

# Serologic evidence of the recent circulation of *Saint Louis encephalitis virus* and high prevalence of equine encephalitis viruses in horses in the Nhecolândia sub-region in South Pantanal, Central-West Brazil

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*As in humans, sub-clinical infection by arboviruses in domestic animals is common; however, its detection only occurs during epizootics and the silent circulation of some arboviruses may remain undetected. The objective of the present paper was to assess the current circulation of arboviruses in the Nhecolândia sub-region of South Pantanal, Brazil. Sera from a total of 135 horses, of which 75 were immunized with bivalent vaccine composed of inactive Eastern equine encephalitis virus (EEEV) and Western equine encephalitis virus (WEEV) and 60 were unvaccinated, were submitted to thorough viral isolation, reverse transcriptase polymerase chain reaction (RT-PCR) and neutralization tests for Saint Louis encephalitis virus (SLEV), EEEV, WEEV and Mayaro virus (MAYV). No virus was isolated and viral nucleic-acid detection by RT-PCR was also negative. Nevertheless, the prevalence of neutralizing antibodies in horses older than seven months was 43.7% for SLEV in equines regardless of vaccine status, and 36.4% for WEEV and 47.7% for EEEV in unvaccinated horses. There was no evidence of MAYV infections. The serologic evidence of circulation of arboviruses responsible for equine and human encephalitis, without recent official reports of clinical infections in the area, suggests that the Nhecolândia sub-region in South Pantanal is an important area for detection of silent activity of arboviruses in Brazil.*

Key words: arbovirus - Pantanal - RT-PCR - neutralization - equines - Brazil

The arboviruses are maintained in nature due to the biological transmission of an infected vertebrate host to another through hematophagous arthropods and, with few exceptions, are etiological agents of zoonoses that depend on animal species other than humans for maintenance in nature (WHO 1985). Humans and horses generally are terminal or incidental hosts that usually do not contribute to the transmission cycle by developing viraemia and infecting arthropods (Gubler 2002).

Wetland ecosystems like the Brazilian Pantanal, which have high movement of migratory birds (Nunes & Tomas 2004), diversity of arbovirus vectors (Pauvolid-Corrêa et al. 2010) and arbovirus circulation (Iversson et al. 1993), present favorable conditions for reservoirs, which can sustain arbovirus circulation in imperceptible enzootic cycles (Pauvolid-Corrêa & Varela 2008).

Encompassing approximately 140.000 km<sup>2</sup> across the territories of Brazil, Bolivia and Paraguay, Pantanal is one of the world's largest freshwater wetland ecosystem and the most significant South American waterbird breeding area (Lopes et al. 2006). In Brazil, where 85% of the total area is located, 65.5% is situated in the state of Mato Grosso do Sul (MS), known as South Pantanal (Alho et al. 1988, Moraes et al. 2001). The region is ecologically classified into sub-regions that differ in degree of vegetation, flooding and physiognomy (Adámoli 1987). The Nhecolândia sub-region located in South Pantanal occupies 19.48% of the total area and is one of the largest of the floodplain (Silva & Abdon 1998). The main economic activity of the region is beef cattle breeding characterized by vast areas of native grassland and low human population density with little impact on the Pantanal environment (Junk & Cunha 2005).

In the 1990s, a serologic survey for antibodies to arboviruses in equine sera was conducted in South Pantanal and detected the circulation of some arboviruses, including equine encephalitis viruses (Iversson et al. 1993). Since then, some properties have conducted a vaccination program to prevent viral equine encephalitis (Santos et al. 2005).

Nevertheless, there is no recent data concerning local arbovirus circulation in the area and the main objective of the present study was to conduct a virologic and serologic survey for antibodies to *Saint Louis encephalitis virus* (SLEV), *Western equine encephalitis virus* (WEEV), *Eastern equine encephalitis virus* (EEEV) and *Mayaro virus* (MAYV) as well as a molecular investigation for the presence of alphaviruses and flaviviruses to update

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our knowledge of arbovirus circulation in equines from the Nhecolândia sub-region of South Pantanal.

The field stations used in this study were two local ranches comprising a total area of 30,000 ha or 300 km<sup>2</sup>. Ranch A (RA), a research station on federal property of approximately 4,500 ha and Ranch B (RB), a privately owned beef cattle ranch within approximately 25,500 ha, are both located in the Nhecolândia sub-region (18°20'–19°40'S and 57°54'–55°00'W) approximately 150 km east of Corumbá, MS.

In February of 2007 during the rainy season, serum samples were obtained from 135 equines, 93 from RA and 42 from RB. From 93 equines sampled at RA, 75 were immunized on May 2006 with a bivalent vaccine composed of inactive EEEV and WEEV as part of an annual vaccination program for equine encephalitis prevention. None of the 42 horses from RB was vaccinated as the vaccination program was not conducted on this property. All 60 unvaccinated horses from the two properties had no records of travel outside South Pantanal. The mean age of RA horses was 5.9 years (range, 20 days–20 years). All horses from RA had a known date of birth, whereas the animals of RB, mostly untamed, had no records available. Thirty-four horses were intact males, 43 were geldings and 58 were females. All animals from RA were thoroughbreds of the Cavalo Pantaneiro breed and the entire flock of RB was crossbred. All 135 samples were submitted to viral isolation, reverse transcriptase polymerase chain reaction (RT-PCR) and neutralization tests (NT).

Aliquots of 200 µL of 1:10 dilution from equine sera were inoculated into Vero cell monolayer tubes (5 x 10<sup>5</sup> cells/mL). Briefly, the tubes were kept at 36°C for 1 h with discrete motion every 15 min to optimize virus adsorption. At the end of this period, 1,800 µL of medium 199 containing 2% of bovine fetal serum was added followed by incubation for seven days. Tubes were daily observed under an inverted microscope for evidence of cytopathic effects (CPE). Three consecutive passages of seven days each, for a total of 21 days, were performed. The absence of CPE after three consecutive inoculations was considered absence of cytopathogenic viruses in a given sample.

All samples were submitted to RNA extraction using a commercial kit (QIAamp<sup>®</sup> Viral RNA Mini Kit QIAGEN) in accordance with the manufacturer instructions. Reverse transcription was conducted on extracted RNA with Superscript II (Invitrogen) system and random primers (Promega).

Aliquots of 10 µL of cDNA served as templates in subsequent Duplex PCR followed by Semi-Nested PCR assay for the alphaviruses *Venezuelan equine encephalitis virus* (VEEV), WEEV and MAYV and for the flaviviruses *Ilheus virus* (ILHV), *Rocio virus*, *Yellow fever virus* and SLEV detection, as previously described (de Morais Bronzoni et al. 2005).

Five microliter aliquots of cDNA of each sample were also subjected to nested PCR for the flaviviruses *Japanese encephalitis virus* (JEV), *Dengue virus* and *West Nile virus* (WNV) genomes detection, as previously described (Pauvolid-Corrêa et al. 2010).

All antigens used for NT were previously submitted to partial nucleotide sequencing of NS5 gene of SLEV

and NS1 gene of alphaviruses. High identity scores were obtained with the following sequences deposited at Genbank: SLEV (access EU099354.1), MAYV (access DQ001069.1), WEEV (access AF214040.1) and EEEV (access X63135.1).

After heat inactivation at 56°C for 30 min, sera were two-fold serially diluted in medium 199 containing 2% bovine fetal serum on 96-well plates. Dilutions were mixed with equal amounts of viral suspensions at 100 CCID<sub>50</sub> of SLEV, MAYV, WEEV and EEEV and incubated for 1 h at 36°C. After incubation, the Vero cell suspension (100 µL of 2.5 x 10<sup>5</sup> cells/mL) was added to each well and the plates were covered with plastic film. The plates were incubated at 36°C and daily observed under an inverted microscope for 5–10 days. End-point titres were determined as the higher serum dilution with total inhibition of CPE in the cell monolayer induced by 100 CCID<sub>50</sub> of the viral suspensions, as previously described (Lennette 1995). NT was performed initially at a dilution of 1:8 for screening and the reactive samples were further tested at higher dilutions (1:8–1:1024) to determine the endpoint titres. Samples were considered positive when a serum dilution of at least 1:8 was able to neutralize totally 100 CCID<sub>50</sub> of the viral suspensions.

No CPE was observed in Vero cells after three consecutive passages and viral nucleic-acid detection using different PCR-based conditions were also negative. The arbovirus infections, with some exceptions as VEEV, are characterized by low viraemia in equines, which are terminal hosts in most arbovirus cycles (Radostits et al. 2002). Virus detection in blood would require venopuncture during acute infection, a condition that may be difficult to detect in extensive breeding systems. For this study, the sera were collected from animals in the absence of clinical signs, which may have influenced and possibly explain the negative results for viral isolation and nucleic-acid detection. From a total of 135 horses, 75 had been vaccinated with EEEV/WEEV bivalent inactivated vaccine. Of these, 34 (45.3%) were seropositive for SLEV, 67 (89.3%) for EEEV and 44 (58.6%) for WEEV (Fig. 1). Out of 60 unvaccinated horses, 24 (40%) were positive for SLEV, 34 (56.6%) for EEEV and 25 (41.7%) for WEEV (Fig. 2).

However, to confirm EEEV and WEEV circulation, we studied unvaccinated horses that were older than seven months, as there would be no possible passive immune response. Of 60 unvaccinated horses, 44 met these criteria, and neutralizing antibodies for EEEV were detected in 21 (47.7%) and for WEEV in 16 (36.4%). Out of 135 horses assayed for SLEV, 119 horses were older than seven months regardless of vaccination; neutralizing anti-SLEV antibodies were detected in 52 (43.7%). The positive titres ranged from 1:8–1:256 for SLEV, from 1:8–≥1:1024 for EEEV and from 1:8–1:128 for WEEV. Four samples of unvaccinated animals show titres to SLEV, EEEV and WEEV and eight samples showed titres to EEEV and WEEV simultaneously. All samples had neutralizing antibody titres < 1:8 to MAYV and were considered negative for this *Alphavirus* species.

The positivity of 36.4% for WEEV differed from the results of similar serologic studies conducted in Brazil.

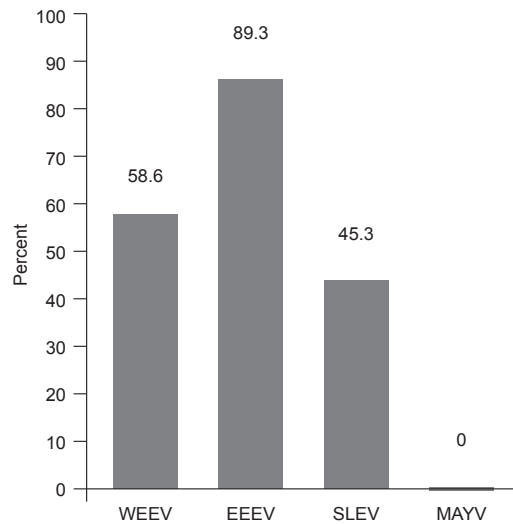


Fig. 1: prevalence by neutralization tests of 75 horses vaccinated with *Eastern equine encephalitis virus* (EEEV)/*Western equine encephalitis virus* (WEEV) inactivated vaccine. MAYV: *Mayaro virus*; SLEV: *Saint Louis encephalitis virus*.

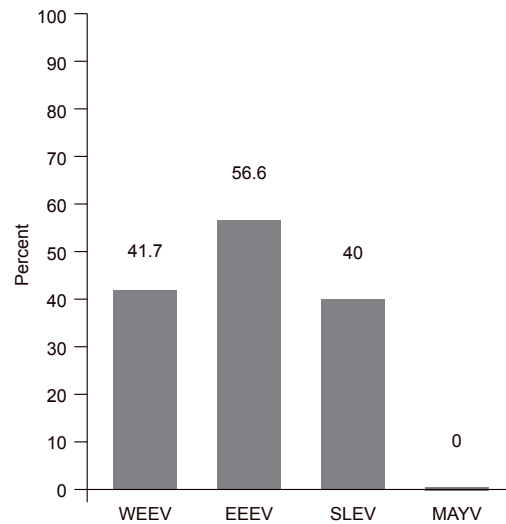


Fig. 2: prevalence by neutralization tests of 60 horses unvaccinated with *Eastern equine encephalitis virus* (EEEV)/*Western equine encephalitis virus* (WEEV) inactivated vaccine. MAYV: *Mayaro virus*; SLEV: *Saint Louis encephalitis virus*.

The first evidence of WEEV circulation in the country was obtained by Bruno-Lobo et al. (1961) in the city of Rio de Janeiro through an equine serologic investigation where 22.2% of the animals evaluated were positive. A serologic survey carried out in horse sera collected at the same region of our studied area showed only 2% positivity to WEEV in the sera assayed by hemagglutination inhibition (HI) and NT (Iversson et al. 1993). The difference in seroprevalence for WEEV among the surveys conducted in the same area may suggest a relevant increase of circulation of this equine encephalitis virus member in the region. Nevertheless, other possible causes such as differences in methodology or interpretation of results must be considered.

The seroprevalence of 47.7% to EEEV is also higher than that observed in most serologic surveys carried out in South America. In the 1970s, positivity to EEEV in Argentina was detected in 11.1% of 892 equines assayed (Monath et al. 1985). In Brazil, Lennette and Fox (1943), during the first investigation of arboviral encephalitis in horses conducted in the country, found that 18.4% of equines assessed were positive for EEEV in the state of Minas Gerais. In the state of Rio de Janeiro, Travassos et al. (1961) detected antibodies to EEEV by HI in 10% of the samples assessed. In the previously cited serologic survey conducted in the same region in the 1990s, seropositivity to EEEV was 6.7% of 432 animals assayed (Iversson et al. 1993). On the other hand, our results were similar to a recent investigation carried out by Fernández et al. (2000) after epizootic encephalitis in the state of Paraná (PR) in which 54.5% of the horses assayed had anti-EEEV antibodies by HI. Unfortunately, we could not clarify the timing of the circulation of equine encephalitis viruses, since the date of birth of nonvaccinated animals was unavailable.

The detection in this study of a high seropositivity to WEEV and EEEV was associated with a previous report of

a clinical infection of EEEV in MS in the 1990s and more recently in PR, a neighboring state. This finding reinforces the need for implementation of local surveillance of equine encephalitis viruses. The circulation of MAYV or even its potential vectors in South Pantanal are unknown. Because no positive results for MAYV antibodies were detected in all equines assessed, the following assumptions must be considered: the absence of MAYV circulation in the area; the biological characteristics of potential local vectors, as the non use of horses as a food source and finally, there are no MAYV vectors in the area.

Despite *Haemagogous* spp, the main known vectors of MAYV (Causey 1962), display an acrodendrophilic characteristic of preferring the canopy in the forest, this species can fly over large areas and descend to bite animals (de Thoisy et al. 2004) and humans mainly in disturbed forests and when they reach high population density (Consoli & Lourenço-de-Oliveira 1994). Because its biological plasticity, the hematofagia on horses by this acrodendrophilic species was considered and MAYV investigation in horses was conducted. However, in a recent preliminary investigation conducted in the same sub-region of our study, of a total of 22 Culicidae species identified, the *Haemagogous* spp were not captured (Pauvolid-Corrêa et al. 2010).

However, many more entomological studies are needed, because these vectors are intensely active during the daytime, a feature little explored in the only Culicidae species study conducted in the area. Despite our negative results, the serologic evidence of MAYV infection in Indians (Neel et al. 1968) and an outbreak registered (Vasconcelos et al. 1998) in the two neighboring states associated with the report of three cases of MAYV fever in men infected in MS (Coimbra et al. 2007) are evidence of MAYV circulation in the Central-West region of Brazil. Moreover, MAYV was detected in migratory birds in the state of Rio Grande do Sul (MS 2003) and six of these avian species

were already observed in South Pantanal (Nunes & Tomas 2004). These data are particularly important because, although vertebrate hosts of MAYV are nonhuman primates, birds that can cover large distances in a short time can act as secondary hosts carrying MAYV in the viraemia phase (Vasconcelos et al. 1998). The absence of seropositivity for MAYV in all samples suggests a lack of cross reaction between the alphaviruses assayed.

Regarding the NT results for SLEV, this is the first serologic evidence of SLEV circulation in South Pantanal. The detection of SLEV antibodies in 43.7% of animals older than seven months was relatively similar to the result described in another serologic survey carried out in Argentina, where the positivity to SLEV was detected by Monath et al. (1985) in 57.9% of equines studied. In Brazil, our results were also similar to a recent report of an equine serosurvey conducted in 2005 in the Amazon region and Maracajú, a municipality of MS outside South Pantanal. In this inquiry, neutralizing antibodies to a chimeric SLEV were detected in 50.9% of 1,401 animals assessed (Rodrigues et al. 2010). Despite the similarity of these results, it is important to consider the different criteria used for serologic diagnosis. Considering the vast co-circulation of flaviviruses in Brazil and consequently its cross-reactivity, our positive samples that neutralized 100% of 100 CCID<sub>50</sub> of sylvatic SLEV aimed to restrict but not completely prevent cross-reactions.

Although NT is the most specific serologic assay for *Flavivirus* diagnosis, the previous report of ILHV circulation (Iversson et al. 1993) and the peculiar ecologic conditions of the South Pantanal that are propitious to arbovirus circulation suggest that another *Flavivirus* species must be considered in future serosurveys.

Neutralizing antibodies to SLEV in animals aged 3-21 years without history of travel outside of Pantanal suggests that SLEV circulated between 2004-2007. This data could be particularly interesting as the human outbreak of SLEV occurred in 2005 in Córdoba province, Argentina (Spinsanti et al. 2008). Recent SLEV circulation has also been reported in Brazil in birds, human and nonhuman primates on different regions of the country (Rocco et al. 2005, Santos et al. 2006, Mondini et al. 2007, Cruz et al. 2009, Nunes et al. 2009). The seropositivity for SLEV in five animals with history of travel to Poconé, a municipality of Mato Grosso (MT), indicates possible infection outside of MS, which should be taken into account by arbovirus surveillance in MT.

Regarding the 75 equines with vaccination history, Waldridge et al. (2003) studied the immune response of equines to the equine encephalitis virus vaccine and revealed that some animals had lower levels of WEEV antibody and, in some cases, total depletion of the antibodies was observed seven months after vaccination. Their result is in agreement with our study where 31 (41.4%) of 75 animals vaccinated in May 2006 did not show neutralizing antibodies to WEEV at the time of venopuncture, which occurred nine months after vaccination.

The high prevalence of EEEV and WEEV was associated with nonofficial reports of horses with clinical signs typical of neurologic diseases and reinforces the need for the maintenance of the viral encephalitis vaccination program, which is already carried out by some properties,

as well as its implementation, mainly for EEEV, by other properties located in the Nhecolândia sub-region.

Despite unapparent SLEV infection in horses and the high specificity of NT, it is important to consider that the close genetic relationship among other species of the JEV group could foil the recognition of previously undetected related species of the group, which occurred in New York City (USA) in the 1990s during the introduction of WNV (CDC 1999).

In Pantanal, the extensive cattle system favors a close contact of the livestock as well as the equines with the local sylvatic fauna and vector populations. Therefore, the horses of Pantanal should be considered as important markers for arbovirus circulation in the area.

In this peculiar cattle livestock, the veterinary assistance is often scarce and most horse deaths are visually attributed by the rural workers to snake bite or equine infectious anemia (EIA), a horse infectious disease well-known by locals. In fact, EIA is highly prevalent in the horses of Pantanal (Silva et al. 2001) and its economic importance may direct the eventual laboratory diagnosis of this *Lentivirus* infection, which could foil the circulation of other important etiological agents, including those previously not detected in Brazil.

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