Isolation and Characterization of the Tubular Organelles Induced by Fumarate Reductase Overproduction in *Escherichia coli*

By M. LYNN ELMES, DOUGLAS G. SCRABA AND JOEL H. WEINER

Department of Biochemistry, University of Alberta, Edmonton, Alberta T6G 2H7, Canada

(Received 7 October 1985; revised 30 December 1985)

Strains of *Escherichia coli* amplifying the intrinsic membrane enzyme fumarate reductase accommodate the overproduced enzyme by increasing the amount of membrane material, in the form of intracellular tubular structures. These tubules have been observed in strains harbouring multicopy *frd* plasmids and in ampicillin hyper-resistant strains. A procedure has been developed for isolation of tubules nearly free of cytoplasmic membrane. Using protein A–gold labelling and optical diffraction of electron micrographs, a model for tubule structure is proposed. The tubules have a lower lipid/protein ratio than the cytoplasmic membrane, with the enzyme accounting for greater than 90% of the protein in the tubules. Both cytoplasmic membranes and tubules from amplified strains are enriched in cardiolipin and have a more fluid fatty acid composition than wild-type strains. Mutants defective in cardiolipin synthesis produce tubules in response to excess fumarate reductase, but these tubules have an altered appearance, indicating that lipid–protein interactions may be important for tubule assembly.

**INTRODUCTION**

Fumarate reductase is a complex iron–sulphur flavoenzyme which serves as the terminal electron transfer enzyme when *Escherichia coli* is grown anaerobically on fumarate-containing medium (Haddock & Jones, 1977). The enzyme, composed of a membrane-extrinsic catalytic domain and a membrane-intrinsic anchor domain, consists of four non-identical subunits in equimolar amounts (Lemire et al., 1982, 1983). The expression of fumarate reductase is greatly amplified in strains harbouring multicopy *frd* plasmids (Cole & Guest, 1980; Grundström & Jaurin, 1982; Lemire et al., 1982; Lohmeier et al., 1981) and in ampicillin hyper-resistant strains due to tandem duplications of the chromosomal *amp-frd* region (Cole & Guest, 1979a, b). In amplified strains, the enzyme can account for up to 70% of the inner membrane protein (Weiner et al., 1984). The cells respond to the excess fumarate reductase by increasing phospholipid biosynthesis, thus maintaining a relatively constant lipid/protein ratio (Weiner et al., 1984). The excess lipid and protein is organized in long tubular structures within the cytoplasm of the host bacteria (Lemire et al., 1983; Weiner et al., 1984).

In a recent paper (Weiner et al., 1984), we examined the lipid and protein composition of a fumarate reductase-enriched crude membrane fraction composed of tubules and cytoplasmic membrane. A large increase in the cardiolipin content, as well as an increase in fatty acids of lower melting temperature, was observed. In this paper we describe a method for isolating an enriched tubule fraction and report a structural and chemical characterization. Tubule formation in a cardiolipin-biosynthesis defective mutant and an *amp-frd* tandem duplication strain has also been examined.

**METHODS**

*Strains and plasmids.* *E. coli* HB101 (F− HsdR− HsdM− Pro− Leu− Gal− Lac− Thi− RecA−) was from our laboratory stock collection. *E. coli* G800 (Ilv− MetB− *ampA1*) was provided by Professor J. R. Guest (University of Sheffield, UK). *E. coli* T1GP (F− Cls− MdoA− Ilv− Met− LacI−) was provided by Dr E. P. Kennedy (Harvard University, USA).
Medical School, Cambridge, Mass., USA). Plasmid pFRD63, carrying the frd operon in pBR322, has been described previously (Lemire et al., 1982).

**Growth conditions.** Cells were grown to late stationary phase (72 h) anaerobically on glycerol fumarate medium (Spencer & Guest, 1974) in medium containing 100 µg ampicillin ml⁻¹ and necessary vitamin and amino acid requirements. G800 was grown in medium containing 800 µg ampicillin ml⁻¹.

**Transformation of T1GP.** E. coli T1GP cells were treated with calcium and transformed with pFRD63 DNA as described previously (Lemire et al., 1982). Transformants were selected on Luria broth agar plates, containing 100 µg ampicillin ml⁻¹.

**Isolation of tubules and membranes.** Late stationary phase cells (1 g wet weight) were washed once with 50 mM-Tris/HCl, pH 7.5, and suspended in 20 ml 50 mM-Tris/HCl, pH 7.5, 20% (w/v) sucrose (see Fig. 1). Lysozyme was added to 200 µg ml⁻¹. The solution was swirled slowly at 37 °C for 30 min, and centrifuged at 10000 g for 10 min. EDTA was not routinely added, as it was not essential for spheroplast formation (Owen et al., 1982) under the conditions used. This procedure resulted in some cell lysis; thus the supernatant contained cytoplasmic constituents, including a portion of the tubules. The spheroplasts and broken cells were suspended in a minimal volume of 50 mM-Tris/HCl, pH 7.5, containing 20% (w/v) sucrose, and the spheroplasts lysed by rapid dilution into 40 vols 50 mM-potassium phosphate buffer, pH 6.6, followed by stirring for 10 min at room temperature. DNAase was added to 20 µg ml⁻¹, and the suspension was put on ice for 10 min and then centrifuged at 10000 g for 10 min. The supernatant from the broken cell preparation also contained tubules. The broken cells, still containing some tubules, were suspended in 5 ml 50 mM-potassium phosphate buffer/0.05% deoxycholate and stored at −20 °C for 18 h. The suspension was thawed and centrifuged at 10000 g for 10 min. The residual tubules...
were removed from the cells by repeating the deoxycholate–freezing steps two to four times. The tubules were collected from all of the low-speed supernatants by centrifugation at 100 000 g for 1 h at 4°C and suspended in 1 to 2 ml Tris/HCl. The supernatant from the high-speed spin contained small tubule fragments, enzyme aggregates and soluble fumarase reductase.

Lyced spheroplasts were suspended in 4 ml 50 mM-potassium phosphate buffer, pH 6.6, and crude membranes were prepared by two passages through a French pressure cell at 16000 lbf in⁻² (110 MPa). The membranes were collected by centrifugation at 100 000 g for 1 h and suspended in 1 to 2 ml Tris/HCl. The high-speed supernatant from the French press sample contained soluble components.

Tubules and inner membranes were further purified by a modification of the procedure of Yamato et al. (1975). Fractions (1 ml) of tubules or envelopes were layered on 6 ml 44% (w/w) sucrose, 50 mM-Tris/HCl, pH 7.5, and centrifuged at 100 000 g for 1 h at 4°C in a Beckman Ti50 fixed angle rotor. Both tubules and inner membranes formed an amber band near the top of the sucrose layer which was removed with a bent Pasteur pipette. The outer membranes pelleted through the sucrose. The tubules and membrane fractions were suspended in 2 vols 50 mM-Tris/HCl, pH 7.5, collected by centrifugation at 100 000 g for 1 h and each suspended in 1 ml 50 mM-Tris/HCl, pH 7.5, and stored frozen at −70°C.

Enzyme assays. Fumarate- and nitrate-dependent oxidation of reduced benzyl viologen was monitored as previously described (Dickie & Weiner, 1979). One unit of activity corresponds to 1 μmol benzyl viologen oxidized min⁻¹.

ATPase was assayed by measuring the release of inorganic phosphate as described by Owen et al. (1982). One unit corresponds to 1 μmol P<sub>i</sub> released min⁻¹.

Protein determination. Protein was estimated by an SDS modification of the Lowry method (Markwell et al., 1978), using Bio-Rad protein standard.

SDS-PAGE. Electrophoresis on 12 to 17% (w/v) acrylamide gradient gels was done using the Laemmli buffer system (Laemmli, 1970), with the addition of 20% (w/v) sucrose to the lower gel solution. Gels were stained, destained and scanned as described previously (Lemire et al., 1982).

Extraction of membrane lipids. Total lipids were quantitatively extracted from purified cytoplasmic membranes prepared from strains HB101(pBR322), HB101(pFRD63), TiGP(pFRD63), and tubules from strain HB101(pFRD63) by a modification of the method of Bligh & Dyer (1959) as outlined previously (Weiner et al., 1984).

Analysis of fatty acids by GLC. Methyl esters of fatty acids were prepared by transesterification of lipids (Gander et al., 1962) as described previously (Weiner et al., 1984).

Quantification of membrane phospholipids. Individual phospholipids were separated by two-dimensional TLC on glass plates coated with silica gel G (BDH Chemicals Canada) in solvent systems of (i) chloroform/methanol/7 M-ammonium hydroxide (60 : 35 : 5, by vol.) and (ii) chloroform/methanol/acetic acid (65 : 25 : 8, by vol.). Lipids were located by exposure of the plates to iodine vapour. The area corresponding to each lipid was removed and eluted with chloroform/methanol (2 : 1, v/v). Phospholipid phosphorus was determined by the method of Raheja et al. (1973).

Protein A-gold labelling and negative-stain electron microscopy. Samples were labelled with colloidal gold using the procedure of Roth et al. (1978) with the following modifications. The colloidal gold was prepared as described by Frens (1973). The protein A-gold complex was prepared by mixing 9 ml colloidal gold with 1 mg protein A (Sigma) dissolved in 0-1 ml H<sub>2</sub>O, followed by the addition of polyethylene glycol (M, 20000), as outlined by Roth (1983). Samples were diluted with 10 mM-potassium phosphate buffered isotonic saline (PBS; pH 7-4), 1% (w/v) bovine serum albumin (BSA; Miles Laboratories) and allowed to adhere to hydrophilic carbon films on grids. The grids were floated, sample side down, on 100 μl (2-2 μg protein) drops of rabbit anti-fumarase reductase antibody (IgG fraction) raised against catalytic dimer, as described by Lemire et al. (1983), or 10 mM-PBS/BSA (control) for 2 h at room temperature in a moist chamber. The samples were washed twice with 10 mM-PBS/BSA, floated as before on a drop of 10 mM-PBS/BSA for 2 min, washed again with 10 mM-PBS/BSA, and dried by touching the edge of the grid to filter paper. The samples were then floated as before on 100 μl drops of protein A-gold for 1 h. The grids were washed four times with 10 mM-PBS. Samples were negatively stained with 1% sodium phosphotungstate, pH 7-0. The samples were examined in a Philips EM420 electron microscope operated at 100 kV.

Optical diffraction. Selected areas of micrographs of well preserved tubules were printed by contact onto Kodak electron microscope film (SO-163), masked, and their diffraction patterns recorded on a Polaron optical diffractometer (Johansen, 1975). Diffraction spots were measured and used to calculate the lattice dimensions for the tubule model.

RESULTS

Tubule isolation

We have developed a procedure for the separation of the tubular fraction from the inner and outer membrane fractions of E. coli harbouring the frd plasmid (Fig. 1). The procedure involves...
Table 1. Enzyme activities of membranes and tubules from E. coli HB101(pFRD63)

Specific activity values [units (mg protein)$^{-1}$] are means of two determinations. PPB, Potassium phosphate buffer.

<table>
<thead>
<tr>
<th>Source</th>
<th>Fumarate reductase</th>
<th>Nitrate reductase</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Specific activity</td>
<td>ATPase specific activity</td>
</tr>
<tr>
<td></td>
<td>200 mm-PPB (a)</td>
<td>30 mm-HEPES (b)</td>
</tr>
<tr>
<td>Tubules</td>
<td>234</td>
<td>180</td>
</tr>
<tr>
<td>Inner membranes</td>
<td>196</td>
<td>164</td>
</tr>
</tbody>
</table>

* With 50 mm-PPB.

the gentle elution of tubules from a broken cell fraction prepared by lysozyme–deoxycholate–freeze-thaw treatment of stationary phase cells. Lysozyme-lysis was insufficient to release all the tubules, making it necessary to treat the vesicles with a low concentration of deoxycholate (0.05%) followed by freeze-thawing to improve the yield. Approximately 50% of the fumarate reductase activity in the overproducing cells is localized in the tubules. We have been unable to devise a biochemical assay to distinguish tubules from inner membranes but electron microscopy and the SDS-PAGE profile (see Fig. 4) indicate a substantially enriched tubular fraction. Inner membrane contamination, as judged by the detection of low levels of typical inner membrane enzyme activities (Table 1), appears to be in the range of 15–20%. However, it must be noted that the specific activities for the membrane-bound enzymes are reduced in the amplified membranes owing to dilution by the large amount of fumarate reductase (Weiner et al., 1984).

As outlined in Fig. 1, inner membranes are isolated from broken cells after the tubules are removed. The purified inner membranes contain approximately 30% of the total fumarate reductase. The supernatant fraction contains 18% of the fumarate reductase activity (Fig. 1). This activity consists of small fragments of tubules and aggregates of enzyme which do not sediment during centrifugation at 100000 g for 1 h. In this procedure, as with the original cytoplasmic membrane purification procedure of Yamato et al. (1979), the outer membrane fraction was less than 5% contaminated with cytoplasmic membranes.

Properties of fumarate reductase in tubules

Previous reports from this laboratory (Lemire et al., 1982; Robinson & Weiner, 1982) have documented two forms of fumarate reductase: a soluble catalytic dimer composed of the FRDA and FRDB subunits and a tetrameric holoenzyme composed of equimolar amounts of the FRDA, B, C and D subunits. These two enzyme forms can be distinguished by the anion dependence, and the alkali- and thermo-lability of the dimer form. We compared the inner membrane-bound form with the enzyme in the tubules; the two behaved similarly with respect to stability and anion stimulation (Table 1).

Protein A–gold labelling of tubules and vesicles

Tubules and inner membrane vesicles were specifically labelled with protein A–gold, using antibody (IgG fraction) raised against the catalytic dimer of fumarate reductase. The protein A–gold labelling is very specific, with few gold particles attached to non-tubule or non-membranous material (Fig. 2a, b). Control samples labelled with PBS/BSA showed few non-specifically bound gold particles, indicative of non-specific binding (data not shown). The antibody-specific attachment of the gold particles to the tubules confirms that these structures are assemblies of fumarate reductase, with the FRDA and B subunits on the surface.
Overproduction of fumarate reductase

Fig. 2. Negative stain electron micrograph of (a) tubules and (b) inner membrane vesicles, treated as described for protein A–gold labelling. Bars, 200 nm.

**Tubule structure**

We have used negative stain electron microscopy coupled with optical diffraction to develop an approximate model of tubule structure. The tubules are linear cylinders (Fig. 3a) densely covered with 5.3 ± 0.3 nm spherical protein structures packed in a regular helical arrangement of 10 units per turn (Fig. 3b, c). From the model proposed by Lemire et al. (1983), it is presumed that these large knobs on the surface of the tubules are the 69 kDa subunit. Also, the specificity of the protein A–gold labelling with the antibody raised against the catalytic dimer of fumarate reductase indicates that fumarate reductase is arranged in the tubules with the catalytic heads on the surface and with the anchor polypeptides embedded in the lipid interior. This arrangement
is also supported by the observation that treatment of the tubules with 4.5 M-urea, which extracts the 69 and 27 kDa subunits (Lemire et al., 1983) results in removal of the large spherical structures, leaving smooth, slightly swollen tubules (data not shown).

Protein profile of isolated tubules

The isolated tubules had an SDS-PAGE profile (Fig. 4a) similar to that reported for the Triton X-100 extract (Lemire et al., 1982), and highly purified for fumarate reductase compared to the initial cell envelopes (Fig. 4b). Based on the integration of Coomassie blue-stained SDS-polyacrylamide gels of tubules, up to 90% of the protein could be attributed to the four subunits of fumarate reductase. As reported previously (Lemire et al., 1982) fumarate reductase is a four-subunit enzyme with an equimolar ratio. A molar ratio of 1:1:0.8:1 was determined for the enzyme associated with the tubules, indicating that the unique assembly of the enzyme in the tubules is not due to an altered subunit composition. Based on the regular helical packing arrangement of fumarate reductase in the tubules it is unlikely that the contaminating polypeptides seen on the gel in Fig. 4(a) are actually components of the tubules; rather, we believe these represent the small amount of contamination by inner membrane in the preparation.

Lipid:protein ratio of cytoplasmic membrane and tubules

Samples of purified inner membranes and tubules were assayed for protein and lipid content (Table 2). It is apparent that the fumarate reductase-enriched inner membranes and tubules
have a lower lipid/protein ratio than normal membranes (Table 2; Kaback, 1971). In fact, the tubules are over 75% protein, which presumably contributes to the unique structure.

**Phospholipid composition of cytoplasmic membranes and tubules**

The membranes from anaerobically grown *E. coli* HB101 contain phosphatidylethanolamine and phosphatidylglycerol as major phospholipids (Table 2). Crude membranes from strains overproducing fumarate reductase are enriched in cardiolipin; it accounts for 15–20% of the total lipid compared to 4% in normal strains (Weiner et al., 1984). Both isolated tubules and inner membrane fractions are enriched in cardiolipin, although the inner membranes are slightly more enriched (Table 2). These data indicate that the unique structure of the tubules is not due solely to the lipid composition, i.e. is not due to sequestration of the cardiolipin in the tubule fraction.

**Fatty acid composition**

As reported previously (Weiner et al., 1984; see Table 3), the fumarate reductase-enriched inner membranes showed decreases in the saturated fatty acid myristate (14:0), as well as in cyclopropanenonadecanoate (19:cyc), compared to wild-type membranes. The tubules also showed decreases in these fatty acids, and in palmitate (16:0) (Table 3). Small increases were observed in the unsaturated fatty acid palmitoleate (16:1) and in cyclopropaneheptadecanoate (17:cyc) in both tubules and inner membranes (Table 3). The tubules isolated from the overproducing strains had a fatty acid profile slightly different from that of inner membranes from the same cells. The changes in the inner membrane and tubule fatty acid composition would tend to lower the phase transition temperature and fluidize the lipid component (Jain & Wagner, 1980) compared to the membranes of the non-overproducing strain of HB101.

**Effect of a cardiolipin synthesis mutation on tubule structure**

The altered levels of cardiolipin in amplified strains prompted us to examine tubule formation in *E. coli* T1PG, a strain deficient in cardiolipin synthesis (Pluschke et al., 1978). The mutant has only 4% of total lipid as cardiolipin, compared to 20% in normal overproducing strains, and also
Table 2. Phospholipid composition of E. coli membranes containing normal and amplified levels of fumarate reductase

For HB101(pBR322) and HB101(pFRD63) the results of the lipid analyses are means of two determinations. The T1Gp(pFRD63) and G800 lipid analyses were done once. The results for HB101(pBR322) are from Weiner et al. (1984).

<table>
<thead>
<tr>
<th>Strain</th>
<th>Percentage of:</th>
<th>Lipid concn (mg ml⁻¹)</th>
<th>Protein concn (mg ml⁻¹)</th>
<th>Lipid/protein ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PE</td>
<td>PG</td>
<td>DPG</td>
<td>Minor components</td>
</tr>
<tr>
<td>HB101(pBR322)</td>
<td>64</td>
<td>27</td>
<td>9</td>
<td>-</td>
</tr>
<tr>
<td>Total membrane</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>HB101(pFRD63)</td>
<td>62</td>
<td>8</td>
<td>27</td>
<td>3</td>
</tr>
<tr>
<td>Cytoplasmic membrane</td>
<td>63</td>
<td>12</td>
<td>21</td>
<td>4</td>
</tr>
<tr>
<td>Tubules</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T1GP(pFRD63)</td>
<td>88</td>
<td>6</td>
<td>4</td>
<td>2</td>
</tr>
<tr>
<td>Total membrane</td>
<td>75</td>
<td>2</td>
<td>23</td>
<td>-</td>
</tr>
<tr>
<td>G800</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

ND, Not determined.

* PE, phosphatidylethanolamine; PG, phosphatidylglycerol; DPG, diphosphatidylglycerol (cardiolipin).

Table 3. Percentage fatty acid content of anaerobically grown E. coli

The GLC analysis was done three times. The results obtained were similar; representative results are shown. The results for HB101(pBR322) are from Weiner et al. (1984). The fatty acids are designated by the number of carbon atoms followed by the number of double bonds; cyc, cyclopropane.

<table>
<thead>
<tr>
<th>Strain</th>
<th>14:0</th>
<th>15:cyc</th>
<th>16:0</th>
<th>16:1</th>
<th>17:cyc</th>
<th>18:0</th>
<th>18:1</th>
<th>19:cyc</th>
</tr>
</thead>
<tbody>
<tr>
<td>HB101(pBR322)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Total membranes</td>
<td>27:9</td>
<td>1:3</td>
<td>38:4</td>
<td>3:3</td>
<td>15:0</td>
<td>1:0</td>
<td>1:9</td>
<td>11:1</td>
</tr>
<tr>
<td>HB101(pFRD63)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Cytoplasmic membrane</td>
<td>19:8</td>
<td>0:8</td>
<td>38:5</td>
<td>10:1</td>
<td>25:3</td>
<td>0:4</td>
<td>2:6</td>
<td>1:9</td>
</tr>
<tr>
<td>Tubules</td>
<td>17:5</td>
<td>1:4</td>
<td>35:0</td>
<td>10:9</td>
<td>29:5</td>
<td>0:4</td>
<td>2:6</td>
<td>2:0</td>
</tr>
</tbody>
</table>

has lower levels of phosphatidylglycerol. These decreases were compensated by an increase in phosphatidylethanolamine to 88%.

Although the mutant has only 4% cardiolipin, tubules are still formed to accommodate the overproduced fumarate reductase. However, the fumarate reductase is randomly oriented in these tubules (Fig. 5) and is not packed in the helical arrangement seen in HB101(pFRD63) (Fig. 3a). Although the SDS-PAGE profile indicates that the mutant tubules are highly enriched in fumarate reductase (data not shown), the enzyme has a low specific activity of approximately 50 units (mg protein)⁻¹. This low specific activity may be due to the instability of these tubules to freeze-thaw manipulations and to the presence of inactive enzyme.

Tubule formation by an ampicillin hyper-resistant mutant

We have reported the synthesis of a novel tubular organelle in response to high level amplification of plasmid-coded fumarate reductase (Weiner et al., 1984). G800, a β-lactamase-hyper-producing strain of E. coli that is resistant to 800 μg ampicillin ml⁻¹, has amplified levels of fumarate reductase due to tandem duplication of the amp-frd region of the chromosome (Cole & Guest, 1979a, b). We have now found that this method of amplification also results in tubule formation. When the cytoplasmic membrane-binding capacity for fumarate reductase becomes saturated, the G800 cells produce the long tubular assemblies of helically packed enzyme and lipid. During the exponential phase of growth (24 h) these structures appear indistinguishable from the tubules observed in strains harbouring a frd multicopy plasmid (Weiner et al., 1984), as judged by negative stain electron microscopy of lysed cells. However by 72 h (stationary phase)
Overproduction of fumarate reductase

Fig. 5. Negative stain electron microscopic views of tubules from a cardiolipin-synthesis mutant, E. coli T1GP(pFRD63). Bars, 50 nm.

only small fragments of tubules are visible in the cytoplasm and the total activity of fumarate reductase is reduced by up to 50%. This loss of activity accompanied by tubule breakdown is not observed with the plasmid-coded enzyme, indicating an inherent difference in the two forms of amplified fumarate reductase. The membranes from these cells have an increased amount of cardiolipin, as seen in the normal overproducing strains (Table 2), indicating the lipid composition is not responsible for the instability of the G800 tubules.

DISCUSSION

Little information is available on the response of bacterial cells to very high levels of expression of integral membrane proteins. Our studies with the terminal electron transfer
enzyme fumarate reductase suggest that the cells respond to this stress by synthesizing a novel intracellular organelle composed of phospholipid and enzyme (Weiner et al., 1984).

The method we have devised for isolating tubules, relatively free of inner and outer membrane, relies on gentle hypotonic lysis of bacterial cells. Tubules not released in the soluble fraction were eluted from the spheroplasts by a combination of mild detergent and freeze-thaw treatments. Our inability to separate tubules from inner membranes routinely by sucrose density gradient centrifugation required that the initial preparation contained as few inner membranes as possible. The isolated tubules were further purified by centrifugation on a sucrose cushion. The yield of fumarate reductase in the tubules is about 50%, a value in good agreement with the doubling of membrane material observed in overproducing cells; 90% of the tubular protein is fumarate reductase. The physical properties and catalytic behaviour of tubule-bound fumarate reductase are indistinguishable from cytoplasmic membrane-bound enzyme or soluble holoenzyme. Tubules have a phospholipid composition and fatty acid profile similar to inner membranes from amplified strains. The only observed difference is in the lipid to protein ratio. Tubules have a much lower ratio (0.25) than the cytoplasmic membranes (0.35). This difference would suggest that the tubules are much denser than the cytoplasmic membranes, but we have been unable to separate tubules from vesicles by sucrose density gradient centrifugation.

Isolated tubules display a regular helical packing arrangement of exposed spherical protein structures when observed by electron microscopy. Based on protein A-gold labelling and other experiments (Lemire et al., 1983) we have shown that the spheres are the catalytic subunits of fumarate reductase. These results suggest that the anchor subunits are buried within the tubules together with the phospholipid fatty acyl side chains.

The mechanism of tubule assembly remains to be elucidated, but as a first step we have examined the role of cardiolipin, since the tubules and cytoplasmic membranes of overproducing cells are highly enriched in this phospholipid. A transformed mutant, E. coli T1GP(pFRD63), with a defect in cardiolipin biosynthesis still produced tubules, but these tubules differed morphologically from the wild-type, displaying a variable and irregular arrangement of the fumarate reductase. Also, the enzyme in these structures was unstable to freeze-thaw manipulations. These results suggest that cardiolipin plays a role in the assembly of fumarate reductase in the tubules and regulates the packing of the protein in these structures.

The expression of fumarate reductase can be amplified in at least two ways. First, selection of ampicillin hyper-resistant strains, in an ampA1 genetic background, leads to tandem duplications of the amp-frd region, resulting in an increased frd gene dosage (Cole & Guest, 1979a, b). A second approach is to use multicopy plasmids expressing the cloned frd operon (Cole & Guest, 1980; Lohmeier et al., 1981; Lemire et al., 1982, 1983). In this paper we have shown that both methods of amplification result in tubule formation. However, the tubules formed in response to ampicillin hyper-resistance are qualitatively more unstable than those formed in response to the plasmid-coded genes. Indeed it did not prove possible to isolate tubules from G800 by the protocol reported in this paper. The reason for the instability is unclear but may be related to the ampA1 mutation. This mutation is a frame shift in the overlapping amp attenuator-frdD region (Jaurin & Grundström, 1981), and causes the hydrophobic FrdD subunit to be covalently-linked to β-lactamase. We have been unable to identify the frdD-ampC fusion protein in membrane preparations (B. D. Lemire, S. T. Cole & J. H. Weiner, unpublished), even though the β-lactamase can be detected in the membrane protein profile.

The correlation of tubule formation with intrinsic membrane protein amplification remains to be determined, but similar structures have been observed with F0/F1 ATPase (Von Meyenburg et al., 1984) and in strains overproducing glycerol-3-phosphate acyl transferase and pyridine nucleotide transhydrogenase (R. Bell & P. D. Bragg, personal communications). The ability to isolate tubules, consisting of a nearly homogeneous membrane protein and lipid, may be an important biotechnological approach for isolating cloned membrane proteins.

We thank Professor J. R. Guest for providing strain G800 and Dr E. P. Kennedy for providing strain T1GP. We also thank Dr R. N. McElhaney, Dr R. Lewis and Dr M. Poznansky for help with phospholipid analysis, L. Rodseth for the optical diffraction analysis and R. D. Bradley for assistance with the electron microscopy and photography. We also thank Gillian Shaw for excellent technical assistance.
REFERENCES


This work was funded by grants from the Medical Research Council of Canada to J. H. W. and D. G. S. M. L. E. received support from the Alberta Heritage Foundation for Medical Research.