DIAGNOSTIC TESTS FOR AMOEBIC LIVER ABSCESS: COMPARISON OF ENZYME-LINKED IMMUNOSORBENT ASSAY (ELISA) AND COUNTERIMMUNOELECTROPHORESIS (CIE)

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The liver abscess is the most frequent extraintestinal complication of intestinal amoebiasis: its diagnosis is suggested by the clinical picture but it must be confirmed by paraclinical tests. The most stringent diagnosis requires identification of E. histolytica. But this is possible only in a few cases. Serological tests greatly improve the diagnosis of this severe complication of amoebiasis. We compared the Enzyme Lided Immunosorbent Assay and the Counterimmunoeletrophoresis techniques. Both techniques were used to detect amoebic antibodies in 50 control patients, 30 patients with liver abscess and 30 patients with intestinal amoebiasis. All the sera from control patients gave negative results in both techniques. When analysing the sera from patients with intestinal amoebiasis, 10% of them were positive by ELISA but not by CIE. The sera of patients with liver abscess, we found that 90% were positive by the ELISA method and 66.6% by the CIE technique. In patients with amoebic liver abscess, the results showed that the ELISA was more sensitive than the CIE, as it presented a higher sensitivity (100%) than that of the CIE technique (66%).

Key-words: Amoebic liver abscess. Diagnosis. ELISA. CIE.

Amoebiasis is an infection caused by Entamoeba histolytica, an intestinal protozoan pathogenic for humans17. It is commonly found in tropical countries, specially in underdeveloped ones where sanitation and hygiene are deficient.

It is believed that 2 to 50% of the world's population and 12.1% of Colombia population according to the second Colombian National Morbidity Survey is infected by Entamoeba histolytica. The parasite colonises the intestinal tract and form here it spreads to other organs, depending on its invasive capacities8.

Amoebic infection is considered to be third most important cause of parasite-related deaths in the world, after malaria and schistosomiasis18. The most frequent extraintestinal complication of amoebic infection is the invasion of the liver by the parasite, a condition known as amoebic liver abscess.

The diagnosis of amoebic abscess is suggested by clinical features, but further paraclinical tests are required in order to identify the lesion in the liver: Ultrasonography, Computarised Tomography, Isotopic Scan, Magnetic Resonance Imaging, diagnostic aspiration, liver biopsy and surgical exploring are employed. Liver biopsy and diagnostic aspiration are, however, the methods that allow identification of E. histolytica in the necrotic material of the abscess or in liver tissue10. This method is the most specific of all and provides a differential diagnosis with pyogenic liver abscess, a condition which requires cultured identification of the causative bacteria1. Since the identification of E. histolytica is feasible only in very few case, serological techniques that could help in the diagnosis of amoebic liver abscess have been implemented. They all evaluate the individual's immunological response detecting circulating immune complexes or detecting antibodies in the serum by a diversity of serological tests7 18.

Our study evaluates the diagnostic sensitivity of two serological tests: Counterimmunoelectrophoresis (CIE) and Enzyme Linked Immunosorbtent Assay (ELISA)
MATERIAL AND METHODS

Study population. This population was composed by 110 patients which were distributed in 3 groups as follows:

a) 30 patients with symptomatic intestinal amoebiasis without acute dysentery, all confirmed by the presence of *E. histolytica* cysts or trophozoites in microscopic stool examination.

b) 50 control patients, asymptomatic presumably healthy persons; with no recent or past medical history of amoebic infection in the last 2 years, and 3 negative stool examination on 3 different days.

c) 30 patients with suspected amoebic liver abscess as suggested by clinical features and ultrasonographic findings.

Blood samples were obtained from all patients in 10ml tubes with no anticoagulant, to obtain serum and this was stored at -20°C until serological testing.

CIE technique. This HM1 clone of *E. histolytica* strain (obtained from Instituto Nacional de Salud - INS, Bogotá, Colombia) was cultured in the axenic BIS-33 medium of Diamond, and then we obtained a purified antigen as described by Gandhi et al. The CIE test used purified antigen of *E. histolytica* placed in the right column wells and problem serum in the left column wells. Electrophoresis was run in barbital buffer (pH 8.6) for 45min in a chamber at a constant voltage of 110 to 140 volts and 4 to 8 milliampers. The slide was incubated at 37°C until completely dry. The presence of a precipitin line between the antigen and serum wells was interpreted as a positive reaction.

ELISA technique. The antigen was prepared from cultures of *E. histolytica* clone strain HM1, grown axenically in medium TYIS-33. Harvested trophozoites were washed 3 times with saline. Then we applied the method recommended by Gandhi et al. to obtain the antigen. The trophozoites were frozen once and thawed, sonicated for 5 minutes at 20 kilocycles. Finally centrifuged at 10,000 x g for 30 minutes at 4°C. We used the clear supernatant as antigen with a pure protein concentration of 6mg/ml. This antigen was distributed in 100 microliter aliquots to each well of the polystyrene microplates (Dynatech (R)), incubated overnight at 40°C in a moist chamber. The microplates was rinsed with Tween-20 in PBS for 3 minutes, three consecutive times; 200 microliter of casein (w/v) in PBS were added to each well and afterwards the microplate was incubated for 1 hour at 37°C in a moist chamber. The plate was rinsed again with PBS-Tween 20 before adding 100 microliters of serum diluted 1/100 in PBS-Tween 20; then incubated for 1 hour at 37°C and rinsed again. Next step, 100 microliters of conjugate anti-human IgG peroxidase (Sigma (R)) was used at a dilution of 1:400 in 0.1 M PBS, pH 7.2. The microplate was incubated at 37°C in a moist chamber for 1 hour. Finally, 100 microliters of substrate (0.4mg of o-phenylene diamine per ml of 0.1M phosphate citrate buffer, pH 5.0) were added to each well left to rest for 20 minutes in the dark. The enzyme reaction was stopped by the addition of 50 microliters of HCL 2.5N.

Readings were performed with a 492nm filter to determine 2 optical densities (O.D.) per patient and from these, an average of O.D. values were calculated: three negative and two positive control sera were additionally processed.

We are using as the Gold Standards the parasitological diagnosis of intestinal amoebiasis in the stools and ultrasonographic findings for amoebic liver abscess.

The average cut-off point of the ELISA technique was 0.156, the absorbance reading needed to differentiate between positive and negative samples was established by calculating the mean value plus S.D. (p < 0.05) of the absorbance values of samples of healthy patients, amoebiasis intestinal patient and those patients with amoebic liver abscess.

Determination of test parameters were determined using a 2 x 2 contingency table that included the following calculations: for sensitivity (S): a/a+c; a) true positives, is the number of patients with parasitological diagnosis of intestinal amoebiasis and positive serological test; b) false positives corresponded to healthy asymptomatic persons without parasites in the stools (control group) but positive serological test. For specificity (E): d/b+d; d) true negatives, is healthy persons without amoebiasis and negative serological test; and c) false negatives, the patients with intestinal amoebiasis and negative serological test. For positive predictive value (PPV): a/a+b and negative

predictive value (NPV): d/c+d. The calculation was applied for both ELISA and CIE tests.

RESULTS

From the 30 patients with intestinal amoebiasis, three presented positive ELISA test, but all of them were negative in the CIE test (Tables 1 and 2). None of the 50 sera from control patients gave positive results in the two tests used (Tables 1 and 2).

In patients with intestinal amoebiasis, the sensitivity of the ELISA technique was only 10% but its specificity was 100% (Table 3). The CIE test was never found to be positive in this clinical form of amoebiasis. Only 3 sera from patients with symptomatic intestinal amoebiasis without dysentery were positive by ELISA. The predictive values of positive test were 100% for ELISA and 0% for CIE. The predictive values of negative test was 65% for ELISA and 62.5% for CIE (Table 3).

In the group of patients with liver abscess, 27 (90%) out of 30 were positive by ELISA and 20 (66.6%) by CIE (Tables 1 and 2). The sensitivity for ELISA and CIE tests were 100% and 66% respectively (Table 3). The specificity and the predictive value of positive tests with both reactions were 100%. The predictive value of negative test was 100% for ELISA and 83% for CIE (Table3).

DISCUSSION

In this study, we confirmed the clinical suspicion of extraintestinal amoebiasus by serological tests. Indeed, all cases of amoebic liver abscess could be detected by the presence of antibodies against E. histolytica by

CIE and ELISA technique. Our results validate the use of these techniques as a tool in the diagnosis of invasive amoebiasis, e.g. amoebic liver abscess (ALA)\textsuperscript{11 18}, ELISA technique is very useful in determination of different serotypes of antibodies (IgG, IgM, IgE, IgA) in sera of patients infected by E. histolytica as was
shown by Hock et al. This method showed high sensitivity and specificity in those cases where the parasite was very invasive such as intestinal (dysentery) or liver amoebiasis. The ELISA was reliable compared with healthy controls.

Analyzing the ELISA results in the diagnosis of 30 patients with liver abscess, we found that only 3 patients were negative by this serological test. Reviewing these cases we concluded that two of the patients had liver abscesses secondary to stabbing injuries, which responded favorably to combined antimicrobial therapy (against gram positive, gram negative and anaerobic bacteria) and surgical drainage. The third patient did not respond to the anti-amoebic therapy, then a surgical drainage was performed and appropriate antibiotic treatment was indicated, since Staphylococcus aureus was isolated from the purulent material. Thus, the 3 patients who were negative by ELISA and CIE were also negative for amoebic liver abscess, and actually had a pyogenic liver abscess. On the other hand, 3 patients with intestinal amoebiasis and without liver abscess were positive by ELISA but negative by CIE. Although our study did not take in consideration the difference between dysenteric and acute form of intestinal amoebiasis, the seropositivity on ELISA test could indicate a tissue invasion with E. histolytica. In fact, most of our patients without clinical evidence of liver abscess had a chronic form of intestinal amoebiasis.

The specificity of ELISA testing in the diagnosis of amoebic liver abscess was supported by the fact that amoebic antibodies were absent in serum from healthy controls. However some troubles may arise when the optical densitics are in the limit of the cut-off (+1 SD). Longitudinal studies are needed to assess the antibody responses variability along the evolution of the disease. This could also allow a better therapeutic follow-up.

The CIE for detection of antiamoebic antibodies in the patients sera has been carried out and correlated with the routine diagnostic microscopic examination of stools and pus samples from patients with a characteristic clinical case. In the 30 patients with liver abscess evaluated by CIE test, 7 cases were negative. Those cases had clinical and ultrasonographic findings of liver abscess, of which just 3 were the patients mentioned above (non-amoebic liver abscess) and 4 cases were false-negative. These 4 patients did respond and improve after anti-amoebic treatment.

Our results are in agreement with of Bapat and Bhave, who found in 20 of 30 proved cases of ALA a seropositivity by CIE. Moreover, Samrejrongroj et al compared CIE and ELISA for E. histolytica antibodies determination in patients with amoebiasis and reported a sensitivity of 93.5% and 100% respectively. Sathar et al evaluated the ELISA test for serodiagnosis of amoebic liver abscess, and found the presence of IgG antibodies with a sensitivity of 99%. Similar results were obtained by Shetty et al. Finally, Hock et al, comparing both tests found ELISA more sensitive than CIE (97.4% vs 88.5% respectively). Altogether, these results show that ELISA is more sensitive than CIE: and that both tests are highly specific for amoebic liver abscess.

The gold standard for the diagnosis of ALA is the liver biopsy. However this procedure is not currently done in every case. Otherwise, the percentage of positively for trofozoites in the analysis of pus in the liver abscess is low. Therefore the E. histolytica antibodies determination by ELISA is a good diagnostic tool in ALA, as a non-invasive and cost-effectiveness procedure.

In conclusion, the diagnosis of liver amoebiasis in tropical countries can be attempted by the search of E. histolytica antibodies by both ELISA and CIE techniques. Currently we are routinely using ELISA to confirm ALA diagnosis. This will allow us to study the seroprevalence as well as the reliability of this test in therapeutic follow-ups. Other cases of liver abscess should be included (e.g. hydatid cyst, metastatic or hepatic carcinomas), as well as other parasitic diseases, in order to assess the possibility of cross-reaction with diseases clinically similar to extra-intestinal amoebiasis.

RESUMO

O abscesso hepático é uma complicação mais frequente da amebíase intestinal: o seu diagnóstico sugere-se pelo quadro clínico, mas é confirmado pelos estudos parasitológicos. Para confirmar o diagnóstico precisa-se identificar a E. histolytica, o
que é apenas possível em muito poucos casos. As pratas sorológicas melhoram notadamente o diagnóstico das complicações severas da amebiase. Em nosso estudo comparávamos o teste de ELISA e a Contraimunoeletroforese (CIE). Ambas as técnicas foram utilizadas para detectar anticorpos contra ameba em 50 pacientes sem amebiase, 30 pacientes com abscesso hepático e 30 com amebiase intestinal. Todos os soros dos pacientes sem amebiase foram negativos por ambas as técnicas. Quando analisamos os soros dos pacientes com amebiase intestinal, encontrou-se que 10% destes, foram positivos para ELISA, enquanto que nenhum o foi para CIE. Nos soros dos pacientes com abscesso hepático, encontrou-se uma positividade de 90% para ELISA e 66.6% para CIE. Estes resultados mostram que o teste de ELISA foi de maior sensibilidade (100% de sensibilidade) no diagnóstico do abscesso hepático amebiano, quando comparado com a Contraimunoeletroforese (66.6%).

Palavras-chave: Abscesso hepático amebiano, Diagnóstico, ELISA. CIE.

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