The Ultrastructure of the Capsules of *Diplococcus pneumoniae* and *Klebsiella pneumoniae* Stained with Ruthenium Red

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SUMMARY

Capsules of *Diplococcus pneumoniae* type III and *Klebsiella pneumoniae* type I were examined in the electron microscope using ruthenium red in combination with osmium tetroxide. The ability of this combination stain and fixative to stain polysaccharides has been well established. A slightly acid pH was necessary to obtain satisfactory capsular polysaccharide staining.

The capsules of both *Diplococcus pneumoniae* and *Klebsiella pneumoniae* were stained by the ruthenium red before alcohol dehydration and embedding in Epon 812. The capsule of *D. pneumoniae* had the appearance of a tightly woven mat. The capsule of *K. pneumoniae* had a fibrous appearance. Ruthenium red was found inside the plasma membrane of *D. pneumoniae*, but remained outside the cell membrane of *K. pneumoniae*.

INTRODUCTION

Early efforts to examine bacterial capsules by electron microscopy made use of whole organisms that were either stained with metal salts or were metal shadowed. Mudd, Heinmets & Anderson (1943) published electron micrographs of whole *Diplococcus pneumoniae* stained with lead acetate or copper sulphate; only a hazy indistinct outline of the capsule was visible. After palladium shadowing, Labaw & Mosley (1954) observed spindle shapes in the capsules of the Lisbonne strain of *Escherichia coli*. Studies of the cellulose capsule of whole cells of *Acetobacter xylinum* using palladium or palladium–gold shadowing revealed microfibrils comprising the capsule (Colvin & Beer, 1960; Millman & Colvin, 1961).

Part of the problem in preserving and staining the bacterial capsule has been the lack of a suitable fixative and stain specific for acid mucopolysaccharides. In 1964, Luft used ruthenium red to stain acid mucopolysaccharides in animal tissues examined by electron microscopy. Pate & Ordal (1967) used ruthenium red to demonstrate peripheral fibrils in *Chondrococcus columnaris*. Jones, Roth & Sanders (1969) modified the procedure of Pate & Ordal (1967) to visualize polysaccharide fibres in aquatic slime layers.

Ruthenium red might be useful in studying the ultrastructure of bacterial capsules. The purpose of this work was to examine, by electron microscopy, the ultrastructure of the capsules of *Diplococcus pneumoniae* type III and *Klebsiella pneumoniae* type I using ruthenium red in combination with osmium tetroxide and several methods of dehydration and embedding.


**METHODS**

**Culture**

*Diplococcus pneumoniae* type III (ATCC 6303) and *Klebsiella pneumoniae* type I obtained from the Center for Disease Control, Atlanta, Georgia, were inoculated into starter cultures of brain heart infusion broth (Difco) supplemented with 2% (w/v) sucrose. After 16 h of incubation at 37°C, fresh media was inoculated and allowed to incubate under static conditions. The turbidity of the culture and the production of acid were followed using a Bausch and Lomb Spectronic 20 spectrophotometer at 660 nm and a Corning model 10 pH meter. The bacteria were harvested by centrifugation when late log phase was reached and the maximum amount of capsule was present.

**Staining and fixation**

The pellet was suspended in 1 ml of ruthenium red (K & K Laboratories, Plainview, New York, U.S.A.) (1.5 mg/ml) in water, 1 ml 3.6% (w/v) glutaraldehyde (Ladd Research Industries Burlington, Vermont, U.S.A.) and 1 ml of 0.2 M-cacodylate buffer (Ladd Research Industries), pH 6.5, and held for 1 h at 0°C.

Stainless controls were incubated with an additional ml of buffer in place of ruthenium red.

**Dehydration and embedment**

*Luft's method.* The bacteria were dehydrated in successive 10 min steps of 30% (v/v), 50% (v/v), 70% (v/v), 90% (v/v) ethanol and two washings with absolute ethanol for 15 min each. They were next suspended in a mixture of three parts ethanol to one part of an Epon 812 mixture. The Epon 812 mixture contained 3 ml of mixture A (62 ml of Epon 812 and 100 ml of dodecenyl succinic anhydride; Ladd Research Industries) and 7 ml of mixture B (100 ml Epon 812 and 89 ml of methylnadic anhydride; Ladd Research Industries) and was incubated 30 min at room temperature. The bacteria were transferred to a solution that was 1 part alcohol and 1 part Epon mixture and held for 30 min at room temperature. They were then harvested by centrifugation and transferred to a BEEM capsule (Poly-sciences, Warrington, Pennsylvania, U.S.A.) and a fresh Epon mixture containing the amine accelerator, tridi-methylamino-methyl phenol (Ladd Research Industries; 0.15%, v/v), was added. The embedding was then cured for 24 h at 60°C (Luft, 1961).

*Craig's method.* In the Craig, Frajola & Greider's (1962) procedure using Epon 812 dehydration and embedding in Epon 812, the bacteria were dehydrated in ascending concentrations of Epon 812 in 0.2 M-cacodylate buffer, pH 6.5, following this schedule: 25% (v/v) buffer and 75% (v/v) Epon 812 for 1 h, transferred to a mixture of 55% (v/v) Epon 812 and 45% (v/v) hexahydrophthallic anhydride (HHPA) (Polysciences) for 2 h at room temperature. Next the bacteria were placed in a fresh solution of 55% (v/v) Epon 812 and 45% (v/v) HHPA now including 2% (v/v) benzylidemethylamine (BDMA; Ladd Research Industries) for 2 h. After 2 h the bacteria were placed in a fresh solution of Epon 812, HHPA and BDMA (in the same concentrations as before) and cured 12 h at 55°C.

*Staubli’s method.* In Staubli's (1963) method for dehydration and embedding, the water-soluble resin Durcupan is used to dehydrate the bacteria and the water insoluble Araldite 502 is used for embedding. The bacteria were dehydrated at 0°C with Durcupan (Fluka AG, Buchs SG, Switzerland) in successive 15 min steps of 50% (v/v), 70% (v/v) and
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90% (v/v) Durcupan in water with two washings of 45 min each in 100% Durcupan. The bacteria impregnated with Durcupan were embedded in Araldite 502 (Ciba Products, Fairlawn, New Jersey, U.S.A.) at 50°C as follows: 70% (v/v) Durcupan at 30% (v/v) Araldite I (one part Araldite 502 and one part dodecenyl succinic anhydride) for 1 h; 50% (v/v) Durcupan and 50% (v/v) Araldite I for 1 h; three washes of 30 min each with Araldite II (Araldite I + 2%, v/v, 2,4,6-tris-dimethylaminoethyl phenol; Ladd Research Industries) and placement in a BEEM capsule with fresh Araldite II to polymerize at 50°C for 24 h.

Sections were cut with an LKB Ultratome Type 4801A with glass knives made on a LKB Knifemaker Type 7801A. The sections were lifted from distilled water on copper grids. Contrast of some sections was enhanced by treatment with saturated aqueous uranyl acetate for 15 min as suggested by Watson (1958). All of the sections were examined with a JEM (Japan Electron Optics Laboratory, Tokyo, Japan) Model 6-C electron microscope.

RESULTS

Diplococcus pneumoniae stained with ruthenium red

Bacteria dehydrated with alcohol and embedded in Epon 812 showed dense fibrous capsules characterized by a woven or mat-like structure. Additional staining also occurred in and beyond the plasma membrane (Fig. 1). In Fig. 2, some of the capsule has sloughed off showing the fibrous structure of the capsule more clearly; again intense staining of the plasma membrane and part of the cytoplasm has occurred. A well preserved mesosome is also present. Since the contrast in neither organism has been enhanced by additional staining, the increased density of the capsule and the plasma membrane can be attributed to the ruthenium red-osmium coupled reaction. Bacteria dehydrated either in Epon 812 and embedded in Epon 812 or dehydrated in Durcupan and embedded in Araldite 502 did not appear to retain capsule and resembled the stainless control which was dehydrated with alcohol and embedded in Epon 812 (Fig. 3).

Klebsiella pneumoniae stained with ruthenium red

Bacteria dehydrated with alcohol and embedded in Epon 812 display well preserved and stained capsule with no increased ruthenium red staining of the plasma membrane and cytoplasm like that seen in pneumococcus treated identically. A variety of capsule forms are visible within the same micrograph. This probably indicates multiple effects of dehydration on the capsular gel (Fig. 4). The spike-like structure of the bacterial capsule is clearly evident in Fig. 4. Other bacteria in the same micrograph have overlapping spikes creating a net-like appearance which is most evident in the central bacterium in Fig. 5. Some of the fibrils of capsule seem to have cross-linking structures between them as in Fig. 6. To the left of the organism cut longitudinally, a transverse section through the capsule showing the fibril tips is seen.

Dehydrating and embedment with Epon 812 caused separation of the spiked capsular fibrils into thin strands (Fig. 7). In bacteria dehydrated with Durcupan and embedded in Araldite 502, better preservation of capsular fibrils was noted than was evident after alcohol dehydration and embedment in Epon 812. The fibrils are nearly uniform in length and width (Fig. 8). A section through the capsule portion of the organism shows the individual fibrils, as well as some of the wall. Bacteria fixed in the absence of ruthenium red and dehydrated and embedded by all the three described methods showed faint fibrils exterior to the wall (Fig. 9). However, since contrast was poor without ruthenium red, aqueous uranyl acetate was used to enhance contrast.
Fig. 1, 2. *Diplococcus pneumoniae* stained with ruthenium red, dehydrated with alcohol and embedded in Epon 812. The mat-like capsule (C) surrounds each bacterium. Ruthenium red penetration is evident at the plasma membrane (PM) and into the cytoplasm. The marker represents 0.5 μm.
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Fig. 3. *Diplococcus pneumoniae*, stainless control dehydrated and embedded as cells in Fig. 1 and 2. No capsular material is evident external to the wall nor is the plasma membrane heavily stained as in Fig. 1 and 2. The marker represents 0.5 µm.

Fig. 4. *Klebsiella pneumoniae* stained with ruthenium red, dehydrated with alcohol and embedded in Epon 812. Capsular fibrils (C) are seen at regular intervals along the wall. The marker represents 0.5 µm.
Fig. 5, 6. *Klebsiella pneumoniae* treated as Fig. 4. Fig. 5: The central bacterium has overlapping capsular fibrils (C) creating a net-like appearance. Fig. 6: Capsular fibrils (C) seem to have cross-linking structures between them. The spike-like nature of the dehydrated capsule (C) is seen in the tangentially cut bacterium. The markers represent 0.5 μm.
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Fig. 7. *Klebsiella pneumoniae* stained with ruthenium red, dehydrated with Epon 812 and embedded with Epon 812. The capsule (C) is composed of slender, thread-like fibrils. The marker represents 0.5 μm.

Fig. 8. *Klebsiella pneumoniae* stained with ruthenium red, dehydrated with Durcupan and embedded in Araldite 502. Capsular fibrils (C) are seen at regular intervals along the wall. A bacterium tangentially cut through the capsule (C) also reveals the wall (CW). The marker represents 0.5 μm.
Fig. 9. *Klebsiella pneumoniae* stainless control representative of all three dehydration and embedding procedures. Some faintly electron-dense capsular fibrils were preserved by this fixation procedure at pH 6.5. The marker represents 0.5 μm.

Comparison of *Klebsiella* capsule preservation after either alcohol, Epon 812 or Durcupan dehydration

The appearance of the capsule after any of the three dehydration procedures was fairly consistent (i.e. repetitive fibrils external to the wall). However, differences in fibril length and width and tendency to overlap varied considerably with the dehydrating procedure used. Capsules dehydrated with alcohol (Fig. 4) or Durcupan (Fig. 8) had fibrils which averaged 156 nm by 32 nm; whereas capsules dehydrated in Epon 812 (Fig. 7) had long (235 to 236 nm) and thin (7 to 8 nm) fibrils. The fibrils resulting from Epon 812 dehydration of capsules were 60% longer and only 23% was wide as those dehydrated with alcohol or Durcupan (Table 1). These ranges were established by making ten measurements on specimens dehydrated by each procedure.

Discussion

Although the technique of Mudd *et al.* (1943) for preserving and increasing the electron-density of the capsule of *Diplococcus pneumoniae* permitted visualization of the capsule in the electron microscope for the first time, fine structural detail was not visualized and the metal salts used precipitated the capsule polysaccharide and destroyed the gel matrix. Metal shadowing used by Ohad, Danon & Hestrin (1962) showed that the capsule of *Acetobacter xylinum* was composed of many fibrils which extended outwards from the bacterial surface. Neither of these reports included studies of ultrathin sections.
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Table 1. **Range of dimensions of capsule fibrils of Klebsiella pneumoniae**

<table>
<thead>
<tr>
<th>Dehydrating procedure</th>
<th>Fibril length</th>
<th>Fibril width</th>
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<tbody>
<tr>
<td>Alcohol</td>
<td>146 to 160 nm</td>
<td>37 to 40 nm</td>
</tr>
<tr>
<td>Durcupan</td>
<td>153 to 166 nm</td>
<td>24 to 27 nm</td>
</tr>
<tr>
<td>Epon 812</td>
<td>253 to 267 nm</td>
<td>7 to 8 nm</td>
</tr>
</tbody>
</table>

In *Diplococcus pneumoniae* and *Klebsiella pneumoniae* fixed and stained with ruthenium red–osmium tetroxide in 0·2 M-cacodylate buffer at pH 7·3 as used by Pate & Ordal (1967), no staining of the capsule occurred. The pH of the buffer was lowered to 6·5 as for uptake of ruthenium red by polygalacturonic acid (Stoddart & Tipton, 1968). This pH also falls within the recommended limits (pH 6·5 to 8·0) for optimal fixation of biological specimens (Millonig & Marinozzi, 1968), and is satisfactory for retention of the capsule on the bacterial surface, as Bernheimer (1953) reported that below pH 6·5 the capsule of *D. pneumoniae* is lost. The electron-density of the capsule of both organisms was increased after fixation and staining at pH 6·5, making visualization of the capsule fibres possible. The action of ruthenium red, as suggested by Luft (1971), involves the formation of an electron dense deposit with OsO₄ at the location of polysaccharides. The ability of ruthenium red to stain polysaccharides has been well established in studies with both mammalian and plant cells. Since the capsules of *Diplococcus* and *Klebsiella* are polysaccharides, the electron-dense fibrils external to the wall of both organisms, which were stained with ruthenium red–osmium tetroxide, are considered to be capsular fibrils.

When water is removed from specimens by displacement with Durcupan or Epon 812 (Craig *et al.* 1962) instead of alcohol dehydration, the chemical effects of dehydration are quite different. The epoxides react with amino, imidazole, and carboxyl groups of proteins, as well as sulphhydryl groups of denatured proteins (Millonig & Marinozzi, 1968) but not with carbohydrates. If epoxides react with the carbohydrate components of the cells, the reaction would be unlike that with alcohol. Indeed, the epoxy resins may maintain the dielectric constant between adjacent polysaccharide polymers (Fig. 7, 8) and thereby prevent the overlapping of capsule fibrils seen in alcohol dehydration (Fig. 4, 5, 6). The preservation of the bacteria following Durcupan dehydration was as good as with alcohol dehydration; however, the capsule fibrils of *Klebsiella* revealed less overlapping (Fig. 8). The tendency of the fibrils to overlap seemed to be completely overcome with Epon 812 dehydration. Since both Epon 812 and Durcupan are epoxides, the difference in preservation of the capsular fibrils may reside in the degree to which the dielectric constants between the fibril polymers were lowered, permitting attraction between polymers and eventual annealing. According to Wilkinson, Duguid & Edmunds (1954), the capsule of *Klebsiella aerogenes* contains almost 95% water. *Klebsiella pneumoniae* has a capsule with a fibrous structure capable of binding large amounts of water. The preservation of many long, thin fibrils of capsular material more probably represents the natural hydrated state of the capsule than capsule fibrils which are short, thick and offer little surface for the binding of water. Thus, these micrographs of *K. pneumoniae* offer ultrastructural evidence in support of the finding of Wilkinson *et al.* (1954).

The slender fibrillar structures in Fig. 7 and 8 are also in keeping with the proposal of Heidelberger, Kendall & Scherp (1936) that capsular carbohydrate is in thread-shaped polymers. Hershey (1940) postulated that the surface of capsulated bacteria is covered with projecting molecules or fibrils after his experiments with the absorption of antibody to the
capsule surface revealed that a greater uptake of antibody occurred than was necessary to form a single surface layer of closely packed molecules. Electron micrographs of metal-shadowed Acetobacter xylinum (Colvin & Beer, 1960) and Escherichia coli Lisbonne strain (Labaw & Mosley, 1954) also, show the respective capsules to be composed of many fibrils.

In the micrographs of Diplococcus pneumoniae (Fig. 1 and 2) stained with ruthenium red, dehydrated with alcohol and embedded in Epon 812, there is intense staining of the plasma membrane and cytoplasm adjacent to it; while Klebsiella pneumoniae treated identically shows no such staining of the plasma membrane. In the only other study of the effect of ruthenium red on a pure culture of bacteria, Pate & Ordal (1967) found no increased staining of the plasma membrane of the myxobacterium Chondrococcus columaris. Since Chondrococcus is considered to have a Gram-negative type wall (Lautrop, Reyn & Birch-Anderson, 1964); Klebsiella would be expected to respond to ruthenium red in a similar manner, i.e. the ruthenium red would remain outside the wall. The fact that the ruthenium red penetrated the plasma membrane of D. pneumoniae indicates a greater porosity of this membrane as compared with mammalian plasma membranes.

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REFERENCES


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