INTRODUCTION

Trichomonas vaginalis is the most common non-viral sexually transmitted diseases (STDs) agent on a world scale. One hundred and eighty millions new infections are estimated every year (20). According to the population studied, the prevalence of infection in females varies from 5 to 50% in patients attending sexually transmitted disease (STD) clinics (16). The incidence is rapidly growing, particularly in Sub-Saharan Africa, South-East Asia, and East Europe. The incidence is highest among women with multiple sexual partners and in groups with high rates of other STDs (7). This agent causes trichomoniasis and has been linked to preterm birth, acquisition of human immunodeficiency virus, infertility, and nongonococcal urethritis (3, 19). Patients harboring the protozoon are at risk of, and should be screened for, other pathogens, such as Neisseria gonorrhoeae, Chlamydia trachomatis, and HIV. These can be more worrisome, even if clinically silent at the time of clinical examination. In clinical practice the direct microscopic examination of vaginal secretions using the wet mount technique is the most commonly used approach. To identify the protozoon it is necessary to recur to an expert.

FULL PAPERS

Real-Time Polymerase Chain Reaction Detection of Trichomonas vaginalis from vaginal swabs: Validation of a Diagnostic Method and Preliminary Epidemiological Application

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Key Words: Trichomonas vaginalis, real-time PCR, vaginal swab

SUMMARY

Background

Trichomonas vaginalis is the most common non-viral sexually transmitted diseases (STDs) agent. For females, the diagnostic gold standard is the culture of vaginal swab, which is labor-exacting. The direct microscopic examination of vaginal secretions is the most used approach, but its sensitivity depends on the skill of the observer.

Objectives

We evaluated an original real-time TaqMan-based Polymerase Chain Reaction (PCR) technique. The scope of the study was to confirm the effectiveness of the molecular approach in a clinical context and to explore its relevance to an epidemiological investigation.

Study Design

a-ß-tubulin gene was chosen as target sequence. The assay was designed to exploit the quantitative potential of the TaqMan procedure. The population sample was 583 adult females presenting at the Service from January 2005 to December 2005. Three vaginal swabs were collected from each patient, one for wet mount microscopy, one for broth culture, and one for the molecular assay.

Results

The prevalence was 3.3% (culture), 3.1% (microscopy), 3.8% (PCR). An excess risk was detected in the immigrant population (risk ratio by PCR = 28).

Conclusions

The molecular approach was the most accurate way to detect the protozoon. The real-time PCR is convenient in a busy laboratory, provided the necessary equipment is available, and it is suitable for epidemiological investigation.

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microscopist. As with all microscopic diagnoses, the accuracy of the direct examination is highly dependent on the skill and diligence of the observer. The sensitivity of the direct examination depends strictly on timeliness of the microscopic reading which must be done immediately after the swab collection. Indeed *T. vaginalis* quickly loses its motility and cannot be distinguished from white blood cells. Culture is still regarded as the “gold standard” for determining the performance of other diagnostic tests, but it is not clear which medium formulation and culture system is best for *T. vaginalis*. Moreover, all liquid culture systems for *T. vaginalis* require repeated media sampling and examination up for seven days, making the procedure unpractical for population screening programs (14).

A DNA 2.3 kb probe was cloned by Rubino et al. (18) for detection of *T. vaginalis* by a dot blot hybridization technique. A molecular PCR-based approach was developed by Riley and coworkers in 1992 (17). Since then, several *T. vaginalis*-specific assays have been described. Among the PCR targets are the ferredoxin gene (17,8), highly repeated DNA sequences (11), the family of adhesion proteins (1), β-tubulin gene (13), and 18S ribosomal gene (15). A real-time TaqMan-based detection of ferredoxin gene was designed and evaluated in 2001, using a previously described set of primers in conjunction with a novel, double-labeled fluorescent probe (9). Later, an existing PCR method for β-tubulin gene was adapted into a real-time PCR assay based on fluorescence resonance energy transfer (FRET) probe (6).

Epidemiological data of this curable disease are scarce for Italy. The prevalence of *Trichomonas vaginalis* infection was 4.2% in 15,213 infections of the cervico-vaginal tract reported in Italy in the period September 1990-December 1999 (4) and it was 5.0% in 861 Italian female teenagers affected by STDs in the period September 1990-December 1997 (5).

In order to contribute to this promising line of evolution in diagnostic parasitology, and hoping to ease an epidemiologic investigation in our area, we developed an original real-time PCR TaqMan-based technique. Since in the last few years there have been several reports concerning nucleic acid amplification-based approach, but this has never reached a wide acceptation, the aim of the present study was to investigate the effectiveness of the approach to the diagnosis of *T. vaginalis* infection by means of a molecular method. Compared to the traditional techniques, it was reasonable to expect an increase in the diagnostic accuracy by using a PCR procedure. Moreover, a step was done toward the exploitation of the potential to furnish quantitative information residing in real-time PCR. Indeed, in several instances concerning the infectious diseases the quantitative assay of the infecting agent is often predictive of the clinical context. In order to achieve this target, a calibration curve was set up. Moreover, to assess the adequacy of the pathologic sample, an endogenous DNA sequence (part of the β-globin gene) (2) was amplified within the same reaction tube for each sample, as an inner control.

**MATERIALS AND METHODS**

**Patient population, sample collection, and diagnostic tests**

The population sample under study was formed by 583 consecutive adult female patients presenting at the Microbiology and Virology Service of Padua’s Hospital, during the period January 2005-December 2005. All had symptoms or personal history pointing to a possible genital infection. Three vaginal swabs were collected from each patient under speculum control. One swab was employed for wet mount microscopy, one for broth culture, and one for molecular approach. *Trichomonas vaginalis* was identified by means of a real-time PCR TaqMan. As a gold standard, the protozoan search was done by broth culture. Moreover, a microscopic wet mount search was performed.

**Broth culture**

The composition of the medium, *Trichomonas* Selective Broth (Labobasi srl, Milano, Italy) was peptone, liver extract, yeast extract, maltose, agar, chloramphenicol, methylene blue, horse serum. The culture was done by inoculating the swab into the broth and by incubating it at 37 °C up to five days. The culture fluid was microscopically examined (magnification 40x) daily. Identification of *T. vaginalis* was achieved by observation of motile organisms in wet-mount preparations.

**Wet mount microscopy**

By one hour from the collection of the samples, one vaginal swab for each patient was inoculated into 1 ml of physiologic solution. Then, one drop of this solution was examined by microscopy (magnification 40X) in order to detect *T. vaginalis* by this characteristic shape and motility.

**DNA processing and TaqMan real-time PCR method**

After collection, the swab for molecular approach was eluted in 1 ml of physiologic solution.
The eluate was centrifuged at 8,500 rpm for 5 minutes. The pellet was suspended in 400 ml of saline and divided in two equal aliquots, one of which was submitted to DNA extraction, the other was stored at -80°C. The DNA, extracted by proteinase K and phenol-chloroform method, was resuspended in 100 µl of water and stored at -20°C until use. Primers and probe sequences were selected from the β-tubulin gene. The forward (TricF) and reverse (TricR) primer sequences are 5'-CACCTCAAGCTCAACAACCCAACA-3' and 5'-TGCCGGACATAACCATGGA-3' respectively. The minor groove binder probe TricP (5'-ACGGCGATCTTAAC-3') contained a fluorescent reporter dye (6-carboxy-fluorescein, FAM) at the 5'-end. The reaction was performed in a final volume of 50 µl containing 5 µl of DNA solution, 600 nM TricF, 600 nM TricR, 200 nM TaqMan Probe, and 25 µl TaqMan Universal PCR Master Mix (Applied Biosystems, Foster City, CA). Real-time PCR reaction was carried out with the ABI PRISM 7900 HT sequence detection system according to the following program: one cycle of 95°C for 10 min, followed by 45 cycles of 95°C for 15 s and 60°C for 1 min.

DISCUSSION

We had the opportunity to collect the relevant casuistic at the Clinics for STDs outpatients, part of the Service of Microbiology and Virology of the Padua’s Azienda Ospedaliera. The feasibility of the study was increased by the facilities concerning the technology of molecular diagnostic medicine, connected to the University Department of Histology, Microbiology and Medical Biotechnology. The method developed by our group was designed to target a well known gene, β-tubulin. Primers were selected in order to amplify a shorter stretch of DNA (66 bp) than in a former system (13), aiming at a more efficient duplication process. Nonetheless, the specificity for T vaginalis was maintained, as confirmed by comparing the amplicon with a gene bank.
In order to obtain a quantitative assay of the protozoan load, a procedure based on interpolation with a calibration curve was set up. Another enhancement with respect to the methods published formerly was the addition of an inner standard, a real-time PCR amplification system targeted to a human gene, ß-globin. This was co-amplified within the same reaction tube. So doing, the adequacy of the sample and the possible presence of inhibitors were routinely assessed, easing the conversion of the assay into a quantitative molecular detection. In this context, the present paper is the first report on a quantitative real-time PCR detection of *T. vaginalis* in clinical samples. The molecular approach resulted to be very accurate. Three apparent false positives by real-time PCR probably reflected a superior sensitivity compared to culture. Ease of execution and short processing time were additional advantages. The accuracy of the method confirms previous reports based on real-time PCR methods (9,6). The quantitative potential of the TaqMan method was also documented. The prevalence of *Trichomonas vaginalis* in the population presenting at our diagnostic center was reliably evaluated. By real-time PCR the prevalence of *T. vaginalis* in the female population was 17.9% (immigrants), 0.6% (natives), and 3.8% (overall). The present report is the largest evaluation of the prevalence of the infection done in Italy with the aid of the most accurate diagnostic tool presently available. Moreover, the risk factor weighing on the immigrant population is apparent.

In conclusion, the target of confirming the effectiveness of the molecular approach was achieved. The real-time PCR was clearly the most accurate way to detect the protozoon. The method appears to be feasible and convenient in a busy laboratory, provided the necessary equipment is available. Moreover, our real-time PCR is likely to be suitable also to urethral swabs of male patients. Because of difficulties with diagnosing trichomoniasis in men, the implementation of a molecular method to detect *Trichomonas vaginalis* could

**Table 1. Sensitivity, specificity, PPV and NPV of real-time PCR and microscopy**

<table>
<thead>
<tr>
<th>Method</th>
<th>Sensitivity</th>
<th>Specificity</th>
<th>PPV</th>
<th>NPV</th>
</tr>
</thead>
<tbody>
<tr>
<td>real-time PCR</td>
<td>100%</td>
<td>99.5%</td>
<td>86.4%</td>
<td>100%</td>
</tr>
<tr>
<td>Microscopy</td>
<td>94.7%</td>
<td>100%</td>
<td>100%</td>
<td>99.8%</td>
</tr>
</tbody>
</table>

The qualitative accuracy indexes are reported for two tests and for the overall, unsegmented population. The gold standard is liquid culture detection of *Trichomonas vaginalis*.

PPV: positive predictive value. NPV: negative predictive value

**Table 2. Positivity for Trichomonas vaginalis in foreigners, natives, and overall casuistic**

<table>
<thead>
<tr>
<th>Immigrants</th>
<th>Natives</th>
<th>Overall</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>positives %</td>
<td>positives %</td>
</tr>
<tr>
<td>real-time PCR</td>
<td>19</td>
<td>17.9%</td>
</tr>
<tr>
<td>microscopy</td>
<td>14</td>
<td>13.2%</td>
</tr>
<tr>
<td>culture</td>
<td>15</td>
<td>14.2%</td>
</tr>
</tbody>
</table>

Risk ratios, indicating the increase of the base risk due to immigrant condition, and the related p-values, are indicated. Data are stratified for the three diagnostic tests under scrutiny

RR: Risk ratio
increase the reliability of estimates of prevalence of this infection in the male population. Indeed, the sensitivity of microscopic examination of wet mounts or cultures, for male samples, was reported to range from 60% to 80% (12), whereas the sensitivity of urethral swab culture or urine sample was 62.2% or 61.6% , respectively (10). Moreover, the potential application of PCR to urine samples could avoid the urethral swab to male patients.

The present study also explored the potential of the technique as an epidemiological investigation tool, furnishing preliminary data on females in a local geographical context, situated on North East Italy.

REFERENCES