The response of selected members of the archaea to the Gram stain

Terry J. Beveridge and Susanne Schultze-Lam

Archaea possess a broader range of cell envelope structural formats than eubacteria and their cell walls do not contain peptidoglycan. Some archaea have only a single S-layer as their cell wall (e.g. Methanococcus jannaschii and Sulfolobus acidocaldarius), whereas others have multiple layers (e.g. Methanospirillum hungatei). Sometimes there can also be a high proportion of tetraether lipids in membranes to make the envelope more resilient to environmental stress (e.g. Methanococcus jannaschii and Sulfolobus acidocaldarius grown at 70 °C). Since the Gram reaction depends on both the structural format and the chemical composition of the cell envelope of eubacteria, it was important to determine if the same is true for archaea. Methanospirillum hungatei, Methanosarcina mazei, Methanobacterium formicicum, Methanococcus jannaschii and Sulfolobus acidocaldarius, chosen because of their different envelope formats and chemistries, were subjected to a Gram stain that can be used for transmission electron microscopy. In this staining regimen, the iodine is replaced by potassium trichloro(ethylene)platinate(lI) as the mordant, and the platinum of the new compound is the electron-scattering agent for electron microscopy. Of all these archaea, only Methanobacterium formicicum stained Gram-positive since its pseudomurein wall remained intact; the platinum compound formed large electron-dense aggregates with the crystal violet that were located in the vicinity of the cell wall and the plasma membrane. All but the terminal filament cells of Methanospirillum hungatei stained Gram-negative because the limiting porosity of its external sheath was so small that the Gram reagents could not enter the cells. The terminal cells of filaments stained Gram-positive because the staining reagents gained entry through the terminal plugs. All other archaea stained Gram-negative because their cell walls were so disrupted during staining that the crystal violet-platinum complex could not be retained by the cells. Methanococcus jannaschii was grown at both 50 °C and 70 °C so that the tetraether lipids in its plasma membrane could be increased from 20% (50 °C) to 45% (70 °C) of the total lipids; in both cases the cells stained Gram-negative.

Keywords: archaea, Gram stain, walls, cell envelopes, ultrastructure

INTRODUCTION

Microbiologists rely on the Gram stain for the initial screening of new isolates so that they can be classified as either Gram-positive or Gram-negative. Since the turn of the century, the correlation between Gram stain and structural detail of cell envelopes has been derived from years of experience by many microbiologists but is based, primarily, on eubacteria. Traditionally this has meant that Gram-negative bacteria possessed a bilayered outer membrane, a thin peptidoglycan layer and a bilayered plasma membrane (such as found in Escherichia coli) as their essential structural elements for a cell envelope (Beveridge, 1981; Beveridge & Graham, 1991). Gram-positive

Abbreviations: EDS, energy-dispersive X-ray spectroscopy; LM, light microscopy; TEM, transmission electron microscopy; TPt, trichloro(ethylene)platinate.
bacteria (such as *Bacillus subtilis*) possessed thick amorphous cell walls overlying the plasma membrane (Beveridge, 1981), although more recent work using cryo-electron microscopy has now shown these envelopes to be more complex (Beveridge & Graham, 1991; Graham & Beveridge, 1994). The staining response for eubacteria has usually been clear-cut with few exceptions (Beveridge & Davies, 1983; Beveridge, 1990).

Surprisingly, little work has been done on the archaea and their response to the Gram stain. Most representatives of this relatively recently recognized Domain stain Gram-negative, but the rationale for this is still unclear. Since the structural formats of archaeal cell envelopes are dramatically different from those of eubacterial envelopes and because their chemical makeup also differs (Beveridge, Patel, 1980) supplemented with 5 mM NiCl₂, *Methanosoccus jannaschii* JAL-1 (DSM 2661) was grown in defined medium under the same H₂/CO₂ atmosphere at either 50 °C or 70 °C as described by Ferrante et al. (1990). *Methanobacterium formicum* was grown in the medium described by May et al. (1988). *Methanosarcina mazei* S6 (DSM 2053) was grown at 35 °C on methanol (0.6%, v/v) under N₂ in Balch medium no. 3 in which the yeast extract and tryptone were replaced by 0.1 g L-isoleucine 1⁻ and 0.05 g L-leucine 1⁻; the NH₄Cl concentration was raised to 0.5 g 1⁻, and Na₂CO₃ was replaced by NaHCO₃. *Sulfobolus acidocaldarius* 98-3 (ATCC 33900) was grown under aerobic conditions at 70 °C using the medium described by Brock et al. (1972).

**METHODS**

**Bacteria and growth conditions.** *Methanospirillum hungatii* GP1 (DSM 1101) was grown at 35 °C under an atmosphere of H₂/CO₂ (80:20, v/v) in mineral salts medium SA (Breuil & Patel, 1980) supplemented with 5 mM NiCl₂. *Methanosoccus jannaschii* JAL-1 (DSM 2661) was grown in defined medium under the same H₂/CO₂ atmosphere at either 50 °C or 70 °C as described by Ferrante et al. (1990). *Methanobacterium formicum* was grown in the medium described by May et al. (1988). *Methanosarcina mazei* S6 (DSM 2053) was grown at 35 °C on methanol (0.6%, v/v) under N₂ in Balch medium no. 3 in which the yeast extract and tryptone were replaced by 0.1 g L-isoleucine 1⁻ and 0.05 g L-leucine 1⁻; the NH₄Cl concentration was raised to 0.5 g 1⁻, and Na₂CO₃ was replaced by NaHCO₃. *Sulfobolus acidocaldarius* 98-3 (ATCC 33900) was grown under aerobic conditions at 70 °C using the medium described by Brock et al. (1972).

**RESULTS**

Table 1 shows the results of the analyses of the archaea by LM, TEM and EDS; Figs 1–4 provide representative images on which Table 1 is based. Fig. 5 shows the typical
TEM-EDS results for a control Gram-positive eubacterium (Streptococcus pyogenes).

The archaea used for our study were carefully chosen so that they represented the diverse number of cell envelopes that exist in this Domain. These can be very simple (i.e. the cell wall can consist of a single S-layer as in Sulfolobus acidocaldarius and Methanococcus jannaschii) or they can be amongst the most complex of all prokaryotes (i.e. the multiple layers in Methanospirillum hungatei). Fig. 1 shows the cell envelope formats that our archaea possessed. Methanococcus jannaschii (Fig. 1a) and Sulfolobus acidocaldarius (Fig. 1b) had the simplest envelopes, consisting of a plasma membrane and a single proteinaceous (Methanococcus jannaschii) or glycoproteinaceous (Sulfolobus acidocaldarius) S-layer. There was no change in the envelope profile of Methanococcus jannaschii when it was grown at either 50 °C or 70 °C. Methanobacterium formicicum did not possess an S-layer, but it had a relatively thick (~15 nm) amorphous wall (Fig. 1c) composed of pseudomurein (Kandler & König, 1985; König, 1988), whereas Methanosarcina mazei possessed a wall of variable thickness (15–30 nm; Fig. 1d) composed of amorphous methanochondroitin (Kandler & König, 1985; König, 1988) which overlies a single S-layer (Aldrich et al., 1986). This S-layer was not seen in any of our thin sections (Fig. 1d) even when aligned correctly at right angles to the electron beam or when tilting experiments were performed. The most complex envelope profile was that of Methanospirillum hungatei, which consisted of an S-layered sheath (Stewart et al., 1985), an S-layered wall (Firtel et al., 1993), and a multilayered cell spacer (Firtel et al., 1994) that separated the cells from one another within a filament (Fig. 1e; see Beveridge et al., 1987, for more structural details).

Light microscopy (LM)

When the cells were subjected to either the conventional regimen or the TPt regimen of the Gram stain, LM showed most of the archaea to be Gram-negative (Table 1) when compared to the eubacterial controls. The two staining regimens were indistinguishable from one another on all bacteria. It was our impression that Methanococcus jannaschii (at both growth temperatures), Sulfolobus acidocaldarius and Methanosarcina mazei did not have the same colour intensity (as Gram-negatives) as our eubacterial E. coli control had when we viewed the samples by LM. Light micrographs of these stains, after the unstained backgrounds of each were adjusted to similar intensities, confirmed our visual evaluation (data not shown). Presumably, this was an indication of the

![Fig. 1. Thin sections of cell envelopes of the archaea used in this study.](image-url)
Table 1. Description of the selected archaea and the results of the Gram stain as seen by LM, TEM and EDS

<table>
<thead>
<tr>
<th>Bacterium</th>
<th>Cell envelope structure*</th>
<th>Gram reaction</th>
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<tbody>
<tr>
<td><em>Sulfolobus acidocaldarius</em></td>
<td>Single S-layer; plasma membrane with high concentrations of tetraether lipids</td>
<td>Gram-negative; Cells lysed; no TPt</td>
</tr>
<tr>
<td><em>Methanococcus jannaschii</em></td>
<td>Single S-layer; plasma membrane with low amounts of tetraether lipids</td>
<td>Gram-negative; Cells lysed; no TPt</td>
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<tr>
<td>At 50 °C</td>
<td></td>
<td></td>
</tr>
<tr>
<td>At 70 °C</td>
<td>As above, with high amounts of tetraether lipids</td>
<td>Gram-negative; Cells lysed; no TPt</td>
</tr>
<tr>
<td><em>Methanobacterium formicicum</em></td>
<td>Amorphous cell wall of pseudomurein; plasma membrane with intermediate amounts of tetraether lipids</td>
<td>Gram-positive; Cells intact; TPt associated with cell wall and plasma membrane</td>
</tr>
<tr>
<td><em>Methanosarcina mazei</em></td>
<td>Amorphous cell wall of methanochondroitin over top of a single S-layer; few tetraether lipids in plasma membrane</td>
<td>Gram-negative; Cells lysed; no TPt</td>
</tr>
<tr>
<td><em>Methanospirillum bungatei</em></td>
<td>S-layered sheath overlying a single S-layered wall; multilayered spacer plugs between cells; intermediate amounts of tetraether lipids in plasma membrane</td>
<td>Gram-negative except for the terminal cells of each filament, which were Gram-positive; Cells intact; TPt found only in terminal cells of each filament</td>
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amount of carbol fuchsin (red) which complexed to the cells as a counterstain; all three archaea complexed less counterstain than *E. coli*.

*Methanobacterium formicicum* stained Gram-positive (Table 1); yet, when compared to *Streptococcus pyogenes* or *B. subtilis* as eubacterial controls, it was apparent that *Methanobacterium formicicum*’s staining intensity was less than that of the two eubacteria (compare Fig. 2a and b). *Methanobacterium formicicum*’s colour was definitely purple and the cells were stained darker than *E. coli* (compare Fig. 2b and c) but it was not as dark as that of *B. subtilis* (Fig. 2a). The difference in staining intensity between *Methanobacterium formicicum* and *B. subtilis* could have simply been due to the narrower cell diameter of the former, but *S. pyogenes* (as another Gram-positive eubacterium), which has a similar cell diameter to *Methanobacterium formicicum*, was also more intensely stained. In

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Fig 2. Gray-tone light micrographs of Gram-stained bacteria showing the different stain intensities. *B. subtilis* stained deep purple (a), *Methanobacterium formicicum* was lighter purple (b), and *E. coli* was deep red (c). All micrographs are the same magnification and the cell diameter of *E. coli* is 0.5 μm.

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Fig. 3. Thin sections of single S-layered archaea before and after the Gram stain. (a, b) *Sulfolobus acidocaldarius* before (a) and after (b) the Gram reaction. Both of these sections are stained by heavy metal salts to heighten contrast and to show that most of the cellular substance has been lost during the reaction. The unstained section of *Sulfolobus acidocaldarius* shows that there are no electron-dense TPI-crystal violet aggregates in the cell, which was confirmed by EDS (c; arrow points to cellular substance within a lysed cell). Panel (d) is a heavy-metal-contrasted section of *Methanococcus jannaschii* (grown at 70 °C) and demonstrates that there is also little cellular substance in this cell after Gram staining. Bars, 0.5 µm.
this case, it seemed that *Methanobacterium formicicum*’s low staining intensity was due to a low retention of crystal violet (as a TPT or iodide complex) as compared to the two Gram-positive eubacterial controls.

The staining of *Methanospirillum bongatei* was the same as in our former results (Beveridge et al., 1991). The filaments were typically 7–15 cells long and only the terminal cells stained Gram-positive (see Fig. 3 of Beveridge et al., 1991). The remaining cells in each filament were Gram-negative (Table 1).

**Transmission electron microscopy (TEM) and EDS**

TEM was performed at every step of the Gram stain and it was apparent that both the initial crystal violet solution (which contains ethanol) and the ethanol decolorization step were disruptive to the cell envelopes of *Sulfolobus acidocaldarius* and *Methanococcus jannaschii*. By the end of the staining regimen, there was not much substance left to the cells (compare Fig. 3a to Fig. 3b and d) and no electron-dense TPT deposits could be seen within the bacteria (Fig. 3c), nor could EDS detect TPT (data not shown). Even when *Methanococcus jannaschii* was grown at 70 °C to ensure the presence of tetrathyl lipid, which strengthens and confers rigidity to bilayered membranes (Beveridge et al., 1993; Choquet et al., 1994), the results were the same (Table 1, Fig. 3d).

*Methanobacterium formicicum* was not disrupted by the staining regimen (Table 1). The cells remained intact and the division sites (septa) were not subject to ‘blow-out’ as are some Gram-positive eubacteria (Beveridge, 1990). Interestingly, there were few crystal violet–TPT aggregates within the cytoplasm since most were found at the plasma membrane–cell wall interface (Fig. 4a, b). EDS clearly identified the platinum within the aggregates (Fig. 4b). Because these stain aggregates were only found at the cell periphery, the overall number of precipitates was lower than for eubacterial cells (in which the precipitates are spread throughout the cytoplasm; Beveridge &...
Fig. 5. Heavy-metal-contrasted thin section of a *S. pyogenes* cell as a Gram-positive eubacterial control. TPt-crystal violet aggregates are spread throughout the cytoplasm, as is confirmed by the EDS spectrum of an unstained cell (arrows point to Pt lines). Bar, 0·5 μm.

Davies, 1983), and this could account for the lower staining intensity that was seen by LM (Fig. 2b). Presumably, the crystal violet and TPt entered the cell as soluble components and, once in the cytoplasm, reacted together to form a precipitate that was too large to migrate through the cell wall substance and be removed by the decolorization regimen. The precipitates therefore lodged themselves underneath the cell wall. *Streptococcus pyogenes* (as a Gram-positive eubacterial control) had staining aggregates scattered throughout its cytoplasm (Fig. 5) as did *B. subtilis* (data not shown). In total per cell, the number of these in the two eubacteria was greater than those found in *Methanobacterium formicicum*.

We were surprised by the Gram-negative results for *Methanosarcina mazei* since this archon is usually considered to be Gram-positive because of its cell envelope structure by TEM and its methanochondroitinous cell wall (Kandler & König, 1985; Aldrich et al., 1986). This bacterium usually forms large cell aggregates in which the wall of one cell seems continuous with those of neighbouring cells (Sprott & Beveridge, 1993). Therefore, *Methanosarcina mazei* usually has robust cell walls which should retain the crystal violet–TPt complex of our Gram stain. At mid-exponential growth phase, our cultures contained cell aggregates of only 4–6 cells; this could be due to the action of the *Methanosarcina mazei* dis-aggregatase, which is an enzyme situated at the cell periphery that degrades the cell wall methanochondroitin (Xun et al., 1990; de Macario et al., 1993). The cell walls of our culture were uneven and ranged from 10 to 15 nm in thickness (Fig. 1d); this unevenness (presumably a result of wall breakdown) made these cells sensitive to the decolorization step of the Gram stain. These bacteria stained Gram-negative because the cells lysed and liberated their crystal violet–TPt aggregates (Table 1).

TEM and EDS confirmed earlier work on *Methanospirillum hungatei* (Beveridge et al., 1991). TEM revealed that all cells within the filaments were intact and, essentially, unaffected by the Gram stain. However, crystal violet–TPt aggregates could only be seen in the terminal cells of each filament (Table 1; see also Fig. 4 of Beveridge et al., 1991). EDS confirmed the presence of platinum in these terminal cells (Table 1). It seemed that the staining reagents entered only through the terminal plugs (i.e. the spacer plug at the end of each filament; see Beveridge et al., 1987, for more details of the structure of *Methanospirillum hungatei*), thereby staining the terminal cells. Once crystal violet–TPt aggregates were formed, they were too large to be washed out through these same plugs by the ethanol decolorization step (see Firtel et al., 1994, for a description of plug organization and subunit periodicity). The more internal cells of each filament did not stain Gram-positive because the S-layered sheath (which is their outermost envelope layer) is so impermeable that the staining reagents could not pass through and enter these cells (Beveridge et al., 1991). To our knowledge this remarkable staining phenomenon has, so far, never been identified in another bacterium.
DISCUSSION

All of the archaea that were used in this study possessed cell envelopes very different from those found in eubacteria. Methanococcus jannaschii and Sulfolobus acidocaldarius possess very simple cell walls consisting of a single paracrystalline array, or S-layer, composed of protein (Methanococcus jannaschii) or glycoprotein (Sulfolobus acidocaldarius) (Deatherage et al., 1983; Michel et al., 1980; Sprott & Beveridge, 1993; Weiss, 1974). Both of these archaea produce plasma membranes with high concentrations of membrane-spanning tetraether lipids when grown at elevated temperature (i.e. 70 °C). Methanococcus jannaschii can also be grown at 50 °C and, at this temperature, these lipids are reduced from ~45% of the membrane lipid content to 20% (Sprott et al., 1991; Beveridge et al., 1993). At the higher concentration, the membranes are so greatly strengthened by tetraether lipids that they cannot be fractured through their hydrophobic domains during freeze-etching (Beveridge et al., 1993). Using these two archaea and two growth temperatures for Methanococcus jannaschii, we were able to determine if simple S-layered walls containing either protein or glycoprotein, and plasma membranes containing either high or low concentrations of tetraether lipids, had any effect on the action of the Gram stain. Neither of these archaea could retain enough cell envelope integrity during staining to entrap crystal violet–TPT precipitates at any temperature (Table 1). These bacteria stain Gram-negative because their S-layers and plasma membranes (even with a high tetraether content) are disrupted and dissolved (Fig. 3b, d). Presumably, this is because their S-layers are held together by only weak bonding forces (i.e. ionic bonding and hydrophobic interaction) and their plasma membranes are sensitive to the ethanol of the decolorization regimen of the Gram stain. Additional experiments whereby these cells were subjected to only ethanol (50–100%, v/v) also lysed these cells (T. J. Beveridge, unpublished results). In both these ethanol and Gram-stain experiments, the cellular residue sustained the natural spherical shape of the cells (this is the minimal energy shape of the vesicular remains) and carbol fuchsin ionically bonded to the spheres to stain them red (see Beveridge & Davies, 1983, for more details).

Methanosarcina mazei and Methanospirillum hungatei also stained Gram-negative but for entirely different reasons. Under our growth conditions Methanosarcina mazei did not develop large clusters of cells with thick cell walls. Instead, the walls were 10–15 nm thick and aggregates of 4–6 cells were common, presumably because of the hydrolytic action of disaggregatase, an enzyme which is found in the cell envelope and breaks down wall material (Xun et al., 1990; de Marcario et al., 1993). Like the envelopes of Methanococcus jannaschii and Sulfolobus acido-caldarius, those of Methanosarcina mazei could not withstand the action of the Gram stain (Table 1). Laboratory cultures of Methanosarcina mazei which have entered stationary growth phase, or cells in natural microbial communities, typically result in aggregates of 20 or more cells, each with thick (25–50 nm) robust walls (Sprott & Beveridge, 1993). Most cells in these aggregates will stain Gram-positive by LM (T. J. Beveridge, unpublished data). It is important to recognize that growth phase, activity of disaggregatase, and cell wall thickness can affect this archean’s response to the Gram stain. Thus, for Methanospirillum hungatei, the Gram stain can be an unreliable diagnostic technique.

The most remarkable staining response was seen with Methanospirillum hungatei. Most of the cells of this culture were Gram-negative because the staining reagents were too large to pass through the outermost cell envelope layer, the S-layered sheath, and enter the cytoplasm. Only the terminal cells of each filament stained Gram-positive since the reagents, in this case, could enter through the terminal plugs. These results confirm our earlier report (Beveridge et al., 1991) and we will not reiterate the expanded explanations here. Research has continued to verify the extraordinary physical and chemical resilience of Methanospirillum hungatei’s S-layered sheath. For example, only the most harsh chemical conditions (e.g. extremely high pH or phenol) are capable of disrupting its fabric (Beveridge et al., 1985; Southam & Beveridge, 1992) and it can withstand almost 400 atm of physical pressure (Xu et al., 1996). The sheath’s minute pore-size and its strong physical nature are unaffected by the Gram stain and these properties ensure this archaeon’s unique staining response.

Archaea possess a remarkable variety of cell envelope structural formats and chemical compositions. The present study has shown that in these organisms, as in eubacteria, the Gram stain is dependent on these cell envelopes, especially the cell walls. The physical nature of the wall and, particularly, its bonding structure are of utmost importance. Simple, single S-layered walls (such as those of Methanococcus jannaschii and Sulfolobus acidocaldarius) possess subunits that are weakly bonded to one another (Messner & Sleytr, 1992) and intimately associated with the underlying plasma membrane; these walls are breached by the action of the Gram stain. More robust walls (such as those of Methanosarcina mazei) can be subject to the lytic action of disaggregatase so that they, too, are disrupted during staining. Highly bonded cell walls (such as those of Methanobacterium formicicum), containing an analogue of peptidoglycan (i.e. pseudomurein), are strong enough to ensure that the cells stain Gram-positive even though the stain distribution differs from that found in Gram-positive eubacteria. Remarkably, at least one archean (Methanospirillum hungatei) has such an impermeable envelope layer (the sheath) that the staining reagents cannot enter the majority of the cells.

The Gram stain may depend on the cell envelope in archaea, but the exceptional variability of these envelopes between genera ensures that the essential mechanism of Gram reaction can be quite different from that seen in eubacteria.

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