In situ hybridisation and direct fluorescence antibodies for the detection of *Chlamydia trachomatis* in synovial tissue from patients with reactive arthritis

J Berlau, U Junker, A Groh, E Straube

Abstract

Background—*Chlamydia trachomatis* is associated with Reiter’s syndrome and reactive arthritis but the form in which the organism survives in synovial cells is unclear.

Aim—To compare in situ hybridisation with direct fluorescence in the detection of inapparent chlamydial infection in synovial tissue.

Methods—Synovial tissue from four patients with reactive arthritis patients was examined using biotin labelled probes for chlamydial DNA and fluorescein isothiocyanate (FITC) labelled monoclonal antibodies against the major outer membrane protein.

Results—In two of the four patients, evidence of chlamydial infections was detected by in situ hybridisation in parallel sections but not with FITC labelled monoclonal antibodies.

Conclusions—Detection of chlamydial DNA by in situ DNA hybridisation may be a better way to identify chlamydial infection in synovial tissue than phenotype targeting with FITC conjugated antibodies, which is used as a standard procedure for screening clinical specimens for chlamydia.

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Keywords: *Chlamydia trachomatis*; reactive arthritis; in situ hybridisation.
The fifth patient had post-traumatic arthritis and was operated on after a severe fall onto the right knee. This patient did not show any signs of chlamyidal infection (synovial fluid PCR negative, no IgM and IgG antibodies to chlamydia), and sections of synovial tissue from this patient were used as a standard negative control.

Specimens were embedded in paraffin, cut into 5 µm sections, and deposited on silane treated slides in the Institute of Pathology, University of Jena. Between 10 and 12 slides, each containing three biopsy sections from each patient, were provided and analysed.

McCoy cell cultures grown on cover slips infected with various concentrations of C trachomatis were used as standard positive controls and prepared as described previously.11

IN SITU HYBRIDISATION

After removal of paraffin as described by Lewis et al.,12 fixation in 4% paraformaldehyde (Sigma) for 15 minutes, and washing in phosphate buffered saline (PBS), the slides were incubated for 20 minutes with 0.2 M HCl. They were washed again followed by incubation for five minutes at 37°C with a 1 µg/ml proteinase K solution (Sigma), then washed in 0.1 M glycine in 0.1 M PBS, and once in 2 × SSC (NaCl/sodium citrate). Specimens were prehybridised at 42°C for six hours in a solution containing 6 × SSC, 45% formamide deionised, 5 × Denhardt’s solution, and 100 µg/ml salmon sperm DNA (all from Sigma). The sections on the slides were denatured for four minutes at 94°C on a heating block and hybridised with denatured probe. Probe denaturation was performed by boiling for five minutes, followed by cooling in ice. Hybridisation was performed by boiling for 18 hours overnight in the same prehybridisation buffer containing 250 ng/ml probe. After hybridisation slides were washed in 2 × SSC at room temperature, followed by a further wash in 2 × SSC and another in 0.2 × SSC, both at 42°C. Free binding sites were blocked with 3% dried milk and the slides were washed in 0.1% Tween 20 in PBS. Following this, streptavidin–alkaline phosphatase (Vector) at 1:100 in 0.1% Tween 20 in PBS was applied to all slides for 20 minutes, followed by one wash in PBS and a second in carbonate buffer (pH > 8.5) containing 0.01 M sodium carbonate and 0.01 M sodium bicarbonate (both from Sigma). The slides were then incubated with 5-bromo-4-chloro-3-indolyl phosphate and nitroblue tetrazolium substrate (Sigma), prepared according to the manufacturer’s instructions. Slides were washed in water, counterstained with fast green (Sigma), mounted in glycerol gelatine (Serva), and read by light microscopy. At least 10 slides each containing three tissue sections were investigated for each patient.

DIRECT FLUORESCENCE CYTOLOGY

Slides and cells on coverslips were prepared as described above. The cells were fixed with absolute methanol for 10 minutes and incubated at 37°C for 30 minutes with a fluorescein isothiocyanate conjugated monoclonal antibody (Syva) targeting the MOMP of C trachomatis, washed in PBS, and mounted as recommended by the manufacturer. Slides were read the same way as in situ hybridisation slides.

Initial studies of the newly developed in situ hybridisation system addressed the question of sensitivity. McCoy cells were infected with dilutions of infection forming units of C trachomatis D IC Cal 8 ranging from 0 to 103 as described.11 After 48 hours, these infected cells were hybridised with both probes and incubated with labelled monoclonal antibodies as described above.

Results

The initial study comparing the sensitivity of in situ hybridisation and direct fluorescence antibodies in infected McCoy cells showed similar numbers of signals for both methods (infection of 105 cells with 104 IFU: 8–10 inclusions per coverslip detectable). Positive signals were confined to inclusions and uninfected cells did not show any signals.

MOMP-PCR examination of synovial fluid of patients 2 and 4 showed positive results, and patients 1 and 3 showed negative results. Examination of synovia of all five patients with FITC labelled monoclonal antibodies against the MOMP showed negative results (data not shown).

The results of the investigation of synovial tissue from four patients are shown in table 1. In two of the samples from patients with reactive arthritis there were unequivocal hybridisation signals in the synovial tissue. These signals were observed in the subsynovial cell layer just below the lining cells (fig 1) or in deeper layers (fig 2) and seem to be located within cells rather than being distributed through extracellular compartments (fig 1). Five or six sections, depending on the probe used, of the 36 sections from patient 2 showed such positive signals just below the lining cells. The five parallel sections were positive with both probes. In patient 4 there were slightly fewer positive signals, all in deeper layers. Three of the parallel sections were positive with both probes (table 1). None of the 24 parallel sections of the negative control (patient 5) showed any hybridisation signal with either probe. When tested with FITC conjugated monoclonal antibodies, none of 9–15 parallel sections from each patient gave positive results.

Table 1 Comparison of gene probes p-Ctm and p-Ctp with FITC labelled monoclonal antibodies for the detection of Chlamydia trachomatis in synovial tissue of patients with reactive arthritis

<table>
<thead>
<tr>
<th>Patient (C trachomatis IgG titre)</th>
<th>Diagnosis and duration (years)</th>
<th>p-Ctm</th>
<th>p-Ctp</th>
<th>FITC mAB</th>
</tr>
</thead>
<tbody>
<tr>
<td>(1) Male (1:32)</td>
<td>ReA (4)</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>(2) Male (1:32)</td>
<td>ReA (1)</td>
<td>+ (5)</td>
<td>+ (6)</td>
<td>-</td>
</tr>
<tr>
<td>(3) Male (1:64)</td>
<td>ReA (1)</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>(4) Female (1:32)</td>
<td>ReA, RA (2)</td>
<td>+ (3)</td>
<td>+ (5)</td>
<td>-</td>
</tr>
</tbody>
</table>

FITC mAB, fluorescein isothiocyanate conjugated monoclonal antibody; RA, rheumatoid arthritis; ReA, reactive arthritis; +, positive; −, negative; number of positive tissue sections in brackets.
Detection of Chlamydia trachomatis in synovial tissue

Discussion

It has been known for a long time that, along with other bacteria, C trachomatis plays a role in diseases such as reactive arthritis and Reiter’s syndrome. Nevertheless, owing to the failure to isolate microorganisms from the affected joint, the term “reactive” was used in 1969 to describe an obviously sterile synovitis following an infection localised elsewhere in the body. The apparent absence of viable organisms, together with the demonstration of bacterial antigens in synovium and synovial fluid, led to the view that both reactive arthritis and Reiter’s syndrome are immune mediated diseases. Intra-articular chlamydial antigen has been detected in inflamed joints, and intracellular persistence of antigens such as major outer membrane protein and lipopolysaccharide in cells has also been shown, the detection of RNA indicating living organisms at the site of infection. Some investigators have therefore postulated an inapparent infection which is characterised by persistence of the whole bacterium in a viable state. In this state, the bacterium is undetectable by phenotypic screening methods but can be seen using nucleic acid directed methods. Such organisms may play a crucial role in recurrent arthritis as well as in an obviously sterile synovitis.

To investigate the hypothesis that inapparent chlamydial infections occur in the joints of patients with reactive arthritis, we employed an in situ hybridisation method consisting of two PCR generated probes directed against the endogenous plasmid and the MOMP gene. In two of the four patients, evidence of chlamydial infection was detected by in situ hybridisation in parallel sections. In both the cases, the signals were concentrated in a subsynovial layer but not in the synovial lining itself. The size of the inclusions indicates that the organisms might be in a reticulate body-like state rather than in an elementary body-like state. Despite having equal sensitivity, direct fluorescence antibodies—which are viewed as the gold standard for screening for chlamydia—failed to detect these organisms. This might indicate the suppression, or even the absence, of synthesis of the major outer membrane protein, which is normally found in both reticulate and elementary bodies. Thus the results of our study may partly explain the failure to culture chlamydia from synovial tissue of patients with reactive arthritis, and they confirm the view of an inapparent chlamydial infection in at least some patients with reactive arthritis, as postulated by other investigators. This might even suggest a previously undescribed stage of the chlamydial life cycle. Such inapparent organisms could be critical in the pathogenic process leading to arthritis. It will be the task of further investigations to screen larger numbers of patients and characterise these unusual organisms to determine the specific cell types harbouring them.

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