Influence of canine brain decomposition on laboratory diagnosis of rabies

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Abstract  Canine brains infected with rabies virus were submitted to decomposition by being left at room temperature of 25 to 29°C for up to 168h. At 24h intervals, brain fragments were analyzed by immunofluorescence (IF) and by the mouse intracerebral inoculation (MI) test to confirm the diagnosis of rabies and to measure the putrefaction effect on the accuracy of the diagnosis. Forty eight h after the beginning of the experiment, the MI test showed signs of impairment with four negative results, while after 72h, 100% of the results were negative to the MI test and only one result was negative to the IF test, indicating that the threshold period for accurate diagnosis is 24 to 48h before putrefaction. The authors recommend the shipment of suspected cases of rabies to the laboratory for confirmation, but the use of putrid materials for diagnosis is meaningless because of false-negative results.


Resumo  Cérebros de cães infectados com o vírus da raiva foram submetidos à decomposição, deixando-os à temperatura ambiente de 25 a 29°C por até 168 horas. A cada 24 horas, fragmentos de cérebros foram analisados pela imunofluorescência (IF) e inoculação intracerebral em camundongos (IC) para confirmar o diagnóstico de raiva e medir o efeito da putrefação na acurácia do teste. Após 48 horas do início do experimento o teste de IC começou a ser prejudicado, detectando-se quatro resultados negativos, enquanto que, após 72 horas, 100% dos resultados foram negativos para IC e apenas um foi negativo para IF, indicando que o período limite para um diagnóstico seguro está entre 24 e 48 horas antes da putrefação. Os autores recomendam o envio de material suspeito para o diagnóstico laboratorial, no entanto, o uso de materiais em adiantado estado de decomposição não é adequado, devido à ocorrência de resultados falsos-negativos.


According to information of Boletín de Vigilancia Epidemiologica de la Rabia en las Americas⁴, Brazil is indicated with the highest cases of human rabies in Latin American countries; although reports of bats provoking human rabies have been increasing nowadays, but dogs are still the main reservoirs. Isolations of rabies virus from non-hematophagous bats...
have been increasingly common in several regions of the world\textsuperscript{2,7,9}, probably these viruses belong to genotype 5 or 6 of EBL\textsubscript{1} or EBL\textsubscript{2}\textsuperscript{2}.

The livestock industry in these regions is severely affected by rabies, causing heavy economic losses due to high mortality rates especially in bovines\textsuperscript{1,8}.

The laboratory diagnosis of rabies is essential for the guiding of control programs, and for the establishment of epidemiological surveillance and for the effective monitoring of one geographical area, as well as for orientation of prophylactic measures. Among the methods recommended by WHO experts on rabies\textsuperscript{14}, the immunofluorescent (IF) technique accompanied by mouse intracerebral inoculation (MI) test are the best choices for this purpose, because results could be used whether to continue or not the rabies treatment, with or without indication of rabies anti-serum, as specified in the Manual Atualizado de Normas Técnicas de Profilaxia da Raiva em Humanos\textsuperscript{6}.

The results of laboratory diagnosis must be accurate and safe, one factor that impairs rabies diagnosis could be the state of conservation of brain tissues. Fluorescent antigen may be seen in putrid rabies specimens, but one cannot predict at which time the diagnostic method begins to fail\textsuperscript{13}. The use of decomposed specimens could imply in false positive or false negative results. In many rabies diagnostic centers the reception of decomposed brain specimens are still very frequent (FH Ito: personal communication, 1997), and due to important questions concerning accurate confirmation of rabies, the present paper has been designed to experimentally assesses the influence of the state of conservation of brain specimens submitted to rabies diagnosis and to determine the time the diagnostic method of IF and MI tests begin to fail.

MATERIAL AND METHODS

Brains. Among the brain materials shipped for routine laboratory diagnosis of rabies, ten positive canine brains were selected for this study. The direct IF technique was according to Dean et al\textsuperscript{3}, using for each specimen, two microscopic glass slides for making smears by the impression method and for already decomposed brains, thin film smears were prepared.

Anti-nucleocapside rabies polyclonal conjugate used was from SANOFI Diagnostics, Pasteur/France, and the microscope used was the Carl Zeiss, Jannalumar model.

The MI technique was according to Koprowski\textsuperscript{5}, using a 20% brain suspension by weight, and the diluent was prepared by using sterile distilled water added to 2% normal horse serum, with 500IU of penicillin and 500µg of streptomycin per ml and centrifuged at 1500rpm for ten minutes.

After 48h of storage at room temperature of 25-29°C, 10% brain suspensions were obtained from tissues already in decomposition and the suspensions were kept in ice bath soon after centrifugation. Each brain suspension was inoculated in each group of 8 mice weighing 11 to 14g, through intracerebral route, and observed for 21 days and searched for rabies signs and symptoms, and mice found in agonizing state or dead were again submitted to IF test for rabies confirmation.

Procedures. The brains selected had been previously confirmed as rabid, by means of IF and MI techniques and stored frozen until the beginning of the experiment. The criteria for determination of IF positivity were based on fluorescence, and brightness of inclusion bodies and presence of sandlike particles and registered as + to +++++. The + degree was registered for a positive material showing very few inclusion bodies with weak brightness and absence of sandlike particles. The +++++ was used for materials showing numerous round and oval shaped inclusion bodies with typical brightness. The other degrees were used to indicate intermediate results. For evaluation, each brain was simultaneously defrosted and kept in Petri dish at room temperature varying from 25 to 29°C for up to 168h in order to provoke tissue decomposition. Searching for the presence of rabies antigens and the isolation of virus were made by means of MI and IF tests, at each 24h interval.

RESULTS

The results of IF test are summarized in Table 1; the fluorescent brightness has decreased as the time of exposure at 25 to 29°C has evolved. After 72h of exposition, 90% of materials
have been found positive, however after 96h from the beginning of the experiment two samples were found negative, and after 120h, only two out of ten materials examined were positive, and after 144 h, only one specimen was positive. After 7 days (168h), none of the materials were found to have any fluorescent particles.

In relation to MI test (Table 1), mice inoculated with suspension made of brain left at room temperature for 24h showed symptoms and signs of rabies in 100% of mice examined. After 48h of temperature treatment, six specimens still were found positive through MI test. However, 100% of inoculated mice were negative with materials stored for 72h, this result occurred in the following intervals until 168h.

**DISCUSSION**

The results found in this experiment corroborate to that reported by Winkler and Adams14 who stated that fluorescent antigen may still be seen in putrid rabies specimens after mouse inoculation, but one cannot predict the time the diagnostic method begins to fail. Valentini et al10 and Vasconcellos et al11, using mice brains, reported that the fluorescence of rabies antigens could still be identified in materials showing advanced stage of putrefaction. The findings of the specific fluorescence in these materials varied according to the state of decomposition. The impairment of MI results could be felt after 48h from the beginning of the experiment. Between 48 to 72h, the degree of putrefaction has increased, the brains became pasty, and for IF test, after this period, glasses were prepared through smears. In fact, the IF results, after 72h showed only one negative brain while by means of MI test, all materials examined were negative, increasing the discrepancy between the tests. After this period, both IF and MI test presented false negative results, being no more accurate.

Wachendorfer12, Valentini et al10 and Vasconcellos et al11 also found higher frequency of disagreeing results between MI and IF test in materials previously known to be rabid. In this respect, Valentini et al10 searching the presence of rabies antigens through IF test in mice inoculated with CVS strain of rabies virus and having been submitted to carcass decomposition at a temperature of 25°C, for a period of 18h, still detected high positivity. In mice carcasses the putrefaction could be accelerated by post-mortem migration of intestinal microorganisms to the whole carcass. In this experiment, however, we have used canine brains, greater than to that of mice and the materials had been stored separately from the carcasses, thus preventing the post-mortem migration and proliferation of intestinal microorganisms.

The putrefaction of brain tissues could alter the rabies diagnosis, increasing the discrepancies between the IF and MI results. The false negative results were first observed after 48h through MI test. The authors encourage the shipment of specimens for diagnosing rabies, but the use of decomposed carcasses, especially when death has elapsed more than 48h and in the absence of adequate means of conservation, the negative results are meaningless, particularly in cases of
disintering buried carcasses of suspected animals.

REFERENCES


