Morphological analysis of *Chlamydia pneumoniae*

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In recent years we have succeeded in isolating 18 strains of *Chlamydia pneumoniae* strains, which we designated KKpn-1~KKpn-18, from the nasopharynx of patients with acute respiratory tract infections. In the present study, we examined the morphology of these clinical isolates and compared their morphology with that of *C. pneumoniae* TW-183, AR-39, AR-388 (TWAR), IOL-207, Kajaani-6, YK-41, *Chlamydia psittaci* Frt-Hu/Cal 10, *Chlamydia trachomatis* L2/434/Bu and *Chlamydia pecorum* Bo/E56. The results indicated that the "pear-shape" of the elementary body (EB) is not a valid morphological criterion for recognizing a member of *C. pneumoniae*, although the basic morphological properties, such as location of the nucleus, presence of projections on the surface of the organisms, and the dimension of hexagonally arrayed regular structures on the inner surface of the EB outer membrane, were similar in all members of the genus *Chlamydia*. It was, therefore, concluded that the morphology of the KKpn-1~KKpn-18 strains was identical to that of *C. trachomatis*, *C. psittaci* and *C. pecorum* organisms, and that the EB profiles are different from those of *C. pneumoniae* TWAR strains.

Key words: *Chlamydia pneumoniae*, pear- or round-shaped EB, Button structure, surface projection, hexagonal structure

Introduction

The chlamydiae, obligate intracellular parasites, have been assigned to a single genus, *Chlamydia*, which comprises four species, *Chlamydia pneumoniae*

1), *Chlamydia trachomatis* 2), *Chlamydia psittaci* 2) and *Chlamydia pecorum* 3). These organisms multiply through a common, unique developmental cycle in which there are two morphologically and functionally distinct forms: one is the infectious elementary body (EB) and the other is the reproductive reticulate body (RB). The EB penetrates a susceptible host cell by being phagocytized and is converted into an RB, which multiplies by binary fission. The RB then undergoes maturation through an intermediate body, to form a mature progeny EB 4).

*C. pneumoniae* is well recognized as a major respiratory pathogen 5). The organism is a common cause of pneumonia, bronchitis, sinusitis, and pharyngitis, as well as being responsible for up to 10% of cases of community-acquired pneumonia 6,7). The prevalence of *C. pneumoniae* infections has a worldwide status similar to that of *C. trachomatis* infections 8). In addition, recent investigations have suggested a possible association of *C. pneumoniae* infection with atherosclerotic cardiovascular disorders 9,10), acute exacerbations of chronic obstructive pulmonary disease 10,11), and bronchial asthma 12,13).

In the last four years, we succeeded in isolating 18 *C. pneumoniae* strains, designated KKpn-1~KKpn-18 11,14,15), from the nasopharynx of patients with acute respiratory tract infections. In the present study, we examined the morphology of the clinical isolates (KKpn-1~KKpn-18) and compared it with the morphology of established strains of *C. pneumoniae*, *C. trachomatis*, *C. psittaci* and *C. pecorum*.

Materials and Methods

*Chlamydial strains*

The following chlamydial strains were used: *C. pneumoniae* TW-183, AR-39, AR-388 (TWAR), IOL-207 16), Kajaani-6 17), YK-41 18), KKpn-1 19), KKpn-15 19), KKpn-16 19) and 15 clinical isolates 19*, *C. psittaci* Frt-Hu/Cal 10 19*, *C. trachomatis* L2/434/Bu 20)
and C. pecorum Bo/E58 strains. Strains TW-183, AR-39 and AR-388 were purchased from the Washington Research Foundation, Seattle, WA, USA. Strains IOL-207 and Kajaani-6 were supplied by P. Saikku, University of Helsinki, Finland. Strain YK-41 was supplied by Kanamoto Y, Hiroshima Prefectural Institute of Public Health, Japan. KKpn-1, KKpn-15, KKpn-16 and 15 other clinical isolates of the C. pneumoniae strains were isolated in our own laboratory. These C. pneumoniae strains were grown in HeLa 229 or HL cell cultures and harvested on day 3 postinoculation. C. pecorum E58 was supplied by Hirai K, Faculty of Agriculture, Gifu University. C. trachomatis L2 was supplied by Yamasaki S, National Institute of Health, Japan. C. psittaci Cal 10 has been maintained in our laboratory for more than 20 years. C. trachomatis L2 and C. psittaci Cal 10 were grown in L929 suspension or monolayer cultures and HeLa 229 cell cultures and then harvested on day 2 or 3 postinoculation. C. pecorum E58 was propagated continuously in a MDBK cell line.

**Cultivation of infected cells in the experiments**

HeLa 229 cells were used for infection with C. pneumoniae strains. After brief sonication, all inocula were centrifuged (900 × g for 60 min at room temperature) onto the cell monolayers. Infected cells were then incubated at 35°C in an atmosphere of 5% CO₂ in minimum essential medium (MEM, Nissui, Tokyo) containing 1 µg/ml cycloheximide and supplemented with 10% heat-inactivated fetal calf serum. The Cal 10 and L2 strains were inoculated onto L929 cells without centrifugation. Similarly, MDBK monolayer cells were inoculated with the E58 strain and then incubated at 37°C in an atmosphere of 5% CO₂ in Dulbecco-modified MEM (Nissui, Tokyo) supplemented with 10% serum.

**Light microscopy**

To examine the morphology and stainability of the intracytoplasmic inclusions, a monolayer of L929 cells or HeLa 229 cells on a cover slip (14 mm in diameter) was inoculated and incubated as described above. After incubation for 48~72 h, the infected cells were air-dried, fixed in absolute methanol, and stained with iodine and May–Gruenwald–Giemsa staining solutions. The cells were also fixed with absolute ethanol and stained with C. pneumoniae species-specific monoclonal antibodies (MABs), RR-402 (Washington Research Foundation, Seattle, USA) and SCP-53, C. trachomatis species-specific MAb, MicroTrak (Syva Co.), genus-specific MAB, Cultureset (Ortho Diagnostics Systems, Inc.) and XC-60, which was obtained in our laboratory and confirmed to be an MAB directed to the species-specific epitope of C. pneumoniae 60 KDa heat-shock protein.

**Preparation of EBs**

EBs of the C. psittaci and C. pecorum strains were purified by the method of Tamura and Higashi. Our preliminary experiments indicated that EB preparations of C. pneumoniae and C. trachomatis strains purified by the method of Tamura and Higashi or Caldwell et al. contained a number of RBs and their fragments. We therefore established a new method for the purification of C. pneumoniae and C. trachomatis EBs. At 72 h postinoculation, infected cells were collected in sucrose-phosphate-glutamate solution (SPG: sucrose 7.5%, KH₂PO₄ 0.052%, Na₂HPO₄ 0.1529%, glutamic acid 0.072%) and homogenized with a Teflon homogenizer. After brief centrifugation at 900 × g for 10 min at room temperature to remove cell debris, the supernatant obtained was layered onto a two-layer cushion (bottom layer, 50% wt/vol sucrose solution; top layer, 30% vol/vol Urografin [3,5-diacetamido-2,4,6-triisobenzoic acid; Schering AG, Berlin/Bergkamen, Germany] in 30 mM Tris-HCl buffer [pH 7.3]); and then centrifuged at 8,000 × g for 60 min at 4°C with an RPS-25 swing rotor (Hitachi, Tokyo, Japan). The precipitate and the turbid bottom layer were suspended together in SPG and centrifuged at 12,000 × g for 30 min at 4°C. The precipitate obtained was suspended in SPG and then treated with DNase and RNase (final concentration; 20 µg/ml each) for 30 min at 37°C. The suspension was layered onto a continuous Urografin gradient column (40 to 52% vol/vol) and centrifuged at 8,000 × g for 60 min at 4°C. Two distinct bands were formed in the gradient column. The presence of a number of complete EBs was confirmed the lower band by electron microscopy. After washing with SPG, the EBs were suspended in SPG and stored at -70°C until required.

**Preparation of EB outer membranes**

Four milliliters of purified EB suspension in Tris buffer was mixed with 6 g of glass beads (0.1 mm in diameter; M and S Instruments Inc., Osaka, Japan) and shaken vigorously on a Cell Mill shaker (Edmund Bübler, Tübingen, Germany) at 70 CPS for 4 min. After centrifugation at 10,000 × g for 60 min at 4°C on
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The specimens were examined with a Hitachi H-500 or JEM-2000 transmission electron microscope at 75 or 80 kV.

To determine the location of lipopolysaccharide (LPS) on the EB surface of all strains, immunogold labeling was performed by the method of Birkelund et al.20. Briefly, the specimen was mounted on a grid covered with a collodion membrane and exposed to anti-LPS MAb (supplied by Hirai K, Faculty of Agriculture, Gifu University, Gifu, Japan), washed with phosphate-buffered saline, and then mixed with a suspension of protein A-gold particles (15 nm in diameter, Amersham Co.). After several washings, the specimen was examined with the Hitachi H-500 transmission electron microscope at 75 kV.

Determination of the spacing of regular structure by Fourier transformation

To determine the spacing of the hexagonal regular structure in the EB outer membrane, diffraction images were obtained by a formula programmed in an IBAS 2000 image analysis system (Zeiss, Germany)20. Briefly, the information in a micrograph was in put into the image memory of the system through a camera, and the Fourier integral was calculated in the computer system, so that the original image in the micrograph was displayed as a diffraction pattern. The average spacing of the regular structure was calculated by measuring of the distance from the origin of the transform (frequency 0) to the diffraction spots. In some cases, the images were reconstituted from the selected spots in the diffraction pattern by the inverse Fourier transform to obtain a clear image without the “noise” present in the original micrographs.

Results

Inclusion morphology and staining properties of clinical isolates of C. pneumoniae

The inclusions of KKpn-1, KKpn-15, KKpn-16 and other 15 clinical isolates positively stained with C. pneumoniae-specific MAbs and the genus-specific MAb (Cultureset), but not with C. trachomatis-specific MAb (MicroTrak) or iodine stain. Their morphology in HeLa 229 cells closely resembled that of C. psittaci and C. pecorum, which are oval and dense in the early stages in the Giemsa staining preparations. The inclusions expanded without compression of the host nucleus. These properties of the 18 clinical isolates obtained in our laboratory were identical with those of TWAR strains, indicating they are the members of the species C. pneumoniae.

Morphology of C. pneumoniae EBs in thin sections and shadowcast preparations

The chlamydial bodies in in situ inclusions of the
C. pneumoniae strains at 60 h postinoculation are shown in Fig. 1. A number of EBs and RBs are seen in the inclusions. There were no distinct differences in the RB morphology of the different strains. The EBs of KKpn-1, KKpn-15, KKpn-16, IOL-207, Kajaani-6, YK-41, and the 15 clinical isolates (Fig. 1a), however, were round and their morphology was identical to that of C. pecorum, C. psittaci, and C. trachomatis EBs. In contrast, the EBs of the TWAR strains had a wide periplasmic space limited by a wavy outer membrane, creating “pear-shaped” profiles (Fig. 1b). A similar morphological difference between TWAR and other strains was seen when the purified EBs were dried in air and shadowcast (Fig. 2). The EBs of C. pecorum, C. psittaci and C. trachomatis strains had a round “fried egg” shape that was indistinguishable from the EBs of strains KKpn-1, KKpn-15, KKpn-16, IOL-207, Kajaani-6, YK-41, and the 15 clinical isolates (Fig. 2a). However, the EBs of

**Fig. 1.** Thin sections of KKpn-15 and AR-39 strains in HeLa 299 cells at 60 h postinoculation. a, Chlamydia pneumoniae KKpn-15; b, C. pneumoniae AR-39 at higher magnification. KKpn-15 EBs have a narrow periplasmic space and are round in shape, whereas AR-39 EBs are enclosed by a wavy outer membrane and are “pear-shaped” in profile. c, KKpn-15 inclusion. d, AR-39 inclusion. Arrowheads point to mitochondria located close to inclusions, but unassociated with the inclusion membrane. Bars; 500 nm.

**Fig. 2.** Shadowcast of purified EBs of Chlamydia pneumoniae KKpn-15 (a) and TW-183 (b) strains. EBs were air-dried on a smooth-surfaced agar plate, transferred to a specimen grid by the pseudoreplica method with collodion, and then shadowcast with Pt-palladium alloy. Bars; 1 μm.
the TWAR strains exhibited the "pear-shaped" morphology with a wide outer-membrane area even by the air-drying method (Fig. 2b).

Further observations on the distribution of mitochondria in the host cells infected with all strains of C. pneumoniae revealed the absence of any mitochondrial association with the inclusion membrane (Fig. 1c, 1d arrowheads). This strongly suggested that the absence of the mitochondria-inclusion association is common in C. pneumoniae multiplication, although mitochondrial association with C. psittaci inclusions is a natural occurrence during the reproductive stage.

**Surface projections and related structures**

To examine the EB envelope and inclusion membrane, freeze-replicas of the inclusion-bearing cells were prepared at 60 h postinoculation. As shown in Fig. 3, the cleaved faces of the EBs and RBs were exposed as convex or concave faces in the replica membrane. Button structures (B structures) or craters were clearly seen in a limited area of the concave faces (Fig. 3, arrowheads). When the inclusion membrane was replicated, many fine particles grouped together within several areas were frequently observed on the convex faces (Fig. 4, arrowheads). The morphological properties of the B structures on the concave faces of in situ chlamydial bodies and the particles on the convex faces of inclusion membrane were identical with those mentioned previously. Therefore, the results obtained in the present study strongly suggested the presence of projections on the surface of the C. pneumoniae EBs and RBs.

To directly confirm the presence of projections on the surface of C. pneumoniae EBs, purified EBs were dried by the critical point drying method and examined by scanning electron microscopy. Projections were observed on a limited area of the EB surface of the C. pneumoniae strains (Fig. 5). The EBs of KKpn-15 were globular in shape (Fig. 5a), whereas those of the TWAR strains (Fig. 5b) had irregular shapes as a result of the presence of a wide periplasmic space. The number of projections on the EBs of each of the strains was not determined.

**Regular structure in the outer membranes**

In a preliminary experiment, it was found that the

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**Fig. 3.** Freeze-replica images of in situ chlamydial bodies and inclusion membranes of strain KKpn-15 at 60 h postinoculation. Chlamydial bodies are cleaved into convex or concave faces. Many B structures are seen (arrowheads). Bar; 1 μm.

**Fig. 4.** Freeze-replica images of an in situ chlamydial inclusion membrane containing strain KKpn-15 at 60 h postinoculation. Fine particles in groups are indicated by arrowheads. The face is rugged over the RB outlines. Bar; 1 μm.

**Fig. 5.** Scanning electron micrograph of EBs of Chlamydia pneumoniae KKpn-15 (a) and AR-39 (b). Both micrographs show hexagonally arrayed projections in a limited area of the surface. Bars; 300 nm.
EB outer membranes of *C. pneumoniae* strains broke into small fragments when their envelopes were treated with 0.5% SDS to remove the inner membrane, whereas those of *C. psittaci* and *C. trachomatis* strains did not. This suggested that the outer membrane of *C. pneumoniae* EBs is fragile in SDS solution. Therefore, the SDS concentration was reduced from 0.5% to 0.125% to isolate the outer membranes of *C. pneumoniae*.

When the outer membranes were negatively stained, the hexagonally arrayed structures were clearly seen despite the difference in chlamydial species (Fig. 6). The dimensions of the structures appeared to be similar in all strains examined. Such regular structures were also seen in the shadowcast preparations only on the inner surface exposed by disintegration of the outer membrane (data not shown). These results were in good agreement with those reported previously, indicating that the inner surface of the EB outer membrane of *C. pneumoniae* strains is also composed of hexagonally arrayed structures.

A computer Fourier transform was carried out to determine the spacing of the hexagonal structure. Fig. 7 shows an example. As a result of the Fourier transform, the fragment of the KKpn-15 outer membrane shown in Fig. 6a yielded the diffraction spots in Fig. 7a. The spots show three different structural units of about 176, 90 and 50 Å, respectively. When reconstruction by the inverse Fourier transform was performed for the innermost spots (176 Å), a clear hexagonal pattern was obtained (Fig. 7b), and the periodicity in the outer membrane of all strains used in the present study was measured in this manner. The results are summarized in Table 1. The average periodicity of all strains was very similar despite the difference in morphology, i.e., the round or pear-shaped profile, in thin sections. These results indicate that the regular structure composing the inner layer of the EB outer membrane is a morphological property common to the EB outer membranes of the genus *Chlamydia*, although the outer membranes of the TWAR EBs show peculiar profiles in thin sections.

**Immunolabeling of LPS on EBs and the diversity of the LPS of *C. pneumoniae***

When immunogold-labeling with anti-LPS MAb was carried out, the surface of the EBs of all chlamydial strains was heavily labeled with protein A-gold particles (Fig. 8). This finding indicated that the immunogold-labeling method was accurate and that the LPS was located on the EB surface.

The IDEIA chlamydia (DAKO Diagnostics Co.) test kit is designed to detect *C. trachomatis* LPS.

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**Table 1. Comparison of periodicity in chlamydial species**

<table>
<thead>
<tr>
<th>Species</th>
<th>Strain</th>
<th>Mean periodicity ± SD</th>
</tr>
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<tr>
<td><em>C. pneumoniae</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td>round-shape</td>
<td>KKpn-15</td>
<td>179 ± 8 Å</td>
</tr>
<tr>
<td></td>
<td>KKpn-16</td>
<td>178 ± 8 Å</td>
</tr>
<tr>
<td></td>
<td>KKpn-1</td>
<td>176 ± 7 Å</td>
</tr>
<tr>
<td></td>
<td>10L-207</td>
<td>177 ± 7 Å</td>
</tr>
<tr>
<td></td>
<td>Kajaani-6</td>
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</tr>
<tr>
<td></td>
<td>YK-41</td>
<td>176 ± 8 Å</td>
</tr>
<tr>
<td>pear-shape</td>
<td>TW-183</td>
<td>173 ± 7 Å</td>
</tr>
<tr>
<td></td>
<td>AR-39</td>
<td>174 ± 8 Å</td>
</tr>
<tr>
<td></td>
<td>AR-388</td>
<td>173 ± 8 Å</td>
</tr>
<tr>
<td><em>C. trachomatis</em></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>L2/434/Bu</td>
<td>176 ± 7 Å</td>
</tr>
<tr>
<td><em>C. psittaci</em></td>
<td>Frt-Hu/Cal 10</td>
<td>175 ± 8 Å</td>
</tr>
</tbody>
</table>

* Periodicity measured from center to center by Fourier transform.

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Fig. 6. Negatively stained EB outer membrane of *Chlamydia pneumoniae* KKpn-15 strain. b, higher magnification of the micrograph in a. The hexagonal structures are seen throughout the outer membrane. Bar: 200 nm.

Fig. 7. Image processing of the hexagonal structures in the outer membrane of strain KKpn-15 EB. a, Diffraction spots obtained by Fourier transformation from the original micrograph shown in Fig. 6b; b, Image reconstituted by inverse Fourier transformation from the six spots of the diffraction shown in Fig. 7a.
Fig. 8. Immunogold-labeling on the surface of EBs of the Chlamydia pneumoniae KKpn-15 (a) and TW-183 (b) strains. The samples were treated with monoclonal antibody directed to Chlamydia psittaci LPS, and then labeled with protein A-gold particles. Intact EBs and EB envelopes are heavily labeled with the protein A-gold particles. Bars; 500 nm.

Table 2. Diversity of Chlamydia pneumoniae strains in reaction intensity to the IDEIA test kit

<table>
<thead>
<tr>
<th>Strain</th>
<th>Shape of EB</th>
<th>EB number per assay</th>
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<tbody>
<tr>
<td>IOL-207</td>
<td>round</td>
<td>$6.0 \times 10^6$</td>
</tr>
<tr>
<td>TW-183</td>
<td>pear-shape</td>
<td>$7.0 \times 10^6$</td>
</tr>
<tr>
<td>AR-388</td>
<td>pear-shape</td>
<td>$8.4 \times 10^6$</td>
</tr>
<tr>
<td>Kajaani-6</td>
<td>round</td>
<td>$1.2 \times 10^6$</td>
</tr>
<tr>
<td>AR-39</td>
<td>pear-shape</td>
<td>$2.4 \times 10^6$</td>
</tr>
<tr>
<td>KKpn-1</td>
<td>round</td>
<td>$2.8 \times 10^6$</td>
</tr>
<tr>
<td>YK-41</td>
<td>round</td>
<td>$4.0 \times 10^6$</td>
</tr>
</tbody>
</table>

The numbers of EBs are shown in increasing order.

which is a major genus-specific (genus-common) antigen, and consequently the LPS of C. psittaci and C. pneumoniae also reacts with the IDEIA kit. When the IDEIA kit was tested with particle number of purified C. pneumoniae EBs as a parameter, the reaction occurred positively at different intensities. No correlation was found between reaction intensity and EB morphology (Table 2). However, it was not clarified whether the different reactivities of C. pneumoniae strains resulted from differences in the quality or quantity of LPS on the EB surface, although the chemical structure of the antigenic epitope in LPS has been proposed.

**Discussion**

The morphology of C. pneumoniae TWAR strains was compared with that of C. trachomatis L2 and C. psittaci Cal 10. The TWAR EBs were shown to be "pear-shaped", with a large periplasmic space, in thin sections, whereas those of C. trachomatis and C. psittaci were observed to be round. This morphological difference was confirmed again in the present study. However, the EBs of KKpn-1~18 and YK-41 strains, which were recognized as C. pneumoniae on the basis of their properties, including reactivity with C. pneumoniae-specific MAbs and stainability with iodine, were found to have a narrow periplasmic space in sections and shadowcast preparations.

Carter et al. demonstrated that the EB morphology of C. pneumoniae IOL-207 strain was also round in thin sections and indistinguishable from that of C. trachomatis and C. psittaci EBs. Similarly, Popov et al. demonstrated round EBs of C. pneumoniae Kajaani-6 strain. These results, together with the results obtained in the present study, strongly suggest that the "pear-shape" profiles of EBs in thin sections are not common among C. pneumoniae strains, and thus that the pear-shape is not valid as a morphological criterion for identifying C. pneumoniae species.

Based on a series of electron-microscopic studies on C. psittaci EBs, Matsumoto proposed a possible diagram showing the projections grouped together within a limited surface area and their related internal structures. In a scanning electron microscopy study, Gregory et al. and Matsumoto et al. found the presence of projections on the EBs of many strains of C. trachomatis and C. psittaci, and in our study, scanning electron microscopy showed projections not only on TWAR EBs but on KKpn-1, KKpn-15, KKpn-16, and YK-41 EBs as well. Moreover, the presence of B structures on the concave face of EBs in freeze replicas strongly suggests that the projections are common to all members of the genus Chlamydia, thereby supporting the proposition by Gregory et al. that the projections are a phenotypic marker for recognizing members of the genus Chlamydia.
Matsumoto examined *C. psittaci* Cal 10 inclusions isolated from infected cells by thin sectioning and freeze-replica techniques and concluded that the hexagonally arrayed particles on the convex face of the inclusions were projections directly connecting the RBs to the host cytoplasm. Matsumoto also reported the presence of identical particles on the convex face of *in situ* inclusions of *C. trachomatis* L2 and *C. psittaci* strains. In the present study, the particles were again seen on the convex face of the replicated inclusions of *C. pneumoniae* strains. It is, therefore, very likely that the RBs of *C. pneumoniae* strains also possess projections and that the direct connection between RBs and inclusion membranes with the projections is a common phenomenon during multiplication of the genus *Chlamydia*.

Matsumoto examined the outer membranes of *C. psittaci* EBs and reported that the dimensions of the hexagonal structures had a periodicity of 167Å by an optical transformation method using laser beams. However, in the present study, the periodicity in the Cal 10 outer membranes was estimated to be 175Å by the Fourier transform in the computer system. This difference (approximately 5%) appeared to be due to the difference in methods of measurement and seemed unavoidable. It was noteworthy that the EBs of the *C. pneumoniae* strains examined possessed hexagonal structures in the inner surface of the outer membrane and that no marked difference in periodicity was recognized between the round-shaped EBs (KKpn-1, KKpn-15, KKpn-16, IOL-207, Kajaani-6, YK-14) and the pear-shaped EBs (TW-183, AR-39 and AR-388). Based on these findings, it is very likely that the hexagonal structure composing the inner layer of the EB outer membrane is a morphological property common to the genus *Chlamydia*, although the outer membranes of the TWAR EBs show pear-shaped profiles. On the basis of a comparative study of EB and RB of *C. psittaci* Cal 10 strain, Matsumoto and Manire suggested a correlation between the presence of hexagonal structures and the rigidity of the chlamydial bodies. In addition, outer membranes obtained from abnormal, giant RBs formed in the presence of penicillin were also fragile and defective in the hexagonal structure. Routine observations on the multiplication of many chlamydial strains with round EBs, such as *C. psittaci* and *C. trachomatis*, suggest that a large RB outer membrane may be diminished in size into a small EB outer membrane. If this were not so, a number of fragments or vesicles derived from the RB outer membranes would be regularly seen in the inclusion during the conversion from RB to EB during maturation. A similar reduction of the RB outer membrane should occur during the maturation of the KKpn-1—KKpn-18 strains, because their EBs are enclosed tightly with a round outer membrane. This led us to speculate, on the other hand, that this reduction in size of the RB outer membrane is incomplete in the strains with pear-shaped EBs, such as *C. pneumoniae* TWAR. Thus, the relationship between the formation of the hexagonal structure and the reduction in size of the RB outer membrane is still unclear.

No mitochondria associated with the inclusion-containing *C. pneumoniae* strains used were detected. This finding is in good agreement with the results reported previously. Therefore, it is likely that the absence of the mitochondria/inclusion association is also a common phenomenon in *C. pneumoniae* species.

*C. pneumoniae* was classified in 1989 as a new species on the basis of EB morphology, serology, and DNA analysis of the TWAR strains. Since then, several *C. pneumoniae* strains, including our isolates, have been established from patients with respiratory diseases. It is noteworthy that in spite of the striking difference in EB morphology, only one serotype has been identified. Although humans have been thought to be the natural reservoir of *C. pneumoniae*, similar chlamydiae have been isolated from a horse and koala bear. These findings, together with the variety of reactivity of many *C. pneumoniae* isolates in the IDEIA kit led us to speculate that different *C. pneumoniae* serotypes exist.

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Chlamydia pneumoniae の形態学的解析

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Chlamydia pneumoniae は1989年に新種となったクラミジアであり、従来その分離培養は困難とされていた。最近我々は、急性呼吸器感染症患者の咽頭スワブから計18株の分離に成功した（KKpn-1からKKpn-18と命名）。今回これら臨床分離18株を中心に超微構造を、既存のC. pneumoniae TW-183, AR-39, AR-388（以上TWAR株）、IOL-207, Kajaani-6, YK-41株、Chlamydia psittaci Frt-Hu/Cal10株、Chlamydia trachomatis L2/434/Bu株ならびにChlamydia pecorum Bo/E58株と比較検討した。TWAR株の基本小体（elementary body; EB）は、広いperiplasmic spaceをもつ西洋梨状を呈し、C. pneumoniaeを新種としたクライテリアの1つとして挙げられている。しかし、今回対象としたKKpn-1～KKpn-18株 EBは円形でC. psittaci, C. trachomatis, C. pecorum のEB同様に狭いperiplasmic spaceを有していた。また同様にIOL-207, Kajaani-6, YK-41株 EBも円形であったことから、EBの広いperiplasmic spaceはC. pneumoniaeの形態学的特徴とはいえないことが判明した。しかし、クラミジア独特の表面突起（surface projection; SP）およびB構造、封入体膜上のSP集団が用いたクラミジア全株に認められ、EBの基本形態に差がないことが判明した。したがってC. psittaci, C. trachomatis同様SPを介するhost-Chlamydia interactionのあることが示唆された。EBの剛性は外膜に依存していると考えられ、C. psittaci, C. trachomatisの外膜内面はhexagonalな規則的構造（hexagonal structure; HS）で形成され、脆弱な網様体（reticulate body; RB）外膜にはHSがないことから、このHSと剛性の間の相関性が強く示唆されている。TWAR株のEBが西洋梨状を呈していることや、超音波処理・界面活性剤（SDS）処理に対する脆弱性・感受性が異なることからHSに相違がみられると予想されたが、いずれの株においてもHSが認められ、その周期性は種間、株間で有意差はなかった。