BACTERIAL STRUCTURE AND PATHOGENICITY

Structural studies of the surface projections of Chlamydia trachomatis by electron microscopy

J.-J. CHANG, K. R. LEONARD* and Y.-X. ZHANG†

Institute of Biophysics, Academia Sinica, Beijing 100101, China, *European Molecular Biology Laboratory, Postfach 10.2209, D-69012 Heidelberg, Germany and †Beijing Institute of Ophthalmology, Beijing, China

Rod-like projections on the surface of Chlamydia trachomatis have been studied by a combination of computer image analysis and electron microscopy. The rods, c. 60–80 Å in diameter and c. 500 Å in length, were found on the surface of prokaryocells of C. trachomatis inserted in the cytoplasmic membrane through a ring-like structure in the outer membrane. The rod-like structures were found at all stages of the life cycle, even in very small elementary bodies (EBs) of C. trachomatis and in vesicles < 0.2 μm. Computer image analysis of isolated rods indicated that they comprise helically arranged subunits with a periodicity of c. 50 Å. From their localisation and distribution, these structures may be related to the proliferation, or to the infectivity, of chlamydiae.

Introduction

Chlamydia trachomatis is a member of the genus Chlamydia which comprises obligate intracellular bacteria causing various diseases in both animals and man (reviewed in [1]). As with other chlamydia species, C. trachomatis can be distinguished functionally during its developmental cycle between two different cell types, namely the elementary body (EB) which is the infectious stage and the reticulate body (RB) which is non-infectious. EBs may exist as small cocci, c. 0.3 μm in diameter; the RBs may be larger, c. 1 μm in diameter.

The best identified feature of chlamydiae is the specific structure of their envelopes. Like other species, C. trachomatis has been shown to have a regular hexagonal lattice within the outer membrane of the cell envelope [2]. In negatively stained preparations of strain TE55 of C. trachomatis, folds and grooves on the surface of EBs were also observed [3].

Some reports also describe the presence of short stick-like structures attached to the outer surface of chlamydiae. For example, projections on the surface of C. psittaci were found after thin-sectioning and staining with ruthenium red by Matsumoto [4], who suggested that they were cytoplasmic-membrane sites where the projections were connected to DNA molecules. By negative staining and freeze-fracturing, the presence of projections on the surface of C. psittaci was confirmed [5, 6] and the surface projections were found not only in EBs but also in RBs. Each elementary body had 18–23 projections. Stick-like protuberances were also described [2].

Nichols et al. [7] observed two kinds of surface projections, spike-like rods and hemispheric projections in C. trachomatis by use of thin-sectioning techniques. Again, when Miyashita et al. [8] compared isolated C. pneumoniae TWAR, C. trachomatis L3/434/Bu, C. psittaci Call 10 and C. pneumoniae YK-41, they found that all strains possessed very similar morphological features, including surface projections. Matsumoto [9], in a preparation of isolated surface projections of C. trachomatis, reported that stain penetration occasionally outlined fine subunits and a central channel in the projections, suggesting that they are tubular with a helical arrangement of subunits. Although it has been proposed that the surface projections are related to multiplication of chlamydiae, their function, whether to anchor the parasites to host cells or to play a role in nutrient absorption, is still unknown (reviewed in [10]).

This paper describes the structure of the surface projections of C. trachomatis, studied in more detail by combining electron microscopy with computer image analysis. Finally, some possibilities of their functions related to proliferation and infection are discussed.
Materials and methods

Cultivation and purification of C. trachomatis

Prokaryotic cells of C. trachomatis strain TE55 were cultured in yolk sac membranes [11] and the cultured cells were purified [3] as described previously. To visualise the hexagonal lattice on the envelopes as an identifying characteristic of C. trachomatis, some purified chlamydia cells were digested in sodium dodecyl sulphate 1% w/v or in sodium deoxycholate 1% w/v in phosphate-buffered saline (PBS, pH 7.2). Some cells were also disrupted with a microprobe-ultrasonicator to break the envelopes into single layers.

Electron microscopy

Negatively stained specimens were prepared as follows: c. 5 μl of chlamydial suspension was applied to hydrophilic carbon-coated grids freshly glow-discharged in air. After adsorption for 1 min, a drop of phosphotungstic acid (PTA, 1% w/v, pH 7.2) was added. Excess liquid was removed with filter paper and grids were dried in air.

Freeze-dried shadow-casting samples were prepared as follows: after removal of liquid with filter paper, grids with chlamydiae were frozen immediately in liquid
nitrogen and transferred to a cryochamber (Reichert Cryofract) under liquid nitrogen. Grids, after freeze-drying in the cryochamber, were shadowed with platinum at an angle of 25° in an evaporator (Balzers TCP-121). As an internal size standard, tobacco mosaic virus (TMV) was mixed with chlamydiae before being applying to grids.

Micrographs were taken in an electron microscope (Phillips EM 400) operating at 80 kV and at a magnification of ×28 000 with a low dose unit (Philips) for off-specimen focusing and astigmatism correction.

Computer image analysis
Images were digitised at 7 µm raster on a scanning system (Zeiss Phodis/SCAI). The scanned pixel size was 4 Å. Fast Fourier transforms were calculated, after boxing and floating image data, by use of 1024 × 1024 arrays.

Results
Distribution of projections
Images of platinum-shadowed cells and broken envelopes are shown in Fig. 1. The hexagonal lattice is

![Image](image-url)

**Fig. 2.** Negatively stained intact EBs (a) and RB (b) showing rods inserting into the surface of the prokaryotic cells (arrowed). a, a region containing several rods which traverse the two cell membranes is outlined by a box (shown at higher magnification in Fig. 6). Scale bars a, 100 nm, b, 0.5 µm.
clearly visible on the inner surface of a broken envelope. The outer surface of intact cells is smoother in appearance, with small rods lying irregularly over the envelope. Images of negatively stained intact EB and RB are shown in Fig. 2. In each case, surface projections or rods are indicated at the edge of cells. In addition, rods in profile are shown lying on the surface of one cell (Fig. 2a).

Structure of surface projections

A negatively stained sample of disrupted envelopes with rod-like projections broken away from the cell envelopes is shown in Fig. 3. The hexagonal cell envelope lattice is clearly visible, as are several ring-like structures superimposed irregularly on the lattice. The rings are c. 150 Å in width, with a central stain-filled hole c. 80–100 Å in diameter.

Selected images of negatively stained fragments of rods are presented in Fig. 4. A fine stain-filled channel is just visible along the centre of the rods (Fig. 4a, b). A full-length rod, c. 500 Å in length and 70 Å in diameter, is shown in Fig. 4f, with a computer-calculated diffraction pattern from that particle shown in Fig. 4g. The diffraction patterns from 11 samples

---

Fig. 3. Image of negatively stained patches of broken cells after SDS treatment and ultrasonication showing the hexagonal lattice of the outer membrane and a number of ring-like structures distributed randomly across the patches. Broken rods at the edges of the patches and in the background are arrowed. Scale bar, 0.5 μm.
Fig. 4. Isolated projections after SDS treatment and ultrasonication. Broken fragments c. 200 Å in length are shown (a–e). A ring-like structure is visible at the base of the rod (e). A full length rod (c. 400 Å) is shown in (f) from which the computer-calculated diffraction pattern (shown in g) has been obtained. Two layer lines at c. 50 Å and 100 Å reciprocal spacing are visible in g. Scale bar, a–f, 100 Å.

Localisation of rod-like surface projections

A ring-like structure is seen at the end of a broken rod (Fig. 4e), with a central hole of the same diameter as the width of the rod. In size and shape, this ring corresponds to those seen in disrupted envelopes (Fig. 3). Fig. 6 shows images of rods (enlarged from the boxed area shown in Fig. 2a). These negatively stained samples show two cell membranes in profile. There appears to be a ‘collar’ around the rod, just below the inner face of the outer membrane (Fig. 6b, c). Taken together, these observations suggest that the ring or collar structures are present at the point where the rods pass through the outer membrane. The rods traverse the inter-membrane space and extend through the outer membrane. As may be seen, the distance between the two membranes varies; thus, the rods appear to be strongly anchored to the inner membrane but are loosely attached as they pass through the outer membrane.

Fig. 5. Radial projection of a 70 Å diameter helix with two principal helical lines of pitches 50 Å and 100 Å defining the surface lattice (a), corresponding to the layer lines seen in Fig. 4g. Schematic view of this lattice on the surface of the rod is shown (b).

were weak, as only very short lengths of particle could be selected. In all cases, a layer line was visible at about 50 ± 4 Å. A second layer line at c. 100 Å is also visible (Fig. 4g).

A simple model to explain this result is shown (Fig. 5), taking into account the diameter of the rods and the presence of two main helical directions separating the subunits. A radial projection of the surface lattice of a simple helix, with a radius of 35 Å, and two principal helical directions corresponding to the 50 Å and 100 Å layer lines are shown in Fig. 5b. This model has c. six subunits per 100 Å length. A solid cylinder of protein with 35 Å radius would correspond to a volume of 385 000 Å³/100 Å length. Taking the density of protein to be 0.8 Da/Å³, this would give a mass of c. 51 kDa per subunit. Allowing a 7 Å radius channel at the centre of the rod, as suggested by the penetration of stain, the subunit mass would be reduced to c. 50 kDa.

Fig. 6. Computer-scanned and enlarged images from the boxed area in Fig. 2. Negative stain has penetrated the region between the inner and outer membranes. Rods are anchored to the inner plasma membrane at one end and pass through the outer membrane. Just below the outer membrane a structure is visible (arrowed) which appears to be attached to the rod. The width of this structure is the same as that of the ring seen in Fig. 4e and may be a collar surrounding the rod. The length of rod between the inner and outer membranes is variable, suggesting that rods can pull through this collar. Scale bar a, 0.5 μm; b, c, 10 nm.
Discussion

The results of the present study support a model in which the rod-like projections on the surface of C. trachomatis are helical tubes with a narrow, almost invisible, central channel. The overall length of the rod is c. 500 Å. One end is anchored to the inner membrane with the remainder passing relatively freely through the outer membrane, such that a segment (c. 300 Å) projects from the surface. Broken rods found after ultrasonication are usually c. 300 Å in length, suggesting that they have sheared off at the level of the outer membrane. These dimensions are similar to those found for C. psittaci [5], but less than those reported for C. trachomatis [7].

The approximate estimate of 50 kDa for the size of the protein subunit making up the surface projections is based on a simple model which does not take into account the shape of the subunit. At present no direct data (e.g., measurements in mol wt from SDS gels) exist for the subunit, which is a very minor component of the outer membrane. However, it is possible that a protein of this size, which should be present in both EB and RB stages, may be antigenic or may be available for surface labelling by chemical methods. Two possible candidates are the antigenic 59-kDa protein present in both EB and RB [12] or the 38-kDa cytadhesin protein [13] shown to be present on the surface of EBs.

The distribution of rods varied on the cell surface among different cells as well as on different areas of the same cell surface. The number and distribution of ring-like structures seen in images of the hexagonal inner surface of the outer membrane was similar to that seen for the rods on the outer surface. This observation would be consistent with the rings forming an opening in the outer membrane through which the rods might pass relatively freely. The function of the rod-like projections remains unclear. The stain-filled central channel of the rod that appears faintly in some images, is probably <15 Å in diameter. Unless this is a result of shrinkage during specimen preparation, the channel would be too small for DNA or protein to pass through. However, it may be an ion channel. Alternatively, the rods may be involved in binding to target cells or in transmitting signals to the inner (cytoplasmic) membrane on binding. However, they are found at all developmental stages, even in the very small EBs, suggesting that their presence is necessary for the proliferation of chlamydial cells or to contribute to the infectivity of these obligate intracellular bacteria.

References