Neospora caninum as causative agent of bovine encephalitis in Brazil

Neospora caninum como agente causal de encefalite bovina no Brasil

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Abstract

For supporting the Brazilian bovine encephalitis surveillance program this study examined the differential diagnosis of Neospora caninum in central nervous system (CNS) by histological analysis (HE staining), immunohistochemistry (IHC), and nested-PCR using a set of primers from the Nc5 region of the genomic DNA and ITS1 region of the ribosomal DNA. A sample of 302 cattle presenting neurological syndrome and negative for rabies, aged 0 to 18 years, from herds in 10 Brazilian states was evaluated for N. caninum from January 2007 to April 2010. All specimens tested negative with IHC and nested-PCR using primers from the ITS1 region of ribosomal DNA, while two positive cases (0.66%) were found using primers from the Nc5 region of genomic DNA: a 20 month-old male and a 72 month-old female, both from São Paulo State. Only the male presented severe multifocal necrotizing encephalitis associated with mononuclear cell infiltration, a pathognomonic lesion caused by parasites of the family Sarcocystidae, and only this case was associated with N. caninum thus representing 0.33% positivity. Future studies should explore the association of IHC and nested-PCR with real-time PCR, a quantitative method that could be standardized for improving the detection of N. caninum in bovine CNS specimens.

Keywords: Histopathology, immunohistochemistry, nested-PCR, Nc5, ITS1.

Resumo

Este estudo contribuiu para o programa de vigilância epidemiológica de encefalite bovina no Brasil realizando o diagnóstico diferencial de Neospora caninum no sistema nervoso central (SNC) por análise histológica (coloração HE), imunohistoquímica (IHC) e nested-PCR utilizando-se primers da região Nc5 do DNA genômico e da região ITS1 do DNA ribossomal. Um total de 302 amostras de bovinos com síndrome neurológica, negativos para raiva, na faixa etária de zero a 18 anos, provenientes de rebanhos de 10 estados brasileiros foi avaliada para N. caninum no período de janeiro/2007 a abril/2010. Todas as amostras foram negativas na IHC e na nested-PCR usando-se primers da região ITS1 do DNA ribossomal, enquanto dois casos (0.66%) foram positivos à nested PCR, usando-se primers da região Nc5 do DNA genômico: um macho de 20 meses de idade e uma fêmea de 72 meses de idade, ambos do Estado de São Paulo. Apenas o macho apresentou severa encefalite multifocal necrotizante associada com infiltrado inflamatório mononuclear, lesão patognomônica causada por parasitas da família Sarcocystidae, mostrando que apenas este caso de encefalite foi associado à infecção por N. caninum, representando 0,33% de positividade. Sugere-se em estudos futuros utilizar também a PCR em tempo real para detecção do parasito.

Palavras-chave: Histopatologia, imunohistoquímica, nested-PCR, Nc5, ITS1.

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Introduction

*Neospora caninum* is a protozoan belonging to the phylum Apicomplexa, family Sarcocystidae, that mainly causes abortion in cattle, but its impact as causative agent of neurological syndromes has been poorly documented (Dubey; Schares, 2006).

Infected calves may be born underweight, unable to rise and with no clinical signs of disease. Hind limbs, forelimbs or both may be flexed or hyper-extended. Neurologic examination may reveal ataxia, decreased patellar reflexes, and loss of conscious proprioception. Calves may show exophthalmia or asymmetric eyes. *N. caninum* occasionally may cause birth defects, including hydrocephalus and narrowing of the spinal cord (Barr et al., 1991, 1993; Dubey et al., 1990, 1998a; Dubey; Lahunta, 1993; De Meerschman et al., 2005).

There are few reports of occurrence of the parasite in congenitally infected calves with neurological signs (Dubey; Schares, 2006). Encephalomyelitis was the predominant lesion in live born calves but clinically affected or that developed clinical disease soon after birth and were necropsied by 2 weeks of age (Parish et al., 1987; O’Toole; Jeffrey, 1987; Dubey et al., 1989, 1992; Barr et al., 1991, 1994; Magnino et al., 1999; De Meerschman et al., 2005).

In Brazil, cases of neurological syndrome associated with *N. caninum* in calves have been reported. Locatelli-Dittrich et al. (2003) isolated the protozoan in a 3-month-old Jersey calf in Paraná State, which presented with blindness. Santos et al. (2006), also in Paraná State, reported the occurrence of encephalitis in a 21 day-old Jersey calf infected with *N. caninum*.

The World Organization for Animal Health requires the diagnosis of causative agents of encephalitis for compliance with international sanitary standards for trade of animals and animal products (OIE, 2010). Many countries have implemented new diagnostic techniques and applied them in epidemiological studies in order to participate in the global marketplace and to guarantee food safety.

The Brazilian Ministry of Agriculture, Livestock, and Food Supply (MAPA) coordinates the National Program for Herbivore Rabies Control (PNCRH), and for encephalitis and encephalopathy epidemiological surveillance (BRASIL, 2009). In order to comply with the PNCRH, laboratories that perform rabies diagnoses are encouraged to conduct differential diagnosis of other etiological agents causing neurological syndromes when the CNS is negative for rabies virus. A preliminary study using PCR of neosporosis as a neurological syndrome in cattle in São Paulo State found 0.7% (1/131) of CNS specimens to be positive (Del Fava; Pituco, 2010). Patricio (2008) assessed 253 brain specimens from 161 adult cattle and 92 calves with neurological signs from various regions of Paraná State, and detected 28 (11.07%) positive specimens by PCR (primers Np21 and Np6) as follows: five 0–12 months old, six 12–24 months old, two 24–36 months old, and 15 > 36 months old.

Immunohistochemistry (IHC) is highly specific when used to identify *N. caninum* in tissue sections (Lindsay; Dubey, 1989), but must be interpreted in conjunction with histological analysis, i.e., the presence of the parasite in tissue must be associated with inflammatory lesions and tissue necrosis (Barr et al., 1994; Anderson et al., 2000). To improve the sensitivity and specificity of *N. caninum* diagnosis, several PCR protocols have been developed focusing on the ITS1 and NC5 specific sequences to these protozoa, such as hemi-nested and nested-PCR (Collantes-Fernández et al., 2002; Cabral et al., 2009).

The aim of this study was to examine the differential diagnosis for *N. caninum* in CNS of cattle with neurological syndrome that tested negative for rabies using hematoxylin-eosin (HE) staining of paraffin embedded tissue, immunohistochemistry (IHC), and nested-PCR using primers from the NC5 region of the genomic DNA and ITS1 region of the ribosomal DNA.

Material and Methods

All procedures conformed to animal experimentation ethical principles adopted by the Brazilian College of Animal Experimentation (COBEA), and approved by the Animal Experimentation Ethics Committee of the Instituto Biológico (CETEA-IB nº 092/09).

Three hundred and two specimens from bovid CNS sent from January 2007 to April 2010 to the Centro de Pesquisa e Desenvolvimento de Sanidade Animal do Instituto Biológico were initially examined for rabies by immunofluorescence (FAT) according to Dean et al. (1996) using anti-rabies conjugate labeled with fluorescein isothiocyanate (Sanoft®). Specimens that were negative for rabies were examined for *N. caninum* infection by histological methods (HE and IHC), and two protocols of nested-PCR.

Tissue specimens were fixed in 10% buffered formalin, embedded in paraffin, and 4-μm sections were prepared and stained with HE for routine histological examination. A second set of sections were mounted on slides, along with controls, and processed for IHC according to Cabral et al. (2009) using avidin-biotin-complex peroxidase (Vector Elite ABC-peroxidase®). As a positive control, goat fetal heart was taken from a pregnant goat that had been inoculated with the Illinois-NC strain of *N. caninum*. The IHC negative control was not exposed to the primary antibody. All slides were hydrated, and endogenous peroxidase was blocked by a combination of 10 mL of 30% hydrogen peroxide, 10 mL methanol, and 80 mL buffer. Slides were placed in a microwave oven at 800 watts for 10 minutes for antigen retrieval. Non-specific reactions were blocked with 10% milk powder dissolved in PBS.

The primary polyclonal antibody, anti-*N. caninum* rabbit hyper-immune serum, was diluted at 1:2000. The DAB chromogen (Data Sheet-Liquid DAB Substrate Pack, concentrated – VECTORS) was used for display and disclosure of the reaction. Sections were counterstained with Mayer’s hematoxylin and examined microscopically.

Genomic DNA was extracted from fresh and frozen tissue using Wizard Genomic DNA Purification Kit – Promega®, following the manufacturer’s instructions. One hundred μL of DNA hydration solution was added to the pellet, which was then heated for 1 hour at 64 °C. Specimens were stored at –80 °C for nested-PCR. Two nested-PCR protocols were conducted using primers from the NC5 region of the genomic DNA (genomic
nested-PCR) and ITS1 region of the ribosomal DNA (ribosomal nested-PCR). For the genomic nested-PCR, pairs of NP21 and NP6 primers (5’ GGTTGTCGCTCCAACCTCTGTGAAC 3’ - 5’ CTCGCACGCTAAGCTCGTCTCT 3’) were used to amplify the 337 bp DNA fragment and NP6-NP7 (5’ CATGCTACGTTGTTTCTCTC 3’ - 5’ GGTTGAACCGAGGGAGTTG 3’), which amplified 227 bp of the Nc5 gene of *N. caninum* (Hughes et al., 2006). Five µL of the DNA sample were added to the PCR mix making a final 20 µL solution with commercial buffer from a Promega PCR Master Mix™ kit (M7502) and 0.2 µM of each primer. The amplification was a modification of Hughes et al. (2006). The first amplification was 1 cycle of 95 °C for 5 minutes, 40 cycles of 94 °C for 1 minute, 60 °C for 1 minute, 72 °C for 1 minute, 1 cycle of 72 °C for 10 minutes, and maintenance at 4 °C. The second amplification was the same using 5 µL of the first amplification product. For ribosomal nested-PCR, pairs of JB1 - JB2 primers (5’AGGAAGGAGAAGTCGTAACAGG3’ - 5’GAGCCAAGACATCCATTGC3’) were used to amplify the 500 bp DNA fragment and SF1 - SF2 (5’ CCTGTAGATCGTTATCGCCTT3’ - 5’TCTTTTCCCTAAACCGCTATCC3’) to amplify 250 bp of the ITS1 region of ribosomal DNA from *N. caninum* (Barratt et al., 2008). Five µL of the DNA sample were added to the PCR mix, and a final 20 µL solution was made with buffer from the commercial kit Promega PCR Master Mix™ (M7502) and 0.5 µM of each primer. The amplification was a modification of Barratt et al. (2008). The first amplification was: 1 cycle of 95 °C for 5 minutes, 40 cycles of 94 °C for 1 minute, 60 °C for 1 minute, 72 °C for 1 minute, 1 cycle of 72 °C for 5 minutes, and maintenance at 4 °C. The second amplification using 5 µL of the first amplification product was: 1 cycle of 95 °C for 5 minutes, 30 cycles of 94 °C for 1 min, 60 °C for 1 minute, 72 °C for 1 minute, 1 cycle of 72 °C for 5 minute, and maintenance at 4 °C. The product of each amplification was applied to 2% agarose gel prepared in TAE 1X buffer (LGC®) and subjected to electrophoresis at a constant voltage of 100 V for 1 hour. The amplicons were visualized with ultraviolet light after ethidium bromide (5 µg.mL−1) staining. The DNA fragment size was compared with a standard molecular weight (100 bp DNA ladder - FERMENTAS®).

**Results, Discussion and Conclusions**

A total of 302 CNS specimens collected from cattle with neurological syndrome were analyzed. Convenience sampling was used, and diagnosis of encephalitis due to *N. caninum* was based on the presence of histological lesions, positive reaction with IHC, and two protocols of nested-PCR. According to Dubey et al. (1998b), tachyzoites or cysts of *N. caninum* are rarely sufficiently numerous to be found in all histological sections from an animal, requiring the use of specific and sensitive tests to detect the parasite. For this reason, histological findings should be interpreted as complementary to tests such as IHC and nested-PCR to identify the causative agent. *N. caninum* causes multifocal necrosis in the CNS associated with mononuclear inflammatory infiltrate (Pescador et al., 2007; Cabral et al., 2009), and histological examination was used as a first screening method for lesions in the specimens. The most frequent histopathological changes found by HE were non-purulent meningoencephalitis in 24% (72) of the specimens studied, with none of them IHC-positive for *N. caninum*. The IHC protocol used negative control (positive block without primary antibody), and it did not show non-specific staining, while the control evidence disclosed cysts with typical morphological structures intensely stained by DAB, confirming it as a very specific method, as reported by Cabral et al. (2009).

The genomic nested-PCR Nc5 revealed two positive cases (0.66%) (Figure 1), while the ribosomal nested-PCR ITS1 was negative. The sequence of primers from the NC5 region was not consistent with the genomes of *Toxoplasma gondii*, *Sarcocystis caprata*, *S. cruzi*, *S. mierchiana*, *S. moulei*, *S. tenella*, or *Hammondia hammondi* (Hughes et al., 2006), showing the specificity of the primers for *Neospora*.

The low frequency of CNS positive specimens for *N. caninum* found in this study agrees with that reported by Del Fava and Pitucu (2011), who found 0.7% of neosporosis by PCR in CNS of 131 cattle with neurological syndrome that tested negative for rables. In a similar study in the Paraná State, using a set of primers of the NC5 region (Np21/Np6) and rables negative bovine CNS specimens, Patrício (2008) found 11.07% positivity for *N. caninum*, though a histological evaluation to differentiate inflammatory lesions characteristic of *N. caninum* was not performed. It is thus not possible to determine if the positive nested-PCR represented disease (neurological syndrome) or latent infection.

In the present study, specimens came from 10 Brazilian states, of which 88% (265) were from São Paulo, including the two animals testing positive with nested-PCR of the NC5 region.

The most predominant breeds were zebu (*Bos indicus*, 42.0%, 126) and cross-breeds (28.0%, 86), followed by *Bos taurus* (13.0%, 38). In 17.0% (52) of the cases, the breed was unknown. The two animals (0.66%) positive by genomic nested-PCR were cross-breeds. Most cattle were female (69.0%, 208 versus 27.0%, 82 males); gender was unknown in 4.0% (12). The two positive animals were a male (0.33%) and a female (0.33%).

The animals were aged 0 to 18 years. Most were adults over two years old (59.3%, 179), with 34.4% (104) younger than two years, and only 1.7% (5) were <1-month– old calves. Age was unknown for 6.3% (19). The low prevalence of disease in this study may be associated with the low number of animals at the age of risk, given that several authors report that calves with clinical neosporosis die within the first four weeks of life (Dubey et al., 1992; Barr et al., 1993; Dubey; Lindsay, 1996).

The female testing positive by PCR, a 72 month-old cross-breed, did not exhibit pathological lesions in the CNS, and this may be considered a *N. caninum* latent infection. The positive PCR 20 month-old cross-breed male showed clinical signs of neurological syndrome such as sialorrhea, incoordination, recumbency and paddling, and opisthotonus, and died in four days. Histological examination of the CNS revealed severe multifocal necrosis in the neuropil (Figure 2), meningeal mononuclear infiltrate (Figure 3), and mononuclear perivascular cuffing (Figure 4). The clinical signs, histopathology, and PCR positivity implied that only the young male had encephalitis associated with *N. caninum*; therefore, the
proportion that best represents the positive cases of neurological syndrome caused by *N. caninum* in this study is 0.33% (1/302). This is the first report in Brazil of a bovine older than one year diagnosed with *N. caninum*-associated encephalitis.

Recent studies have associated inflammatory changes in the CNS identified by histology, positivity to molecular techniques and IHC to confirm active infection of *N. caninum* (OKEOMA et al., 2004; DUBEY; SCHARES, 2006; CABRAL et al., 2009). When neosporosis is studied in mature animals, histopathologic findings should be combined with molecular analysis since the parasite may cause unrecognized chronic infection, or the animal may be an asymptomatic carrier, and no lesions are found because the animal is immunologically protected. The balance between the tachyzoite’s ability to penetrate and multiply into host cells and the host’s ability to inhibit parasite multiplication was discussed by Buxton et al. (2002) and Hemphill et al. (2006). The low prevalence of *N. caninum* in the CNS of young and adult cattle suggests that the immune system responds efficiently to *N. caninum* through humoral immune response mediated by secreted antibodies, and by cellular immune response mediated by Th1 and Th2 (YAMANE et al., 2000; INNES et al., 2005; PINHEIRO et al., 2006; NISHI et al., 2009).

The present study applied a protocol suggested by Baszler et al. (1999) and Cabral et al. (2009) for routine diagnosis of *N. caninum* in aborted bovine fetuses using histopathology, PCR, and IHC. The majority of CNS specimens sent to the Instituto Biológico for histopathological examination of encephalitis were fixed in formalin, but some were chilled, or frozen, preventing histopathological and IHC diagnosis. Freezing and autolysis destroys the architecture of parasite cells, which can explain the non-visualization of parasitic cysts or tachyzoites by IHC,
a technique that combines morphology and antigen binding antibodies with marked with chromogen. Immunohistochemistry is a specific technique that has low sensitivity and can be effective when applied to intact tissues and those fixed in formalin (EPERON et al., 1999). Tissue fixation with formalin forms multiple connections with proteins (methylene bridges), which masks _N. caninum_ epitopes, blocking the binding of antibodies to the antigen. Enzymatic or heat methods should be used for antigen retrieval to destroy methylene bridges and improve immunoreactivity (BOENISCH, 2005; KEY; BOENISCH, 2006; RAMOS-VARA et al., 2008). In the present study antigen retrieval from CNS-tissue containing paraffin blocks was improved by heating the slides in microwave oven, as described by Cabral et al. (2009).

Ugglå et al. (1998) did not find either histopathological lesions or the parasite in brain specimens using IHC in inoculated calves; however, parasite DNA was detected, confirming that the test has greater sensitivity. An additional factor that may cause inconsistent detection rates of IHC and nested-PCR is the preservation of CNS specimens. Chilling, freezing and autolysis affect histological analysis (HE and IHC), and nested-PCR is a useful tool under these conditions. Because veterinarians use different methods for preservation of CNS specimens that are sent to laboratorial analysis, it is required for the detection of the parasite the association of techniques that are specific, such as IHC, or nested-PCR that are both specific and sensitive, preferably in conjunction with identification of histopathological lesions (SAGER et al., 2001). As autolysis is accelerated in the CNS, future studies should explore the association of IHC and nested-PCR with real-time PCR, a quantitative method (COLLANTES-FERNÁNDEZ et al., 2002) that could be standardized for improving the detection of _N. caninum_ in CNS specimens of bovine with neurological syndrome. A manual for collecting, preserving and shipping CNS specimens for clinical encephalitis cases recommends that those from anatomic areas of CNS for microbiological and parasitological examination should be refrigerated, whereas those for histopathology examination must be fixed in 10% formalin, and only after that specimens should be sent for laboratory analysis (PITUÇO et al., 2010).

The need for diagnosing causal agents of encephalitis is supported by the number of histopathological lesions that were found in CNS specimens undergoing routine analysis at the Instituto Biológico, i.e., 40% (690) of cattle testing negative for rabies presented non-purulent meningoencephalitis (DEL FAVA et al., 2007). Histological, microbiological, and parasitological analyses are important for the differential diagnosis of etiological agents of encephalitis, and veterinarians should be encouraged to increase the number of CNS specimens sent for screening.

With respect to _N. caninum_, based on the results of this study, sampling should include animals aged 0 to 2 years to provide new epidemiological data, support the encephalitis control program (BARROS; MARQUES, 2003; DEL FAVA et al., 2007; BRASIL, 2009), and determine the actual occurrence of _N. caninum_ as a causal agent of encephalitis in Brazilian herds, contributing to the PNCRH surveillance.

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