18S rRNA Gene-Based Piroplasmid PCR: An Assay for Rapid and Precise Molecular Screening of *Theileria* and *Babesia* Species in Animals

Binod Kumar1 · Biswa Ranjan Maharana2 · Bhupendrakumar Thakre1 · Nilima N. Brahmbhatt3 · Joice P. Joseph4

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**Abstract**

*Purpose* The parasites of genera such as *Babesia* and *Theileria* are called piroplasmids due to the pear-shaped morphology of the multiplying parasite stages in the blood of the vertebrate host. Because of the enormous number of parasite species and the challenges of multiplex PCR, initial screening of samples using piroplasmid-specific PCR may be a more cost-effective and efficient technique to identify parasite species, especially during epidemiological studies. Accordingly, 18S rRNA PCR was standardized and optimized on common piroplasmids of different animals like cattle, buffaloes, sheep, goats, dogs, horses, and leopards.

*Methods* Blood samples from 1250 animals were collected from different animals in Junagadh district of Gujarat, India. 18S rRNA PCR was standardized and optimized as a primary method for molecular screening of piroplasms in domestic and wild animals. The method was checked for its analytical sensitivity and specificity. Parasite species-specific PCR and sequencing was used to validate the test. Moreover, in-silico restriction enzyme (RE) analysis was also done to assess its applicability in PCR–RFLP.

*Results* Piroplasm infections were recorded in 63.3% of animals in Junagadh. The 18S rRNA PCR detected the piroplasm DNA in as low as 39 picograms (pg) of whole blood genomic DNA isolated from microscopically *Theileria* positive blood samples and no reactivity was recorded from common but unrelated haemoparasites viz., *Trypanosoma evansi*, *Hepatozoon* spp., *Anaplasm* spp., and *Ehrlichia canis* was observed. The 18S rRNA PCR assay findings were confirmed by species-specific PCR and sequencing. Analysis of different sequences generated using 18S rRNA PCR revealed that the amplicon size of *Babesia* spp. is nearly 400 bp (393–408 bp) whereas *Theileria* spp. were more than 400 bp (418–424 bp). The percentage of sequence divergence among *Babesia* and *Theileria* spp. was 7.3–12.2% and 0.7–12.2%, respectively. *In-silico* restriction enzyme (RE) analysis reveals the presence of at least one site for a commercially available RE in 18S rRNA fragments of every parasite, which can differentiate it from its congeners.

*Conclusions* The presented universal oligonucleotide-based PCR assay provides a highly sensitive, specific, cost-effective, and rapid diagnostic tool for the initial screening of piroplasmids infecting domestic and wild animals and is potentially helpful for large-scale epidemiological studies.

**Keywords** Piroplasmids · *Theileria* · *Babesia* · 18S rRNA PCR · Restriction enzyme · Phylogenetic analysis

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1 Department of Veterinary Parasitology, College of Veterinary Science and Animal Husbandry, Kamdhenu University, Junagadh, Gujarat 362001, India
2 Referral Veterinary Diagnostic and Extension Centre, LUVAS, Uchani, Karnal 132001, India
3 Animal Diseases Diagnostic Laboratory, Veterinary Clinical Complex, College of Veterinary Science and Animal Husbandry, Kamdhenu University, Junagadh, Gujarat 362001, India
4 Department of Veterinary Medicine, College of Veterinary Science and Animal Husbandry, Kamdhenu University, Junagadh, Gujarat 362001, India
Introduction

In general, piroplasmosis is a disease condition in vertebrate hosts caused by the parasites belonging to the class Piroplasmea and order Piroplasmoridea and is transmitted by ticks [1–4]. Piroplasms are pear-shaped intraerythrocytic stages. They primarily consist of hemoprotozoan parasites belonging to four related genera: Babesia, Theileria, Cytauxzoon, and Rangelia infecting both wild and domestic animals [1]. A variety of parasite species belonging to these genera are common and widespread blood parasites worldwide, with significant economic, medical, and veterinary consequences [1, 2, 5]. Different piroplasms like B. bigemina, B. gibsoni, B. canis vogeli, T. equi, T. annulata, T. orientalis, T. levenshuni, T. ovis and in some extent B. caballi, B. bovis, and T. equi have been reported among domesticated animals from India [6–14]. Although some are benign or low pathogenic, others can cause a wide range of symptoms and even death. A range of prevalence of the piroplasms in different animals was recorded worldwide [reviewed by 4, 15–21].

Fever, anaemia, malaise, lethargy, and anorexia are common symptoms of piroplasmid infections in animals [1, 2, 5, 8, 22]. However, the pathogenesis and accordingly clinical signs and symptoms vary with parasite species and the animal-associated factors such as species, breed, age, immunological status, concurrent infections with other pathogens, and/or genetic factors. In most cases, clinical signs and symptoms caused by piroplasmid infections are confused with other protozoal, bacterial, or viral infections that can be identified through laboratory testing. On the diagnostic front, microscopic examination of blood smear has been widely used to detect the piroplasm stage of parasites in erythrocytes. However, they have serious limitations like very low sensitivity, the requirement of skilled person, inability to detect the subclinical and carrier animals, and inability to identify the species [16, 23]. Other modern diagnostic techniques like serodiagnostics utilize to detect parasite-specific antibodies have also suffered from some limitations like cross reactivity, inability to discriminate active from past-infection, and cannot be used as a test of cure [24]. However, the antigen-based serodiagnostics can overcome certain limitations but have challenging to identify the suitable antigens and compromised with sensitivity.

Alternatively, with the advancement of molecular biology, PCR-based diagnostics now give researchers and diagnosticians the ability to detect and identify numerous parasites down to the subspecies or strain level in clinical samples and their natural carrier [23]. The method can overcome the significant limitations of both microscopy and serology and has proven to be the most sensitive and specific method for detecting agents and new strains and being an essential tool for evaluating therapeutic efficacy [25, 26].

Various regions of DNA were targeted for molecular identification of Theileria and Babesia parasites, either at the species or genus level that includes 18S rRNA, internal transcribed spacer-1 (ITS-1), Apical membrane antigen-1 (AMA-1), equi merozoite antigen-1 (EMA1), equi merozoite antigen-2 (EMA2), 16S rRNA, major piroplasm surface protein (MPSP), Heat shock protein70 (HSP70), T. annulata merozoite surface antigen-1 (Tams-1), cytochrome b, rhophtry-associated protein-1c (rap-1c), thrombospondin-related anonymous protein (TRAP), spherical body protein-4 (SBP-4) [5, 27–31], etc. 18S rRNA is the most common target for PCR-based detection and identification in eukaryotic species. Piroplasmids have been identified at the genus and species levels using different regions of the 18S rRNA gene [27, 32–35].

At least 111 valid Babesia and 39 valid Theileria species had been described with morphological and biological characteristics. However, many more are still waiting to get species status after introducing molecular biology [5]. Because there are so many different species of Theileria and Babesia, identifying individual parasites at the species level using a single PCR is difficult, especially during epidemiological investigations. Recently, few multiplex PCR assays for the simultaneous and rapid detection of multiple pathogens have been achieved. However, this technique also has its inherent problems like challenges of designing effective primers to avoid the cross-primer dimer formation or false priming amplification, limitations in the size of target amplification (short-length amplification is more effective), and variable amplification efficiency to different targets/templates [36]. Even if these constraints are solved, PCR multiplexing can only be done on a limited number of organisms. So, instead of going straight to species identification, screening the samples for piroplasmids first will be more cost-effective and efficient.

Moreover, the same amplicons can be used for species identification through restriction enzyme (RE) and sequencing analysis. If not, species-specific PCR can be performed solely on piroplasmid-positive samples. Accordingly, the present investigation was designed to optimize and validate the short-length 18S rRNA PCR for rapid screening of animals suspected of piroplasmid infections.

Materials and Methods

Ethical Statement

Approval and necessary guidelines of Institute Animal Ethics Committee (IAEC), College of Veterinary Science and Animal Husbandry, Junagadh, Gujarat, India were obtained and followed during this study (F. No. 25/15/2018-CPCSEA
Sample Collection and Isolation of Whole Genomic DNA

The blood samples collected from different animal species presented at Veterinary Clinical Complex (VCC), Veterinary College, Junagadh were used in the present investigation. About 2–3 ml jugular vein blood was collected from each animal in a vial containing EDTA, and various animal parameters like age, sex, breed, and species were recorded. During 2015–2020, 1250 samples were collected from cattle, buffaloes, horses, dogs, sheep, goats, and wild felids. Randomly, 468 samples were chosen for microscopic examination (Supplementary Table 1). The samples found positive in microscopy were used as the positive control. The samples collected from apparently healthy cow calves maintained in the institute farm were used as the negative control.

The whole blood genomic DNA was isolated from 200 µl of blood sample using GeneJET Whole Blood Genomic DNA Purification Kit (Thermo Scientific, Lithuania) following the manufacturer’s protocol. Finally, DNA was eluted in 200 µl of elution buffer and stored at −20 °C till further use.

Microscopic Examination

Thin blood smear was prepared on the microscopy glass slide (Borosil, India), dried, and fixed by methanol (Merck, India) for 2–3 min. Subsequently, the smear was stained 20 times diluted Giemsa’s stain (Himedia, India) in distilled water for 30 min. The slide was washed over tap water, dried, and observed under 100 × oil emersion objectives of a compound light microscope (Labomed, USA).

Piroplasmid 18S rRNA PCR

The oligonucleotide primers used in the present investigation were previously used to identify the common piroplasmids found in this region (Table 1). Three of the seven pairs of primers were previously published, while the remaining four were designed specifically for this study using protein coding genes like rhoptry-associated Protein 1c (rap-1c) gene for Babesia bigemina (Accession: AY146987), Major Piroplasm Surface Protein (MPSP) gene for Theileria orientalis/buffali (Accession: D11047), erythrocyte/equine merozoite antigen-1 (EMA1) for Theileria equi (Accession: MF447154) and Cytochrome b gene in Theileria lusitaurdi (Accession: MN481239). The alignment of sequences for primers design are presented in Supplementary Figs. 1–4. Standard 25 µl PCR reaction was set up in 200 µl tubes using 12.5 µl of 2x DreamTaq Green PCR master mix (Thermo Scientific, Lithuania), 1 µl each of forward and reverse primers (10 mM), 4 µl (15.8 to 28.5 ng/µl) genomic DNA, and 6.5 µl of nuclease-free water (NFW). The reaction mixture was loaded in a Thermal Cycler (Applied Bio Systems, USA) optimized for amplification of the 18S rRNA sequence and the cycling protocol was programmed as: initial denaturation at 95 °C for 3 min followed by 35 cycle of denaturation (96 °C for 15 s), annealing (60 °C for 20 s), and extension (72 °C for 25 s). The final extension was kept at 72 °C for 1 min at the end of the PCR cycle. Finally, to resolve the amplified products, 10 µl of the PCR product along with DNA ladder were electrophoresed in 1.5% agarose gel containing 0.5 µg/ml of Ethidium bromide (EtBr) in 1× Tris–Acetate–EDTA (TAE) buffer at 120 V for 20 min. Subsequently, amplicons were visualized and documented in the Gel-documentation system (Syngene, UK).

Identification of Parasite Species Using Species-Specific PCR and Sequencing

Randomly, 5–6 piroplasmid-positive samples (confirmed by 18S rRNA PCR) from each animal species such as cattle, buffaloes, horses, dogs, and sheep were chosen for species-specific PCR. In-house standardised species-specific PCR was used to identify the common piroplasmids found in this region (Table 1).
for amplification of targeted gene to the commercial house (Eurofins Genomics India Pvt. Ltd., Bengaluru) for bi-directional Sanger sequencing. Upon receiving of sequences, the quality of sequences was checked in BioEdit programme, both forward and reverse sequences were aligned, and a correct consensus sequence was obtained. Subsequently, the sequence analysis and similarity searches were performed with the basic local alignment search tool available in GenBank data in National Centre for Biotechnology Information, USA, and species were confirmed. The sequences generated from different species of parasites were submitted to GenBank (NCBI, USA), and accession numbers were obtained.

### Sensitivity and Specificity of Piroplasmid 18S rRNA PCR

The analytical sensitivity of 18S rRNA PCR was done on whole blood genomic DNA positive for *T. annulata*. The concentration of DNA was measured by Qubit® dsDNA BR assay kit in Qubit™ 4 Fluorometer (Thermo Scientific, Singapore) as per the manufacturer’s instructions. The sample with 10 ng/µl genomic DNA was serially double-fold diluted in NFW, and 2 µl was used as a template DNA from each dilution in an optimized PCR to amplify 18S rRNA DNA fragments.

The analytical specificity of the 18S rRNA PCR was determined by including the total genomic DNA in the reactions extracted from whole blood of different animals infected with other common haemoparasites viz., *Trypanosoma evansi, Hepatozoon* spp., *Anaplasma* spp., and *Ehrlichia canis* maintained in the laboratory.

### Piroplasmid 18S rRNA Sequence Analysis

Randomly, 18S rRNA PCR positive samples from each species of animals, whether confirmed through species-specific PCR or not, were used to amplify the targeted DNA using BaF and BaR primers in a 50 µl reaction at optimized PCR conditions. The amplified product was purified using GeneJET Gel Extraction Kit (Thermo Scientific, Lithuania), and 10 µl was submitted along with primers used for amplification of targeted gene to the commercial house (Eurofins Genomics India Pvt. Ltd., Bengaluru) for bi-directional Sanger sequencing.

Upon receiving the sequences, the quality of sequences was checked in the BioEdit programme. Both forward and reverse sequences were aligned to achieve the right consensus sequence. Subsequently, sequences were analysed on the BLASTn programme (NCBI, USA) and confirmed their identity. The sequences generated from different species of parasites were submitted to GenBank (NCBI, USA), and accession numbers were obtained.

### Table 1 Details of PCR and primers for species-specific identification of piroplasmids

| Parasites                  | Primers (5′ → 3′) | Amplicon size (bp) | Target DNA | Annealing Temp. (X) | References         |
|----------------------------|-------------------|--------------------|------------|---------------------|--------------------|
| *Theileria annulata*       | Tctb1-accttgccgtaatgttaaact Tctb2-ctctggaccaactgtttgg | 313 | Cytochrome b | 60 ºC               | [37]               |
| *T. buffaloi/orientalis*   | 878TO_F-atgttgccagagaagatc 878TO_R-tegatatagggactcagctgc | 878 | MPSP | 55 ºC               | Present study*     |
| *Babesia bigemina*         | MRCF-cgctggctctatagcag MRCR-cctcccctctggcagaagctattgt | 462 | Rap-1c | 60 ºC               | Present study*     |
| *T. equi*                 | TEEMA1F-5'aagcagctgccagagagcAA3' TEEMA1R-5'tctgggaaggtctgtgtg3' | 595 | EMA1 | 58 ºC               | Present study*     |
| *B. gibsoni*              | BgTRAP1-aagcccaatacagagagc BgTRAP2-tctgtatcgcagcatacgta | 679 | TRAP | 56 ºC               | [11]               |
| *B. canis vogeli*         | BcvF-5'gtgaaccttatcacttaagg3' BcvF-5'caactctccacagcactgg3' | 610 | 18S rRNA | 56 ºC               | [38]               |
| *T. lestoquardi*          | TCLCybF-accttgccgtaatgcttcattcagcata TGAC  TCLCybR-tctggaccaactgttataa | 313 | Cytochrome b | 50 ºC               | Present study*     |

*The new oligonucleotide primers described here were designed from species-specific signature genes, and then selected the region which was highly specific for a particular parasite species based on nucleotide alignment and finally tested in Primer-BLAST (NCBI) where programme was set as Search mode: automatic. Database: nr, Exclusion: none, Organism: kept blank and Primer specific stringency kept default for Primer Pair Specificity Checking Parameters. Further the specificity of these primers was checked based on its reactivity to the samples microscopically positive for infection with targeted parasite species, sequencing and nBLAST analysis (NCBI, USA).
In-Silico Restriction Enzyme Analyses of 18S rRNA DNA Fragments of Piroplasms

The newly generated 18S rRNA piroplasm sequences of bovines, horses, dogs, sheep, and goats (MZ573175, MZ573172, MZ573176, MZ573174, MZ573177, MZ573173, MZ573171, MZ701980, MZ573178) as well as available sequences in GenBank (MN629354, MH257729) were used to identify the unique restriction enzymes (RE) sites (GenScript Restriction Enzyme Map Analysis Tools, https://www.genscript.com/tools/restriction-enzyme-map-analysis). The group of parasites affecting an animal species was further analyzed to identify the unique RE enzymes for easy identification and differentiation.

Results

Piroplasmid 18S rRNA PCR and Microscopy

Primer_BLAST analysis revealed reactivity of published primers (BaF/BaR) to wider range of piroplasms. However, primers showed matching with the sequences of few species of Cryptosporidium and Sarcocystis, and Besnoitia besnoiti under the apicomplexan group of mammals. Alignment of 18S rRNA sequences of piroplasmid with other apicomplexan parasites revealed that the sequences of reverse primer region is conserved but forward primer region of other apicomplexan parasites varied by only a few nucleotides. (Supplementary Fig. 5). The PCR assay successfully amplified the expected size of targeted DNA from blood samples found positive for various piroplasms collected from different animals (Fig. 1) that was confirmed by sequencing and BLASTn analysis. The amplicon size of Babesia spp. was somewhat smaller than Theileria spp., especially B. bigemina, compared to T. annulata. Among the 1250 samples collected, 63.3% of animals were detected positive for piroplasmids infections through 18S rRNA PCR. In contrast, 23.3% of animals were found positive by microscopy in 468 samples examined (Supplementary Table 1). The time required for PCR reaction was merely 58 min.

Species Identification Through Species-Specific PCR

The three steps confirmation of species have been done first, by the use of species-specific PCR primers; second, by sequencing and NCBI–nBLAST analysis and third, the primers were reacted on microscopically and piroplasmid 18S rRNA PCR positive samples. Species-specific PCR assays successfully detected the common parasites available in this region like B. bigemina, T. annulata, T. buffali/orientalis, T. equi, B. canis vogeli, B. gibsoni, and T. lestoquardi (Supplementary Fig. 6). To further confirm the species of parasites, sequences were generated, submitted to GenBank, and accession numbers were obtained (MZ665956, MZ665960, MZ720762, MZ665961, MZ646048, MZ665957, MZ665959).

Sensitivity and Specificity of Piroplasmid 18S rRNA PCR

The PCR was able to amplify the 18S rRNA gene fragment of T. annulata from as low as 39 picograms (pg) of whole blood genomic DNA microscopically positive for the same parasites. On the other hand, no amplification of such gene fragments was recorded from common but unrelated haemoparasites viz., Trypanosoma evansi, Hepatozoon spp., Anaplasma spp., and Ehrlichia canis positive samples (Fig. 2).

Sequence Analysis of 18S rRNA Gene Fragments of Piroplasms

The sequence analysis revealed two different size ranges of 18S rRNA gene fragments. The Babesia spp. sequences were nearly 400 bp (393–408 bp), whereas; Theileria spp. were more than 400 bp (418–424 bp). BLASTn analysis revealed more than 99% identity with distinct parasite species of Theileria and Babesia. The resultant consensus sequences were submitted to GenBank (National Center for Biotechnology Information, USA), and accessions were acquired as B. bigemina (MZ573175), T. annulata (MZ573172), T. buffali/orientalis (MZ573176), T. equi (MZ573174), B. canis vogeli (MZ573177), B. gibsoni (MZ573173), T. luwenshuni (MZ701916). After blast
analysis, the sequences obtained from 18S rRNA gene fragments and species-specific PCR amplicons indicated the same parasite species. The sequence divergence among the different species of *Babesia* and *Theileria* were recorded as 7.3–12.2% and 0.7–12.2%, respectively (Fig. 3A). Sequence alignment report revealed that the forward and reverse primer regions of the 18S rRNA gene of various piroplasmids are highly conserved. In phylogenetic analysis, sequences of all members of Theileriade and Babesidae formed two distinct clades. *T. annulata* and *T. lestoquardi* form a close group. Similarly, *T. orientalis* and *T. luwenshuni* are very close to each other. *T. ovis* is very much closer to *T. annulata* compared to *T. orientalis*. Similarly, in the Babesidae group, *B. gibsoni* is closer to *B. canis vogeli* than *B. bigemina* but farther away from *Babesia* sp. Leopard. *Toxoplasma gondii*, an apicomplexan parasite with a unique lineage, forms a separate clad along with Theileriade group while *Plasmodium vivax* showed almost equal distance to both Theileriade and Babesidae groups and form a separate clad (Fig. 3B).

**In-Silico RE Analysis of 18S rRNA Gene Fragments of Piroplasmids**

Based on *in-silico* analysis of 18S rRNA gene fragments of various piroplasmid, few unique RE sites were identified for an individual parasite of an animal species. None of the RE sites for *T. luwenshuni* of sheep and goat was recorded those were available commercially, which can differentiate it from *T. ovis* and/or *T. lestoquardi*. Otherwise, each parasite has at least one commercially available unique RE that distinguishes it from its congener in the same animal species (Table 2).

**Discussion**

*Babesia* and *Theileria* spp. are apicomplexan parasites transmitted by ticks. Traditional morphological and contemporary molecular studies substantiated these genera as close relatives and, correspondingly, constitute the order Piroplasmida [39]. Piroplasmid infections are detrimental to the health of various wild and domestic animals. They are a major cause of economic losses in the animal husbandry industry in tropical and subtropical countries worldwide [40].

Animals suffering from acute babesiosis or theileriosis can have various symptoms such as fever, haemolytic anaemia, haemoglobinuria, oculo-nasal discharge, anaemia, malaise, lethargy, increased heart rate, increased respiratory rate, and even death in severe case. Although these symptoms are very typical, they are not pathognomonic, and animals with chronic infections can be asymptomatic carriers [22, 41]. Carrier animals with no clinical symptoms are thought to be a key reservoir of infection for ticks that can spread the infection to other animals.

The laboratory diagnosis of piroplasmosis was based on the microscopic examination of Giemsa-stained blood smears to detect piroplasmid inclusions in erythrocytes. However, species identification by microscopy is challenging because different parasites have similar morphologies, making identification even more difficult if mixed infections occur. Furthermore, identifying parasites in carrier animals with low parasite counts and in acute instances at the onset of the disease might be difficult. Additionally, it needs special diagnostic knowledge [16]. Some serological assays, such as the complement fixation test (CFT) and the indirect fluorescent antibody test (IFAT), can help to
detect prior infections, however, there have been reports of cross-reactions between species [42, 43]. Furthermore, these tests can produce erroneous positive and negative results. Molecular techniques, such as polymerase chain reaction (PCR) and its variants, have shown to be the most sensitive and specific tools for diagnosis and study in recent decades and have been widely employed for detection and discrimination between *Theileria* and *Babesia* species [44–48]. Although uniplex PCR assays are designed to detect single species at a time, they can be time-consuming and expensive when applied to many samples co-infected with more than one pathogen species. On the contrary, multiplex PCR (mPCR) shows unparalleled advantages, including detecting multiple pathogens in a single reaction and saving time [49]. However, this technique also has inherent problems like designing effective primer, limitation in the target size amplification, and variable amplification efficiency [36]. Reverse line blotting method had also been used for the simultaneous detection and differentiation of *Babesia* and *Theileria* spp. infecting ruminants [45, 50, 51]. However, Reverse line blot (RLB) assay requires expertise, specialized equipment, and the protocol is very labor-intensive and due to its high cost, this technique is not feasible [52]. None of these approaches can be deemed superior to the others. There is a need to develop efficient diagnostic strategies to deal with this problem in resource-poor countries and developed countries facing piroplasmosis as an extensive burden. So, screening the samples for piroplasms first will be more cost-effective and efficient than proceeding straight to species identification. The present study focused on standardisation and validation of a very convenient and universal PCR assay that can detect piroplasms infective to domestic and wild animals, which can be used for diagnosis, quarantine, and epidemiological survey. Accordingly, the current study

Fig. 3 Analysis of nucleotides sequences of 18S rRNA gene fragments of various piroplasms of animals generated in the present study. A sequence identity/divergence analysis and B sequence phylogenetic analysis using MEGA X software (statistical Method: Maximum-likelihood; model: Tamura 3-parameter model+Gamma distribution (+G); bootstrap value:500; Neighbor-Join and BioNJ algorithms)
was planned to optimize and validate the short-length 18S rRNA PCR for rapid screening of animals suspected of piroplasm infections. The eukaryotic 18S rRNA gene has both conserved and variable regions. Due to its high specificity and sequence conservation, it has been used as a universal biomarker to screen closely related species and biodiversity studies \[7, 53–55\]. Universal oligonucleotide primers based on 18S rRNA has also been used for initial screening of piroplasmids infecting horses and Bactrian camels north eastern Mongolia \[56\].

Most of the earlier studies on 18S rRNA gene screening commonly prevalent \textit{Babesia/Thelileria} spp. were limited to their related host species \[32\]. The novelty of the current investigation is to standardize and optimize 18S rRNA PCR for the initial screening of common piroplasmids in different animals like cattle, buffaloes, horses, dogs, and sheep were chosen at random for species-specific PCR. Species-specific PCR assays successfully detected the common parasites available in this region like \textit{B. bigemina}, \textit{T. annulata}, \textit{T. buffaloi/orientalis}, \textit{T. equi}, \textit{B. canis vogeli}, \textit{B. gibsoni}, and \textit{T. lestoquardi} further confirmed through sequence analysis.

Sequence analysis of 18S rRNA gene fragments revealed two different size ranges of amplicons. The \textit{Babesia} spp. sequences were nearly 400 bp (393–408 bp), whereas \textit{Theileria} spp. were more than 400 bp (418–424 bp). After blast analysis, the sequences obtained from 18S rRNA gene fragments and species-specific PCR amplicons revealed the same parasite species. Phylogenetic analysis and sequence distance report further confirm the current findings. Similar observations based on molecular phylogenetic tree of 18S rRNA gene sequences were reported by previous researchers among different \textit{Theileria} and \textit{Babesia} spp. infecting domestic animals \[55\]. These studies further reveal that neither \textit{Theileria} nor \textit{Babesia} are monophyletic taxonomic units and systematic re-examination is required to determine the generic diversity of the piroplasmids \[54, 59–61\].

| Animals        | Piroplasms            | Amplicon size | Unique RE cuts/position |
|----------------|-----------------------|---------------|-------------------------|
| Cattle & Buffaloes | \textit{B. bigemina}_{MZ573175} | 393 bp        | Ban I 120/ EcoRII 215/ Mval 217/ ScrF1/217 |
|                | \textit{T. annulata}_{MZ573172} | 419 bp        | NspI 235/              |
|                | \textit{T. orientalis}_{MZ573176} | 418 bp        | AcelII 262/ HpalI 263/ MnlI 96/ MspI 263/ SmaI 209/ |
|                | \textit{B. bovis}_{MH257729*} | 378 bp        | Ava I 94/ HinfI 194/   |
| Dogs           | \textit{B. gibsoni}_{MZ573173} | 405 bp        | HinfI 210/            |
|                | \textit{B. vogeli}_{MZ573177} | 408 bp        | BstEII 77/ BstXI 224/  |
| Horses         | \textit{T. equi}_{MZ573174} | 421 bp        | Ddel 243/ Drl 82/ MboI 80/ |
|                | \textit{B. caballi}_{MN629354*} | 396 bp        | MnlI 98/              |
| Sheep and Goats| \textit{T. ovis}_{MZ573178} | 424 bp        | HinfI 224/            |
|                | \textit{T. luwenshuni}_{MZ573171} | 422 bp        | –                      |
|                | \textit{T. lestoquardi}_{MZ701980} | 421 bp        | Ncil 223/ NspI 237/ ScrF1 223/ |

*Previously published GenBank sequences
several piroplasm species [62]. The PCR assay standardised in the present study was able to amplify the 18S rRNA gene fragment of *T. annulata* from as low as 39 pico grams (pg) of whole blood genomic DNA microscopically positive for the same parasites. On the other hand, no amplification of such gene fragments was recorded from common but unrelated haemoparasites. However, apart from piroplasms, primer-BLAST research revealed plausible primer reactivity to a few species of *Cryptosporidium*, *Sarcocystis*, and *Besnoitia besnoiti* in mammals. Yet, the likelihood of these parasites in circulation is remote. These results support the reliability of 18S RNA-specific oligonucleotides used in the present study as universal primers for screening piroplasms infecting livestock. The current findings are consistent with the previously published report [7, 56]. Sequence alignment report revealed that both the forward and reverse primers region of the 18S rRNA gene of various piroplasms are highly conserved.

Moreover, BLASTn analysis, further demonstrated that the sequences obtained from 18S rRNA gene fragments and species-specific PCR amplicons indicated the same piroplasm. In-silico RE analysis of 18S rRNA gene fragments revealed that each parasite has at least one commercially available unique RE that distinguishes it from its congener in the same animal species. These findings support the hypothesis that 18S rRNA-based PCR assay is a highly specific, sensitive, cost-effective, and rapid diagnostic tool for initial screening of piroplasms infecting domestic and wild animals and can be helpful for large-scale epidemiological studies.

**Conclusion**

This study standardized and optimized an 18S rRNA PCR assay to detect common piroplasms of different animals like cattle, buffaloes, sheep, goats, dogs, horses, and leopards. The presented universal oligonucleotide-based PCR assay provides a highly sensitive, cost-effective, and rapid diagnostic tool for the initial screening of piroplasms infecting domestic and wild animals and is potentially helpful for large-scale epidemiological studies.

**Supplementary Information** The online version contains supplementary material available at https://doi.org/10.1007/s11686-022-00625-2.

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**Declarations**

**Conflict of Interest** On behalf of all authors, the corresponding author states that there is no conflict of interest.

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