Measurement of IgG antibodies to *Chlamydia trachomatis* by commercial enzyme immunoassays and immunofluorescence in sera from pregnant women and patients with infertility, pelvic inflammatory disease, ectopic pregnancy, and laboratory diagnosed *Chlamydia psittaci/Chlamydia pneumoniae* infection

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**Technical Report**

Background: Screening for *Chlamydia trachomatis* specific antibodies is valuable in diagnosing asymptomatic pelvic inflammatory disease (PID) and tubal damage following repeated episodes of PID. The assays in current use are unsuitable for screening large numbers of samples so there is a need to develop more suitable assays.

Aims: To compare the performance of several commercial *C trachomatis* enzyme immunoassays (EIAs) (SeroCT, C *trachomatis* IgG EIAs using major outer membrane protein (MOMP), an inactivated organism EIA (Genzyme Virotech EIA), and a genus specific EIA (Platelia Chlamydia IgG) with the whole cell inclusion immunofluorescence (WIF) assay. In addition, to adapt, using time resolved fluorescence technology, the assay showing the highest correlation with WIF.

Methods: Ninety sera from patients presenting with ectopic pregnancies, 187 sera from those with a variety of types of infertility, 33 sera from cases of PID where a fourfold rise in WIF titre occurred, and 90 sera from antenatal clinic attenders were tested. A panel of 36 sera from laboratory diagnosed cases of *Chlamydia psittaci/Chlamydia pneumoniae* infection was also tested.

Results: The Genzyme Virotech EIA showed the highest rank correlation coefficient (0.82) with WIF, particularly at high WIF titres. The MOMP specific assays varied in their correlation with WIF, with rank correlation coefficients ranging from 0.70 (Medac p-EIA) to 0.80 (Vircell EIA). The Genzyme Virotech assay showed poor specificity (5.6%; 95% confidence interval [CI], 0.68% to 18.7%)—it was reactive with 34 of the 36 C *psittaci/ C pneumoniae* positive sera. The MOMP based EIAs showed high specificity, particularly the Medac p-EUSA (97.2%; 95% CI, 85.5% to 99.9%)—only one serum was reactive. In view of the good correlation between WIF and the Genzyme Virotech EIA, a time resolved fluorescence immunooassay (TRFIA) was developed using the Genzyme Virotech antigen. Using an appropriate cut off the TRFIA assay showed excellent correlation with WIF.

Conclusions: The TRFIA assay may be useful as a screening assay, possibly in conjunction with one of the highly specific EIAs studied (for example, Medac p-EIA) to confirm the antibody specificity of sera selected by the screening assay.

*Chlamydia trachomatis* infection is the most common sexually transmitted bacterial disease in England, Wales, and Northern Ireland, with 64 000 diagnoses made in the year 2000. Most *C trachomatis* lower genital tract infections are asymptomatic and the most common clinical presentation in women is mucopurulent cervicitis, and in men urethritis. For lower genital tract infection, the detection of specific antibodies in a single serum specimen is held to be of little value because such antibodies are frequently found in sera from women who do not have active infection. Despite the difficulty of differentiating between previous and current lower genital tract infection, there is a considerable amount of evidence that the presence of *C trachomatis* specific antibody is significantly associated with upper genital tract infection, particularly when the antibody is at a high titre. Screening for *C trachomatis* specific antibodies is valuable in diagnosing asymptomatic pelvic inflammatory disease (PID) and tubal damage following repeated episodes of PID, particularly because it has been shown that *C trachomatis* is rarely isolated from the upper genital tract and clinical diagnosis requires invasive procedures not routinely available in general practice.

There are two accepted reference assays for measuring *C trachomatis* specific antibodies, the microimmunofluorescence assay (MIF) of Wang and colleagues and the whole cell inclusion immunofluorescence assay (WIF) of Richmond and Caul. The WIF assay is a single antigen immunofluorescence test in which cytochalasin B treated McCoy cells infected with an LGV type 2 strain of *C trachomatis* are placed in wells on slides coated with polytetrafluoroethylene. In this system, the whole chlamydial inclusion acts as the antigen, in contrast to the MIF test in which elementary bodies act as the antigen. The WIF test detects both genus specific (lipopolysaccharide; LPS) antibody and species specific major outer membrane protein (MOMP) antibody and, like MIF, it is a subjective, labour intensive assay not suited to screening large numbers of sera. Our laboratory uses the WIF assay because we have found it to be more reliable for the diagnosis of upper genital tract infection than MIF, and also because inclusions are easier to visualise than cell free elementary bodies.

**Abbreviations:** CI, confidence interval; EIA, enzyme immunoassay; LPS, lipopolysaccharide; MIF, microimmunofluorescence assay; MOMP, major outer membrane protein; PID, pelvic inflammatory disease; TRFIA, time resolved fluorescence immunoassay; WIF, whole cell inclusion immunofluorescence
Time resolved fluorescence immunoassay is suitable for the measurement of low and high amounts of antibody, even with single dilutions of specimen, because of its large linear dynamic range.

The effectiveness of intervention measures to reduce *C. trachomatis* infection in designated female populations can be measured through determining population prevalences of chlamydia mediated upper genital tract infection. For screening activities, in which large numbers of sera need to be tested, we need alternative assays to MIF and WIF and commercial *C. trachomatis* antibody assays may prove to be a useful alternative. Furthermore, for population screening the utilisation of non-invasive sampling techniques, such as collection of oral fluid, is our desired goal, and this approach requires the adoption of specialised technology with potential application to oral fluid screening. We favour the use of time resolved fluorescence immunoassay (TRFIA), because in our hands this technology has been found to be highly sensitive. TRFIA is suitable for the measurement of low and high amounts of antibody, even with single dilutions of specimen, because of its large linear dynamic range, and has previously been assessed for population screening of anti-*C. pneumoniae* IgG, where its objectivity, reproducibility, and amenability to automation were distinct advantages compared with MIF.

Our study had two objectives: (1) to compare the performance of several commercial *C. trachomatis* antibody assays in relation to WIF, with a view to assessing their specificity and sensitivity for *C. trachomatis*; and (2) to adapt, using time resolved fluorescence technology, the commercial assay showing highest correlation with WIF.

**MATERIALS AND METHODS**

**Sera tested**

A set of 310 sera was assembled, comprising 90 sera from ectopic pregnancies, 187 sera from patients with infertility, and 33 sera from cases of PID shown by at least a fourfold increase in titre. In addition, a set of 36 sera from WIF diagnosed cases of *C. psittaci* or *C. pneumoniae*, as shown by rising complement fixation test/WIF titre, and a control set of 90 antenatal sera were also assembled to produce a test panel of 436 sera.

**Figure 1** Correlation between commercial enzyme immunoassay (EIA) results (log10 scale) and whole cell inclusion immunofluorescence (WIF) results. (A) Medac pEIA v WIF; (B) Labsystems v WIF; (C) Patelia Chlamydia IgG v WIF; (D) C trachopp v WIF; (E) SeroCT v WIF; (F) Vircell v WIF; (G) Genzyme Virotech EIA v WIF.
Whole cell inclusion immunofluorescence assay (WIF)

This assay was performed as described previously. WIF testing was performed before the other tests to avoid potential bias in the results. *Chlamydia trachomatis* L2 was used for *C. trachomatis* specific antibody detection, enzootic abortion ewes *C. psittaci* was used for *C. psittaci* specific antibody determination, and *C. pneumoniae* TW183 was used for *C. pneumoniae* specific antibody determination.

**EIA kits tested**

The following EIA kits were tested:

- **Platelia Chlamydia IgG (Sanoﬁ Pasteur)**, catalogue number 62766, batch number 9K058U. Supplied by Sanoﬁ Diagnostics Pasteur Ltd, Guildford, UK.
- **SeroCT™-IgG** (Savyon Diagnostics Ltd), catalogue number A181–01M, batch number 181–912A. Supplied by Brownes Ltd, Reading, UK.
- **C trachomatis** IgG/EIA (PBS Organics), catalogue number HX.CTG.096, batch number 990905. Supplied by Quest Biomedical, Knowle, UK.
- **Chlamydia trachomatis** IgG/IgM EIA (Vircell SL), catalogue number G/M1017, batch number 99ECTR104. Supplied by Microgen Bioproducts Ltd, Camberley, UK.
- **Chlamydia trachomatis-IgG-pEIA** (Medac Diagnostica), catalogue number 497/TMB, batch number CTG08. Supplied by Medac Diagnostica, Wedel, Germany.
- **Chlamydia trachomatis** EIA (Genzyme Virotech), catalogue number EC 120.00, batch number 90817–01. Supplied by Diasorin, Wokingham, UK and Genzyme Virotech, Russelsheim, Germany.
- **Chlamydia trachomatis** IgG EIA (Labsystems), catalogue number 6111–101, batch number 112TK2–3. Supplied by Quest Biomedical.

**EIA methods**

All kits used were of the same batch and testing was performed before their expiry dates. The manufacturers instructions were followed, in full, when performing the assays and only IgG values were determined. In each instance, kits were stored at 4°C and allowed to stand at room temperature for one hour before use. Incubations at 37°C were performed in moist chambers unless specifically indicated otherwise. Microtitre plate washing was performed using a Labsystems (Thermo Life Sciences Ltd, Basingstoke, UK) Wellwash 4 Mk 2 and plates were read using a Labsystems Multiskan RC reader. Assays were only regarded as valid if all manufacturers’ validation criteria were satisfied.

**Time resolved fluorometric immunoassay (TRFIA)**

Sera for testing were diluted 1/64 in DELFIA assay buffer (Wallac Oy, Turku, Finland) and 100 µl was loaded on to Genzyme Virotech *C. trachomatis* coated microwell plates (Genzyme Virotech). The plates were incubated for two hours at 37°C in a moist chamber and then washed four times with DELFIA wash solution (Wallac Oy) using a DELFIA 1296–026 Platewash (Wallac Oy). Europium labelled antihuman IgG conjugate (Wallac Oy), diluted 1/500 in DELFIA assay buffer, was added (100 µl/well) to the plates, which were then incubated for one hour, at 37°C, in a moist chamber. The plates were washed four times, DELFIA enhancement solution (Wallac Oy) was added (100 µl/well), and the plates were then shaken incubated for 10 minutes at room temperature using a Stuart mini-orbital shaker (Bibby Sterilin, Stone, UK) set at 125 rpm. The plates were then read using a DELFIA 1234 fluorometer (Wallac Oy) and the counts processed by multicalc software, version 2.5 (Wallac Oy).

**Statistical methods**

Assays were compared on a log scale using Spearman’s coefficient of rank correlation (r) with 95% confidence intervals (CI) calculated by bootstrapping. Best fit regression lines were also plotted. For the 36 sera diagnosed by WIF as *C. pneumoniae* or *C. psittaci*, the specificity of the assays was calculated with 95% confidence intervals.

**RESULTS**

Figure 1 shows the correlation between commercial EIA results (log₁₀ scale) and WIF results. From the graphs, a clustering effect of EIA values could be seen with WIF low and high titre sera to varying degrees, depending on the EIA kit used. A substantial number of sera had low titre WIF and high titre EIA results (false positives) and high titre WIF with low titre EIA results (false negatives). Overall, the Genzyme Virotech assay, which had the highest rank correlation coefficient of 0.82, correlated most closely with WIF, particularly with WIF high titre sera. The Vircell (species specific) and Sanoﬁ Pasteur (genus specific) assays had the next highest rank correlation (0.80), and again showed good clustering with WIF high titre sera. The lowest rank correlation coefficient (0.70) was obtained with the Medac p-EIA.

Using a panel of WIF diagnosed *C. psittaci* and *C. pneumoniae* sera, the *C. trachomatis* antibody specificity of the commercial assays was evaluated (table 1). The Medac p-EIA had the highest specificity when compared with WIF (97.2%; 95% CI, 85.5% to 99.9%), with only one serum testing *C. trachomatis* antibody positive. Both Labsystems and PBS Organics assays had specificities of 94.4% (95% CI, 81.3% to 99.3%), followed by the Savyon Sero CT assay with a specificity of 91.7% (95% CI, 77.5% to 98.3%). Despite having the highest rank correlation coefficients with WIF, the Genzyme Virotech and Vircell assays showed the lowest specificities—5.6% and 58.3%, respectively—of the species specific assays tested. The Sanofi Pasteur EIA had a specificity of 0%, which reflects the fact that it is a genus specific assay.

**Table 1 Specificity of the commercial enzyme immunoassays (EIAs) for detecting Chlamydia trachomatis antibody when tested against a panel of Chlamydia psittaci/Chlamydia pneumoniae antibody positive sera as diagnosed by whole cell immunofluorescence assay**

<table>
<thead>
<tr>
<th>Commercial EIA</th>
<th>Number negative out of 36</th>
<th>Specificity (95% CI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Medac Diagnostica</td>
<td>35</td>
<td>97.2% (85.5% to 99.9%)</td>
</tr>
<tr>
<td>Labsystems</td>
<td>34</td>
<td>94.4% (81.3% to 99.3%)</td>
</tr>
<tr>
<td>PBS Organics (Tracho PEP)</td>
<td>34</td>
<td>94.4% (81.3% to 99.3%)</td>
</tr>
<tr>
<td>Savyon Diagnostics (SeroCT)</td>
<td>33</td>
<td>91.7% (77.5% to 98.3%)</td>
</tr>
<tr>
<td>Vircell SL</td>
<td>2</td>
<td>58.3% (40.8% to 74.5%)</td>
</tr>
<tr>
<td>Genzyme Virotech</td>
<td>2</td>
<td>5.6% (0.68% to 18.7%)</td>
</tr>
<tr>
<td>Sanoﬁ Pasteur (Platelia)</td>
<td>0</td>
<td>0% (0% to 9.7%)</td>
</tr>
</tbody>
</table>

CI, conﬁdence interval.
Figure 2 Modified Genzyme Virotech assay showing the excellent discrimination between the 45 sera with whole cell inclusion immunofluorescence (WIF) titres the same, or higher, than the cut off (≥ 512) and 22 sera with titres below the cut off (< 512).

For application to screening activities we considered the Genzyme Virotech assay most suitable for further development because it showed a good correlation with WIF, particularly for sera having WIF titres of 512, or higher, which is the cut off titre used in our laboratory for predicting upper genital tract infection. A limitation of the Genzyme Virotech assay was that the cut off titres stipulated by the manufacturer resulted in many sera that had low WIF titres (that is, ≤ 64) being classified as C trachomatis antibody positive. Antibody to C trachomatis could have been present at low levels; however, for our purposes we needed to recalibrate the Genzyme Virotech assay so that the cut off used correlated more closely with the WIF cut off titre of 512. Such a recalibration was achieved by using time resolved fluorescence technology. Forty five sera with WIF titres the same, or higher, than our chosen cut off (that is, WIF titres ≥ 512) and 22 sera with titres below the cut off (that is, WIF titres < 512) were tested (fig 2). A cut off point of 100 000 counts with the modified Genzyme Virotech assay was chosen to differentiate between high WIF titre and low WIF titre sera because it gave excellent discrimination.

DISCUSSION
A wide range of antibodies has been reported to be produced following C trachomatis infection because the organism has a unique biphasic life cycle, alternating between infectious elementary bodies and a replicating, reticulate body. There are three species of chlamydia that cause infection in humans, namely: C trachomatis, C pneumoniae, and C psittaci and several serovars of C trachomatis (D–K) that infect the urogenital tract. Any serological test that is used to detect C trachomatis infection must differentiate between infection by other species of chlamydia, yet retain the capacity to detect antibodies produced at different stages of the life cycle and by different serovars. One advantage of the microimmunofluorescence and whole cell inclusion immunofluorescence assays is that because they use whole organisms/native antigens they detect a wide variety of antibodies and therefore confer enhanced sensitivity. A potential limitation of these assays is that they may detect antibodies unrelated to C trachomatis infection, thereby compromising specificity. For example, crossreaction between the genus specific LPS antigen of Chlamydia spp and the lipopolysaccharides of Porphyromonas gingivalis, Escherichia coli O19, and Salmonella newington has been reported for both immunofluorescence and EIAs utilising chlamydial LPS.

“the specificity of the Medac p-EIA may be attributed to its use of a highly immunogenic species specific epitope, which shares no sequence homology with Chlamydia pneumoniae”

MOMP has been used extensively in EIA based tests for C trachomatis because this antigen is considered to be species and serovar specific, and the Savoy SeroCT, PBS Orgenics C trach+ , Labsystems C trachomatis, and Medac C trachomatis p-EIA assays use peptides mimicking regions of this protein. From fig 1 it can be seen that the species specific peptide assays performed variably in relation to WIF, with rank correlations ranging from 0.70 to 0.76. The Vircell assay, which uses extracted and purified MOMP had a higher rank correlation with WIF (0.80) than the synthetic MOMP peptide assays, which may reflect the fact that the antigen used is native, and therefore possesses more epitopes. The assay that had the best correlation with WIF, particularly at high WIF titres, was the Genzyme Virotech C trachomatis EIA, which uses C trachomatis LGV type II strain, cultured in mouse L cells, and inactivated using γ irradiation. It is possible that the reduced correlation of the synthetic peptide assays results from conformational differences in the epitopes presented by the peptides compared with native antigens.

The Medac p-EIA showed the highest specificity for C trachomatis specific antibody (97.2%) with WIF when tested against sera from WIF diagnosed C psittaci and C pneumoniae infection. The specificity of the Medac p-EIA may be attributed to its use of a highly immunogenic species specific epitope, which shares no sequence homology with C pneumoniae. The Labsystems C trachomatis EIA was the next most specific assay, and this test is based on four synthetic peptides derived from the variable domain IV of the MOMP of C trachomatis serotypes C, G, E, and L2. The Genzyme Virotech EIA, although showing the best correlation between high WIF titre and low WIF titre sera because it gave excellent discrimination.

The absence of complement fixing antibody or its presence at low titres in sera with WIF titre ≥ 512 is suggestive of tubal damage with no evidence of active infection; however, if complement fixing antibody is found at titres ≥ 12, PID is suspected. The Genzyme Virotech EIA was selected as our screening assay; however, we needed to refine the assay so that the cut off used correlated with that used for WIF (≥ 512) in our laboratory. The assay was modified using time resolved fluorescence technology because this technique, as a result of its large linear dynamic range, has particular application to screening single serum dilutions. This approach appeared to be highly successful (fig 2) in differentiating between WIF positive sera (titres ≥ 512) and WIF negative sera (titres < 512). Using the modified, time resolved Genzyme Virotech assay, high chlamydial antibody titre sera can be selected from our screening programme for further testing for C trachomatis specific MOMP antibody. Our studies of specificity have shown that the Medac p-EIA is the most appropriate assay to use; however, there are occasions when sera have high screening antibody titres but are not p-EIA positive. Such incidents may be the result of genuine C psittaci/C pneumoniae infection, crossreactivity with LPS from Gram negative bacteria, or a failure of C trachomatis antibodies in the serum to recognise the MOMP epitopes in the Medac p-EIA.

Our ultimate goal is to develop a non-invasive screening assay for chlamydial infection which, with appropriate refinement, will be specific for C trachomatis mediated upper genital tract disease. The amount of IgG in whole saliva is about one thousandth of that in plasma, and using the TRFIA cut off
count of 100 000 (when testing sera at a 1/64 dilution), a theoretical cut off would be 100 counts. We normally test saliva at a 1/4 dilution; therefore, the theoretical cut off would be 1600 counts, which is measured with ease by TRFIA.

“Our ultimate goal is to develop a non-invasive screening assay for chlamydial infection which, with appropriate refinements, will be specific for Chlamydia trachomatis mediated upper genital tract disease”

To conclude, we have evaluated several commercial C. trachomatis specific EIAs against WIF and the correlation has ranged between 0.70 and 0.80. The best correlation (0.82) was seen with an EIA using a cultured, inactivated organism and this assay has been modified, using time resolved fluorescence technology, for use as a screening assay. Some of the EIAs studied (for example, Medac p-EIA) were highly specific for C. trachomatis antibody and can be used to confirm the antibody specificity of sera selected by the screening assay. Further studies have been initiated to assess the usefulness of this screening approach.

Take home messages

- Screening for Chlamydia trachomatis specific antibodies is valuable in diagnosing asymptomatic pelvic inflammatory disease (PID) and tubal damage following repeated episodes of PID
- Because the currently used assays are unsuitable for screening activities in which large numbers of sera need to be tested, we evaluated several commercial C. trachomatis specific enzyme immunoassays (EIAs) against the most commonly used assay (WIF)
- The best correlation (0.82) was seen with an EIA using a cultured, inactivated organism and this assay has been modified, using time resolved fluorescence technology, for use as a screening assay
- Some of the EIAs studied (for example, Medac p-EIA) were highly specific for C. trachomatis antibody and can be used to confirm the antibody specificity of sera selected by the screening assay
- Further studies have been initiated to assess the usefulness of this screening approach

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