A multiplex PCR assay for the simultaneous detection of *Chlamydia trachomatis*, *Neisseria gonorrhoeae*, and *Trichomonas vaginalis*

Ahmad N. Abou Tayoun, Paul R. Burchard, Angela M. Caliendo, Axel Scherer, Gregory J. Tsongalis

**1. Introduction**

An estimated 499 million new cases of curable sexually transmitted infections (STIs) occur every year, with around 19 million in the United States alone (http://www.cdc.gov/std/stats10/default.html; Coleman et al., 2013; Mahony et al., 1995; Blake et al., 2008). Such infections (STIs) occur every year, with around 19 million in the United States alone (http://www.cdc.gov/std/stats10/default.html; Manam et al., 2013; Blake et al., 2008). Such STIs often appear asymptomatic and, if untreated, can have devastating consequences on reproduction, maternal and newborn health, as well as increase the likelihood of acquisition and transmission of HIV (http://www.cdc.gov/std/stats10/default.html; Katusiime et al., 2013; Beharry et al., 2013). The World Health Organization (WHO) has identified rapid, low-cost, and accurate point-of-care diagnostic tests as a key point for action in the global strategy for the prevention and control of infectious diseases-related to travel; sexually transmitted diseases; http://www.who.int/mediacentre/factsheets/fs110/en/index.html; Owusu-Edusei et al., 2013; Chesson et al., 2004). For developing countries, traditional methods for diagnosing STIs are laborious, often not very sensitive, and have a long turnaround time with most recent commercially available diagnostic tests targeting one or, at most, two of these STIs at a time. Here, we describe the development of a highly sensitive, rapid and affordable sample-to-answer multiplex PCR-based assay for the simultaneous detection of *Trichomonas vaginalis*, *Neisseria gonorrhoeae*, and *Chlamydia trachomatis*.

**Materials and Methods:** We designed a multiplex PCR assay for the detection of 4 targets (CT, TV, NG, and process/PCR control) using melt curve analysis. To establish the limit of detection (LOD) for each pathogen, we used previously extracted and quantified TV, NG, and CT genomic DNA (Vircell, Spain). For each target, the LOD was determined by lowering its copy number while increasing the other two STI loads in a stepwise fashion. The process/PCR control remained constant in the optimized assay and was spiked into each sample before extraction. For a concordance study, we tested urine, vaginal and rectal swab specimens from 26 patients positive for one or more of the tested STIs. In addition, 56 liquid cytology specimens (Thinprep) were used to assess specificity.

**Results:** This assay has a turnaround time of less than 2 h and has a limit of detection as low as 7–31 copies for each STI in the presence of the other 2 targets. Our assay also demonstrated 100% concordance with 26 known clinical samples from urine, vaginal and rectal swab specimens. TV, NG, CT, and our process/PCR control were consistently identified at 78 °C, 82.3 °C, 85.7 °C, and ~92 °C, respectively. When applied to DNA extracted from residual Thinprep specimens, the assay was negative in 54/56 samples. Two samples were found to be co-infected with CT.

**Conclusions:** Our multiplex assay combines a rapid and cost-effective approach to molecular diagnostics with the versatility required for use within a variety of laboratory settings. These performance characteristics make this multiplex STI assay highly suitable for use in a clinical laboratory.

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Keywords: Chlamydia, Neisseria, Trichomonas, PCR

In developing countries, traditional methods for diagnosing STIs are laborious, often not very sensitive, and have a long turnaround time with most recent commercially available diagnostic tests targeting one or, at most, two of these STIs at a time (http://www.cdc.gov/std/stats10/default.html; Mahony et al., 1995; Marangoni et al., 2012; Doseeva et al., 2011; Choe et al., 2013). However, studies have shown that 45.7% of persons infected with NG, are co-infected with CT (Datta et al., 2007). With approximately 106 million new Neisseria gonorrhoeae cases occurring every year, such co-infections are abundant in the population (http://www.who.int/mediacentre/factsheets/fs110/en/index.html). In addition, the incidence of co-infection with either CT, NG or both accounts for over 2% of STI cases or roughly 11 million cases per year (Ginocchio et al., 2012; van Veen et al., 2010). This combined with the high prevalence of TV signifies the necessity for testing all three targets in any given specimen to ensure the highest level of sensitivity as well as reducing the number of unnecessary treatments (Marangoni et al., 2012; Doseeva et al., 2011; Choe et al., 2013). However, studies have shown that 45.7% of persons infected with NG, are co-infected with CT (Datta et al., 2007).

2. Materials and methods

2.1. Samples

A total of 83 patient samples were used in this study. In the concordance study, 26 samples including 10 urine, 9 vaginal and 7 rectal swab specimens were obtained from patients that have previously tested positive for CT and NG using the Abbott m2000 real-time CT/NG assay, and TV using a laboratory developed Real-Time PCR assay (Caliendo et al., 2005). For the sensitivity study, 26 samples including 10 urine, 9 vaginal and 7 rectal swab specimens were obtained from patients that have previously tested positive for CT and NG using the Abbott m2000 real-time CT/NG assay, and TV using a laboratory developed Real-Time PCR assay (Caliendo et al., 2005). For the sensitivity study, limit of detection was established using genomic DNA extracted from known numbers of TV, NG and CT organisms (Vircell, Spain). For the specificity study, we used 57 female patient samples previously tested for Human Papillomavirus (HPV) with the Roche COBAS® 4800 HPV Assay.

2.2. Extraction

For each patient sample, 2 ml (urine) or 600 μl buffer (swabs) was pelleted by centrifugation at 6000 rpm for 10 min and re-suspended with 195 μl of water. Each sample was then spiked with 5 μl of 2.5 pg/μl internal control DNA (IC2M, see Table 1). DNA was then extracted with the Qiagen EZ1 robotic system using the bacterial DNA card and the EZ1 tissue extraction kit/cartridges (Qiagen, Valencia, CA). DNA was eluted in 50 μl elution buffer, of which 10 μl was used for the 25 μl PCR reactions.

2.3. PCR

Internal control (IC2M) and primer sequences (Table 1) were synthesized by Integrated DNA Technologies (Coralville, Iowa). The primer sequences were compared in silico to human, viral, and bacterial genomes within the NCBI database. No significant similarities between the targeted sequences and any of the tested genomes that could theoretically lead to non-specific amplification were found.

For patient samples, we prepared 25 μl PCR reactions consisting of 1× SsoFast™ EvaGreen® Supermix, 10 μl extracted DNA (this should theoretically contain ~2.5 pg IC2M, see above), 100 nM forward and reverse CT primers, 175 nM forward and reverse NG primers, 125 nM forward and reverse TV primers and 150 nM IC2M forward and reverse primers. For the limit of detection studies, DNA extracted from a known copy number of each target was spiked into the PCR reaction, in addition to 2.5 pg IC2M.

PCR amplification was performed on the SmartCycler® (Cepheid, Sunnyvale, CA) using a touchdown PCR protocol (Fig. 1A) that includes a 5 minute hot start (95 °C) followed by 2 cycles of 5 second denaturation (95 °C) and 30 second annealing/extension at 70 °C, 2 cycles of 5 at 95 °C and 30 s at 67 °C, 2 cycles of 5 s at 95 °C and 30 s at 65 °C, 2 cycles of 5 s at 95 °C and 30 s at 63 °C, and finally 32 cycles of 5 s at 95 °C and 30 s at 60 °C. A melt protocol was also included from 60–95 °C at 0.2 °C/s (Fig. 1A).

3. Results

3.1. Multiplex STI assay

Our PCR-based assay was developed for the simultaneous detection of three major STIs (CT, NG, and TV) in addition to a novel internal control DNA sequence (IC2M). The latter was spiked into each patient sample before extraction and thus served as both a process and PCR control (see Materials and methods). All primer sequences were carefully designed and selected to clearly resolve the characteristic intercalating dye-based melt peaks for the four different targets in our multiplex assay (Fig. 1B). In our primer design, we targeted multi-copy regions within the three STI genomes to achieve highest sensitivity. For
Chlamydia trachomatis, we targeted the multi-copy cryptic plasmid thus, not only enhancing sensitivity, but also ensuring the detection of all Chlamydia trachomatis serovars including the Swedish nvCT serovar (Caliendo et al., 2005; Palmer and Falkow, 1986; Unemo & Clarke, 2011). After extensive PCR optimization, we devised a touchdown PCR protocol with extension temperatures gradually decreasing from 70 °C to 60 °C as shown in Fig. 1A. With this optimal protocol, we were able to precisely resolve the melt peaks for TV at 78 °C, NG at 82.3 °C, CT at 85.7 °C, and IC2M at 92 °C (Fig. 1B).

3.2. Sensitivity or limit of detection

To establish the limit of detection (LOD) for each pathogen, we spiked into our PCR reactions genomic DNA extracted from known copy numbers of TV, NG, and CT (Vircell, Spain). For each target, the LOD was determined by lowering its copy number while increasing the other two STI loads in a stepwise fashion (Tables 2–4). For example, the LOD for Neisseria gonorrhoeae was 7 or 15 copies/reaction even when either TV or CT was present at a high concentration (3000 copies/reaction), respectively. This LOD was increased to 31 copies/reaction when both TV and CT were each present at 3000 copies/reaction (Table 2 and Fig. 2). Although the LOD for either TV or CT was 7 copies/reaction in the absence of the other two pathogenic targets, this LOD ranged between 7 and 31 copies/reaction as the other STIs were increasingly spiked (up to 3000 copies) into the PCR reaction (Tables 3 and 4).

Overall, our assay demonstrates a very high sensitivity for the detection of each of the STI targets even when the other targets were present at very high copy numbers.

3.3. Specificity

We assessed the specificity of the assay by testing specimens which had been previously submitted for HPV screening. We extracted DNA from 56 female patient Thinprep liquid cytology specimens. Twelve of these specimens were positive for high risk HPV. Of the 12 positive HPV samples, we identified 2 as also positive for CT. This was not a surprising finding as co-infection of HPV and CT occurs in approximately 14% of HPV positive women (Verteramo et al., 2009), which is

![Fig. 1. A. Representation of touchdown real-time PCR reaction profile. B. SmartCycler real-time PCR melt curve analysis results for Neisseria gonorrhoeae (NG), Chlamydia trachomatis (CT), and Trichomonas vaginalis (TV) with equal copy numbers of each in the reaction mixture.](image-url)

**Table 2**

<table>
<thead>
<tr>
<th>Chlamydia trachomatis (LOD)</th>
<th>Neisseria gonorrhoeae</th>
<th>Trichomonas vaginalis</th>
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<tbody>
<tr>
<td>7</td>
<td>0</td>
<td>0</td>
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<tr>
<td>7</td>
<td>Low&lt;sup&gt;a&lt;/sup&gt;</td>
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<tr>
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<td>High&lt;sup&gt;b&lt;/sup&gt;</td>
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<tr>
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<tr>
<td>31</td>
<td>High</td>
<td>High</td>
</tr>
</tbody>
</table>

<sup>a</sup> 7 copies per reaction.  
<sup>b</sup> 3000 copies per reaction.

**Table 3**

<table>
<thead>
<tr>
<th>Neisseria gonorrhoeae (LOD)</th>
<th>Chlamydia trachomatis</th>
<th>Trichomonas vaginalis</th>
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<tbody>
<tr>
<td>7</td>
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<td>31</td>
<td>High</td>
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<sup>a</sup> 7 copies per reaction.  
<sup>b</sup> 3000 copies per reaction.
consistent with our findings. The other HPV positive samples as well as the 44 HPV negative samples showed only amplification of the internal control. This suggests proper DNA extraction and no cross reactivity between positive HPV samples or genomic DNA with our assay. In addition, we tested five gram-positive and five gram-negative bacteria samples representing normal vaginal flora. Our assay identified all ten of these bacteria samples as negative for our STI targets, which was expected. This suggests no cross reactivity between our assay and common vaginal flora.

Furthermore, of the 26 known positive swab and urine samples (see below), our assay never gave an unexpected melt peak for any of the tested targets. This further supports our primer sets as specific for each target and that they do not cross react with one another or the internal control, even at high concentrations.

### 3.4. Accuracy

To determine the accuracy of our assay, we extracted DNA from 26 samples for patients previously shown to be positive for CT, NG, and/or TV and tested them using our assay. The specimens used for this accuracy study were from urine, vaginal swabs, and rectal swabs. Our assay correctly identified 26 of the 26 previously tested patient samples. TV and CT infections were found in 8 samples and NG infection was found in 6 samples. Co-infections with CT and TV, NG and TV, and CT and NG, were found in 1 sample each. These results were 100% concordant with what was expected and demonstrate a high level of accuracy for our assay from a variety of specimen sources.

### 3.5. Turnaround time

The total turnaround time for this assay is less than 2 h. The assay begins with a 10 minute centrifugation, followed by a 15 minute automated extraction. PCR amplification takes approximately 1 h to set up and run. The results of the melt-curve are available within a few minutes of completion of the amplification and easy to interpret.

### Table 4

<table>
<thead>
<tr>
<th>Trichomonas vaginalis (LOD)</th>
<th>Chlamydia trachomatis</th>
<th>Neisseria gonorrhoeae</th>
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</thead>
<tbody>
<tr>
<td>7</td>
<td>0</td>
<td>0</td>
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<tr>
<td>15</td>
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<td>15</td>
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\(^a\) 7 copies per reaction.  
\(^b\) 3000 copies per reaction.

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**Fig. 2.** SmartCycler real-time PCR melt curve analysis results for Neisseria gonorrhoeae (NG) limit of detection. Plotted as the fluorescence with respect to temperature. Copy number refers to the copies of NG in the reaction mixture. High Chlamydia trachomatis (CT) and Trichomonas vaginalis (TV) correspond to 3000 copies of each in the reaction mixture.
4. Discussion

In this study, we report proof of principle for a new multiplex method for the simultaneous detection of the three most common STIs; TV, CT, and NG from various specimen sources including Thinprep liquid cytology samples. This multiplex PCR-based assay uses an intercalating dye and melt-curve analysis to determine the presence of each STI. Such an assay can be made affordable, highly sensitive and can be performed on real-time PCR platforms common to any molecular diagnostics laboratory. Multiplexing provides the advantage of detecting multiple targets at the same time and from the same specimen. Furthermore, the ability to limit the diagnostic process to one test minimizes both cost and time.

Unlike fluorescently labeled probes, the use of intercalating dyes and melt curve analysis greatly lowers the cost of this assay while maintaining high specificity and sensitivity. Our assay also uses an automated DNA extraction system, which minimizes the required hands-on time. This not only limits the possibility of contamination but also makes for a safer extraction process. The turnaround time for our assay is less than 2 h, which drastically reduces the turnaround time as compared to more traditional culture based methods.

Our results demonstrate this assay as highly sensitive, specific, and accurate. The LOD for this assay was very low for each STI, even in the presence of high copy numbers of one or both of the other STIs. Our assay will therefore be able to easily identify co-infection amongst all three of these STIs. In addition, our results were 100% concordant with what was expected for the previously tested patient specimens. Specimens containing one or more of the desired STIs were equally identifiable, which was expected. In addition, these specimens exhibit the wide range of samples that can be used in our assay, such as urine, vaginal swabs, and cervical cytology samples. This assay combines a rapid and cost-effective approach to molecular diagnostics with the versatility required for use within a variety of laboratory settings. These performance characteristics make this multiplex STI assay highly suitable for use in a clinical laboratory.

Acknowledgments

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