Phenotypic and pathological characterizations of different *Sarcocystis* species in donkeys

**Abstract:**

This study aimed to record the prevalence of different *Sarcocystis* spp., demonstrate their morphological characters by laboratory, histopathological and immunohistochemical techniques, also to investigate the pathological lesions in esophagus, heart, and brain samples collected from naturally infected donkeys in Egypt. The current study identified four *Sarcocystis* spp. (*S*. *bertrami*, *S*. *equicanis*, *S*. *asinus* and *S*. *fayeri*) in meat samples and *S*. *neurona* in the brain of the examined donkeys. Examination of meat samples showed macroscopic tissue cysts of *S*. *bertrami* and microscopic tissue cysts of *S*. *equicanis*/*S*. *asinus* and *S*. *fayeri*. According to the thickness of the cyst wall, *Sarcocystis* spp. were divided into smooth thin walled *S*. *bertrami*, ciliated thin walled *S*. *equicanis*/*S*. *asinus* and striated thick walled *S*. *fayeri*. Also, immunohistochemical (IHC) examination showed strong labeled schizonts and cyst wall of *S*. *neurona*, while the bradyzoites appeared negative staining. Pathological examination revealed degeneration of muscles with intermuscular edema and few mononuclear inflammatory cells, but brain tissue showed perivascular lymphocytic cuffing and satellitosis with developing schizonts. The current study concluded that the accurate recognition about *Sarcocystis* spp. in donkeys would be helpful in their treatment, prevention and control measures.

**Keywords:** *bertrami*, donkey, *equicanis*, *fayeri*, immunohistochecmical, *neurona*.

**INTRODUCTION**

*Sarcocystis* spp. cysts were noticed by Miescher (1843) for the first time in mouse muscles in Switzerland. Such nomenclature was derived from finding them encysted (cyst) and localized in the muscles (sarco) of the intermediate hosts Dubey et al. 2016a; Dubey et al. 1989. The cysts were formed through endodyogeny of merozoites and could reproduce through endodyogeny (production of metrocytes with a pair of...
daughter nuclei) and endopolygeny (production of multinucleated metrocyes) Slapeta et al. 1999. Sarcocystis spp. is a common protozoan parasite and distributed worldwide in muscles of mammals and birds. Four species; S. bertrami, S. equicanis/S. asinus, S. fayeri and S. neurona were recorded in equine Levine and Tadros 1980. Dogs act as final hosts for S. bertrami, S. equicanis/S. asinus, while opossums act as final hosts for S. neurona Dubey et al. 2016a.

S. neurona investigated in a wide range of hosts. If their schizonts were detected in the host, it is considered as an aberrant host, while if the mature cysts were investigated, the host is considered intermediate Dubey et al. 2015. The schizonts had been found only in brain Dubey et al. 2001, but their cysts might be found in brain, cardiac or skeletal muscles Miller et al. 2009. S. neurona is the most pathogenic species in equine where it induce equine protozoal myelonecephalitis (EPM). The manifested clinical signs vary according to the involved area in CNS from weakness, facial paralysis, dysphagia, and gait abnormalities to seizures Dubey et al. 2001; Dubey et al. 2015. Also, Aleman et al. 2015 reported that S. fayeri infection in equine induced eosinophilic myositis and cardiomyopathy, which manifested by weight losses, stiffness, lethargy, muscle pain, swollen tongue and dysphagia.

Determination of different Sarcocystis spp. in equines is still confused Dubey et al. 2016a. They included S. bertrami, S. equicanis, S. fayeri and S. neurona. Zeng et al. 2018 found that Sarcocystis spp. in horses and donkeys were closely related based upon their phenotypic characters and coxl sequences and so they suggested both horses and donkeys had the same Sarcocystis spp. Many authors studied sarcocystosis in horses, but little data was known about donkeys and so this study is dealing with Sarcocystis spp. prevalence, morphological characterization of different Sarcocystis spp., and pathological findings in meat and brain samples of naturally infected donkeys in Egypt.

**MATERIAL AND METHODS**

1. **Samples collection**

Meat and brain samples were collected from 80 and 20 slaughtered donkeys, respectively in Giza Zoo, Giza Governorate, Dokki, Egypt. Samples were transported in an ice tank to Zagazig University, Faculty of Veterinary Medicine, Parasitology Department for further examinations. This study was approved with number ZU-IACUC/2/F/75/2018 by ZU-IACUC Committee, Zagazig University, Egypt

2. **Laboratory examination**

By naked eye, meat samples were examined for macroscopic cysts. Small pieces of meat were crushed between two glass slides
and examined under the light microscope to detect microscopic cysts. Also, small pieces of brain were squeezed on glass slides and then stained with Giemsa stain to observe the released bradyzoites or metrocytes under the oil immersion lens.

3. **Pathological examination**

Meat and brain samples were collected from slaughtered donkeys for detection of *Sarcocystis* spp. The collected specimens were fixed in 10% buffered neutral formalin solution, processed, stained and prepared for microscopical examination according to the descriptions of Suvarna et al. 2013.

4. **Immunohistochemical (IHC) examination**

The de-paraffinized sections of brain and meat samples were incubated with polyclonal goat-antibodies against *S. neurona* (VMRD, Inc. Pullman, WA) Soldati et al. 2004 and A streptavidin-immunoperoxidase labeling (Sigma Chemicals, St. Louis, US) was used to distinguish positive immune reactions Kiupel et al. 2003. Positive control was prepared from positively infected tissue sections incubated with polyclonal antibodies against *S. neurona*, while the negative control was prepared from sections incubated with non-immune sera. The procedures were completed as the description of Dubey et al. 2016a. The measurements of length, width, and cyst wall thickness of *Sarcocystis* spp. were performed by a calibrated eye piece micrometer.

**RESULTS**

1. **Prevalence**

Visual and light microscopical examination of fresh and stained sections (H&E or IHC) of meat and brain samples showed detection of *S. bertrami*, *S. equicanis/S. asinus*, *S. fayeri*, and *S. neurona*. Their prevalence were listed in Tables (1-3).

2. **Morphological identification**

Examination of meat samples showed macroscopic tissue cysts of *S. bertrami* and microscopic tissue cysts of *S. equicanis/S. asinus* and *S. fayeri*. The mature cysts appeared septated, spindle-shaped, contained numerous bradyzoites and surrounded by cyst wall. According to the thickness of the cyst wall, *Sarcocystis* spp. were divided into smooth thin walled *S. bertrami*, ciliated thin walled *S. equicanis/S. asinus* and striated thick walled *S. fayeri*. In *S. fayeri*, the immature cysts contained numerous metrocytes and surrounded by granular cyst wall. The stained brain smears showed developed metrocytes with granular basophilic nucleus. The morphological identification of *Sarcocystis* spp. in donkey’s meat was recorded during light microscopical examination of fresh and stained sample sections with H&E (Fig. 1&2).

3. **Immunohistochemical (IHC) examination**

The deparaffinized brain samples from donkeys are the only
Sarcocystis spp. positive immune reactions against polyclonal antiserum of S. neurona. The irregular-shaped schizonts, the cyst wall and the septa of mature cysts appeared strongly labeled, while the bradyzoites showed negative staining (Fig. 2). The measurements of different Sarcocystis spp. in the examined samples from brain and meat of donkeys were recorded and showed in Table 4.

4. Pathological examination

The heart exhibited multiple basophilic mature cross and elongated intra muscular cysts containing banana shape bradyzoites. Few mononuclear inflammatory cells were present around some cysts. Acute degeneration of the cardiac muscle and intermuscular edema were detected. Hyaline thickening of the blood vessel with endotheliosis were observed (Fig. 3).

The examined oesophagus tissues showed microscopic, elongated and cross cysts containing basophilic bradyzoites. Immature cysts were also observed between the muscles. Degeneration of muscles with intermuscular edema was detected (Fig. 4).

The examined brain tissues revealed perivascular lymphocytic cuffing with scattered glia cells in the white and gray matter (Fig. 5A). Satellitosis and demyelination were detected in the white matter (Fig. 5B). Encophalomalacia represented by depletion of certain part of brain tissue with gliosis were observed (Fig. 5C). Developing schizonts were noticed (Fig. 5D).
Table 1: Total prevalence of *Sarcocystis* spp. in the examined donkeys

<table>
<thead>
<tr>
<th>Item</th>
<th>Examined numbers</th>
<th>Positive numbers</th>
<th>Positive %</th>
</tr>
</thead>
<tbody>
<tr>
<td>Meat samples</td>
<td>80</td>
<td>67</td>
<td>83.75% (67/80)</td>
</tr>
<tr>
<td>Brain samples</td>
<td>20</td>
<td>6</td>
<td>30% (6/20)</td>
</tr>
<tr>
<td>Total</td>
<td>100</td>
<td>73</td>
<td>73% (73/100)</td>
</tr>
</tbody>
</table>

Table 2: Prevalence of single and mixed infections with muscular *Sarcocystis* spp. in donkeys

<table>
<thead>
<tr>
<th>Infection type</th>
<th>Positive %</th>
<th>Single</th>
<th>Mixed</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>83.75 (67/80)</td>
<td>25.37% (17/67)</td>
<td>74.63% (50/67)</td>
</tr>
</tbody>
</table>

Table 3: Prevalence of different *Sarcocystis* spp. in the examined donkeys

<table>
<thead>
<tr>
<th>Item (numbers)</th>
<th><em>S. bertrami</em></th>
<th><em>S. equicanis</em></th>
<th><em>S. fayeri</em></th>
<th><em>S. neurona</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Prevalence</td>
<td>80.60% (54/67)</td>
<td>85.07% (57/67)</td>
<td>50.75% (34/67)</td>
<td>30% (6/20) for schizonts 5% (1/20) for cysts</td>
</tr>
</tbody>
</table>

Table 4: Measurements of different *Sarcocystis* spp. in the examined donkeys

<table>
<thead>
<tr>
<th>Item</th>
<th><em>S. bertrami</em></th>
<th><em>S. equicanis</em></th>
<th><em>S. fayeri</em></th>
<th><em>S. neurona</em></th>
</tr>
</thead>
<tbody>
<tr>
<td>Length (mm)</td>
<td>5-8mm</td>
<td>0.5-2.3 mm</td>
<td>450.8-600 µm</td>
<td>175µm</td>
</tr>
<tr>
<td>Width (mm)</td>
<td>0.4-1mm</td>
<td>50-150 µm</td>
<td>47.2-75.5 µm</td>
<td>40 µm</td>
</tr>
<tr>
<td>Cyst wall thickness (mm)</td>
<td>0.3-0.8 µm</td>
<td>0.7-1.5 µm</td>
<td>3.4-6.5 µm</td>
<td>0.5 µm</td>
</tr>
</tbody>
</table>
Fig. 1: A&B: Photographs showing macroscopic cysts of *S. bertrami* in donkey esophagus (arrow); C: Fresh cysts of *S. bertrami* under light microscope (X40); D: higher magnification of *S. bertrami* showing well developed septa (X100); E: Fresh cysts of *S. equicanis/S. asinus*(X100) and F&G: Fresh cysts of *S. fayeri* demonstrated the thick cyst wall (arrowheads, X100, Digital camera).

Fig. 2: Histopathological sections stained with H&E for *Sarcocystis* spp. in donkeys. A: longitudinal section of *S. bertrami* with clear septa and thin walled
cyst (X100); B: Cross section of *S. equicanis* showing ciliated and thin walled cyst (X100); C: Longitudinal section of *S. fayeri* mature cyst in meat samples showing thick and striated cyst wall (X100); D: Cross section of *S. fayeri* mature cyst (X100); E: Cross section of *S. fayeri* developed cyst (X400); F: Cross section of *S. fayeri* immature cyst contained metrocytes and surrounded by granular cyst wall (X400); G: stained brain smear with Giemsa showing developed metrocytes with granular basophilic nucleus (X1000); H&I: Strong labeled schizonts of *S. neurona* obtained by IHC (X400) and J: IHC stained brain section showed strong labeled cyst wall and negative stained bradyzoites of *S. neurona* in donkeys ((X1000, Digital camera).

**Fig.3:** A: Heart showing multiple basophilic mature intramuscular cysts (arrow) between degenerated cardiac muscle (arrow head) with intermuscular edema (oblique arrow, bar=100); B: Heart showing cross section of *S. fayeri* (arrow) with acute muscle degeneration (arrow head, bar=20); C: Heart showing *S. equicanis* (arrow) among degenerated muscles (white arrow) with few number of mononuclear cells around cyst (arrow head, bar=20) and D: Heart showing two thin walled *S. equicanis* intramuscular cyst (arrow) with intermuscular edema (arrow head) and hyaline thickening of blood vessel wall together with endotheliosis (oblique arrow, bar =100).
Fig. 4: A: Oesophagus showing elongated *S. equicanis* between muscle fibers (arrow) with intermuscular edema (arrow head, bar=100); B: Higher magnification of *S. equicanis* cyst (arrow) with degenerated muscle fibers (white arrow) and intermuscular edema (arrow head, bar=20); C: Oesophagus showing *S. equicanis* (arrow) with myofiber degeneration (arrow head, bar=20) and D: Oesophagus showing small sized developed cyst of *S. fayeri* (arrow) among degenerated muscles (arrow head, bar =20).

Fig. 5: A: Brain showing perivascular lymphocytic cuffing (arrow) with scattered glia cells in white and gray matter (arrow head, bar=20); B: Brain
showing satellitosis (arrow) and demyelination in the white matter of brain tissue (arrowhead, bar=20); C: Brain showing encophalomalacia (arrow) with gliosis bar=20 and D: Brain showing developed merozoites (arrow

Discussion

The current study revealed that the total prevalence of Sarcocystis spp. in donkeys reached 73% by laboratory and IHC examination, while that obtained by Abdel- Maogood et al. 2015 reached 12% and 32% by PM examination and ELISA technique, respectively in Egypt. Also, it reached 66.66 % in Egypt by Attia et al. 2018, 16.6% in Ethiopia by Woldemeskel and Gebreab, 1996.

Regarding S. bertrami, the recorded prevalence was 80.60%, while in Italy it was 28.57% Passantino et al. 2019. The current study showed prevalence of 85.07% for S. equicanis, nearly similar rate was previously recorded in Egypt by Hilali and Nassar 1987 to be 90%. This study is considered the first record of the prevalence of S. fayeri in donkeys which reached 50.75%. In addition, the current study showed that the prevalence of 30% for S. neurona by IHC technique, while it was 2.5% by ELISA in Mexico Alvarado-Esquivel et al. 2017, 61.1% by immunoblot in USA by Saville et al. 1997, 3% and 21% by indirect fluorescent antibody test (IFAT) and direct agglutination tests, respectively by Gennari et al. 2016 in Brazil.

The differences in the prevalence rates is not only due to different diagnostic methods (PM examination, ELISA, IFAT or IHC) or climatic changes, but also due to the differences in intermediate host species (horse or donkey), host susceptibility to infection, and exposure to the final hosts (dogs for Sarcocystis spp. in meat or opossums for S. neurona). The more opportunity of donkeys became in contact with final hosts, the more chance to acquire infection with Sarcocystis spp. Alvarado-Esquivel et al. 2017; Dubey et al. 2016a.

The obtained morphological descriptions for S. bertrami, S. equicanis/S. asinus, S. fayeri were similar to those noticed by Aleman et al. 2015; Dubey et al. 1989; Hilali and Nassar 1987; Saville et al. 2004; Zeng et al. 2018. Zeng et al. 2018 found that horses and donkeys had the same Sarcocystis spp. They had been included S. bertrami, S. equicanis, S. fayeri and S. neurona. The previous study of Dubey et al. 2016b on donkey’s sarcocytosis in Egypt recorded only S. bertrami in their meat, while the current study recorded four Sarcocystis spp. The current study described S. bertrami in donkeys as macroscopic cysts in spite of the previous description of S. bertrami in donkeys by Dubey et al. 2016b in Egypt as microscopic cysts. They also stated that smooth and ciliated parts were noticed in the same cyst of S. bertrami, while the currently studied appeared smooth thin walled macroscopic cysts in S. bertrami, ciliated microscopic thin walled cysts in S. equicanis/S. asinus and striated...
thick walled microscopic cysts in *S. fayeri*. *S. equicanis* cysts were noticed in the esophageal and cardiac muscles of donkeys, however inability of Sakran et al. 2013 to detect them in the cardiac muscles of horses in Egypt. Zeng et al. 2018 described 2 types of *S. bertrami* in horses and donkeys in China, where Type I cysts were large without protrusions (smooth) and Type II cysts were smaller with protrusions (ciliated) and referred that to the Type I might be aged cysts. However, Sheffield et al. 1977 stated that Sarcocystis spp. cysts increased in size with age, but their wall structure remained stable once the cysts become mature (contained bradyzoites). This confirmed our record for *S. bertrami* and *S. equicanis* as different species and they have not the same description as the previous ones Dubey et al. 2016a; Odening et al. 1995; Rommel and Geisel 1975. Also, we suggested that the previously described *S. equicanis* in horses is a synonym to the currently obtained *S. asinus* in donkeys depending upon their similarity with morphological characters.

In the current study, the cysts in brain were difficult to be identified in sections stained with H & E, but it was more easily noticed in sections reacted against polyclonal *S. neurona* antibody. Also, the schizonts were more easily identified in sections reacted against polyclonal *S. neurona* antibody than those in histological sections. These findings were similar to those of Lindsaya et al. 2000. The noticed morphological features of *S. neurona* in brain were agreed with those described by Seguel et al. 2019 in California sea lions. The observed variability in staining pattern between *S. neurona* cyst wall and bradyzoites resembled the recorded observations of Butcher et al. 2002 in domestic cats and Miller et al. 2009 in southern sea otters. Sakran et al. 2013 reported that *S. neurona* schizonts were observed only within the neural cells of brain in horses, while the current study observed both schizonts and cyst in brain of donkeys by IHC technique. Alvarado-Esquível et al. 2017 reported the seroprevalence (2.5%) of *S. neurona* in donkeys in Mexico, while the current study reported them for the first time in the brain of donkeys in Egypt. As mentioned in a previous study by Dubey et al. 2015 that the presence of mature cysts and schizonts in the brain of the same host indicated it as I.H. and not an aberrant host and so from our obtained results, the donkeys were considered I.H. for *S. neurona*, however horse considered an aberrant host Ma et al. 2020.

*Sarcocystis* spp. had been associated with myocarditis reaction. Hyaline thickening of the blood vessel with endotheliosis were observed due to localization of their schizonts in the endothelia of blood vessels (Yarim et al. 2004). In the current study, the heart and esophagus exhibited multiple basophilic mature round and elongated intra muscular cysts containing banana shape bradyzoites. Some cysts had thin wall and other had thick wall. These findings were completely agreed with Faghiri et al. 2019 who observed thin and thick walled cysts inside the cardiac and esophageal muscles. Also the obtained results detected few mononuclear inflammatory cells were present around the cysts. Acute
degeneration of the cardiac muscle and intermuscular edema were detected. These findings were completely agreed with Caspari et al. 2011 who observed multifocal degeneration and necrosis of myocardial fibers with interstitial edema. Microscopical features of the brain revealed perivascular lymphocytic cuffing with scattered glia cells in the white and gray matter. Satellitosis and demyelination were detected in the white matter. Encophalomalacia represented by depletion of certain part of brain tissue with gliosis were observed. The developing schizonts of Sarcocystis spp. were also detected. These findings were partially agreed with (Özmen et al. 2009) where their histopathological examinations showed focal areas of perivascular cuffing in the brainstem, cerebellum, and medulla spinalis with presence of Sarcocystis spp. schizonts.

The current study concluded that the donkeys were considered I.H. for S. neurona in brain and for S. bertrami, S. equicanis/S. asinus, and S. fayeri in meat. Also, S. equicanis in horses is a synonym for S. asinus in donkeys. As the donkeys had great economic importance in Africa, the accurate recognition about Sarcocystis spp. in donkeys would be helpful in performance of prevention and control measures against future infections.

References


المملوح العربي

الخصائص المظهرية والنسيجية لمختلف أنواع الساركوسومستيس في الحمير

على محمد إبراهيم عبد الرحمن – ** هبه محمد عبدالغنى

قسم الطفيليات و ** الباثولوجيا كلية الطب البيطري جامعة الزقاق.

درس العديد من المؤلفين السابقين طفيل الساركوسومستيس في الحيوان، ولكن لا يتوفر سوى القليل من الدراسات عنها في الحمير ولذلك فقد هدفت هذه الدراسة إلى تسجيل معدلات انتشار الأنواع المختلفة من طفيل الساركوسومستيس وتوضيح خصائصها المورفولوجية من خلال التقنيات المخبرية والفحص الباثولوجي والكيميائي المناعي وكذلك دراسة التغيرات النسيجية الباثولوجية في عينات المريء والقلب والدماغ التي تم تجميعها من الحمير المصابة طبيعيا في مصر.

وقد تمكنت الدراسة الحالية من التعرف على أربعة أنواع من ساركوسومستيس بيرترامي و فاييري و أكوي كانيز في عضلات الحمير و اضافة النوع نيرونا في دماغ الحمير المصابة وقد تم تصنيفها طبقاً لمسك حاجز الحويصلات حيث كانت ذات جدران رقيقة ناعمة للنوع ساركوسومستيس بيرترامي و رقيقة ذات اهداف في النوع اكوي كانيز و سميكه في ساركوسومستيس فاييري وقد أوضح الفحص الكيميائي المناعي للنوع نيرونا تفاعل إيجابي لكل من الفيرونين والجذور الخارجية للحويصلات بينما البراديزويت كانت ذات تفاعل سلبي. وقد أوضح الفحص النسيجي عن انحلالات العضلات ذات الوذمة العضلية وعدد قليل من الخلايا الالتهابية أحادية النواة ولكن أنسجة الدماغ أظهرت تضخم المفاوتات حول الأوعية الدموية وداء الامطار الصناعي وبالتالي فإن الدراسة الحالية تستخلص أن الحمير تعد عائل وبيئي للنيرونا على عكس الحيوان ولكن أيضاً قد يشكل التعرف الدقيق على أنواع حويصلات الساركوسومستيس أهمية كبيرة في تحديد العلاج اللازم وكذلك الوقاية منها والسيطرة عليها.