Molecular and microscopic detection of *Theileria equi* and *Babesia caballi* in horses in Kurdistan Province, Iran

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ABSTRACT
Equine piroplasmosis is a tick-borne disease caused by intra-erythrocyte protozoa, *Theileria equi* and *Babesia caballi*. The present study aimed to detect piroplasm infection in horses in Kurdistan Province, Iran, through molecular and microscopic approaches. In this study, 186 blood samples were randomly collected from horses of five regions of Kurdistan Province. The Tbs-S/Tbs-A primer set was used for amplification of *Theileria equi* and *Babesia caballi* DNA through polymerase chain reaction. Blood smears of each case were also examined by Giemsa staining method. During microscopic examination of Giemsa-stained blood smears, 3 out of 186 (1.61%) blood samples were positive for piroplasm infection. The product of only one blood sample yielded 426-430 bp-sized fragments on a 1.5% agarose gel electrophoresis, and BLAST analysis of the sequenced sample indicated a 100% similarity with *T. equi* 18S rRNA gene sequences in GenBank. The results indicated that one out of 186 blood sample was positive (0.54%) for *Theileria equi* and none of them was positive for *Babesia caballi*.

Keywords: *Theileria equi*, *Babesia caballi*, PCR, Horse, Iran

Détective moléculaire et microscopique de *Theileria equi* et *Babesia caballi* chez les chevaux de la province du Kurdistan, Iran

Résumé: La piroplasmosis équine est une maladie transmise par les tiques et causée par les protozoaires intra-érythrocytaires *Theileria equi* et *Babesia caballi*. L’objectif de cette étude était de détecter les piroplasmoses affectant les chevaux de la province du Kurdistan iranien par une approche moléculaire et à l’aide d’analyses microscopiques. A cet effet, 180 échantillons sanguins ont été prélevés de façon aléatoire à partir de chevaux originaires de cinq régions de la province du Kurdistan. La série d’amorce Tbs-S/Tbs-A a été utilisée pour l’amplification en chaine par polymérase de l’ADN de *T. equi* et *B. caballi*. De plus, des frottis de chaque échantillon sanguin ont été analysés après coloration au Giemsa. Parmi les 186 frottis examinés au microscope, 3 étaient atteints d’une piroplasmosis (1.61%). Les échantillons positifs ont été ensuite soumis à une analyse électrophorétique sur gel d’agarose (1.5%). Une recherche BLAST sur les séquences des fragments obtenus a révélé 100% de similitude entre un fragment d’une taille de 426-430 bp issu de l’un des échantillons avec la séquence ARNr 18S de *T. equi* répertoriée dans GenBank. Ces résultats montrent que parmi les 186 échantillons sanguins analysés, un seul était positif au *Theileria equi* alors qu’aucune contamination au *Babesia caballi* n’a été détectée.

Mots clés: *Theileria equi*, *Babesia caballi*, PCR, Cheval, Iran
INTRODUCTION

Equine piroplasmosis, caused by *Theileria equi* (recently re-classified from *Babesia equi* (Mehlhorn and Schein, 1998) and *Babesia caballi*, is known as an important protozoan infection from the veterinary and economic viewpoints (Schein, 1988). Equine piroplasmosis, which occurs throughout the tropical and subtropical areas of the world, imposes great economic losses to endemic countries (Friedhoff et al., 1990; de Waal, 1992). Tick species of the genera *Boophilus*, *Dermacentor*, *Hyalomma*, and *Rhipicephalus* are the biological vectors of equine piroplasmosis (Battsetseg et al., 2001). Equine piroplasmosis is characterized by fever, anemia, dyspnea, icterus, hepato- and hemorrhages in mucosal surfaces, hemoglobinuria, and even death (Schein, 1988; Uilenberg, 2006). Infection caused by *T. equi* may be suppressed by anti-theilerial drugs, but only some of them can eliminate infection. After recovery from theileriosis, horses may remain carriers for years or for life without further clinical diseases; these animals can act as a source of infection to tick vectors and naïve equids (de Waal, 1992; Brüning, 1996). The Giemsa staining of the blood smears is the common method for the identification of piroplasm infection, but this method has low sensitivity and provides some false positive or false negative results (Shayan and Rahbari, 2005). In addition, serological procedures are usually employed in determining subclinical infections. False positive and false negative results are also commonly observed in serological tests due to cross-reactions or sometimes inadequate antibody level in carriers due to long-term infection (Gubbels et al., 2000). The previous studies performed in Iran on equine piroplasmosis are mostly in the form of case reports (Nouri and Lotpholahzadeh, 1993; Aslani, 2000; Seifi et al., 2000; Sakha, 2007; Erfêi Akhole et al., 2013). Recently, serological and molecular methods have been applied for diagnosis of equine piroplasmosis (Seifi et al., 2000; Malekifard et al., 2011; Abedi et al., 2013; Bahrami et al., 2014). The polymerase chain reaction (PCR) method has more sensitivity and specificity in comparison with the serological tests. The PCR method can detect parasitic infections with or without clinical signs (D’oliveira et al., 1995; Almeria et al., 2001; Dumanli et al., 2005). Herein, we aimed to estimate the prevalence of equine piroplasmosis caused by *Theileria equi* and *Babesia caballi* in Kurdestan Province, Iran.

MATERIALS AND METHODS

Study area and sampling. Kurdestan Province, a mountainous region with a temperate climate, is located in the west of Iran. The study was conducted during May-September (the tick activity season) 2012 and 2013, in five areas (north, south, east, west, and center) of Kurdestan Province. Regardless of age and gender, a total of 186 horses were randomly selected from each area. Blood samples were collected from the jugular vein of each horse into sterile vacuum tubes with anticoagulant agent (EDTA). Additionally, two thin blood smears were prepared from each sample, and immediately 500 µl of each collected blood sample was mixed with 1 ml ethanol in 1.5 ml sterile eppendorf tube; the specimens were transferred to the Laboratory of Parasitological of the School of Veterinary Medicine, Azad University of Sanandaj, Iran. The blood samples were stored at -20 °C for further analysis. Data of the sampled animals (e.g., gender, age, and area) were recorded during sample collection.

Microscopic examination. The fixed blood smears were stained with Geimsa method in order to determine the presence of haemoprotozoal parasites. The stained smears were microscopically examined to identify haemoprotozoal parasites.

DNA extraction and PCR. For detection of *Theileria* spp. and *Babesia* spp, primers derived from 18S rRNA were used (Shayan, P. and S. Rahbari 2005). Tbs-S/Tbs-A primer set was employed for PCR amplification of 18S rRNA of *Theileria* spp. and *Babesia* spp. Weights of the amplified sequences by this primer set for *Theileria* spp. and *Babesia* spp. were 426-430 bp and 389-402 bp, respectively (Shayan and Rahbari, 2005). Distilled water was used as negative control in each PCR reaction. The positive control was
prepared from the School of Veterinary of Tehran University and Ferdowsi University of Mashhad. The positive sample was sent to Bioneer Laboratory in South Korea for sequencing (3-49, Munpeong-Dong, Daedok-Gu, Daejeon 220-306, Korea), and the result was assessed by BLAST analysis in National Center for Biotechnology Information (NCBI).

RESULTS AND DISCUSSION

Out of 186 examined horses, 62 were female and 124 male; all the horses were aged more than 1 year. During microscopic examination of Giemsa-stained blood smears, 3 out of 186 (1.61%) blood samples were positive for *Theileria equi*. The genomic DNA from blood samples was amplified with Tbs-S/Tbs-A primer. The product of only one blood sample yielded 426-430 bp-sized fragments on a 1.5% agarose gel electrophoresis (Figure 1), and BLAST analysis of the sequenced sample indicated a 100% similarity with *T. equi* 18S rRNA gene sequences in GenBank (T. equi isolate Sample2, KJ573370.1, Brazil; T. equi genotype A, KC465785.1, Tunisia). The sequenced sample was in alignment with *T. equi* 18S rRNA gene sequences in PubMed including numbers 23542456 (Tunisia), 23399005 (US), 19019541 (South Africa), and 12719133 (Southern Europe). Therefore, the result indicated that one out of 186 blood sample was positive (0.54%) for *Theileria equi* and none of them was positive for *Babesia caballi*. The previous studies on equine piroplasmosis conducted in Iran are mostly case reports (Nouri and Lotpholahzadeh, 1993; Aslani, 2000; Seifi et al., 2000; Sakha, 2007; Erfei Akhole et al., 2013). In this study, the result of microscopical observation showed an infection rate of 1.61%, while the obtained result by PCR was 0.54%, and BLAST analysis of the sequenced sample indicated a 100% similarity with *T. equi* 18S rRNA gene sequences in GenBank. Recently, serological and molecular methods have been applied to diagnose equine piroplasmosis in Iran. Abedi et al. (2014) reported that 45% of horses from North Khorasan Province are infected with *T. equi*. On the other hand, Malekifard et al. (2011) reported that 10.83% of horses are infected with *T. equi* in the suburbs of Urmia in the northwest of Iran. Similar studies conducted in Turkey (neighboring northwest of Iran) described the high prevalence of equine piroplasmosis with *T. equi* (Akkana et al., 2003; Oncel et al., 2007; Acici et al., 2008).

Figure 1. Agarose gel electrophoresis: analysis of polymerase chain reaction amplification for detection of *Theileria sp.* by using Tbs-S/Tbs-A primer set derived from the 18S rRNA encoding gene M:100 bp DNA marker; NC: Negative control; PC: Positive control for *T. equi* (426-430 bp) S: Sample 116

These differences between Urmia and Kurdistan might be associated with the restriction that has been implemented to control the animals' movement between the border of Iraq and Iran. It seems that the entry of horses across the border of Turkey and Iran can play a major role in causing higher babesial infection in those regions. In addition, Kurdistan is located in a mountainous area, where climatic factors may influence tick population; accordingly, the lower prevalence rate of infection compared to previous result can be acceptable. Bahrami et al. (2014) examined 105 blood samples from healthy horses from different stables in Yazd Province, Iran. They demonstrated that the prevalence of *T. equi* infection by Giemsa staining method was 4.76%, while by PCR assay this rate was 22.86% (Bahrami et al., 2014). Malekifard et al. (2011) reported that the positive infection rates with *T. equi*
and *B. caballi* by Giemsa staining method were 6.25% and 2.80%, respectively, while 10.83% and 5.83% of cases were infected with *T. equi* and *B. caballi*, respectively, by PCR (Malekifard et al., 2011). Abedi et al. (2014) examined 100 blood samples of horses from Turkoman breed only by microscopy and indirect immunofluorescent antibody test. They reported that the prevalence rate of *T. equi* infection by Giemsa staining method was 5%, while by indirect immunofluorescent antibody, this rate was 48% (Abedi et al., 2013; Abedi et al., 2014) and by PCR was 45% (Abedi et al., 2014). In conclusion, this study showed that the prevalence of *Theileria equi* is low in the horses of Kurdestan Province and there was no infection with *Babesia caballi*.

**Ethics**

We hereby declare all ethical standards have been respected in preparation of the submitted article.

**Conflict of Interest**

The authors declare that they have no conflict of interest.

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