

PLATELIA™ RABIES II KIT

Ad Usum Veterinarium

192 TESTS

Ref.: 355-0180

KIT FOR *IN VITRO* DETECTION AND TITRATION OF IgG ANTI RABIES VIRUS GLYCOPROTEIN IN DOGS, CATS AND FOXES SERUM.

All manufactured and commercialized reagents are under complete quality system starting from reception of raw materials to the final commercialization of the product.

Each lot is submitted to a quality control and released on the market when conforming to the acceptance criteria.



Fitness for purpose validated and certified by OIE
Registration number: 20070101

BIO-RAD

User's manual

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1 - INTENDED USE

The PLATELIA™ RABIES II kit is an *in vitro* diagnostic ELISA test allowing detection and titration of IgG anti rabies virus glycoprotein in animal (dogs, cats and foxes) serum.

2 - INTEREST OF THE PLATELIA™ RABIES II KIT

Rabies is one of the oldest viral diseases afflicting humans and animals. Rabies virus is a highly neurotropic virus, usually causing fatal encephalitis in mammals. The virus is mostly transmitted through the saliva via bites from rabid animals. Immunization and control programs for the protection of animal populations from rabies virus have resulted in the elimination of the disease in several Western-European countries and in some countries in South America.

As of July 3rd 2004, a new European regulation applies to international movements of domestic carnivores from rabies-infected to rabies-free countries. Domestic carnivores are allowed to travel if their serum contain at least 0.5 IU/ml (International Units/ml) of rabies neutralising antibodies.

Anti-rabies antibodies titration has several practical applications :

- Individual serology for international trade purposes: anti-rabies antibodies are measured in specialized laboratories in order to determine the degree of immunity of vaccinated animals (cats and dogs). A level of antibody equal to or superior to 0.5 IU/ml is considered by OIE experts as an acceptable seroconversion level above which the vaccination could be considered as successful.
- Confirmation of vaccinated status during a vaccination campaign: control of the anti-rabies antibodies enables indirect assessment of the effectiveness of oral vaccination campaign in wildlife (foxes).

3 - PRINCIPLE OF THE PLATELIA™ RABIES II ASSAY

PLATELIA™ RABIES II is an immuno-enzymatic technique for the detection of rabies virus anti-glycoprotein antibodies. This assay can be carried out on the serum of several species of animals - dog, cat and fox.

The test is based on the use of a solid phase enzyme immunoassay technique referred to as an indirect ELISA. A microplate is coated with rabies glycoprotein extracted from the inactivated and purified virus membrane. The enzymatic conjugate consists of a protein A from *Staphylococcus aureus* coupled with peroxidase. Positive controls, calibrated against OIE standard, allow the qualitative or quantitative determination of anti-rabies antibody titre in the serum.

Implementation of the test comprises the following reaction steps:

- 1 - The unknown sera as well as the calibrated Positive Controls or the Quantification Standards are distributed in the glycoprotein coated wells of the microplates. During incubation of one hour at 37°C, anti-rabies antibodies present in the sample bind to the glycoprotein coated to the microplate wells. After incubation, unbound antibodies and other serum proteins are removed by washings.
- 2 - The conjugate (protein A labelled with peroxidase) is added to the microplate wells. During a second incubation of one hour at 37°C, the labelled protein A binds to the anti-rabies-antibody-antigen complexes attached to the microplate wells. The unbound conjugate is removed by washings.

- 3 - The presence of immune complex is demonstrated by the addition of a solution containing a peroxidase substrate and a chromogen, initiating a colour development reaction.
- 4 - After 30 min. incubation at room temperature, the enzymatic reaction is stopped by addition of a solution H₂SO₄ 1N. The optical density reading obtained with a spectrophotometer set at 450 - 620 nm is proportional to the amount of anti-rabies antibodies present in the samples. A standard curve is constructed using the Quantification standards (S1 to S6), obtained by serial dilutions of the R4b calibrated Positive Controls.

The optical density values for the unknown samples are compared with the Positive Controls. Sera titres in quantification tests are obtained after a direct reading on the standard curve and are expressed as Equivalent units per ml (EU/ml), unit equivalent to the international units defined by seroneutralization.

4 - COMPOSITION OF THE PLATELIA™ RABIES II ASSAY

All reagents are exclusively for *in vitro* veterinary use.

Enough reagents are provided for 2 x 96 assays. On each microplate, 90 samples can be analysed if a qualitative test is performed. Up to 80 samples can be quantified precisely for rabies antibodies if a quantification system is used.

LABELLING	TYPE OF REAGENT	PRESENTATION
R1	Microplate: 12 strips of 8 wells sensitized with the rabies virus glycoprotein	2 plates
R2	Wash solution: 10-fold concentrated Tris-NaCl buffer Preservative: ProClin™ 300 (0.01%)	1 vial (250 ml)
R3	Negative control: Non reactive control TRIS-EDTA Preservative: ProClin™ 300 (0.1%)	1 vial (0.6 ml)
R4a	0,5 EU/ml Positive control: 0.5 EU/ml calibrated positive control Glycine buffer containing BSA and dog serum with anti-rabies IgG. Yellow coloured. Preservative: ProClin™ 300 (0.1%)	1 vial (0.6 ml)
R4b	4 EU/ ml Positive control: 4 EU/ml calibrated positive control Glycine buffer containing BSA and dog serum with anti-rabies IgG. Blue coloured Preservative: ProClin™ 300 (0.1%)	1 vial (0.6 ml)
R6	Sample diluent: Ready-to use TRIS - EDTA buffer for sample dilution. Red coloured. Preservative: ProClin™ 300 (0.1%)	2 vials (2 x 125 ml)
R7	Conjugate: Solution containing Protein A-Peroxidase and purified bovine protein. Concentrated 10 times. Green coloured. Preservative: ProClin™ 300 (0.1%)	1 vial (3 ml)
R8	Peroxidase substrate buffer: Solution of citric acid and sodium acetate containing 0.015% H ₂ O ₂ and 4% dimethylsulfoxide (DMSO)	1 vial (60 ml)
R9	Chromogen: Tetramethylbenzidine (TMB) solution 0.25%	1 vial (5 ml)
R10	Stop solution: 1 N sulphuric acid solution	1 vial (28 ml)
	Adhesive films for microplates	6

5 - PREPARATION OF REAGENTS

Note:

Before use, allow reagents to reach room temperature (+18 to +30°C) for 30 minutes. Homogenize the reagents by gently mixing, before opening.

1 - Ready-to-use reagents

- **Microplates (R1)**

Prior to use, let the microplate adjust to room temperature (+18°C to +30°C) in its protective packaging to avoid any water condensation in the wells. Any unused strips should be immediately returned to the bag, tightly close the bag after expelling any air, then store at +2°C to +8°C.

- **Sample diluent (R6)**

- **Stop solution (R10)**

2 - Reagents to be reconstituted

- **Wash solution (R2)**

Dilute the solution 1/10 in distilled water (example 100 ml of reagent R2 in 900 ml of distilled water). 500 ml are required for a plate.

- **Negative control (R3)**

Dilute the solution 1/100 in R6.

- **0.5 EU/ml Positive control (R4a)**

Dilute the R4a positive control 1/100 in R6

- **4 EU/ml Positive control (R4b)**

Dilute the R4b positive control 1/100 in R6

- **Conjugate (R7)**

The solution is concentrated 10 times. To prepare diluted conjugate solution, add 1 volume of concentrated conjugate to 9 volumes of prepared 1X wash solution (R2). 11 ml is required for a full microplate.

- **Enzymatic development solution (R8 + R9)**

Dilute reagent R9 to 1/11 in the reagent R8 (example: 0.1 ml of reagent R9 in 1 ml of reagent R8) bearing in mind that 11 ml of enzymatic revelation solution is sufficient for 1 microplate. Homogenize gently. Avoid using a Vortex® agitator.

3 - Preparation of the Quantification Standards for a quantification assay

Each quantification assay using PLATELIA™ RABIES II kit should include 6 Quantification standards – from S1 to S6.

The calibrated R4b Positive control (4 EU/ml) corresponds to the S6 Quantification standard. Serial dilutions out of the R4b reagent allow the preparation of S5 to S1 Quantification standards. The dilutions are performed using the sample diluent (reagent R6).

Quantification standards		Concentrations obtained by serial dilutions of the R4b Positive control
S6	R4b diluted to 1/100	4 EU/ml
S5	S6 diluted to 1/2	2 EU/ml
S4	S5 diluted to 1/2	1 EU/ml
S3	S4 diluted to 1/2	0.5 EU/ml
S2	S3 diluted to 1/2	0.25 EU/ml
S1	S2 diluted to 1/2	0.125 EU/ml

6 - STORAGE - SHELF LIFE

As supplied, all the components of the PLATELIA™ RABIES II kit are stored at +2° to +8°C and can be used until the expiry date indicated on the kit. Reagent R2 (10 X) can be stored at +2°C to +25°C before reconstitution.

The shelf-lives of the reagents after preparation are as follow:

	REAGENT	REMARKS	SHELF-LIFE
R1	Microplate	After opening the sealed foil lined bag, the unused strips should immediately be put back in the bag and the bag again sealed. The desiccant should remain in the foil bag.	1 month at +2°C to +8°C
R2	Diluted wash solution		2 weeks at +2°C to +8°C
R7	Diluted conjugate solution	The diluted conjugate solution should preferably be used immediately	8 hours at +2°C to +8°C
R8 + R9	Reconstituted development solution		6 hours at room temperature (+18°C to +30°C) always protected from light

7 - SAMPLES

- The PLATELIA™ RABIES II test has been developed for application to animal (dogs, cats and foxes) serum.
- The assay is carried out on sera after a 1/100 dilution in R6 reagent.
- The samples are stored at +2° to +8°C if the detection is carried out within 24 hours or can be stored at -20°C for 6 months. The samples can be submitted to 3 cycles of freezing and thawing. Previously frozen samples should be thoroughly mixed after thawing prior to testing.

NB: Eliminate, if necessary by centrifuging, the fibrin particles or aggregates in suspension that can provide false positive results.

8 - ASSAY PROTOCOL

Strictly follow the recommended protocol.

Use the negative and the positive controls for each test run and on each microplate in order to validate the quality of the detection in the qualitative assay. If quantification is performed, the negative control and the Quantification standards should be deposited on each plate. In both cases, follow the recommended microplate set-up.

Use good laboratory practices.

1. Remove the microplate rack and the required number of rows (R1) from the protective packaging. Replace the unused rows with the desiccated bag in the microplate sachet and hermetically close it.
2. Carefully establish the sample distribution and identification plan as follow:

Microplate set-up for Qualitative assay:												
	1	2	3	4	5	6	7	8	9	10	11	12
A	R3	E3										
B	R3	E4										
C	R4a	E5										
D	R4a	E6										
E	R4b	E7										
F	R4b	E8										
G	E1	E9										
H	E2	E10...										

Microplate set-up for Quantitative assay:												
	1	2	3	4	5	6	7	8	9	10	11	12
A	R3	S4	E1	E9								
B	R3	S4	E2	E10...								
C	R4a	S3	E3									
D	R4a	S3	E4									
E	S6	S2	E5									
F	S6	S2	E6									
G	S5	S1	E7									
H	S5	S1	E8									

3. For the qualitative assay, dilute the R3, R4a and R4b controls and the unknown sera to 1/100 in R6 reagent (e.g.: 10 µl of sample in 990 µl of dilution solution).
4. For the quantitative assay, prepare the quantification standards (refer to chapter 5.3) and dilute the R3, R4a controls and the unknown sera to 1/100 in R6 reagent (e.g. 10 µl of sample in 990 µl of dilution solution).
5. Distribute 100 µl of diluted samples, controls and Quantification standards in the correspondent microplate wells according to the pre-established distribution plan.
6. Cover the microplate with adhesive film (cut the sheet if necessary). Press firmly all over the plate to ensure a tight seal.
7. Incubate the strips at 37 ± 2°C for 60 minutes ± 5 minutes.
8. Prepare the wash solution (R2) [refer to chapter 5]
9. Prepare the conjugate solution (R7), as described in chapter 5 before the end of the first incubation.
10. Remove the adhesive film. Perform 3 wash cycles. Optimal washing conditions are obtained with PW40, PW41 or 1575 Bio-Rad plate washers with program TSE 3. Do not let the microplate stand for more than 5 minutes after the last wash cycle. Dry by inversion on absorbent paper before the following step.
11. Distribute 100 µl of the conjugate solution (R7) into each well. Cover with a new adhesive film and incubate for 60 minutes ± 5 minutes at 37 ± 2°C.
12. Prepare the enzymatic development solution (R8+R9) as described in chapter 5, just before use.
13. Remove the adhesive film, perform 5 wash cycles. Optimal washing conditions are obtained with PW40, PW41 or 1575 Bio-Rad plate washers with programs TSE 5. Do not let the microplate stand for more than 5 minutes after the last wash cycle. Dry by inversion on absorbent paper before the following step.
14. Away from direct light, quickly distribute 100 µl of the enzymatic development solution (R8 + R9) into each well and incubate the plate in darkness at room temperature (+18 to +30°C) for 30 minutes ± 5 minutes.
NB: Do not use adhesive film during this incubation.
15. Add 100 µl of the stop solution (R10) to each well according to the same sequence and same distribution rate as for the enzymatic development solution.
16. Thoroughly wipe the bottom of the plate. Read the optical density at 450 - 620 nm (Dual wavelength mode) within 30 minutes after stopping the reaction (the strips must always be protected from light before reading).
17. Before recording the results, check that the reading complies with the distribution and identification plan for the plates and samples.

MICROPLATE WASHER PARAMETERS

NAME: TSE 3

EDIT Mode friction	PLATE	Manifold	STRIPS	Met. [Method]	CROS. VOLUME	ASP. TIME	OVER FLOW	LIQUID FLOW	BOT. WASH NUMBER	BOTTOM TIME	BOT. ASP NUMBER	SHAKE TIME	Nr. OF CYCLES	SOAKING	MET. INTER	Nr. OF KITS	KIT INTER
Main parameter	Flat 01 [PN/40/PW41] Flat 03 [1575]	1*8 [PN/40/1575] 2*8 [PW41]	1,2,3,4, 5,6,7,8,9, 10,11,12	-	-	-	-	-	-	-	-	-	-	-	1	-	
Method 1	-	-	-	WASH	Plate	Yes	0.3	800	2.5	W1	0	-	-	3	30 [PW41] 45 [PN/40/1575]	0	-
Method 2	-	-	-	BOTTOM ASP.	Plate	Yes	0.3	-	-	-	-	1	-	1	0	-	-

NAME: TSE 5

EDIT Mode friction	PLATE	Manifold	STRIPS	Met. [Method]	CROS. VOLUME	ASP. TIME	OVER FLOW	LIQUID FLOW	BOT. WASH NUMBER	BOTTOM TIME	BOT. ASP NUMBER	SHAKE TIME	Nr. OF CYCLES	SOAKING	MET. INTER	Nr. OF KITS	KIT INTER
Main parameter	Flat 01 [PN/40/PW41] Flat 03 [1575]	1*8 [PN/40/1575] 2*8 [PW41]	1,2,3,4, 5,6,7,8,9, 10,11,12	-	-	-	-	-	-	-	-	-	-	-	1	-	
Method 1	-	-	-	WASH	Plate	Yes	0.3	800	2.5	W1	0	-	-	5	30 [PW41] 45 [PN/40/1575]	0	-
Method 2	-	-	-	BOTTOM ASP.	Plate	Yes	0.3	-	-	-	-	1	-	1	0	-	-

PLATE NAME: FLAT 01 (PNW40/PW41) - FLAT 03 (1575)

BOT. SHAPE	ASP. HOR. POS.	CENTERING	ASP. VER. POS.	BOT. VERT. POS.	B.W. VERT. POS.	HORIZONTAL SPEED	VERTICAL SPEED	ASP. DOWNW. SPEED	DISP. UPW. SPEED	BOT. DOWNW. SPEED	BOT. UPWARD SPEED	SHAKING AMPLITUDE	SHAKING SPEED	SHAKING SPEED	
Flat	1.4	0.3	13.5	9.5	9.5	6	8	6	9	6	9	1	9	1	9

9 - CALCULATION AND INTERPRETATION OF THE RESULTS

The results are given in Optical Densities (ODs) after reading of the microplate at 450 - 620 nm.

1. Qualitative determination

For the qualitative determination, include all the controls (R3, R4a and R4b) for each test run.

a) Conditions of validation

Criteria	Validation
OD R3(i) < 0.05	The absorbance of each individual negative control must be lower than 0.05. The test must be repeated if at least one value lies outside of this limit.
0.300 ≤ OD R4a(i) ≤ 1.200	Each individual value of the R4a positive control OD must be between 0.300 and 1.200. The test must be repeated if at least one of the R4a OD value lies outside of this limit.
1.500 ≤ OD R4b(i) ≤ 3.500	Each individual value of the R4b positive control optical densities must be between 1.500 and 3.500. However, a maximum of one individual aberrant value can be eliminated when its optical density is lower than 1.500 or higher than 3.500. The test must be repeated if the two R4b positive control OD lie outside of this limit.

b) Interpretation of the results

The Threshold Value is equal to the mean of the two R4a Positive controls ($\overline{\text{OD R4a}}$) and corresponds to the seroconversion threshold value at 0.5 EU/ml.

High seroconversion Value is equal to the mean of the two R4b Positive control ($\overline{\text{OD R4b}}$), or to the single R4b positive control if one aberrant value has been eliminated.

Optical Density of each sample is compared to this High seroconversion and Threshold Value.

Criteria	Result	Validation
Seroconverted +++	$\text{OD sample} > (\overline{\text{OD R4b}})$	Samples with OD value greater than the High Seroconversion Value originate from individuals who have highly seroconverted after vaccination according to PLATELIA™ RABIES II test.
Seroconverted	$(\overline{\text{OD R4a}}) \leq \text{OD sample} \leq (\overline{\text{OD R4b}})$	Samples with OD value greater than or equal to the Threshold Value and lower than or equal to the High Seroconversion Value originate from individuals who have seroconverted after vaccination according to PLATELIA™ RABIES II test.
Not seroconverted	$\text{OD sample} < (\overline{\text{OD R4a}})$	Samples with OD value lower than the Threshold Value originate from individuals presenting a level of seroconversion insufficient to assess the efficiency of the vaccination according to PLATELIA™ RABIES II test.

2. Quantitative determination

For the quantitative determination, include all the standards and controls (S1 to S6, R3 and R4a) for each test run.

a) Conditions of validation

Criteria	Validation
$\text{OD R3(i)} < 0.05$	The absorbance of each individual negative control must be lower than 0.05. The test must be repeated if at least one value lies outside of this limit.
$0.300 \leq \text{OD R4a(i)} \leq 1.200$	Each individual value of the R4a positive control optical densities must be between 0.300 and 1.200 . The test must be repeated if at least one of the R4a OD value lies outside of this limit.
$\overline{S1} < \overline{S2} < \overline{S3} < \overline{S4} < \overline{S5} < \overline{S6}$	Calculate the mean OD for S1 to S6 in the following way: $\overline{S1}$ = mean of the two S1 ODs (corresponds to 0.125 EU/ml). $\overline{S2}$ = mean of the two S2 ODs (corresponds to 0.25 EU/ml) etc. The signals of the standards must increase in the following way: $\overline{S1} < \overline{S2} < \overline{S3} < \overline{S4} < \overline{S5} < \overline{S6}$.
$0.7 \leq \overline{S3/R4a} \text{ ODs} \leq 1.3$	The ratio between $\overline{S3}$ (mean S3 standard) and $\overline{R4a}$ (mean R4a) must be between 0.7 and 1.3.

b) Interpretation of the results

The Threshold Value is equal to the mean of the two OD of the S3 Quantification standard ($\overline{S3}$). The S3 Quantification standard corresponds to the seroconversion threshold value 0.5 EU/ml.

The quantity of anti-rabies antibodies in a sample is determined by comparing the optical density of the sample to a standard curve.

Sera titres are expressed as Equivalent units per ml (EU/ml), unit equivalent to the international units defined by seroneutralization.

Drawing the standard curve:

For manual data reduction, use graph paper and plot the mean values of the OD readings of the Quantification standards (S1 to S6) on the vertical (y) axes. Plot the corresponding concentrations in EU/ml on the horizontal (x) axes. Draw a series of line segments which goes through the 6 points.

For automated data reduction, a "point-to-point" function is used to construct a curve from the OD readings obtained for the standards.

A quantitative result for the anti-rabies antibodies titre for an unknown sample can be delivered if the OD value for the sample ranges between the S1 (0.125 EU/ml) and the S6 (4 EU/ml) Optical Density mean values. In this case, the unknown sample titre is found using the Standard curve. On the Standard curve, find the value corresponding to the OD reading of the sample on the y-axis and draw a horizontal line to the standard curve. At the point of the intersection with the standard curve, draw a vertical line to the x-axis. Read the corresponding concentration in EU/ml.

If the OD value of the unknown sample is higher than the mean Optical Density value for the S6 standard, a precise quantification cannot be performed. If a precise quantification is required, proceed to 1/10 dilution or more of the sample and perform the assay again in order to have an Optical Density in the interval of the Standard curve.

Note: An interpretation and conversion tool of OD values into titres is available on the CD (Rabies QT-ELISA BIORAD-Vers.200712.A.XLS) for users who are not equipped with Bio-Rad readers.

General comprehension of the results:

Results in Optical Density for the unknown sample	Titre for the unknown sample (X)	Interpretation of the result
OD sample > $\overline{S6}$	X > 4 EU/ml	High seroconversion level. If a precise titre is required, the sample must be diluted before repeating the assay.
$\overline{S3} \leq$ OD sample $\leq \overline{S6}$	X in EU/ml (0.5 - 4 EU/ml)	Sufficient seroconversion level.
$\overline{S1} \leq$ OD sample < $\overline{S3}$	X in EU/ml (0.125 - 0.5 EU/ml)	Insufficient seroconversion level according to PLATELLA™ RABIES II test.
OD sample < $\overline{S1}$	-	Undetectable seroconversion.

10 - LIMITATIONS OF THE TEST

Insufficient or undetectable seroconversion can possibly be reported in recently vaccinated animals, due to delayed immune response. Similarly to the official neutralising antibody titration method and in accordance with regulations (EC) No 998/2003 effective July 3, 2004, the PLATELIA™ RABIES II test should be carried out on samples taken at least thirty days after vaccination.

11 - EQUIPMENT AND MATERIAL REQUIRED BUT NOT SUPPLIED

Equipment

- Microplate reader equipped with 450 and 620 nm filters (*)
- Microplate incubator set at 37°C ± 2°C.
- Manual, semi-automatic or automatic microplate washer (*)
- Vortex® mixer.

(*) Contact us for detailed information about Bio-Rad instruments validated by our technical departments.

Material

- Automatic or semi-automatic adjustable or preset pipettes or micropipettes to measure and deliver 10 to 1000 µl and 1, 2 and 10 ml.
- Graduated cylinders of 25 ml, 50 ml, 100 ml, and 1 000 ml capacity.
- Disposable tubes.
- Distilled or deionized water.
- Sodium hypochloride (bleach).
- Absorbent paper.
- Goggles or safety glasses.
- Disposable latex gloves.
- Container for biohazard waste.

12 - PRECAUTIONS

1. Precautions

The quality of the results depends on correct implementation of the following good laboratory practices:

- Reagents must be stored at +2°C to +8°C.
- Do not use expired reagents.
- Do not mix reagents from different lots within a given test run with the exception of generic reagents R2, R8, R9, R10.
- Before use, wait 30 minutes for the reagents to adjust to the laboratory temperature.
- Thoroughly reconstitute reagents, avoiding any contaminations.
- Do not perform the test in the presence of reactive vapors (acids, alkalis, aldehydes) or dust that could alter enzyme activity of the conjugate.
- Use glassware thoroughly washed and rinsed with deionized water or preferably, disposable material.
- Do not leave the microplate more than 5 minutes between the end of washing operation and the reagent distribution.
- Check the accuracy of the pipettes and the good operation of the apparatus used.
- Never use the same container to distribute conjugate and the development solution.

- The enzyme reaction is very sensitive to metal or metallic ions. Consequently, do not allow any metal element to come in contact with the various conjugate or substrate solution.
- The development solution (substrate buffer + chromogen) must be colourless. The appearance of a blue colour within a few minutes after reconstitution of the development solution indicates that the reagent can not be used and must be replaced. Preparation of this solution can be made in a clean disposable plastic tray or glass container that has first been pre-washed with 1N HCl, and rinsed thoroughly with distilled water and dried. Store the solution in the dark.
- Use a new pipette tip for each sample.
- Do not change the assay procedure.
- Washing of the wells is a crucial step in the procedure: respect the recommended number of washing cycles and make sure that all wells are completely filled, then emptied. Inadequate washing can give incorrect results.

2. Hygiene and safety instructions

Generally, hygiene conditions, biosafety measures and good laboratory practices must be in agreement with recommendation of regular authorities of the country.

All the reagents of the kit are intended for "in vitro" veterinary diagnostic use - dog, cat and fox sera.

- Wear disposable gloves when handling reagents and samples. Thoroughly wash your hands after sample and reagent handling.
- Do not pipette with the mouth.
- Consider any material directly in contact with the samples and the reagents, as well as the wash solutions, as infectious material.
- Avoid spilling samples or solutions containing samples.
- Spills may be rinsed with bleach diluted to 10% Chl. If the contaminating fluid is acid, spills must be initially neutralized with sodium bicarbonate, then cleaned with bleach and dried with absorbent paper. The material used for cleaning must be discarded after decontamination.
- Samples and contaminated material and products should be discarded after decontamination
 - either by immersion in bleach at the final concentration at 5% of sodium hypochloride for 30 min.
 - or by autoclaving at 121°C for 2 hours minimum.

Caution: Do not introduce solutions containing sodium hypochloride into the autoclave.

- Chemicals should be handled and disposed of in accordance with Good Laboratory Practice.
- Reagents containing 0.1% ProClin™ 300 are classified as irritating preparations according to European legislation.



Xi
(0.1% ProClin™ 300)

R: 43 May cause sensitisation by skin contact.

S: 28-37 After contact with skin, wash immediately with plenty of water. Wear suitable gloves.

13 - LITERATURE

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