Application of different techniques to detect *Toxoplasma gondii* in slaughtered sheep for human consumption

Utilização de diferentes técnicas para detecção de *Toxoplasma gondii* em ovinos abatidos para consumo humano

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Abstract

The aim of this study was to investigate occurrence of *Toxoplasma gondii* in sheep slaughtered in the state of Alagoas, Brazil, by means of different diagnosis techniques. Serum samples and tissues from 100 slaughtered sheep were used. To detect antibodies, the indirect immunofluorescence antibody test (IFAT) was used, and tissues from seropositive animals (cut-off $\geq 1:64$) were submitted to Polymerase Chain Reaction (PCR) and immunohistochemistry (IHC). To assess the concordance between the direct techniques, the kappa test was used. In the IFAT, it was observed that $14\% (14/100)$ of the ovine samples were serum-positive. In the PCR, $21.43\% (3/14)$ of the animals were positive and in IHC, it was observed that $7.14\% (1/14)$ were positively stained for *T. gondii* in cerebral tissue. Histopathologically, the predominant finding was the presence of mononuclear cell infiltrate in the heart and a perivascular cuff in the cerebrum and cerebellum. The concordance between the direct diagnosis techniques was moderate ($k=0.44$). Thus, it is important to use different direct techniques in diagnosing toxoplasmosis in naturally infected sheep.

Keywords: Sheep, *Toxoplasma gondii*, Polymerase Chain Reaction, histopathology, immunohistochemistry.

Resumo

O objetivo deste estudo foi pesquisar a ocorrência de *Toxoplasma gondii* em ovinos abatidos no Estado de Alagoas, Brasil por meio de diferentes técnicas de diagnóstico. Foram utilizadas amostras de soros e tecidos de 100 ovinos abatidos. Para a pesquisa de anticorpos foi utilizada a Reação de Imunofluorescência Indireta (RIFI), e os tecidos dos animais soropositivos (ponto de corte $\geq 1:64$) foram submetidos às técnicas de Reação de Cadeia da Polimerase (PCR) e Imunohistoquímica (IHQ). Para o estudo da concordância entre as técnicas diretas foi empregado o teste *Kappa*. Na RIFI, $14\% (14/100)$ das amostras foram soro-positivas. Na PCR, $21.43\% (3/14)$ dos animais foram positivos e, na IHC, $7.14\% (1/14)$ apresentaram marcação positiva para *T. gondii* no tecido cerebral. Na histopatologia, o achado predominante foi o infiltrado celular mononuclear no coração e manguito perivascular no cérebro e cerebelo. A concordância entre as técnicas diretas de diagnóstico foi moderada ($K=0.44$). Desse modo, é importante utilizar diferentes técnicas diretas no diagnóstico da toxoplasmose em ovinos naturalmente infectados.


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Introduction

Economic losses caused by *Toxoplasma gondii* infection in sheep are difficult to evaluate because this disease occurs sporadically (DUBEIY, 2009). Animals such as sheep and goats kept on pastures have an increased risk of *T. gondii* infection due to the environmental contamination with oocysts this parasite (TENTER et al., 2000).

Toxoplasmosis in sheep can cause abortions, neonatal mortality, mummified fetuses or weak lambs at birth (DUBEY & JONES, 2008; OIE, 2008). In the chronic form of the disease, most cases are not diagnosed or are confused with other similar diseases (BRAGA FILHO et al., 2010).

The diagnosis of toxoplasmosis can be made by means of indirect methods such as serological evaluation to detect specific antibodies. There are direct techniques to identify tachyzoites and cysts in animal tissues by means of: histological sections stained with hematoxylin and eosin (HE), immunohistochemistry (IHC) techniques and bioassays in mice (ROSA et al., 2001). In histological sections stained with HE, it is difficult to identify cysts of *T. gondii* because the parasite can be confused with nuclei or nuclear fragments that stain similarly (TSUNEMATSU et al., 1964; BARBOSA, 1988). This technique is generally used to look for histological lesions consistent with infection by this parasite, with or without associated use of immunohistochemistry (UGGLA et al., 1987).

Different Polymerase Chain Reaction (PCR) techniques are also used to detect the fetal DNA of the parasite in tissues and fluids from aborted fetuses (OWEN et al., 1998; HURTADO et al., 2001).

There is no much information about the most appropriate direct technique to evaluate the presence of *T. gondii* in tissues from sheep naturally infected after a prior serological screening. The aim of this study was to investigate the occurrence of *Toxoplasma gondii* in sheep slaughtered, by means of different diagnosis techniques.

Materials and Methods

Samples

The present study was developed according to Brazilian laws and was approved by the Animal Use Ethics Committee (protocol number 59/2013).

One hundred blood samples (50 males and 50 females) were collected during bleeding of the sheep and tissues samples (cerebrum, cerebellum, pons, spinal cord and heart) were collected along of the slaughter line of the abattoir located in state of Alagoas, northeastern Brazil.

The sera samples were collected in microtubes and stored at –20 °C until used. The tissue samples (20g) were stored at –80 °C and tissue fragments (0.5 × 2 × 2 cm) in 10% buffered formalin.

Experimental layout

The steps of the present study are summarized below (Figure 1).

Detection of antibodies

The indirect immunofluorescence antibody test (IFAT) was performed to detect IgG antibodies against tachyzoites of *T. gondii*, in accordance with the method proposed by Camargo (1974), with a serum dilution of 1:64 (cut-off). In cases with a positive reaction, serum serial dilutions at two-fold increments were tested up to the endpoint titer. Ten microliters of sera were diluted in Phosphate Buffered Solution (PBS), pH 7.2, at room temperature and were placed in wells on *T. gondii* slides containing RH tachyzoites as the antigen, and incubated. The slides were washed in PBS (pH=7.2), three times during ten minutes, followed by an incubation with rabbit antibody anti-sheep IgG conjugated for fluorescein isothiocyanate (Sigma Chemical, USA), diluted 1/250 in PBS containing 0.001% of Evans Blue (Sigma Chemical, USA). The slides were washed again, covered with buffered glycerin (pH 8.0) and a cover slip and then examined under a fluorescent microscope (Nikon Eclipse; objective: 40x) by one trained person. Positive and negative control was included in all the slides. For positive serum samples, diffuse or continuous peripheral membrane fluorescence was considered to be positive, but apical fluorescence alone was not. Serum samples that were positive with titers ≥ 1:64 were considered to be positive. Tissues from seropositive animals were submitted to PCR, histopathology and IHC.

DNA extraction and Polymerase Chain Reaction

DNA was extracted from tissues (cerebrum, cerebellum, pons, spinal cord and heart) using the commercial kit Wizard Genomic DNA Purification (Promega®, USA), in accordance with the manufacturer's instructions.

Samples for the polymerase chain reaction (PCR) were prepared for a final volume of 12.5 μL containing: 6.25 μL of Top Taq Master Mix (Qiagen; Germany), 0.5 μL of each primer (10 μM), 2.75 μL of ddH₂O and 2.5 μL of DNA. The positive control for *T. gondii* was obtained from a suspension of intraperitoneal lavage material from mice previously infected with the RH strain. For the negative control, 2.5 μL of ddH₂O was added, replacing the DNA. The primers used were TOX4 (5’-CGCTGAGG-GAGGAAGACGAAAGTTG-3’) and TOX5 (5’-CGCTGCAGACACAGTGCATCTGGATT-3’), thus obtaining a 529 bp fragment that is repeated 200- to 300-fold in the genome of *T. gondii*.

PCR was performed as described by Homon et al (2000) with some modifications: an initial cycle of 94 °C for 7 min (initial denaturation), followed by 35 cycles of 94 °C for 1 min (denaturation), 60 °C for 1 min (annealing) and 72 °C for 1 min (extension), followed by a final extension of 72 °C for 10 min. The PCR amplification was performed using an XP thermal cycler (Bioer Technology Co. Ltd., China).

The PCR product was subjected to electrophoresis. It was loaded into wells containing 2% agarose gel stained with blue-green dye I (LGC Biotecnologia, Brazil). The bands were viewed under ultraviolet light in a documentation system (L.PIX model, Locus Biotecnologia®).
**Immunohistochemistry**

The immunohistochemistry technique was performed as described by Galiza et al. (2014), with some modifications: we used the labeled streptavidin biotin (LSAB) alkaline phosphatase technique (Dako Corporation, USA). The histological sections with 4 μm fixed on silanized slides were dewaxed and hydrated. Inactivation of endogenous peroxidase was performed using 3% hydrogen peroxide solution in distilled water at room temperature for 30 minutes and washed in PBS (0.01 M; pH 7.4). For antigen retrieval, the sections were subjected to heating in a water bath with citrate buffer (pH 6.0; 10 mM) and were then placed in a microwave oven for 20 minutes at high power and washed twice in PBS afterwards. Non-specific binding was blocked by means of incubation with 5% skim milk in PBS for 30 minutes at room temperature. After the slides had been washed with distilled water, they were immersed in PBS and then incubated with primary *T. gondii* antibody (Goat polyclonal antiserum #210-70, VMRD, USA) diluted 1:200 in PBS (pH 7.4), in a moist chamber at 4 °C for overnight. After this period, the slides were washed in PBS and the commercial LSAB alkaline phosphatase (Dako Corporation, USA) kit was used. Subsequently, the sections were washed in PBS and the
Permanent Red (Dako Corporation, USA) (chromogen) was used in accordance with the manufacturer’s recommendations. Finally, the samples were counterstained with Harris hematoxylin (Sigma Chemical, USA), dehydrated, cleared in xylene (Vetec®, Brasil) and mounted using balsam (Labsynth, Brasil), under a cover slip.

**Histopathology**

Different histological sections of the cerebrum, cerebellum, spinal cord and pons were cut and dehydrated in alcohols with increasing concentrations (from 70% to 100%). These were then embedded in paraffin using routine processing. From each block, sections of 4 μm thickness were cut, deparaffinized, rehydrated and stained with hematoxylin-eosin. All the slide was analyzed for presence of tissue cysts or histopathological lesions (Olympux BX40, objective 40x).

**Statistical analysis**

The Cohen’s kappa coefficient (kappa) was used to assess the agreement between the techniques used. The kappa values were interpreted as described by Landis & Koch (1977): 0.00-0.20 = weak agreement; 0.21-0.40 = regular agreement; 0.41-0.60 = moderate agreement; 0.61-0.80 = good agreement; 0.81-1.00 = very good agreement. Negative values were interpreted as equal to 0.00.

**Results**

The overall seroprevalence of *T. gondii* to IgG antibodies was 14% (14/100). Six males and eight females were seropositive with the following titers: 64 (6/14), 128 (1/14), 256 (4/14) and 1024 (3/14). Among the IFAT-positive sheep, three males were PCR-positive for *T. gondii*. Parasite DNA was detected in the spinal cord, cerebellum, cerebrum and heart. Among the IFAT-positive samples, the histopathological predominant findings were mononuclear cell infiltration in heart (Table 1) and a perivascular cuff in cerebrum and cerebellum (Figure 2). Only one sample of cerebellar tissue showed positive IHC staining. There was a moderate agreement between PCR and IHC results (kappa = 0.44).

**Discussion**

The seroprevalence results from the present study are in accordance with other results previously described in Brazil (PEREIRA et al., 2012; ANDRADE et al., 2013). By means of IHC, *T. gondii* was found in central nervous system from asymptomatic sheep confirming the findings previously described by Esteban-Redondo et al. (1999), Motta et al. (2008) and Benavides et al. (2011). Weissmann (2003) recorded that the brain was considered the best tissue for sampling for immunohistochemical evaluation and confirmation of the diagnosis of toxoplasmosis. In the case of tissues from animals that were slaughtered or necropsied, it is useful to examine more than one tissue type, such as the brain, heart or other tissue in which the parasite is frequently encysted. In this study, *T. gondii* was also detected in heart of sheep, in accordance with other studies that suggest that this organ may be the most appropriate tissue for detecting *T. gondii* infection using IHC (SILVA, 2011; VILLENA et al., 2012). Rosa et al. (2001) stated that during the asymptomatic phase of infection in experimentally infected goats, the cerebrum and the spinal cord are the most suitable tissues for isolating *T. gondii*. In the present study (which examined cases of natural infection), the PCR and IHC techniques respectively detected parasite DNA and *T. gondii* cysts, in the cerebrum, cerebellum and spinal cord. It is not possible to compare results relating to natural and experimental infections, given that in cases of natural infection, the infection phase and the strain might not be determined and the number of cysts is lower.

Although the histological findings in the present study were nonspecific, such findings have been widely described in cases of *T. gondii* infection (MCGAVIN & ZACHARY, 2007). These histological alterations may differ between studies, especially with regard to the intensity of the mononuclear infiltrate observed in tissues targeted by the parasite. Silva (2011) reported congestion, followed by presence of inflammatory mononuclear cells or mild

<table>
<thead>
<tr>
<th>Sample</th>
<th>Number of positive samples/total number of animals</th>
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<tbody>
<tr>
<td>Heart</td>
<td></td>
</tr>
<tr>
<td>Mononuclear infiltration between the myocytes</td>
<td>8/14 (57.14%)</td>
</tr>
<tr>
<td>Cerebellum</td>
<td></td>
</tr>
<tr>
<td>Perivascular cuff</td>
<td>2/14 (14.28%)</td>
</tr>
<tr>
<td>Cyst</td>
<td>1/14 (7.14%)</td>
</tr>
<tr>
<td>Cerebrum</td>
<td></td>
</tr>
<tr>
<td>Focal congestion</td>
<td>1/14 (7.14%)</td>
</tr>
<tr>
<td>Perivascular cuff</td>
<td>4/14 (28.57%)</td>
</tr>
</tbody>
</table>

Figure 2. *Toxoplasma gondii* cysts in cerebellum section from IFAT-seropositive sheep for *T. gondii* IgG antibodies. Arrows: tissue cysts DAB-stained (objective 100x).
focal polymorphonuclear infiltrate. The complementary examination of the cerebrum may be useful, given that primary and secondary lesions by *T. gondii* may be found there. Gliosis around a central necrosis, sometimes mineralized with mild lymphocytic meningitis, represents a response to direct tissue damage caused by the parasite (MCGAVIN & ZACHARY, 2007).

In one case of presence of a cyst in the cerebellum of a male sheep, as confirmed by means of IHC and PCR, there was no inflammatory reaction in the vicinity. Dubey et al. (1998) reported that intact tissue cysts probably do not cause any damage to the tissue and can persist without causing any inflammatory response by the host. According to Weiss & Kim (2000), rupturing of tissue cysts induces a strong inflammatory response that results in formation of glial nodules in the cerebrum of chronically infected hosts.

Histopathological findings combined with IHC and clinical history allows for a definitive diagnosis of toxoplasmosis in most of cases (DAGLEISH et al., 2010).

Tachyzoites or cysts are difficult to identify in histological sections, especially when present in low numbers. Immunohistochemistry may facilitate the viewing of parasites in tissues (MCGAVIN & ZACHARY, 2007; WAREE et al., 2007), but in cases of natural infection, this technique may have low sensitivity, as observed in the present study, in which most of the samples that were positive in serological and PCR evaluations were negative in IHC. The sensitivity of IHC is affected by the parasite inoculation routes, the infective form used, the species of the host, the strain virulence and the random distribution of the parasite in different tissues of the host. Thus, it is important to include samples from different tissues in order to increase the sensitivity of the technique.

Using IHC, positive staining for *T. gondii* was observed in the brain tissue of 7.14% (1/14) of the seropositive sheep. This differed from result obtained by Silva (2011), among naturally infected sheep, who reported that the positive staining rate was 52% (12/25), using IHC. Among sheep in Río de Janeiro, Brazil, Silva et al. (2013) concluded that IHC was proven to be effective in making the definitive diagnosis of toxoplasmosis. This was not confirmed in the present study, because the agreement between the PCR and IHC results was moderate (k = 0.44). We therefore consider that the results were insufficient for this technique to be adopted in routine diagnostic laboratories. We recognize that there is some difficulty in identifying the infective forms of *T. gondii* in naturally infected animal tissues and that it is therefore more appropriate to use an association of direct diagnostic techniques in order to increase the chances of identifying the parasite.

The size of the sample examined has been described as one of the main limitations for detecting *T. gondii* infection in sheep tissue (ESTEBAN-REDONDO & INNES, 1998). It is possible that the parasite is present in unanalyzed parts of tissues and, consequently, a negative result in the sample does not necessarily mean that the tissue is negative.

Esteban-Redondo & Innes (1998) used PCR to detect the DNA of *T. gondii* in sheep inoculated with oocytes and observed positivity in 8/10 samples, especially among the sheep that received high infective doses. According to Dubey (2010), the PCR technique is able to detect only one parasite, thus providing a quick and efficient diagnosis, but the positive result depends on the protozoa in the sample.

**Conclusion**

This study identified the presence of *T. gondii* cysts in tissues from seropositive sheep that had been slaughtered in Alagoas state, northeastern Brazil. The use of different direct techniques in association enhanced the chances of finding the parasite in tissue samples.

**References**


Homan WL, Vercammen M, De Braekeleer J, Verschueren H. Identification of a 200- to 300-fold repetitive 529 bp DNA fragment in *Toxoplasma*


