Analysis of Diagnostic Efficiency of Excretory-Secretory Antigens for *Trichinella spiralis* and *Trichinella nativa* in ELISA

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Abstract

The comparative studies of diagnostic efficiency of excretory-secretory antigens of *Trichinella spiralis* and *Trichinella nativa* were performed using blood sera of rats from Wistar line experimentally infected with Arctic trichinellae. For animal infection and antigen preparation *Trichinella* from muscles of wild carnivorous mammals from Arctic regions of Russia were used. When antigen from *T. nativa* larvae was used to analyze titers of sera of rats experimentally infected with Arctic *Trichinella*, a significant increase in efficacy of ELISA was detected. E.g., sera of rats infected with trichinellae from ringed seals retained in ELISA with *T. nativa* antigen values higher than diagnostic level at titers of 1:6400 - 1:12800, while titer of those same sera when using *T. spiralis* antigen was no higher than 1:200 - 1:400.

Keywords

ELISA, *Trichinella nativa*, Diagnostic Efficiency, Laboratory Infection, Wistar Rats, Blood Sera, Excretory-Secretory Antigen

1. Introduction

Epidemiological situation regarding trichinellosis in several regions of Russian Federation (RF) is a stable one,

though the problem remains serious. It is especially pronounced in regions of Arctic North where human trichinellosis is clearly connected with that of land and sea mammals. Hunting and fishing are historically the main source of sustenance for natives of the regions, they determine social and economic structure of coastal settlements on Arctic and Pacific ocean shores. An analysis of data on epizootology of trichinellosis at Chukotka territory shows an advisability of broad seroepidemiological studies among natives of the region [1] [2].

It’s well known that the success of in Vivo immunological diagnostics is mainly determined by quality of an antigen used in enzyme-linked immunosorbent assay (ELISA). Currently at all Russian territories where trichinellosis is endemic, test systems (immune enzyme analysis, non-direct gem agglutination) based on antigens obtained from infective larvae of European strains of *T. spiralis* circulating throughout biocenosis of North Caucasus, Krasnodar Region, and Belarus Republic are used [3]. However, significant differences in results of seroepidemiological monitoring in infection foci of South and North territories of RF suggest that blood of people infested with “Arctic” trichinellae contains immune complexes which are not revealed by test systems based on *Trichinella spiralis* antigens [3] [4]. Accordingly, in this study we’ve decided to precede studies of blood sera of Chukotka natives for antibodies specific to *T. nativa* antigens with comparative analysis of diagnostic efficiency of excretory-secretory antigens of European and Arctic strains of trichinellae in ELISA regarding blood sera from laboratory animals experimentally infected with Arctic isolates of trichinellae.

To work out ELISA parameters we have performed a comparative analysis of efficiency of two immune-enzyme test systems based on *T. spiralis* and *T. nativa* antigens, respectively. The sensitivity of these methods is measured through results of the reaction with the same group of blood sera from experimentally infected Wistar line rats. Necessity of such studies is due to the fact that no respondents among the Arctic regions natives whose blood was tested had clinical pattern of trichinellosis, so objective assessment of efficiency of the test using different antigens in ELISA is not possible.

2. Materials and Methods

2.1. Animals

50 rats from Wistar line, weight 160 - 180 g; 40 golden hamsters, weight 100 - 120 g.

2.2. Materials

- *Trichinella* larvae initially obtained from muscles of naturally infected carnivores from Arctic Region and larvae of standard laboratory strain of *Trichinella spiralis* (from domestic pig of Belarus) passed for many years through laboratory rodents in Skrjabin Institute’s vivarium;
- Excretory-secretory antigens of *Trichinella spiralis* and *Trichinella nativa*;
- Blood serum of rats from Wistar line—clinically healthy and experimentally infected with Arctic trichinellae and other helminths (*E. granulosus*, *E. multilocularis*, *Hymenolepis nana*);
- Anti-species conjugate antibodies to rat IgG marked with horseradish peroxidase;
- Necessary reagents and assays for ELISA prepared according to European Union Reference Laboratory for Parasites recommendations.

2.3. Methods

2.3.1. Obtaining Infective *Trichinella* Larvae

To exclude a possibility of antigenic mimicry affecting an analysis results, larvae for secretion of Arctic *Trichinella* antigen were obtained from hamsters exclusively, while blood serum for ELISA was taken from rats infected with *T. nativa*. For simultaneous obtainment of necessary volume of viable infective *Trichinella* larvae able to produce the antigen all hamsters used were arbitrarily divided into 6 groups of 10 hamsters each. Five groups were infected with Arctic trichinellae (from a blue polar fox (*Alopex lagopus*), a sled dog (*Canis familia-ris*), Polar bear (*Ursus maritimus*), wolverine (*Gulo gulo*) and a ringed seal (*Phoca hispida*) from Chukotka and Yakut Region)) and one with standard *T. spiralis* strain (from domestic pig of Belarus). The inequality in distribution of the animals between supra-groups (Arctic *Trichinella* and *T. spiralis*) was due to very low reproduction potential of all Arctic *Trichinella* strains (relative to *T. spiralis*) shown earlier in 4 passages of specimen of these isolates through laboratory rodents [5].
Golden hamsters and rats were infected per os with larvae of *T. nativa* and *T. spiralis* at dose of 10 larvae per 1 g of live weight. After 1.5 - 2 months the animals were sacrificed through cervical dislocation, skinned, internal organs removed and carcasses digested in artificial gastric liquid. Larvae obtained were rinsed multiple times with sterile saline with added antibiotics according to European Union Reference Laboratory for Parasites recommendations [6].

Muscle tissues of the experimentally infected animals were studied using method of trichinellloscopy.

### 2.3.2. Incubation of Infective Larvae

Further work on obtaining the larvae excretory-secretory products was performed in sterile conditions. *Trichinella* larvae were incubated in Petri dishes in Dulbecco’s Modified Eagle’s Medium (DME) substrate with L-glutamine (40 μl/ml) and above-mentioned antibiotics at 38.5°C in common thermostat with larvae concentration of 5000 per 1 ml of medium.

### 2.3.3. Obtaining Excretory-Secretory *Trichinella* Antigens

Protein products were obtained from the medium where larvae were incubated in sterile conditions. The medium with *Trichinella* excretory-secretory products was sampled once per 24 hr for 3 - 5 days; after each product sampling initial volume was restored through addition of new medium with L-glutamine. *Trichinella* larvae viability was daily assessed under microscope, if more than 30% of the larvae were dead, cultivation was stopped. Excretory-secretory products obtained were combined, then dialyzed for 48 hr in distilled water with NaCl added (1:10) at 4°C, with buffer solution changed each 6 - 8 hrs. After the dialysis the antigen was concentrated using polyethylene glycol (PEG)-8000 at 4°C for 6 - 8 hrs. The protein concentration was determined using spectrophotometer at wave length of 280 nm after which the antigen was passed through a bacterial membrane with pore diameter of 22 - 24 μm. Thereafter the antigens obtained were used for sensitization of polystyrene plates in ELISA at 5 μg/ml in carbonate buffer saline (pH 9.6) for 16 hours. The antigen solution was kept at −18°C, or at −70°C in kelvinator [7]-[9].

### 2.3.4. Obtaining and Maintenance of Diagnostic Blood Serum

No less than 1.5 - 2 ml of blood serum were taken from each animal and placed into a thermostat for 1 hour at 37°C. The resulting blood concentrate was contoured with sterile glass stick and transparent liquid phase sampled. These serum probes were placed into microtubes, marked and kept at +4°C - +8°C for no more than 4 days before an analysis. In cases of erythrocytes sediment present, the serum was centrifuged at 3000 rotations/minute for 10 minutes.

### 2.3.5. Determining an Optimal Titer of Anti-Species Peroxidase Conjugates

Immunoglobulins from blood serum of intact rats to be used in experiment and clinically healthy donors were obtained through sedimentation using ammonium sulphate (50% - 70% saturation). For titration of anti-species conjugates twofold dilutions in phosphate-saline buffer were prepared, starting with 1:100 and up to 1:25600, ELISA was performed according to standard scheme using rat immunoglobulins adsorbed on polystyrene as an “antigen”. As a result, specific titer of anti-species conjugate against *IgG* of rats had been found to be 1:6000.

### 2.3.6. Conditions for Solid-Phase ELISA

The analysis was performed by using indirect variant with polystyrene 96-pits strip microplates from Gosniipolimer, Moscow. For solid phase sensitization excretory-secretory antigens with molecular mass of 29 - 63 kDa were used. Measuring of protein concentration in initial solution was performed using spectrophotometer at wave length of 280 nm. *Trichinella* antigen was placed onto polystyrene strips at 5 μg/ml concentration. A solid phase sensitization continued for 16 - 18 hours at 4°C, after which pits contents were removed and pits rinsed 3 - 5 times with phosphate-saline buffer. Preparation of non-specific reagents for immune-enzyme reaction: phosphate-saline buffer, buffer solutions for sera and conjugate, 3.3',5.5'-tetramethylbenzidyne-based substrate solution (TMB), stop-reagent (0.5 M sulphuric acid solution)—were performed by using traditional methods with some modifications.

Results were assessed on spectrophotometer 340/ATC by STL-Labsistems (Austria) with automated reader and vertical light ray at wave length of 450 nm.
2.3.7. Analysis of Sensitivity and Specificity of Excretory-Secretory Antigens from *T. nativa* and *T. spiralis* Obtained through ELISA

The activity and specificity of excretory-secretory antigens from *T. nativa* and *T. spiralis* were determined through ELISA reaction using blood sera of rats from Wistar line experimentally infected with *Trichinella* from Arctic isolates obtained from muscles of carnivores from Chukotka and Yakutia, as well as those of rats infected with other helminths (*E. granulosus*, *E. multilocularis*, *Hymenolepis nana*) and clinically healthy rats; samples of the latter two categories were taken from serum bank of Skrjabin’s Institute.

Initially, to work out ELISA parameters we had performed a comparative analysis of efficiency of two immune-enzyme test systems: the first was based on *T. spiralis* antigen, while the second on that of ‘Arctic’ *T. nativa*. For this comparative study, ELISA was performed on two plates in parallel, using the same rat sera and reagents but different antigens—*T. nativa* and *T. spiralis*.

2.3.8. Detection of Anti-*Trichinella* Antibodies

Detection of the antibodies was performed in general accordance to methods recommended by European Union Reference Laboratory for Parasites, with some changes and notions shown above [6].

2.3.9. Statistical Analysis

Extinction value (%) was calculated according to formulae given by Community Reference Laboratory for Parasites [6].

3. Results and Discussion

A *Trichinella* larvae count after digestion of carcasses of experimentally infected animals in artificial gastric liquid had shown that parasite load (mean for a group) in hamsters infected with laboratory strain of *T. spiralis* was 2317 larvae per 1 g of bone-meat mince. The same parameter was 1.86 larvae/g from ringed seal, 76.12 larvae/g from blue polar fox, 656.08 larvae/g from Polar bear and 23.75 larvae/g from wolverine. All larvae at muscle stage from carcasses of hamsters infected with Arctic isolates were pooled into one sample for a total of 2.6 ml of viable trichinellae ready for cultivation in artificial medium.

Totally we had obtained about 100 ml of purified excretory-secretory antigen of *T. spiralis* with protein contents of 580 μg/ml and 23 ml of one from *T. nativa*, with protein contents of 286 μg/ml. These antigen proteins were later used for sensitization of polystyrene plates and subsequent serological studies of blood sera aimed at finding specific antibodies to *T. nativa* antigens using ELISA.

Microscopic studies of experimental rats sacrificed after 45 - 60 days p.i. (Figure 1) have shown that rats infected with Arctic *Trichinella* isolates had most larvae in muscles at various stages of destruction, mainly mumified, the helminths were surrounded with extensive cell infiltrates. There were few living *Trichinella* larvae in the muscles. Most parasites were at the stage of cell-enzymatic resorption.

As a result of comparative studies of efficiency of immune-enzyme test systems based on *T. nativa* and *T. spiralis* antigens significant differences in optical density for the same serum samples when using antigens of different *Trichinella* species as solid phases have been shown (Figure 2). In such cases mean optical density
Figure 2. Indicators of instrumental control in ELISA with *T. nativa* and *T. spiralis* antigens (OD given is mean for series combined from blood sera of rats experimentally infected with *Trichinella* obtained from the host species shown on X-axis). K-mean is an indicator of optical density (OD) for negative controls of the reaction.

(OD) of sera in groups of negative controls and rats infected with other helminths were practically the same: 0.127/0.124; 0.120/0.125; 0.099/0.112; 0.175/0.160; 0.303/0.310 (for blood serum of the rat with extensive *Echinococcus* cyst). All serum probes from rats infected with Arctic trichinellae were considered a “positive control”. They were these probes exactly that have shown significant differences in optical density values when antigens from two *Trichinella* species were used in ELISA. In all cases OD<sub>450nm</sub> values when using *T. spiralis* laboratory strain were significantly lower than in cases of *T. nativa* antigen (see Figure 2). So, sera of rats infected with trichinellae from ringed seals retained in ELISA with *T. nativa* antigen values higher than diagnostic level at titers of 1:6400 - 1:12800, while titer of those same sera when using *T. spiralis* antigen was no higher than 1:200 - 1:400. The similar pattern was seen with titers of sera of rats infected with trichinellae from sled dog and polar fox. Of course, most sera of line rats experimentally infected with large dosage of trichinellae (10 larvae/gram) were ELISA-positive at significantly higher than diagnostic level on plates with antigens from both *Trichinella* species (Table 1).

The results obtained are in good agreement with several earlier works. E.g., Serrano, Perez, Reina, Navarrete [10] had studied a diagnostic efficiency in ELISA of antigens of 5 Spanish *Trichinella* isolates. The study’s goal was a development of a test system to detect homologous and heterologous infections. Using blood sera from 243 pigs the authors have registered cross reactions between all *Trichinella* isolates, but homologous sera have shown higher efficiency in ELISA.

Smith and Snowdon [11] have studied diagnostic efficiency of ELISA using blood sera of pigs vaccinated and infected with *T. nativa* and *T. spiralis*. In all cases humoral cross reaction had been detected. However, in all cases of homologous sera being used, diagnostic levels of OD five or more times higher than negative control level had been shown, while heterologous blood sera had shown OD level high, but not reaching this diagnostic level.

Sukura *et al.* [12] have shown using Western blot with *T. spiralis* and *T. nativa* antigens obvious cross reactivity with sera of infected raccoon dogs. However, antigens with molecular mass of 40 - 76 kDa reacted with homologous sera much stronger. It should be noted that this experiment have shown impossibility of early detection of *Trichinella* infection using heterologous antigens. For example, usage of blood sera with homologous antigens allowed detecting specific antibodies two weeks earlier than with heterologous antigens in Western blot and one week earlier in ELISA. At later stage (12 weeks p.i.) indexes of OD in ELISA were significantly higher when using homologous components than with heterologous ones.
Table 1. Diagnostic efficiency of ELISA with sera of Wistar rats experimentally infected with Arctic *Trichinella*.

<table>
<thead>
<tr>
<th>Blood sera of rats experimentally infected with</th>
<th>Numbers of animals in the group</th>
<th>ELISA results</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td><em>T. nativa</em> isolate from sled dog, Chukotka</td>
<td>5</td>
<td>5</td>
<td>100% 5 -</td>
</tr>
<tr>
<td><em>T. nativa</em> isolate from ringed seal, Chukotka</td>
<td>5</td>
<td>5</td>
<td>100% 5 -</td>
</tr>
<tr>
<td><em>T. nativa</em> isolate from Polar fox, Chukotka</td>
<td>5</td>
<td>5</td>
<td>100% 5 -</td>
</tr>
<tr>
<td><em>T. nativa</em> isolate from Polar bear, Yakutia</td>
<td>5</td>
<td>5</td>
<td>100% 4 2</td>
</tr>
<tr>
<td><em>T. nativa</em> isolate from wolverine, Chukotka</td>
<td>5</td>
<td>5</td>
<td>100% 3 1</td>
</tr>
<tr>
<td>E. granulosis</td>
<td>3</td>
<td>1</td>
<td>2 1</td>
</tr>
<tr>
<td>E. multilocularis</td>
<td>3</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Hymenolepis nana</td>
<td>3</td>
<td>-</td>
<td>3</td>
</tr>
<tr>
<td>Clinically healthy rat</td>
<td>34</td>
<td>34</td>
<td>34</td>
</tr>
<tr>
<td>Total</td>
<td>50</td>
<td>100% 95.35%</td>
<td>80% 95.35%</td>
</tr>
</tbody>
</table>

4. Conclusion

So, both experimental and literature data clearly demonstrate that in large-scale seroepidemiological studies at conditions far from laboratory standard, such a difference between sensitivity when using heterologous and homologous antigens may lead to a significant decrease in the test efficiency, with many positive sera samples falling behind the method’s sensitivity threshold or becoming “doubtful”. So, it is necessary when performing epidemiological monitoring in trichinellosis foci in Arctic to use an excretory-secretory antigen from larvae of Arctic natural isolates of *T. nativa* as an active component for ELISA.

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References


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