Intracellular persistence of chlamydial major outer-membrane protein, lipopolysaccharide and ribosomal RNA after non-productive infection of human monocytes with Chlamydia trachomatis serovar K

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Summary. The replication of Chlamydia trachomatis serovar K was studied in human peripheral blood monocytes (PBMo). The intracellular fate of the bacteria was examined by determining the presence of chlamydial major outer-membrane protein (MOMP), lipopolysaccharide (LPS) and ribosomal RNA (rRNA). In-vitro infection of PBMo with C. trachomatis serovar K was not productive. However, chlamydial MOMP antigen, demonstrated by immunofluorescence, was present in PBMo for up to 14 days. Infected monocytes also contained chlamydial rRNA, measured by in-vitro hybridisation, and LPS, measured by enzyme immunoassay, for up to 14 days. These data are compatible with the hypothesis that the infection of PBMo with C. trachomatis may play a role in the systemic distribution of chlamydial antigens, leading to systemic manifestations of urogenital chlamydial infection.

Introduction

Chlamydia trachomatis is an obligately intracellular bacterium whose various serotypes cause several ocular, pulmonary and genital diseases in man. A common feature of chlamydial infections is their tendency to persist if not adequately treated. C. trachomatis biovar II (serovars L1-L3) causes lymphogranuloma venereum (LGV), characterised by penetration of the epithelial barrier and infection of inguinal lymph nodes with subsequent ulceration. In contrast, infection with C. trachomatis biovar I (serovars A-K) tends to be locally restricted to epithelial mucosal cells and is generally thought to be characterised by its lack of invasiveness. Recently, however, chlamydial antigen has been demonstrated intra-articularly in the inflamed joints of patients with Reiter's syndrome or reactive arthritis after urogenital infection with C. trachomatis serovars D–K. The intra-articular chlamydial antigen was observed primarily in synovial macrophages.

Therefore, mononuclear phagocytes may play a key role in the systemic distribution and extra-urogenital persistence of C. trachomatis in chlamydial diseases in man. Nevertheless, the mechanisms of chlamydial antigen persistence in mononuclear cells are poorly understood. In this study we investigated the replication of C. trachomatis serovar K in human monocytes and the fate of chlamydial components.

Materials and methods

Preparation of infectious elementary bodies (EB)

Infectious EB of C. trachomatis serovar K (UW/31/Cx; Washington Research Foundation, Seattle, USA) were grown in HEp-2 cells (human larynx carcinoma epithelial line) in RPMI 1640 medium, supplemented with fetal calf serum (Biochrome, Berlin, Germany) 10% v/v, L-glutamine 1% w/v, vancomycin 0.1% w/v, gentamicin 0.1% w/v, i.e., “growth medium”, and containing glucose 1% w/v and cycloheximide 1 μg/ml, i.e., “infection medium” (all chemicals were from Sigma). Cultures were incubated for 48 h at 37°C in an atmosphere of CO2 5% w/v. The organisms were purified in a discontinuous gradient of Urografin (Schering, Berlin, Germany) 10% v/v, L-glutamine 1% w/v, vancomycin 0.1% w/v, gentamicin 0.1% w/v, i.e., “growth medium”, and containing glucose 1% w/v and cycloheximide 1 μg/ml, i.e., “infection medium” (all chemicals were from Sigma). Cultures were incubated for 48 h at 37°C in an atmosphere of CO2 5% w/v. The organisms were purified in a discontinuous gradient of Urografin (Schering, Berlin, Germany) by ultracentrifugation as described by Caldwell et al.12 Purified EB were resuspended in sucrose phosphate buffer (0.01 M sodium phosphate, 0.25 M sucrose, 5 mM L-glutamic acid, pH 7.2), frozen in 0.1-ml volumes and stored at −80°C.

Determination of chlamydial infectivity

The infectivity of purified EB or infected monocytes

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was determined by titration on HEp-2 cells. Infected mononuclear phagocytes were treated to release bacteria by thawing the frozen samples in an ice-bath over a 15-min period followed by sonication for 2 min. Suspensions containing HEp-2 cells $3 \times 10^6$/ml in growth medium were seeded as 0.1 ml/well in 96-well microtitration plates (Nunc, Wiesbaden, Germany) and grown for 2 days at 37°C in an atmosphere of CO$_2$, 5% v/v to confluent monolayers. Ten-fold dilutions of infective samples in infection medium were added as 0.05 ml/well and the cultures were incubated for 48–60 h at 37°C. For detection of typical large inclusion bodies by light microscopy, the cells were fixed with absolute ethanol and an indirect immunoperoxidase assay$^{15}$ was performed as follows. Cells were first incubated for 1 h with the serum of a *C. trachomatis* antibody-positive patient with a specific IgG titre of $> 256$ (Ispazyme test, Medac, Hamburg, Germany) and further incubated with a peroxidase-conjugated goat-anti-human IgG antibody (Sigma) diluted 1 in 40 in PBS. After addition of the substrate 4-chloro-1-naphthol (Savyon, Beer-Sheva, Israel), 1 in 2 in PBS, inclusion bodies were seen as black dots which could be easily identified and counted by light microscopy. The numbers of inclusion bodies/volume were expressed as inclusion forming units (IFU)/ml.

Unabsorbed chlamydiae were removed 2 h post-infection (p.i.) by washing five times in HBSS. Fresh growth medium was added to the cells. A control microscopic examination was performed daily and the viability of PBMo was checked by frequent trypan blue dye-exclusion tests. At least 80% of the PBMo were alive on day 14. The cells were harvested at intervals (0, 1, 7 and 14 days p.i.) by scraping carefully with a rubber policeman. Samples were stored at $-70°C$.

To exclude any interference of plastic-adherent extracellular chlamydia with the numbers of infective particles inside PBMo, a control experiment was performed. PBMo were first cultured and infected in 250-ml culture flasks. After incubation for 2 h at 37°C, infected adherent PBMo were washed five times with HBSS, but then scraped carefully from the plastic base of the culture flask. The PBMo suspensions were washed three times with HBSS and then (1–2) $\times 10^6$ cells/ml were seeded in 2-ml volumes in a six-well plate. At days 0, 1, 7 and 14, re-adherent cells were harvested and washed three times with HBSS.

**Immunofluorescence microscopy of chlamydial particles**

Chlamydial major outer membrane protein MOMP was detected by a direct immunofluorescence technique.$^{14}$ After harvesting infected PBMo, cytocentrifuge smears, each containing c. $5 \times 10^5$ cells, were prepared (Cytospin, Shandon). They were fixed for 10 min with methanol 100% and were further incubated with a fluorescein-conjugated murine monoclonal antibody (MAb; Sva, Palo Alto, CA, USA). This MAb is directed against a common MOMP epitope of EB and the detection limit was 10 IFU/ml.

**Determination of chlamydial lipopolysaccharide (LPS)**

The presence of chlamydial LPS in infected PBMo was determined by an enzyme immunoassay (IDEEA* Chlamydia test, Novo BioLabs, Cambridge) with genus-specific Fab-fragments of a MAb against chlamydial LPS and an alkaline–phosphatase substrate with an alcohol-oxidoreductase-diaphorase amplification system. PBMo were harvested and suspended in 1 ml of EIA transport medium (IDEEA Chlamydia test). The determination of chlamydial LPS was

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Fig. 1. Infectivity of C. trachomatis. One-step growth curves of chlamydiae after infection of PBMo with C. trachomatis at MOI 1 (▲; n = 6), 0·5 (■; n = 3) and 0·1 (○; n = 4). The numbers of chlamydiae were determined by titration on HEp-2 cells and expressed as IFU/ml. In control experiments (n = 4), growth in HEp-2 cells infected with C. trachomatis at MOI 1(○) was determined. Results are indicated as means, bars represent SD.

performed according to the manufacturer’s instructions. Relative concentrations of LPS in infected monocytes are expressed as extinction at 492 nm. The cut-off for each test procedure was calculated as the mean of three negative controls plus 0·2. The sensitivity of the EIA to detect chlamydial LPS in cell cultures, as evaluated by a serial dilution analysis of EB suspended in transport medium, was 100 IFU/ml.

Determination of chlamydial ribosomal RNA

Chlamydial rRNA was measured by in-vitro hybridisation with a single-stranded chemiluminescent-labelled DNA probe. The commercially available DNA probe (PACE 2, Gene-probe, San Diego, USA) recognises all 15 human serovars of C. trachomatis and reacts specifically with complementary rRNA sequences of C. trachomatis. After harvesting mononuclear cells, samples containing (3–6) x 10⁶ cells were transferred to 1·5-ml reaction tubes and centrifuged at 30000 g (Biofuge, Heraeus, Hamburg, Germany). The pellets were resuspended in 100-μl volumes of lysis buffer (Gene-probe) and incubated for at least 3 days. For the test procedure, samples were transferred to test tubes and 100 μl of the acridinium ester-labelled probe reagent was added to each specimen. After a differential hydrolysis to inactivate the non-hybridised DNA, light emission from the DNA-RNA hybrid was measured in a luminometer (Leader I®, Biermann, Bad Nauheim, Germany). Concentrations of rRNA are expressed as relative light units. Three negative and one positive control were included in each test kit. Non-infected
PBMo were used as additional negative controls. According to our experience, the test has a detection limit for chlamydial rRNA of at least 50 IFU/ml.17

Results

To study the fate of C. trachomatis in PBMo the presence of specific MOMP, LPS and rRNA was investigated over a period of 14 days after infection.

Infectivity of C. trachomatis in PBMo cultures

PBMo could not be infected productively with C. trachomatis serovar K. The one-step growth curves of C. trachomatis in PBMo at MOI 0.1, 0.5 and 1 are illustrated in fig. 1. After the removal of unabsorbed bacteria at 2 h p.i. (i.e. day 0), 3–5% of the initial inoculum could be detected by titration on HEP-2 cells. The number of infectious particles further decreased during the first day. After day 1, with an MOI of 0.5 or 1, inclusions were scarcely detectable; no IFU were found at MOI 0.1. In none of the experiments was a productive infection observed on day 7 or 14 p.i. In contrast, infection of HEP-2 cells with C. trachomatis at MOI 1 was productive. After an initial decline in the number of infective particles after 1 day, indicating the transformation of infectious EB into non-infectious RB in the life-cycle of chlamydia, the number of infective particles had reached a high level by day 4. Typical inclusion bodies were visible 1–2 days p.i., indicating a productive infection with the formation of new infectious EB.

Thus, C. trachomatis did not infect PBMo productively, in contrast to its ability to replicate in HEP-2 cells. The experiments to exclude the possible influence of plastic-adherent extracellular C. trachomatis on the infectivity for PBMo demonstrated no alteration in these kinetics.

Detection of chlamydial MOMP by immunofluorescence

In contrast to the absence of chlamydial replication, the determination of chlamydial MOMP revealed a considerable number of fluorescent particles (f.p.)/cell at MOI 0.5 and 1 throughout the cultivation period (fig. 2). Mostly single and occasionally multiple round, bright green particles were identified in infected PBMo (fig. 3), compared with only non-specific staining in uninfected PBMo. No clusters of f.p. suspected of being inclusion bodies were found and free
extracellular f.p. could be seen only rarely. The number of f.p. decreased considerably during the first day (paired Student's t test, MOI 1: p < 0.005; MOI 0.5: p < 0.02) and from days 1–7 p.i. (MOI 1: p < 0.01; MOI 0.5: n.s.). A constant amount of MOMP was observed from day 7 to day 14 p.i. with MOI 1 and 0.5. On day 14, an average of 10% of the f.p. at day 0 could still be detected in infected PBMo. With MOI 0.1, no persistence of MOMP was observed. Detectable numbers of f.p. could be seen only during the first day p.i. Control experiments, as described above, to exclude any influence of plastic-adherent extracellular EB showed no alterations in these kinetics.

Thus, in infected PBMo, a persistence of MOMP could be observed over the entire culture period of 14 days.

**Determination of chlamydial LPS**

The determination of chlamydial LPS in infected PBMo showed dose-dependent concentrations of this cell-wall constituent for up to 14 days p.i. (fig. 4). LPS was identified in PBMo on the first day at all MOI. The concentrations of LPS in PBMo tended to decrease during the cultivation period. At day 14 p.i., however, significant amounts of LPS were still detectable in cells with an MOI of 1 and 0.5, but not at MOI 0.1. No chlamydial LPS was found in non-infected PBMo.

**Determination of chlamydial rRNA**

To determine whether chlamydial cell-wall components, or chlamydial nucleic acids, or both, persisted in PBMo, the amount of chlamydial rRNA was determined. After infection of PBMo, a significant decline in the concentrations of rRNA was found during the first day p.i. (MOI 1: p < 0.05; MOI 0.5: p < 0.05; fig. 5). From day 1 to day 7 a further decrease in chlamydial rRNA concentrations (MOI 1: p < 0.01; MOI 0.5: p < 0.01) was demonstrated but the amount of rRNA did not decline further from day 7 to day 14. Approximately 10% of the input rRNA concentrations could be detected at 14 days p.i. In contrast, at MOI 0.1, detectable amounts of rRNA could only be detected during the first day p.i. No chlamydial rRNA was detected in controls of uninfected PBMo.
Fig. 5. Determination of kinetics of chlamydial rRNA by hybridisation with a single-stranded DNA probe in PBMo infected with C. trachomatis at MOI 1 (▲; n = 6), 0.5 (■; n = 3) and 0.1 (●; n = 4). Results are indicated as means, bars represent SD.

Discussion

Clinical and in-vitro experimental evidence for persistent infection by C. trachomatis has been described frequently. A possible involvement of human mononuclear phagocytes in latent and persistent chlamydial infection has been recognised, but the mechanisms of bacterial or antigen persistence are still poorly understood. So far, there have been few studies of the in-vitro interaction between monocytes or macrophages and chlamydia. Limited replication of C. psittaci has been described in human monocytes and their number was reported to increase during the differentiation of the monocytes into macrophages. C. trachomatis serovars B and L₂ have been shown to grow to a limited extent in mouse peritoneal macrophages. C. trachomatis serovars B and L₂ have been shown to grow to a limited extent in mouse peritoneal macrophages. With human monocytes and monocyte-derived macrophages it was found that serovar L₂ could infect productively in-vitro matured macrophages, while infection in non-differentiated PBMo was non-productive. In a similar study, Bard and Levitt confirmed these observations, although they found some (i.e. < 0.5%) inclusion bodies in monocytes, in contrast to an absolute lack of productive infection described previously.

In contrast to the productive infection of macrophages with the LGV biovar, Yong et al. demonstrated a non-productive infection with various trachoma biovar I strains (B, C and I). Comparing the fate of C. trachomatis serovars L₂ and K in human monocyte-derived macrophages we came to the same conclusions. Yong et al. suggested, from their ultrastructural investigations with monocytes cultured for < 1 week, that both biovars were degraded after phagosome-lysosome fusion. However, the possibility of a persistent or latent infection with the trachoma biovar I in monocytes and macrophages has not been investigated in detail. Therefore, in the present study we examined whether PBMo may be infected persistently by C. trachomatis serovar K.

In accordance with previous studies, serovar K, as a member of biovar I, did not infect PBMo productively. The yield of infective particles during the first day p.i. has to be interpreted as the remnant of the initial inoculation. Nevertheless, chlamydial MOMP antigen was identified intracellularly during the entire cultivation period of 14 days. The fluorescent particles in PBMo were mostly single and rarely multiple. As would be expected in the presence of inclusion bodies, intracellular agglomerates could not be detected. The
observed immunofluorescence staining pattern is consistent with a lack of or insufficient degradation of chlamydial cell walls expressing MOMP antigens. It has long been known that chlamydial cell walls—especially those of the EB—are highly resistant to environmental conditions and intracellular degradation. However the persistence of chlamydial envelopes consisting of MOMP and LPS would not explain the present finding of the persistence of the more fragile chlamydial rRNA in PBMo. Owing to the presence of eukaryotic ribonucleases, free rRNA is assumed to be a target for rapid enzymatic digestion.

Our data suggest that infection of human monocytes with C. trachomatis serovar K in vitro leads to an intracellular persistence of complete chlamydial organisms. The findings have several implications for the understanding of systemic manifestations of urogenital infections with C. trachomatis. The in-vitro observations may explain recent studies demonstrating the presence of non-infectious particles in samples from joints of patients with Reiter’s syndrome, sexually-acquired reactive arthritis and undifferentiated arthritis. Furthermore, the most recent detection of rRNA in synovial membrane biopsies and synovial fluid cells from such patients is in full accord with the in-vitro persistence of rRNA in mononuclear phagocytes shown in this study. Although the results obtained in vitro do not necessarily reflect the situation in vivo, they support the hypothesis that monocytes take up C. trachomatis in the genital tract and distribute the organism in a Trojan-horse manner through the blood circulation into the joints. Subsequently, the infected PBMo, as precursors of sessile tissue macrophages such as synovial membrane cells, may induce arthritis by production and secretion of pro-inflammatory cytokines, recently described for in-vitro infection of macrophages with C. trachomatis serovar K.

Further studies are needed to elucidate whether the chlamydiae persist as viable organisms only temporarily arrested in a non-productive or latent state of the growth cycle, or, alternatively, as defective non-viable organisms resistant to degradation by the phagocytic host cell.

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References


