Use of a commercial PCR kit for detecting *Chlamydia trachomatis*

I W Smith, C L Morrison, C Patrizio, A McMillan

**Abstract**

**Aims**—To evaluate a commercial polymerase chain reaction (PCR) kit for the detection of *Chlamydia trachomatis*.

**Methods**—Two hundred and fifty seven genital specimens, which had been submitted in 2SP medium for chlamydial isolation and subsequently stored at −70°C, were retrospectively examined by a commercial PCR kit which detects chlamydial plasmid DNA. Culture negative, PCR positive specimens were examined by immunofluorescence and an in-house major outer membrane protein (MOMP)-PCR.

**Results**—All 49 specimens which were culture positive were also PCR positive. Another 14 specimens were also PCR positive. After resolution of these results by immunofluorescence and a PCR assay for MOMP the sensitivity for PCR was 98-4% and that of culture 79%. The specificities were 99-5% and 100%, respectively.

**Conclusions**—This kit, which is highly sensitive and specific, is straightforward to use and has a built-in safeguard against cross contamination. The role of this test in the examination of routine genital specimens from patients with uncomplicated chlamydial infection is questionable due to its expense. It may have a role in the investigation of trachoma or infertility, however, where it has been shown that DNA can be detected when culture is unsuccessful.

(J Clin Pathol 1993;46:822–825)

*Chlamydia trachomatis* is the commonest cause of urethritis in Edinburgh, being seven times more common than gonorrhoea. Gonococcal urethritis declined from 58 to 10 per 100 000 men between 1984–1987; the incidence of chlamydial urethritis has levelled out or is rising slightly. Although they can be symptomless, especially in women, chlamydial infections can produce salpingitis with subsequent infertility and an increased risk of ectopic pregnancy. In men, especially those under 35 years of age, the infection can also cause epididymitis. Neonates are also at risk of infection from infected parents. As the infection is amenable to antibiotic treatment it is important to identify infected patients and their partners to eradicate infection. Various methods have been used to detect the presence of chlamydia but none is ideal. Cell culture detects viable organisms but requires special transport and storage of specimens; antigen detection by immunofluorescence is subjective, requiring expertise to interpret the smears, and antigen detection by ELISA, though objective, can give rise to false positive results due to cross-reactions with other micro-organisms unless confirmatory tests are carried out. Polymerase chain reaction (PCR) has been used in research and seems to be both sensitive and specific: commercial kits are now being developed for the amplification and detection of chlamydiae. A PCR kit manufactured by Roche has been applied to stored culture specimens in a diagnostic laboratory.

**Methods**

Two hundred and fifty seven genital specimens (from 157 women and 100 men) were submitted in 2SP transport medium for chlamydial culture and stored at −70°C. Of these, 49 (19-8%) were culture positive (27 from women, 22 from men). Culture had been carried out in McCoy cells in the presence of cycloheximide, but had been incubated for 72 hours before being stained with iodine to demonstrate the presence of glycoprotein containing inclusions.

POLYMERASE CHAIN REACTION

**Roche plasmid amplification for 2SP specimens**

The kits were kindly supplied by Roche Diagnostic Systems. All specimens were tested at 1 in 20 dilution and those which were negative were subsequently tested at 1 in 10 to detect any weakly positive reactions. Ten or 20 µl of specimen were transferred to either 190 or 180 µl of Amplicor specimen transport medium (STM), giving a 1 in 20 or 1 in 10 dilution. After vortexing in capped tubes the suspension was held at room temperature for 5-10 minutes. Amplicor STD specimen diluent (200 µl) was added to this suspension, again vortexed and left 10 minutes at room temperature before use.

The amplification of the DNA was carried out according to the manufacturer’s instructions. Briefly, 50 µl of prepared sample, positive or negative control (in triplicate), was added to 50 µl of Master Mix plus Amplerase (uracil-N-glycosylase) and subjected to 30 cycles of amplification in a Perkin-Elmer Cetus Gene Amplifier PCR System, 9600 thermal cycler. As the Master Mix contained biotinylated primers the resulting product...
Table 1  Laboratory and clinical findings on PCR negative culture patients

<table>
<thead>
<tr>
<th>Case No</th>
<th>Sex</th>
<th>Roche PCR (1 in 20)</th>
<th>Immunofluorescence</th>
<th>Chromosome PCR</th>
<th>Clinical presentation</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>F</td>
<td>1:227</td>
<td>&lt;10</td>
<td>+</td>
<td>Asymptomatic: no evidence of chlamydial infection</td>
</tr>
<tr>
<td>2</td>
<td>F</td>
<td>&gt;2:00</td>
<td>+</td>
<td>ND</td>
<td>Genital warts: no evidence of chlamydial infection +3 weeks later</td>
</tr>
<tr>
<td>3</td>
<td>F</td>
<td>&gt;2:00</td>
<td>&lt;10</td>
<td>+</td>
<td>Genital warts: mucous at cervix Culture +4 months later</td>
</tr>
<tr>
<td>4</td>
<td>F</td>
<td>&gt;2:00</td>
<td>+</td>
<td>ND</td>
<td>Vaginal discharge: no evidence of chlamydial infection</td>
</tr>
<tr>
<td>5</td>
<td>M</td>
<td>&gt;2:00</td>
<td>+</td>
<td>ND</td>
<td>NSG1</td>
</tr>
<tr>
<td>6</td>
<td>M</td>
<td>&gt;2:00</td>
<td>+</td>
<td>+</td>
<td>NSG1</td>
</tr>
<tr>
<td>7</td>
<td>M</td>
<td>&gt;2:00</td>
<td>+</td>
<td>+</td>
<td>NSG1</td>
</tr>
<tr>
<td>8</td>
<td>M</td>
<td>&gt;2:00</td>
<td>unknown</td>
<td>+</td>
<td>Asymptomatic: contact of chlamydia infected female</td>
</tr>
<tr>
<td>9</td>
<td>M</td>
<td>&gt;2:00</td>
<td>+</td>
<td>+</td>
<td>NSG1: ELISA positive 9 months ago Boyfriend not treated until 8 months later</td>
</tr>
<tr>
<td>10</td>
<td>M</td>
<td>&gt;2:00</td>
<td>+</td>
<td>+</td>
<td>NSG1: culture positive 7 months and 12 months ago</td>
</tr>
<tr>
<td>11</td>
<td>F</td>
<td>&gt;2:00</td>
<td>&lt;10</td>
<td>+</td>
<td>Partially treated 7 months ago</td>
</tr>
<tr>
<td>12</td>
<td>F</td>
<td>0:982</td>
<td>+</td>
<td>ND</td>
<td>ELISA positive 11 days ago: not treated at that time</td>
</tr>
<tr>
<td>13</td>
<td>F</td>
<td>&gt;2:00</td>
<td>+</td>
<td>ND</td>
<td>ELISA positive 10 days ago: not treated at that time</td>
</tr>
<tr>
<td>14</td>
<td>F</td>
<td>1:094(1:10)</td>
<td>+</td>
<td>ND</td>
<td>Vaginal discharge No evidence of chlamydial infection</td>
</tr>
</tbody>
</table>

ND not done; NA not available; NSG1 non-specific genital tract infection.

was biotinylated and was denatured by adding 100 µl denaturation solution for 10 minutes at room temperature.

The presence of any biotinylated amplified DNA was detected according to manufacturer's instructions. The labelled product was captured on the probe coated plate by allowing 25 µl of sample or control to interact with the plate for 1 hour at 37°C in a moist box. The plate was carefully washed with Roche wash buffer, 300 µl being added to each well for 30 seconds, on each of 5 occasions. Avidin-horseradish peroxidase conjugate (100 µl) was added to each well for 1 minute at 37°C. After washing as above, 100 µl of tetramethylbenzidine-peroxide substrate was added, the plate incubated at room temperature for 10 minutes in the dark, and the reaction stopped by the addition of 100 µl stop reagent before reading the results at 450 A. An optical density of 0.25 or over was taken as positive provided the positive and negative controls fell within the correct limits.

In-house MOMP amplification

2SP specimen (100 µl) was spun in the wind shielded Biofuge 15000 at 13 000 rpm for 10 minutes. The resulting pellet was extracted, amplified using nested primers, and the amplicon detected by ethidium bromide staining of an agarose electrophoresis gel following the method of Simmonds et al. Briefly, the DNA was phenol-chloroform extracted, the amplification carried out on a Techne PHC-2 machine with a programme of 94°C for 36 seconds, 50°C for 42 seconds, 68°C for 3 minutes for 25 cycles, and 68°C for 5 minutes to complete the strands. The outer nested primers were 420M (sense) 5’ CAG GA c/t ATC TTC TCT GGC TTT AA3 and 423M (antisense) 5’ CGg/a ATG CTg/a ATA GGC TCA CAC AAA GT 3’, and the inner ones which gave a product of 109 base pairs were CT3 (sense) and CT2 (antisense) (Dutilh et al). All the primers were produced by Oswe DNA Service, Department of Chemistry, University of Edinburgh.

IMMUNOFLUORESCENCE

Preparations were prepared by depositing material from 150 µl of 2SP specimen in a wind shielded Biofuge 15000 at 15 000 rpm for 15 minutes. The pellet was resuspended in sterile distilled water and 5–10 µl spread on the well of a Teflon coated slide. Alternatively, cytopsin preparations were prepared from 150 µl of specimen by spinning at 2000 rpm for 10 minutes in a Shandon 3 Cytospin. Both types of preparation were allowed to dry, fixed in methanol for 10 minutes, and stained by Syva Micro Trak reagent according to the manufacturer’s instructions.

Results

Initially the 2SP specimens were spun down and resuspended in specimen diluent. One hundred and eighty were PCR negative and as they were also culture negative the results were accepted and the specimens subsequently discarded. The remainder of the specimens, some of which were discrepant, were retested, as discussed in methods section. Fourteen had an optical density reading of 0.077–0.172 at a 1 in 10 dilution and 0.058–0.211 at a 1 in 20 dilution, and so

Table 2 Comparison of PCR and culture with total resolved specimens

<table>
<thead>
<tr>
<th>Test and result</th>
<th>No of resolved specimens</th>
<th>Sensitivity (%)</th>
<th>Specificity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>PCR</td>
<td>Positive</td>
<td>98.4</td>
<td>99.5</td>
</tr>
<tr>
<td>Negative</td>
<td>1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Culture</td>
<td>Positive</td>
<td>99.0</td>
<td>98.3</td>
</tr>
<tr>
<td>Negative</td>
<td>0</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

J Clin Pathol: first published as 10.1136/jcp.46.9.822 on 1 September 1993. Downloaded from http://jcp.bmj.com/ on April 19, 2022 by guest. Protected by copyright.
were considered PCR negative. These 14 specimens were all culture and immunofluorescence negative and the 11 available were MOMP PCR negative, so they were considered to be resolved negative results. The initial “positive” results were thought to be due to excess magnesium in the reaction mixture because the specimen diluent, which is the source of magnesium, was effectively at double strength (Dr E Dragon, personal communication). Sixty three samples had an optical density of 0.978–2.0 (only six being below 2.0), so were PCR positive, confirming the culture results in 49 instances.

The 14 Roche PCR positive, culture negative specimens were also examined by immunofluorescence, in-house PCR, and the clinical presentation recorded (table 1). In all but specimen 11 the Roche PCR was confirmed by either more than 10 elementary bodies or in-house PCR, or both. There was insufficient specimen 11 to carry out an in-house PCR and less than 10 elementary bodies were present in the immunofluorescence preparation. Compared with specimens 1 and 3, however, it seemed likely that the Roche PCR result was confirmed and specimen 11 has been regarded as resolved. Specimen 14 yielded an optical density of 1.094 in the Roche PCR, but did not have any elementary bodies in the immunofluorescence preparation, and was negative by the in-house PCR test, so it was considered to be an unconfirmed Roche PCR.

The comparison of the PCR and culture with total (n = 257) resolved specimens is shown in table 2 where the sensitivity of PCR is 98.4% and culture 79% and specificity 99.5% and 100%, respectively.

Discussion
As the chlamydia isolation rate is about 8% in the specimens received in this laboratory it was decided to use stored specimens which had been cultured before. In the absence of a method for testing specimens, unwittingly, double the concentration of magnesium was incorporated in the amplification mixture, resulting in 14 “positives”. These results disagree with those of culture, immunofluorescence, in-house MOMP-PCR and clinical observations. When, however, these 14 specimens were retested using the correct reaction mixture, they were all negative, indicating that the method described here must be used for 2SP specimens.

Loeffelholz et al1 reported on the development of the Roche PCR used here. They examined duplicate (2SP and STM) endocervical specimens from 503 women attending a clinic where the incidence of C trachomatis was 17% at the time of study. The culture sensitivity against total resolved specimens was 85.6% compared with 97.0% for PCR. The specificities were 100% and 99.7%, respectively. In this study two fifths of the 257 patients were men and only stored 2SP specimens were available. The culture sensitivity was 79% compared with 98.4% for PCR with specificities of 100 and 99.5%, respectively. These results were very similar and showed that the Roche kit could be applied to both male and female specimens. The level of culture sensitivity is just below the published 80–90%.10 The lower sensitivity of the culture found here might have been influenced by the fact that the specimens were neither blind passaged nor confirmed by immunofluorescence. The results do, however, compare favourably with those of Ossewaarde et al,11 whose 220 patients were predominantly male, and who found that with a different plasmid PCR the culture and PCR sensitivity with reference to confirmed PCR was 72.3% and 100%, with specificities of 98.8% and 98.3%, respectively.

The issue of clinical importance of PCR positivity in the absence of culture has been raised by the previous authors11 and Näher et al12 whose culture sensitivity against unresolved plasmid PCR was 65%. C trachomatis has been isolated from asymptomatic male and female patients,13,14 and in this series eight of the 49 culture positive patients were classified as asymptomatic. Only one of the 14 PCR positive, culture negative patients was asymptomatic, all the others had evidence of sexually transmitted disease or were a contact of a patient with a chlamydial infection. One of the patients with condylomata had a swab taken at follow up three weeks later and this was culture positive so the index specimen was either false culture negative or the infection was in its very early stage.15 Certainly, very few elementary bodies were observed. Five of the patients had a history of chlamydial infection, two in the previous 10–11 days. These patients were treated on day 10 and 11 based on the confirmed ELISA positive result and were culture negative (one also PCR negative) at test of cure. It would therefore seem logical to treat PCR positive, culture negative patients and to undertake contact tracing.

Another reason for treating PCR positive, culture negative patients was thought that the organism was “live” but was incapable of replication, perhaps due to the presence of a cytokine. In such cases test of cure should be done on follow up to ensure that they were PCR negative. This would detect treatment failure in patients with non-replicating chlamydia. Bobo et al17 found that nine of 112 PCR positive, “culture-not-done” patients with trachoma were still PCR positive after four weeks of treatment. These patients are being followed up as they could be non-compliers or they may be refractory to treatment and continue to be a source of chlamydia. Claas et al18 found that the tests of cure of culture positive patients were PCR negative, and so considered that the organism had been eliminated, but Ossewaarde et al19 found that two of their 29 patients still had evidence of
Use of a commercial PCR kit for detecting Chlamydia trachomatis

825

1 Ross JDC, Scott GR. Seasonal variation in gonorrhoea. 
3 Threlfall JE, Ballard RC. The expanding spectrum of the 
4 Chlamydia—a microbiological and clinical appraisal. 
6 Ridgway GL, Taylor-Robinson D. Current problems in 
7 microbiology: I Chlamydial infections: which laboratory 
9 Taylor-Robinson D, Thomas BJ. Laboratory techniques for 
11 Thomas BJ, Evans RT, Hutchinson GR, Taylor-Robinson 
12 D. Early detection of Chlamydia infections combining 
13 the use of cycloheximide treated McCoy cells and 
14 immunofluorescence staining. J Clin Microbiol 1977;6: 
16 Evans RT, Woodland RM. Detection of chlamydiae by 
19 Leefholtz MJ, Lewinski CA, Silver SR, et al. Detection of 
20 Chlamydia trachomatis in endocervical specimens by 
21 polymerase chain reaction. J Clin Microbiol 1992;30: 
22 2447-51.
23 Simmonds P, Balfé P, Peutherer JF, Ludlam CA, Bishop 
24 JO, Leigh Brown AJ. Human immunodeficiency virus- 
25 infected individuals contain provirus in small numbers of 
26 peripheral mononuclear cells and at low copy numbers. 
28 Dutilh B, Bébéar C, Rodriguez P, Vevers A, Bennet J, 
29 Garnet M. Specific amplification of a DNA sequence 
30 common to all Chlamydia trachomatis serovars using 
32 Mahony JB, Luijstra KE, Sellors JW, Jang D, Chrensky 
33 MA. Confirmatory polymerase chain reaction testing for 
34 Chlamydia trachomatis in first-void urine from asymptom- 
36 2241-5.
37 Osewaarde JM, Rieffe M, Rozenberg-Arks M, Ossenkoppeme PM, Nawrocki RP, van Loon AM. 
38 Development and clinical evaluation of a polymerase 
39 chain reaction test for detection of Chlamydia trachomatis. 
41 Näher H, Drzeznik H, Wolf J, von Knebel Doeberitz M, 
42 Perzoldt D. Detection of C trachomatis in urogenital 
43 specimens by polymerase chain reaction. Genitourin Med 
45 Hunter JM, Smith JW, Peutherer JF, MacAslauy A, Tuach 
46 S, Young H. Chlamydia trachomatis and Ureaplasma 
47 urealyticum in men attending a sexually transmitted dis- 
49 Hunter JM, Smith JW, Peutherer JF, MacAslauy AJ. 
50 Chlamydia trachomatis infection of the cervix: the need 
52 Ostergaard L, Truelson J, Birkeland S, Christiansen G. 
53 Evaluation of ureogestant Chlamydia trachomatis infec- 
54 tions by cell culture and the polymerase chain reaction 
55 using a closed system. Eur J Clin Microbiol Infect Dis 
57 Holland SM, Hudson AP, Bobo L, et al. Demonstration 
58 of chlamydial RNA and DNA during a culture-negative 
60 Bobo L, Munos B, Viscidi R, Quinn T, Mbooa H, West 
61 S. Diagnosis of Chlamydia trachomatis eye infection in 
63 Claas HJC, Wagenvoor JHT, Niesters HGM, Tio TT, 
64 van Rijnvort-Vor J, Quins WGV. Diagnostic value of the 
65 polymerase chain reaction for Chlamydia detection as 
67 Osewaarde JM, Planteuma FHF, Rieffe M, Nawrocki RP, 
68 de Vries A, van Loon AM. Efficacy of single-dose 
69 azithromycin versus doxycycline in the treatment of cer- 
70 vical infections caused by Chlamydia trachomatis. Eur J 
73 methods for diagnosing chlamydial infection in patients 
74 with trachoma: a clue to the pathogenesis of the disease? 
76 Campbell LA, Patton DL, Moore DE, Cappuccio AL, 
77 Mueller DA, Wang S. Detection of Chlamydia tracho-
78 miti deoxyribonucleic acid in women with tubal infertility. 