Surface Density of Major Outer Membrane Proteins in Salmonella typhimurium in Different Growth Conditions

By MARTI ALDEA, ENRIQUE HERRERO,* M. ISABEL ESTEVE AND RICARDO GUERRERO

Department of Microbiology and Institute of Fundamental Biology, Autonomous University of Barcelona, Bellaterra, Barcelona, Spain

(Received 30 November 1979; revised 11 March 1980)

The amount of major proteins per unit of surface area in the outer membrane (OM) of Salmonella typhimurium LT2 remained constant during steady-state growth in different media. Between growth rates of 2.40 doublings h⁻¹ and 0.31 doublings h⁻¹, the surface density of major OM proteins varied between 0.9 × 10⁵ and 1.2 × 10⁵ molecules per μm², while surface area per cell more than halved. The accumulation of molecules of the major OM proteins was not affected by addition of cyclic AMP to the growth medium. When exponentially growing cells were subjected to shift-up transitions, cell dimensions began to increase after a lag period of 20 min. Accumulation of major OM proteins followed the same pattern as total protein; this created a transitory imbalance of major OM protein density in the shift from acetate minimal medium to LB medium, before the steady situation was reached. After shift-down transitions, cell dimensions began to decrease immediately, cells eventually becoming shorter than in steady-state conditions. No fluctuations in major OM protein density were observed during the shift-down, although final stable levels differed from those in steady-state conditions. All these results indicate that bacteria adapt the accumulation of major proteins into the OM according to the amount of surface. Thus, no large differences exist at different cell sizes, although transitions between media can lead to transitory or stable changes in the composition of the OM.

INTRODUCTION

The outer membrane (OM) of Gram-negative bacteria contains at least 20 different polypeptides, as resolved by sodium dodecyl sulphate–polyacrylamide gel electrophoresis (SDS–PAGE) (Schnaitman, 1970; Ames et al., 1974; Kamio & Nikaido, 1977; DiRienzo et al., 1978), while the number of types of protein in the whole envelope has been estimated as about 150 by two-dimensional gel electrophoresis (Ames & Nikaido, 1976). Four or five of the OM proteins (OMP) of Escherichia coli are present in rather higher amounts than the others, so they are called 'major proteins' (Schnaitman, 1970; DiRienzo et al., 1978). Some of these major OMP of E. coli are peptidoglycan-associated and have been alternatively named matrix protein (Rosenbusch, 1974), proteins 1a and 1b (Bassford et al., 1977), proteins 1a and 1b (Schmitges & Henning, 1976), proteins b and c (Lugtenberg et al., 1975) and porins (Kamio & Nikaido, 1977). Studies with cross-linking agents have revealed the close spatial association between at least some molecules of porins (Palva, 1979), supporting previous observations with the electron microscope which indicated that porins were arranged as trimers with a central pore (Steven et al., 1977). Proteins b and c of E. coli K12 are biochemically different (Henning et al., 1977; Lee et al., 1979), although they
seem to be commonly regulated by the product of the \textit{ompB} gene (Verhoeff \textit{et al.}, 1979). With the usual SDS–PAGE methods, porins appear in \textit{Salmonella typhimurium} in three bands at the region 34 to 36 kilodaltons (Ames \textit{et al.}, 1974; Kamio & Nikaido, 1977; Gmeiner & Schlecht, 1979) and their role in the penetration of the OM by hydrophilic molecules smaller than 600 daltons has been demonstrated (Decad & Nikaido, 1976; Nakae, 1976). The 35 kilodalton protein of \textit{S. typhimurium} is similar to protein b of \textit{E. coli} K12 (Sato & Yura, 1979). In addition, \textit{Salmonella} OM contains another major protein of 33 kilodaltons, although no pore activity has been clearly demonstrated for it.

During the cell cycle of \textit{E. coli} B/r porins are synthesized at a constant rate, with a doubling of the rate 10 min before division (Boyd & Holland, 1979). However, very little is known about the factors regulating the synthesis of porins and other major proteins of the OM. Culture conditions such as the richness of the medium, temperature and salt concentration influence the relative amounts of the major proteins (Lugtenberg \textit{et al.}, 1976; van Alphen & Lugtenberg, 1977), but few data exist about the total amounts present at different growth conditions. The level of another major protein, II*, in the OM of \textit{E. coli} K12 seems to be regulated by a feedback mechanism since partial diploids involving the gene \textit{ompA}, which codes for protein II* (Henning \textit{et al.}, 1976; Manning \textit{et al.}, 1976), do not show any gene dosage effect (Datta \textit{et al.}, 1976). This mechanism regulating the incorporation of OMP into the envelope could simply be determined by the amount of surface available for occupation by these proteins (Lugtenberg \textit{et al.}, 1976; Boyd & Holland, 1979). If this is correct, the amount of the major OMP per unit of cell area would be expected to remain constant independently of the growth rate.

The above prediction can be tested by transferring bacteria from one growth medium to another, in so-called shift experiments. Such studies have previously shown that bacterial size is dependent upon the richness of the medium, in such a way that cell length and width increase as the growth rate increases (Schaechter \textit{et al.}, 1958; Donachie \textit{et al.}, 1976; Grover \textit{et al.}, 1977; Pierucci, 1978; Rosenberger \textit{et al.}, 1978). Therefore, when bacteria are transferred from poorer to richer media (shift-up) or from richer to poorer media (shift-down), cell dimensions change to adapt to the new conditions (Maaløe & Kjeldgaard, 1966; Sloan & Urban, 1976; Loeb \textit{et al.}, 1978). After a shift-up transition, cell division continues at the former rate for some time, while the rate of synthesis of biomass increases almost immediately to that characteristic of the new conditions (Maaløe & Kjeldgaard, 1966; Loeb \textit{et al.}, 1978). This leads to the immediate increase of cell size, probably both in length and width. However, some strains of \textit{E. coli} have been shown to behave in a more complex way, particularly during drastic changes of media. In these conditions, slowing (Loeb \textit{et al.}, 1978) or acceleration (Sloan & Urban, 1976) of cell division can occur after the transition. Shift-down transitions lead to an immediate reduction of the rate of accumulation of biomass, which in some strains can stop completely for a period. The rate of division is affected only after some time in the new medium. Consequently, the average cell size diminishes immediately after the shift-down transition (Maaløe & Kjeldgaard, 1966).

In this paper we report on cell dimensions, both length and diameter, and on the amount of major OMP per unit of surface area when \textit{S. typhimurium} LT2 grows in steady-state conditions and when it is subjected to shift-up and shift-down treatments, in order to investigate the control of OMP synthesis and assembly. Results indicate that the surface density of major OMP changes only slightly during different growth conditions that cause large differences in cell dimensions.

\textbf{METHODS}

\textit{Bacteria, growth conditions and culture transitions.} Wild-type \textit{Salmonella typhimurium} LT2, obtained from the collection of the Department of Microbiology of the University of California in Davis, was employed. Bacteria were grown at 37°C in LB medium or in minimal AB medium (Clark & Maaløe, 1967) supplemented with a source of carbon. As carbon sources, glucose was added at 0.2% (w/v) and glycerol,
sodium acetate or L-alanine at 0-4 % (w/v). Amino acids were added at 25 µg ml⁻¹, vitamins at 2 µg ml⁻¹ and casein hydrolysate at 0-4 % (w/v). In order to obtain steady-state conditions of growth, all bacterial cultures were grown exponentially for at least four generations before beginning the experiments. When necessary, bacteria were diluted in the same prewarmed medium to keep the culture density less than 10⁴ cells ml⁻¹ during the experiments. Growth kinetics were monitored by measuring the absorbance of the culture (at 550 nm for cultures in LB medium and at 450 nm for minimal medium), using a Beckman spectrophotometer, model DU-2 (1 cm light path). The total number of bacteria was determined with a Petroff-Hauser counter, from samples fixed in 0.2 % (v/v) formaldehyde at 0 °C.

Culture transitions were carried out by filtering 150 ml of exponentially grown bacteria at a concentration of 3 x 10⁹ to 5 x 10⁹ cells ml⁻¹ using a Sartorius membrane filter (pore size 0-45 µm), washing the filter with 50 ml of the new medium prewarmed at 37 °C and transferring it to a flask containing 150 ml of the new medium at 37 °C. The whole operation was completed in no more than 3 min.

Electrophoresis of total cell proteins in SDS-polyacrylamide slab gels. Samples were taken from the cultures and KCN was added to a final concentration of 2 mM. Bacteria were washed with 50 mM-Tris/HCl pH 6-8 at 0 °C and resuspended in the same buffer at 0 °C to obtain a final protein concentration of 3 to 5 mg ml⁻¹. This suspension could be kept at -20 °C for at least 2 months. Proteins from the washed organisms were solubilized by boiling them for 2 min in a solution containing 2 % (w/v) SDS, 5 % (v/v) mercaptoethanol, 10 mM-EDTA, 0-015 % (w/v) Bromophenol Blue and 5 % (w/v) glycerol in 0-5 M-Tris/HCl pH 6-8. Samples (10 µl, corresponding to about 20 µg protein) were then applied to each slot of an SDS-polyacrylamide gel. Preparation of the gels and buffer system were as described by Laemmli (1970). The concentration of the stacking gel was 5 % (w/v) acrylamide and 0-15 % (w/v) bisacrylamide, while that of the separating gel was 10 % acrylamide and 0-3 % bisacrylamide. Electrophoresis was carried out at a constant current of 15 mA through the stacking gel and 30 mA through the separating gel. Gels were stained for 1 h in a solution of 0-05 % (w/v) Coomassie Blue in 25 % (v/v) isopropyl alcohol plus 10 % (v/v) acetic acid. Destaining was done by three successive washings for 1 h periods in the following solutions: (i) 0-005 % Coomassie Blue in 10 % isopropyl alcohol plus 10 % acetic acid, (ii) 0-0025 % Coomassie Blue in 10 % acetic acid and (iii) 10 % acetic acid. Stained gels were scanned using a Pye Unicam densitometer, model SP1809, and the relative areas under the peaks corresponding to the proteins being studied were measured. Absolute amounts of individual proteins per cell were calculated considering the relative areas of the peaks and the value of total protein per cell. Protein concentration was calculated by the Lowry method, using a 50 mM-Tris/HCl pH 6-8 solution as standard.

In all the electrophoresis experiments the following molecular weight standards were run in parallel: bovine pancreatic RNAase (125000), egg white lysozyme (14000), human γ-globulin L (215000), ovalbumin (43000), human γ-globulin H (50000) and bovine serum albumin (68000).

Fractionation of cells and protein analysis of fractions. About 2 x 10⁸ bacteria were resuspended in 2 ml of ice-cold 10 mM-Tris/HCl pH 7-6 containing 0-75 M-sucrose. Lysozyme was added to a final concentration of 0-5 mg ml⁻¹ and, after 3 min on ice, EDTA was added dropwise to a final concentration of 5 mM. The suspension was incubated on ice for 2 h with stirring and the resulting spheroplasts were lysed by addition of 3 vol. of cold distilled water. Unbroken cells were removed by low speed centrifugation and the supernatant was again centrifuged (48000g, 2 h). The resulting supernatant was kept as the fraction containing cytoplasmic proteins, while the envelope pellet was washed once in 50 mM-Tris/HCl pH 6-8 and separated into inner and outer membrane fractions by treatment with Sarkosyl NL97 at 0-5 % (w/v) in a final volume of 100 µl (Filip et al., 1973). After 30 min incubation at room temperature inner membrane proteins were solubilized, while outer membranes remained insoluble and were subsequently sedimented by centrifugation at 9 °C (48000g, 2 h). The outer membrane pellet was resuspended in 100 µl 50 mM-Tris/HCl pH 6-8. Proteins from the different fractions were analysed by SDS-PAGE, using the method described above for total cell lysates.

Measurement of cell dimensions. Samples (2 ml) from the cultures were fixed with formaldehyde at a final concentration of 2 % (v/v) and kept at 0 °C for 1 h. They were washed with 0-2 % formaldehyde (5 ml) and resuspended in 10 µl of the same solution. A small drop of the suspension was placed on a Formvar film, allowed to dry, stained with 1 % (w/v) phosphotungstic acid and the bacteria were observed with a Hitachi electron microscope, model HU-12A, with an enlargement of 3000. Electron micrographs were taken and the measurement of cell dimensions (length and diameter) was done on the corresponding prints with a final enlargement of 10000. At least 100 cells from each sample were measured. In order to calculate surface area and volume, bacteria were considered as rods with two hemispherical caps.

Chemicals. Sugars, amino acids, vitamins and inorganic chemicals were obtained from Merck. Casein hydrolysate was from Oxoid, acrylamide was from Cambrian Chemicals and bisacrylamide from Eastman Kodak.
Table 1. Cell dimensions of S. typhimurium LT2 at different growth rates

Media used were as follows: 1: LB medium. 2: AB medium plus casein hydrolysate (0.4%, w/v), glucose (0.2%, w/v), tryptophan (25 μg ml⁻¹), adenine, cytosine, guanine, uracil and thymidine (each 20 μg ml⁻¹) and thiamin and biotin (each 2 μg ml⁻¹). 3: AB medium plus casein hydrolysate (0.4%, w/v), glucose (0.2%, w/v) and tryptophan (25 μg ml⁻¹). 4: AB medium plus glucose (0.2%, w/v) and methionine, histidine, arginine, proline, threonine and tryptophan (each 25 μg ml⁻¹). 5: AB medium plus glucose (0.2%, w/v). 6: AB medium plus glycerol (0.4%, w/v). 7: AB medium plus sodium acetate (0.4%, w/v). 8: AB medium plus l-alanine (0.4%, w/v).

All values are means of at least two different experiments, the values of individual experiments being represented in Fig. 1(a). Values of length and diameter are the average of at least 100 cells measured in each experiment. The numbers in the parentheses correspond to the standard deviation.

<table>
<thead>
<tr>
<th>Medium</th>
<th>Doubling rate (h⁻¹)</th>
<th>A₄₅₀ per 10⁹ cells</th>
<th>Length (μm)</th>
<th>Diameter (μm)</th>
<th>Surface area (μm²)</th>
<th>Volume (μm³)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>2.40</td>
<td>4.06</td>
<td>2.80 (0.727)</td>
<td>0.731 (0.054)</td>
<td>5.43</td>
<td>1.073</td>
</tr>
<tr>
<td>2</td>
<td>2.18</td>
<td>3.55</td>
<td>2.69 (0.720)</td>
<td>0.676 (0.051)</td>
<td>5.71</td>
<td>0.885</td>
</tr>
<tr>
<td>3</td>
<td>1.85</td>
<td>3.27</td>
<td>2.60 (0.676)</td>
<td>0.650 (0.049)</td>
<td>5.31</td>
<td>0.791</td>
</tr>
<tr>
<td>4</td>
<td>1.60</td>
<td>2.44</td>
<td>2.52 (0.597)</td>
<td>0.598 (0.039)</td>
<td>4.73</td>
<td>0.651</td>
</tr>
<tr>
<td>5</td>
<td>1.15</td>
<td>1.91</td>
<td>2.31 (0.514)</td>
<td>0.581 (0.039)</td>
<td>4.21</td>
<td>0.560</td>
</tr>
<tr>
<td>6</td>
<td>0.77</td>
<td>1.25</td>
<td>2.06 (0.464)</td>
<td>0.546 (0.041)</td>
<td>3.54</td>
<td>0.441</td>
</tr>
<tr>
<td>7</td>
<td>0.56</td>
<td>1.07</td>
<td>1.80 (0.433)</td>
<td>0.528 (0.035)</td>
<td>2.98</td>
<td>0.355</td>
</tr>
<tr>
<td>8</td>
<td>0.31</td>
<td>0.88</td>
<td>1.82 (0.417)</td>
<td>0.507 (0.031)</td>
<td>2.90</td>
<td>0.334</td>
</tr>
</tbody>
</table>

RESULTS

Cell dimensions and amount of major OMP in exponentially growing S. typhimurium LT2

The dependence of cell length and diameter on the growth rate in S. typhimurium LT2 was studied. Exponentially growing cells were both longer and wider in rich media than in poorer media, in the eightfold range of growth rates studied (Table 1). This confirmed results previously obtained by others with several strains of S. typhimurium and E. coli (Schaechter et al., 1958; Donachie et al., 1976; Grover et al., 1977; Pierucci, 1978; Rosenberger et al., 1978). In Fig. 1(a) cell size, measured both as the real cell volume and as the index of A₄₅₀ per 10⁹ cells, is plotted against the doubling rate. Each point corresponds to a different growth experiment. It can be seen that both size parameters are directly proportional to the growth rate and that the proportionality constant is approximately the same for both. That is, the ratio of A₄₅₀ per unit of volume remains constant, independently of cell dimensions and, therefore, the absorbance seems to be a measure of cell volume. However, the variation of cell size with respect to growth rate in the strain studied here was less than that observed in other strains. Whilst at low growth rates the values indicated in Table 1 are similar to those reported for other bacteria, at higher growth rates the cells were not so large. Thus, we observed a near fourfold change in cell volume over the range from 0.31 to 2.40 doublings h⁻¹, while other strains of S. typhimurium (Schaechter et al., 1958) or E. coli B/r (Rosenberger et al., 1978) showed a five- to sevenfold change in volume over the same range of growth rates.

The amount of protein per cell in S. typhimurium LT2 also increased proportionally to growth rate in the whole range of conditions studied (Fig. 1b). However, whilst the amount of protein per unit of cell volume increased in proportion to growth rate at growth rates less than 1.5 doublings h⁻¹, it remained constant at higher growth rates. This fact is in accordance with the observed constancy in the ratio protein/DNA (Maaløe & Kjeldgaard, 1966; Dennis & Bremer, 1974; Shen & Bremer, 1977) and in the amount of DNA per unit of cell volume (Kubitschek, 1974) in rapidly growing bacteria.

The amount of major OMP per unit of surface area in cells growing in different media was next measured in order to determine if it was in any way dependent upon cell size and shape. Total cell proteins were analysed by SDS-PAGE and the relative area under the
Surface density of outer membrane proteins

Fig. 1. Relationship between growth rate and several cellular properties in exponentially growing cells of Salmonella typhimurium LT2. Bacteria were grown exponentially at 37 °C for more than four generations in the media indicated in the legend to Table 1. (a) A_{450} per 10^9 cells (○), surface area (●) and cell volume (□). (To obtain the A_{450} per 10^9 cells from bacteria grown in LB medium, culture samples were fixed at 0 °C, filtered and resuspended in AB medium.) (b) Total protein per cell (○), total protein per unit of cell volume (●) and major OMP per unit of surface area (□).

four peaks in the 33 to 36 kilodalton region was measured from gel scans and taken as the proportion of major OMP relative to total protein. This approximation could be justified because the major OMP appear to be amongst the most abundant proteins at different growth rates (Pedersen et al., 1978). Thus, any other band appearing in this region should constitute only a minor proportion. To confirm this, proteins from different cell fractions were analysed separately by SDS-PAGE. Figure 2 shows the gel scans from cells growing in minimal medium. Porins are described as 34 to 36K and the 33000 kilodalton protein as 33K. Cytoplasmic proteins migrating in the 33 to 36K region accounted for only 5% of those in total cell lysates. A minor amount of the 33, 34, 35 and 36K proteins appeared in gels corresponding to inner membranes, due to the fact that Sarkosyl solubilizes a small proportion of OMP (Filip et al., 1973; Boyd & Holland, 1979). However, two-dimensional gel electrophoresis confirmed that those minor peaks corresponded to the major OMP. Results from cells grown in other media were similar, therefore confirming that no less than 95% of the area under the peaks in the 33 to 36K region is due to the four major OMP. In a second control, the relative area under the 33 to 36K peaks was found to be independent of the concentration of protein in the solubilization mixture in the range 0.5 to 3.5 μg protein μl⁻¹, indicating that solubilization of major OMP by SDS was not influenced by the amount of total protein.

As indicated in Table 2, the proportion of major OMP with respect to total protein was lower in rich media than in poorer media. However, the density of the four major OMP per unit of surface area did not significantly change in the different growth conditions (Fig. 1b and Table 2). When the porins and the 33K protein were analysed separately, the ratio of one to the other did not change over the whole range of growth rates tested, porins representing about 45% of the major OMP (Table 2). The results obtained correspond to 0.9 × 10^5 to 1.2 × 10^6 molecules of major proteins per μm² of OM. It has been proposed that lipopolysaccharide molecules are associated with porin molecules in the OM (DiRienzo et al., 1978). The number of lipopolysaccharide molecules per μm² of OM has been reported as 3.5 × 10^5 (Smit et al., 1975; Nikaido & Nakae, 1979), that is, about eight times that of porins reported here. The lack of dependence of the amount of porins on growth rate is in accordance with results obtained recently with E. coli B/r (Boyd & Holland, 1979).
Fig. 2. Scans of proteins from different fractions of *S. typhimurium* LT2 cells grown in glucose minimal medium at 37 °C. Proteins were subjected to SDS-PAGE and stained gels were scanned as described in the text. (a) Total cell lysates, (b) cytoplasmic fraction, (c) inner membranes and (d) outer membranes. The amount of protein loaded in each slot was obtained from a similar number of cells. Only the region of the gel where the major OMP migrated is represented; the 43K peak corresponds to the EF-Tu protein.

Table 2. Amount of total protein and major OMP in *S. typhimurium* LT2 at different growth rates

Growth media are given in Table 1. Results are the mean of at least two different experiments and the values of individual experiments are represented in Fig. 1(b).

The protein per volume was calculated from the values of average cell volume (indicated in Table 1) and the amount of protein per cell. The proportion of individual proteins with respect to total protein was obtained as indicated in the text. The major OMP per surface area was calculated from the ratio of major OMP to total protein, the amount of protein per cell and the average cell surface.

<table>
<thead>
<tr>
<th>Doubling rate (h⁻¹)</th>
<th>10⁵ × Protein per cell (µg cell⁻¹)</th>
<th>10⁵ × Protein per volume (µg µm⁻³)</th>
<th>Porins per total protein (%)</th>
<th>33K protein per total protein (%)</th>
<th>Porins/ Major OMP</th>
<th>Major OMP per total protein (%)</th>
<th>10⁵ × Major OMP per surface area (µg µm⁻²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>2:40</td>
<td>5.21</td>
<td>4.86</td>
<td>3.2</td>
<td>3.4</td>
<td>0.49</td>
<td>6.6</td>
<td>5.34</td>
</tr>
<tr>
<td>2:18</td>
<td>4.34</td>
<td>4.90</td>
<td>3.1</td>
<td>4.3</td>
<td>0.42</td>
<td>7.4</td>
<td>5.58</td>
</tr>
<tr>
<td>1:85</td>
<td>3.84</td>
<td>4.85</td>
<td>3.6</td>
<td>4.8</td>
<td>0.43</td>
<td>8.4</td>
<td>6.07</td>
</tr>
<tr>
<td>1:60</td>
<td>3.54</td>
<td>5.44</td>
<td>3.7</td>
<td>5.3</td>
<td>0.41</td>
<td>9.0</td>
<td>6.72</td>
</tr>
<tr>
<td>1:15</td>
<td>2.33</td>
<td>4.16</td>
<td>4.5</td>
<td>6.2</td>
<td>0.42</td>
<td>10.7</td>
<td>5.84</td>
</tr>
<tr>
<td>0:77</td>
<td>1.52</td>
<td>3.45</td>
<td>5.8</td>
<td>5.8</td>
<td>0.50</td>
<td>11.6</td>
<td>5.00</td>
</tr>
<tr>
<td>0:56</td>
<td>1.24</td>
<td>3.45</td>
<td>5.8</td>
<td>7.1</td>
<td>0.45</td>
<td>12.9</td>
<td>5.19</td>
</tr>
<tr>
<td>0:31</td>
<td>0.99</td>
<td>2.97</td>
<td>6.7</td>
<td>9.0</td>
<td>0.43</td>
<td>15.7</td>
<td>5.33</td>
</tr>
</tbody>
</table>

**Effect of cyclic AMP on the amount of major OMP**

A catabolite repression effect on several OMP of *E. coli* has been observed (Aono *et al.*, 1978; Alderman *et al.*, 1979), although no results for major OMP have been described. Therefore, we considered the possibility that the level of major proteins in the OM in the different conditions of growth was regulated by cyclic AMP. However, no effect of catabolite repression on major OMP was observed, since cyclic AMP at 1 mM did not alter the relative proportion of these proteins (Table 3), although bacteria grew more slowly and the total amount of protein per cell was reduced in the presence of cyclic AMP in both LB medium and minimal medium plus glucose. This latter effect was not observed in minimal medium plus acetate.
Table 3. Effect of cyclic AMP on the relative amount of major OMP in *S. typhimurium* LT2 growing in different media

Results in the absence of cyclic AMP are the same as in Table 2.

Without added cyclic AMP | With 1 mM-cyclic AMP
---|---
**Doubling rate** (h⁻¹) | **Doubling rate** (h⁻¹)
**10⁷ × Protein per cell** (µg cell⁻¹) | **10⁷ × Protein per cell** (µg cell⁻¹)
**Major OMP per total protein** (%) | **Major OMP per total protein** (%)

<table>
<thead>
<tr>
<th>Medium</th>
<th>2.40</th>
<th>5.21</th>
<th>6.6</th>
<th>2.00</th>
<th>4.5</th>
<th>6.6</th>
</tr>
</thead>
<tbody>
<tr>
<td>LB</td>
<td>1.15</td>
<td>2.33</td>
<td>10.7</td>
<td>0.98</td>
<td>2.12</td>
<td>10.6</td>
</tr>
<tr>
<td>AB plus glucose</td>
<td>0.56</td>
<td>1.24</td>
<td>12.9</td>
<td>0.95</td>
<td>1.42</td>
<td>12.3</td>
</tr>
</tbody>
</table>

**Changes in major OMP surface density during a shift-up transition**

We next studied the effect of medium transitions on the amount of major OMP and the parallel changes in cell dimensions and, correspondingly, in surface area. As major OMP accumulation was not affected by catabolite repression we assumed that alterations in the former during a shift should be exclusively due to the mechanisms which regulate the accumulation of those proteins in the OM in relation to cell size.

Figure 3(a) shows the effect of a shift-up from minimal medium plus glucose (doubling time of 50 to 55 min) to LB medium in *S. typhimurium* LT2. Absorbance began to increase immediately at higher growth rate than in the pre-shift medium, although the doubling time (29 min) throughout the experiment was slightly longer than in steady-state cultures (24 to 25 min). Bremer & Dennis (1975) have predicted – on a theoretical basis – a delay in the attainment of the steady-state conditions after a shift-up, but these were not reached in the shifts reported. Changes in cell number followed a more complex pattern, with a delay in the rate of cell division between 10 and 25 min after the shift. After this interval all the parameters increased in parallel.

Figure 3(b) shows the effect of a more drastic change from minimal medium plus acetate (doubling time of 105 to 115 min) to LB medium. The pattern of changes in the different parameters was similar to that following the shift from glucose to LB medium, although the increase in absorbance did not immediately adjust to that equivalent to LB medium. During the first 50 min after the shift, the doubling time was 41 min and after that a doubling time of 29 min was achieved, still longer than that of steady-state conditions. Cell number increased during the first 10 min after the transfer, followed by a lag before resuming at a rate which paralleled absorbance. Both total protein and major OMP also showed a lag of 20 to 25 min in their accumulation, followed by a sharp increase until reaching an amount per cell close to that characteristic of the new medium.

Changes in cell dimensions in both shifts were followed by direct measurement of electron micrographs of cells from samples taken at different times. Figure 4(a) shows the changes in cell length and diameter during both shifts. It should be noted that cells became smaller immediately after the transfer, so cell dimensions at time 0 in Fig. 4 are not comparable with those of steady-state conditions in the original media; this change in cell dimensions after transfer has also been observed by Woldringh et al. (unpublished observations) in shift-up experiments not involving filtration and may be due to the effect of osmotic stress on the cells. Whereas in the acetate to LB medium shift 80 min were required to stabilize cell dimensions, in the glucose to LB medium shift no stabilization of these parameters was reached after 80 min from the transition. The most interesting result was that in each case length and diameter did not begin to increase immediately after the shift. There was a lag period of 20 min during which cell dimensions were those of the previous medium. This period coincided with the deceleration of cell division previously described (see Fig. 3).
Fig. 3. Effect of shift-up transitions from (a) AB medium plus 0·2% (w/v) glucose, or (b) AB medium plus 0·4% (w/v) acetate, to LB medium on the pattern of cell division and on the amount of total cell protein and major OMP. *Salmonella typhimurium* LT2 was grown exponentially at 37 °C in the original medium to a density of about 5 × 10^8 bacteria ml^-1; at time 0, a sample of cells was filtered, washed with prewarmed LB medium and resuspended in the original volume of LB medium at 37 °C. At intervals, samples were taken from the culture and the following measurements were done: \( A_{560} \) (○), total number of cells per ml (●), total cell protein per ml (□) and major OMP per ml (■).

Fig. 4. (a) Changes in cell length (○, □) and diameter (●, ■) after shift-up transitions of *S. typhimurium* LT2 from glucose minimal medium (○, ●) or acetate minimal medium (□, ■) to LB medium. Transitions were done as described in the legend to Fig. 3. Relative values of 1·0 correspond to the following absolute values: ○, 2·23 μm; ●, 0·518 μm; □, 1·56 μm; ■, 0·464 μm. (b) Changes in the major OMP per unit of cell surface after shift-up transitions from glucose minimal medium (○) or acetate minimal medium (●) to LB medium. Relative values of 1·0 correspond, respectively, to absolute values of 7·99 × 10^{-8} and 9·07 × 10^{-9} μg major OMP per μm^2 of cell surface. Bars with the corresponding symbols (on the right) indicate the levels of the different parameters in steady-state cultures in LB medium (see Tables 1 and 2).
Surface density of outer membrane proteins

Fig. 5. Effect of shift-down transitions from LB medium to (a) AB medium plus 0.2% (w/v) glucose, or (b) AB medium plus 0.4% (w/v) acetate, on the pattern of cell division and the amount of total cell protein and major OMP. *Salmonella typhimurium* LT2 was grown exponentially at 37°C in LB medium; at time 0, a sample of cells was filtered, washed with prewarmed medium and resuspended in the original volume of new medium at 37°C. At intervals samples were taken from the culture and the following measurements were done: $A_{450}$ ($\bigcirc$), total number of cells per ml ($\bigcirc$), total cell protein per ml ($\square$), and major OMP per ml ($\blacksquare$).

The close correspondence between the accumulation of total protein and of major OMP during both shift-up transitions (Fig. 3) indicated that the synthesis of major OMP followed the same pattern as that of total protein. As cell dimensions did not follow the same kinetics, there was a transitory uncoupling between the amount of major OMP and cell surface. A plot of the major OMP surface density against time after the shift indicates that the former value did not remain constant (Fig. 4b). During the abrupt shift from acetate to LB medium, the density of major OMP apparently increased by more than 50% before reaching the equilibrium value about 80 min after the shift (this value was about the same as in steady-state conditions). This sharp increase was mainly due to the slow increase in cell dimensions compared with the rapid acceleration in the accumulation of major OMP at around 40 min after the shift (Fig. 4b). Only when cell length and diameter reached the final stable value did major OMP density also stabilize.

Changes in major OMP surface density during a shift-down transition

The effect of a mild (LB to glucose minimal medium) and abrupt (LB to acetate minimal medium) shift-down on the change of cell dimensions and the accumulation of major OMP was also studied. Shift-down transitions are more difficult to interpret than shift-up transitions, because more or less prolonged transitory inhibitions of metabolic processes can occur, which would mask or indirectly influence the phenomena being studied.

Results of a transition from LB to glucose minimal medium are shown in Fig. 5(a). Absorbance and cell number followed a simple pattern. The latter increased initially at a rate characteristic of the former medium and it achieved the rate corresponding to the new medium only after 40 min from the transition time. Total protein and major OMP accumulated in parallel after a 20 min lag. Shift-down from LB to acetate minimal medium (Fig. 5b) showed a slightly different pattern. There was no parallelism between absorbance and cell number until 70 min after the shift. In contrast to the shift from glucose minimal medium, total protein and major OMP accumulation did not completely stop after the transfer. The reasons for this difference are unknown.
Fig. 6. (a) Changes in cell length (○, □) and diameter (●, □) after shift-down transitions of *S. typhimurium* LT2 from LB medium to glucose minimal medium (○, ●) or acetate minimal medium (□, □). Transitions were done as described in the legend to Fig. 5. Relative values of 1·0 correspond to the following absolute values: ○, 2·71 μm; ●, 0·736 μm; □, 2·65 μm; □, 0·743 μm. (b) Changes in the major OMP per unit of cell surface after shift-down transitions from LB medium to glucose minimal medium (○) or acetate minimal medium (●). Relative values of 1·0 correspond to absolute values of 5·49 x 10⁻⁸ and 7·13 x 10⁻⁸ μg major OMP per µm² of cell surface, respectively. Bars with the corresponding symbols (on the right) indicate the steady-state levels of the different parameters in the respective media (see Tables 1 and 2).

In contrast to experiments involving shift-up, cell dimensions began to change immediately after the shift-down (Fig. 6a), decreasing to a final cell size in which cells were shorter than in steady-state conditions. The density of major OMP in the surface gradually increased in both shifts, the values eventually reached being higher than those observed during steady-state growth in the same media (Fig. 6b). This may reflect an imbalance between accumulation of major OMP, which is parallel to total protein accumulation, and the smaller cell size attained in these shift-down conditions.

**DISCUSSION**

Since the bacterial cell size depends on the growth rate, we have studied both the amount of major OMP in steady-state conditions and its evolution when the envelope area per cell is modified after transition between different media, as an approach to understanding the regulation of major OMP synthesis. The level of major OMP was obtained after analysing total cell lysates by SDS–PAGE. It is known from studies with toluene-treated cells that major OMP are synthesized in the form of a larger precursor which is then processed to the mature form (Sekizawa *et al.*, 1977). However, this precursor does not exist *in vivo*, so its processing and incorporation into the OM must accompany its synthesis. Therefore, all the major OMP observed in total cell lysates must correspond to molecules incorporated into the OM.

The observed constancy in major OMP surface density supports the idea that the formation of a regular pattern of molecules of these proteins in the OM is controlled by the amount of surface area available. Although the cross-sections of the major OMP are not known, the calculated number of molecules of major OMP per unit area (0·9 x 10⁵ to
1.2 \times 10^5 \text{ per } \mu m^2 should imply that a considerable fraction of the OM is covered by major OMP, confirming electron microscope studies (Smit et al., 1975; Steven et al., 1977). It has been shown in *E. coli* B/r that the number of lipopolysaccharide (LPS) molecules per unit area increases with growth rate (Zaritsky et al., 1979). No numerical data exist on phospholipids, although Cronan & Vagelos (1972) reported that the amount of phospholipid with respect to dry weight is constant at different growth rates. As the dry weight per cell increases with growth rate while the ratio of surface area per dry weight decreases (Maaløe & Kjeldgaard, 1966), the constancy in the proportion of phospholipids should imply an increase in phospholipid surface density with increasing growth rates. These facts suggest that the amount of other OMP should decrease at high growth rates because less physical space should be available. LPS-defective mutants of *S. typhimurium* are known which also have decreased amounts of OMP per unit of surface area, this defect being compensated by an increasing amount of phospholipid (Muhlradt et al., 1974; Smit et al., 1975). Also, some LPS-chain defective mutants increase the number of LPS molecules per area while reducing that of proteins (Gmeiner & Schlecht, 1979). All these data, therefore, point to a competition between OMP molecules, on the one hand, and LPS and phospholipid, on the other, for the physical space in the OM.

Shift-up and shift-down transitions with the strain employed here showed some differences with respect to other strains. Thus, during the shift-up transitions, cell dimensions began to increase only 20 min after the shift. Others have reported that cell size increases immediately after the transition (Maaløe & Kjeldgaard, 1966; Sloan & Urban, 1976). The discrepancy could be due to differences between strains or between the methods of measuring cell size: we have measured cell dimensions directly from electron micrographs. A delay has also been observed in *E. coli* B/r using similar methods to those employed here (Woldringh et al., unpublished observations). The increase in cell dimensions at a particular time after the shift could be due to the increase in the number of sites for envelope growth or to the increase in the rate of accumulation of envelope material into a fixed number of sites. Either of these possibilities raises the question of the cell event that accelerates the growth of the envelope. During steady-state conditions cell elongation seems to be linear with a doubling in rate at a time of the cell cycle (Pritchard, 1974; Donachie et al., 1976; Pierucci, 1978; Rosenberger et al., 1978). It has been proposed that this time is related to the attainment of a critical maximum density (Rosenberger et al., 1978). During the shift-up, cell density probably increases after the transition, because the immediate increase in the rate of absorbance increase is not accompanied by an increase of mean cell volume (Figs 3 and 4). A critical value of density would be achieved after a time, which would induce an increase in the number of sites of envelope growth or in the rate of envelope growth at each site. Interestingly, shortly after the transition to the new medium there is a lag in cell division. This period, which also has been observed in *E. coli* strains (Loeb et al., 1978), occurs immediately before and after initiation of cell size increase. This is consistent with the idea that formation of cell constrictions leading to division might be due to an excess of envelope produced after doubling of the rate of envelope elongation (Pritchard, 1974; Rosenberger et al., 1978). If bacteria employ the envelope material in increasing both length and width there would be no excess of envelope and no division would occur. Division would resume only after cell density had fallen below a critical level as a consequence of the increase in cell volume.

With respect to major OMP accumulation after shifts, the results show that it follows the same pattern as total cell protein. As the latter does not immediately adjust to changes in cell volume, there is a fluctuation in the ratio of major OMP to cell surface, more pronounced in the shift from acetate minimal medium to LB medium. What happens to the OM structure during the shifts is not known. However, after the shift from acetate minimal medium to LB medium, the density of major OMP increases more than 50\% over the initial and final values. As these proteins can cover more than 60\% of the external OM surface
We thank Drs I. B. Holland and A. Zaritsky for critically reading the manuscript. M.A. is a recipient of a grant from the Spanish Ministry of Universities and Research.

REFERENCES


Surface density of outer membrane proteins

LUGTENBERG, B., MEIJERS, J., PETERS, R., HOEK, P.


