Detection the infection with Babesia spp. Cytauxzoon felis and Haemobartonella felis in stray cats in Mosul

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Abstract

The current study was included to examination of 50 blood samples which collected from 50 stray cats in Mosul city for diagnosis Babesia spp., Cytauxzoon felis and Haemobartonella felis, different diagnostic techniques were used for differentiation between these parasites, they were: Impression smear, fine needle aspiration and histological examination of the liver and spleen. Some clinical signs and pathological changes were observed in the affected cats which may be result from the infection with these blood parasites. The percentage of infection with Babesia spp. (small type), Cytauxzoon felis, and Haemobartonella felis were 26%, 22%, 42% respectively. The sizes of Babesia spp. was between 1.33-2.15 µm within a mean of 1.53 µm and the size of Cytauxzoon felis ranged between 0.8-2.3 within a mean of 1.64 µm while the sizes of Haemobartonella felis varied between 0.5-2 µm within a mean of 1.15 µm. Tissue stage (schizonts) of Cytauxzoon felis were diagnosed in the cytoplasm of macrophages in the liver and spleen with different stage of replication.

Keywords: Babesia, Cytauxzoon, Haemobartonella, Cat.

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Introduction

Babesia spp are tick–borne hemoprotozoal parasites found in a variety of domestic, wild animals and humans, in domestic cats, babesiosis has primarily been reported in South Africa and Sudan where infected is mainly due to Babesia felis, a small Babesia spp that causes anemia and icterus (1). Others Babesia spp were detected in naturally
infected domestic cats in different countries such as: B. leo from Africa, B. cati from India, B. canis canis from Spain and Portugal, and B. microti – like spp form Portugal (2). In addition sporadic cases of infection in domestic cats by unidentified Babesia spp. (3-6). Feline babesiosis often presents as a chronic low grade disease (1) and the appearance of certain clinical signs (7). C. felis is a protozoan parasite that causes cytauxzoonosis (Cat's theileriosis) (8). C. felis was classified in the order: piroplasmsida, Family: Theileriidae (9), the genus Cytauxzoon has two stages in their life cycle: Erythrocytic stages, signet rings 1-1.5 μm in diameter, bipolar oval (safety pin) 2 μm in diameter and tetrad forms or anaplasmoid round “dots” less than 0.5μm in diameter (10). The second stage occur in the spleen, liver, and lymph nodes and the parasite proliferate in the mononuclear phagocytes under asexual reproduction, resulting obstruction of the blood flow (11,12). Which eventually cause host cell rupture and enter the blood, these intravascular merozoites infects variable numbers of erythrocytes (13). C. felis is believed to be spread by ticks bites of cats (12,14,15). The death are depended on the development of schizonts in mononuclear phagocytes rather than parasitemia (8).

H. felis is a gram negative epicellular parasites of feline Red Blood cells, the organisms was classified in the Family:Anaplasmataceae, but recently was shown to be more closely related to the Genus: Mycoplasma (16,17). Haemobartonella appear microscopically as cocci or short rods on the surface of red blood cells often completely surrounding the margin of the red cells and which are tightly attached to the red cells and are rarely free in the plasma (9). H. felis is a significant cause of hemolytic anemia in young cats recovered cats may remain as carriers (9,16,18). H. felis may be transmitted by a number of routes such as: injection of small volume of blood from an infected cats, blood sucking arthropods like fleas and ticks (16), (19) also reported the H. felis can be transmitted from queen (mother cats) to her kittens. The disease can run the spectrum from being very mild, asymptomatic, or a slight anemia, to avery sever form, (16,19). Because the fewer information about the prevalence of infection with these parasites in cats in Mosul city, therefore to aim of the present study was to determined the infection with Babesia spp, C. felis and H. felis in stray cats.

Materials and methods

In this study, 50 cats, in different ages group and both sexes were obtained from different areas of Mosul City from the period of March 2008 to March 2009, all examined clinically and signs were recorded before blood samples were taken from the cephalic vein during the postmortem, thin blood smears were prepared and stained with Giemza to identify the Babesia spp, C. felis and H. felis (20) and the procedures used for determination the percentage of parasitemia by measuring one thousand of RBC. Later the percentage of parasitemia in the RBCs were calculated according to the following formula (21):

\[
\text{Calculated RBCs} = \frac{\text{The infected RBCs}}{\text{Calculated RBCs}} \times 100
\]

Diagnosis of C. felis was made on the basis of the study of schizonts which is not present in Babesia spp and the H. felis which chains form, the techniques were used includes: Impression smear from liver, spleen (22), fine needle Aspirate of the spleen (23), histological examination of liver, and spleen, these organs fixed with 10% neutral buffered formalin, embedded in paraffin, sectioned 6µm thick and stained with hematoxylin and eosin (24). Differentiation of B. spp, C. felis, and H. felis also made on the basis of specific feature and by microscopic measurement by using ocular micrometer (9,25,26).

Results

In this study, some clinical signs were observed during blood sampling before PM and some pathological lesions when PM performed. The clinical findings included appearance of pale mucous membrane, weakness, rough hair coats, jaundice nasal discharge and diarrhea. PM diagnosis showed generalized pallor of different organs, enlarged lymph nodes, congested lungs, and petechial haemorrhage on the surfaces of abdominal organs and lungs, slight to marked enlargement of liver and spleen, the spleen was dark and the liver in orange-brown colour. The examination of blood smear revealed diagnosis of: B. spp, C. felis and H. felis, with percentage 26%, 22%, 42% respectively (Table 1).

Table 1: The percentage of infection with the B. spp., C. felis and H. felis in 50 examined cats.

<table>
<thead>
<tr>
<th>The parasite</th>
<th>Number of affected cats</th>
<th>The percentage of infection</th>
</tr>
</thead>
<tbody>
<tr>
<td>B. spp</td>
<td>13</td>
<td>26</td>
</tr>
<tr>
<td>C. felis</td>
<td>11</td>
<td>22</td>
</tr>
<tr>
<td>H. felis</td>
<td>21</td>
<td>42</td>
</tr>
</tbody>
</table>

Babesia spp. appeared in blood smear as small intracellular piroplasm, round, oval and in some blood smear appeared as pyriform shape and found within Red blood cells in a single or pairs (Fig. 1).

The sizes are between 1.33-2.15 μm within a mean 1.53 micron and the percentage of parasitemia was 0.6-2.4% with a mean of 1.4% (Table 2).
Table 2: The morphological features, microscopical measurements and percentage of parasitemia from *B. spp.*, *C. felis* and *H. felis*, in 50 blood smears of cats.

<table>
<thead>
<tr>
<th>The parasite</th>
<th>Morphological features</th>
<th>Measurements range (mean) µm</th>
<th>The percentage of parasitemia, the range (mean)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>B. spp.</em></td>
<td>Round, oval,</td>
<td>1.33-2.15 (1.53)</td>
<td>0.6-2.4 (1.4)</td>
</tr>
<tr>
<td></td>
<td>pyriform,</td>
<td></td>
<td></td>
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<tr>
<td></td>
<td>intracelluar</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>piroplasm</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>C. felis</em></td>
<td>Signet rings,</td>
<td>0.8-2.3 (1.64)</td>
<td>0.8-5.9 (2.99)</td>
</tr>
<tr>
<td></td>
<td>safety pin and</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>anoplasmoid</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>H. felis</em></td>
<td>Cocci, rods, chain</td>
<td>0.5-2 (1.15)</td>
<td>2.1-6.1 (3.78)</td>
</tr>
<tr>
<td></td>
<td>on the surface of</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>red blood cells</td>
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</tr>
</tbody>
</table>

*C. felis* diagnosed in the Red blood cells as round to signet rings, safety pin and anoplasmoid round shape, the cytoplasm of *C. felis* was stained light blue while the nucleus was stained dark red or purple in colour, the diameter ranged between 0.8-2.3µu within a mean of 1.64 µu and the Red blood cells appeared as containing one parasite of *C. felis* as well as in pairs or tetrads form (Fig 2). The percentage of parasitemia 0.8-5.9% within a mean of 2.99% (Table 2).

Fig (1): Intracellular piroplasms (small *B spp.*) in blood smear of cats. 1700X. (a) round form (more common), (b) pyriform (less common).

Fig (2): *C. felis* in blood smear of cat. 1400X. (a) signet ring, (b) safety pin, (c) anaplasmoid forms.
H. felis appeared in blood smears as small, blue staining cocci, rods and in some cases appeared as chain forming. These organisms attached to the external surface of the Red blood cells (Fig 3).

The sizes varied between 0.5-2 micron in diameter with a mean of 1.15 micron and the percentage of parasitemia was 2.1-6.1% with a mean of 3.78% (Table 2).

Impression smears from liver and spleen and fine needle aspirate from spleen showed schizont's of C. felis present in the cytoplasm of macrophages (in different replication) and contain a number of merozoites the nucleus of macrophages located at the bottom of the macrophages and in some cases has been disintegrated (Fig 4).

On histopathological examination the merozoites of C. felis appeared in the cytoplasm of phagocytes that are especially attached to the endothelium of vessels of liver and spleen in various stages of replication from the first development in the cytoplasm of macrophages until to demonstrating macrophages with schizont and merozoites in the parenchyma of liver and spleen. The parasite-laden macrophages identified free within the lumen or attached to the vessels leading in partial obstruction of blood vessels (Fig 5).

Fig (3): H. felis, located on the surface of RBCs in blood smear from a cat.1200X. (a) rod form, (b) small blue staining cocci, (c) small aggregates, (d) chain forming (uncommon).

Fig (4): Tissue stage of C. felis (a 1-2-3) Impression smear, spleen, Giemza stain, macro phages contain schizonts and merozoites of C. felis in different replication. 1200X, (b) fine needle aspirate, spleen, Giemza stain, (c) Impression smear, liver, Giemza stain (large macrophages contain large number of C. felis merozoites and the nucleus of the
macrophages is eccentrically located at the bottom of the cells). 1400X.

![Figures](image)

**Fig (5):** Histopathological examination of tissue stage of *C. felis*. (a) Early schizonts of *C. felis* development within macrophages in the liver. H & E. 1400x, (b) the merozoites of *C. felis* within the macrophages and the nucleus of the cell located at the bottom edge (ghost of nucleus). liver. H & E. 1400x, (c) Demonstrating macrophages with schizonts of *C. felis* and merozoites in the parenchyma. liver. H & E. 1200x, (d) Tissue stage of *C. felis* within macrophages which located near to the wall of blood vessels. liver. H & E. 1200x, (e) Partial obstruction of blood vessels by parasite– laden macrophages. liver. H&E. 1200x.

**Discussion**

There are a little information and studies about stray cats, which lives around and contaminate the surrounding, therefore, they may be affected with many different parasitic diseases, such as infection with blood parasites. In this study blood samples were collected and postmortem performed, certain clinical signs and pathological lesions were observed in cats who was microscopically positive for *B. spp, C. felis* and *H. felis*, in addition that these cats were affected with other parasites such as different internal parasites, (27) referred that feline babesiosis often present as chronic low-grade disease and the most clinical findings of symptomatic feline babesiosis are anorexia, lethargy, weakness and a rough hair coat.

The clinical signs which observed in cats with cytauxzoonosis are non specific, no ticks were found and mortalities was nil, in this study, some cats showed splenomegaly, hepatomegaly and the lungs were congested. The results were in agreement with (8) who demonstrated the presence of *C. felis* in Van cats in Turkey and revealed that preimmunity against the parasite may prevent the observation of the clinical sings. Recent studies indicate historically, cytauxzoonosis has been nearly 100% fatal in domestic cats, increasing number of reports of infected cats which demonstrate less and sever disease suggest the existence of different strains of *C. felis* (28).

In naturally affected cats with haemobartonellosis, the clinical signs occur in immunocompetent or immunosuppressed cats and depend on the degree of anemia, stage of infection and the immune status of infected cats (16). Some of the cats recovered from this disease to become carries for the *H. felis* this mean that some cats look healthy but still have small numbers of these parasites in their bodies, if these cats are stressed, it sometimes causes parasites to multiply and produce disease (19). In the infection with *H. felis* the gross necropsy findings are not pathognomonic, but splenomegaly is common and mesenteric lymph nodes may be enlarged (15).

In present study, the percentage of infection with *B. spp* was 26%, (27) who referred that feline babesiosis appear to be less common. Sporadic cases of babesiosis has been reported from various countries such as France, Germany, Thailand, and Zimbabwe (3-6) while studies in South Africa indicate a higher prevalence of *B. felis* infection has been observed in Siamese and Oriental cats (1). In endemic area many cats are exposed to the parasites at any early age, it appears that some of these cats may become chronic asymptomatic carrier, concurrent illness or stress contributes to the exacerbation of clinical babesiosis in carrier cats (1,29).

The mean size of *B. spp* which diagnosed in this study was 1.53 µm and the mean percentage of parasitemia was 1.4%. Classification were done according to the size and morphology of felin *B. spp*, in this study, it concluded that the *B. spp* which described as a small type, (29) who classified feline piroplasms morphologically as small piroplasms < 1.5 µm or as large piroplasms > 2.5 µm. Small feline piroplasms include *B. felis, B. leo* and *Cyttauxzoon* and large feline piroplasms include, *B. cati, B. pantherae, B. canis canis, B. canis presenti* and *B. herpailuri* while in
France, Germany, Thailand, and Zimbabwe un characterized spp has been reported (3–6), (30) revealed that the detection of this organisms in blood smear may be difficult because the level of parasitemia may be low and (1) showed that the feline babesiosis often present as chronic low grade disease.

In this study, during the examination of 50 stray cats, *C. felis* was diagnosed in 11 cats with percentage 22%, by using thin blood smear, impression smear, fine needle aspiration and histological examination, in compared with a study in Van region in Turkey, depending on the examination of 120 blood smears of cats only 9 had *C. felis* with percentage 7.5%. On microscopic examination of blood smears of cats, multiple Red Blood cells had *C. felis* which appear in different forms like signet ring, safety pin and anaplasmoïd form, the mean size of *C. felis* was 1.64, this results was agreement (12,15), they considered that *C. felis* was small felin piroplasms while (31) referred that these bodies ranged between 1.5 µm in diameter. The mean percentage of parasitemia with *C. felis* which recorded in this study was 2.99%, (32) referred that the piroplasms appeared in affected cells from only a few perfilm to 25% before death while the (13) referred that the parasitemia of *C. felis* in the RBCs is low and the number of Red Blood cells parasitized with *C. felis* varies among cats and with the stage of disease, (31) recorded that up to 25% of Red Blood cells may be affected with *C. felis*. In impression smears from liver and spleen and with fine needle aspiration from spleen, showed many macrophages contain large number of schizonts with innumerable *C. felis* merozites and these cells appeared especially at the feathered edge of peripheral smears, this results was agreement with (12,32).

On histopathological examination large number of parasitized macrophages containing large numbers of merozites in different stages of replication free within the lumen of the organs (liver, spleen) or on the walls of the veins these vessels appeared partially or completely occluded this results were in agreement with (12,33,34).

In present study, the percentage of infection with *H. felis* was 42%. (22,35) which they referred that many cats are carrier to *H. felis* while (36) showed that the prevalence of this parasites in the feline population from 0.9% to 28% and the cats without door were more likely to be infected with Haemobartonella. The result of this study revealed that *H. felis* appeared in cocci, rod, chain form on the surface of Red blood cells and the mean size was 1.15 µm this is in agreement with (9,25), the meanpercentage of parasitemia of *H. felis* was 3.78%, (15) referred that the number of RBCs affected varies with the severity of the infection and the stage of life cycle of the parasite. In acute phase numbers of *H. felis* increase gradually than disappear rapidly, while in chronically infected cats, the parasites appear only sporadically and in small numbers, and the parasitemia of *H. felis* is transient or cyclical. The peak of parasitemia may last for few days but occurs just prior to the hemolytic episode, by the time the infected cats is presented, very few or no parasite may be detected on blood film. Thus a single negative findings is not adequate evidence for freedom from infection, it is necessary therefore to examine a number of blood film prepared over a period of 4-10 days, before it may be concluded that the patient is free from infection (37).

Reference


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