Evaluation by ELISA of Anisakis simplex Larval Antigen Purified by Affinity Chromatography

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In order to improve the specificity and sensitivity of the techniques for the human anisakidosis diagnosis, a method of affinity chromatography for the purification of species-specific antigens from Anisakis simplex third-stage larvae (L3) has been developed. New Zealand rabbits were immunized with A. simplex or Ascaris suum antigens or inoculated with Toxocara canis embryonated eggs. The IgG specific antibodies were isolated by means of protein A-Sepharose CL-4B beads columns. IgG anti-A. simplex and -A. suum were coupled to CNBr-activated Sepharose 4B. For the purification of the larval A. simplex antigens, these were loaded into the anti-A. simplex column and bound antigens eluted. For the elimination of the epitopes responsible for the cross-reactions, the A. simplex specific proteins were loaded into the anti-A. suum column. To prove the specificity of the isolated proteins, immunochromical analyses by polyacrylamide gel electrophoresis were carried out. Further, we studied the different responses by ELISA to the different antigenic preparations of A. simplex used, observing their capability of discriminating among the different antisera raised in rabbits (anti-A. simplex, anti-A. suum, anti-T. canis). The discriminatory capability with the anti-T. canis antisera was good using the larval A. simplex crude extract (CE) antigen. When larval A. simplex CE antigen was loaded into a CNBr-activated Sepharose 4B coupled to IgG from rabbits immunized with A. simplex CE antigen, its capability for discriminate between A. simplex and A. suum was improved, increasing in the case of T. canis. The best results were obtained using larval A. simplex CE antigen loaded into a CNBr-activated Sepharose 4B coupled to IgG from rabbits immunized with adult A. suum CE antigen. When we compared the different serum dilution and antigenic concentration, we selected the working serum dilution of 1/400 and 1 µg/ml of antigenic concentration.

Key words: Anisakis simplex - antigen purification - affinity chromatography - ELISA - rabbit antisera

The diagnosis of human anisakidosis can be made from the history of eating raw fish, and a final diagnosis is not easy even with the development of gastrofiberscopical examination of patients (Ishikura et al. 1993), and in most cases it is only correctly made after post-operative pathological examination of the affected part of the gut. In intestinal anisakiasis post-operative diagnosis is made by identification of whole larvae or cross sections of larvae in histological sections (Smith & Wootten 1978). Consequently, anisakidosis has often been misdiagnosed as appendicitis, acute abdomen, gastric tumour or cancer, ileitis, cholecystitis, diverticulitis, tuberculous peritonitis, cancer of the pancreas, or Crohn’s disease (Sakanari & McKerrow 1989). Immunodiagnostic methods have been developed, but several cross-reactivities have appeared. The earliest method for immunodiagnosis of anisakiasis was a complement fixation test (Daniels 1962). A immunofluorescence test was more sensitive than complement fixation, but cross-reactions with sera from toxocariasis patients can occur (Ruitenberg 1970). Immunelectrophoresis on starch demonstrated cross-reactivity to antigens from both Toxocara and Ascaris (Suzuki 1968). Iglesias et al. (1996) observed that the murine sera raised by immunization with Ascaris suum, Toxocara canis and Hysterophylum aduncum somatic antigens reacted even more strongly with the Anisakis simplex antigen than with the corresponding homologous by ELISA and immunoblotting. In fact, extensive homology between both somatic and excretory-secretory antigens of A. simplex and other ascaridoid nematodes mainly A. suum, A. lumbricoides and T. canis has been reported (Kennedy et al. 1988).

In this work the specificity and sensitivity of A. simplex antigens prepared by affinity chromatography were assayed by ELISA using sera from immunized animals.

MATERIALS AND METHODS

Parasites - A. simplex L3 were picked up manually from the viscera, flesh and body cavities of naturally infected blue whiting (Micromesistius poutassou) and exhaustively washed in water (Perteguer & Cuéllar 1998). A. suum and T. canis were obtained from natural infections of swine or dogs, respectively. T. canis eggs were obtained by dissection of gravid females and, after their purification, they were embryonated under sterile conditions and preserved at 4ºC until use (Guillén et al. 1986).

Antigens - For preparation of the crude extract (CE) of A. simplex (A. simplex CE antigen), L3 were placed at 4ºC in PBS. This material was homogenized in a hand-oper-
ated glass tissue grinder at 4°C, followed by sonication for 10 s with a Virstonic 5 (Virtis, NY, USA) set at 70% output power. The homogenate was extracted in PBS at 4°C overnight and, subsequently, delipidized with n-hexane and then centrifuged at 8,497 g for 30 min at 4°C (Biofuge 17RS: Heraeus Sepatech, Gmb, Osterode, Denmark). The supernatant was dialysed overnight at 4°C in PBS. Protein content of the extract was estimated and the extract was frozen at -20°C until used (Verteguer & Cuéllar 1998). CE antigen from *A. suum* adults (*A. suum* CE antigen) was obtained using a modification of the Welch et al. (1983) method by homogenization and extraction in PBS at 4°C overnight (instead of ultrasonic burst). Its protein content was estimated by the Bradford (1976) method, and then the antigen frozen at -20°C until use (Águila et al. 1987, Cuéllar et al. 1990). *T. canis* eggs were obtained by dissection of gravid females, resulting from a natural infection. After purification, they were embryonated under sterile conditions at 37°C and preserved at 4°C until use (Guillén et al. 1986, Fenoy et al. 1987, 1988).

Hyperimmune sera - New Zealand rabbits of about 3 kg body weight were immunized with larval *A. simplex* or adult *A. suum* CE antigen as multiple doses of 3 ml of antigen (1,000 µg/ml in final volume) in Freund’s Complete Adjuvant (FCA) intramuscularly given weekly for 3 weeks. Other New Zealand rabbits were inoculated with multiple doses of 2,000 embryonated eggs of *T. canis* weekly, during 4 weeks, by oral administration with gastric tubing. Animals were bled weekly postimmunization (p.i.) after the 1st inoculation (week 0) (Cuéllar et al. 1990, García et al. 1996).

Purification of *A. simplex* CE products by affinity chromatography - Protein A-Sepharose CL-4B beads (Pharmacia Biotech) columns were prepared according to the manufacturer’s instructions. Rabbit anti-*A. simplex*, or -*A. suum* antibodies, in sample buffer [0.05 M Tris, 0.5 M NaCl (pH 8.0)] were loaded into the columns. Fractions of 1 ml were then collected. Unbound immunoglobulins were washed with washing buffer (0.05 M Tris, 0.5 M NaCl). Bound immunoglobulins then were eluted with glycine buffer (0.2 M glycine, 0.5 M NaCl, pH 2.8). Fractions were collected onto 100 µl of collection buffer (Tris-base 1 M, pH 8.5) and read on a spectrophotometer at A280 for calculating IgG concentration. A column was prepared with Protein A affinity isolated IgG anti-*A. simplex*, at a concentration of 5 mg/ml in NaHCO3 0.1 M with NaCl 0.5 M (pH 8.5) coupled to CNBr-activated Sepharose 4B according to the manufacturer’s instructions (Pharmacia Biotech). *A. simplex* CE antigen in sample buffer was loaded into the column and incubated 3 h at room temperature. Fractions of 1 ml were then collected. Unbound antigens were washed with washing buffer and bound antigens were then eluted with glycine buffer followed by 50 mM diethyleneam in saline, pH 11.5 and collected into glycine to neutralise the eluted fractions. Fractions were read at A280. This antigen was named as *A. simplex* PAK antigen. The same procedure was carried out using columns prepared with rabbit IgG anti-*A. suum*, obtaining the *A. simplex* PAS antigen and the *A. simplex* EAS antigen.

SDS-PAGE - Sodium dodecyl sulphate-polyacrilamide gel electrophoresis (SDS-PAGE) was carried out as described by Laemmli (1970) and revised by Hames (1986) using a Mini Protean® II cell (Bio Rad). The gels consisted of a 4% stacking gel and a 5-20% gradient separating gel. Samples were dissolved in a sample buffer (50 mM Tris-HCl buffer, pH 8.6, containing 2% SDS, 20% glycerol and 0.02% bromphenol blue) diluted 1:1 in electrode buffer (25 mM Tris, 192 mM glycine, pH 8.3), containing 1% SDS. Electrophoresis was performed for 2 h at a constant 100 V in Tris-glycine electrode buffer (see above). Broad range molecular weight markers (6500-205,000 or 7200-209,000 Da; Bio Rad) were incorporated into each electrophoretic run. Gels were stained with silver.

**Determination of specific antibody levels** - The 96 well microtitre plates (Nunc-Immu-no Plate PolySorp™) were sensitized overnight at 4°C by the addition of 100 µl well of antigen diluted at 0.5 or 1 µg/ml in a 0.1 M carbonate buffer, pH 9.6. After washing three times with 0.05% PBS-Tween 20 (PBS-Tween), wells were blocked by the addition of 200 µl per well of 0.1% BSA in PBS, for 1 h at 37°C. After washing, 100 ml of duplicate dilutions of rabbit sera at 1/100, 1/200 or 1/400 in PBS-Tween, containing 0.1% BSA were added and incubated at 37°C for 2 h. Once the plates were washed, 100 µl per well of affinity isolated, peroxidase conjugated, goat anti-rabbit IgG (Caltag Laboratories, San Francisco, CA, USA), at the appropriate dilution in PBS-Tween, 0.1% BSA, were incubated for 1 h at 37°C. After adding substrated (phosphate-citrate buffer, containing 0.04% H2O2 and 0.04% o-phenylenediamine) the reaction was stopped with 3N sulfuric acid and the plates were read at 490 nm (García et al. 1996). Results were expressed as O.D.p-O.D.c indexes by subtracting the mean optical density (O.D.) of the control from the mean O.D. of the test sera once the non-specific reaction with the BSA used in the blocking was subtracted.

**RESULTS**

The electrophoresis comparative study between the *A. simplex* CE antigen and the one represented by the mixed fractions eluted with glycine buffer followed by diethyleneam from the Sepharose 4B-rabbit IgG anti-*A. simplex* column (*A. simplex* PAK antigen) indicates that both preparations are very similar (Fig. 1).

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**Fig. 1:** SDS-PAGE, lane 1: *Anisakis simplex* EAS antigen; lane 2: *A. simplex* PAS antigen; lane 3: *A. simplex* PAK antigen; lane 4: *A. simplex* CE antigen. See arrows for *A. simplex* specific proteins.
Likewise, *A. simplex* PAK antigen was loaded into the column prepared with anti-*A. suum* rabbit IgG. Unbound (*A. simplex* free *A. suum* antigens) antigens were washed and bound (cross-reacting *A. suum* antigens) antigens eluted. After passing across the anti-*Ascaris* column, the proteins of *A. simplex* PAK antigen were maintained but increased in intensity the bands of 40 and 14 kDa. With *A. simplex* PAK antigen a doublet around 60 kDa appeared. With the *A. simplex* PAS antigen, 60 kDa was maintained with a high intensity. Bright bands of high molecular weight of about 209 and 150 kDa were seen. In Fig. 1 electrophoretic patterns obtained with the above mentioned antigens by SDS-PAGE are shown. In the *A. simplex* PAK the previously mentioned doublet is shown. In the *A. simplex* PAS antigen the protein of 120 kDa increased in concentration, as well as, a doublet of about 66-45 kDa, the protein of 40 kDa was maintained but increased in intensity the protein of 25 kDa, and a doublet between 21.5 and 14 kDa. In the *A. simplex* EAS antigen the protein of 120 kDa were observed at 97-66 kDa. Likewise, very bright proteins of about 35 kDa and 7 kDa were present.

Further, the ELISA method has been carried out in order to determine the IgG levels in sera from rabbits experimentally immunized with larval *A. simplex* or adult *A. suum* CE antigen as multiple doses in FCA intramuscularly or multiple doses of *T. canis* embryonated eggs. After they were bled weekly p.i., the sera with strong reactions were selected by ELISA against their homologous antigens. The selected rabbit antisera were tested at the 1/100, 1/200 or 1/400 dilutions and the antigens at 1 or 0.5 µg/mL. The antigenic preparations used were as follows: larval *A. simplex* CE antigen (*A. simplex* CE antigen); *A. simplex* CE antigen after loading into a CNBr-activated Sepharose 4B coupled to IgG from rabbits immunized with *A. simplex* CE antigen (*A. simplex* PAK antigen); *A. simplex* PAK antigen after loading into a CNBr-activated Sepharose 4B coupled to IgG from rabbits immunized with adult *A. suum* CE antigen (*A. simplex* PAS antigen); *A. simplex* PAK antigen eluted from the anti- *A. suum* column (*A. simplex* EAS antigen) and adult *A. suum* CE antigen (*A. simplex* CE antigen). Results were expressed as O.D.p-O.D.c indexes by subtracting the mean optical density of the test sera once the non-specific reaction with the BSA used in the blocking was subtracted (Figs 2, 3).

When the capability of the different antigens to discriminate among the different antisera raised in rabbits was studied, the minimal responses were observed using sera from rabbits inoculated per os with *T. canis* embryonated eggs against all the antigenic preparations used. The highest responses were obtained when the sera from rabbits immunized with larval *A. simplex* or adult *A. suum* CE were tested against their corresponding homologous and heterologous CE from *A. simplex* or *A. suum*. In order to study the different antigenic preparations of *A. simplex* used we have obtained an index as the ratio between the optical density resulting from the anti-*A. simplex* antiserum and the optical density of the anti-*A. suum* or anti-*T. canis* heterologous antisera once their corresponding non-specific reaction with the BSA used in the postcoating were subtracted, as well as, the optical density of the negative controls. In the case of the *A. simplex* CE antigen those indexes were between 0.7 (serum dilution 1/100-antigen concentration 0.5 µg/ml) and 1.7 (serum dilution 1/200 and 1/400-antigen concentration 0.5 µg/ml) using anti-*A. suum* antisera. On the contrary, the discriminatory capability with anti-*T. canis* antisera was good with indexes situated between 5.1 (serum dilution 1/100-antigen concentration 1 µg/ml) and 14 (serum dilution 1/200-antigen concentration 0.5 µg/ml). When *A. simplex* PAK antigen was tested, its capability for discriminate between *A. simplex* and *A. suum* was improved with indexes situated between 1.7
(serum dilution 1/100-antigen concentration 1 µg/ml) and 3 (serum dilution 1/400-antigen concentration 1 µg/ml), increasing in the case of T. canis from 4.3 (serum dilution 1/100-antigen concentration 1 µg/ml) to 103.3 (serum dilution 1/200-antigen concentration 1 µg/ml).

Using A. simplex PAS antigen, with anti-A. suum antisera, we obtained values from 1.8 (serum dilution 1/100-antigen concentration 1 µg/ml) to 3.3 (serum dilution 1/400-antigen concentration 1 µg/ml; serum dilution 1/100-antigen concentration 0.5 µg/ml). In the case of T. canis with the A. simplex PAS antigen the values were from 4.6 (serum dilution 1/100-antigen concentration 1 µg/ml) to 571 (serum dilution 1/200-antigen concentration 1 µg/ml).

In summary, to eliminate the cross-reactivity between A. simplex and T. canis and A. suum it is necessary a sequential step affinity chromatography using IgG from rabbits immunized with A. simplex and A. suum.

**DISCUSSION**

The aim of this work was to assay by ELISA the specificity and sensitivity of larval A. simplex CE antigen purified by means of affinity chromatography using sera from rabbits experimentally immunized.

The first purification step consisted in binding the larval A. simplex antigen to homologous antibodies from rabbits experimentally immunized with this antigen and eluted from the column in order to eliminate the major cross-reactivity molecules. The SDS-PAGE patterns of A. simplex CE antigen showed proteins of 205, 120, 66-45, 40, 31-21 and 14 kDa. In the A. simplex PAK antigen we observed the same proteins but several were in different proportions. The molecular mass of ABA-1, the Ascaris nematode polyprotein, is controversial, it has been previously estimated at 14,000 Da (Christie et al. 1990), but mass spectrometry analysis indicated that there were five componentes of similar size, with the major species being 14,643.2 ± 1.4 Da with a high degree of similarity amongst ascaridid parasites (Christie et al. 1993). Yahiro et al. (1998) cloned the cDNA of TBA-1, the nematode polyprotein allergen of T. canis, and on the basis of the amino acid sequence found it to be most similar to ABA-1. They observed a transient TBA-1 IgG antibody response during the infection that could explain the failure of Kennedy et al. (1989) in the aim of finding anti-TBA-1 antibodies in animals infected with T. canis for a prolonged period. Also different forms of ABA-1 have been reported by Kato and Komatsu (1996), which could explain the different antibody recognition such, also two different forms of TBA-1 may be expressed in a stage specific manner. Also Kennedy et al. (1988) observed evidence that a Mr 14,000 component of A. simplex has a homologue in A. suum, A. lumbricoides and T. canis, but did not elicit an antibody response in anisakiasis.

Zarnowska and Jastrzebska (1994) by SDS-PAGE of larval ES products from T. canis showed polypeptides of molecular weights ranging from 19 to 200 kDa. However, an additional polypeptide, not observed on stained gels, and resolving at 14 kDa, was detected by immunoblotting. Sera from patients with A. lumbricoides recognized polypeptides of 74, 75 and 160 kDa. Iglesias et al. (1996) confirmed by immunoblotting the high degree of cross-reactivity between the somatic antigens of A. simplex and somatic antigens of the ascaridoids A. suum, T. canis and H. aduncum, although several A. simplex components in the 11-18 kDa range were only recognized by sera from mice infected with A. simplex. Tanaka et al. (1983) developed a radioimmunoassay for A. suum protein and observed that T. canis had a high concentration of a substance partially cross-reactive with A. suum protein. Also, small amounts of substance cross-reacting with A. suum protein was also exhibited by Anisakis larvae. High concentrations of A. suum protein were observed in sera from patients with ascariasis (64.5 ± 18.8 ng/ml), anisakiasis

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**Fig. 3: ELISA IgG responses expressed as O.D.p-O.D.c indexes with serum samples. A: at 1/100; B: at 1/200 and C: at 1/400, of rabbits respectively immunized with Anisakis simplex crude extract (CE) antigen, Ascaris suum CE antigen, or Toxocara canis embryonated eggs, against the following antigens diluted at 0.5 µg/ml: larval A. simplex CE antigen (A. simplex CE antigen); A. simplex CE antigen treated in a IgG anti-A. simplex CE column (A. simplex PAK antigen); A. simplex PAK antigen treated in a IgG immunized anti-A. suum CE column (A. simplex PAS antigen); A. simplex PAK antigen eluted from the A. suum column (A. simplex EAS antigen); adult A. suum antigen (A. suum CE antigen).**
(75.2 ± 28.0 ng/ml) and toxocariasis (78.4 ± 31.3 ng/ml). Nunes et al. (1997) detected at least one band with molecular weight around 55-66 kDa that seems to be responsible for the cross-reactivity between T. canis and A. suum once it disappears when previous absorption of serum samples with A. suum antigens was performed. Kennedy et al. (1989) observed, using radioimmunoprecipitation and SDS-PAGE, that there was a significant antigenic similarity between the antigens of A. suum and T. canis. Among the cross-reactive components, these authors found a 14 kDa internal protein which has a homologue in the two parasites, observing that was the subject of an IgG antibody response in Ascaris infection, but there was not measurable response to it in toxocariasis. McWilliams et al. (1987) observed that A. suum cross-reacted allergically with T. canis and that the cross-reacting allergens were predominantly of high molecular weight.

A. simplex PAK antigen was purified from the anti-A. suum rabbit IgG column. In the A. simplex PAS antigen the proteins of A. simplex PAK antigen were maintained but increased in intensity the bands of 40 and 14 kDa. In the A. simplex PAS antigen, the proteins were of 120, 66-45, 40, 31-21 and 14 kDa with elevated concentrations of these specific proteins that those observed in the non-purified samples except in the case of the 14 kDa protein. The eluted fractions which contained the cross-reacting antigens with A. suum showed bands in the ranges of molecular weight situated at 40 and 14 kDa, observing in the latter higher concentrations than in the specific antigen. In our experimental conditions, the minimal responses were observed by ELISA using sera from rabbits inoculated per os with T. canis embryonated eggs against all the antigenic preparations used. Conversely, the highest responses were obtained when the sera from rabbits immunized with larval A. simplex or adult A. suum CE were tested against their corresponding homologous and heterologous CE from A. simplex or A. suum. Further, we studied the different responses to the different antigenic preparations of A. simplex used, observing their capability of discriminating among the different antisera raised in rabbits. The discriminatory capability with the anti-T. canis antisera was good using the larval A. simplex CE antigen. When A. simplex PAK antigen was used, its capability for discriminate between A. simplex and A. suum was improved, increasing in the case of T. canis. The best results were obtained using A. simplex PAS antigen. When we compared the different serum dilution and antigenic concentration, we selected the working serum dilution of 1/400 and 1 μg/ml of antigenic concentration. These conditions insure the maximal differences among the different sera using small quantities of the samples. Works are in progress to study the specificity and sensitivity of A. simplex antigens prepared by affinity chromatography using sera from human patients diagnosed of anisakidosis, ascariosis and visceral larva migrans by ELISA and western-blot techniques.

REFERENCES

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