Microscopic and Molecular Detection of *Theileria annulata* Infection of Cattle in Egypt

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**Abstract** | **Background and objective:** Bovine theileriosis is a global tick-borne disease and it has emerged as one of the major constraints affecting livestock production in Egypt. The disease is caused by the hemoparasite, *Theileria annulata*, which is transmitted by the tick vector, *Hyalomma* species. **Materials and methods:** A total of 376 Egyptian female Balady-Friesian cross cattle were randomly selected in El-Wadi El-Gadid province, Egypt and were examined using microscopic examination of Giemsa-stained thin blood smears and PCR-dependant molecular identification using a *Theileria annulata*-specific primer pair, *Cytob1*, targeting the amplicon of 312 bp of the mitochondrial cytochrome b gene. **Results:** It has been found that 102 (27.13%) animals had clinical symptoms of piroplasmosis, predominantly enlargement of superficial lymph nodes and a sustainable high fever (40-42°C), anorexia, loss of generalized conditions, lacrimation, corneal affections, oculo-nasal discharge, dyspnea and diarrhea. Light microscopy of Giemsa-stained thin blood smears revealed that 35 (9.31%) of examined animals showed the presence of intra-lymphocytic and/or intra-erythrocytic stages of hemoparasites. Various stages including macroschizonts, microschizonts and merozoites were demonstrated in thin blood smears. Application of PCR elucidated that 11.44% (43/376) animals had theileriosis indicated by the appearance of specific diagnostic bands. Most of clinically-positive animals and all those positive by the use of microscopic examination and PCR are associated with the existence of ticks vector. **Conclusion:** The present work revealed the prevalence of bovine theileriosis in El-Wadi El-Gadid, Egypt using microscopic examination of Giemsa-stained thin blood smears (9.31%) and PCR (11.44%). To reduce economic losses induced in cattle due to theileriosis, a considerable attention and veterinary care are mandatory to be done by authorities and veterinarians through elimination of the tick vectors and frequent treatment of infected animals.

**Keywords** | Cattle, *Theileria annulata*, *Hyalomma* spp., PCR, Egypt.

**INTRODUCTION**

Livestock, particularly cattle and buffaloes as well as small ruminants, are major resources for the production of milk and meat through a wide belt of tropical and sub-tropical areas (Nayel et al., 2012). The productivity of such animals is known to be greatly diminished by tick-borne parasitic diseases, particularly babesiosis and theileriosis, causing considerable economic losses on both local and global scales with several complications (Uilenberg, 1995; Ahmed et al., 2002; Al-Saeed et al., 2010; Elsify et al., 2015).

Theileriosis is serious hemoparasitic tick-borne disease caused by the apicomplexan, *Theileria* parasites, infecting a variety of wild and domestic animals and is transmitted trans-stadially by the tick vectors (Preston, 2001; Silva et al., 2010; Nayel et al., 2012). *Theileria annulata* causes tropical theileriosis, a disease transmitted by ticks of the genus *Hyalomma* (Uilenberg, 1981; El-Deeb and Younis, 2009) in North African countries, particularly Egypt (Gharbi et al., 2013; Ayadi et al., 2016). The protozoan is often
present in areas in which the tick species are predominant including a wide range of tropical and subtropical areas, particularly African countries (Purnell, 1978). Accordingly, the disease is commonly proved in northern Africa and Sudan the Middle East, Central Asia, China, India, southern Europe and southern Russia where more than 250 million cattle are estimated to be at risk (McCosker, 1979; Robinson, 1982).

Traditionally, the diagnosis of *Theileria* parasites is based on the clinical symptoms and the detection of morphological characters of piroplasms inside red blood cells in Giemsa-stained thin blood smears (Latif et al., 1977; Muslih et al., 1988). However, such technique is only reliable for the diagnosis in acute cases and has limited value in subclinical cases due to a low parasitemia, when few numbers of piroplasms could be detected in the peripheral blood. Furthermore, high experts are demanded to accurately differentiate among various pathogenic and non-pathogenic piroplasms (Al-Saeed et al., 2010). However, since antibodies may still circulating for some time after the parasite has been cleared from the diseased animals, serological techniques do not often provide information about the actual existence of the parasite associated with the high opportunity of false data as a result of the cross reactions and the weak immune response (Passos et al., 1998; Nayel et al., 2012). The development of molecular techniques has made accurate tools for the detection of the parasite DNA. Such methods are appropriate and successful in neighboring countries which have a similar topographical nature to Egypt; Tunisia (M’ghirbi et al., 2008), United Arab Emirates (Jaffar et al., 2010) and Iran (Zaeemi et al., 2011). The first polymerase chain reaction (PCR) applied for the diagnosis of *T. annulata* in the bovine hosts was based on the *Tams* 1 gene (d’Oliveira et al., 1995). Therefore, the present study was designed to detect the current prevalence of the infection with the hemoparasites, *Theileria annulata*, of cattle in El-Wadi El-Gadid, Egypt. Assessment is based on clinical symptoms, microscopic examination of Giemsa-stained thin blood smears and PCR-molecular diagnosis using the specific primer pair targeting the mitochondrial cytochrome b gene.

**MATERIALS AND METHODS**

**The Study Area and Animals**

A total of 376 Egyptian female Balady-Friesian cross cattle admitted to different veterinary clinics in El-Wadi El-Gadid province (coordinates: 24° 32’ 44” N, 27° 10’ 24” E) in the south-western part of Egypt, were examined for the detection of *Theileria* parasites. Available data concerning the general conditions of animals, the past history of infection, the existence of tick species, clinical symptoms, and other infections were accurately recorded.

**Blood Sampling**

Blood samples were collected for both microscopic examination and PCR (with anticoagulant, sodium salt of EDTA). Blood was collected from the jugular vein and immediately preserved in Eppendorf tubes containing a few drops of EDTA (Hamed et al., 2011).

**Microscopic Examination**

Thin blood smears were fixed in methanol (99.5%) for 5 min and stained for 30 min in Giemsa solution diluted with 5% buffer. Slides were examined for the presence of intracellular hemoparasites. The smears were recorded as negative for piroplasms if no parasites were detected in 50 oil-immersion fields (Moretti et al., 2010).

**DNA Extraction**

DNA was extracted from the blood specimens according to the manufacturer instructions (Sivakumar et al., 2012). Briefly, DNA was extracted from 200 µl of the whole blood from individual cattle using a commercial gSYNCTM DNA extraction kit (Cat. No.GS050) (Geneaid, New Taipei, Taiwan). DNA samples were stored at −20 ºC until use.

**PCR Amplification**

DNA extracted from blood samples obtained from *Theileria*-infected cattle were used for PCR with the use of a specific primer pair, *Cytob* 1, forward (F) 5’-ACTTTGGC-CGTAATGTTAAAC-3’ and reverse (R) 5’-CTCTG-GACCAACTGTTC-3 (Belgic et al., 2010) targeting the amplicon of 312 bp of the mitochondrial cytochrome b gene. Accordingly, 12.5 µl 2X master mix (Geneaid Biotech Ltd., New Taipei, Taiwan), 1 µl 1F (10 pmol/µl), 1 µl R (10 pmol/ µl), 5 µl DNA sample, and 7.5 µl nuclease free water were mixed in a total volume of 25 µl. PCR cycling started with an initial denaturation for 5 min at 95°C followed by 37 cycles of denaturation for 30 second at 95°C, annealing for 60 second at 52°C, and elongation for 1 min at 72°C. The final extension occurred at 72 ºC was allowed to proceed for 7 min. PCR amplicons were electrophoresed on 1.5% agarose gel with ethidium bromide and photographed using a transilluminator.

**RESULTS**

Among 376 examined Balady-Friesian cross cattle, 102 (27.13%) animals had clinical symptoms of piroplasmosis, predominantly enlargement of superficial lymph nodes, particularly parotid, prescapular, and prefemoral lymph nodes, as well as a sustainable high fever (40-42°C) extended along the course of infection. Other clinical symptoms including anorexia, loss of generalized conditions,
lacrimation, corneal affections, ocu-no nasal discharge and diarrhea were consistent (Figure 1).

Figure 1: Clinical symptoms of theileriosis in examined Balady-Friesian cattle in Egypt. a) The perineal area of an infected animal heavily infected with tick species. b) An eye of Theileria-infected cattle showing ticks infestations. c) Severe bovine ocular affections. d) Enlargement of superficial lymph nodes

Among clinically infected cattle, light microscopy of Giemsa-stained thin blood smears revealed that 35(9.31%) specimens showed the presence of intra-lymphocytic and/or intra-erythrocytic stages of hemoparasites. As a result of multiplication of Theileria parasites, both macroschizonts (Figure 2) and microschizonts were found in most of clinically infected cattle. The appearance of a high fever and the generalized lymphadenopathy in such animals proved the infection. Meanwhile, some infected cattle exhibited ruptured schizonts due to prolonged infection. Sporadically, infected red blood cells might contain individual merozoites.

Figure 2: Bovine lymphocytes infected with Theileria parasites. Arrow referred to macroschizont. Scale bar= 20 µm

Application of PCR using a specific primer pair, Cytob1, for *Theileria annulata* elucidated that 43(11.44%) animals had theileriosis targeting the amplicon of 312 bp (Figure 3). Diagnostic bands proved the bovine theileriosis.

Figure 3: PCR amplification of DNA extracted from blood specimens of Theileria-infected cattle using the specific primer pair, Cytob1, targeting the mitochondrial cytochrome b gene. Note the diagnostic band appeared as 312 bp. Lane M revealed 100-bp ladder DNA marker. Lanes 1-10 indicated 10 positive animals (animals are numbered as 23, 44, 56, 60, 76, 85, 91, 92, 93 and 98, respectively). Lanes 11 and 12 indicated negative animals (animals numbered 18 and 79)

It is worth mentioning that most of clinically-positive animals and all those positive by the use of microscopic examination and PCR are associated with the abundance of ticks vector. Highly feverish animals and those with enlarged lymph nodes were heavily parasitized by ticks. Application of both traditional and molecular tools is useful in diagnosis of infection, particularly in subclinical cases.

DISCUSSION

In the current study, Egyptian Balady-Friesian cattle, bred in different farms and admitted to veterinary clinics in El-Wadi El-Gadid province, were surveyed for tropical theileriosis. Subsequently, cattle underwent both traditional diagnosis, by the use of light microscopy of Giemsa-stained thin blood smears, and a molecular technique, using a specific primer pair.

The present investigation revealed that 9.31% of examined cattle had *Theileria* parasites using Giemsa-stained thin blood films. In Egypt, several studies investigated the prevalence of *Theileria* parasites in cattle. Among those, Abdel-Rady et al. (2008) detected that out of 120 cattle, 31 (25.8%) animals in Assiut, Sohage and El-Wadi El-Gadid provinces harboured *T. annulata* parasites in Giemsa-stained thin blood smears. Moreover, Abdel-Rady et al. (2010) revealed an infection rate of 65.6% by micro-
scopic examination of thin blood smears in cattle examined in El-Wadi El-Gadid, Assiut, El-Fayoum, El-Minia and Sohage provinces. In both reports, theileriosis might be related to the predominance of the vector, *Hyalomma* species, however, in the second study, the higher infection rate could be attributed to the use of microscopic examination for such animals in the early acute stage allowing great opportunities for both intracellular piromals and lymphocytic Koch’s blue bodies. Furthermore, Al-Hosary et al. (2015) found that Giemsa-stained thin blood smear revealed 30.98% cattle (145/468) positive for theileriosis in El-Wadi El-Gadid. Several reports in such province indicated the abundance of *Hyalomma* ticks which encouraged in the hot climate, the main topographical feature of El-Wadi El-Gadid province. Meanwhile, Elsify et al. (2015) recorded an infection rate of 9.56% with *Theileria annulata* in cattle in Menoufia, Behera, Giza and Sohage provinces, Egypt by the use of light microscopy of Giemsa-stained thin blood smears. In a similar African country, Algeria, Ayadi et al. (2016) demonstrated that the percentages of *Theileria*-positive Giemsa-stained thin blood smears, before and after the tick season, were 10.4% and 16.4%, respectively.

In the current work, two techniques were used for the diagnosis of *T. annulata*, microscopic examination of Giemsa-stained thin blood smears and PCR. In the former, the number of infected blood cells increases during the acute (early) infection, allowing an ease appearance of intracellular trophozoites and, consequently, a reliable detection of the parasite by microscopic examination in Giemsa-stained thin blood smears. It is well known that PCR is more sensitive and specific than traditional methods. Accordingly, authors noticed that microscopic examination of thin blood smears might be useful in acute infections of theileriosis. However, establishment of the immune response drastically decreases the number of infected erythrocytes. Moreover, during the tick season, the cattle immune system is enhanced by further infections leading to a consequent decline in sensitivity of such traditional method (Ayadi et al., 2016). Several studies reported that PCR is more sensitive and specific than microscopic examination of Giemsa-stained thin blood smears for detecting bovine theileriosis, particularly in carrier animals (Azizi et al., 2008; Safarpour Dehkordi et al., 2012; Ayadi et al., 2016).

Currently, PCR-based technique using *T. annulata*-specific primer pair, Cytob1, reported that 11.44% of animals were theileriosis-positive and had specific diagnostic bands at 312 bp. Similarly, Ibrahim et al. (2009) revealed the percentages of PCR-positive animals for *Theileria annulata* were 13% (7/54). Moreover, Elsify et al. (2015) detected that *T. annulata* was found in 9.56% of examined bovine blood samples using PCR in Behera, Giza, and Sohag provinces, Egypt. On the other hand, Abdel Rady et al. (2010), in El-Wadi El-Gadid, Assiut, El-Fayoum, El-Minia and Sohage provinces, recorded that the sensitivity of PCR using Tams1 primer pair was 58.3%. Meanwhile, in the same province, El-Wadi El-Gadid, Al-Hosary et al. (2015) revealed that Tams1 primer-based PCR proved *Theileria annulata* infection in 343 (73.29%) out of 468 bovine blood specimens. The discrepancy from the present study might be referred to the intensity of ticks infestation. Similar to the current results, Hoghooghi-Rad et al. (2011) found that the semi-nested PCR accurately revealed 12 (7.5%) positive bovine blood specimens in Golestan province of Iran. In Banaskantha district, India, Chauhan et al. (2015) recorded that 24 (46.15%) cows were infected with *T. annulata* using Tams1 gene-based PCR. Concomitantly, Acharya et al. (2017) recovered 68% (34/50) infection rate of theileriosis in cattle in Odisha, India using PCR. In Europe, Brigido et al. (2004) applied PCR on bovine blood and revealed that 3 (2.57%) out of 116 specimens were infected with *Theileria* and/or *Babesia* parasites.

In conclusion, the present investigation has emerged tropical theileriosis, due to *Theileria annulata*, by the use of both traditional methods, microscopic examination of Giemsa-stained thin blood smears, and a molecular assay, conventional PCR based on the use of *T. annulata*-specific primer pair targeting the mitochondrial cytochrome b gene (312 bp) in El-Wadi El-Gadid, Egypt. It is worthy to mention that cattle of local breeds are generally more resistant than exotic animals, explaining that the infection rate in the current breed, Balady-Friesian species, is lower than imported species. For a proper control of theileriosis, it is advisable to apply the appropriate acaricides together with eradication of tick vectors. In such concern, sealing of cracks and plastering and smoothing the inner and outer walls of animal stables could reduce the survival of ticks vector.

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**CONFLICT OF INTEREST**

There are no conflicts of interest.

**AUTHORS CONTRIBUTION**

Experimental design and ideas developed by Khaled Mohamed El-Dakhly. Data and samples collection by Saad Soliman, Omima Abdel-Fatah and Ahmed Anwar. Lab analysis by Waleed Arafa and Khaled Mohamed El-Dakhly. Manuscript preparation by Khaled Mohamed El-Da-
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