Myenteric neuronal plasticity induced by *Toxoplasma gondii* (genotype III) on the duodenum of rats

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**ABSTRACT**

The effects of acute and chronic infection caused by *Toxoplasma gondii* on duodenal myenteric neurons were analyzed. Eighteen rats were assigned into four groups: Acute Control Group (ACG, n=4); Acute Experimental Group (AEG, n=4); Chronic Control Group (CCG, n=5); and Chronic Experimental Group (CEG, n=5). Rats from the AEG and CEG were inoculated orally with 10⁵ genotype III (BTU-II strain) tachyzoites of *T. gondii* isolated from a dog with neurological signs. Acute groups were killed after 24 hours after the inoculation and the chronic groups after 30 days. Whole-mount from the duodenum were stained with Giemsa. The population density of myenteric neurons, as well the body cell, nuclear and cytoplasmic area were analyzed. Both acute and chronic toxoplasmic infection did not provoke neuronal loss. On the other hand, plastic alterations were observed: decreasing of the nuclear and cytoplasmic area during the acute phase and neuronal hypertrophy during the chronic phase.

**Key words:** enteric nervous system, infection, parasite, toxoplasmosis.

**INTRODUCTION**

*Toxoplasma gondii* is an obligate intracellular protozoan, highly spread all over the world, which is responsible for an important zoonosis: toxoplasmosis. This disease is usually asymptomatic in humans; however, they are largely an accidental host. Domestic animals and livestock can be infected by this microorganism. In intermediate hosts (warm-blooded vertebrates except felines), the parasite rapidly disseminates throughout the body, right after the infection, by fast growing tachyzoites. In response to following immune responses, *T. gondii* differentiates into a slow-growing tissue stage named bradyzoites, which inhabit in tissue cysts within low-turnover cells (Weiss and Kim 2007). For this reason, neurons are normally encysted by this parasite, which explains the severe neurological disorders observed in immune-compromised patients.

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In addition, when *T. gondii* invades the host cell, it is known that a parasitophorous vacuole (PV) is formed, which has a membrane containing proteins (ROP2) anchoring the mitochondria of the host cell to the PV. This strategy contributes for the lysosome of the host cells not to merge with the PV in order to protect the intracellular development of the bradyzoites (Sinai and Joiner 2001). Therefore, cells with higher number of mitochondria (such as neurons) seem to be the primary target of this protozoan.

Besides, studies reported changes in the intestinal wall of infected animals (Shiraishi et al. 2009, Bonapaz et al. 2010, Da Silva et al. 2010a, b), possibly because the digestive tube is the route of entry of *T. gondii* (Weiss and Kim 2007). Such changes indicate the both digestion and absorption may be compromised, even in animals showing no clinical signs of infection. Additionally, the literature relates diarrhea in animals (Weiss and Kim 2007, Galli et al. 2008, Odorizzi et al. 2010), which indicates that the enteric nervous system (ENS) - the main autonomic nervous system subdivision that controls the digestive function – can be altered by the toxoplastic infection.

Experiments carried out by our group demonstrated that the myenteric plexus - from rats infected with the same *T. gondii* strain (BTU-II / genotype III) - shows different alterations depending of the analyzed region from the intestine. Thus, it is known for this strain that, during the acute phase, hypertrophy in the jejunal neurons (R.P. Torres et al., unpublished data), atrophy in those from the ileum and the colon, respectively (Sugauara et al. 2008). On the other hand, in the chronic phase, no morphometric alterations in the jejunal myenteric neurons (R.P. Torres et al., unpublished data), and atrophy and hypertrophy in those from the ileum and the colon, respectively (Sugauara et al. 2008). It is perceptible that the effects of acute and chronic infection caused by the BTU-II strain of *T. gondii* (genotype III) for the duodenal myenteric plexus of rats, and this was the objective of this study.

**MATERIALS AND METHODS**

This experimental protocol was previously approved by the Ethics Committee for Animal Experimentation of Univesidade Paranaense in accordance with the norms of Colégio Brasileiro de Experimentação Animal (COBEA). It also meets the orientations presented on the NRC Guide for the Care and Use of Laboratory Animals.

**EXPERIMENTAL OUTLINE**

For this fully randomized study, eighteen male 60-day-old Wistar rats (266.25±21.76 g) were used. They were assigned into four groups: Acute Control Group (ACG, n= 4), Acute Experimental Group (AEG, n=4), Chronic Control Group (CCG, n=5), and Chronic Experimental Group (CEG, n=5). NaCl solution was administered through gavage to the animals in the ACG and CCG. *T. gondii* tachyzoites (10⁴) from a genotype III (BTU-II) strain isolated from dog brains with neurological symptomatology (Da Silva et al. 2005) were orally administered to the AEG and CEG.

**EUTHANASIA AND COLLECTION OF BIOLOGICAL MATERIAL**

Acute groups were died after 24 hours, and the chronic groups after 30 days. In this moment, the animals were weighted and anesthetized by the intramuscular route using following protocol: 2% Acepromazine (1.26 mL/kg) + 10% Ketamine (1.26 mL/kg) + 2.5% Xylazine hydrochloride (0.42 mL/kg), and 1% Atropine sulfate (0.22 mL/kg). Blood samples were collected by puncturing the retro-orbital plexus of each animal of the chronic groups, and then used in serological analysis for antibodies against *T. gondii* by the direct agglutination method (Da Silva et al. 2002). All animals were died by anesthetic deepening, followed by laparotomy for the removal of the duodenum.
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OBTAINING THE WHOLE-MOUNT PREPARATIONS

Theses intestinal segments were measured with respect to length using a milimetric ruler. Then, washed in a 9%-NaCl solution, filled and immersed in a fixation solution containing acetic formol for 48h. The duodenal samples were inspected for gross pathology lesions. Next, they were dissected with the aid of a stereomicroscope by removing the mucosa and the submucosa. Therefore, the whole-mount used for this study were constituted by the external muscle (where the myenteric plexus is located), which was stained according to the Giemsa technique (Barbosa 1978).

QUANTITATIVE ANALYSIS

Duodenum of all animals was divided in three regions for morphometric and quantitative analysis: mesenteric (0°-60°; 300°-360°), antimesenteric (120°-240°), and intermediate region (60°-120°; 240°-300°) considering the mesenteric insertion as 0°. A MOTIC BL220A binocular microscope with a 40x objective was used in order to count neurons in 40 microscopic fields (0.21 mm²) of each region (mesenteric, antimesenteric and intermediate), totaling 25.2 mm² per animal. Neurons positioned at the edges of each field were counted in alternated fields.

MORPHOMETRIC ANALYSIS

The area of the body cell, cytoplasm and nucleus of 300 myenteric neurons (100 cells per region: mesenteric, antimesenteric and intermediate) from each intestinal segment collected (uniformly distributed around the intestinal circumference) was measured with software: Image Motic Plus, version 2.0. A microscope with a digital camera (Moticam 2000, 2.0 megapixel) attached to a computer was used. From these values, the neurons were assigned into classes considering the body cell area (50 µm² intervals) and the nucleus/body cell area ratio (0.05 intervals).

STATISTIC ANALYSIS

All data were initially submitted to the Kolmogorov-Smirnov test for the verification of their distribution type. Normal distribution data were expressed as mean±standard deviation, and dates with free distribution were expressed as median and percentiles 25 and 75 (P25; P75). The ANOVA one-way (data presenting normal distribution) and Kruskal-Wallis (data presenting free distribution) were used in order to compare the groups, by considering p<0.05 significant values.

RESULTS

Animals from the CEG were positive in serological analysis for antibodies anti-*T. gondii*. None of the rats showed diarrhea during the experiment nor pathological lesions were detected during inspection. No significant differences were observed in the body weight and duodenum length of rats in any of the groups (Table I). The neuronal quantitative analysis did not demonstrate alterations (Table II), and the staining technique used did not allow us to found tachyzoites or bradyzoites of *T. gondii* inside the neurons.

The acute infection provoked enlargement of the body cell and reduction of the nuclear and cytoplasm area of myenteric neurons. Additionally, it was observed that the enlargement of the cytoplasmic area and the preservation of the nuclear

<table>
<thead>
<tr>
<th>Group</th>
<th>Weight (g)</th>
<th>Duodenal length (cm)</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACG</td>
<td>219.33±16.74</td>
<td>9.000±0.000</td>
</tr>
<tr>
<td>AEG</td>
<td>227.07±16.25</td>
<td>7.875±0.629</td>
</tr>
<tr>
<td>CCG</td>
<td>368.30±16.83</td>
<td>6.800±1.090</td>
</tr>
<tr>
<td>CEG</td>
<td>368.30±16.83</td>
<td>6.800±1.710</td>
</tr>
</tbody>
</table>

There is no significant difference between control and experimental groups (p > 0.05).
area resulted in the increasing of the body cell area, during the chronic infection (Table II). It should be remarked that specific alterations were observed only in the acute infection: (1) when the distribution of neurons according to different classes of body cell size was evaluated, an increased prevalence of smaller neurons (151-200 µm$^2$) and a reduced prevalence of larger cells (301-350 µm$^2$) was detected (Figure 1); (2) with relation to the nuclear area, animals from the AGE showed an increasing number of neurons in which the nucleus took 26 to 30% of the body cell, and a reduction in the number of those in which the nucleus took more than 36% of the body cell (Figure 2).

### Table II

<table>
<thead>
<tr>
<th>Group</th>
<th>Number of neurons (in 25.2 mm$^2$)</th>
<th>Body cell area (µm$^2$)</th>
<th>Nucleous area (µm$^2$)</th>
<th>Cytoplasm area (µm$^2$)</th>
<th>Nucleous/body cell ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACG</td>
<td>2013.5 (1584.5; 2255.0)</td>
<td>242.4* (223.9; 306.6)</td>
<td>89.9* (75.7; 104.6)</td>
<td>172.10* (144.2; 205.5)</td>
<td>0.34* (0.31; 0.38)</td>
</tr>
<tr>
<td>AEG</td>
<td>2368.0 (1533.0; 3387.5)</td>
<td>228.2* (189.5; 272.7)</td>
<td>72.3* (60.8; 84.2)</td>
<td>159.65* (128.0; 197.2)</td>
<td>0.31* (0.27; 0.35)</td>
</tr>
<tr>
<td>CCG</td>
<td>3530.0 (3470.0; 3752.9)</td>
<td>183.1* (151.8; 218.6)</td>
<td>55.9 (47.1; 67.1)</td>
<td>125.70* (101.0; 154.9)</td>
<td>0.31* (0.27; 0.35)</td>
</tr>
<tr>
<td>CEG</td>
<td>3331.0 (3110.0; 3392.0)</td>
<td>197.0* (158.6; 242.3)</td>
<td>56.4 (47.4; 66.4)</td>
<td>139.50* (107.7; 176.2)</td>
<td>0.29* (0.25; 0.32)</td>
</tr>
</tbody>
</table>

Values presented as median (P25; P75). Values in the same column are significantly different (p <0.05).

![Figure 1 - Histogram of the body cell area of duodenal myenteric neurons of rats from control (ACG and CCG) and experimental (AEG and CEG) groups infected by genotype III (BTU-II strain) tachyzoites of Toxoplasma gondii. Columns of the same class with one asterisco (*p<0.05) and with two asteriscos (**p<0.01) are significantly different.](image-url)
DISCUSSION

Only animals from the CEG showed positive serology for *T. gondii*, demonstrating the efficiency of the experimental model used in this study to induce the toxoplastic infection. The strain is one of the most important factors to determine the infectivity and pathogenicity of the parasite (Dubey 1996). Although using a genotype III strain - considered of low infectivity when compared to genotype I strains (Howe et al. 1997) - it was effectively infective. It is possible consider that the infection was caused by a small number of infective forms once tachyzoites showed low infectivity after gastric digestion, demonstrating low resistance to pepsin (Dubey 1998). However, not all tachyzoites were killed by gastric juice once the presence of antibodies by serological analysis, demonstrating that infection occurred.

*T. gondii* can cause diarrhea on dogs, chickens, pigs (Freyre and Falcon 1989), rodents (Dubey and Frenkel 1973) and immunocompromised humans (Boothroyd and Grigg 2002), with intestinal ulcers evolving to lethal ileitis in mice (Mennechet et al. 2002). However, neither diarrhea nor gross pathology lesions were observed in this study. These results indicate that, although the experimental model used in this study was enough to induce infectivity, the pathogenicity was low. By this way, no changes in the body weight and duodenum length and the absence of disease signs both in the chronic and acute phases of the infection.

Other studies of this same research group using the same experimental conditions showed a reduction of the jejunum-ileum area (R.P. Torres et al., unpublished data) and of the length and width of terminal ileum (Sugauara et al. 2008) caused possibly by atrophy or hypoplasia of the organ. Therefore, comparisons between organ areas obtained in previous and in this study show how singular organs respond differently to toxoplastic infection.

The duodenum is a short organ with pH 6.6, while pH in other parts of the small intestine ranges from 6.9 to 7.5 (Guyton and Hall 2002). The Brünner glands located in the submucosa of
this organ are responsible for the production of a thick mucus layer that covers its internal surface. Therefore, the lower pH and the thick mucus layer may have made it difficult for the parasite to gain access to the enterocytes, reducing in this way the number of microorganisms available to cause direct and indirect lesions.

Alterations were not detected in quantitative analysis of myenteric neurons. Other studies of this same research group also demonstrated that the number of myenteric neurons remains constant in rats infected with *T. gondii*, no matter the duration of the infection, the strain used or the organ studied (Sugauara et al. 2008, 2009, Barbosa et al. 2009, Soares et al. 2009, Pereira et al. 2010). On other hand, in a parallel study during the chronic phase of the toxoplasmic infection, R.P. Torres et al. (unpublished data) observed a reduction of the jejunum-ileum without proportionally greater myenteric neuronal density, reporting a 17.54% reduction in the neuronal population. In the present study, as there was no reduction in intestinal area, the maintenance of density demonstrates that these neurons did not die due to infection. Would duodenal myenteric neurons be protected from the toxoplasmic infection?

Tachyzoites or tissue cysts of *T. gondii* were not detected inside the neurons. It is well known that, during the acute infection, parasitemia is observed after the parasite penetration in the organism, and immunity is activated followed by the formation of cysts in different tissues (Frenkel et al. 1975). Cysts characterize the chronic infection persisting from months to years and sometimes throughout the lifetime of the host. In parallel with the production of tissue cysts parasitemia gradually decreases and disappears, not been found in the blood of rats after 60 days of infection possibly due to macrophage activity characterizing a natural resistance of rats (Guerrero 1995). During the chronic phase of infection, tissue cysts are commonly found in nervous tissue (Guerrero and Chinchilla 1996), however, it is not common to find them in histological sections (De Champs et al. 1998). The absence of tissue cysts in the present study may indicate their low prevalence in the enteric nervous system, possibly due to a series of factors such as the resistance of rats (Guerrero 1995) and the resistance of neurons (Lüder-Carsten et al. 1999).

After ingestion of the parasite, *T. gondii* penetrates the epithelial cells of the intestine, being able to reach different organs such as the liver and the heart in only few hours after infection (Barragan and Sibley 2003). In this way, the immune system quickly responds inducing an answer characterized by the differentiation of a T-cell subpopulation called Th1. Differentiation of CD4 T-cells is activated by antigens of *T. gondii* in Th1 effector cells. When toxoplasmosis ensues, the parasite binds to receptors on macrophages and dendritic cells stimulating rapid production of IL-12 cytokine, essential for the stimulated production of IFN-γ. Rapid induction of IL-12 production by the protozoan may contribute for the control of multiplication and diminishing of virulence (Robben et al. 2004). This may be one of the reasons why the number of enteric neurons in the present study was preserved.

Although plastic responses induced by *T. gondii* in myenteric neurons have already been described (Sugauara et al. 2008, 2009, Barbosa et al. 2009, Soares et al. 2009, Odorizzi et al. 2010, Pereira et al. 2010), our results demonstrate that these cells did not undergo significant changes, suggesting that duodenal myenteric neurons are more resistant to infection or are less accessible to pathogens. Therefore, in order to prove this hypothesis further studies using different strains and techniques and studies upon the cells involved in this process are necessary.

There are evidences that the nervous system of different species may show diverse responses to infection by tachyzoites of *T. gondii* (Halonen et al. 1996). In studies using nervous cells from the brain cortex of rats was proved that neurons and astrocytes
are the main host cells involved in the intracerebral proliferation phase (when tachyzoites are observed), and that the microglial cells inhibit parasite growth, having toxoplasmicidal activity, hence, the propagation of \textit{T. gondii} in the CNS is reduced (Lüder-Carsten et al. 1999). As myenteric ganglia do not have microglial cells, the protection of the nervous system located inside the intestinal wall depends on the immune cells found in this organ (Furness 2006).

Even though neuronal death was not observed, they may have been affected by infection responding with changes in size to the direct stimulation by the parasite or to the indirect stimulation by inflammatory cytokines. These changes were probably a reflex of modifications in cell metabolism. During the acute phase of infection the nuclear area was reduced and the body cell area enlarged. These findings revealed that acute infection may have repressed the synthesis of nuclear and cytoplasmatic proteins leading to cell volume changes, probably indicating cell injury as described by Sugauara et al. (2008, 2009). It was demonstrated that the jejunal myenteric neurons presented enlargement of the body cell area, nuclear area, and cytoplasm area, which was attributed to an acceleration of cytoplasmatic and nuclear metabolism (R.P. Torres et al., unpublished data), probably induced by intracellular \textit{T. gondii} growth which depends on the increasing of the cell metabolism in the host (Carruthers and Suzuki 2007) or by the indirect action of inflammatory cytokines.

In the chronic phase was observed that the body cell and the cytoplasmic area were enlarged, whereas the nuclear area was preserved, indicating that duodenal infection influenced the metabolism of these neurons. Therefore, the interaction between the parasite and the myenteric cell components needs to be further investigated. In previous studies with jejunum and ileum using different strains under chronic infection, results were different from those found in the duodenum. In these studies a reduction of the nuclear and neuronal areas was observed (Sugauara et al. 2008, R.P. Torres et al., unpublished data).

When the neuron frequency assigned into classes according to the body cell area was analyzed most of the cells were concentrated in the classes between 151 and 300 $\mu$m$^2$, in both the control and the experimental group. When the consequences of infection on this distribution, changes were only observed in the acute phase of the experiment in which the prevalence of neurons increased between the classes 151-200 $\mu$m$^2$ ($p<0.05$), and decreased between the classes 301-350 $\mu$m$^2$ ($p<0.01$).

In concern about the distribution of frequency considering the nucleus/body cell area ratio, most of neurons of all groups had nuclei that took from 26 to 35% of the body cell, whereas in the acute infection there was a significant increasing in the number of neurons in the classes between 0.26-0.30, and a reduction in the prevalence of neurons in the classes between 0.36-0.40 ($p<0.05$), and $>0.41$ ($p<0.05$). However, further studies using transmission electron microscopy are necessary in order to elucidate which cell element induced morphometric alterations in enteric neurons infected by \textit{T. gondii}.

In the present study was demonstrated that the acute and chronic infection caused by genotype III tachyzoites of \textit{Toxoplasma gondii} did not provoked alterations in the number of myenteric neurons, but induced to plastic changes such as decreasing of the nuclear and cytoplasmatic area during the acute phase and neuronal hypertrophy on duodenum of rats.

ACKNOWLEDGMENTS

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RESUMO

Foram analisados os efeitos da infecção aguda e crônica provocada pelo \textit{Toxoplasma gondii} sobre os neurônios miêntéricos do duodeno. Dezio ratos foram divididos em quatro grupos: Grupo Controle Agudo (GCA, n= 4), Grupo Experimental Agudo (GEA, n=4), Grupo Controle Crônico (GCC, n=5) e Grupo Experimental Crônico.
(GEC, n=5). Os animais do GEA e GEC receberam por via oral 10⁵ taquizoítos de *Toxoplasma gondii* da cepa BTUII (genótipo III) isolada de um cão com sintomatologia neurológica. Os grupos agudos foram submetidos à eutanásia após 24 horas e os crônicos após 30 dias da inoculação. Preparados totais do duodeno foram corados com Giemsa. A densidade populacional dos neurônios e citoplasma foram analisados. Ambas, as infecções toxoplásmicas aguda e crônica não provocaram a perda neuronal. Por outro lado, alterações plásticas foram observadas: diminuição da área nuclear e citoplasmática durante a fase aguda e hipertrofia neuronal durante a fase crônica.

**Palavras-chave:** sistema nervoso entérico, infecção, parasita, toxoplasmose.

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