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RAPID TITRATION OF RABIES VIRUS INFECTIVITY BY BIOTIN-AVIDIN-PEROXIDASE TECHNIQUE AND ITS APPLICATION TO VIRUS NEUTRALIZATION TEST

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Abstract: The rapid titration method of the infectivity and the neutralizing antibody of rabies virus was established by the use of immuno-peroxidase (biotin-avidin-peroxidase) staining technique in microslide culture chambers. This method offers high sensitivity and reproducibility and would provide a new mean for the rapid diagnosis of rabies and the seroepidemiology of rabies virus.

INTRODUCTION

Many studies have been reported on the quantitation of infectious rabies virus in vitro (Yoshino et al., 1966; Sedwick and Wiktor, 1967; Matsumoto and Kawai, 1969; Schneider, 1973; Buckley and Tignor, 1975; Strating et al., 1975). However, the plaque titration method can be applied, in general, for only laboratory-passaged virus strains and requires a longer incubation period to obtain the results. In contrast, in fluorescent antibody (FA) technique, rapid and reproducible titration can be attained using the culture cells sensitive to rabies virus and even street strains can be quantitated by counting fluorescent foci on the cells (Smith et al., 1977).

Recent progress of immuno-peroxidase techniques provides a new mean to detect viral antigens in the infected cells (Benjamin, 1974; Miller et al., 1974; Gerna et al., 1976) and to quantitate the infectivity of virus (Hahon et al., 1975; Okuno et al., 1977, 1979). This technique has several advantages when compared with FA technique. The samples can be examined with a light microscope and stored for long time after fixation according to the stability of the staining. Thus if the technique is applicable for virus neutralization test, this should be the most suitable method for determining the neutralizing antibodies against rabies virus of large number of specimens.

Among the immuno-peroxidase techniques, the use of biotin-avidin interaction system has been proved to be highly sensitive for the detection of cell-associated antigens and has been extensively employed for immunohistochemistry (Guesdon et al., 1979; Warnke et al., 1980; Hsu et al., 1982).

Therefore, the present study was begun with an application of biotin-avidin-peroxidase (BAP) technique for the infectivity assay of rabies virus and virus neutralization test with an ultimate goal of an application for the immuno-electron-microscopic studies of the virus.

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**Materials and Methods**

**Virus:** The plaque purified CVS strain (Buckley and Tignor, 1975) of rabies virus was grown in murine neuroblastoma cells (N-18 clone) and stored at -75°C until use.

**Cells:** Murine neuroblastoma cells and CER cells (Smith *et al.*, 1977) were grown in Eagle minimum essential medium (MEM) supplemented with 5% fetal calf serum and 5% calf serum and antibiotics. CER cells were used for the infectivity assay of rabies virus throughout the study. Microslide culture chamber (10 chambers, Belco Glass, Inc. NJ, U. S. A.) (Fig. 1) received 0.25 ml/chamber of cell suspensions at a cell density of 8 X 10⁴ cells/ml of medium and the cells were incubated at 37°C in 5% CO₂ incubator until the monolayers of the cell completed.

![Figure 1 Microslide culture chamber (10 chambers per slide). The culture chamber can be used repeatedly only by changing a bottom slide glass.](image)

**Staining of the cells by BAP technique:** CER cell monolayers on microslide culture chambers were inoculated with 50 μl/chamber of serially diluted virus suspensions and incubated for 90 min at 37°C for virus adsorption. Then the inoculum was removed and the cells were incubated for various periods with Eagle MEM supplemented with 5% fetal calf serum and 2% Sephadex G-200. At intervals, the cells were washed with phosphate buffered saline (pH 7.4, PBS) and fixed with cold acetone for 20 min. The cells were first reacted with anti-rabies virus mouse ascitic fluid for 60 min at 37°C. After washing with PBS 3 times, the cells were then stained with biotinyl-anti-mouse immunoglobulin (Ig) G goat serum (E. Y. Laboratories, Inc. CA, U. S. A.) at an appropriate dilution for 60 min at 37°C and finally stained with avidin conjugated with peroxidase (E. Y. Laboratories, Inc. CA, U. S. A.) for 60 min at 37°C. Peroxidase reaction was done with H₂O₂ and 3, 3′-diamino benzidine tetrahydrochloride (Sigma Chemicals, MO, U. S. A.) as described by Graham and Karnovsky (1966). The slide was sealed with buffered glycerin and the stained foci were counted under an ordinary light microscope.

**Kinetics of virus neutralization:** Anti-rabies virus mouse ascitic fluid was pretreated...
with 25% kaolin suspension as described by Smith et al., (1973). Virus suspension containing $10^6$ focus-forming unit (FFU)/0.1 ml was mixed with an equal volume of serially diluted anti-rabies virus mouse ascitic fluid and incubated at 36 °C in water bath. At intervals of incubation, 0.2 ml of the mixture were taken and diluted immediately one hundred fold and kept in ice. The samples were then inoculated onto the cell monolayers in microslide culture chambers and assayed for the infectivity of surviving virus. Control virus suspension with virus diluent only was treated similarly.

**RESULTS**

**Formation of foci:** Development of infected foci on CER cells by rabies virus was examined. Cell monolayers were infected with approximately 200 FFU of the virus and incubated. At intervals, samples of the slides were taken and processed for BAP staining and the number of foci (Fig. 2) was counted. As shown in Fig. 3, the foci became countable at 16 hr after infection. The number of foci increased thereafter until 32 hr after infection and reached a maximum although the size of focus increased even thereafter and became uncountable at 40 hr because of the combinations with adjacent foci. From this observation, the number of foci was counted after 32 hr of incubation in the following experiments.

Relationship between the number of foci and virus dilution was next examined. Aliquots of serial twofold dilutions of appropriately diluted virus were inoculated on the cells and the number of foci was counted at 32 hr after infection. Fig. 4 shows a linear relationship between the logarithms of the virus dilution and the number of foci per chamber, indicating that one focus was produced by one infectious particle of virus.
Simultaneous experiments with FA staining technique showed an identical dose response of fluorescent focus formation by rabies virus (Data not shown).

Kinetics of virus neutralization: The optimum incubation period of virus-antibody mixtures for measuring neutralizing antibody was determined at 36 C. Fig. 5 shows the results of the kinetic studies examined with $10^6$ FFU/ml of virus and different dilutions of anti-rabies virus mouse ascitic fluid. Although the virus was neutralized by more than 90% after 15 min of incubation, the neutralization progressed gradually until 90 min of incubation and completed in each dilution of ascitic fluid. Only 28% of the virus was inactivated after 2 hr of incubation.

Thus, the incubation of 90 min at 36 C was selected as the conditions of the following neutralization tests.

Determination of neutralizing antibody titer of anti-rabies virus mouse ascitic fluid: Two lots of mouse ascitic fluid were diluted by twofold dilution starting 1:800. Each dilution was incubated with an equal volume of virus suspension containing 400 FFU of virus /0.1 ml and incubated at 36 C for 90 min. Fifty $\mu$l of the mixtures were then inoculated onto the cells and assayed for surviving virus by BAP technique. The percent of focus inhibition was obtained by dividing the number of foci in test samples by the number of foci in control virus sample which was done in parallel only with the medium.
Figure 5  Kinetics of neutralization of rabies virus. Rabies virus containing $10^6$ FFU/ml was mixed with an equal volume of anti-rabies virus mouse ascitic fluid diluted 1:250 (○-○), 1:500 (□-□), or 1:1000 (▲-▲) and incubated at 36°C. Virus control was incubated with virus diluent only. At intervals of incubation, two hundred microliters of the mixture were taken and the infectivity of surviving virus was assayed.

The percent of focus reduction was transformed into probit and a probit regression line ($Y=a+bX$) was calculated against the logarithms of antibody dilution ($X$). As shown in Fig. 6, probit of the percent of focus reduction fitted to the obtained probit regression lines at various concentrations of antibodies. The slopes of the probit regression lines of two preparations of anti-rabies virus mouse ascitic fluid were same. Thus the fifty percent focus reduction titer of neutralizing antibody of a given serum can be estimated from the probit regression line.

**DISCUSSION**

An immuno-peroxidase (BAP) technique was applied to the rapid titration of rabies virus infectivity and virus neutralization test. The results indicated that the BAP technique can be replaced by FA technique for the titration of rabies virus infectivity. This application to virus neutralization test would provide a new method especially for determining neutralizing antibody titers of a large number of specimens since immuno-peroxidase staining method has several advantages over the FA technique as mentioned previously.
Figure 7 The principle of BAP staining technique and the comparison with PAP (peroxidase-anti-peroxidase) staining technique.

The principle of BAP technique (Fig. 7) is based on extremely high affinity of avidin (affinity constant, $10^{15} \text{ M}^{-1}$) to the biotin which is a vitamin with a molecular weight of 244 and derived from the diet and intestinal bacteria. Ig or proteins can be covalently coupled with many molecules of biotin without loss of biological activities of the proteins. Avidin is a basic glycoprotein of approximately 68000 of molecular weight which is present in eggs. Each avidin molecule possesses four binding sites for biotin while each biotin molecule is capable of binding to only one avidin molecule (Moss et al., 1971). The binding is extremely resistant to dissociation. One molecule of avidin can be conjugated with more than two molecules of horseradish peroxidase. Thus the use of antiviral Ig (first antibody), biotinyl anti-Ig serum (second antibody) and peroxidase conjugated avidin for the staining of viral antigens in the cells should provide higher sensitivity and specificity than the peroxidase-labelled antibody techniques or peroxidase-anti-peroxidase (PAP) staining technique as has been studied and proven by Guesdon et al. (1979). In addition, BAP technique has one more advantage over the PAP technique, i.e. the peroxidase conjugated avidin can be used to biotinyl anti-Ig prepared in any species of animals, while in PAP technique, PAP complex should be prepared each in corresponding animal to which antiviral serum (first antibody) is prepared (Fig. 7).

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ビオチン、アビジン、ペルオキシダーゼ法（免疫酵素抗体法）による狂犬病ウイルス感染質の迅速測定法とそのウイルス中和抗体測定への応用

七條 明久1・三舟 求真1・林 魁君2

マイクロスライドチェンバーを使用し、免疫酵素抗体法（ビオチン、アビジン、ペロキシダーゼ法）による狂犬病ウイルス感染質およびウイルス中和抗体価の迅速測定法を確立した。この方法は迅速であるのみならず、感度、再現性も優れている。従来から使用されている蛍光抗体法に代って狂犬病の迅速診断あるいは血清学的に有効な新しい方法となるものである。

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