartic acid/asparagine, and tyrosine; L-tryptophan for cystine, diaminopimelic acid, histidine, and arginine. One microlitre of the derivative was injected in the pulsed splitless mode and analysed by a gas chromatograph (model HP 5890A; Hewlett-Packard, Wilmington, DE) equipped with a fused silica capillary column (SE-54; Nordian Instruments, Helsinki) and coupled directly with a TRIO-1 mass spectrometer (VG Instruments,Manchester, UK). The oven temperature for sugars was started at 50°C, programmed to 270°C at a rate of 10°C/min, and held for one minute; finally, the oven was heated to 290°C for five minutes. For amino acids the oven temperature was started at 85°C, programmed to 280°C at a rate of 10°C/min, and held for one minute; finally, the oven was heated to 290°C for five minutes.

### Table 1 Gas chromatography-mass spectrometry analysis of the cell wall preparations

<table>
<thead>
<tr>
<th>Component</th>
<th>E aerofaciens</th>
<th></th>
<th>E limosum</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Crude cell wall</td>
<td>PG-PS*</td>
<td>PG</td>
<td>PS</td>
</tr>
<tr>
<td>Carbohydrates</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sugars</td>
<td></td>
<td></td>
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<tr>
<td>Rhamnose</td>
<td>18.5</td>
<td>18.0</td>
<td>—</td>
<td>30.3</td>
</tr>
<tr>
<td>Glucose + galactose</td>
<td>30.9</td>
<td>27.0</td>
<td>1.0</td>
<td>50.1</td>
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<tr>
<td>Amino sugars</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>N-Acetylglucosamine</td>
<td>1.9</td>
<td>9.2</td>
<td>11.5</td>
<td>0.6</td>
</tr>
<tr>
<td>N-Acetylgalactosamine</td>
<td>7.6</td>
<td>14.3</td>
<td>9.1</td>
<td>11.1</td>
</tr>
<tr>
<td>Amino acids</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Alanine</td>
<td>5.0</td>
<td>4.9</td>
<td>20.4</td>
<td>0.3</td>
</tr>
<tr>
<td>Glutamic acid/glutamine</td>
<td>6.0</td>
<td>8.1</td>
<td>32.8</td>
<td>1.0</td>
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<tr>
<td>Lysine</td>
<td>9.7</td>
<td>6.4</td>
<td>24.2</td>
<td>0.3</td>
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<tr>
<td>Aspartic acid/asparagine</td>
<td>4.8</td>
<td>6.9</td>
<td>17.0</td>
<td>0.4</td>
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<tr>
<td>Ornithine</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
<td>0.3</td>
<td>0.1</td>
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<tr>
<td>Serine</td>
<td>&lt;0.1</td>
<td>0.1</td>
<td>&lt;0.1</td>
<td>12.1</td>
</tr>
<tr>
<td>Leucine</td>
<td>0.4</td>
<td>0.4</td>
<td>0.4</td>
<td>0.1</td>
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<tr>
<td>Isoleucine</td>
<td>0.4</td>
<td>0.7</td>
<td>—</td>
<td>2.0</td>
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<tr>
<td>Phenylalanine</td>
<td>0.2</td>
<td>&lt;0.1</td>
<td>0.7</td>
<td>&lt;0.1</td>
</tr>
<tr>
<td>Proline</td>
<td>0.3</td>
<td>&lt;0.1</td>
<td>0.5</td>
<td>1.4</td>
</tr>
<tr>
<td>Tyrosine</td>
<td>0.3</td>
<td>&lt;0.1</td>
<td>0.3</td>
<td>1.0</td>
</tr>
<tr>
<td>Valine</td>
<td>0.4</td>
<td>—</td>
<td>—</td>
<td>1.6</td>
</tr>
</tbody>
</table>

Each component is expressed as a percentage of the dry weight of cell wall. Figures in bold indicate PG amino sugars and amino acids. Norleucine, methionine, tryptophan were applied as internal standards for amino acid analysis. Cystine, glycine, threonine, diaminopimelic acid, cysteine, arginine, and histidine were not found.

*PG-PS = peptidoglycan-polysaccharide.

**Figure 2** (A) A single intraperitoneal (IP) injection of *E aerofaciens* peptidoglycan-polysaccharide (PG-PS; 150 µg/g body weight of rat) induced a more severe chronic arthritis than that induced with a crude cell wall (140 µg/g body weight of rat) (p<0.02 by non-parametric Mann-Whitney U test). Each black symbol represents a mean (SEM) of six rats. Each white symbol represents a mean (SEM) of 11 rats. (B) Rats injected with *E limosum* PG-PS (200 µg/g body weight of rat) IP or with the crude cell wall (200 µg/g body weight of rat) developed only mild acute arthritis. Each black symbol represents a mean (SEM) of eight rats. Each white symbol represents a mean (SEM) of six to 12 rats.

**GAS CHROMATOGRAPHY-MASS SPECTROMETRY (GC-MS)**

The chemical composition of the cell wall preparations was analysed by GC-MS as described previously. The derivatised molecules were ionised by the electron impact method and analysed in the selected ion monitoring mode using single positive ions at a mass to charge ratio (m/z). Sugars were analysed as alditol acetate derivatives with the same ions as monitored by Gilbart et al, except m/z 289 for rhamnose. Fucose was used as internal standard for rhamnose, allose for glucose/galactose, and N-methyl-D-glucamine for muramic acid. Amino acids were analysed as butyl heptafluorobutyl derivatives with the same ions as used by Gilbart et al. L-Norleucine was used as internal standard for glycine, serine, alanine, leucine, cysteine, threonine, isoleucine, and valine; L-methionine for phenylalanine, glutamic acid/glutamine, proline, ornithine, lysine, aspartic acid/asparagine, and tyrosine; L-tryptophan for cystine, diaminopimelic acid, histidine, and arginine. One microlitre of the derivative was injected in the pulsed splitless mode and analysed by a gas chromatograph (model HP 5890A; Hewlett-Packard, Wilmington, DE) equipped with a fused silica capillary column (SE-54; Nordian Instruments, Helsinki) and coupled directly with a TRIO-1 mass spectrometer (VG Instruments, Manchester, UK). The oven temperature for sugars was started at 50°C, programmed to 270°C at a rate of 10°C/min, and held for one minute; finally, the oven was heated to 290°C for five minutes. For amino acids the oven temperature was started at 85°C, programmed to 280°C at a rate of 10°C/min, and held for one minute; finally, the oven was heated to 290°C for five minutes.

**ANIMALS AND INDUCTION OF ARTHRITIS**

All the animal experiments were done in concordance with the national and international
laws and policies (Order No 1076/85, Finland; EEC Council Directive 86/609, OJL 358, Dec 1987). Inbred pathogen-free female Lewis rats weighing approximately 150 g were purchased from Harlan Sprague Dawley, Inc, Indianapolis, IN, USA. The animals were kept in Macrolon III cages with disposable filter tops (Scanbur, Denmark) and given autoclaved standard diet and water; all handling was performed in a laminar-flow hood. Arthritis was induced by a single IP injection of the cell wall preparations suspended in sterile phosphate buffered saline (PBS) on day zero. Control rats were injected with an equal volume of sterile PBS. To monitor development of arthritis, each limb was assigned a score of 0–4, based on the degree of erythema, oedema, painfulness, and functional disorder of the ankle and metatarsal joints (wrist and metacarpal joints) and judged by two independent observers as described previously. The arthritis score used in the “Results” is the sum of the score of each limb. The animals were observed for at least four weeks. Rats were killed under anaesthesia by Metofane (Pitman-Moore, Inc, Washington Crossing, NJ, USA).

**Sensitivity to lysozyme**

The sensitivity of different cell wall preparations to lysozyme was tested as described previously. Suspensions of cell wall preparations (4 mg/ml) in 0.1 M sodium acetate buffer (pH 5.0) were incubated with lysozyme (400 µg/ml) at 37°C. The optical density of the suspension was measured at wavelength 560 nm after five and 24 hours’ incubation.

**Statistical analysis**

Differences of arthritis development between the experimental groups were analysed with the non-parametric Mann-Whitney U test. Differences were considered significant at p<0.05.

**Results**

**Identification of bacteria**

The gene coding for 16S ribosomal RNA is generally used to identify bacterial strains. To confirm the identity of the bacterial strains used in this study the genes coding for this region were partially sequenced (320–452 base pairs). Both for *E. aerofaciens* and for *E. limosum*, a 100% identity with respective sequences in the GenBank data base was confirmed. In addition, a 72.6% identity between the strains of *E. aerofaciens* and *E. limosum* was obtained.

**Analysis of cell wall preparation by GC-MS**

All bacterial cell wall preparations were analysed by GC-MS before injection into the rats (table 1). The total amount of protein amino acids is 2.2% in the *E. aerofaciens* crude cell wall, and less than 0.6% in its PG-PS polymer, in which the amount of individual protein amino acids is less than 0.1%. The amount of protein amino acids is 12.6% in the *E. limosum* crude cell wall, and less than 0.7% in its PG-PS polymer. Both bacterial PSs contain rhamnose and glucose/galactose. *E. aerofaciens* PS also contains N-acetylglucosamine. N-Acetylgalactosamine is present in the *E. limosum* cell wall and missing in the *E. aerofaciens* cell wall. PG of both cell walls contains the amino sugars N-acetylmuramic acid and N-acetylgalactosamine. Arthritogenic *E. aerofaciens* has four PG amino acids: alanine, glutamic acid, lysine, and aspartic acid. Non-arthritogenic *E. limosum* cell wall has five PG amino acids: alanine, glutamic acid, lysine, ornithine, and serine (table 1). These findings are in concert with the previously reported PG types—A4u for *E. aerofaciens* and B2a for *E. limosum.*

**Induction of arthritis**

For induction of arthritis a single IP injection of the cell wall components was applied. *E. limosum* PG-PS appeared non-arthritogenic, similar to the crude cell wall of *E. limosum* (fig 2B). A high dose (200 µg/g body weight) of PG-PS isolated from the arthritogenic *E. aerofaciens* crude cell wall was toxic, leading to death within three days (data not shown here), probably owing to the high PG content. When a lower dose (150 µg/g body weight of rat) was used, *E. aerofaciens* PG-PS caused a significantly more severe chronic arthritis than the equivalent dose of the crude cell wall (fig 2A), suggesting that the PG-PS complex possesses...
Characterisation of Eubacterium cell wall

Table 2  Degradation of bacterial cell wall preparations by lysozyme

<table>
<thead>
<tr>
<th>Substrate</th>
<th>Lysozyme 5 h</th>
<th>Lysozyme 24 h</th>
</tr>
</thead>
<tbody>
<tr>
<td>E. aerofaciens</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Crude cell wall</td>
<td>2.0</td>
<td>15.6</td>
</tr>
<tr>
<td>De-O-acetylated</td>
<td>5.3</td>
<td>31.0</td>
</tr>
<tr>
<td>PG-PS</td>
<td>1.9</td>
<td>15.8</td>
</tr>
<tr>
<td>PG</td>
<td>0.0</td>
<td>75.9</td>
</tr>
<tr>
<td>E. limosum</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Crude cell wall</td>
<td>0.7</td>
<td>24.4</td>
</tr>
</tbody>
</table>

Results are given as the percentage decrease of the OD_{560} at time 0. Mean values from triplicate five hours and 24 hours' incubation with or without lysozyme, experiments are given.

Discussion

It has been shown that synovial tissue of patients with RA may harbour traces of bacterial PG.32 The results presented in this paper, obtained with a normal bacterial habitant (E. aerofaciens) of the human intestine, indicate that the PG moiety of the bacterial cell wall contains a structure decisive for induction of chronic arthritis in rats. They also show that the PG-PS complex is necessary for the arthritis induction; the PS moiety most probably protects PG from enzyme digestion through steric hindrance, allowing prolonged tissue persistence and leading to the chronic synovial inflammation. Similar conclusions have previously been reached in studies on streptococcal cell wall arthritis.16 Up to this point, our observations indicate that the same principles apply for the chemical structure, determining arthritogenicity here also, independently of the bacterial species used. However, in contrast with the streptococcal cell wall, in which the O-acetylation is known to be associated with the arthritogenicity and resistance against biodegradation,15 O-acetylation does not seem to be a prerequisite for the arthritogenicity or biodegradation of the E. aerofaciens cell wall. Our findings suggest that O-acetylation does not occur in E. aerofaciens cell wall or only occurs to a small degree. The gonococcal cell wall seems to resemble the streptococcal cell wall in this respect because O-acetylation is required for its arthritogenicity.43 20 Lysozyme susceptibility is usually thought to be associated with arthritogenicity of the cell wall. Almost all bacterial strains studied for their ability to induce cell wall arthritis have been found to be lysozyme resistant; this includes species of Lactobacillus, Streptococcus, Bifidobacterium, and Eubacterium.53 The only exception is the E. limosum cell wall, which is non-arthritogenic and which has been reported to be lysozyme resistant.1 However, our results obtained by using the same strain of E. limosum indicate both non-arthritogenicity and lysozyme sensitivity (table 2). Furthermore, because PG is degraded in vivo by other enzymes also (for example, N-acetylmuramyl-l-alanine amidase, N-acetylglucosaminidase, and others)19 34 35 it is unlikely that lysozyme resistance alone would be a decisive requirement for the ability of a cell wall to induce chronic arthritis.

It has so far remained undetermined whether CWPs attached to PG-PS contribute or are required for induction of chronic arthritis. Our results indicate that their presence is not necessary for arthritis induction. Removal of CWPs by proteolytic enzymes resulted in PG-PS which induced an even more severe chronic arthritis than that seen after injection of an equivalent amount of the crude cell wall (fig 2A). These findings parallel those showing that genetic mutation of the streptococcal M protein attached to PG does not change arthritogenicity of the cell wall.36 Otherwise, results from GS-MS analyses (table 1) show very little, if any, further information about chemical or structural requirements for arthritogenicity. For instance, the presence of rhamnose does not correlate with this ability, even though such a correlation has previously been suggested.37

In a previous study on four Eubacterium strains it was shown that the chemical structure of PG present in the bacterial cell walls is decisive in determining arthritogenicity/non-arthritogenicity.53 The present results, based on characterisation of PG-PS complexes, further strengthen that conclusion by indicating that the source of a cell wall plays no part in the chemical or structural requirements of PG arthritogenicity; it does not matter whether the cell wall is isolated from a normal inhabitant of the human intestine or from a pathogenic strain of Streptococcus. The structure of PG is decisive. Studies with other bacterial species—for example, with Clostridia and Bifidobacteria, are needed to elucidate further the detailed structural determinants of the PG arthritogenicity.

We thank Leena Kivistö, Marja-Riitta Terasjärvi, Karolina Karlson, and Heli Näättymäki for excellent technical assistance, Jari Jalava for 16S rDNA sequence analysis, and Yong Zhang for help in evaluating the arthritis.

This work was supported by EVO of Turku University Central Hospital.


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*Ann Rheum Dis* 2001 60: 269-274
doi: 10.1136/ard.60.3.269

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