Review

From Wet Mount to Nucleic Acid Amplification Techniques: Current Diagnostic Methods and Future Perspectives Based on Patenting of New Assays, Stains, and Diagnostic Images for *Trichomonas vaginalis* Detection

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Abstract: Trichomoniasis is the most common non-viral sexually transmitted infection (STI) in the world. The estimated global prevalence in 2016 was 156 million adults aged 15–49. However, these data are underestimated, since the most used diagnostic method is the wet mount, which has low sensitivity, the information regarding the estimated duration of infection is limited and there is evidence of undiagnosed asymptomatic cases in both sexes. Currently 80% of cases—including both sexes—are asymptomatic, which makes the disease silent and chronic in course, leading to complications. The aim of this review was to discuss the diagnostic methods for *T. vaginalis* detection that are currently available and applicable in the clinical laboratory routine. Overall, nucleic acid amplification techniques are the best option for *T. vaginalis* detection, with higher sensitivity and specificity than other tests. Although these techniques present higher cost, their implementation should be supported to ensure correct trichomoniasis diagnosis and treatment beyond contributing to questions on epidemiology and control.

Keywords: *Trichomonas vaginalis*; trichomoniasis; diagnosis; nucleic acid amplification test (NAAT’s)

1. Introduction

Trichomoniasis is notably the most predominant non-viral sexually transmitted infection (STI) worldwide. In 2016, the World Health Organization (WHO) estimated 156 million cases of *Trichomonas vaginalis* infections worldwide, constituting almost half of the total global incidence of STIs among adults aged 15 to 49 years [1]. However, these data are underestimated because the most commonly used diagnostic method is the wet mount examination, which has variable sensitivity (50%), reaching 50% to 80% sensitivity in the absence or presence of *colpitis macularis* [2]. *T. vaginalis* can cause symptomatic infections in the vulvar and urethral areas of the genital tract. However, 80% of cases—including both sexes—are asymptomatic, which makes the disease silent and chronic, leading to complications such as pelvic inflammatory disease, cervical and prostate cancer, premature birth, and low birth weight in newborns [3,4]. The association of trichomoniasis with the increased transmission and acquisition of HIV/AIDS in a bidirectional relationship is well established, supporting the HIV epidemic in populations where trichomoniasis is endemic [5,6]. Trichomoniasis is currently treated with nitroimidazoles, mainly metronidazole, tinidazole, and secnidazole, which are approved by the Food and Drug Administration (FDA/USA) [7]. Although these are low-cost drugs and most cases are curable, approximately 2.0 to 28% are caused by drug-resistant *T. vaginalis* isolates, which contributes to therapy failures [8–10].

The impact of trichomoniasis on public health has become increasingly important and better understood, encompassing both the direct and indirect costs associated with
its treatment [11]. Data analyzed from 2016 to 2018 demonstrated a higher treatment cost for female patients (USD 220) than for male patients (USD 158), which is 70% higher than the costs reported in previous studies from 2001–2005 [12]. This increased cost can reach USD 167 million per year when comparing cases of HIV facilitated by the presence of the protozoan [13]. Therefore, accurate and early diagnosis is crucial for interrupting transmission and preventing complications associated with the infection. Trichomoniasis remains an underreported disease due to the inability to fulfill all the necessary criteria. The underreporting of cases, the occurrence of asymptomatic individuals, the increasing reports of *T. vaginalis* isolates resistant to nitroimidazoles, and the absence of public policies for prevention, detection, and treatment contribute to the lack of control of this STI. This situation will gradually become an unsustainable public health strategy. Efforts are needed to identify alternatives to mitigate the spread of trichomoniasis, along with attention towards and investment in diagnostic laboratory methods for detecting *T. vaginalis* and accessing qualified professionals to carry out an accurate diagnosis.

This review addresses laboratory diagnostic methods for *T. vaginalis* detection as well as an update on new methods that have been developed through patent filings around the world. The main methods currently available, described in Table 1, include: wet mount examination [14], cultural exam [15], staining [16], rapid tests [17,18], and molecular techniques based on nucleic acid amplification test (NAAT), such as conventional PCR [19,20], as well as other more recent tests for rapid identification that can often detect more than one sexually transmitted pathogen [21,22].

**Table 1.** Most common diagnostic methods for the detection of *T. vaginalis* approved by the Food and Drug Administration (FDA/USA) [7,18].

<table>
<thead>
<tr>
<th>Assay</th>
<th>Equipment Requirements</th>
<th>Sample Type</th>
<th>Cost 1</th>
<th>Relative Sensitivity 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Wet mount</td>
<td>Microscope</td>
<td>vaginal and endocervical swabs, urine sediment</td>
<td>$</td>
<td>+</td>
</tr>
<tr>
<td>Staining method</td>
<td>Microscope</td>
<td>vaginal and endocervical swabs, urine sediment</td>
<td>$</td>
<td>++</td>
</tr>
<tr>
<td>Culture</td>
<td>Incubator, microscope</td>
<td>vaginal and endocervical swabs, urine sediment</td>
<td>$$</td>
<td>++</td>
</tr>
<tr>
<td>Sorology (antigen)—OSOM&lt;sup&gt;®&lt;/sup&gt;</td>
<td>None</td>
<td>vaginal and endocervical swabs</td>
<td>$$</td>
<td>+++</td>
</tr>
<tr>
<td>Nucleic acid test (NAAT)—Affirm VPIII&lt;sup&gt;®&lt;/sup&gt;</td>
<td>Affirm VPIII instrument</td>
<td>vaginal and endocervical swabs</td>
<td>$$</td>
<td>++</td>
</tr>
<tr>
<td>Nucleic acid test (NAAT)—Hologic Aptima TV&lt;sup&gt;®&lt;/sup&gt;</td>
<td>Tigris or Panther automated system</td>
<td>endocervical swabs, vaginal swabs, endocervical specimens collected in PreservCyt&lt;sup&gt;(®)&lt;/sup&gt; (Thin Prep, Hologic Incorporated, MA, USA) solution and female urine specimens</td>
<td>$$$</td>
<td>+++</td>
</tr>
<tr>
<td>Nucleic acid test—Solana Trichomonas Assay&lt;sup&gt;®&lt;/sup&gt; (Quidel)</td>
<td>Solana’s platform</td>
<td>vaginal swabs and urine specimens</td>
<td>$$</td>
<td>+++</td>
</tr>
<tr>
<td>Nucleic acid test (PCR) Max CTGCTV2 assay&lt;sup&gt;®&lt;/sup&gt; (Becton Dickinson)</td>
<td>Benchtop instrument, BD CTGCTV2</td>
<td>vaginal swab specimens and male or female urine specimens</td>
<td>$$$</td>
<td>+++</td>
</tr>
</tbody>
</table>
Table 1. Cont.

<table>
<thead>
<tr>
<th>Assay</th>
<th>Equipment Requirements</th>
<th>Sample Type</th>
<th>Cost 1</th>
<th>Relative Sensitivity 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nucleic acid test (PCR)—Cepheid®6</td>
<td>GeneXpert Instrument</td>
<td>Vaginal and endocervical swabs, female or male urine sediment</td>
<td>$$$</td>
<td>+++</td>
</tr>
<tr>
<td>Nucleic acid test (PCR real time multiplex) The Allplex™ STI Essential assay</td>
<td>Seegene®7</td>
<td>vaginal and endocervical swabs, urine sediment</td>
<td>$$$</td>
<td>+++</td>
</tr>
</tbody>
</table>

Legend: 1. The symbols represent different price ranges, $—low cost; $$—intermediate cost; $$$—high cost. 2. The symbols indicate a range of sensitivity, +—low sensitivity; ++—intermediate sensitivity; +++—high sensitivity. 3. Hologic, Inc., San Diego, United States. 4. Quidel Corporation, Ohio, United States. 5. Becton Dickinson, New Jersey, United States. 6. Cepheid, Sunnyvale, United States. 7. Seegene, Inc., Seul, Republic of Korea.

2. Methods

For this study, scientific databases such as Pubmed (https://pubmed.ncbi.nlm.nih.gov, accessed on 10 September 2023), Clinical Trials (https://clinicaltrials.gov, accessed on 22 September 2023), and various patent databases covering different regions, Europe (Espacenet, https://worldwide.espacenet.com, accessed on 22 September 2023), Brazil (INPI, https://www.gov.br/inpi/pt-br, accessed on 22 September 2023), Canada (CIPO, https://www.ic.gc.ca, accessed on 22 September 2023), the United States (USPTO, https://www.uspto.gov/, accessed on 22 September 2023), Australia (AUSPAT, https://pericles.ipaustralia.gov.au/, accessed on 21 February 2024), LATIPAT (Latin America, https://lp.espacenet.com/, accessed on 21 February 2024), and China (CPO, accessed on 21 February 2024) have been used. The search was limited to articles published within the last ten years (2013–2023) using the keywords “Trichomoniasis” OR “Trichomonas vaginalis” AND “Diagnosis”. The inclusion criteria were: (1) articles that aimed to compare or to evaluate the performance of methods in diagnosing trichomoniasis; (2) articles that used methodologies applicable to routine clinical laboratory practice; (3) articles that discussed diagnostic techniques for Trichomonas vaginalis and other associated STIs. The exclusion criteria were languages other than English, Portuguese, and Spanish, and year of publication before 2013.

3. Consolidated Methods for Trichomoniasis Diagnosis

3.1. Wet Mount

The accuracy of diagnosing T. vaginalis has notably improved over the past decade, with the availability of a broader range of tests. Wet mount is a traditional and common method for diagnosing trichomoniasis. The method is low-cost, low-tech, and easy to prepare, but it has low sensitivity, especially in men. The wet mount examination involves collecting material through a swab and visualizing the parasite in secretions (vaginal, endocervical, urethral, urine sediment) using a conventional microscope. This is based on trophozoite morphological characteristics such as pear shape, the presence of flagella, an undulating membrane, and asynchronous motility.

The main limitation of this technique is its immediate execution requirements, which cannot exceed a few hours since the organisms lose motility ex vivo due to temperature differences. Therefore, the slides must be prepared and analyzed as soon as possible after clinical sample collection to avoid false-negative results. In fact, the microscopic examination should be conducted within 10 min after collection for the most accurate results [23]. To achieve a successful diagnosis using a wet mount, it is also crucial to have a qualified professional present who can identify the parasite, even under abnormal conditions regarding morphology and motility. Furthermore, the parasite load required to make an accurate diagnosis is not yet standardized [24]. Such limitations compromise the sensitivity of the technique, which is generally lower than that of other methods. Sensitivities range from 50 to 70%, depending on the reader’s experience [23]. A cross-sectional study evaluated the diagnostic precision of the wet mount and PCR methods.
against culturing and considered reference standard for directly diagnosing *T. vaginalis* among symptomatic women. The swabs were tested for *T. vaginalis* with wet mount microscopy (WMM) in-house PCR and *T. vaginalis* culturing. The sensitivity and agreement kappa of the WMM were observed to be lower compared to PCR. However, the specificity for both methods was high, with 100% (95% confidence interval 97–100) for WMM and 99.3% (95% confidence interval 96–100) for PCR, respectively. Among the *T. vaginalis*-positive women, a decrease was observed in the sensitivity of the WMM, compromising treatment of two-thirds of the patients. The authors recommend further research to integrate PCR tools into diagnostic algorithms for trichomoniasis [25].

3.2. Staining Techniques

Staining techniques can be combined with wet mount examination to enhance the diagnosis of trichomoniasis. Among the most commonly used staining technique in clinical laboratories for the identification of *T. vaginalis* trophozoites is the Papanicolaou smear, which can be easily applied to urine sediment samples. Urine sediment is a clinical sample that is readily available and abundant. It is a relatively rapid technique (staining time of application, fixing, drying, and reading) and is reasonably cost-effective for implementation in laboratory routines. A meta-analysis conducted on research published between 1976 and 1998 concerning the efficacy of Papanicolaou staining in detecting vaginal trichomoniasis revealed a cytological evaluation specificity of 97%. Other staining and fixation methods have also been described in the literature, such as Giemsa and Gram staining. However, it has been demonstrated that Giemsa staining does not allow for the identification of the characteristic structures of the parasite, while Gram staining does not provide satisfactory fixation performance and the identification of morphological features [16]. Indeed, staining methods are not recommended in the clinical laboratory routine due to the potential for false negatives and false positives [7]. A presumptive diagnosis of *T. vaginalis* in a Papanicolaou staining smear can be identified by a perinuclear halo in epithelial cells. It is worth noting that even with well-trained professionals present for identification, the absence of the parasite does not necessarily exclude the possibility of the patient being infected [26]. While *T. vaginalis* may incidentally appear in a Pap test, it is essential to understand that neither conventional nor liquid-based Pap smears serve as diagnostic tests for trichomoniasis. According to the Sexually Transmitted Infections Treatment Guidelines from CDC, women who test positive for *T. vaginalis* on a Pap smear should undergo retesting using sensitive diagnostic methods, and treatment should be administered if the infection is confirmed [27].

3.3. Culture Exam

The culture examination for *T. vaginalis* demonstrates superior sensitivity compared to the wet mount test. Samples from women (vaginal, cervical, endocervical secretions or urine sediment) or men (urethral secretion or urine sediment) must be immediately inoculated into the culture medium after collection. Cultures are maintained at 37 °C and meticulously observed under a microscope daily for up to 5 days until motile trophozoites are detected [28]. Typically, cultures from women with trichomoniasis yield positive results within the initial 3 days following inoculation. Nonetheless, male cultures require daily examination for a period of 5 days or longer before being deemed negative. Extended incubation times are frequently necessary to facilitate the growth of a discernible quantity of organisms from male specimens [28]. In contrast to the sensitivity of Nucleic Acid Amplification Techniques, the sensitivity of culture examination varies from 44% to 75% for detecting *T. vaginalis* in female samples. For men, culture sensitivity ranges from 40% to 56% for detecting this pathogen. Notably, urine from men demonstrates higher sensitivity for culture compared to a urethral specimen. Liquid culture media are relatively affordable; however, their cost is augmented by the necessity of extended incubation time and daily scrutiny by a skilled microscopist, leading to results taking up to a week [29]. Historically, culture techniques such as the InPouch system (BioMed Diagnostics, White Vite, OR, USA)
were deemed the gold standard for diagnosing *T. vaginalis* infection prior to the advent of NAAT [7].

### 3.4. Point-of-Care Tests

Rapid or point-of-care (POC) tests for the detection of *T. vaginalis* have also gained prevalence in clinical practice. The OSOM Trichomonas Rapid Test is based on the detection of antigens from *T. vaginalis*. This immunochromatographic capillary-flow enzyme immunoassay employs membrane proteins that can detect the parasite in up to 10 min. Although it features an increase in the cost related to the material, a reduction is observed in monthly labor costs and the time devoted to microscopy. Moreover, the test can be performed as part of high-volume laboratory analysis, and shows specificity in cases of infection with low trichomonads charge loads [30]. Compares with the wet mount method, the OSOM test has a better result, with sensitivities ranging from 83 to 90%. Furthermore, it does not require special instruments for its analysis, and therefore its use is quite common in gynecological clinics, emergency rooms, and in self-test programs [31].

In 2008, an observational study conducted at the University of Pittsburgh, identified by Clicantrials.gov ID NCT00682851, evaluated, validated the accuracy of two rapid tests (OSM Trichomonas and OSOM BVBlue test) for diagnosing trichomoniasis and bacterial vaginosis, respectively. The sensitivity for symptomatic women was 92% (confidence interval: 78 to 98), while for asymptomatic women, it was 91% (confidence interval: 71 to 99). Regarding specificity, it was 99% (confidence interval: 97 to 100) for both symptomatic and asymptomatic women [32].

Point-of-care tests can also be based on acid nucleic detection. AffirmTM VPIII (Becton Dickinson, MD, USA) is an unamplified acid nucleic probe use as a POC test to detect *T. vaginalis*, *Gardnerella vaginalis* and *Candida albicans*. The Affirm VPIII is approved by the FDA and uses specific oligonucleotide probes to detect nucleic acid of *T. vaginalis*. The processing and test samples require a heating unit and a processor [28]. The performance results of this assay are better than those of wet mount and culture examination, and present excellent specificity to *T. vaginalis*, but are significantly less sensitive than NAAT [33]. The Solana Trichomonas assay (Quidel) is an additional rapid test designed for the qualitative detection of *T. vaginalis* DNA, which can deliver results in under 40 min following specimen collection. This assay is FDA-approved for diagnosing *T. vaginalis* from both female vaginal and urine specimens, and is applicable to asymptomatic and symptomatic women. It boasts a sensitivity exceeding 98% when compared to NAAT for vaginal specimens and over 92% for urine specimens. The Amplivue Trichomonas assay (Quidel) is another rapid test allowing the qualitative detection of *T. vaginalis*. It has been FDA-cleared for vaginal specimens from both symptomatic and asymptomatic women, boasting a sensitivity of 90.7% and a specificity of 98.9% when compared to NAAT. It is important to note that neither the Osom assay nor the Affirm VP III test have FDA clearance for use with specimens from men [34].

### 3.5. Molecular Based Methods

Due to the precision of molecular biology tests, these techniques have revolutionized laboratory diagnostics by elucidating the genes of microorganisms as well as the products they encode. Technologies based on nucleic acid amplification are surprising as regards their high performance in STI diagnosis, thus reducing analysis time and enabling the detection of infections in non-invasive ways. In this sense, the search for new diagnostic alternatives has been undertaken using innovative molecular techniques or new detection targets. In this scenario, a study employed a multiplex PCR assay to assess its capacity for concurrently detecting *T. vaginalis*, *N. gonorrhoeae* and *C. trachomatis* using urine, liquid cytology, and swabs from vaginal and rectal sites. Its results can be obtained within 2 h, showcasing a low detection limit even when other targets are present, consistent with findings from previously tested patient samples. This versatile multiplex STI assay offers a rapid and cost-effective approach to molecular diagnostics, catering to diverse laboratory
settings. These attributes collectively render it exceptionally well-suited for deployment in clinical laboratories [35]. The Max CTGCTV2 assay (Becton Dickinson, New Jersey, NJ, United States) represents an advanced iteration of molecular triplex techniques for identifying *Chlamydia trachomatis*, *Neisseria gonorrhoeae*, and *Trichomonas vaginalis*. It is FDA-approved for use in detecting *T. vaginalis* in both vaginal swab and urine specimens, offering sensitivity and specificity of 86.6% to 97.9% and 99.2% to 99.8%, respectively, depending on the type of specimen, be it vaginal swab (97.8% and 99.6%), endocervical swab (89.9% and 99.8%), and male urine (97.9%) [36]. A comparative study of the diagnostic performance of the BD MAX vaginal panel (Becton, Dickinson and Company, BD Life Sciences—Diagnostic Systems, New Jersey, NJ, United States) molecular test versus the clinician assessment of vaginitis was performed, using Amsel’s criteria for bacterial vaginitis, the presence of pseudohyphae or budding yeast for candidiasis, and wet mount microscopy for *T. vaginalis*. The authors illustrated a notably elevated sensitivity and negative predictive value of molecular testing compared to clinician-administered tests. This enhancement aids in the precise identification of vaginitis [37]. The Allplex™ STI Essential assay (Seegene®, Seoul, Republic of Korea) utilizes a multiplex Real-Time PCR (RT-PCR) technique as its foundation [38]. This in vitro diagnostic (IVD) system, bearing European Conformity (EC) marking, was engineered for the concurrent identification of seven pathogens—*Chlamydia trachomatis*, *Neisseria gonorrhoeae*, *Mycoplasma hominis*, *Mycoplasma genitalium*, *Ureaplasma parvum*, *Ureaplasma urealyticum*, and *T. vaginalis*—using genital swabs, urine, and cytology liquid. In a comparative study conducted by Salazar et al. in 2019, Aptima® assays (Hologic®, San Diego, United States) demonstrated greater sensitivity across various sample types compared to the Allplex™ STI Essential assay (Seegene®) [39].

Techniques currently recognized as the reference for detecting *T. vaginalis* include those that amplify and detect the parasite’s nucleic acids (DNA or RNA). The first FDA-approved NAAT was the APTIMA® (Hologic, Inc., San Diego, United States) *Trichomonas vaginalis* Assay, which was introduced in 2011 and has been utilized ever since [39,40]. This technique shows sensitivity ranging from 95.2% in urine samples to 100% in vaginal and cervical secretion samples. The parasite is identified in urine sediment and samples collected for oncolytic cytology. This technology relies on extracting rRNA from the specimens being analyzed, followed by the transcription-mediated amplification of the captured rRNA. Utilizing this target naturally enhances the detection limit, given that each organism contains over 100 copies of rRNA. These amplified products are detected by a chemiluminescent reaction [41]. Recently, the performances of Hologic Aptima BV and CV/TV assays were compared to those of other methods of vaginitis diagnosis using more than 600 samples from a local health system. The authors described 100% of positive and negative agreement for *T. vaginalis* detection between Aptima CV/TV and Aptima TV technologies [42]. The Xpert® TV Assay (Cepheid, Sunnyvale, CA, United States) technology applies the real-time PCR technique through disposable cartridges for the qualitative detection of *T. vaginalis* and several STIs, such as chlamydia and gonorrhea, with positive results in approximately 40 min of analysis at the POC. First-catch urine, endocervical, and vaginal swab samples can be analyzed. Moreover, the specificity and sensitivity were found to be high and comparable when using both female and male samples [17,43]. The Probe Tec TV Qx Amplified DNA Assay (Becton Dickinson) is FDA-cleared for detecting *T. vaginalis* from vaginal (patient-collected or clinician-collected) swabs, endocervical swabs, or urine specimens from women. It boasts a sensitivity of 98.3% and a specificity of 99.6%, compared to wet mount and culture methods [22]. Like the Aptima *T. vaginalis* assay, this test is exclusively FDA-cleared for female use and requires internal validation before application in male specimens.

Loop-mediated isothermal amplification (LAMP) is a molecular biology technique used for the rapid and specific amplification of DNA under isothermal conditions. The widespread applications in several fields, including clinical diagnostics, environmental monitoring, and research, relate to its high sensitivity and specificity; reactions typically produce results within 30 min to a few hours, and it also shows versatility [44]. A novel
detection approach for *T. vaginalis* has been developed utilizing loop-mediated isothermal amplification targeting the adhesion protein 65 (AP65) gene. This rapid detection method was fine-tuned to optimize the reaction system and the conditions for ideal performance. Analyses of sensitivity indicated that the LAMP assay, focusing on the AP65 gene, displayed a sensitivity level surpassing that of the commonly employed nested PCR, which targets the actin gene for detecting *T. vaginalis*, by a factor of 1000. The LAMP assay was found to have a minimum detection limit as low as 10 trichomonads. Furthermore, the amplification of the target gene AP65 via the LAMP assay exhibited outstanding specificity, yielding products solely from *T. vaginalis*. Importantly, this LAMP detection technique did not exhibit any cross-reactivity with common pathogens, such as *Candida albicans*. Based on the findings of this study, the LAMP assay directed at the AP65 gene emerges as an effective approach for the early detection of *T. vaginalis* infections. Consequently, the researchers propose the LAMP assay as a valuable point-of-care diagnostic tool, although it is not commercially available yet, and it offers an alternative molecular approach with significant potential for enhancing the treatment, control, and prevention of trichomoniasis transmission and related complications [45]. Considering the global health concern of cervical cancer, primarily driven by high-risk human papillomavirus (HPV) infections and the connections with trichomoniasis, researchers have introduced a novel approach using a microfluidic-chip-based system paired with loop-mediated isothermal amplification (LAMP), enabling the swift and simultaneous identification of *T. vaginalis*, as well as specific HPV types (HPV16, HPV18, and HPV52). The system offers enhanced sensitivity, cost-effectiveness, and facility of use, making it particularly applicable in resource-constrained settings. Additionally, its capacity to detect multiple pathogens’ positions makes it a versatile tool with potential applications beyond cervical cancer diagnostics.

4. Clinical Trials

Clinical Trials is a USA website and online database that provides information about current clinical studies around the world. Among these, five completed studies have specifically focused on diagnostic techniques. The first multi-center study was an interventional type study entitled “Clinical study of a Single-Use, Point-of-Care Molecular Diagnostic Device for the Detection of *N. gonorrhoeae*, *T. vaginalis*, and *C. trachomatis* utilizing vaginal swabs” (NCT03596151). The hypothesis of this study was that the Click Diagnostics Sexual Health Test performs comparably to the NAAT predicate system, and the identification of each organism in self-collected vaginal swabs by women using the Click device will show high sensitivity and specificity, aligning with the Patient Infected Status (PIS). The primary aim of this study was to evaluate the effectiveness of the Click device in detecting *Chlamydia trachomatis*, *Neisseria gonorrhoeae*, and *T. vaginalis* in self-collected vaginal specimens as compared to PIS. The sensitivity of the Click device was 96.7%, while the specificity was 94.2% [46]. Another observational study (NCT02566447) was conducted to establish the clinical performance of the Solana® Trichomonas Assay for the detection of *T. vaginalis* in both vaginal swabs and urine samples. This assay is an NAAT that utilizes helicase-dependent amplification (HDA) for the diagnosis of trichomoniasis. The results of this study were not available on the website [47]. Another observational study used the APTIMA® assay (NCT01728103), which qualitatively detects the ribosomal RNA from *C. trachomatis* and/or *N. gonorrhoeae* and *T. vaginalis* in female specimens [48]. NCT02203942 was a study that validated NAAT testing by comparing it with conventional methods for the diagnosis of vaginitis, including the Amsel criteria, Nugent score, yeast culture, and trichomonad culture. However, at this moment there are no results available on the website [49]. Another study (NCT02641717) validated patient-collected wet mounts by comparing them with clinician-collected specimens from symptomatic women. Currently, there is limited literature on this topic. Therefore, self-collection can enhance the performance and efficiency of diagnosing vaginitis [50].
5. Patents

The patent databases from the following countries and continents were consulted: Brazil (INPI, n = 1), Canada (CIPO, n = 5), Korea (KIPO, n = 3), United States (USPTO, n = 19), Europe (Espacenet, n = 12), Australia (AUSPAT, n = 1), Latin America (LATIPAT, n = 1), and China (CPO, n = 8) (Figure 1). The identified techniques included staining, culturing, NAATs, and novel approaches (Table 2).

![Figure 1. Worldwide distribution of registered patents by country. ( ): Number inside parentheses indicates the number of patents on the respective methodology. NAAT: Nucleic Acid Amplification Tests.](image-url)

5.1. Staining Technique

Only one patent was found that describes the staining technique for diagnosing atypical trichomoniasis when an infection occurs in the bloodstream, which is very rare. In summary, these procedures are divided into the collection of peripheral blood from the patient, sample preparation, the enzymatic destruction of blood elements, a smear on a slide, staining with 1% aqueous methylene blue solution, and finally, analysis under a microscope. This method is an efficient technique for diagnosis and has high sensitivity [51].

5.2. Culture Exam

Before the advent of NAATs, the isolation of the protozoa in specific culture media was the gold standard for the diagnosis of trichomoniasis. Regarding this approach, a technique has been developed that combines cultural examination—to detect the presence of *T. vaginalis*—with the preparation of smears and Giemsa staining—to assess the patient’s leukocytosis. The culturing of the urogenital tract sample is performed in a selective medium and incubated at 37 °C for three days. Subsequently, the stained smears are analyzed using methylene blue and the Feulgen method [52]. Another patent that has been developed describes a new liquid medium for the isolation of the parasite. Some of the components of the medium include thioglycolate, glycine, human blood plasma, and glucose, which promote the growth of *T. vaginalis* [53].

5.3. Nucleic Acid Amplification Techniques—NAATs

NAATs are the diagnostic approach with the most patents in all consulted databases. One invention presents nucleic acid-based tests for detecting vaginitis and/or vaginosis-causing pathogens in samples from symptomatic patients. Samples can be collected from various areas including the urethra, penis, anus, throat, cervix, or vagina. The tests can identify several pathogens including *Trichomonas vaginalis, Atoleopium vaginae, Gardnerella vaginalis, Lactobacillus spp.*, and *Candida spp.* [54–56]. Furthermore, *T. vaginalis* can be detected along with *Chlamydia trachomatis, Neisseria gonorrhoeae*, and *Mycoplasma spp.* through an assay based on the amplification of oligonucleotide primers and probes [57,58]. Another
invention that specifically detects *T. vaginalis* by multiplex PCR in a single test tube and discriminates *T. tenax* and *Pentatrichomonas hominis* was also patented [59,60].

One nested polymerase chain reaction (PCR) method can detect *T. vaginalis* in urine. It uses a specific gene sequence, AP51-3, to design a primer. The method involves amplification and electrophoretic steps. It is straightforward to use, highly sensitive, and specific [61]. This patent that presents the primer composition of F3, B3, FIP, and BIP primers, and can be utilized for LAMP amplification in *T. vaginalis* detection; it also exhibits high sensitivity and specificity [62]. Another method for diagnosing trichomoniasis involves the detection of the TV 40S ribosomal protein gene through PCR, which is carried out in mostly substantially automated manner. The sample can be taken from an endocervical, vaginal, or urethral swab [63,64]. A diagnostic kit for diagnosing vaginitis at low costs was developed in the Republic of Korea. The device is divided into a sample introducing unit, a nucleic acid extraction unit, and a PCR unit [65].

Oligonucleotides are short nucleic acid sequences that can be used to determine the presence of *T. vaginalis* in a biological sample, as described in a patent. The determination occurs through the multi-phase amplification of a target nucleic acid sequence [66,67]. Another patent provides oligonucleotides and methods for the simultaneous detection of *T. vaginalis* and *Mycoplasma genitalium* through the multiplex detection of nucleic acids using mixed reporters [68].

Different methods allow the simultaneous detection of pathogens causing sexually transmitted infections. A system for generating diagnostics based on detecting microbiome targets has been patented. It includes a sampling kit, diagnostic analyses to generate a microbiome sequence dataset from microorganism nucleic acid sequences, and therapy recommendations [69]. In addition, next-generation sequencing (NGS) can analyze IST pathogens and human papillomavirus (HPV) in any sample [70]. Another method uses loop-mediated isothermal amplification (LAMP), whereby released nucleic acids are amplified by LAMP using specific primers targeting these pathogens’ nucleic acids [71].

### 5.4. Biomarker Tests

Peptidases are important enzymes in *T. vaginalis*, aiding in protein breakdown. Peptidases are divided into types such as cysteine (CP), metallo (MP), serine (SP), threonine (TP) and aspartic (AP) [72]. Methods for the production of antibodies against metallopeptidase (TvMP50) and TvMP50 recombinant contribute in diagnosing trichomoniasis in men [73,74]. TvCP39, another peptidase, is used for *T. vaginalis* detection [75]. An inhibitor of TvCP39 can serve as a diagnostic marker for this infection, such as trichocystatin 2 protein, a cysteine proteinase inhibitor [76]. Another useful cysteine proteinase for *T. vaginalis* diagnosis is TvCP4 [77]. TvCP2 levels, useful for immunodiagnostics and as markers of trichomoniasis, can also be detected [78]. Pyruvate-ferredoxin oxidoreductase (PFOR) is a key enzyme in flagellated protozoa, aiding pyruvate oxidation and acetyl-coA production [79]. A patent has introduced an immunological method for detecting *T. vaginalis* in bodily secretions and urine, identifying PFOR and adhesion proteins as markers [80]. Another patent utilizes polymerase chain reactions for the amplification and detection of the *T. vaginalis* AP65-1 gene by exposing a biological sample to an oligonucleotide probe [81].

### 5.5. Novel Approaches

#### 5.5.1. Devices of Images

In the USA’s database, new methods for diagnosing trichomoniasis can be found. The first invention is a urine specimen analyzer, which obtains information about the counts of *T. vaginalis* trophozoites, squamous epithelium cells, and white blood cells. This equipment consists of a detector that identifies urine particles and an analysis unit responsible for counting [82]. Another invention is an imaging platform for the detection of motile objects in a fluid sample. In this context, *T. vaginalis* is a flagellated protozoan; therefore, it can be detected through a computing device that receives images from the sensors and light sources used to analyze the sample [83]. Furthermore, another patent describes a device
with a microfluidic module associated with an image sensor and a processor. This device can be utilized for motion-based pathogen detection [84].

5.5.2. Other Devices

One characteristic of vaginitis is the high concentration of amines at the infection site. For this reason, a personal care product with an amine-sensitive dye was created to visually identify this infection. This invention can be useful in the diagnosis of vaginitis if the indicator is placed in products used for feminine hygiene [85]. Another invention offers a method for detecting and diagnosing STIs using a specific string of epitopes (SOE). In the case of diagnosing trichomoniasis, the SOE can detect the following proteins in vaginal fluid, semen, or prostatic fluid: aldolase, glyceraldehyde-3-phosphate dehydrogenase (GAPDH), α-enolase, and actinin [86].

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INPI—Instituto Nacional da Propriedade Industrial (Brazil); CIPO—Canadian Patents Database; KIPRIS—Korea Intellectual Property Rights Information Service; USPTO—United States Patent and Trademark Office; STIs—sexually transmitted infections; LAMP—loop-mediated isothermal amplification; PCR—Polymerase Chain Reaction; SOE—string of epitopes; HPV—human papillomavirus; NGS—next-generation sequencing; BR—Brazil; CA—Canada; USA—United States of America; LA—Latin America; AU—Australia; EU—Europe; CN—China; ROK—Republic of Korea. * Biological marker.

6. Conclusions

Trichomoniasis caused by *Trichomonas vaginalis* is a highly prevalent STI associated with an increased risk of the acquisition and transmission of HIV, as well as premature birth, pelvic inflammatory disease, cervical and prostate cancers, and infertility. Recent advances in the diagnosis of this underdiagnosed infection, especially using molecular methods, are of paramount importance for the epidemiological control of this pathology and its comorbidities, especially in asymptomatic cases. As the application of more sensitive diagnostic techniques based on nucleic acid detection tests has been seen, more cases of *T. vaginalis* infection will be detected, and more infections that appeared clinically cured may be identified as present in an asymptomatic form. Among symptomatic women, wet mount microscopy is recommended, with special attention paid to ensure immediate execution in a few hours so as to avoid the loss of the organism’s motility ex vivo, followed by death. The inclusion of trichomoniasis in screening tests is an important step as a strategy for the reduction and control of this and related ISTs, along with cost reduction, through drug treatments. In this review, our search for clinical trials related to trichomoniasis diagnosis granted us access to both ongoing and completed studies evaluating the accuracy, sensitivity, and specificity of diagnostic tests. Additionally, we identified 40 patents related to trichomoniasis diagnosis within the last 10 years, indicating a significant presence in the STI diagnostics scenario. This suggests a competitive landscape among numerous companies and institutions striving to offer novel inventions with potential industrial applications soon. Our assessment underscores the importance of researchers meticulously considering the entire translational process concerning infection management. This process encompasses the entire evolution from laboratory investigations to the refinement of clinical testing methodologies, alongside the innovation of novel patents and solutions aimed at addressing prevailing challenges. Considering that the clinical tests demonstrate similarities to other approved methods, our article emphasizes the necessity of continued studies concerning diagnostic targets in *T. vaginalis*, as well as the pursuit of innovative approaches.

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