Use of culture and immunochromatographic technique for diagnosis of trichomoniasis in Sri Lanka

H Banneheke, R Fernandopulle, S Prathapan, G de Silva, N Fernando, R Wickremasinghe

(Index words: trichomoniasis, Sri Lanka, laboratory diagnosis)

Summary
As a majority of the trichomoniasis patients are asymptomatic, laboratory tests are crucial in case detection. The usefulness of culture and immunochromatographic technique (ICT) compared to microscopy for detection of trichomoniasis in Sri Lanka was assessed. Females (16-45 years) from Colombo district were screened for *Trichomonas vaginalis* using three laboratory tests namely, microscopy of wet smear, Trichomonas liquid culture and ICT (OSOM® trichomonas rapid test). Trichomoniasis by at least one test being positive was 4.8%. Microscopy, culture and ICT detected 2.8%, 4.2% and 10% cases respectively. Microscopy missed 32% of culture positives. ICT is a simple, practical and reliable alternative to microscopy in laboratory diagnosis of trichomoniasis.

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Introduction
Trichomoniasis caused by *Trichomonas vaginalis* is a sexually transmitted infection, accounting for an annual incidence of 174 million cases with 44% (76.5 million) occurring in South and South East Asia [1]. The prevalence of *T. vaginalis* in clinic based populations in Sri Lanka has varied from 4.4%-7.2% [2,3]. The prevalence of asymptomatic infection may vary from 50%-85% in females, thus screening is vital for case detection [4]. Wet mount microscopy with sensitivity of 52%-85% in clinic based populations is used commonly in routine practice worldwide [5,6]. Other investigations include microscopy of stained preparations, culture, polymerase chain reaction (PCR) and immunochromatographic technique (ICT) for *Trichomonas* antigens.

No studies have been done on the use of ICT for the diagnosis of trichomoniasis in Sri Lanka. The objective of this study was to evaluate the usefulness of culture and ICT compared to wet mount in the Sri Lankan setting.

Methods
A total of 601 females aged 16-45 years from the Colombo district were screened for *T. vaginalis*, irrespective of the presence or absence of symptoms. Study participants were recruited over a period of 18 months in 2007-2009. Those who refused vaginal examination, were on antibiotics or having any form of vaginal bleeding were excluded. Using speculum, three vaginal swabs from the posterior fornix were obtained. *Trichomonas* liquid culture “Trichomonas modified CPLM medium base” (“HIMEDIA®” Ref M 460) was inoculated at the bed side and incubated at 37°C. They were examined microscopically daily till the 5th day for motile trichmonads. ICT OSOM® trichomonas rapid test (genzyme Diagnostics, Kent, ME194AF) was performed according the manufacturer’s instructions for only 100 (17%) participants due to financial constraints. Microscopic examination of the wet smear was done at the bed side under ×10 and × 40 objectives for motile trichomonads. The study was approved by the Ethical Review Committee of Faculty of Medical Sciences, University of Sri Jayewardenepura.

Results
The mean age of the participants was 31.7 years. The confirmed cases of trichomoniasis by at least one positive laboratory test were 4.8% (29/601) with 2.8% (17/601) by wet mount microscopy, 4.2% (25/601) by culture and 10% (10/100) by ICT. Wet mount was unable to detect 32% (8/25) of culture positive infections (Table 1). Out of the 10 ICT positives, four were culture negative (Table 2). The sensitivity and the specificity of vaginal smear were 68% and 100% and the positive and negative predictive values were 100% and 98% respectively. The ICT had 100% sensitivity, 96% specificity, 60% positive and 100% negative predictive values. In all these instances culture was taken as the gold standard.

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ICT showed a higher detection rate than microscopy. Infection would have been missed in 32% if only microscopy was performed. The sensitivity and specificity of microscopy in our study is comparable with other studies [5-8]. The manufacturer states 83% sensitivity and 99% specificity of ICT compared to 'culture and PCR combination' and 96% sensitivity and 95% specificity compared to wet mount. ICT showing a higher sensitivity than the culture in this study can be attributed to false positivity. ICT is based on the detection of *T. vaginalis* antigens rather than the live whole parasite, therefore such false positivity results are likely to occur [9]. ICT, having high levels of sensitivity and specificity, has been recommended by others as a good screening test especially for low prevalence settings like in Sri Lanka [9-10].

Microscopy has retained its place as the sole diagnostic test in Sri Lanka and many other countries, because it is easy to perform, inexpensive and available at the point of care. When compared with laborious culture techniques in microbiology laboratories, the trichomonas liquid culture is simple and easy to prepare, store and process in the laboratory. However, ICT seems an easy test which can be done at the point of care when compared with the less sensitive microscopy or culture that requires 72 to 120 hours to obtain a result. This study shows that wet mount should not be used as the sole diagnostic technique due to low sensitivity. Instead, a second test should be added to the routine microscopy. The decision regarding the selection of the second test can be taken by the individual laboratory considering all aspects. In conclusion, we recommend ICT as a simple, practical and reliable alternative to wet mount. We also recommend culturing to be established along with the microscopy at least in tertiary laboratories.

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### References


### Table 1. Comparison of vaginal smear with liquid culture of participants

<table>
<thead>
<tr>
<th>Culture positives (n=25)</th>
<th>Culture negative (n=576)</th>
<th>Total (n=601)</th>
</tr>
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<tbody>
<tr>
<td>Wet smear</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive</td>
<td>17</td>
<td>0</td>
</tr>
<tr>
<td>Negative</td>
<td>8</td>
<td>576</td>
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</tbody>
</table>

### Table 2. Comparison of immunochromatographic technique (ICT) with liquid culture of participants

<table>
<thead>
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<th>Culture positives (n=6)</th>
<th>Culture negative (n=94)</th>
<th>Total (n=100)</th>
</tr>
</thead>
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<tr>
<td>ICT</td>
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<tr>
<td>Positive</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td>Negative</td>
<td>0</td>
<td>90</td>
</tr>
</tbody>
</table>

### Discussion

ICT showed a higher detection rate than microscopy. Infection would have been missed in 32% if only microscopy was performed. The sensitivity and specificity of microscopy in our study is comparable with other studies [5-8]. The manufacturer states 83% sensitivity and 99% specificity of ICT compared to ‘culture and PCR combination’ and 96% sensitivity and 95% specificity compared to wet mount. ICT showing a higher sensitivity than the culture in this study can be attributed to false positivity. ICT is based on the detection of *T. vaginalis* antigens rather than the live whole parasite, therefore such false positivity results are likely to occur [9]. ICT, having high levels of sensitivity and specificity, has been recommended by others as a good screening test especially for low prevalence settings like in Sri Lanka [9-10].