

***Mycoplasma suis* in naturally infected pigs: an ultrastructural and morphometric study¹**

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ABSTRACT.- Portiansky E.L., Quiroga M.A., Machuca M.A. & Perfumo C.J. 2004. ***Mycoplasma suis* in naturally infected pigs: an ultrastructural and morphometric study.** *Pesquisa Veterinária Brasileira* 24(1):1-5. Instituto de Patología, Facultad de Ciencias Veterinarias, Universidad Nacional de La Plata, Calle 60 y 118, C.C. 296, (1900) La Plata, Buenos Aires, Argentina. E-mail: elporti@fcv.unlp.edu.ar

Swine eperythrozoonosis is a haemotrophic disease caused by *Eperythrozoon suis*, actually called *Mycoplasma suis*, an extracellular bacterial organism that apparently adheres to pig erythrocyte membrane, inducing its deformation and damage. Since little is known about the ultrastructural and morphometrical aspects of this microorganism, the present work aimed to deal with these issues. The ultrastructural study revealed the presence of structures corresponding to tubules disseminated throughout the soma of *M. suis*. A variable separation between the microorganism membrane and that of the erythrocyte was also observed. The structural and positional attitude of *M. suis* could allow speculation about its mechanism of action.

INDEX TERMS: Eperythrozoonosis, *Mycoplasma suis*, ultrastructure, morphometry, pathogenesis.

RESUMO.- [*Mycoplasma suis* em suínos naturalmente infectados: um estudo ultra-estrutural e morfométrico.]

A eperitrozoonose suína é uma doença hemotrópica causada por *Eperitrozoon suis*, atualmente denominado *Mycoplasma suis*, uma bactéria extracelular que, aparentemente, adere à membrana dos eritrócitos suínos, induzindo sua deformação e lesionando-os. O presente trabalho busca estabelecer os aspectos estruturais e ultra-estruturais, pouco conhecidos, deste microorganismo. O estudo ultra-estrutural revelou a presença de estruturas correspondentes a túbulos disseminados no soma bacteriano. Observou-se também uma separação variável entre a membrana do microorganismo e a parede do eritrócito. O estudo morfométrico e a localização de *M. suis* pode permitir especulação sobre seu mecanismo de ação.

TERMOS DE INDEXAÇÃO: Eperitrozoonose, *Mycoplasma suis*, ultra-estrutura, morfometria, patogenia.

INTRODUCTION

Eperythrozoonosis is a haemotrophic disease in swine caused by a bacterium formerly classified as the rickettsial agent *Eperythrozoon suis* (Hoelzle et al. 2003). The disease is clinically characterized by anemia, jaundice and a variety of associated syndromes (Hsu 1986, Henderson et al. 1997, Machuca et al. 1999).

Eperythrozoon suis, now allocated to the genus *Mycoplasma*, is an extracellular bacterial organism that attaches to and causes deformity and damage to porcine red blood cells (Hoelzle et al. 2003). The accepted actual name of the agent is *Mycoplasma suis* (Neimark et al. 2002). It has a restrictive cellular membrane, lacking organelles and nucleus. Its cytoplasm contains as much DNA as RNA (Gwaltney 1995, Gresham 1996). *Mycoplasma suis* can be identified by optic microscopy in Romanowsky stained blood smears (Gwaltney 1995). It may be round, oval or ring-shaped with diameters ranging from 0.8 µm to 2.5 µm or even larger (Kreier & Ristic 1984, Gwaltney 1995).

Although the pathogenesis of the disease is not completely known, it is considered that once *M.suis* comes into contact with the erythrocyte, it uses the plasmatic glucose for its own metabolism. On the other hand, upon damaging the cellular membrane of the red blood cell, it would induce the

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production of autoagglutinins, favoring agglutination of these cells and their removal by way of the phagocyte-mononuclear system (Gwaltney 1995, Gresham 1996).

Since little is known about the structural (morphometric) and ultrastructural aspects of this microorganism (Pospichil & Hoffmann 1982), the goal of the present work was to obtain more details regarding its morphology that could explain its pathogenesis.

MATERIALS AND METHODS

Blood samples were obtained from 27 naturally infected feeder male and female pigs, from a semi-intensive establishment located in the town of Junin, Buenos Aires Province, Argentina. The samples were obtained by venipuncture of the anterior cava vein, using ethylenediaminetetraacetic acid (EDTA) as anticoagulant, in a concentration of 0.1 mg/ml of blood. The samples thus obtained were centrifuged at 800 rpm, for 5 min. Later, the cells were washed 3 times with PBS, for 5 min each time. The washed samples were fixed in 2% glutaraldehyde solution. After 30 min, the pellet was broken into small pieces, which were left in contact with the fixer for another 30 min. The fragments were later washed 3 times with PBS and post-fixed with osmium tetroxide, for 30 min. After dehydration of the samples, they were infiltrated and embedded with epoxy resins (Quetol 812, Nisshin EM Co. Ltd, Tokyo, Japan). Blood smears were stained with May-Grünwald-Giemsa and Wright stains.

Semi-thin 1 mm thick cuts were obtained using an ultramicrotome. The samples were stained with methylene blue and a "Multiple Stain" solution (Polysciences Inc., Warrington, USA) and observed with an optic microscope. Ultra-thin cuts were stained with uranyl acetate/lead citrate and observed with a transmission electronic microscope JEM-1200 (JEOL Co. Ltd, Tokyo, Japan).

Images of blood smears and semi-thin cuts were captured from a microscope (Olympus BX50 system microscope, Tokyo, Japan) with a microscopic magnification of $\times 100$, through an analogical RGB video-camera (Sony DXC-151A CCD, Tokyo, Japan) and digitized with a frame grabber (Flashpoint 128, Integral Technologies Inc., Indianapolis, IN, USA) connected to a computer. Images were processed using the Image-Pro Plus v4.50 software (Media Cybernetics, Silver Spring, MA, USA), with a depth of pixel of 24 bits, RGB and TIFF format. The resolution of the microscopic images was of 640×480 pixels with a spatial calibration yield of $0.13 \mu\text{m}/\text{pixel}$. The photographs obtained with the electron microscope were digitized through a full-page scanner (Microtek Scanmaker E6, Rotterdam, Holland), with a resolution of 300 ppp. In order to separate the desired object from the rest of the tissue, color segmentation, based on the optical density of the object, was issued. Once separated from the rest of the image, the object was quantitatively characterized based on its area (in μm^2), major and minor axis (to obtain the aspect of the object), maximum and minimum diameter, perimeter and roundness, all of them measured in lineal μm . Table 1 describes the morphometric parameters evaluated.

RESULTS

The blood smears revealed the presence of *Mycoplasma suis* in close contact with the red blood cells (Fig. 1a). The stained semi-thin cuts showed a regular quantity of spherical elements appearing alone or in groups of 2 or 3, stuck to the membrane of the erythrocyte or free among the red blood cells (Fig. 1b).

Table 1. Morphometric parameters used to evaluate *Mycoplasma suis* characteristics

Parameter	Description
Area	Reports the area of each object
Aspect	Reports the ratio between the major axis and the minor axis of the ellipse equivalent to the object (i.e., an ellipse with the same area, first and second degree moments), as determined by Major Axis/Minor Axis
Major Axis	Reports the length of the main axis of the ellipse equivalent to the object (i.e., an ellipse with the same area, first and second degree moments)
Minor Axis	Reports the length of the minor axis of the ellipse equivalent to the object (i.e., an ellipse with the same area, first and second degree moments)
Max. Diameter	Reports the length of the longest line that can be drawn to pass through the centroid position and join two points on each object's perimeter
Min. Diameter	Reports the length of the shortest line that can be drawn to pass through the centroid position and join two points on each object's perimeter
Max. Distance	Reports the maximal separation between <i>Mycoplasma suis</i> and the red blood cell membrane
Min. Distance	Reports the minimal separation between <i>M. suis</i> and the red blood cell membrane
Perimeter	Reports the length of the outline of each object
Roundness	Reports the roundness of each object, as determined by the following formula: $(\text{perimeter}^2) / (4\pi * \text{area})$. Circular objects will have a roundness = 1; other shapes will have a roundness > 1

Fig.1c shows a magnification of *M.suis* in a red blood cell membrane. Fig.1d shows a 3D-like image obtained by a filtering process, in which a depression of the erythrocyte membrane surrounding *M. suis* can be observed. A higher magnification of the microorganism and the depression it produces on the red blood cell surface membrane can be observed in Fig.1e.

The ultra-structural studies revealed some morphological details of the *M. suis*. The organisms were observed as round or oval structures, limited by a membrane (Fig.2a). Neither cellular organelles nor nuclear elements were observed. In some cases, widespread electron-dense granules were identified in the cytoplasm. Some microorganisms evidenced structures similar to microtubules. Although *M. suis* was present on the plasma membrane of the erythrocyte, in most cases a separation between both structures was observed. No cellular membrane damage was detected in the red blood cells, although some irregularities in the shape of the infected erythrocyte were observed on many occasions (Fig.2b).

Table 2 shows the morphometrical values obtained from the light and electron microscopy studies. These values are the average of at least 30 images obtained from each analyzed sample. It was observed that there is coincidence between the major and minor axes and the maximum and minimum diameters, respectively. The electron microscopic data demonstrate that a distance of 37 to 64 nm separates those parasites that are not touching the erythrocytes from these cells. The real and calculated data for the inner tubule-like

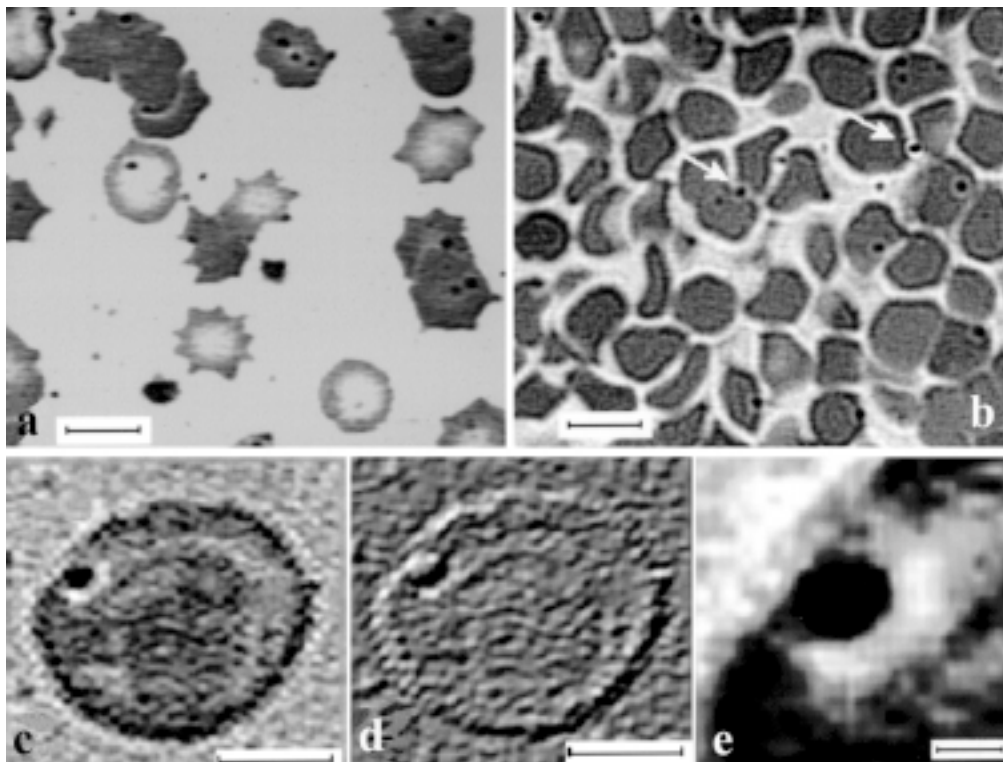


Fig. 1. Blood smears (a) and semi-thin cuts (b) showing spherical elements corresponding to *Mycoplasma suis* in close contact with the red blood cells (100x). Arrows point to the microorganism attached to two erythrocytes; Magnification (2x) of an erythrocyte appearing in (a) infected with *M. suis* (c); The same cell as in (c) with a 3D-like aspect obtained by filtering process; Magnification (4x) of *M. suis* (a) and the depression it produces on the erythrocyte surface membrane. Bars represent either 5 μm (a, b); 1 μm (c, d) or 0.25 μm (e).

structures of *M. suis* (Fig. 3a,b), maintain certain proportion. The calculated values were obtained from measurements

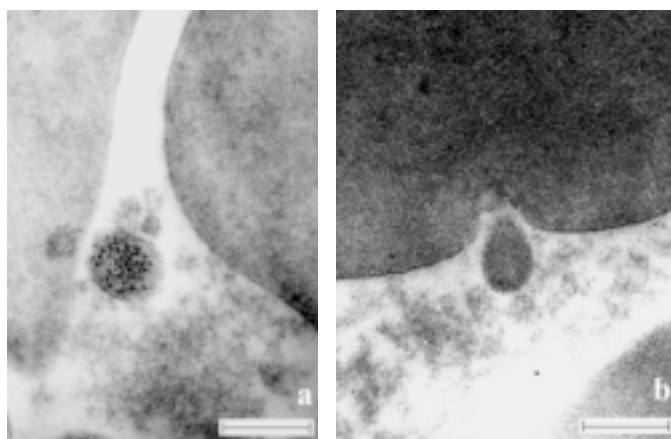


Fig. 2. Electron microscopic photographs showing a spherical structure corresponding to *Mycoplasma suis* (a). Widespread electron-dense granules were identified in the cytoplasm that resemble microtubules. *M. suis* could also be observed as an oval structure (b). Although no cellular membrane damage was detected some irregularities in the shape of the infected erythrocyte could be observed. In neither case is *M. suis* touching the red blood cell it infects. Bars represent 400 nm.

taken after a filtration process on the images from the electron microscope. The filtration process preserved just the desired objects for measuring at the expense of some image modifications. Since certain proportion exists, a correction index can be established for the filtered images. The area fraction of the real and calculated values of the tubules is shown in Table 3. In both cases the values are quite similar.

DISCUSSION

Swine eperythrozoonosis is a contagious disease characterized by a marked paleness of the mucous surfaces, presence of blood of watery aspect, delay of growth and increase of

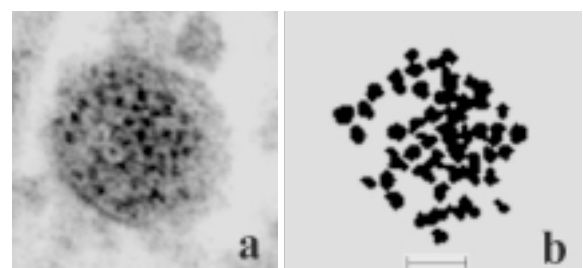


Fig. 3. *Mycoplasma suis* showing electron-dense structures resembling tubules (a). The same dense structures after thresholding the previous image (b). Bar represents 100 nm.

Table 2. Morphometrical values of *Mycoplasma suis* obtained from light and electron microscopy

Measurements ^a	<i>M. suis</i> soma (LM values)		Tubules-like structures (EM Values)		
	Blood smears	Semi-thin cuts	Grid	Real values	Calculated values
Area	0.33±0.07	0.18±0.06	0.17±0.09	221.47±126.55	304.08±169.20
Major axis	0.65±0.06	0.57±0.11	0.49±0.27	20.94±8.00	23.14±7.79
Minor axis	0.57±0.08	0.39±0.09	0.44±0.21	12.95±4.73	16.11±4.60
Aspect	1.14±0.31	1.49±0.38	1.11±0.24	1.79±0.68	1.46±0.30
Max. Diameter	0.66±0.04	0.57±0.11	0.49±0.29	20.31±7.04	22.78±7.32
Min. Diameter	0.57±0.08	0.39±0.09	0.45±0.26	12.76±4.63	15.51±3.66
Avrg. Diameter	0.61±0.06	0.48±0.09	0.47±0.28	16.55±5.37	19.17±4.82
Perimeter	1.23±0.19	1.07±0.35	1.47±0.86	42.58±21.61	51.71±24.23
Roundness	1.00±0.00	1.00±0.00	1.01±0.05	1.03±0.10	1.02±0.13
Max. Distance			64		
Min. Distance			37		

^aArea values are expressed either in μm^2 (*M. suis* soma) or nm^2 (tubules). All other values are expressed either in μm (*M. suis* soma) or nm (tubules). Tubules values represent each tubule characteristic. Maximal and minimal distance between *M. suis* and a red blood cell membrane are expressed in nm . Values represent the average of at least 30 images \pm SD. LM = Light microscopy. EM = electron microscopy.

Table 3. Area fraction of inner tubules-like structures of *Mycoplasma suis*

	Total number of tubules	Total tubule area ^a	<i>M. suis</i> area	Area fraction
Real values	44±11	25437.09±1392.05	117911.60±21758.02	21.57%
Calculated values	35±5	23406.58±846.00	117911.60±21758.02	19.85%

^aArea values are expressed in nm^2 .

mortality in feeder pigs, with a death toll that ranges between 3.1% and 18.7%, monthly. *Mycoplasma suis* can be detected by light microscopy in Romanowsky stained blood smears (Gresham 1996). In our work, round elements were observed ranging from 0.25 to 0.40 μm , stuck to the membrane of the erythrocyte in coincidence with those described by other authors (Hsu 1986, Solignac et al. 1996). In the 1 μm semi-thin cuts similar structures were observed but with smaller dimensions. This could be attributed to the stretching of the cells in the smears.

The ultrastructural study revealed some details of the morphology of *M. suis*. Although an apparent contact was observed between the microorganism and the erythrocyte in the semi-thin cuts, the ultrastructural studies revealed a separation ranging from 37 to 64 nm between the red blood cell membrane and *M. suis*. The closeness of the structures persisted even after the blood samples had been washed, lasting few free elements among the cells. Most of the infected erythrocytes showed variations in the contour of their membrane at the point of contact with the microorganism, but did not show any damage. These observations coincide with those described by Pospichil & Hoffmann (1982).

According to the morphometrical values obtained, a concordance between the major and minor axes and the maximum and minimum diameters, respectively, was observed.

This fact endorses the roundness equal to 1. This means that *M. suis* has a spherical form although the ultra-structural study revealed that *M. suis* presents an oval shape on certain occasions.

As described for other microorganisms, the microtubules observed in the cytoplasm of some *M. suis* could be a part of its cytoskeleton (Ghadially 1988). On the other hand, and keeping in mind that this microorganism uses the plasma glucose for its own metabolism (Gwaltney 1995), the microtubules could demarcate channels through which the transport of necessary products would take place. This allows us to speculate that *M. suis* does not need the physical atmosphere of the red blood cells to live and reproduce, but simply its nutrients, and that the destruction of the erythrocyte by exhaustion of its reserves would not necessarily signify the destruction of the agent, which would simply move in search of its next target cell.

The described morphometric details constitute an interesting fact for themselves, as they had never been depicted for *M. suis*. The use of an automated image analyzer allows objective appreciation, repetition of the measurements and simple operation. Data related to *M. suis* can thus be analyzed starting from blood smears as well as from semi-thin cuts. This allows a good diagnosis at a relative speed. The fine detail, represented by the presence of microtubules and the separation of *M. suis* from the erythrocyte membrane, adds to the precision of the diagnosis and leaves an open door to the analysis of a possible pathogenesis of the infection.

There are bacteria that interact with their target cells through a heterotypic union of surface glycoproteins to further internalize them by attaching to their cytoskeleton (Cossart & Bierne 2001). Although no surface proteins have been identified in the bacteria yet, it could be speculated that *M. suis* might interact with the erythrocyte in a similar fashion but without entering into the cell.

In conclusion, the use of the optic and electron microscope

in combination with an image analyzer would allow not only the diagnosis of an infection, as in eperythrozoonosis, but also the analysis of possible mechanisms of action.

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