Molecular detection of *Trichomonas foetus* in bovine samples: a novel real-time polymerase chain reaction (PCR) assay targeting *EF1-alpha-Tf1* and a comparative study of published PCR techniques

Coral Polo1,2 · Teresa García-Seco1 · Víctor Fernández3 · Marta Hernández4 · Victor Briones1,5 · Alberto Díez-Guerrier2,5 · Lucas Domínguez1,5 · Marta Pérez-Sancho1,5

Received: 11 October 2021 / Accepted: 9 March 2022 / Published online: 29 March 2022 © The Author(s) 2022

Abstract

The parasite *T. foetus* causes trichomonosis in cattle but is generally asymptomatic in males. Thus, many bulls carrying the disease go unnoticed, making the detection of *T. foetus* in bulls an important aspect for its control. Due to drawbacks posed by its cultivation, PCR is a preferred option for diagnostic laboratories. Most published PCR protocols target the genomic region comprising the 18S, 5.8S, and 28S rRNA genes and internal transcribed spacers 1 and 2 (rRNA-ITS region), homologous to that of other *Trichomonas* species. There is minimal information on alternative genetic targets and no comparative studies have been published. We compared a protocol based on the microsatellite *TfRE* (called H94) and five protocols based on the rRNA-ITS region (called M06, M15, G02, G05, and N02). We also designed and evaluated a novel PCR-based assay on the *EF1-alpha-Tf1* gene (called V21). The analytical sensitivity and specificity assays for the PCR protocols were performed according to the World Organisation for Animal Health (OIE) directives and the comparative study was performed with a widely used PCR (M06) on clinical samples from 466 breeding bulls. V21 showed a high degree of agreement with our reference M06 (kappa = 0.967), as well as M15 (kappa = 0.958), G05 (kappa = 0.948), and H94 (kappa = 0.986). Protocols H94 and V21 appear to be good approaches for confirming clinical cases in preputial bull samples when genomic regions alternative to rRNA-ITS are required. By contrast, N02 gave false negatives and G02 false positives.

Keywords *Trichomonas foetus* · Real-time PCR · *EF1-alpha-Tf1* · Bull

Introduction

The protozoan *Trichomonas foetus* is a flagellated pathogen that causes trichomonosis in cattle, cats, and pigs (Dąbrowska et al. 2019b). In bulls, trichomonosis is an asymptomatic and persistent venereal disease, whereas in cows, it causes cervicitis, endometritis, foetal death during the first trimester of gestation, a delayed return to estrus, and infertility (Michi et al. 2016). In addition, *T. foetus* is transmitted through both natural breeding and artificial insemination (Givens 2018). In this scenario, the control of *T. foetus* in males is crucial for its control within cattle herds. Due to its relevance in production, bovine trichomonosis is a notifiable disease in certain countries.

The detection of *T. foetus* is generally carried out by microscopy, culture, and molecular techniques, such as PCR (OIE 2018). However, sampling and transport conditions can affect the detection rate of *T. foetus*, primarily via direct microscopy and culture (Yao 2013). In recent years,
molecular detection has become one of the most widely used options in diagnostic laboratories (OIE 2018). The OIE cited real-time PCR, mentioning the work of McMillen and Lew (McMillen and Lew 2006) (here called the M06 protocol) as a suitable method for the detection of *T. foetus* in clinical samples (OIE 2018). This widely used PCR assay (Effinger et al. 2014; Dąbrowska et al. 2019a; Meggiolaro et al. 2019) is based on the genomic region comprising the 18S, 5.8S, and 28S rRNA genes and internal transcribed spacers 1 and 2 (rRNA-ITS region), a multicopy target (Chakrabarti et al. 1992) used for the internal transcribed spacers 1 and 2 (rRNA-ITS region), region comprising the 18S rRNA genes and 5.8S rRNA genes and internal transcribed spacers 1 and 2 (rRNA-ITS region), a multicopy target (Chakrabarti et al. 1992) used for the design of various published PCR assays (Felleisen et al. 1998; Gookin et al. 2002, 2005; Nickel et al. 2002; McMillen and Lew 2006; Mueller et al. 2015; Ginter Summarell et al. 2018; Dąbrowska et al. 2019a), most of them used for the diagnosis of clinical samples (Köster et al. 2015; Casteriano et al. 2016; Li et al. 2016; Dąbrowska et al. 2020). Despite the value of this diagnostic approach, to our knowledge, no systematic comparative study of all the PCR assays for the detection of *T. foetus* has been published. In addition, similarities with rRNA-ITS sequences present in other *Tritrichomonas* species, such as *Trichomonas mobilenensis*, have been described, resulting in potential cross-reactions (Dąbrowska et al. 2019a). Alternative genetic regions have been studied (Reinmann et al. 2012; Šlapeta et al. 2012; Sun et al. 2012) for different purposes. Nevertheless, two alternatives PCR targets for the diagnosis of *T. foetus* directly in clinical samples, which may overcome these potential drawbacks, have been described: (i) *TfRE* microsatellite-PCR (Ho et al. 1994), with minimal information (only one reference sequence in GenBank (AY435432.1) that comes from an unpublished study), and (ii) a commercial kit that targets the *beta-tubulin 1* gene that lacks published data (OIE 2018). In this context, it would be useful to explore new molecular targets for the detection of *T. foetus* as a confirmatory test, for example, in the event of doubtful results. Oyhenart et al. recently published a novel loop-mediated isothermal amplification (LAMP) assay targeting the *EF1-alpha-Tf1* gene for the detection of *T. foetus* on bull preputial wash samples (Oyhenart 2018). This LAMP assay showed high analytical sensitivity (0.5 trophozoites/mL) and specificity, making this genetic target a valuable alternative for the confirmation of *T. foetus* in clinical samples (Oyhenart 2018). However, no PCR assay for *T. foetus* detection directly in clinical samples based on this target has yet been published.

Here, we aimed to perform a comparative study of all molecular target-PCR techniques currently published for the detection of *T. foetus* to assess their diagnostic performance on a cohort of breeding bulls raised in an extensive regimen. In addition, we propose a new real-time PCR assay targeting the *EF1-alpha-Tf1* gene, developed as an alternative method for the detection of *T. foetus* in bull preputial wash samples.

### Materials and methods

#### Systematic review and selection of PCR protocols

We conducted a systematic bibliographic search in PubMed, search string: (Tritrichomonas foetus) AND (PCR)); Web of Science, search string: TS:=(Tritrichomonas AND foetus AND PCR); and Scopus, search string: TITLE-ABS-KEY (“Tritrichomonas foetus”) AND (“PCR”). Among the papers found (*n* = 614), duplicates were removed using EndNote X8 software and the abstracts were manually screened. All publications that mentioned the use of a PCR technique for *T. foetus* detection (41 publications) were further evaluated according to the following inclusion criteria: (i) abstracts must contain information about the *T. foetus* genetic target, (ii) primers and probe sequences must be published, (iii) PCR and thermal cycler conditions must be provided, and (iv) due to the technological limitations of many diagnostic laboratories, the PCR design should be based on DNA detection (not RNA due its fragility in field samples) and, just in case of real-time PCR protocols, use the most common fluorescent systems (SYBRgreen or Taqman probes) for *T. foetus* identification. When the same primer set was used in more than one study, only one was included in the present study based on the following prioritization criteria: (i) those that were assessed on clinical samples and/or (ii) those adapted to real-time protocols, in particular, those using specific probes. Finally, six published protocols (five targeting the rRNA-ITS genomic region and one *TfRE*) were included in the study (Table 1). The locations of the selected primers within the rRNA-ITS region are shown in Fig. 1.

All PCR protocols were carried out under the conditions of the original paper with minor changes for their adaptation to our laboratory (optimal conditions of the Qiagen PCR kits used: QuantiFast Pathogen PCR, QuantiFast SYBR Green PCR, and Qiagen Multiplex PCR kits). For real-time PCR, a probe specific for an internal PCR control was used (provided by the QuantiFast Pathogen PCR kit).

#### Design and optimization of a novel real-time PCR based on the EF1-alpha-Tf1 gene

The design of an in-house PCR based on the *EF1-alpha-Tf1* gene (called V21, Table 1) was carried out using Oligo (Oligo 7 Primer Analysis Software) and HM217356.1 from the GeneBank database as the reference sequence. The specificity of the primers and probe were verified in silico by blastn using BLAST online software from the NCBI website (https://www.ncbi.nlm.nih.gov/). The primers were
| Target | PCR type | Reference (code\textsuperscript{a}) | Primers and probes (5' to 3') | PCR conditions\textsuperscript{c} | Fw (µM) | Rv (µM) | Pr (µM) | ID. (C°/min) | D. (C°/s) | A. (C°/s) | E. (C°/s) | Nº | FE. (C°/min) | AS. (bp) |
|--------|----------|-------------------------------------|-------------------------------|----------------------------------|--------|--------|--------|-------------|--------|---------|----------|-----|------------|--------|
| ITS    | Real time (probe) | McMillen and Lewis 2006 (M06) | TFF2: GCGGCTGGA TTAGCCTTCTTT TFR2: GGCGGGCAAA TGTCAT TRICHIP2: 6-FAM- ACAAG TTCGAT CCTTG-MGB-BHQ | | 0.90 | 0.90 | 0.08 | 95/5\textsuperscript{d} | 95/20 | 60/45 | 40 | 57 |
| ITS    | Real time (SYBR) | Mueller et al. 2015 (M15) | TFR3: CGGGTCTTC CTATAT GAGACA GAACC TFR4: CCTGC CGTT GATCAT TCTTG TAA | | 0.20 | 0.20 | 95/5 | 95/30 | 63/20 | 40 | 348 |
| EF     | Real time (probe) | Current study (V21) | FtEF1A1: AGTCGG CGGCC AAATCAA RfEF1A1: CTCTTC AACCTCGG GCTGT GA PtEF1A1: 6-FAM- ATCATCAAGTAC GCCTGA GAT- MGB- NFQ | | 0.50 | 0.50 | 0.40 | 95/5 | 95/20 | 61/35 | 45 | 76 |
| ITS    | Conventional (nested) | Gookin et al. 2002 (G02) | TFR3: CGGGTCTTC CTATATGAGACA GAACC TFR4: CCTGC CGTT GATCAT TCTTG TAA TFFTS F: CTGC CGTT GATCAT TCTTG TAA TFFTS R: GCAATG TGC ATCTAAAGA TGG | | 1.25 | 1.25 | 95/15 | 95/30 | 57/30 | 72/30 | 50 | 72/10 | 347 |
| ITS    | Conventional | Gookin et al. 2005 (G05) | 499F: GCTCG TAGT CAGAA ACTGC 1140R: CCCAATTAG AACTCT ATCTC | | 0.30 | 0.30 | 95/15 | 95/60 | 60.1/60 | 72/120 | 35 | 72/10 | 642 |
### Table 1 (continued)

| Target | PCR type | Reference (code\(^b\)) | Primers and probes (5’ to 3’) | PCR conditions\(^c\) |  |
|--------|----------|-------------------------|-----------------------------|---------------------|---|
|        |          |                         |                             | Fw (µM) | Rv (µM) | Pr (µM) | ID. (°C/min) | D. (°C/s) | A. (°C/s) | E. (°C/s) | Nº | FE. (°C/min) | AS. (bp) |
| ITS    | Conventional | Nickel et al. 2002 (N02) | TF211A: CCTGCCGGTGGAAGTTTGTTTCGTTTA |
|        |          |                         |                             | 0.20 | 0.20 | 95/15 | 94/60 | 60/60 | 72/60 | 35 | 72/10 | 211 |
|        |          |                         | TF211B: GCGCAATGTTGTCAGATTCAAAAGATT CG |
| TIRE   | Conventional | Ho et al. 1994 (H94) | TF1: CATTATCCCCAAATGGTATAAC |
|        |          |                         |                             | 0.38 | 0.365 | 95/15 | 94/60 | 45/60 | 72/120 | 41 | 72/10 | 162 |
|        |          |                         | TF2: GTGATTAAGTACATAATTTC |

\(^a\)Genomic region that comprises the 18S, 5.8S, and 28S rRNA genes and the internal transcribed spacer 1 and 2 of *T. foetus* (ITS), EF1-alpha-Tf1 gene (EF), and microsatellite TfRE sequence (TIRE)

\(^b\)Internal code that references the protocols used in the current study

\(^c\)PCR were performed in a 25 µL of final reaction volume for M06, V21, G02, G05, N02, and H94 protocols and 20 µL for M15 protocol: Fw (µM), final concentration of forward primer; Rv (µM), final concentration of reverse primer; Pr (µM), final concentration of probe; ID. (°C/min), initial denaturation; D. (°C/s), denaturation step per cycle; A. (°C/s), annealing step per cycle; E. (°C/s), elongation step per cycle; Nº, number of cycles; FE. (°C/min), final elongation step; AS. (bp), amplicon size

\(^d\)A previous step at 50 °C during 2 min was added

\(^e\)It is a one-step nested PCR
synthesized by Eurofins Genomics (Eurofins Genomics, GmbH) and the probe by ThermoFisher (Applied Biosystems, UK). The primer/probe sequences and PCR conditions are shown in Table 1. The QuantiFast Pathogen PCR kit (Qiagen, Germany) and an internal PCR control provided in the kit were used to verify the lack of PCR inhibition.

Analytical sensitivity and specificity tests

Conventional PCR protocols were performed using C1000 and T100 thermocyclers (Bio-Rad, USA) and the PCR product detected following gel electrophoresis. For all conventional PCR assays, electrophoresis was performed in 2% agarose gels (Biotools MB agarose) at 70 mV and 400 mA for 55 min. Real-time PCR assays were performed in a CFX96 thermocycler (Bio-Rad, USA) using specific conditions (Table 1).

The validation data from the original studies (and subsequent studies), which assessed the specificity of each PCR assay, were taken into account for the analytical specificity test (Gookin et al. 2002, 2005; Nickel et al. 2002; McLennan and Lew 2006; Effinger et al. 2014; Mueller et al. 2015; Ginter Summarell et al. 2018; Dąbrowska et al. 2019a; Meggiolaro et al. 2019). In addition, information about the specificity of the genetic target on which the novel PCR protocol was based was considered (Oyhenart 2018). A panel of pathogens (field or reference strains) associated with bovine infertility or potentially present in preputial samples was also included (one strain/species). The field isolates were part of the VISAVET-UCM collection (Universidad Complutense de Madrid, Spain) and type strains came from the ATCC collection: Arcobacter butzleri, Escherichia coli, Aeromonas hydrophila, Brucella abortus, Toxoplasma gondii, Salmonella enterica, Yersinia enterocolitica, Campylobacter fetus, C. fetus subsp. venerealis, C. fetus subsp. fetus (ATCC-27374 TM reference), and C. fetus subsp. testudinum (ATCC-BAA-2539 TM reference).

The ATCC-30232 strain was used as a control for the analytical sensitivity test of each evaluated PCR. The analysis was performed according to the OIE instructions (OIE 2019). The limit of detection (LOD) and corresponding confidence interval (CI) was calculated for each selected protocol using a battery of serial tenfold dilutions (triplicate test for real-time PCR and duplicate test for conventional PCR) starting from 1 ng of DNA/reaction (corresponding to approximately 5750 copies of the T. foetus genome (Benchimol et al. 2017)). The DNA concentration was adjusted using a High Sensitivity DNA Quantitation Kit (Invitrogen, USA) and a Qubit 4 Fluorometer device (Invitrogen, USA). For real-time protocols, the cut-off was established based on the arithmetic mean value of the C_T of the 20 replicates from the LOD test in which the highest dilution of the LOD had a CI ≥ 95% (OIE 2019). For conventional protocols, the LOD and CI were established according to the visual detection of bands in the agarose gel.

Preputial washes and DNA extraction

A selection of 466 samples recovered for routine diagnosis from breeding bulls raised in extensive regimens was included. Samples were selected based on previous molecular results using an in-house PCR designed by Genetics PCR Solutions (GPS, Spain). The samples consisted of 15 mL of preputial washes in PBS, which were centrifuged at 1512 × g for 10 min and stored at −80 °C until analysis. Nucleic-acid extraction was performed using the QIAamp DNA Mini Kit (Qiagen, Germany), with slight modifications: a cellular lysis buffer (20 mM Tris–Cl (pH 8), 2 mM sodium EDTA, 1.2% Triton X-100, and 20 mg/mL lysozyme in a 360 µL final volume) was first added to the sample pellets and the mixture incubated at 37 °C for 1 h. The DNA extraction was subsequently performed according to the manufacturer’s instructions.
Clinical diagnosis classification of the results

In the current study, we used the M06 PCR protocol (McMillen and Lew 2006) as the reference assay, as previously reported (Meggiolaro et al. 2019). We considered non-specific results to be those for which *T. foetus*-positive results were obtained with only one PCR protocol (and negative with M06). The internal PCR control included in this protocol allowed us to verify the lack of inhibition.

The concordance between the results of each protocol and those of the M06 protocol was estimated with Cohen’s kappa coefficient using SPSS software (IBM SPSS Statistics 22.0) according to the following interpretation: 0.0–0.2: insignificant, 0.2–0.4: low, 0.4–0.6: moderate, 0.6–0.8: good, and 0.8–1.0: very good. The percentage of agreement of positive and negative samples of each protocol with the reference was calculated, along with the positive predictive value (PPV) and negative predictive value (NPV).

Results

In vitro analysis: analytical sensitivity and specificity tests

The analytical specificity test revealed no amplification with any control isolated strain for any PCR protocol (including the in-house V21 protocol), being only positive for the *T. foetus* isolate (ATCC-30232). The LOD (CI ≥ 95%) of the M06, M15, G02, G05, and H94 PCR protocols was $1.10^{-4}$ ng of DNA/reaction (approximately 0.575 *T. foetus* genome copies), whereas the N02 protocol had a LOD (CI ≥ 95%) of $1.10^{-3}$ ng of DNA/reaction (approximately 5.75 *T. foetus* genome copies). Further results related to the LOD of each PCR protocol are included in Table 2. After the analytical sensitivity assay, we established a cut-off for real-time protocols. The cut-off for the M06 protocol (here the reference assay) was a $C_T \leq 34$ and that for the M15 protocol a $C_T \leq 32$.

Comparative study on field samples

A summary of the results on the field samples is shown in Table 2. The panel of included field samples was composed of 168 positive samples and 298 negative samples. The percentage of agreement among the PCR protocols targeting the rRNA-ITS region was variable. The results of M15 and G05 showed high agreement with those of M06 (95.8% and 94.6%, respectively, for positive samples and 99.3% for negative samples). The concordance of the results obtained with M15 with those of M06 was very good according to Cohen’s kappa value (0.958), and the PPV and NPV were 0.98 (Table 2). Similarly, Cohen’s kappa value for G05 was 0.948, with a PPV of 1 and a NPV of 0.97 (Table 2). Protocols G02 and N02 showed poorer results. The Cohen’s kappa value for the G02 protocol was 0.587 (moderate concordance), with 70.1% agreement for negative samples and a PPV of 0.64, whereas Cohen’s kappa value for the N02 protocol was 0.700 (good concordance), with 74.4% agreement for positive samples and a NPV of 0.87 (Table 2). The percentage of non-specific results for the rRNA-ITS—PCR protocols also varied; for the M15 protocol, only 2 of 466 total

| PCR protocol | Concordance of positives | Concordance of negatives | PPV | NPV | Non-specific results | LOD (ng/μL) | Cohen’s kappa | Genetic target | Reference |
|--------------|--------------------------|--------------------------|-----|-----|----------------------|-------------|---------------|----------------|----------|
| M06 Ref.*    | 161/168 (95.8%)          | 296/298 (99.3%)          | 0.98| 0.98| 0/466 (0%)           | $4.10^{-6}$ | 0.958         | rRNA-ITS       | McMillen and Lew 2006 |
| M15          | 161/168 (95.8%)          | 296/298 (99.3%)          | 0.98| 0.98| 2/466 (0.4%)         | $5.10^{-6}$ | 0.958         | rRNA-ITS       | Mueller et al. 2015 |
| G02          | 160/168 (95.2%)          | 299/298 (70.1%)          | 0.64| 0.96| 87/466 (18.6%)       | $4.10^{-6}$ | 0.587         | rRNA-ITS       | Gookin et al. 2002 |
| G05          | 159/168 (94.6%)          | 296/298 (99.3%)          | 1   | 0.97| 0/466 (0%)           | $4.10^{-6}$ | 0.948         | rRNA-ITS       | Gookin et al. 2005 |
| N02          | 125/168 (74.4%)          | 295/298 (98.9%)          | 0.99| 0.87| 1/466 (0.2%)         | $4.10^{-5}$ | 0.700         | rRNA-ITS       | Nickel et al. 2002 |
| H94          | 166/168 (98.8%)          | 296/298 (99.3%)          | 1   | 0.99| 0/466 (0%)           | $4.10^{-6}$ | 0.986         | TfRE            | Ho et al. 1994 |
| V21          | 161/168 (95.8%)          | 298/298 (99.3%)          | 1   | 0.98| 0/466 (0%)           | $4.10^{-5}$ | 0.967         | EF1-alpha-Tf1   | Current study |

See Table 1 to know all PCR references. From left to right in the header row is indicated: The PCR protocol (PCR); the agreement of positives with M06 (Concordance of positives) and the concordance of negatives where 168 are positive samples and 298 negative samples for M06 protocol; the positive predictive value (PPV) and the negative predictive value (NPV); the non-specific results (are those samples negative to the reference protocol and positive only by one of the others) where 466 are the total samples; the limit of detection (LOD) with a confidence interval ≥ 95%; the Cohen’s kappa value; the genetic target where it is indicated the genomic region that comprises the 18S, 5.8S, and 28S rRNA genes and the internal transcribed spacer 1 and 2 (rRNA-ITS), TIRE microsatellite, and EF1-alpha-Tf1 gene; and the reference of the original work.

*The method used as reference (Ref.) was the M06 protocol
samples (0.4%) were falsely positive, 87 of 466 (18.6%) for the G02 protocol, and 1 of 466 (0.2%) for the N02 protocol.

The H94 protocol based on the microsatellite TfRE of *T. foetus* showed a Cohen’s kappa coefficient of 0.986 (very good concordance), with high agreement with the M06 protocol for positive and negative samples: PPV = 1 and NPV = 0.99 (Table 2).

**In-house PCR protocol based on the EF1-alpha-Tf1 gene**

The analytical sensitivity test showed a LOD (CI ≥ 95%) of 1.10⁻³ ng of DNA/reaction (approximately 5.75 T. foetus genome copies) and the cut-off was established at a C_T ≤ 40. In terms of the performance of V21 on field samples, this PCR assay showed a Cohen’s kappa value of 0.967 (very good concordance), with 95.8% concordance for positive samples and 100% for negative samples with our reference PCR (M06 protocol) and a PPV of 1 and NPV of 0.98 (Table 2). These results are in accordance with the analytical specificity results of the V21 protocol, which showed no amplification with the isolates included in the specificity test (except *T. foetus* isolates, Table 2).

**Discussion**

*T. foetus* is the etiological agent of trichomonosis, which is generally an asymptomatic and persistent venereal disease in bulls, whereas it can cause a number of reproductive disorders in cows (Michi et al. 2016). In addition, this parasite can be transmitted through natural breeding and artificial insemination (Givens 2018). In this scenario, the detection of *T. foetus* in males is an essential step for the control of trichomonosis. The OIE thus established real-time PCR as a recommended technique for the detection of *T. foetus* in clinical samples (OIE 2018). The OIE guidelines of trichomonosis diagnosis include the PCR protocol of McMillen and Lew (McMillen and Lew 2006) as a suitable assay for *T. foetus* identification in clinical samples. Here, we used this protocol (called M06) as the reference method (as previously considered by Meggiolaro et al. 2019) to compare five different PCR protocols currently available for *T. foetus* detection (Table 1). This is the first report of a systematic comparison of PCR techniques available for *T. foetus* detection directly on bull preputial samples. Five protocols based on rRNA-ITS were selected (Table 1): (1) M06 (McMillen and Lew 2006) (the reference assay in our study), (2) M15 (Mueller et al. 2015), (3) G02 (Gookin et al. 2002), (4) G05 (Gookin et al. 2005), and (5) N02 (Nickel et al. 2002). This genomic region is a multicopy target (Chakrabarti et al. 1992) that appears to be highly similar in other *Trichomonas* species, such as *Trichomonas mobilensis*, resulting in potential cross-reactions, as previously reported for primers TFR3/TFR4 (M15 protocol) (Dąbrowska et al. 2019a).

Although *T. mobilensis* has only been described in the intestine of *Saimiri sciureus* and *Saimiri boliviensis* squirrel monkeys (Scimeca et al. 1989) to date, potential target homology suggests a potential cross-reaction with other *Trichomonas* spp., which could pose a problem in the analysis of field samples. According to our results, G02 (TIFIT-F/TFIT5-R set primers) showed cross-reactions with *Simplicimonas* spp. by in silico analysis of sequences from PCR amplicons sequencing (data not shown). In a previous work by Frey et al. (2017), similar cross-reactions were noted in vaginal swabs of cows and heifers with other set of primers and probes. This data reinforces the idea that PCR protocols based on rRNA-ITS region may provide false positives (Frey et al. 2017; Dąbrowska et al. 2019a). Thus, PCR techniques based on alternative targets could become a useful approach to detect *T. foetus* as a confirmatory test.

In terms of alternative genetic targets for the detection of *T. foetus* directly on clinical samples, one PCR protocol based on the microsatellite TfRE is available: H94 (Ho et al. 1994). This protocol was designed based on a barely studied repetitive region of the *T. foetus* genome, with a single available reference sequence in GenBank (AY435432.1). Finally, according to the OIE (2018), a commercial PCR kit targeting the beta-tubulin 1 gene is also available. However, the information to perform the assay is not published, which is why it was not included in our comparative study. As a consequence, the number of alternatives to rRNA-ITS-based PCR protocols is very limited. Recently, a LAMP technique targeting the EF1-alpha-Tf1 gene showed very good results for *T. foetus* identification in bull preputial samples, being able to detect 0.5 trophozoites/mL (Oyhenart 2018). However, no PCR protocol based on this target has been published. We therefore propose a novel real-time PCR assay (here called the V21 protocol) targeting the EF1-alpha-Tf1 gene as an alternative target for the confirmation of the presence of *T. foetus* in preputial samples (Table 1).

In terms of the protocols targeting the rRNA-ITS genomic region, our results from the comparative study showed very good concordance for the M15 protocol, for which Cohen’s kappa value was 0.958, and the G05 protocol, for which Cohen’s kappa value was 0.948 (Table 2). The M15 protocol is a real-time adaptation (Mueller et al. 2015) of the conventional PCR protocol from the study of Felleisen et al. (Felleisen et al. 1998). Thus, their primers TFR3/TFR4 (Table 1) have been evaluated and used by several studies for *T. foetus* identification (Casteriano et al. 2016; Dąbrowska et al. 2019a), with satisfactory results. On the contrary, it is difficult to find publications that have used the primer pair 499F/1140R (protocol G05, Table 1) (Tolbert et al. 2012). However, we found the diagnostic performance of the G05 protocol to be very similar to that of the M15 and M06 protocols. Thus, the G05 protocol
appears to be relatively good for the detection of *T. foetus* in bull preputial wash samples.

On the other hand, Cohen’s kappa value for the G02 protocol was 0.587 (PPV = 0.6), with 18.6% non-specific results. These results suggest the presence of a high number of false positives, whereas for the N02 protocol, Cohen’s kappa value was 0.700, with a NPV of 0.87, suggesting the presence of false negatives (Table 2). Our results differ from those of the original publications of both protocols (Gookin et al. 2002; Nickel et al. 2002). The absence of additional diagnostic studies using these protocols hampers a proper discussion of this point. Nevertheless, according our results, G02 PCR is a limited technique for the identification of *T. foetus* directly on clinical samples (PPV = 0.64, Table 2). Further studies will be needed to confirm the diagnostic performance of the G02 and N02 protocols.

The H94 protocol, based on the microsatellite *TfRE*, showed a Cohen’s kappa coefficient of 0.986, with no observed non-specific amplification (Table 2), contradicting the results cited in the study of Felleisen et al. (Felleisen et al. 1998). Nevertheless, our results are similar to those of the study of Ho et al. (Ho et al. 1994) and Riley et al. (Riley et al. 1995), in which the use of the primer pair TF1/TF2 appeared to provide high performance for the detection of *T. foetus*. Our study shows that the diagnostic performance of the H94 protocol on field samples appears to be good, with a PPV of 1 and NPV of 0.99.

Finally, Cohen’s kappa coefficient for the in-house V21 protocol (based on the *EF1-alpha-Tf1* gene) was 0.967 (with a PPV of 1 and NPV of 0.98), and there was no non-specific amplification, as for the H94 protocol (Table 2). Thus, V21 protocol seems to be highly specific compared with other PCR designs based on rRNA-ITS genomic region (Table 2) (Frey et al. 2017; Dąbrowska et al. 2019a). Nevertheless, the LOD of the V21 protocol should be considered, as it was an order of magnitude lower than that of M06 (the reference protocol), detecting approximately 5.75 *T. foetus* genome copies per reaction. Based on our results, the use of the H94 and V21 protocols appears to be a good approach for the confirmation of clinical cases when alternative genomic regions to rRNA-ITS are required. An important aspect of the V21 protocol relative to the H94 protocol, for *T. foetus* detection on clinical samples, is the use of an internal PCR control to detect inhibition, the absence of non-specificities, and the rapid obtention of results, as it is a real-time PCR assay with a specific probe.

**Conclusions**

Published PCR protocols based on the rRNA-ITS region showed a high degree of agreement (M15 and G05) with the reference protocol (M06) to identify *T. foetus* in preputial bull samples. However, the rRNA-ITS genomic region appears to be highly similar in phylogenetically close species, which may translate into potential cross-reactions. Thus, protocols based on alternative molecular targets may be useful. The H94 protocol based on the microsatellite *TfRE* (AY435432.1 GeneBank) showed excellent concordance with the reference protocol (Cohen’s kappa coefficient of 0.986), with no non-specific amplification detected. Our study shows the diagnostic performance of H94 in preputial bull samples to be good, capable of detecting approximately 0.575 copies of *T. foetus* genome per PCR reaction.

Our in-house V21 protocol based on the *EF1-alpha-Tf1* gene also showed very high concordance with the reference protocol (Cohen’s kappa coefficient of 0.967), with no non-specific amplification, to detect 5.75 copies of *T. foetus* genome per PCR reaction. Both the H94 and V21 protocols show promise for confirming clinical cases of *T. foetus* when another molecular target is required. An advantage of the V21 over the H94 protocol, for *T. foetus* detection on clinical samples, is the use of an internal PCR control to detect inhibition and the speed to obtain results as it is a specific probe real-time PCR.

**Funding** Open Access funding provided thanks to the CRUE-CSIC agreement with Springer Nature. This work was co-financed by the Community of Madrid (Spain) (IND2018/BIO-9246). In addition, this work was co-funded by a R&D Agreement between the Instituto Tecnológico Agrario de Castilla y León, VISAVET-Universidad Complutense de Madrid, and Universidad de Burgos to carry out research activities on infectious infertility in extensively raised cattle.

**Declarations**

**Ethics approval** The authors confirm that the ethical policies of the journal, as noted on the journal’s author guidelines page, have been adhered to. No ethical approval was required due to sample collection from animals has been gathered.

**Conflict of interest** The authors declare no competing interests.

**Open Access** This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons licence, and indicate if changes were made. The images or other third party material in this article are included in the article’s Creative Commons licence, unless indicated otherwise in a credit line to the material. If material is not included in the article’s Creative Commons licence and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this licence, visit http://creativecommons.org/licenses/by/4.0/.
References

Benchimol M, de Almeida LGP, Vasconcelos AT et al (2017) Draft genome sequence of *Tritrichomonas foetus* strain K. Genome Announc 5:16–17. https://doi.org/10.1128/genomeA.00195-17

Casteriano A, Molini U, Kandjumbwa K et al (2016) Novel genotype of *Tritrichomonas foetus* from cattle in Southern Africa. Parasitology 143:1954–1959. https://doi.org/10.1017/S003318201600115X

Chakrabarti D, Dame JB, Gutell RR, Yowell CA (1992) Characterization of the rDNA unit and sequence analysis of the small subunit rRNA and 5.8S rRNA genes from *Tritrichomonas foetus*. Mol Biochem Parasitol 52:75–83. https://doi.org/10.1016/0166-6851(92)90037-K

Dąbrowska J, Karamon J, Kochanowski M et al (2019a) Development and comparative evaluation of different LAMP and PCR assays for coprological diagnosis of feline trichomoniasis. Vet Parasitol 273:17–23. https://doi.org/10.1016/j.vetpar.2019.07.014

Dąbrowska J, Karamon J, Kochanowski M et al (2019b) *Tritrichomonas foetus* as a causative agent of trichomoniasis in different animal hosts. J Vet Res 63:533–541. https://doi.org/10.2478/jvetres-2019-0072.

Dąbrowska J, Karamon J, Kochanowski M et al (2020) *Tritrichomonas foetus*: a study of prevalence in animal hosts in Poland. Pathogens 9:1–10. https://doi.org/10.3390/pathogens9030203

Effinger L, Peddiredi L, Simunich M et al (2014) Pooling of cultured animal hosts. J Vet Res 63:533–541. https://doi.org/10.2478/jvetres-2019-0072.

Felleisen RSJ, Lambalet N, Bachmann P et al (1998) Detection of *Tritrichomonas foetus* by PCR and DNA enzyme immunoassay based on rRNA gene unit sequences. J Clin Microbiol 36:513–519. https://doi.org/10.1128/JCM.36.2.513-519.1998

Frey CF, Müller N, Stäuber N et al (2017) Simplicomma-like DNA in vaginal swabs of cows and heifers cross-reacting in the real-time PCR for *T. foetus*. Vet Parasitol 237:30–36. https://doi.org/10.1016/j.vetpar.2017.02.024

Ginter Summarell CC, Hairgrove TB, Schroeder ME et al (2018) Improvements in *Tritrichomonas foetus* molecular testing. J Vet Diagnostic Investig 30:603–608. https://doi.org/10.1177/1040387173510003

Givens MD (2018) Review: Risks of disease transmission through semen in cattle. Animal 12:165–171. https://doi.org/10.1017/S1751731118000708

Gookin JL, Birkenheuer AJ, Breitschwerdt EB, Levy MG (2002) Simultaneous detection of *Tritrichomonas foetus* in feline feces. J Clin Microbiol 40:4126–4130. https://doi.org/10.1128/JCM.40.11.4102.

Gookin JL, Birkenheuer AJ, St John V et al (2005) Molecular characterization of trichomonads from feces of dogs with diarrhea. J Parasitol 91:939–943. https://doi.org/10.1645/GE-478R.1

Ho MSY, Conrad PA, Conrad PJ et al (1994) Detection of bovine trichomoniasis with a specific dna-probe and per amplification system. J Clin Microbiol 32:98–104. https://doi.org/10.1128/JCM.32.1.98-104.1994

Köster LS, Chow C, Yao C (2015) Trichomoniasis in cats with diarrhea in Hong Kong, China, between 2009 and 2014. JFMS 424 Open Rep 1(2):2055116915623561. https://doi.org/10.1177/2055116915623561

Li W-C, Wang K, Zhang W et al (2016) Prevalence and molecular characterization of intestinal trichomonads in pet dogs in East China. Korean J Parasitol 54:703–710. https://doi.org/10.3347/kjp.2016.54.6.703

McMillen L, Lew AE (2006) Improved detection of *Tritrichomonas foetus* in bovine diagnostic specimens using a novel probe-based real time PCR assay. Vet Parasitol 141:204–215. https://doi.org/10.1016/j.vetpar.2006.06.012

Meggiolaro MN, Roebel F, Kobylski V et al (2019) Comparison of multiplexed-tandem real-time PCR panel with reference real-time PCR molecular diagnostic assays for detection of *Giardia intestinalis* and *Tritrichomonas foetus* in cats. Vet Parasitol 266:12–17. https://doi.org/10.1016/j.vetpar.2018.12.009

Michi AN, Favetto PH, Kastelic J, Cobo ER (2016) A review of sexually transmitted bovine trichomoniasis and campylobacteriosis affecting cattle reproductive health. Theriogenology 85:781–791. https://doi.org/10.1016/j.theriogenology.2015.10.037

Mueller K, Morin-Adeline V, Gilchrist K et al (2015) High prevalence of *Tritrichomonas foetus* “bovine genotype” in fecal samples from domestic pigs at a farm where bovine trichomoniasis has not been reported for over 30 years. Vet Parasitol 212:105–110. https://doi.org/10.1016/j.vetpar.2015.08.010

Nickel DD, Olson ME, Schultz GA (2002) An improved polymerase chain reaction assay for the detection of *Tritrichomonas foetus* in cattle. Can Vet J 43:213–216

OIE (2018) Trichomonosis. Terrestrial Manual. Chapter 3.4.15.: 1210–1221. https://www.oie.int/fileadmin/Home/eng/Health_standards/tahm/3.04.15_TRICHOMONOSIS.pdf. Accessed 11 January 2022.

OIE (2019) Chapter 1.1.2. Principles and methods of validation of diagnostic assays for infectious diseases, https://www.oie.int/fileadmin/Home/eng/Health_standards/tahm/current/chapitre_valid_ation_diagnostics_assays.pdf. Accessed 11 January 2022.

Oyhenart J (2018) Direct detection of *Tritrichomonas foetus* in cattle genital fluid trough loop mediated isothermal amplification of elongation factor 1 alpha 1. Vet Parasitol 261:67–72. https://doi.org/10.1016/j.vetpar.2018.08.011

Reinmann K, Müller N, Kuhnert P et al (2012) *Tritrichomonas foetus* isolates from cats and cattle show minor genetic differences in unrelated loci ITS-2 and EF-1α. Vet Parasitol 185:138–144. https://doi.org/10.1016/j.vetpar.2011.09.032

Riley DE, Wagner B, Polley L, Krieger JN (1995) PCR-based study of conserved and variable DNA sequences of *Tritrichomonas foetus* isolates from Saskatchewan, Canada. J Clin Microbiol 33:1308–1313. https://doi.org/10.1128/JCM.33.5.1308-1313.1995

Scimeca JM, Culberson DE, Abec CR, Gardner WA (1989) Intestinal trichomonads (*Tritrichomonas mobilensis*) in the natural host *Saimiri sciureus* and *Saimiri boliviensis*. Vet Pathol 26:144–147. https://doi.org/10.1177/031005378902600207

Slapeta J, Müller N, Stack CM et al (2012) Comparative analysis of *Tritrichomonas foetus* (Riedmüller, 1928) cat genotype, *T. foetus* (Riedmüller, 1928) cattle genotype and *Tritrichomonas suis* (Davaine, 1875) at 10 DNA loci. Int J Parasitol 42:1143–1149. https://doi.org/10.1016/j.ijpara.2012.10.004

Sun Z, Stack C, Slapeta J (2012) Sequence differences in the diagnostic region of the cysteine protease 8 gene of *Tritrichomonas foetus* parasites of cats and cattle. Vet Parasitol 186:445–449. https://doi.org/10.1016/j.vetpar.2011.12.001

Tolbert MK, Leutenegger CM, Lobetti R et al (2012) Species identification of trichomonads and associated coinfections in dogs with diarrhea and suspected trichomoniasis. Vet Parasitol 187:1210–1221. https://doi.org/10.1016/j.vetpar.2011.12.031

Yao C (2013) Diagnosis of *tritrichomonas foetus*-infected bulls, an ultimate approach to eradicate bovine trichomoniasis in US cattle? J Med Microbiol 62:1–9. https://doi.org/10.1099/jmm.0.047365-0

Publisher's note Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.