



# Identification of *Trichinella* spp. proteins recognized by specific IgG in serum of infected pigs by western blotting

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#### 1 AIM AND FIELD OF APPLICATION

This document defines an immune-enzymatic method, western blotting, for the qualitative detection of anti-*Trichinella* spp. IgG in swine sera.

The method can be used as confirmatory test to screen pig populations on the exposure of swine to *Trichinella* spp. during sero-epidemiological surveys.

#### 2 PRINCIPLE OF THE METHOD

Partially purified *Trichinella spiralis* excretory/secretory antigens are separated by SDS–PAGE electrophoresis. After running, proteins are transferred to a nitrocellulose membrane, which is then cut into 20 strips. Each strip is blocked with skimmed milk to saturate the sites free from antigens and singularly located in plastic trays.

Then strips are incubated with diluted control and test pig serum samples, allowing anti-*Trichinella* IgG that may be present in sera, to bind to the antigens on the nitrocellulose membranes.

The antibodies not binding to the antigens are eliminated by washing; goat anti-swine IgG conjugated with horseradish peroxidase (HRP) is then added to each strip. This second incubation allows the conjugate to bind to the swine antibodies that were bound to the antigens onto the strip surface.

The excess conjugate is eliminated by washing, and the activity of the enzyme HRP bound to the swine antibodies, is measured by adding a chromogen substrate. This allows to display a pattern of stained bands corresponding to those proteins that reacted specifically with anti-IgG antibodies to *Trichinella* spp., present in the pig serum (figure 1).



Figure 1. Schematic representation of the method

#### 3 REFERENCES

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#### 4 ACRONYMS

Antigen

- Ab Antibody
- Ag E/S Excretory/secretory antigens
- BSA Bovine serum albumin
- Wb Western blotting

#### **5 DEVICES/INSTRUMENTS**

- 5.1 Power supplies
- 5.2 Thermomixer
- 5.3 pH meter
- 5.4 Analytical balance or technical balance
- 5.5 Refrigerator, 1÷8°C
- 5.6 Freezer, ≤ 50°C
- 5.7 Freezer,  $\leq -15^{\circ}$ C
- 5.8 Magnetic stirrer
- 5.9 Vortex mixer
- 5.10 Micropipettes (0.5-10 µL, 5-100 µL, 15-300 µL, 50-1000 µL)
- 5.11 Water analytical grade
- 5.12 Tray X Cell SureLock Mini-cell for vertical electrophoresis
- 5.13 Mini Trans-Blot Module Biorad
- 5.14 Scalpel
- 5.15 Pipette aid
- 5.16 Serological pipettes

#### 6 REAGENTS AND CHEMICALS

#### Electrophoresis

6.1	Preparation of the test and control sample for electrophoretic run					
	Ag E/S	100 µg				
	NuPAGE LDS Sample Buffer (4X)	50 µL				
	NuPAGE Reducing Agent(10X)	20 µL				
	Analytical Grade water	up to 200 µL				
<u>.</u>						

Stir using a magnetic stirrer until completely dissolved.

 6.2 Gel for electrophoresis : NuPAGE Novex 10% Bis-Tris Mini Gels 1.0mm X2D well Invitrogen
6.2 Drosteined melecular weight merkeres

6.3	Prestained molecular weight markers:	
F	restained SDS-PAGE Standards Low range Biorad	10 µL
6.4	Antioxidant : NuPAGE	500 µL



6.5 Running Buffer:	
Mops SDS Running Buffer (20X)	50mL
Analytical grade water	up to 1000mL
Use the solution within 1 month.	
6.6 Transfer Buffer:	
Tris-Glycine (25X)	40mL
Metanol	200mL
Analytical grade water	up to 1000mL
Use the solution within 1 month	
6.7 Nitrocellulose membrane 0,2 μm	
6.8 Ponceau S Solution	20 mL
6.9 Mini-Incubation Trays	
Western blotting	
6.10 TBS Buffer	
Tris 0,2 M	2.4 g
NaCI 3M	17.5 g
Analytical grade water	up to 2000mL
Use the solution within 6 months	
Dissolve the compounds in about 1000 mL of analytical	
the pH (7.8 $\pm$ 0.2) and then bring the solution to the final v	volume; store at 1÷ 8°C.
6.11 TTBS Buffer	
Tween 20	1 mL
Tris 0,2 M	2.4 g
NaCI 3M	17.5 g
Analytical grade water	up to 2000mL
Use the solution within 6 months	
Dissolve the compounds in about 1000 mL of analytical the pH (7.8 $\pm$ 0.2) and then bring the solution to the final	
	volume, store at 1÷ 0°C.
6.12 Sera and conjugate diluter	
Skimmed milk powder	1.5 g
TTBS Buffer (pH 7,4)	up to 50 mL
Prepare the solution immediately before use, as follows: mL tube; add 40 mL of TTBS buffer and mix by vorted bring it to volume. If refrigerated, use the solution within	xing until milk is completely dissolved and
6.13 Washing solution	
TTBS Buffer (6.10)	2000 mL
6.14 Blocking solution	
Skimmed milk powder	2.5 g
TTBS Buffer	50 mL
Prepare the solution immediately before use, as follows mL tube; add 40 mL of TTBS buffer and mix by vortexin bring to volume. If refrigerated, use the solution within 24	g until the milk is completely dissolved and
6.15 Phosphate buffered saline (PBS), pH 7.3±0.2	
KH <sub>2</sub> PO <sub>4</sub>	0.34 g
Na <sub>2</sub> HPO <sub>4</sub>	1.21 g
NaCl	8.0 g

Analytical grade water

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up to 1000 mL





Dissolve compounds in 750 mL of analytical grade water under magnetic stirring. Check the pH  $(7.3\pm0.2)$  and then bring the solution to the final volume; refrigerate.

If refrigerated, use the solution within 6 months.

6.16 Anti-Trichinella spp. IgG positive control pig serum samples (positive controls).

Sera collected at day 60 p.i. from pigs experimentally infected with *Trichinella* spp.

Serum samples from Trichinella free pigs (negative controls) 6.17

6.18 Chromogenic DAB (3, 3'diaminobenzidina).

Dissolve a DAB tablet in 15 mL of PBS. Mix by vortexing until the DAB is completely dissolved.

6.19 Hydrogen peroxide

Add to chromogenic immediately before use

12 µL

Conjugate: peroxidase labeled (horseradish peroxidase) anti-swine IgG goat antibodies . 6 20 If the reagent is freeze-dried, it shall be rehydrated before using, by vortexing with analytical grade water until completely dissolved.

Once rehydrated, store the conjugate refrigerated at  $1 \div 8^{\circ}$  C for one week.

To determine the optimal working dilution of the conjugate (that is, the dilution which displays the specific bands in positive controls with a minimum of background), the working dilutions recommended by the manufacturer are tested. If none of these dilutions results optimal, the further dilutions of the reagent shall be tested until the optimal one is obtained.

Distribute the conjugate in aliquots and stored frozen at <-50°C. In these conditions it remains stable for at least 5 years. Before testing, dilute an aliquot of conjugate to the optimum concentration with the dilution solution. Once diluted, store the conjugate refrigerated at 1+8 ° C and use within 24 hours.

### 7 PROCEDURE

#### 7.1 Samples and marker preparation

Use 100µg of Ag E/S for a gel and add all reagents. Stir using a magnetic stirrer until complete dissolution. Incubate 10' at 70°C in the Thermomixer. Take an aliquot of the molecular weight markers containing 10µl and incubate 1' to 40°C in the heating block.

#### 7.2 Gel electrophoresis loading

Put one or two gels in the electrophoresis tray for vertical electrophoresis (Figure 2). Fill half of the electrophoresis tray (about 500 mL) with running buffer. Add 200 mL of running buffer with 500 µL of antioxidant in the central part of the tray.

Load the proteins in the gel by micropipette, connect the electrophoresis tray to the electrophoresis power supply (5.1) and allow the separation of the proteins by applying 150 V for 1h at room temperature (RT).



Figure 2. Representation of the vertical electrophoresis cassette.





### 7.3 Transfer to nitrocellulose membranes

At the end of the electrophoresis running, put the gel containing the proteins on a nitrocellulose membrane, according to the scheme of the Mini Trans-Blot Module Biorad shown in the figure 3.



Figure 3. Scheme of the transfer from gel to nitrocellulose membrane protein.

Place the assembled system in the X cell SureLock (Figure 4) with transfer buffer and ice block, connect the power supply unit for 1 h at  $+ 4^{\circ}$ C at 38 mA, so that proteins migrate from the gel, to the nitrocellulose.



Figure 4. X cell SureLock protein transfer

### 7.4 <u>Verification of proteins transfer</u>

At the end of the transfer open the cassette, remove the nitrocellulose and place it in a tray with dye in order to verify the effective transfer. Wash with tap water to remove excess dye and cut the filter in strips using a scalpel.

### 7.5 <u>Nitrocellulose blocking</u>

Put the strips in the blocking solution overnight at +4°C. Then, wash the strips with TTBS buffer. Once the strips are dried, can be stored at RT up to two months wrapped in filter paper.

### 7.6 <u>Test and control samples preparation</u>

Thaw the test sera, positive and negative control sera by storing at +1-8°C for at least 5 h.

Once thawed, keep them in an ice bath and stir them by vortexing before use.

Dilute 1:100 the test and control sera as follows: in a 1-2 mL conical bottom tube dispense 1 mL of diluting solution with a serological pipette placed on a pipettor and discard 10  $\mu$ L through a micropipette. Alternatively, dispense 990  $\mu$ L diluting solution with a micropipette and add 10  $\mu$ L of serum. Diluted sera can be stored refrigerated (5.5) for up to 24 hs.





## 7.7 <u>Analytical method</u>.

- a. Bring out the serum samples from the refrigerator.
- b. Dispose the nitrocellulose strips in the appropriate Mini-Incubation Trays and wash them with TTBS for 5' at RT.
- c. Incubate the control and test sera for 1 hour at RT.
- d. Wash 3 times for 5' at RT with TTBS solution.
- e. Incubate the diluted conjugate 1h at RT
- f. Wash 3 times for 5' at RT with TTBS solution.
- g. Wash 1 time for 5' at RT with TBS solution.
- h. Add the chromogenic until you see the bands.
- i. Stop reaction with tap water

### 8 INTERPRETATION OF THE RESULTS

The electrophoretic separation of the proteins of *T. spiralis* and their subsequent transfer on nitrocellulose membrane are considered valid when all the markers (or standards) of molecular weight (MW) colored, used in each gel, were:

- electrophoretically separated,
- transferred to nitrocellulose membrane.

Furthermore, the relative mobility of each standard shall be in the range previously established by means of three independent experiments. For standard of 104 proteins, 94, 51, 36, 28 and 19 kD, the relative mobility are: 0.13, 0.24, 0.41, 0.66, 0.80, 0.91, respectively (S.D.  $\pm$  0.2; Figure 5); for the positive control, see the relative mobility as previously published (Gomez Morales et al., 2014, with S.D.  $\pm$  0.2).

If even one relative mobility is not within the specific range, the results shall be considered invalid and the test has to be repeated.

The specific *T. spiralis* E/S proteins reacting with the reference and test sera from pigs are identified by calculating their molecular weight (MW) as follows (Figure 5):

- 1. in a Microsoft Word Excel electronic sheet trace a PM log graph of pre-stained standards to their relative migration distance (Rf);
- 2. interpolate the Rf of *T. spiralis* specific E/S proteins which react with pig test and reference sera, to calculate their molecular weights;
- 3. the specific proteins that react with pig sera, both reference and test sera, must show a triple-band pattern in a region comprised between 48 and 72 kDa (the first band from 48 kDa to 55 kDa, the second band from 59 kDa to 62 kDa and the third band from 64 to 72 kDa).



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	A	В	С	D	E	F	G	н	I
1	MW (kD) of the prestained molecular marker proteins	Migration distance (cm) of the prestained molecular marker proteins		Rf prestained molecular marker proteins			Rf Mean	Log MW	
2		experiment 1	experiment 2	experiment 3	experiment 1	experiment 2	experiment 3	x	У
3	104	0,6	0,9	0,7	=PRODUCT(B3;1/B10)	=PRODUCT(C3;1/C10)	=PRODUCT(D3;1/D10)	=MEAN(E3:G3)	=LOG10(A3)
4	94	1,2	1,5	1,3	=PRODUCT(B4;1/B10)	=PRODUCT(C4;1/C10)	=PRODUCT(D4;1/D10)	=MEAN(E4:G4)	=LOG10(A4)
5	51	2,2	2,5	2,3	=PRODUCT(B5;1/B10)	=PRODUCT(C5;1/C10)	=PRODUCT(D5;1/D10)	=MEAN(E5:G5)	=LOG10(A5)
6	36	3,6	3,8	3,7	=PRODUCT(B6;1/B10)	=PRODUCT(C6;1/C10)	=PRODUCT(D6;1/D10)	=MEAN(E6:G6)	=LOG10(A6)
7	28	4,4	4,7	4,5	=PRODUCT(B7;1/B10)	=PRODUCT(C7;1/C10)	=PRODUCT(D7;1/D10)	=MEAN(E7:G7)	=LOG10(A7)
8 9	19	5	5,2	5,1	=PRODUCT(B8;1/B10)	=PRODUCT(C8;1/C10)	=PRODUCT(D8;1/D10)	=MEAN(E8:G8)	=LOG10(A8)
9 10 11 12	Migration distance of the front	5,5	5,8	5,6					
13	-	Migration distance of the unknown							
14		proteins			Rf	unknown proteins		Rf Mean	Log MW
15		1.8	2,1	1.9	=PRODUCT(B15;1/B10)	=PRODUCT(C15;1/C10)	=PRODUCT(D15;1/D10)	=MEAN(E15:G15)	=TREND(I3:18;H3:H8;H15;TRUE)
16		2,2	2,4	2,3	=PRODUCT(B16;1/B10)	=PRODUCT(C16;1/C10)	=PRODUCT(D16;1/D10)	=MEAN(E16:G16)	=TREND(I3:18;H3:H8;H16;TRUE)
17		2,5	2,8	2,6	=PRODUCT(B17;1/B10)	=PRODUCT(C17;1/C10)	=PRODUCT(D17;1/D10)	=MEAN(E17:G17)	=TREND(I3:18;H3:H8;H17;TRUE)
18 19									
20									=POWER(10:115)
21						Molecular weight of the	unkown proteins	=POWER(10;116)	
22									=POWER(10;117)



Figure 5. The above table shows the results of experiments carried out to determine the relative mobility of each molecular weight marker. The graph on the left shows the regression line of the relative distance of the reference sera as a function of their molecular weight. The graph on the right shows the pattern of specific proteins and the E/S Ag of *T. spiralis*, which react with pig sera in

#### Wb.

### 9 CHARACTERISTICS OF THE METHOD

This method has been characterized in terms of sensitivity, specificity and reproducibility. The results were used to confirm that the method is suitable for the intended purpose and are reported in the relevant validation file, to which reference is made.

### **10 SAFETY MEASURES**

This test method can be performed only by experienced personnel. As potentially infected sera with zoonotic pathogens are handled, analysts are provided with personal protective equipment, such as disposable gloves and lab coats. For safety measures to be adopted during testing, refer to the WHO Laboratory Biosafety Manual – Third Edition (http://www.who.int/csr/resources/publications/biosafety/WHO CDS CSR LYO 2004 11/en/).