

The emergent importance of rabies in wild animals in Brazil demonstrates the necessity of continuous epidemiological surveillance in these animal species aiming the development of better strategies for the prevention and control of the disease. The use of blood serum samples from several wild species captured in a native Rainforest area in the North coast of São Paulo State, Brazil, was an excellent opportunity for the research of rabies virus circulation among wildlife in the region, and also to compare different techniques for antibodies detection. In this study we used the “Rapid Fluorescent Focus Inhibition Test – RFFIT, the Simplified Fluorescent Inhibition Microtest – SFIMT and the Enzyme Linked Immunosorbent Assay – ELISA techniques for the detection of rabies virus-specific antibodies in terrestrial wild mammals. Out of 139 samples, 15 (10,8%) presented positive titers for RFFIT (“gold standard” for detection of rabies virus neutralizing antibodies), 50 (35,9%) positive titers for SFIMT and 02 (1,43%) positive titers for ELISA. When comparing RFFIT and SFIMT, 100 (72%) samples presented concordant results when considering positive and negative titers. These results are an evidence of rabies virus circulation between the wild animal species (mainly opossums, capuchin-monkey and coati) in the studied area, even when considering the low concordance between RFFIT and SFIMT. The discordant results between ELISA and RFFIT or ELISA and SFIMT, (99,3%), can be due to the fact that the ELISA kit used was developed for vaccinated foxes, and when considering the Brazilian fauna, which present a great species variety without the use of oral vaccination, the efficacy of the technique could be affected. This result indicates the importance of continuous research regarding a better knowledge of the role presented by wild animals in rabies circulation and transmission in Brazil. Epidemiologic studies in different regions of the Country could provide a valuable information regarding the prevention and control of the disease, and also aiming the standardization and validation of the different diagnostic serologic techniques, especially considering the great and unique variety of animals present in our Country. Acknowledgments: Gaia Consultoria Ambiental, Fundação de Amparo a Pesquisa do Estado de São Paulo (FAPESP) and Coordenação de Aperfeiçoamento de Pessoal de Nível Superior (CAPES)

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ANOMALIES IN THE RABIES INDIRECT FLUORESCENT ANTIBODY TEST CONFOUND ACCURATE ANTEMORTEM DIAGNOSIS OF HUMAN RABIES

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The antemortem diagnosis of rabies in humans employs techniques that require accuracy, speed and sensitivity. A combination of histochemical, *in vitro* virus isolation, immunologic and molecular amplification procedures are utilized in an effort to diagnose the disease. Present day technology offers a potentially life-saving treatment for a disease that was considered invariably fatal once clinical signs develop. This new development adds to the need for a rapid diagnosis as early in the course of clinical signs as possible. The techniques offering diagnosis within hours are the direct fluorescent antibody test on skin and the indirect fluorescent antibody procedure on cerebral spinal fluid and serum. We describe examination by indirect fluorescent antibody assay of cerebral spinal fluid and serum taken from patients with viral encephalitis or a presumed viral infection from an agent other than rabies virus. A total of 135 CSF samples from viral encephalitis patients were tested by the rabies indirect fluorescent antibody procedure. A majority of the spinal fluids tested, from patients with encephalitis, presented immunoglobulins that bound to antigens

present in cell culture substrate. Most notable were the reactions on kidney cells provided from sera or spinal fluid obtained from patients diagnosed with the flavivirus infections Powassan Virus or West Nile Virus. The majority of reaction patterns were recognizably different than what is seen with specific anti-rabies antibodies. However, results indicate that false positive results could occur when interpreting the rabies indirect fluorescent antibody procedure. A staining pattern appearing similar to specific anti-rabies staining was observed in 7 of the 135 spinal fluids examined. The potential for false positive results documented in this work offers weight to the argument that tandem positive results from two diagnostic test platforms are essential when diagnosing rabies in the human patient.

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UNIQUE RABIES VIRUS VARIANT AND GENETIC LINEAGE IN INSECTIVOROUS BATS *Histiotus velatus*, BRAZIL.

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Bats represent approximately one-third of the Brazilian mammal fauna and the Rabies virus has been isolated from 41 of the 167 species of bats present in the country. A *Rabies virus* independent species-specific variant was detected in 16 insectivorous bats *Histiotus velatus* in the Southeast of Brazil from 1995 to 2009. The antigenic characterization was made by monoclonal antibodies panel from Centers for Disease Control and Prevention (CDC – Atlanta, USA) and the genetic characterization was performed by sequencing of carboxi-terminal portion of nucleoprotein followed by Maximum Likelihood (ML) genetic analysis with GARLi software. The antigenic characterization made in 12 of these samples showed a unique profile previously described only for the insectivorous bats *Histiotus velatus* (positive reactivity only with MAB C12 from the utilized panel). The entire 16 samples positive to rabies virus were genetically characterized and they were segregated in the independent monophyletic cluster supported by high bootstrap values. These sequences showed a minimal average intrinsic distance whitening group (1,3%) but they presented low similarity when compared to other lineages circulating in bats and other wild animal lineages from Brazil and worldwide with a range of 8.8% to 25.4%. The antigenic site of the nucleoprotein at residue 377 to 379 (based on PV strain) analysis showed a residue TEV (Thr-Glu-Val) like a some insectivorous bats and different to vampire bats lineage, marmosets lineage and terrestrial cycle related samples. The PV strain shows the amino acids residues TDV (Thr-Asp-Val), *D. rotundus* isolates show AET (Ala-Glu-Thr) and Marmoset lineage show the amino acids residues TEA (Thr-Glu-Ala). This antigenic variant and genetic lineage has been identified in a large area covering various kilometers and different biomes for at least 14 years between the states of Minas Gerais and Sao Paulo exclusively in this bat species. Surprisingly the system documentation not describe this antigenic variant and genetic lineage found before in other bat species and the *Histiotus velatus* bat species never ever found before with a different antigenic variant and genetic lineage. The fact of this lineage has been isolated only in this species besides long temporal space and geographically distal to each other, associated with phylogenetic results and previously antigenic data suggest strongly that this rabies virus lineage is associated to *Histiotus velatus*.