Pathological and molecular study of ovine diaphragms naturally infected by Sarcosystis spp.

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Article information

Abstract

Sarcosystis spp., has a close relationship with muscles due to its unique localization within skeletal muscle in humans and the animals it infects, as the chronic condition of the disease causes significant economic losses, especially in terms of meat production as a result of the formation of cysts, whether macroscopic or microscopic, in their muscle fibers. Sarcosystis tenella and Sarcosystis arieticanis are the most important pathogenic cysts forming in sheep. In this study, 50 samples of diaphragm muscles of sheep slaughtered in the butchers' shops and the Mosul abattoir were examined grossly, histologically, and using PCR technique as a diagnostic tool to identify or diagnose the causative and responsible species of these changes.

The results of the molecular analysis using the nested PCR technique indicated that these diagnosed microscopic cysts belong to Sarcosystis tenella with a reaction product of 800bp and 500bp.

Keywords: Sarcosystis species, Sheep, PCR, Muscle, Sarcosystis tenella

Eosinophilic Myositis (EM) is a general term which used in meat examination to identify diseases in animals clinically appeared healthy with muscle lesions of unknown cause (5).

In Bovine Eosinophilic Myositis (BEM) are a heterogeneous group of muscle disease caused by an autoimmune response, allergic reaction or drugs or by bacterial infection Sarcosystis spp. is consider one of the factors causing this pathology (6).

Although Sarcosystis are well adapted to live in the muscles and central nervous system without any reaction from the host, some Sarcosystis rupture from time to time, producing toxic substances. (7).

The results of several re-searchers indicate that Sarcosystis parasite has a direct relationship with EM as it...
contributes to rejection and decrease in carcass quality in meat processing plants (5). One difficulty with Eosinophilic myositis that it cannot be detected in live animals as there are no tests to diagnose or detect this pathology while the animal is alive (8). Generally, EM occurs only in striated muscles and does not appear in internal organs or smooth muscles, and in the case of mild lesions in the carcass the most active muscles are affected, such as muscles of tongue, heart, and diaphragm. In severe cases, all striated muscles are affected (9).

Hence, the aim of our study design to diagnose the pathological changes in the diaphragm muscles of sheep that infected with *Sarcosystis* and to determine particular microscopic *Sarcosystis* species that causes the eosinophilic myositis by using molecular techniques.

**Materials and methods**

**Sample collections**

Diaphragm muscles were collected from butcher shops and abattoirs in Mosul city, Iraq during the period from June - October of 2019, were investigated for *Sarcosystis* infection. During postmortem inspection of slaughtered animals, tissue samples were isolated from 50 sheep. The samples were collected and stored in plastic bags. Where work was done in the Laboratory of Parasitology, College of Veterinary Medicine, University of Mosul.

**Gross examination**

The diaphragm muscles were examined grossly for detecting gross pathological changes.

**Microscopic examination**

Unstained samples of diaphragm muscles were examined by light microscope for detection of the microscopic *Sarcosystis* species (1,2). Approximately 2-3 mm² of muscles were taken and squashed by two slides and then examined under 40X objective lens.

**Histopathological examination**

Parts of diaphragm muscles from each sample were fixed in 10% neutral buffer formalin at least 48 hours. The fixed muscles samples were trimmed into 0.5 cm³ thick section dehydrated with serial dilutions of ethanol and xylene, then embedded in paraffin block for sectioned to 3-5 mm, then stained with hematoxylin and eosin (H&E) and Masson’s trichrome examined for detection of microcysts with light microscope (10-12).

**Extraction of DNA**

DNA was extracted by using Prime Prep Genomic DNA extraction kit (Genet Bio, Korea). From 11 diaphragm muscle samples following the manufacturer instruction, the DNA pellet was rehydrated by adding 100 µl of rehydration solution and kept at -20°C until further assay.

**Polymerase chain reaction (PCR) assay procedure**

PCR was done to confirm the presence of *Sarcosystis tenella* and *Sarcosystis arieticanis* using the primers according to Pipia et al. (13) (Table 1). The PCR reaction mixtures were prepared in 25 µl containing 12.5 µl of HS Prime Taq Premix (2X) (Genet Bio, Korea) with final concentration of 1X. 1 µl of each primer and 3 µl of DNA template and 7.5 µl of PCR grade water. The PCR was done using thermal cycler (T100, Bio-Rad, USA) and PCR cycles were performed as shown in (Table 2-4). The amplified products were separated using electrophoresis in 1.5% agarose gel (Jena Bioscience, Germany) pertained with 4 µl GelSafe (GeNet Bio, Korea). A 4 µl of each PCR product was loaded into the well of agarose gel. The electrophoresis was carried out at 80 V for 1 hour using power supply MP 300V (Bio-Rad, USA) containing 1X TBE buffer (GeNetBio, Korea).

A 100 bp DNA marker, 4 µl (Genet Bio, Korea) was used as standard molecular weight marker. The gel was examined under UV light using Gel doc Ez system (BioRad, USA) (13).

<table>
<thead>
<tr>
<th>No.</th>
<th>Primer Name</th>
<th>Primer Sequence 5' - 3'</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ST1-F</td>
<td>GGATCGCATTATGGTCTA</td>
<td>External primers</td>
</tr>
<tr>
<td>2</td>
<td>AP2-R</td>
<td>CCCGGATCCAAGCTTGATCTTCTGCAGGTTCACCTAC</td>
<td>Nested primers</td>
</tr>
<tr>
<td>3</td>
<td>8-F</td>
<td>TTTGACTCAACACGGG</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>ST3-R</td>
<td>CGTTGCGCGCGGTAA</td>
<td></td>
</tr>
</tbody>
</table>

**Table 2: Sequences of primers used for detection of *Sarcosystis arieticanis* using nested PCR**

<table>
<thead>
<tr>
<th>No.</th>
<th>Primer Name</th>
<th>Primer Sequence 5' - 3'</th>
<th>Type</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>STA-F</td>
<td>TTT CGC AAC GAA GAG GA</td>
<td>External primers</td>
</tr>
<tr>
<td>2</td>
<td>SA2-R</td>
<td>TGA AAC GGG CGG TAG A</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>2-F</td>
<td>AGG GTT CGA TTC CGG AG</td>
<td>Nested primers</td>
</tr>
<tr>
<td>4</td>
<td>SA1-R</td>
<td>GCG GGA AGA GGA GAA T</td>
<td></td>
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Table 3: Cycling conditions of PCR for amplification of *Sarcosystis*

<table>
<thead>
<tr>
<th>No.</th>
<th>Step</th>
<th>Temp</th>
<th>Time</th>
<th>Cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Polymerase activation</td>
<td>95°C</td>
<td>10 min</td>
<td>1X</td>
</tr>
<tr>
<td>2.</td>
<td>Denaturation</td>
<td>95°C</td>
<td>45 sec</td>
<td>35X</td>
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<td>3.</td>
<td>Annealing</td>
<td>52°C</td>
<td>45 sec</td>
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<tr>
<td>4.</td>
<td>Extension</td>
<td>72°C</td>
<td>75 sec</td>
<td></td>
</tr>
<tr>
<td>5.</td>
<td>Final Extension</td>
<td>72°C</td>
<td>5 min</td>
<td>1X</td>
</tr>
<tr>
<td>6.</td>
<td>Cooling</td>
<td>4°C</td>
<td>5 min</td>
<td>1</td>
</tr>
</tbody>
</table>

Table 4: Cycling conditions of PCR for amplification of *Sarcosystis tenella* and *Sarcosystis arietianis* using internal primers

<table>
<thead>
<tr>
<th>No.</th>
<th>Step</th>
<th>Temp</th>
<th>Time</th>
<th>Cycle</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>Polymerase activation</td>
<td>95 °C</td>
<td>10 min</td>
<td>1X</td>
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<tr>
<td>2.</td>
<td>Denaturation</td>
<td>95 °C</td>
<td>45 sec</td>
<td>35X</td>
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<tr>
<td>3.</td>
<td>Annealing</td>
<td>51 °C</td>
<td>45 sec</td>
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<td>4.</td>
<td>Extension</td>
<td>72 °C</td>
<td>1.15 min</td>
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<tr>
<td>5.</td>
<td>Final Extension</td>
<td>72 °C</td>
<td>5 min</td>
<td>1X</td>
</tr>
<tr>
<td>6.</td>
<td>Cooling</td>
<td>4 °C</td>
<td>4 °C</td>
<td>1</td>
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Results

Grossly the muscles of sheep diaphragm appeared pale, white to gray in color with white streak and no macroscopic cysts were observed. The histological examination of diaphragm muscles of sheep revealed presence many sarcocysts with different size, shape (oval and spherical) and blue in color which embedded in the muscle fibers with marked dispersing edema and sever fragmentation of muscle fibers especially nearing the cysts, some of sarcocysts were ruptured and released bradyzoites in the surrounding tissues (Figure 1).

![Figure 1](image1.png)

Figure 1: Presence of sarcosystis in diaphragm muscles with different shapes and sizes (A), sever distraction of muscle fibers (B) and edema between muscle fibers (C). H&E (145X).

Obvious Zenker’s necrosis of myocytes with slightly cellular infiltration of inflammatory cells especially eosinophils monocytes and macrophages, were seen in the tissue (Figure 2).

Activation of fibroblast between muscle fibers, so the normal architecture was replaced by fibrous tissue (Figure 3 and 4). Masson’s trichrome considered special indicator stain for collagen fibers and it is intensity depended on the ferocity and prolong infections (Table 5), so the fibrotic lesions rich with collagen fibers take blue in color (Figure 5-7).

![Figure 2](image2.png)

Figure 2: Presence of ruptured sarcosystis (A), distribution of bradyzoites in interstitial tissue (B), infiltration of mononuclear inflammatory cells (C), Zenker’s necrosis of myocyte (D) and edema between muscle fibers (E). H&E (240X).
Table 5: Lesions and grade of collagen intensity in the affected muscles

<table>
<thead>
<tr>
<th>Intensity</th>
<th>Grades</th>
<th>Lesions</th>
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</thead>
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<tr>
<td>Mild</td>
<td>+ with present sarcocysts</td>
<td>Deposition of collagen fibers, edema, infiltration of inflammatory cells, Zenker’s necrosis and destruction of muscle fiber.</td>
</tr>
<tr>
<td>Moderate</td>
<td>++ with present sarcocysts and bradyzoites</td>
<td></td>
</tr>
<tr>
<td>Severe</td>
<td>+++ with present sarcocysts and bradyzoites</td>
<td></td>
</tr>
</tbody>
</table>

Figure 3: Proliferation of fibroblast (A) with presence of fibrous tissues (B). H&E (240X).

Figure 4: Presence of Zenker’s necrosis in myocytes (A), atrophy (B), edema (C) and fibrosis (D). H&E (220X).

The results of the molecular analysis using the nPCR technique showed the diagnosis of Sarcosystis tenella in 11 samples of the diaphragm muscles of sheep, which were positive by microscopy, using the external and internal primers and the multiplication bands appeared on the agarose gel of 1.5% concentration, the reaction product was 800 bp and 500 bp (Figure 8 and 9) Whereas, the results of the molecular analysis using the external and internal primer of Sarcosystis arietianis were negative in the affected diaphragm muscles, as no bands were observed on the 1.5% agarose gel.

Figure 5: Mild deposition of collagen fiber take blue in color (A), presence of sarcosystis (B), edema between muscle fibers (C) and Zenker’s necrosis of myocyte (D) and + Masson’s trichrome. H&E (140X).

Figure 6: Moderate proliferation of collagen fibers take blue in color (A), presence of sarcostis between muscle fibers (B), Zenker’s necrosis of myocyte (C), infiltration of mononuclear inflammatory cells (D) presence of bradyzoites between muscle fibers (E) and ++ Masson’s trichrome (165X).
affected muscle, eosinophilic myositis, become is associated with inflammatory cellular debris, a granulomatous reaction or by histiocytic infiltration of inflammatory cells specially eosinophils, macrophages, lymphocytes, monocytes and occasionally giant cells at the site of lesion (16,17). These result supported the theory involved the antigenic effects of Sarcosystis have a significance role in activation of immune response then development of eosinophilic myositis (5,18). The muscle necrosis and atrophy observed in this study, so the toxic effects of toxins cause loss of muscle integrity, rapid utilization of glycogen and accumulation of lactic acid that lead to change in protein structure and subsequent unregulation of calcium flow, followed by increase of cytoplasmic calcium which cause activation of protease and hydrolytic enzymes that induce further muscle damage (19). On the other hand, sarcocyst press on muscle for a long time lead to loss of its mass and then atrophied (18).

Trichrome showed proliferation of collagen fibers stained with blue in color. Connective tissue redesigning is an important and significant stage for muscle regeneration. After the muscle injuries a gauze is formed between the muscle fibers and filled with inflammatory cellular debris, so the late in elimination of these remnants and the continuous irritation by sarcocystis lead to activation of fibroblasts and to secrete chemical agents as transforming growth factor-beta1 (TGF-B1) Which have the ability to stimulate the proliferation of fibroblasts subsequently production of collagen fiber (20-22).The intensity of positive reaction of masson’s trichrome maybe depend on the ferocity and prolong of infection (23).

The nPCR analysis showed 11 samples of diaphragm muscles appeared positive for microscopic sarcocystis cysts and the microscopic cysts related to species of Sarcosystis tenella only and this species responsible of grossly and histopathological changes. This result may be related to the type and size of the sample used in this study, and this is consistent with Ali et al. (24) in Baghdad province/ Iraq, and da Silva et al. (25) in Brazil, while Pipia et al. (13) referred to diagnosis two microscopic species of Sarcosystis tenella and S. arieticanis in heart samples of sheep in Italy by using nested PCR with prevalence 95.5%, 17.8% respectively. Oryan et al. (2) showed that the high prevalence of microscopic species of Sarcosystis can have a significant impact on sheep productivity through its negative effect on the weight and growth of sheep as well.

Discussion

Sarcosystis was considered as the most important disease that causes economic losses for meat producing and packing industries, through its effects on morphological and quality of meat which in turn has a negative impact on public health. In our study the affected muscles appeared white and pale in color, this result was agreement with the result of Sun et al. (14).

The histopathological changes of diaphragm muscles represented by the presence of sarcocysts were ruptured and degenerated. The rupture of cyst may be happened due to a host immune reaction or by over distention that occur during the development or as a result of accumulation of metabolic product and cytotoxin that enhanced osmotic pressure and crossing of fluid in to the cyst which become weakened and then ruptured (15), so that the contents of the rupture cyst (bradyzoites and toxins) into the surrounding tissue cause damage and massively infiltration of inflammatory cells specially eosinophils, macrophages, lymphocytes, monocytes and occasionally giant cells at the site of lesion (16,17). These result supported the theory involved the antigenic effects of Sarcosystis have a significance role in activation of immune response then development of eosinophilic myositis (5,18). The muscle necrosis and atrophy observed in this study, so the toxic effects of toxins cause loss of muscle integrity, rapid utilization of glycogen and accumulation of lactic acid that lead to change in protein structure and subsequent unregulation of calcium flow, followed by increase of cytoplasmic calcium which cause activation of protease and hydrolytic enzymes that induce further muscle damage (19). On the other hand, sarcocyst press on muscle for a long time lead to loss of its mass and then atrophied (18).

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as the high economic losses resulting from abortion. Anja et al. (26) indicated that the diagnosis of Sarcosystis in animals is limited, given that the traditional diagnostic methods based on the detection of specific Sarcosystis antibodies are only genus specific and cannot distinguish between sarcosystis species, in addition to which most of these methods appear good sensitivity to the later stages of the disease, therefore, the diagnosis for acute sarcocystosis was mainly based on postmortem examination. Here, molecular tools are the best diagnostic methods used for the detection of the species of Sarcosystis and in nested PCR assays, the unique small ribosomal RNA gene sequences of the S. tenella and S. arrieticanis subunit are used.

Conclusion

This study concluded demonstration the microsporidian species Sarcosystis tenella parasite in diaphragm muscle of sheep by using microscopy and nested PCR associated with the characteristic histopathological changes in diaphragm muscle included, Zenker’s necrosis and intense infiltration of inflammatory cells. With the formation of fibrous tissue whose intensity was measured (mild, medium and intense) by using the Masson’s trichrome stain.

Acknowledgments

The authors thanks College of Veterinary Medicine, University of Mosul, to supports this research.

Conflict of interest

The authors declare that there are no conflicts of interest regarding the publication of this manuscript.

References

دراسة مرضية وجزيئية للحجاب الحاجز المصاب طبيعيا بطفيلي المكيسات العضلية في الأغنام

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فرع الأمراض وأمراض الدواجن، فرع الأحياء المجهرية، كلية الطب البيطري، جامعة الموصل، الموصل، العراق

الخلاصة

لطفيلي المكيسات العضلية علاقة وثيقة مع العضلات وذلك بسبب إصابتها للعضلات الهيكلية في الإنسان والحيوان، إذ تسبب الحالة المزمنة للمرض خسائر اقتصادية مهمة وخاصة من ناحية إنتاج اللحوم من جراء تكوين الأكياس سواء كانت عيانية أم مجهرية في الألياف العضلية. ويعتبر النوعين Sarcosystis tenella و Sarcosystis arieticanis من أهم الأنواع الممرضة المكونة للأكياس العضلية في Sarcosystis arieticanis الأغنام. إذ تم في هذه الدراسة فحص 50 عينة من عضلات الحجاب الحاجز للأخناء المذبوحة في محلات القصابين ومجزرة الموصل عيانيا ونسجيا وتاقيات البلمرة المتسلا في كلاء تشخيصية لتحديد أو تشخيص النوع السبب والمسؤول عن هذه التغيرات. أظهرت 11 عينة من الحجاب الحاجز بالإضايط وشاحب عند الفحص المجهرية في حين تميزت الأعاف النسجية بوجود آفات طفيلي بأعداد واحجام مختلفة بين الألياف العضلية مما أدى إلى حدوث النخر الزجاجي وارتفاع كثيف للخلايا الالتهابية وخاصة خلايا الحمضة وحيدة النواة والخلايا البلعمية والمغلفة فضلا عن الورم وتكاثر الخلايا الليفية مع تكوين النسيج الليفي الذي استمر على شدة (خفيف ومتوسط وشديد) باستخدام صبغة الماسون ثلاثية اللون. كما أشارت نتائج التحليل المجهرية باستخدام تقنية تفاعل البلمرة المتسلسل المتداخل أن هذه الأكياس المجهرية المشخصة تعود للنوع Sarcosystis tenella ووافق نتائج تفاعل 800 و 500 زوجا قاعيا.