Rabies Virus Antibody ELISA

A virus based ELISA, to detect IgG antibodies against Rabies Virus in serum or plasma.

REF  EVL-EIAD1006-AB03
Σ   96
Effective, January 2011

Please use only the valid version of the package insert provided with the kit.
Verwenden Sie nur die jeweils gültige, im Testkit enthaltene, Arbeitsanleitung.
Si prega di usare la versione valida dell’inserto del pacchetto a disposizione con il kit.
Por favor, se usa solo la versione válida de la metodica técnica incluido aquí en el kit.

Table of Contents / Inhaltsverzeichnis / Tabella die Contenuti / Tabla de Contenidos

1 INTRODUCTION.................................................................................................................. 2
2 INTENDED USE OF THE TESTKIT .................................................................................. 2
3 PRINCIPLE OF THE TEST KIT ......................................................................................... 2
4 CONTENTS ........................................................................................................................ 2
5 HANDLING AND STORAGE OF SPECIMENS ................................................................. 2
6 WASH PROTOCOL ............................................................................................................ 3
7 TEST PROTOCOL .............................................................................................................. 3
8 PRECAUTIONS .................................................................................................................. 3
9 VALIDATION OF THE TEST .............................................................................................. 4
10 INTERPRETATION OF TEST RESULTS ............................................................................ 4
SYMBOLS USED WITH EVL ASSAYS ................................................................................. 5
1 INTRODUCTION

Rabies virus can infect all warm-blooded species and in many species the disease can present itself in two different forms. Furious rabies, in which predominantly the brain is infected and paralytic rabies in which predominantly the spinal cord is involved. When cells of the limbic system are infected the first changes in behavior characteristic of rabies may be observed. It has been suggested that the phase before infecting cells of the nervous system may take a considerable length of time, causing a variable incubation period from 10 days to several years. Hence the virus is present in the saliva, which favors the most natural way of transmission by biting in the various stages of the disease, also sporadic cases of aerosol infections have been documented. Carnivores, especially domestic dogs and cats, and also rodent and recently bats, are usually involved in transmission of infections to dogs and man. Infections of dogs with rabies virus seem to be invariably fatal. Persistent in apparent infection accompanied by virus shedding has been documented in several human and animal species including cats and raccoons. This standardized ELISA test system based on whole-inactivated virus is intended to use as a rapid screening test for the detection of rabies antibodies in serum samples of dogs.

2 INTENDED USE OF THE TESTKIT

This diagnostic test system for the establishment of Rabies infection is intended to identify antibodies against epitopes of rabies virus, in serum samples. In contrast to other test systems this standardized ELISA based on whole-inactivated virus, has a very high sensitivity and specificity.

3 PRINCIPLE OF THE TEST KIT

The test is based on the reaction of whole-inactivated virus with polyclonal dog antibodies.

To this end purified inactivated virus has been coated to a 96-well microtiter strip plate. The dog serum sample is added (diluted 1:100) to the wells of the coated plate. The sample also can be titrated using a 3-step dilution, starting with a dilution 1:50 (150; 450; 1350). After washing, the bound dog antibodies are detected by HRPO conjugated anti-species conjugate.

The color reaction in the wells is directly related to the concentration of rabies virus antibodies in the serum sample.

4 CONTENTS

- 12 x 8 microtiter strips.
- 1 x strip holder.
- 1 x 18 ml ELISA buffer.
- 1 x 12 ml HRPO conjugated anti-species antibodies (READY TO USE)
- 1 x 0,5 ml Positive control (Freeze dried).
- 1 x 1 ml Negative control (Freeze dried).
- 1 x 20 ml Wash-solution (200x concentrated), dilute in de-ionized water before use!
- 1 x 8 ml Substrate A.
- 1 x 8 ml Substrate B.
- 1 x 8 ml Stop-solution.
- 1 x Plastic cover seal

5 HANDLING AND STORAGE OF SPECIMENS.

The kit should be stored at +4°C.

An open packet should be used within 10 days.

Samples may be used fresh or may be kept frozen below -20°C before use.

Positive and negative controls may be stored after reconstitution in aliquots at -20°C and used until the expiry date.

Avoid repeated freezing and thawing as this increases non-specific reactivity.
6 WASH PROTOCOL
In ELISAs, un-complexed components must be removed efficiently between each incubation step. This is accomplished by appropriate washing. It should be stressed that each washing step must be carried out with care to guarantee reproducible inter- and intra-assay results. It is essential to follow the washing procedures outlined below. Washing may be done manually or with automatic equipment. Automatic washing equipment usually gives better results.

Manual washing
1. Empty each well by turning the microtiter plate upside down, followed by a firm vertical downward movement to remove the buffer.
2. Fill all the wells with 250 µl washing solution.
3. This washing cycle (1 and 2) should be carried out at least 4 times
4. Turn the plate upside down and empty the wells with a firm vertical movement
5. Place the inverted plate on absorbent paper towels and tap the plate firmly to remove any residual washing solution in the wells.
6. Take care that none of the wells dry out before the next reagent is dispensed

Washing with automatic equipment
When automatic plate washing equipment is used, check that all wells are aspirated completely and that the washing solution is correctly dispensed, reaching the rim of each well during each rinsing cycle. The washer should be programmed to execute at least 4 washing cycles.

7 TEST PROTOCOL
1. Open the packet of strips and take out the strips to be used. Cover the remaining strips with a part of the provided seal and store them at +4°C. and use them within 10 days. Wash the microtiter strip(s) with washing solution, according to washing protocol.

   **The washing solution provided must be diluted 200x in de-ionized water!**

   Reconstitute directly before use the positive control in 0,5 ml and the negative control in 1 ml de-ionized water.

2. Qualitative:
   Dilute the serum or plasma 1:100 in ELISA buffer. Make also a 1:100 dilution of the positive and negative control.

   **Quantitative:**
   Make 3-step dilutions of each sample in ELISA buffer, starting 1:50 (150; 450; 1350) in a round-bottomed microtiter plate. Make also a 3-step dilution of the positive and negative control.

3. Transfer 100 µl of these dilutions to the (virus-coated) microtiter strips.
4. Incubate 60 min. at 37°C.
5. Wash as pointed out in wash protocol.
6. Dispense 100 µl HRPO conjugated anti-species antibodies to all wells.
7. Seal and incubate 60 min. at 37°C.
8. Wash as pointed out in wash protocol.
9. Mix equal parts of buffer A and buffer B with gentle shaking. Prepare immediately before use! Dispense 100 µl substrate solution to each well. Incubate 10 min. at room temperature (21°C).
10. Add 50 µl stop solution to each well; mix well.
11. Read the absorbency values immediately (within 10 min.) at 450 NM.

8 PRECAUTIONS
− Handle all biological material as though capable of transmitting Rabies.
− Do not pipette by mouth.
− Do not eat, drink, smoke or prepare foods, or apply cosmetics within the designated working area.
− TMB substrate (buffer A/B) is toxic by inhalation, through contact with skin or when swallowed; observe care when handling the substrate.
− Do not use components past the expiry date and do not mix components from different serial lots.
− Optimal results will be obtained by strict adherence to this protocol. Careful pipetting and washing throughout this procedure are necessary to maintain precision and accuracy.
− Each well is ultimately used as an optical cuvette. Therefore, do not touch the under-surface of the microtiter plate and protect it from damage and dirt.
9 VALIDATION OF THE TEST
In order to confirm appropriate test conditions the OD of the positive control should be ≥ 1.000 OD units (450nm). The negative control should be lower than 0.350 OD units (450nm) and give an endpoint titer of ≤ 50.

10 INTERPRETATION OF TEST RESULTS

This test can be used in 2 ways.

1. **Qualitative:** positive – negative
   A sample is scored positive if the OD is higher than the OD of the negative control plus 0.200.

2. **Quantitative:** end point titre
   The titre in IU according RIFFIT can be calculated by constructing a curve from the positive control (dilution 1:50- 150-450 -1350-4050 -12150 etc total 8 dilutions of 3 steps) OD on Y-as and IU on X-as
   The ELISA titre of the positive control is 122,3 IU. The IU titre obtained in this will be close to the RIFFIT titre but final correlation depends on the Lab performing the RIFFIT test. Small Lab to Lab variation in RIFFIT titre will always been seen due to the nature of biological material (cells and virus).
### SYMBOLS USED WITH EVL ASSAYS

<table>
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<tr>
<th>Symbol</th>
<th>English</th>
<th>Deutsch</th>
<th>Français</th>
<th>Español</th>
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<td>📖</td>
<td>Consult instructions for use</td>
<td>Gebrauchsanweisung beachten</td>
<td>Consulter les instructions d'utilisation</td>
<td>Consulte las instrucciones de uso</td>
<td>Consultare le istruzioni per l’uso</td>
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<td>European Conformity</td>
<td>CE-Kennzeichnung</td>
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<td>🇺🇸</td>
<td>In vitro diagnostic device</td>
<td>In-vitro-Diagnostikum</td>
<td>Usage Diagnostic in vitro</td>
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<td>For research use only</td>
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