TOXOCARIASIS: SEROLOGICAL DIAGNOSIS BY INDIRECT ANTIBODY COMPETITION ELISA

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SUMMARY

Toxocariasis is caused by infection of man by Toxocara canis and Toxocara cati larvae, the common roundworm of dogs and cats. Because larvae are difficult to detect in tissues, diagnosis is mostly based on serology. Non specific reactions are observed mainly due to cross-reactivity with Ascaris sp antigens.

This investigation aimed at developing and evaluating an indirect antibody competition ELISA (IACE) employing a specific rabbit IgG anti-Toxocara canis excretory-secretory antigens as the competition antibody, in order to improve indirect ELISA specificity performed for toxocariasis diagnosis. For that, the rabbit IgG was previously absorbed by Ascaris suum adult antigens.

Sensitivity and specificity of IACE were first evaluated in 28 serum samples of mice experimentally infected with T. canis embryonated eggs. Adopting cut-off value established in this population before infection, sensitivity and specificity were 100% after 20 days post-inoculation.

For human population IACE was evaluated using sera from 440 patients with clinical signs of toxocariasis and the cut-off value was established with 60 serum samples from apparently healthy individuals. Using as reference test the indirect ELISA performed by Adolfo Lutz Institute, sensitivity was 60.2%, specificity was 98% and concordance was 77.3%.

Repeatability of IACE was evaluated by the inter-reactions variation coefficient (2.4%).

KEYWORDS: Toxocariasis; Visceral larva migrans; Toxocara canis; Indirect antibody competition ELISA; Immunodiagnosis.

INTRODUCTION

Visceral larva migrans was first described by BEAVER et al.⁷ to define a clinical syndrome of man characterized by hepatomegaly, fever, chronic eosinophilia and hypergamaglobulinemia.⁴ A wide range of zoonotic helminths can cause visceral larva migrans but Toxocara canis and Toxocara cati, the common roundworms of dogs and cats are still the most frequently incriminated agents.

Transmission to human beings occurs by ingestion of contaminated soil or eggs on hands and fomites. Direct contact with infected dogs and cats plays a secondary role in transmission because eggs need an extrinsic period to become infective. Neither worms nor eggs are eliminated in human feces and because larvae are difficult to detect in tissues, diagnosis is mostly based on serology.

Since the studies made by de SAVIGNY,⁹ the antigens mostly used for the immunodiagnostic tests derive from larvae cultivated in vitro and are referred to as Toxocara excretory-secretory (TES) antigens.¹⁰

Not all TES antigens are species or genus specific and serum samples from patients with ascariasis, filariasis, schistosomiasis and strongyloidiasis show reactivity with TES antigens.¹¹,¹²,¹³,¹⁴,¹⁵,¹⁶

Evaluation of the true sensitivity and specificity of serology tests for toxocariasis in human populations is not possible because of the lack of parasitological methods to definitively diagnose the disease and to exclude infections in controls.¹⁷

Nevertheless, the introduction of enzyme-linked immunosorbent assay (ELISA) based upon TES antigens has resulted in greatly increased specificity.¹⁸

In areas like Brazil where Ascaris infection is endemic, cross-reactions may be mostly removed by previously absorbing each serum sample with Ascaris suum antigens.¹⁹ The use of A. suum antigens for
both pure and applied work on toxocariasis is justified because of the close homology between *A. suum* and *A. lumbricoides* antigens 11.

Few reports have been published in Brazil in order to characterize the different forms of toxocariasis but most of them apply serological diagnosis by ELISA using TES antigens and pre-absorption with *Ascaris* antigens 12,13.

In the present study, in order to improve indirect ELISA specificity performed for toxocariasis diagnosis, we evaluated the specificity and sensitivity of an indirect antibody competition ELISA based upon TES antigen using sera from patients with clinical signs of toxocariasis and sera from mice experimentally infected with *T. canis* eggs. The IgG fraction of a rabbit hyperimmune serum against TES was employed as the competition antibody after absorption with *A. suum* extract immobilized in glutaraldehyde polymer.

**MATERIALS AND METHODS**

**Toxocara canis** excretory-secretory antigens (TES)

Female adults of *T. canis* were collected from feces of puppies treated with piperazine adipate (100mg/Kg). After collection worms were washed in saline and fertile eggs were obtained from the uteri of gravid female and left to embryonate for 30 days at 28°C in formalin solution. Secretory-excretory antigen was prepared by the method of de SAVIGNY 19 with some modifications. Briefly, eggs were hatched mechanically to allow the migration of larvae in a Baerman apparatus and were maintained in Eagle’s Minimal Essential Medium for no more than 5 months. Culture medium was collected every seven days, then concentrated by ultrafiltration (PM10, Amicon, Lexington, USA) and protein content was estimated by a commercial BCA method kit (Pierce-USA).

**Ascaris suum** adult stage extract

Female worms were collected from swine intestine and washed in saline. Adult extract was prepared as described by CAMARGO et al. 5 and protein concentration was estimated by a commercial BCA method kit (Pierce-USA).

**Extracts of other parasites**

One ml of *Strongyloides venezuelensis* filariform larvae were collected from experimentally infected rats feces. After addition of 0.02M phenylmethyl sulphonyl fluoride (PMSF, Sigma) and 0.05M EDTA they were disrupted in an ice-bath using ultrasound (Heat Systems, Microson XL). The homogenate was gently stirred for 2h at 4°C and centrifuged at 1000 x g for 30 min. at 4°C. The supernatant was stored at -20°C.

*Schistosoma mansoni* adult stage of the BH strain were obtained from experimentally infected hamsters and prepared according to PINTO et al. 11.

Metacestode extract of the ORF strain of *Taenia crassiceps* (*Cysticercus longicollis*) was prepared as described by VAZ 21.

**Antiserum**

Two rabbits were immunized by injecting intradermically 300μg of TES antigen in Freund’s complete adjuvant in multiple sites on the back skin. Six weeks later rabbits were subcutaneously injected with 300μg of TES in Freund’s incomplete adjuvant followed 1 month later by another subcutaneous injection of 300μg of TES. Blood was taken 3 weeks later by intracardiac puncture.

**Absorption with Ascaris suum extract**

Antiserum from immunized rabbits were precipitated by 40% ammonium sulfate according to HEBERT et al. 19 IgG fraction was purified by DEAE-cellulose as described by CORTHIER et al. 3 IgG fraction was then absorbed twice with *A. suum* (AS) extract polymerized with glutaraldehyde as described by TERNYNCK & AVRAMEAS 20. This methodology was chosen in order to avoid the presence of AS soluble antigens. Briefly, 50 mg of AS extract and 400 mg of bovine serum albumin were polymerized with 2 ml of glutaraldehyde 2.5%. After 3 h the polymer was centrifuged at 6,000 x g for 15 min. at 4°C, washed 3 times with PBS and 0.2M glycine-HCl buffer, pH 2.8 was added to eliminate proteins that might not have been polymerized. For absorption 1 ml of rabbit specific IgG was added to the polymer and left at room temperature for 3 h. The polymer was removed by centrifugation at 30,000 x g for 20 min. at 4°C.

**Experimental infection**

CH3 Rockfeller mice were infected by stomach tube with 1,100 embryonated *T. canis* eggs. Mice were bled from retroorbital plexus on the day of infection and on days 10, 20, 30 and 43 to 48 when they were euthanased to have their brain tissue examined for larvae presence.

**Human sera**

Sixty serum samples from healthy individuals aged from 20 to 35 and 440 serum samples from patients with at least one of the common features of toxocariasis (eosinophilia, hepatomegaly and/or pulmonary symptoms, hypergammaglobulinemia, geophagia) were evaluated. Of these, 241 samples were positive by indirect ELISA performed at Adolfo Lutz Institute (IEAL) as described by CAMARGO et al. 19 and 199 samples were negative. IEAL requires pre-absorption of each serum sample with *A. suum* extract.

**Mouse sera**

Serum samples of twenty-eight mice were collected before and 10, 20, 30 and 43 to 48 days after experimental infection with *T. canis* eggs. Sera from 2 mice not infected with *T. canis* eggs were used as control.

**Indirect ELISA**

The assay was performed to evaluated rabbit IgG reactivity before and after absorption with *A. suum* extract employing different parasites antigens. Procedures were as described below in the absence of competition antibody. Antigens concentration were 1.2 and 0.6μg/ml for extracts TES and AS; 1:2 and 3 μg/ml for extracts of *S. venezuelensis*, *S. mansoni* and *C. longicollis*. 

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Indirect antibody competition ELISA (IACE)

Maxisorp® microplates (Nunc, Denmark) were employed and all immunoreagents were assayed in 100 μl volumes. Between all steps of the reaction microplates were washed 3 x 5 min. with PBS-T (0.01M phosphate-buffer saline, pH 7.2 containing 0.05% Tween 80). For microplate coating TES antigen was diluted in 0.05M carbonate-bicarbonate buffer pH 9.6 and incubated overnight at 4°C. Testing sera, specific rabbit anti-TES IgG fraction used as competition antibody as well as conjugate (goat IgG anti rabbit IgG labelled to peroxidase, Sigma A-4914) were diluted in PBS-T containing 5% skimmed milk and incubated for 1h at 37°C. Five minutes after the addition of the substrate (orthophenylene diamine 0.04% in 0.1M citrate-phosphate buffer pH 5.0 and 0.03% H2O2), the enzymatic reaction was stopped with 50 μl of 1N HCl. Absorbance was read at 492 nm using an ELISA reader (Multiskan Plus® - Labsystems, Finland). Data were expressed as IACE values (IACEV), as follows:

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\text{IACEV} = 1 - \frac{\text{μA}}{\text{μIgG}}
\]

where μA represents the mean absorbance of the duplicate of the test sample and μIgG represents the mean absorbance of the quadruplicate of the rabbit IgG.

A checkerboard titration employing sera from healthy people and from patients with clinical signs of toxocariasis was performed to determine the best antigen protein concentration for coating the microplates (0.6 and 1.2 μg/ml) and the best dilution of sera (1:20, 1:40 and 1:80), rabbit IgG (1:1250, 1:2500 and 1:5000) and conjugate (1:1250, 1:2500 and 1:5000).

Cut-off value was calculated by the arithmetic mean of the absorbance of the 60 serum samples from the healthy population plus 2 standard deviation.

Repeatability of IACE was measured by repetition of a serum sample from a patient with clinical signs of toxocariasis in 25 plates.

RESULTS

Figure 1 shows results of rabbit anti-TES IgG reactivity by indirect ELISA before and after absorption with A. suum extract (AS) employing different parasite antigens. Cross-reactivity with AS extract can be seen which decreases after absorption but reactivity to TES antigens still remains. Figure 1 also shows absence of cross-reactivity between specific IgG and extracts of S. mansoni or C. longicollis but certain degree of cross-reactivity with S. venezuelensis extract can be seen which is absent after absorption.

All mice experimentally infected showed live larvae in brain tissue after 43 to 48 days of infection while negative control mice did not.

After checkerboard titration the best results of indirect antibody competition ELISA (IACE) were obtained by employing TES antigen at 0.6 μg/ml, sera diluted at 1:40, rabbit IgG at 1:2500 and conjugate at 1:2500.

Sensitivity and specificity of IACE were first evaluated in 28 serum sample of mice experimentally infected with T. canis eggs. Adopting cut-off value as arithmetic mean of absorbance of mice sera before experimental infection plus 2 standard deviation,
Using IEAL as the reference test sensitivity was 60.2%, specificity was 98% and concordance was 77.3%.

Repeatability of ICAE was evaluated by the inter-assay variation coefficient which was 2.4%.

**DISCUSSION**

Although some authors references have not related cross-reactivity of TES antigens to other parasites antigens this cross-reactivity has been shown by different means and authors\(^{1,14,15}\). A wide variety of helminths like *Ancylostoma*, *Strongyloides*, *Schistosoma*, and *Taenia* can show immunological cross-reactivity with *Toxocara* but the commonest is cross-reactivity between *Ascaris* and *Toxocara*\(^{10,16}\).

A major problem of using sera to study cross-reactivity between *Toxocara* and *Ascaris* is the uncertain infection histories so we preferred to evaluate the reactivity of specific rabbit IgG raised against TES antigens with other parasite extracts by indirect ELISA. Before absorption, this IgG showed cross-reactivity mostly to *Ascaris suum* extract (AS) although certain degree of cross-reactivity with *S. venezuelensis* extract could be seen which is absent after absorption (Figure 1), as observed before by KENNEDY et al.\(^{17}\). Cross-reactivity with AS extract decreases after absorption but reactivity to TES antigens still remains, showing the existence of cross-reactivity and necessity of absorption.

Difficulties on obtaining human sera from patients truly positive to toxocariasis lead us to perform a experimental infection of mice with *T. canis* infective eggs. Infection was confirmed by live larvae observed in brain tissue after 43 to 48 days of infection in all mice experimentally infected. Although in natural condition parthenic hosts are exposed to small doses of infective *T. canis* eggs, we used high doses of infective eggs to assure humoral immune response. IgG anti-TES antigens could be detected by indirect antibody competition ELISA (IACE) as early as 10\(^{th}\) day post-infection in 19 of 28 mice but detection of IgG anti-TES in all experimentally infected mice was possible only by the 20\(^{th}\) day. KAYES et al.\(^{12}\) have showed that antibody level peaks at 14 days post inoculation and remain more or less constant thereafter. We did not evaluate persistence of anti-*Toxocara* antibodies but it remained for at least 48 days post-infection.

Indirect ELISA for VLM performed at Adolfo Lutz Institute (IEAL) according to CAMARGO et al.\(^{1}\) requires pre-absorption of each serum sample with *A. suum* extract and had a sensitivity of 95.4% and specificity of 89.3%. In our study, although we had to immunize rabbits to obtain specific IgG and we still had to absorb it with *A. suum* extract, it has the advantage that absorption is performed in a polymer that will not remain soluble antigens interfere in the test besides the fact that it will be done only for 2 times, not for each serum sample. Lack of relative sensitivity of IACE (60.2%) here presented compared with IEAL, can be explained by the fact that IACE is based on a more specific competition between antibodies from sera of patients and rabbit anti-TES IgG. Another reason to explain the lack of relative sensitivity of IACE could be insufficient absorption of antibodies from patients with AS extract while performing IEAL.
Data of specificity and sensitivity of diagnostic tests performed by most authors are based on serum samples from patients with toxocariasis signs and from patients with other parasitic infection. In this matter our study gives different results because we tested the reactivity of some parasite extracts to the specific rabbit IgG anti-TES besides the fact that it was performed with experimentally infected sera truly positive to VLM.

IACE here presented has not yet been performed with sera of patients with covert or ocular toxocariasis but we think this new test will improve the diagnostic capacity of laboratories for the classical form of toxocariasis because its specificity is higher than the indirect ELISA so far performed in Sao Paulo State-Brazil. Therefore, cross-reactivity with Ascaris should not misdiagnose toxocariasis.

Besides good specificity and repeatability of IACE one advantage of this system is that it can be used in experimental studies of different animal species since the conjugate will always be against the rabbit IgG used as competition antibody.

RESUMO

Diagnóstico sorológico da toxocariase através do método de ELISA indireto de competição

A toxocariase, uma emergente zoonose, é uma síndrome clínica decorrente da infecção humana por larvas de *Toxocara canis* e *Toxocara cati*, parasitas intestinais de cães e gatos, respectivamente.

A dificuldade de detecção das larvas nos tecidos e em fluidos, e a inespecificidade dos sinais clínicos tornam os testes sorológicos os meios diagnósticos mais adequados. Os antígenos excretores-secretóres de *T. canis* empregados nos testes sorológicos, embora tenham contribuído para melhorar a especificidade destes, apresentam reações cruzadas com diversas parasitas, particularmente com *Ascaris* sp.

Com o intuito de aumentar a especificidade do diagnóstico sorológico da toxocariase que rotineiramente é feito através do método de ELISA indireto, no presente trabalho desenvolveu-se um método de ELISA indireto de competição (EIC), utilizando IgG anti-TES produzida em coelhos e absorvida com um polímero de extrato antitérgico de adultos de *A. suum*.

A avaliação da sensibilidade e especificidade do EIC proposto foi inicialmente feita em camundongos experimentalmente infectados com *T. canis*. Adotando-se ponto de corte estabelecido nesta população previamente à infecção, os valores de sensibilidade e especificidade foram de 100% a partir de 20 dias pós-inoculação.

Para a população humana, adotou-se um ponto de corte estabelecido a partir de 60 amostras de soro de indivíduos saudáveis e sem suspeita clínica de LMV. A avaliação do EIC foi realizada em amostras de soro de pacientes com suspeita clínica da doença e tomando como referência a classificação previamente feita pelo ELISA indireto realizado pelo Instituto Adolfo Lutz-SP. O EIC apresentou valores de sensibilidade relativa de 60,2 %, especificidade relativa de 98% e concordância de 77,3%.

Na avaliação da repetibilidade do EIC obteve-se coeficiente de variação inter-reações de 2,4%.

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