Detection of *Toxoplasma gondii* in the reproductive system of male goats

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**Abstract**

Male goats of mating age serologically negative for *Toxoplasma gondii* were divided into three groups: GI – controls (placebo) (n = 2); GII – infected with 1 × 10⁶ tachyzoites (RH strains) (n = 2); and GIII – infected with 2 × 10⁵ oocysts (P strains) (n = 2). Clinical, hematological, parasitological and serological tests and studies of parasites in the semen through bioassay and polymerase chain reaction (PCR), and in reproductive organs (bioassay) were performed to assess toxoplasma infection. Serological titers peaked at 4096 in two animal groups infected with the protozoan. The bioassays allowed an early detection of protozoa in semen samples of tachyzoite-inoculated animals. *T. gondii* DNA was identified through PCR in the semen in five (Days 5, 7, 28, 49, and 70) and two (both at day 56) different days post-inoculation in GII and GIII animals, respectively. It was also possible to detect *T. gondii* DNA in reproductive organs (prostate pool, testicles, seminal vesicle and epididymis) of goats inoculated with either tachyzoites or oocysts. The present study suggests the possibility of venereal transmission of *T. gondii* among goats and it should be further assessed.

**Keywords:** Toxoplasma gondii, goats, reproductive systems, PCR and bioassay.

**Introduction**

*Toxoplasma gondii* is the causative agent of toxoplasmosis, a cosmopolitan zoonosis of medical and veterinary relevance leading to miscarriage and a congenital disorder in intermediary hosts (TENTER et al., 2000; MASSALA et al., 2003).

Brazil has a goat herd of over 13 million animals but little is known about the effects of toxoplasmosis among goats. Munday and Mason (1979) were the first to describe toxoplasmosis as an important cause of reproductive losses in goats. Although often unnoticed, this infection can cause significant damage in both young and adult animals (DUBEY, 1987). The main route of infection is ingestion of the parasite's sporulated oocysts present in the environment (DUBEY; BEVERLEY, 1988). Risk factors for *T. gondii* infection in goats include age, number of cats in the farm, and either no use of feeding troughs or use of wooden feeding troughs (CAVALCANTE et al., 2008).
The objective of this study was to detect the presence of *T. gondii* in semen samples of experimentally infected goats using bioassay as well as molecular and histopathology techniques.

### Material and Methods

#### 1. Experimental infection of goats with *T. gondii*

This study used “P” (JAMRA; VIEIRA 1991) and “RH” (SABIN, 1941) *T. gondii* strains kept at the Animal Health Research Center (CPPAR), School of Agrarian and Veterinary Sciences, Universidade Estadual de São Paulo, Jaboticabal campus, São Paulo, Brazil. “P” strains were genotypically characterized by using PCR-RFLP segment of locus SAG2 located in chromosome VIII as a genetic marker and classified as Type III. (BRESPIANI et al., 2009). “RH” strains were previously characterized as Type I by using the same analysis of loci SAG1, SAG2, new SAG2, SAG3, BTUB, GRA6, c22-8, c29-2, L358, PK1 and Apico (PENA et al., 2008; HERRMANN et al., 2010).

Six male goats of undefined breed serologically negative for *T. gondii* aged between one and two years were selected. Serology titers obtained by indirect immunofluorescence (IFI) were considered positive starting at a 1:16 dilution. The animals were identified, sorted and randomized into three groups: GI – non-inoculated controls (n = 2); GII – subcutaneously infected with 1 × 10^6 RH strain tachyzoites per animal (n = 2); and GIII – orally infected with 2 × 10^7 P strain oocysts per animal (n = 2). All goats were kept in proper individual stalls with water and food ad libitum. Serological tests were carried out during the experimental period as follows: acidified buffered antigen test (COLE et al., 1973) for leptospirosis, with 2 × 10^5 P strain oocysts per animal (n = 2).

Goats from Days 3 to 7 PI and then weekly until the end of the experiment. *T. gondii* presence was assessed by bioassays in mice (DUBIEY; SHARMA, 1980) and positive samples were tested through PCR.

### 3. Molecular detection of *T. gondii* DNA

The methods as proposed by Fuentes et al. (1996) were used for semen and tissue sample standardization. RH strain *T. gondii* tachyzoites were counted on a modified Neubauer chamber in dark field microscopy (400×) with at a concentration of 10^8 parasites.mL^-1. From this first dilution, 500 µL were diluted in 9.5 mL of sterile saline solution (SST), and dilutions of 10^9, 10^8, 10^7, 10^6, 10^5, 10^4, 10^3, 10^2, 10^1 and 10^0 parasites.mL^-1 of the sample were successively obtained. Dilutions were adjusted in 700 µL volume aliquots and DNA extracted from each sample. *T. gondii* DNA of the scales of dilution (positive control) and from semen samples for DNA detection were extracted according to the modified method as described by Leale et al. (1982). At the end of the experiment, goats were euthanized and specimens from testes, seminal vesicles, epididymides and prostates were collected for histopathology tests and for *T. gondii* isolation by bioassay (DUBIEY, 1980).

### Results and Discussion

Following inoculation with *T. gondii* tachyzoites or oocysts, toxoplasma infection of goats was confirmed by parasitemia as well as seroconversion of the inoculated animals. Hyperthermia (40.65 °C) on Day 5 PI was the most remarkable clinical sign seen in the group of animals inoculated with oocysts (Figure 1). Likewise, Nishi et al. (2001) reported that temperature increase is one of the most evident clinical signs in goats orally inoculated with 10^7 oocysts of *T. gondii*. Additionally, anorexia and lethargy were seen from Day 3 to 7 PI in all inoculated animals.

Thirteen parasitemic outbreaks were detected during the experimental period: one on Days 11, 21, 28, 49, 63, two on Day 56 and 70 and four on Day 14 PI. Parasitemic outbreaks during acute stage of disease have been described from Day 7 to 14 (CHHABRA et al., 1982) and Day 5 to 12 PI (NISHI et al., 2001).

Toxoplasma infection elicited a fast immune response in goats with antibody anti-IgG detection by IFI from Day 11 PI. An early humoral response was also detected by Nishi et al. (2001) on Day 10 PI, demonstrated through IFI (anti-IgG) test.

IgG detection started on Day 11 PI in both tachyzoite-inoculated goats (titers of 256 and 1024) and one oocyst-inoculated goat (256). The peak titer (4096) was detected on Days 21 and 28 PI in all animals of GI and GII, respectively. IgG titer decreased from Day 35 PI on, but remained high (1024) in all inoculated animals until the end of the experiment.

These results are partially inconsistent with that reported by Nishi et al. (2001) who detected serological titer peaks (IFI) at a later stage (after Day 35 PI) in goats infected with a different dose and strain of *T. gondii* (10^7 oocysts and AS 28 strain). Corroborating our findings, these authors found that serological titers remained at relatively high levels until Day 56 PI. Persistently high serological (IgG) levels until Day 70 PI found in the present

* Monoclonal anti-goat/sheep IgG-FITC antibody produced in mouse – GT-34- purified immunoglobulin, buffered aqueous solution (Sigma).
study can be explained by the fact that in chronic infections IgG antibodies can remain “active” for a long time compared to IgM immunoglobulins, notably identified only a few weeks after infection, as described in Dubey and Towle (1986).

Dubey and Sharma (1980) used a bioassay in mice to demonstrate the presence of *T. gondii* in the semen of three goats orally inoculated with $1 \times 10^6$ oocysts of GT-1 strain (isolated from goats). Protozoa were detected on Day 7 PI in semen samples from two animals and from Day 12 PI on in the third animal studied. In semen samples of these animals it was possible to identify parasite excretion on Day 59 PI. Despite differences between strains studied, in this study the protozoan was detected on Days 56 PI in semen samples from oocyst-inoculated animals.

Molecular detection of *T. gondii* DNA in semen samples from tachyzoite-infected animals in more experimental days (Days 5, 7, 28, 49 and 70) is similar to that found by Dubey and Sharma (1980). This technique also allowed to detecting *T. gondii* DNA in samples from two sporulated oocysts-infected animals on Day 56 DPI.

Bioassay-positive semen samples from animal 1 on Day 63 PI, animal 11 on Days 28 and 63 PI, and animal 47 on Day 70 PI were not PCR-positive for *T. gondii* (Figure 2). PCR results supported bioassay findings and confirmed that it is an useful ancillary tool for toxoplasma infection diagnosis in semen samples.

In contrast to Pescador et al. results (2007), histopathology tests did not identify any significant change related to *T. gondii* parasitism on tissues of the reproductive system. These authors, when evaluating tissues from goat miscarriage, found significant microscopic and macroscopic alterations like enlarged pale mesenteric lymph nodes with lymphoplasma cells infiltrate in brain and lungs.

Nishi et al. (2001) successfully isolated through bioassay *T. gondii* from brain tissue, lymph nodes, liver, kidneys, skeletal and cardiac muscles but they did not find any macroscopic lesions in these tissues. Similar findings were reported by Cavalcante et al. (2007) regarding protozoan isolation in skeletal muscle samples from goats slaughtered in the State of Ceará, Brazil.

**Conclusion**

*T. gondii* was isolated from semen samples of experimentally infected goats associated to tissue parasitism in specimens of the reproductive system diagnosed by bioassay and PCR techniques, suggesting the possibility of venereal transmission of this coccidium in goats.

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