First Documented Study of Mycoplasma Wenyonii of Cattle in Iraq

Basima. A¹, Baraa. A²

¹ Department of Internal and Preventive Medicine, College of Veterinary Medicine, University of Mosul, Mosul-Iraq
² Nineveh Agricultural Office, Application Studies Laboratory. Mosul -Iraq

Abstract: Mycoplasma wenyonii was studied by PCR and staining blood smears in twelve local breed cattle brought from different farms around Mosul city for examination in teaching hospital of Mosul University. Clinical signs and hematological parameters of all tested animals were compromised with data of another twelve local breed healthy cattle. Clinical signs of hind limb edema, pyrexia, painful swollen udders, pre-femoral and mammary lymph nodes enlargement with milk drop were noticed in dairy cows. Scrotal edema was found to be significantly associated with the infection in calves. Eperythrozoon organisms were seen on traditional microscopic examination of blood smears of anemic cattle by blood smear examination is M. wenyonii by molecular analysis, and secondly to determine if it is a newly observed organism which may be the cause of outbreaks of recumbence, hind limb edema, udder swelling and scrotal edema in cattle herds at various locations across Mosul-Iraq in recent years.

Keywords: Mycoplasma wenyonii, PCR, DGGE, Iraq.

1. Introduction

Mycoplasma wenyonii is a member of the haemomycoplasmas, a group of haemotropic bacteria, which are found in close association with mammalian erythrocyte membranes. Mycoplasma wenyonii is common in cattle worldwide. Formerly known as Eperythrozoon wenyonii, it has been recently reclassified to the Mycoplasma genus based on analysis of the 16S rRNA [1]. Mycoplasma wenyonii is an epicellular prokaryote found in cocci, ring, and rod shapes that parasitizes red blood cells of cattle. The organisms have a single-cell membrane and adhere to the red blood cell membrane, but are not intraerythrocytic [2, 3]. The mode of transmission of M. wenyonii is currently unknown, but most likely results from mechanical transmission by blood-sucking arthropods, direct contact, and iatrogenic transmission. The majority of cattle infected with M. wenyonii do not develop clinical signs. A clinical illness is rare and seen only when concurrent illness results in immunosuppression. Clinical signs associated with infection include anemia, ill-thrift, fever, lymphadenopathy, depression, diarrhea, and decreased milk production [4].

Haemoplasmas have never been cultured in vitro. So, until recently, their detection was only possible with a subjective and insensitive microscopy test for blood smears but is not specific for M. wenyonii and can be difficult to interpret. McAuliffe et al., have described the use of PCR and denaturing gradient gel electrophoresis (DGGE) for the detection and differentiation of Mycoplasma species [5, 6]. DGGE can be used on blood samples as a rapid and specific test for M. wenyonii and can also be used as a screening test for other blood borne pathogens [7]. However M. wenyonii has been recently reported in cattle of Iraq according to the microscopic examination of blood smears of anemic cattle during the first outbreak in 2011[8].

The objectives of this study were, firstly to confirm that structures observed on erythrocytes of anemic cattle by blood smear examination is M. wenyonii by molecular analysis, and secondly to determine if it is a newly observed organism which may be the cause of outbreaks of recumbence, hind limb edema, udder swelling and scrotal edema in cattle herds at various locations across Mosul-Iraq in recent years.

2. Materials and Methods

Twelve local breed cattle (six calves of 11 to 15 months of age and six dairy cows 2-7 years old) were brought from different farms around Mosul city to the Veterinary Teaching Hospital at Veterinary Medicine College of Mosul University, Mosul Iraq, and the calves were suffering of poor appetite, fever, tachycardia, tachypnea, emaciation, exhaustion, anemia, hind limbs and scrotal edema. Where dairy cows showed clinical signs of hind limbs and udder edema, pyrexia, pre-femoral and mammary lymph nodes enlargement with milk drop. Some of the cattle were treated at local veterinary clinics with Imidocarb dipropionate (3mg/kg/day for 2 days) or Oxytetracyclin (20 mg/ kg for 3 days) by intramuscular injections. The clinical signs and hematological parameters of all tested cattle were recorded. Another twelve cattle (Six calves and six dairy cows) were clinically normal and negative to hemomycoplasma in blood smears and then by PCR/ (DGGE) which served as uninfected controls.

As a material for this research, whole blood with EDTA and peripheral blood smears have been used. Two blood smears were prepared immediately after each blood collection. Thick blood smears were made and stained in Wright’s -staining solution. Light microscopy was used to look for the presence of M. wenyonii on the erythrocyte surface. Also the fluorescence microscopy (BX51 Olympus U-RFL-T-Japan) was used to look for the microbes by mixing a drop of infected blood with acridine orange solution (0.1 mg/mL).
Then, thin smears were made and examined under 100X [9]. Parasitaemia was confirmed on blood smears of all cattle showed clinical signs; Positive results with *Mycoplasma wenyonii* infection were considered if one infected erythrocyte was found in 200 observed RBC [10]. We determined the percentage of reticulocytes by mixing equal parts of the anticogulated blood with New methylene blue, allowing the mixture to incubate at room temperature for 10 minutes, and preparing air-dried blood films from the mixture; the reticulocytes per 1000 erythrocytes were counted [11].

PCR, Denaturing Gel Electrophoresis (DGGE) and sequencing were performed at the Animal Health and Veterinary Laboratory Agency (AHVLA), (Weybridge), Woodham Lane, Addlestone, UK, from twelve anticogulated EDTA blood samples of animals showing clinical signs to confirm diagnosis. Total Genomic DNA was extracted from 100 μL of each blood sample using a tissue DNA extraction kit (Sigma). Amplification of the V3 region of the 16S RNA gene was performed according to the method of as described previously [5], using universal bacterial primers GC-341F 50-CGC CCG CCG CGC GGC GCG GGC GGG GCA CGG GGG GCC TAC GGG AGG CAG CAG and 534R 50-ATTACC GCG GCT GGG GGC GGG GCA CGG GGG GCC TAC GGG AGG CAG CAG and 534R 50-ATTACC GCG GCT GCT GG. DGGE was performed using the Ingeny PhorU 2 _ 2 apparatus (GRI Molecular Biology, Essex, UK). Samples were loaded onto 10% polyacrylamide/bis (37.5:1) gels with linear denaturing gradients from 30–60% (where 100% is 7 M urea and 40% (v/v) deionized form amide) in 1 _ TAE electrophoresis buffer (Severn Biotech Ltd., Worcestershire, UK). Electrophoresis was performed at 100 V at a temperature of 60 8C for 18 h. Gels were then stained with SYBR Gold (Cambridge ioScience, Cambridge shire, UK) in 1 _ TAE for 30 min at room temperature and visualized under UV illumination. Almost full-length 16S rRNA genes were amplified from isolates using the 63F and 1387R [12] primers under the conditions described above. PCR products were cleaned up using a Qiagen QiAquick PCR purification kit and products were quantified by running on 2% agarose E-gels with a quantitative E-gel ladder (Invitrogen). 16S rDNA was sequenced in both the forward and reverse direction and sequencing reactions were performed using a DTCS Quick Start sequencing kit (Beckman Coulter) according to the manufacturer’s instructions. Samples were ethanol precipitated and run on a Beckman Coulter CEQ8800 Sequencer. Contigs were assembled using the CAP Contig program in Bioedit. These sequences were aligned with the closest matches found in the Ribosomal Database Project with the ClustalW function of the BioEdit package. Neighbor joining phylogenetic trees were constructed with the Molecular Evolutionary Genetics Analysis package (MEGA Version 2.1) using unweighted pair group mean analysis (UPGMA), and the robustness of the phylogeny was tested by bootstrap analysis with 500 iterations.

The arithmetic means (±SD) of hematological parameters in different groups were calculated. Further analysis was done using analyses of variance (ANOVA) technique by SPSS statistical program and means were separated by Duncan's multiple range test. The level of significance was determined at P<0.05.

3. Results

The main clinical signs of *M. wenyonii* infection in dairy cows in the present study were udder and hind limbs edema (Figure 1: A). Scrotal and hind limb edema was associated with infection in bull calves (Figure 1: B).

![Figure 1: Udder and hind limb edema in cow (A), Scrotal and hind limb edema in a bull (B) testing positive for *Mycoplasma wenyonii* on denaturing gradient gel electrophoresis.](image)

Animals showed either acute illness characterized by recumbence, weakness, depression, pyrexia (39.9±0.3°C in dairy cows and 40.5±0.8°C in calves), tachycardia (96±8.3 beats/min in dairy cows and 132±11 beats/min in calves) and tachypnea (38±5.6 breaths/min in dairy cows and 42± 3.9 breaths/min), or may demonstrate chronic illness characterized by weight loss, anorexia, emaciation, enlargement of prefemoral and supra-mammary lymph nodes.

Examination of peripheral blood smears revealed numerous small basophilic staining coccids structures 0.5-1 μm in diameter with a number ranging from 1 to 90 organisms on the surface of the erythrocytes attached to cell membranes singularly or in clusters centered on a single membrane focus which caused different levels of RBC deformation, some erythrocytes were not infected (Figure 2).

![Figure 2: Light micrograph of *Mycoplasma wenyonii*. Note presences of numerous small basophilic structures under oil immersion 160X, Wright’s stain.](image)

Sometimes the organisms were observed free in the background; they were bigger and stained darker. The delicate fibrils of the organism could be seen even under a light microscope (Figure 3).

![Figure 3: Light micrograph of *Mycoplasma wenyonii*. Note presences of numerous small basophilic structures under oil immersion 160X, Wright’s stain.](image)
On hematology, anemia had indicated by values of hemoglobin concentrations, erythrocytes counts and hematoctrit in the clinically diseased cattle. The dairy cows had significantly lower hemoglobin (58.6 g/l ± 6.1), erythrocytes (3.51 x10^{12}/L ± 0.63) and hematoctrit (0.20 L/L ± 0.07) levels than those in healthy control dairy cows, which exhibited values of (105.6 g/l ±12.8 g/l), (8.6 x10^{12}/L ± 0.03) and (0.34 L/L ± 0.04) respectively. In calves the hemoglobin concentration was 78.0 g/l ±13.1, erythrocytes counts was 2.8 x10^{12}/L ± 0.25 and hematoctrit was 0.22 L/L ± 0.02 in comparison with normal values in healthy calves. Macrocytic hypochromic and macrocytic normochromic anemia was revealed in infected dairy cows and calves respectively (Table 1). There was also a leukocytosis with mild neutrophilia (Table 2).

The statistics of each hemomycoplasma-positive group were compared with those of the negative group. * P<0.05; (vs. the negative group).

The statistics of each hemomycoplasma-positive group were compared with those of the negative group. * P<0.05; (vs. the negative group).

Erythrocyte features included moderate anisocytosis and poikilocytosis, polychromasia, reticulocytosis, basophilic stippling, nucleated erythrocytes (NRBC) with presence of Howell-Jolly bodies and Heinz bodies (Figures 6-9). The reticulocyte percentage was about 2-3% and the absolute reticulocyte concentration was about 27800/L in infected dairy cows and 40,200/L in calves on a new methylene blue stained slides in comparison with normal values 10,000/L in healthy cattle. About 5-10 NRBC/100WBC were counted compared with mostly metarubricytes and fewer polychromatophilic rubricytes. Low numbers of neutrophils exhibited mild toxic changes with cytoplasmic basophilia and Döhle bodies (Figures 6-9).

In acridine orange-stained blood smears of infected cattle, numerous M. wenyonii as yellow to orange particles were seen on and between the erythrocytes (Figure 5). A cridine orange-staind blood smears of infected cattle, numerous M. wenyonii adhered on the surface of erythrocytes (MW) and on platelets (PL) with erythrophagocytosis of the infected RBC (E). Note the delicate fibrils of free M. wenyonii in the background (F) under oil immersion 100X.

Hemomycoplasmas attached to the RBCs and platelets membrane surfaces in cases of high parasitemia.

Erythrophagocytosis of the infected red blood cell were also seen in the blood smears of severely affected animals (Figure 4).

In acidine orange-stained blood smears of infected cattle, the hemomycoplasmas attached to the RBCs and platelets membrane surfaces in cases of high parasitemia. Erythrophagocytosis of the infected red blood cell were also seen in the blood smears of severely affected animals (Figure 4).

The hemomycoplasmas attached to the RBCs and platelets membrane surfaces in cases of high parasitemia. Erythrophagocytosis of the infected red blood cell were also seen in the blood smears of severely affected animals (Figure 4).

The hemomycoplasmas attached to the RBCs and platelets membrane surfaces in cases of high parasitemia. Erythrophagocytosis of the infected red blood cell were also seen in the blood smears of severely affected animals (Figure 4).
Figure 6(A): Wright’s stain blood smear of anemic cow infected with *Mycoplasma wenyonii*. Note anisocytosis with Howell-Jolly bodies (HJ) under 160X.

Figure 6(B): The infected erythrocytes have multiple punctate; bacillary, or ring forms and poikelocytosis under 100X.

Figure 7(A): Blood smears from anemic calf with eperythrocytic haemomycoplasma organisms, Note the reticulocytosis (R) in New methylene blue stain.

Figure 7(B): A nucleated erythrocyte (metarubricyte) is present in the upper right (NRBC) of blood smear. Wright-stain, under oil immersion 160X.

Figure 8(A, B): Blood smears from anemic calf with eperythrocytic haemomycoplasma organisms (M). Note that the Howell-Jolly bodies (HJ) in A and Heinz bodies (H) in B, stain with acridine orange, under oil immersion 100X.

Figure 9(A): Numerous intensely stained *Mycoplasma wenyonii* (MW) can be observed on the surface of erythrocytes. Note the macrocytic cells with polychromasia (MP) and basophilic stippling (BS).

Figure 9(B): Mild toxic neutrophilia with cytoplasmic basophilia and Döhle bodies (DB). 160X, Wright’s stain.

Table 3: Erythrocyte inclusions in blood smear of cattle which infected with hemomycoplasma positive and negative groups

<table>
<thead>
<tr>
<th>Erythrocyte inclusions (%)</th>
<th>Positive cows (n=6)</th>
<th>Negative cows (n=6)</th>
<th>Positive calves (n=6)</th>
<th>Negative calves (n=6)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Reticulocyte</td>
<td>2%</td>
<td>0</td>
<td>3%</td>
<td>0</td>
</tr>
<tr>
<td>Nucleated RBC / 100WBC</td>
<td>5</td>
<td>0</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>Basophilic stippling/100</td>
<td>1-5 (1+)</td>
<td>0</td>
<td>1-5 (1+)</td>
<td>0</td>
</tr>
<tr>
<td>Microscopic field</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Heinz bodies / 100</td>
<td>3-5 (1+)</td>
<td>0</td>
<td>5-10 (2+)</td>
<td>0</td>
</tr>
<tr>
<td>Microscopic field</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Howell-Jolly bodies</td>
<td>1-2 (1+)</td>
<td>0</td>
<td>1-2 (1+)</td>
<td>0</td>
</tr>
<tr>
<td>/ 100 Microscopic field</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
**Mycoplasma wenyonii (MW)**-specific DNA was detected by PCR and Denaturing Gradient Gel Electrophoresis (DGGE) in blood samples of calves and dairy cows showing clinical signs. Although epi-erythrocytic organisms were seen on blood smears from all sampled cattle, only five samples (two cows and three calves) were PCR positive. **Mycoplasma wenyonii** was identified by PCR/DGGE in samples 1 (Lane 6), 4 (Lane 9), 5 (Lane 10), 9 (Lane 14) & 10 (Lane 15). According to 16S rDNA PCR and DGGE an identical band was seen in all five samples which did match with bands of positive control samples (Lanes 3 and 19) of the known bovine mycoplasma controls; the control isolate of Mycoplasma wenyonii 599. The samples 5 (Lane 10) and 10 (Lane 15) gave a strong high band on the DGGE whereas a weaker high band were seen in the samples 1 (Lane 6), 4 (Lane 9) and 9 (Lane 14). The samples 2, 3, 6, 7, 8, 11, 12 had negative results. Unidentified multiple bands were also detected in samples 2, 3, 4, 7, 8, 9 & 11. For those samples with unidentified bands we will need to see if sequencing gives any more information. Usually multiple bands do not give a good sequence, but we will have to see what we get. These bands were not consistent with **M. wenyonii** and had a low identity to a variety of Pseudomonas and Klebsiella species on sequencing most likely due to contamination (Figure 10).

Figure 10: DGGE of cattle Hemotrophic Mycoplasmas. Lanes: 1 and 22 Negative control; Positive control of **Mycoplasma wenyonii** 2 and 18 **M. wenyonii** 620, 3 and 19 **M. wenyonii** 599, 4 and 20 **M. wenyonii** 993, 5 and 21 **M. haemobos** 231B11; Samples: Lane 6, 9, 10, 14, 15, **M. wenyonii** 599. Samples (1-6) are of calves and samples (7-12) from dairy cows.

4. Discussion

This study represents the first documented report of **Mycoplasma wenyonii** infection in cattle of Iraq. A combination of hind limb, udder or scrotal edema, pyrexia, in appetite and prefemoral lymphadenopathy is considered highly suggestive of **M. wenyonii** infection in cattle [13, 14]. How a hemomycoplasma with a tropism for the erythrocyte membrane causes or contributes to clinical signs of hind limb and udder edema is not exactly clear at present. Based on biopsies of edematous skin, suggested that the edema might be caused by vasculitis, possibly an Arthus-type reaction with deposition of immune complexes in the vascular endothelium. Immune-mediated disease is a feature of Mycoplasma infections in other species and humans. This theory sounds plausible, although further work is required [15].

Mycoplasma wenyonii has been previously reported to cause clinical anemia in young or splenectomized cattle [16, 17]. Complete blood counts performed on hemomycoplasma infected cattle revealed macrocytic anemia and marked alterations in erythrocyte morphology, however, there was evidence on blood smears (such as reticulocytosis, polychromatic macrocytes, Howell-Jolly bodies, Heinz bodies and basophilic stippling) that anemia in most cattle with the haemomycoplasma organisms was strongly regenerative. The bacteria attached to the RBC membrane surface, which caused different levels of RBC deformation, increased membrane injury associated with various pathologic disorders which can result in increased phagocytosis of erythrocytes by macrophages. Lysis of erythrocytes within macrophages after phagocytosis is sometimes referred to as extravascular hemolysis. We suggest that intimate contact of hemmycoplasma organisms with RBCs leads to cell injury through immune-mediated, erythrophagocytosis and non-specific factors, which increase red cell fragility and may be the cause of anemia. Further work is needed to characterize the mechanism of anemia in cattle with **M. wenyonii**. A mild toxic change of some neutrophils is likely due to acute inflammation [18]. **Mycoplasma wenyonii** was first described in 2011 in Iraq using blood smear examination to the organism formerly known as *Eperythrozoon wenyonii* (**Mycoplasma wenyonii**) by [8]. The PCR/DGGE were able to identify the presence of the bacteria, even in very small amounts. Only five samples were positive in PCR from total transmitted samples, however, most of the ill cows and calves had received extensive treatment with Oxy-tetracycline or Imidocarb in local clinics before the samples collection, which might have led to the underestimation of **M. wenyonii** infection in ill cattle. Samples 5 and 10 from cattle not treated gave a strong high band on the DGGE whereas a weaker high band and a much lower band were seen in samples 1, 4 and 9 from cattle treated previously with Oxytetracycin and negative results with samples treated with imidocarb because hemomycoplasma blood loads can be reduced by application of these therapies [19, 20].

We also suggest that long storage of blood samples or DNA may have led to DNA degradation, and therefore, samples with a long history of storage may have appeared PCR negative by mistake. This may explain why several samples (2, 3, 6, 7, 8, 11 and 12) from 2011 reacted negative in the PCR (performed in 2012/2013) even though, the samples were positive in 2011, as determined by microscopy. Blood had been stored at -20°C, but DNA can degrade due to repeated freezing and thawing. Blood samples, which were posted by DHL, had a long “travel history” to the Veterinary Laboratory.
Laboratory Agency (AHVLA), UK where they were processed. Taking this into account, the calculated blood loads in the samples with a longer storage or shipment history may have been too low or possibly PCR may have reacted negative in error. *M. wenyonii* 599 was thought to be a very serious infection of cattle. We need further studies about the epidemiology of disease in Iraq. We need further studies about the epidemiology of disease in cattle of Iraq to determine whether *M. wenyonii* is the only hemomycoplasma present in Iraqi cattle or another species is the causative agent of the disease, if cattle is the only animal species that is affected with hemomycoplasma, and if *M. wenyonii* 599 is the only strain present or other strains play a role in the epidemiology of disease in cattle.

5. Conclusions

This is the first report of *Mycoplasma wenyonii* in calves, and dairy cows in Iraq based on molecular evidence. Further work is required in order to determine the true mechanism of anemia in *Mycoplasma wenyonii* infection of cattle as well as the epidemiology of disease in Iraq.

References


