Molecular detection and occurrence of equine theileriosis in Arabian horses in Al-Najaf province/Iraq

Detecção molecular e prevalência de teileriose equina em cavalos árabes na província de Al-Najaf/Iraque

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ABSTRACT
This study was designed to detect equine piroplasmosis using the molecular technique in Al-Najaf province during the season that showed an increment in tick activities. Blood samples were collected from 110 horses with more than two signs of piroplasmosis. After DNA extraction, the product was examined by a polymerase chain reaction to amplify 18SrRNA. The results showed that the overall percentage of equine theileriosis was 38.18%. According to gender, the percentage of infection was 43.48% and 29.27% in females and males, respectively. Significant variations appeared between infected horses according to age, and the percentage of infection was 50% and 35.22% in less than 2 years and more than 2 years age, respectively. Moreover, the percentage of infection was 62.5% and 19.35% in animals with and without acariasis, respectively. Significant variations were also seen in equine theileriosis according to geographical areas, and the higher percentage was reported in Hera district (60.87%), while the lowest percentage was in the center of Al-Najaf (21.43%). This difference may be due to different distribution of vector of disease (tick), which may be the availability of the suitable weather that helped in the multiplication of the intermediate vectors. In conclusion, this study proved the variations in the occurrences of equine piroplasmosis according to gender, age, and geographical areas.

Keywords: Equine theileriosis. PCR. Al-Najaf horses. Piroplasmosis.

RESUMO
Este estudo foi desenvolvido para detectar piroplasmo equina usando a técnica molecular na província de Al-Najaf durante o período do ano com maior ocorrência de carrapatos. Foram coletadas amostras de sangue de 110 cavalos que apresentaram mais de dois sinais de piroplasmo. Após a extração do DNA, o produto foi examinado por reação em cadeia da polimerase para amplificar o 18SrRNA. Os resultados mostraram que a porcentagem geral de teileriose equina foi de 38, 18%. De acordo com o sexo, o percentual de infeção foi 43,48% e 29,27% no sexo feminino e masculino, respectivamente. Apareceram variações significativas entre os cavalos infectados de acordo com a idade, e a porcentagem de infeção foi 50% e 35,22% em menos de 2 anos e mais de 2 anos, respectivamente. Além disso, as porcentagens de infeção foram 62, 5% e 19, 35% em animais com e sem acariasis, respectivamente. Também foram observadas variações significativas na teileriose dos equídeos, de acordo com as áreas geográficas, e o maior percentual foi relatado no distrito de Hera (60, 87%), enquanto o menor percentual foi no centro de Al-Najaf (21,43%). Essa diferença pode ser devido à distribuição diferente do vetor da doença (carrapato), que pode ser a disponibilidade do clima adequado que ajuda na multiplicação dos vetores intermediários. Em conclusão, este estudo provou as variações nas ocorrências de piroplasmose equina de acordo com sexo, idade e áreas geográficas.

Palavras-chave: Teileriose equina. PCR. Cavalos de Al-Najaf. Piroplasmosis.

Introduction
Equine theileriosis caused by apicomplexan Theileria equi is a disease of economic importance facing the equine population in endemic areas (Kumar et al., 2007). According to the occurrence of biological vectors (21 tick species of genera Rhipicephalus, Dermatocentor, Hyalomma, and Boophilus), equine theileriosis is distributed in Asia, Africa, Europe and South America (Schein, 1988; Wise et al., 2013). The disease is manifested by clinical and subclinical forms, the clinical form characterized by hemolytic anemia, in addition to systemic signs including fever, anorexia, jaundice, and dyspnea, edema of limbs, colic, hemoglobinuria and death (Butler et al., 2012; Waal, 1992; Zobba et al., 2008). However, subclinically infected horses have represented a reservoir infecting vectors (Friedhoff et al., 1990).

Acute equine theileriosis is easily diagnosed by microscopic examination of blood smears (Mahdy et al., 2016), while in the chronic or carrier stage, the serological technique is required for diagnosis. Lack of sensitivity is one of the disadvantages of complement fixation test (CFT) and indirect immunofluorescent antibody tests (IFAT) (Khalil, 2012). Conversely, the cross-reactivity with Babesia caballi is the disadvantage of the enzyme-linked immunosorbent assay (ELISA) (Mahmoud et al., 2015). Recently, polymerase chain reaction (PCR) was able to detect the level of parasitemia at 0.0083% (Ibrahim et al., 2011). A review of the literature revealed a scarcity of publication regarding diagnosis and occurrence of equine theileriosis in Iraq. Therefore, this study intended to establish the occurrence of equine theileriosis in Al-Najaf province using molecular tools.

Materials and Methods

Samples collections
This study was carried out in 8 districts of Al-Najaf province during tick activity season. Blood samples were collected from 110 horses with more than two signs (mostly fever and jaundice) of piroplasmosis. Collected blood was immediately placed in tubes with anticoagulant, then submitted to the laboratory and stored in -20 °C until extraction of DNA.

Genomic DNA extraction
The DNA was extracted from stored blood Mini Kit (Geneaid Biotech Ltd., Taiwan). The process was carried out according to the instructions of company by using protocol of frozen blood extraction with the Proteinase K. The extracted gDNA product was then checked by Nanodrop spectrophotometer and stored at -20°C at refrigerator until amplification.

Polymerase Chain Reaction (PCR)
PCR assay was conducted by a specific primer, designed by Alhassan et al. (2005), to amplify a 392bp portion of highly conserved regions of 18S ribosomal RNA in Theileria equi 18SrRNA. The forward primer (TCGAAAGAGTACGATAACCGTG), and reverse primer (TGCCCTAAACTCTTTGCGAT) 18SrRNA were provided by Bioneer Corporation (Republic of Korea). The PCR master mix is prepared by AccuPower® PCR PreMix kit manufactured by Bioneer. The tubes of PCR premix comprise pellets of freeze-dried Taq DNA polymerase 1U, Tris–HCl (pH 9.0) 10mM, dNTPs 250µM, MgCl2 1.5mM, KCl 30mM, stabilizer, and tracking dye. The PCR reaction of master mix is prepared in total volume of 20µl by added purified genomic DNA (5µl), forward primer (1µl of 10pmole) and reverse primer (1µl of 10pmole). The PCR premix tube was completed with 20µl deionizer PCR water, then briefly mixed by vortex centrifuge (Bioneer). The thermocycler (T100™ Thermal Cycler, Bio-Rad, Hercules, CA USA) was set as; temperature of initial denaturation at 95 °C for 5 min; followed by denaturation 30 cycles at 95 °C for 30 sec, annealing 60 °C for 30 sec, and extension 72 °C for 30 sec and, finally, extension at 72 °C for 7 min. The products of PCR were inspected by 1% agarose gel electrophoresis, stained with ethidium bromide, and examined under UV illumination.

Results
Out of 110 samples, 42 showed a band of approximately 392pb, which were considered positive to T. equi 18S rRNA gene in a percentage of 38.18% (Figure 1). According to gender, the percentages of infection were 43.48% and 29.27% in females and males, respectively. The percentages of infection according to the age of horses were 50% and 35.22% in less than 2 years and above 2 years, respectively. The percentages of infection in the horses according to tick infestation...
were 62.5% and 19.35% in horses with acariasis and without acariasis, respectively. There was significant geographical variation in equine theileriosis, with the highest occurrence in Hera district (60.87%), while the lowest was in the center of Al-Najaf (21.43%), as presented in Table 1.

### Discussion

Equine piroplasmosis has economic importance in the equestrian industry due to low performance in work and sports activities, restriction of transboundary movement, cost of hospitalization and high mortality rate of infected animals (Bahrami et al., 2014; Friedhoff et al., 1990; Malekifard et al., 2014). The current study proved the detection of *Theileria equi* of the 18S rRNA gene by PCR in 42 out of 110 samples, with percentages reaching 38.18% in horses suffering from main disease signs (pyrexia and icterus). This is close to the results of Aziz et al. (2019), who recorded the rate of occurrence of equine theileriosis diagnosed by PCR (41.91%) in Erbil, Iraq, as well as the results being somewhat consistent with those of Saleem & Al-Samarai (2018), who recorded 32% in central Iraq. In the neighboring countries of Iraq Saudi Arabia (Alanazi et al., 2012), Jordan (Abutarbush et al., 2012), Iran (Abedi et al., 2014) and Turkey (Guven et al., 2017), the percentages were 10.14%, 14.6%, 48%, and 8.8%, respectively.

The diagnosis of equine theileriosis was a challenge for veterinary field practice as a microscopic examination of the suspected blood smear is unreliable, particularly in the low parasitic stage. Many serological tests have been used for the diagnosis of equine theileriosis, including ELISA, IFAT, and CFT. However, these techniques have low sensitivity and cross-reactivity with other blood protozoa (Abedi et al., 2014). Rampersad et al. (2003) found high accuracy of the molecular identification of *T. equi*, as the parasite can be diagnosed by PCR (18S ribosomal RNA genes) in 25 µl blood with the parasitemia percentage of 0.00008%.

The result of the current study showed gender variation in the occurrence of equine theileriosis. The female's percentage was 43.48%, which was higher than the male's percentage of 29.27%. These findings were similar to that concluded by Sray et al. (2019) and might be explained by the high exposure of mares to stress than males due to pregnancy or racing, whereas most local racing and religious rituals depend on mares. Many authors suggested that horses' hard exercise may compromise their immunity, resulting in the
development of acute theileriosis in animals with carrier state (Hodgson, 2002).

The results of the present study demonstrated that the diseases were 50% more prevalent in immature horses. This finding may be due to placental transportation or bad management regime, whereas the foals aged less than 2 years were highly infested with ticks due to tick control neglect. This result agrees with many previous recorded studies that showed the occurrence of equine theileriosis in foals more than adult horses due to the exposure of foals to stress, which may lead to reduced immunity (Hodgson, 2002; Sevinc et al., 2008). This study showed variation in the percentage of equine theileriosis infection due to acarasis infestation. The high occurrence of disease (62.5%) occurred in animals with tick infestation in comparison to tick-free animals. This result is compatible with previous studies that also documented many species of ticks as the biological vectors to equine theileriosis (Alanazi et al., 2012; Nava et al., 2009).

The result of this study also revealed variations in the occurrence of the disease according to a geographical area. The highest occurrence was recorded in rural district of Hera (21.43%), while the lowest was in center of Al-Najaf (urban district). This difference may be due to different distribution of vector of disease (tick), which may be the availability of the suitable weather that helped in the multiplication of the intermediate vectors (Javed et al., 2014).

Conclusion

This study proved the detection of Theileria equi of the 18S rRNA gene by PCR in Arabian horses in Al-Najaf province/Iraq. Moreover, the variations in the occurrences of equine piroplasmosis according to gender, age, and geographical areas were also reported.

Conflict of Interest

The authors declare that there is no conflict of interest.

Ethics Statement

Ethics approval was approved by Faculty of Veterinary Medicine/University of Kufa.

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