# ORIGINAL PAPER

# The use of different diagnostic tools for *Babesia* and *Theileria* parasites in cattle in Menofia, Egypt

Mohamed Nayel • Khaled Mohamed El-Dakhly • Mahmoud Aboulaila • Ahmed Elsify • Hany Hassan • Elsayed Ibrahim • Akram Salama • Tokuma Yanai

Received: 17 March 2012 / Accepted: 10 April 2012 / Published online: 29 April 2012 © Springer-Verlag 2012

**Abstract** Bovine piroplasmosis is caused by tick-borne hemoprotozoans of the genera *Babesia* and *Theileria* and is the most prevalent in tropical and subtropical countries, causing a major economic impact worldwide. In the current study, a total of 405 cattle of different ages, sexes, and breeds were randomly sampled for surveying and diagnosis of babesiosis and theileriosis using three methods: direct microscopy (blood smears), indirect fluorescent antibody test (IFAT) and polymerase chain reaction (PCR). Giemsastained blood smears revealed that, out of 405 examined cattle, 33 (8.15 %) were infected with *Babesia* sp. and 65 (16.05 %) with *Theileria* sp. (total number of infected cattle was 98). Mixed infection was seen in 11 (2.72 %) animals.

M. Nayel · A. Elsify · H. Hassan · A. Salama Department of Animal Medicine and Infectious Diseases, Faculty of Veterinary Medicine, Menofia University, Sadat City, Egypt

K. M. El-Dakhly Department of Parasitology, Faculty of Veterinary Medicine, Beni-Suef University, Beni-Suef 62511, Egypt

M. Aboulaila Department of Parasitology, Faculty of Veterinary Medicine, Menofia University, Sadat City, Egypt

E. Ibrahim Department of Animal Medicine and Infectious Diseases, Faculty of Veterinary Medicine, Benha University, Benha, Egypt

K. M. El-Dakhly · T. Yanai (⊠)
Department of Veterinary Pathology,
Faculty of Applied Biological Science, Gifu University,
1-1 Yanagido,
Gifu 501-1193, Japan
e-mail: yanai@gifu-u.ac.jp

Moreover, application of the three diagnostic assays on 158 randomly sampled cattle indicated that 17 (10.76 %) and 33 (20.89 %) were positive for *Babesia* and *Theileria* spp. by the direct smear technique, 25 (15.82 %) and 33 (20.89 %) by IFAT (fluorescence was greenish yellow for Babesia and yellowish for Theileria), and 20 (12.66 %) and 38 (24.05 %) by PCR. Using primers specific for Babesia and Theileria spp., we found that diagnostic bands appeared at ~350 and ~370 bp, respectively indicating the presence of these piroplasms. Statistically, there was a non-significant difference of the positivity in response to the three techniques; thus, any of these methods can be described as useful for diagnosing blood parasites in both domesticated animals and birds. On the basis of the obtained results, it could be concluded that direct microscopy can be used in acute infections, whereas IFAT and PCR are useful in chronicity.

## Introduction

*Babesia* and *Theileria* species are apicomplexan–hemoprotozoan parasites transmitted by Ixodidae ticks (Preston 2001; Silva et al. 2010) and are viewed to be devastating parasites affecting the production of livestock, mainly cattle and small ruminants. They pose a significant problem for veterinary authorities, occasionally emerging in conjunction with other disease conditions, and thus being difficult to pinpoint (Altay et al. 2008). These infections are of worldwide importance and are characterized by anemia, icterus, hemoglobinuria, and death, and as a result, they have a high economic impact in several parts of the world, including tropical and temperate countries (Wagner et al. 2002).

Bovine babesiosis is caused by multiple species: *Babesia* bigemina, Babesia divergens, Babesia bovis, and Babesia major. Babesia species have the potential for wide

distribution wherever their tick vectors are encountered. Two species, *B. bigemina* and *B. bovis*, have a considerable impact on cattle health and productivity in tropical and subtropical countries. Cattle suffering from theileriosis demonstrate varying clinical signs that range from lymphoproliferative changes with high morbidity and mortality, as seen with *Theileria annulata* and *Theileria parva*, to benign or mild disease, as seen with *Theileria orientalis* (Altay et al. 2008; Safieldin et al. 2011).

Detection of these blood parasites is highly beneficial in early diagnosis. Traditionally, microscopy using Giemsastained blood smears has been considered the "gold standard" for detecting Babesia and Theileria organisms in the blood of infected animals, particularly in acute cases, but not in carriers, where the parasitemia is low, with small numbers of the protozoa in the peripheral blood (Friedhoff and Bose 1994; Bose et al. 1995). Therefore, serological techniques were proposed for detecting circulating antibodies against these parasites, particularly in subclinical infections during epidemiological investigations. In addition, serological diagnosis using the indirect fluorescent antibody test can be used to detect antibodies against the Theileria species (Leemans et al. 1997). One disadvantage of such tests is the occurrence of false-positive and false-negative results, involving cross-reactions or improper specific immune response (Passos et al. 1998), while another is the inability to differentiate between previous and current infections, making sensitive and highly specified diagnostic techniques for Babesia and Theileria. Therefore, the application of PCRbased techniques is imperative in order to detect these hemoparasites in carrier animals (Criado-Fornelio et al. 2009). These methods were used for diagnosis of babesiosis and theileriosis in several species of related countries, showing similar climatic conditions, including Tunisia (M'ghirbi et al. 2008), United Arab Emirates (Jaffar et al. 2010) and Iran (Zaeemi et al. 2011).

The present study was performed for the purpose of surveying and diagnosing both *Babesia* and *Theileria* spp. in cattle in Menofia province, Egypt, using Giemsa-stained blood films, indirect fluorescent antibody test (IFAT)-sero-logical test, and PCR assay.

#### Materials and methods

#### Animals and the study area

and blood samples were collected to confirm clinical diagnosis of both diseased and carrier animals.

## Blood smears

Thin smears were prepared from EDTA-whole blood on clean and dry slides, fixed in methanol, stained with Giemsa stain, and microscopically examined for the detection of intraerythrocytic forms of both *Babesia* and *Theileria* spp. piroplasms at  $100 \times$  objective magnification. The smears were recorded as negative for piroplasms if no parasites were detected in 50 oil-immersion fields (Moretti et al. 2010).

## Blood samples

Blood samples were collected for both PCR (with anticoagulant, sodium salt of EDTA) and IFAT (without anticoagulant). Blood was collected from the jugular vein and immediately preserved in Eppendorf tubes containing a few drops of EDTA.

# Laboratory assays

## Indirect fluorescent antibody test

Available parasite-coated slides for both *Babesia* and *Theileria* antigens were kindly provided by the Veterinary Serum and Vaccine Research Institute, Abassia, Egypt, ready for the indirect fluorescent antibody test as described by Leeflang and Perie (1972). A positive reaction is indicated by a bright fluorescence (Papadopoulos et al. 1996).

### Polymerase chain reaction

*DNA extraction and amplification* DNA extraction was performed according to the Manual Chemical Method (rapid isolation of mammalian DNA) using cell lysis buffer (pH 8.0), ethanol (70 %), isopropanol, potassium acetate solution, red blood cell lysis buffer, and proteinase K (20 mg/ml) as described by Sambrook and Russell (2001).

PCR amplification was performed in a final reaction volume of 50  $\mu$ l containing 200  $\mu$ M of each dNTPs, 0.2 uM of each primer, 2.5 U of Taq DNA polymerase (Fermentas, Germany), 10 mM of TBE (tris, boric acid, and EDTA) buffer pH 8.0 containing 1.5 mM MgCl<sub>2</sub> and 5  $\mu$ l of the DNA template. The designated primers were obtained from Bioneer Corporation (064550), Korea. The oligonucleotide sequences of the primers used were forward strand primer BAB GF2 (5'-GTC TTG TAA TTG GAA TGA TGG-3') and reverse strand primer BAB GR2 (5'-CCA AAG ACT TTG ATT TCT CTC-3') (Adaszek and Winiarczyk 2008) under the following conditions: an initial denaturation at 95 °C for 5 min followed by 40 cycles of



Fig. 1 Diagrammatic scale showing the overall prevalence of *Babesia* and *Theileria* parasites among the examined cattle in Menofia province

denaturation at 95 °C for 1 min, annealing at 55 °C for 1 min, and extension at 72 °C for 1 min, followed by final extension at 72 °C for 10 min.

The amplification reactions were carried out in a PCR thermal cycler Biometra T- personal/Germany S/N 1205334 and the corresponding amplicons were checked on 1.5 % agarose gel, stained with ethidium bromide, examined under UV transilluminator, and photographed using a digital camera.

## Statistical analysis

Data were analyzed using multiple comparisons between different diagnostic methods for *Babesia* and *Theileria* parasites, including direct microscopy, IFAT, and PCR using the general linear model, Tukey test. This was carried out using Minitab statistical software (MTW13) (Raza et al. 2007). *P*<0.05 was accepted to be statistically significant.

## Results

In this study, parasitological examination of 405 randomly selected cattle, by direct microscopy using Giemsa-stained



Fig. 2 Greenish yellow fluorescence (*arrow*) indicating intraerythrocytic stages of *Babesia* sp. piroplasms using IFAT (×100)



Fig. 3 Yellowish fluorescence of *Theileria* sp. piroplasms (*arrow*) indicating positive reaction with IFAT (×100)

thin blood films, revealed that 98 had intra-erythrocytic stages of piroplasms of both Babesia and Theileria spp. with an overall prevalence of 24.2 %. Among those, 33 (8.15 %) were infected by Babesia sp., 65 (16.05 %) had Theileria sp., and 11 cases showed mixed infection (Fig. 1). These results were consistent with the clinical signs and previous case histories taken from the farmers and owners of animals that were collected from different districts in Menofia province. As a result of chronic infection, the developing antibodies reacted positively using the IFAT, producing a fluorescence that was clearly distinct and greenish yellow (Fig. 2) or yellowish (Fig. 3) in color and indicated the presence of the intra-erythrocytic stages of both Babesia and Theileria piroplasms, respectively. PCR findings showed that diagnostic bands were produced at ~350 and ~370 bp, and the fragments were specific for both Babesia and Theileria spp., respectively (Fig. 4). This



**Fig. 4** PCR results showing diagnostic bands at 350-bp for *Babesia* (a) and 370bp for *Theileria* (b). *Lane M* 100-bp ladder=DNA marker (from Fermentas, Germany)

Table 1Comparative detectionof Babesia and Theileria para-sites in cattle using directmicroscopy, IFAT, and PCR

Parasite	Assay $(n=158)$						P value
	Direct microscopy		IFAT		PCR		
	Infected	Percent	Infected	Percent	Infected	Percent	
<i>Babesia</i> sp.	17	10.76	25	15.82	20	12.66	NS
Theileria sp.	33	20.89	33	20.89	38	25.05	NS

finding revealed the presence of both hemoparasites in the blood of examined cattle.

Furthermore, the seroprevalence of *Babesia* and *Theileria* species in cattle using IFAT revealed that out of 158 serum samples examined for the presence of antibodies, 25 (15.82 %) and 33 (20.89 %) were positive for *Babesia* spp. and *Theileria* spp., respectively. Furthermore, PCR findings indicated that 20 (12.66 %) and 38 (24.05 %) were positive for *Babesia* and *Theileria* spp., respectively (Table 1).

The findings in the present study confirmed that the use of direct microscopy, IFAT, or PCR is valuable in the diagnosis of *Babesia* and *Theileria* spp. Based on the confidence intervals calculated using the Tukey method (Minitab), it has been found that the differences between the three methods were not statistically significant, as there was no pairwise variation between infected animals (Fig. 5). This means that in some instances, mainly acute infections, the use of any of these methods may be applicable, although IFAT and PCR are preferable in chronic infections. It is worth mentioning that the methodology in the present investigation is based on collecting a considerable number of animals regardless of studying the effects of sex, age, or season on the infection level.

## Discussion

Babesiosis and theileriosis have extensive prevalence and mortality rates with high economic losses in several countries (Shahnawaz et al. 2011). In Egypt, large numbers of cattle are infected with subclinical piroplasmosis (Adham et al. 2009). In such cases, in addition to the parasitological

Fig. 5 Comparison of the positivity of *Babesia* and *Theileria* spp. using three diagnostic techniques: direct microscopy, IFAT, and PCR

examination of stained blood films for detecting the *Babesia* protozoan parasites, low parasitemia necessitates the use of more advanced diagnostic tools, rather than conventional ones, to detect specific antibodies. In addition, negative microscopic examination does not exclude the possibility of infection (Weiland and Reiter 1988). Subclinical babesiosis and theileriosis lead to the affected livestock, including cattle and small ruminants, becoming chronic carriers of the piroplasms and in turn sources of infection for tick vectors, and cause natural transmission of the disease. Therefore,

latent infections are the target in the epidemiology of the

diseases. In the present study, the results obtained from Giemsastained blood smears revealed that 33 smears (8.1 %) were positive for Babesia. Similar results were obtained by Sevnc et al. (2001) in Turkey and Mazyad and Khalaf (2002) in Egypt. On the other hand, Jeon (1978) in Korea and Osaki et al. (2002) in Brazil detected infection rates of 23 and 64 %, respectively. Fluctuation in the prevalence rates might be due to the variation of environmental conditions that affect both parasites and vectors. For Theileria sp., the present investigation showed that 65 (16.05 %) of 405 cattle were positive. These results agreed with those obtained by Jeon (1978) in Korea (17%), Acici (1995) in Turkey (17%), and those obtained in Egypt, which included Abu El-Magd (1980) in Quena province (11.1 %) and Adel (2007) in Gharbia province (11.31 %). Conversely, our results opposed a number of reports in Egypt, among them El Bahy (1986), who revealed prevalence rates of 65 and 53 % in cattle and buffaloes, respectively, and Gamal EI-Dien (1993) in El-Behera province, who found that the prevalence of T. annulata was 65.4 % using stained blood films.



Variation in prevalence rates could possibly be attributed to an abundance of the vectors as a result of high temperature and humidity. The present investigation showed that mixed infection, using Giemsa-stained blood smears, of *Babesia* and *Theileria* spp. appeared in 11 (2.72 %) of 405 examined cattle. These results coincided with those obtained by Dumanli and Ozer (1987) in Turkey, who found a mixed infection rate for *B. bigemina* and *T. annulata* of 1.5 % in cattle.

Using IFAT, the present study revealed an infection rate of 15.82 % (25/158) for Babesia sp. This finding was in accordance with that obtained by Terkawi et al. (2012), who examined a herd of cattle in the central region of Syria using IFAT and found infection rates of 18.36 and 21.74 % for B. bovis and B. bigemina, respectively. Moreover, Jaffar et al. (2010) in Dubai, UAE recorded infection rates of 10.5 and 33.3 % for equine babesiosis and theileriosis. On the other hand, Singh et al. (2009) recorded a rate of 56.11 % for B. bigemina in India, Iseki et al. (2010) noted rates of 68.8 and 75.8 % for B. bovis and B. bigemina in Thailand, and Sevgili et al. (2010) recorded an infection rate of 43.9 % for B. bigemina in Sanliura, Turkey. The variation in infection rates could be due to differences in climatic conditions and species of cattle. Concerning Theileria sp., the current investigation revealed that out of 158 bovine serum samples, 33 (20.89 %) was found to be positive for Theileria antibodies. This result agreed with that obtained by Sayin et al. (2003), who noted that 34 (21 %) of 155 examined cattle were found to be seropositive for T. annulata. Moreover, Adel (2007) in Gharbia province, Egypt reported that the incidence peaks of T. annulata seropositive native breed cattle using IFAT were 27.8 and 22 % in spring and summer, respectively. On the other hand, Gamal EI-Dien (1993) in EI-Behera province, Egypt detected that the prevalence of T. annulata was 71.9 %, and Caille (1987) in Somalia found that 71.2 % of cattle had antibodies against T. annulata. This discrepancy may be due to variation in the susceptibility of the animal species and could also be due to variance in animal locale.

Results of PCR revealed that 20 (12.66 %) of 158 animals were positive for *Babesia* sp. These findings were similar to those mentioned by Oliveira-Sequeira et al. (2005), who recorded a 10 % infection rate for *Babesia* using PCR and M'ghirbi et al. (2008) who revealed an infection rate of 11.11 % for bovine *Babesia* spp. Similar results were obtained by Figueroa et al. (1993) and Gubbels et al. (2002). On the other hand, De Vos and Potgieter (1994) in France found a 20 % infection rate, Costa-Junior et al. (2006) in Brazil found that 8 of 30 cattle (26.7 %) were positive for *Babesia* infection using PCR, and Rania (2009) in Egypt mentioned that the infection rate of *Babesia* sp. was 25.33 %. These discrepancies could be due to changes in locale. The infection rate of *Theileria* using PCR indicated that 60 animals (21.8 %) were positive. This result was similar to those cited by Ogden et al. (2003), Aktas et al. (2005), Altay et al. (2007) and M'ghirbi et al. (2008). Ogden et al. found an infection rate of 23.4 % for *Theileria* sp. in cows in Tanzania using PCR.

The authors concluded that the three techniques, direct microscopy, IFAT, and PCR, are all methods of choice with slight variation in their results, and this was confirmed by obtaining non-significant findings among them. Consequently, they are used in the detection of prevalence of blood parasites, primarily *Babesia* and *Theileria*. Direct microscopy with Giemsa-stained blood films is the conventional and the more practical method on farms, as it is rapid and inexpensive, but it is useful only in acute infections. Moreover, IFAT and PCR are modern assays that help veterinarians detect protozoan parasites in chronic carrier animals, and they circumvent the problem of false negative results obtained under direct microscopy.

Acknowledgments The authors are grateful to veterinarians and farmers in Menofia province, Egypt, who provided assistance in collecting samples. We also offer our sincere thanks to Dr. El-Shyamaa Nabil for providing statistical guidance in the study.

## References

- Abu El-Magd MM (1980) Some epidemiological studies about theileriosis at Quna Governorate. PhD. thesis, Faculty of Veterinary Medicine, Assuit University, Assuit, Egypt
- Acici M (1995) Prevalence of blood parasites in cattle in the Samsun region, Turkey. Etlik Vet Mikrob Derg 8:271–277
- Adaszek L, Winiarczyk S (2008) Molecular characterization of *Babesia canis canis* isolates from naturally infected dogs in Poland. Vet Parasitol 152:235–241
- Adel EM (2007) Studies on some blood parasites infecting farm animals in Gharbia

governorate, Egypt. PhD thesis, Faculty of Veterinary Medicine, Cairo University, Cairo, Egypt

- Adham FK, Abd-El-Samie EM, Gabre RM, El Hussein H (2009) Detection of tick blood parasites in Egypt using PCR assay I— Babesia bovis and Babesia bigemina. Parasitol Res 105:721–730
- Aktas M, Altay K, Dumanli N (2005) Development of a polymerase chain reaction method for diagnosis of *Babesia ovis* infection in sheep and goats. Vet Parasitol 133:277–281
- Altay K, Dumanli N, Aktas M (2007) Molecular identification, genetic diversity and distribution of *Theileria* and *Babesia* species infecting small ruminants. Vet Parasitol 147:161–165
- Altay K, Fatih Aydin M, Dumanli N, Aktas M (2008) Molecular detection of *Theileria* and *Babesia* infections in cattle. Vet Parasitol 158:295–301
- Bose R, Jorgensen WK, Dalgliesh RJ, Friedhoff KT, De Vos AJ (1995) Current state and future trends in the diagnosis of babesiosis. Vet Parasitol 57:61–74
- Caille JY (1987) Serological survey of the prevalence and seasonal incidence of haemoprotozoa in livestock in Somalia. Thesis, Freie Universitat Berlin, Berlin, Germany
- Costa-Junior LM, Rabelo EML, Filho OAM, Ribeiro MFB (2006) Comparison of different direct diagnostic methods to identify *Babesia bovis* and *Babesia bigemina* in animals vaccinated with live attenuated parasites. Vet Parasitol 139:231–236

- Criado-Fornelio A, Buling A, Pingret JL, Etievant M, Boucraut-Baralon C, Alongi A, Agnone A, Torina A (2009) Hemoprotozoa of domestic animals in France: prevalence and molecular characterization. Vet Parasitol 159:73–76
- De Vos AJ, Potgieter FT (1994) Bovine babesiosis. In: Coetzer JAW, Thomson GR, Tustin RC (eds) Infectious diseases of livestock. Oxford University Press, Cape Town, pp 278–294
- Dumanli N, Ozer E (1987) Elaziğ yöresinde siğirlarda görülen kan parazitleri ve yayihs jan üzerinde araştirmalar. Selcuk Ü nin. Vet Fak Derg 3:159–166
- El Bahy NMA (1986) Some studies on ticks and tick borne disease among ruminants in Fayom Governorate. M.V.Sc. thesis (Parasitology), Faculty of Veterinary Medicine, Cairo University, Cairo, Egypt
- Figueroa JV, Chieves LP, Johnson GS, Buening GM (1993) Multiplex polymerase chain reaction based assay for the detection of *Babesia bigemina*, *Babesia* bovis and *Anaplasma marginale* DNA in bovine blood. Vet Parasitol 50:69–81
- Friedhoff K, Bose R (1994) Recent developments in diagnostics of some tick-borne diseases. In: Uilenberg, G., Permin, A., Hansen, J.W. (eds). Use of applicable biotechnological methods for diagnosing haemoparasites. Proceedings of the Expert Consultation, Merida, Mexico, October 4–6, 1993. Food and Agriculture Organization of the United Nations (FAO), Rome, Italy, pp 46–57
- Gamal EI-Dien HY (1993) Studies on *Theileria* protozoan among cattle in Behera Province. M.V.Sc. thesis, Faculty of Veterinary Medicine, Alexandria University, Alexandria, Egypt
- Gubbels MJ, Yin H, Bai Q, Liu G, Nijman IJ, Jongejan F (2002) The phylogenetic position of the *Theileria buffeli* group in relation to other *Theileria* species. Parasitol Res 88:S28–S32
- Iseki H, Zhou L, Kim C, Inpankaew T, Sununta C, Yokoyama N, Xuan X, Jittapalapong S, Igarashi I (2010) Seroprevalence of *Babesia* infections of dairy cows in northern Thailand. Vet Parasitol 170:193–196
- Jaffar O, Abdishakur F, Hakimuddin F, Riya A, Wernery U, Schuster RK (2010) A comparative study of serological tests and PCR for the diagnosis of equine piroplasmosis. Parasitol Res 106:709–713
- Jeon Y (1978) A survey of babesiosis and theileriosis in Korean cattle. Korean J Vet Res 18:23–26
- Leeflang MJ, Perie CD (1972) Duration of serological response to the indirect fluorescent antibody test of cattle recovered from *Theileria parva* infection. Res Vet Sci 14:270–271
- Leemans I, Hooshmand-Rad P, Uggla A (1997) The indirect fluorescent antibody test based on schizont antigen for study of the sheep parasite *Theileria lestoquardi*. Vet Parasitol 69:9–18
- M'ghirbi Y, Hurtado A, Brandika J, Khlif K, Ketata Z, Bouattour A (2008) A molecular survey of *Theileria* and *Babesia* parasites in cattle, with a note on the distribution of ticks in Tunisia. Parasitol Res 103:435–442
- Mazyad SA, Khalaf SA (2002) Studies on Theileria and Babesia infecting live and slaughtered animals in Al Arish and El Hasanah, North Sinai Governorate, Egypt. J Egypt Soc Parasitol 32 (2):601–610
- Moretti A, Mangili V, Salvatori R, Maresca C, Scoccia E, Torina A, Moretta I, Gabrielli S, Tampieri MP, Pietrobelli M (2010) Prevalence and diagnosis of *Babesia* and *Theileria* infections in horses in Italy: a preliminary study. Vet J 184:346–350
- Ogden NH, Gwakisa P, Swai E, French NP, Fitzpatrick J, Kambarage D, Bryant M (2003) Evaluation of PCR to detect *Theileria parva* in field-collected tick and bovine samples in Tanzania. Vet Parasitol 112(3):177–183
- Oliveira-Sequeira TC, Oliveira MC, Araujo JP Jr, Amarante AF (2005) PCR-based detection of *Babesia* bovis and *Babesia bigemina* in

their natural host *Boophilus microplus* and cattle. Int J Parasitol 35:105–111

- Osaki SC, Vidotto O, Marana ERM, Vidotto MC, Yoshihara E, Pacheco RC, Igarashi M, Minho AP (2002) Occurrence of antibodies against *Babesia bovis* and studies on natural infection in Nelore cattle, in Umuarama municipality, Paraná State, Brazil. Rev Bras Parasitol Vet 11:73–83
- Papadopoulos B, Marie Perie N, Uilenberg G (1996) Piroplasms of domestic animals in the Macedonia region of Greece. 1. Serological cross-reactions. Vet Parasitol 63:41–56
- Passos LM, Bell-Sakyi L, Brown CG (1998) Immunochemical characterization of in vitro culture-derived antigens of *Babesia bovis* and *Babesia bigemina*. Vet Parasitol 76:239–249
- Preston PM (2001) Theilerioses. In: Wallingford MW (ed) Encyclopedia of arthropod-transmitted infections of man and domesticated animals. CABI, Wallingford, pp 487–502
- Rania YE (2009) Some studies on diagnosis on babesiosis. M.V.Sc. thesis, Faculty of Veterinary Medicine, Benha University, Benha, Egypt
- Raza MA, Iqbal Z, Jabbar A, Yaseen M (2007) Point prevalence of gastrointestinal helminthiasis in ruminants in southern Punjab, Pakistan. J Helminthol 81:323–328
- Safieldin M, Abdel Gadir A, Elmalik Kh (2011) Factors affecting seasonal prevalence of blood parasites in dairy cattle in Omdurman locality, Sudan. J Cell Anim Biol 5(1):17–19
- Sambrook J, Russell DW (2001) Molecular cloning: a laboratory manual, 3rd edn. Cold Spring Harbor Laboratory Press, New York
- Sayin F, Dincer S, Karaer Z, Cakmak A, Inci A, Yukari BA, Vatansever Z, Nalbantoglu S (2003) Studies on the epidemiology of tropical theileriosis (*Theileria annulata* infection) in cattle in central Anatolia, Turkey. Trop Anim Health Prod 35(6):521–539
- Sevgili M, Cakmak A, Gokcen A, Altas MG, Ergun G (2010) Prevalence of *Theileria annulata* and *Babesia bigemina* in cattle in the vicinity of Sanliurfa. J Anim Vet Adv 9(2):292–296
- Sevnc F, Sevnc M, Brdane FM, Altnoz F (2001) Prevalence of *Babesia* bigemina in cattle. Rev Med Vet 152(5):395–398
- Shahnawaz S, Ali M, Aslam MA, Fatima R, Chaudhry ZI, Hassan MU, Ali M, Iqbal F (2011) A study on the prevalence of a ticktransmitted pathogen, *Theileria annulata*, the hematological profile of cattle from Southern Punjab (Pakistan). Parasitol Res 109:1155–1160
- Silva MG, Marques PX, Oliva A (2010) Detection of *Babesia* and *Theileria* species infection in cattle from Portugal using a reverse line blotting method. Vet Parasitol 174:199–205
- Singh H, Mishra AK, Rao JR, Tewari AK (2009) Comparison of indirect fluorescent antibody test (IFAT) and slide enzyme linked immunosorbent assay (SELISA) for diagnosis of *Babesia bigemina* infection in bovines. Trop Anim Health Prod 41:153–159
- Terkawi MA, Alhasan H, Huyen NX, Sabagh A, Awier K, Cao S, Goo Y, Aboge G, Yokoyama N, Nihikawa Y, Kalb-Allouz A, Tabbaa D, Igarashi I (2012) Molecular and serological prevalence of *Babesia bovis* and *Babesia bigemina* in cattle from central region of Syria. Vet Parasitol. doi:10.1016/j.vetpar.2011.12.038
- Wagner GG, Holman P, Waghela S (2002) Babesiosis and heartwater: threats without boundaries. Vet Clin Food Anim 18:417–430
- Weiland G, Reiter I (1988) Methods for measurement of the serological response to Babesia. In: Ristic M (ed) Babesiosis of domestic animals and man. CRC, Boca Raton, pp 143–162
- Zaeemi M, Haddadzadeh H, Khazraiinia P, Kazemi B, Bandehpour M (2011) Identification of different *Theileria* species (*Theileria lestoquardi*, *Theileria ovis*, and *Theileria annulata*) in naturally infected sheep using nested PCR–RELP. Parasitol Res 108:837– 843