ABSTRACT

An extract of Triatoma infestans has previously been demonstrated to produce specific IgG and IgE both in animals and in atopic humans with rhinitis/asthma as well as hypersensitivity pneumonitis in guinea pigs aerosolized with T. infestans. We attempted to determine whether the antigen or allergens responsible belonged to the protease group, as occurs with other allergens such as house dust mites and cockroaches.

To do this, T. infestans was studied by SDS-PAGE, Western blots and gelatinolysis with and without the use of specific protease inhibitors such as E-64, TLCK, TPCK, PMSF, leupeptin, o-phenanthroline and pepstatin-A.

These assays revealed serine-like proteolytic and gelatinolytic activities. The presence of 10 to 12 bands of between 14 and 100 kDa was detected. The proteolytic activity pattern of T. infestans was greatest at pH 8.5 and gelatinolytic activity was highly sensitive to PMSF, suggesting that this enzyme could be characterized as a serine protease.

Western blots revealed that two bands of 17 and 58 kDa reacted with the sera of atopic humans with respiratory diseases and anti-IgE. However, whether these bands correlated with allergenicity is unclear since the presence of several proteins in each of these bands does not rule out the possibility that this correlation could exist, especially because cross-reactions with antigens from the cockroach Periplaneta americana and its specific antiserum in animals and atopic humans have been demonstrated.

The role of proteases in the etiopathogenesis of perennial rhinitis and bronchial asthma in inhabitants of the area of Argentina infested by T. infestans requires further investigation.

Key words: Triatoma infestans. Serin-proteases. Cross-reactivity with cockroaches. Inhalant insect allergy.

RESUMEN

Habiendo demostrado previamente que un extracto del Triatoma infestans (Ti) era capaz de generar anticuerpos específicos IgG e IgE tanto en los animales como en los humanos atópicos con rinitis/asma al igual que la producción de una típica neumonitis por hipersensibilidad en cobayos aerosolizados con Ti inten
tamos analizar si el o los antígenos responsables per
tencían al grupo de las proteasas tal como sucede con otros alergenos, como los acaros y los blátidos.

Para ello el Ti fue estudiado por medio del SDS-
PAGE, Western blots y gelatinolisis con y sin el em-
plo de inhibidores específicos de las proteasas, tales como, el E-64, el TLCK, el TPCK, el PMSF, la leupeptina, la o-ferantrolina y la pepstatina-A.

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Una vez comprobada la actividad proteásica y la gelatinolítica se destacó la presencia de 10 a 12 bandas entre los 14 y 100 kDa con un patrón proteolítico con una mayor actividad a pH 8.5 y con una gelatinolisis altamente sensible al PMSF revelando su posible actividad de serina.

Por los Western blots se detectó que las bandas de 17 y 58 kDa eran reactivas con los sueros humanos de atópicos respiratorios y la anti-IgE aunque no queda muy clara su alergenicidad ya que la presencia de varias proteínas en cada una de estas bandas no excluye que dicha correlación pudiera existir más aún cuando se demuestran reacciones cruzadas con antígenos de la cucaracha Periplaneta americana y sus antisueros específicos en animales y en atópicos.

Su papel en la etiopatogenia de la rinitis perenne y del asma bronquial de los habitantes de la zona endémica argentina para el Ti requiere de mayores investigaciones.

**Palabras clave:** Triatoma infestans. Proteasas tipo serina. Reactividad cruzada con cucarachas. Alergia a insectos.

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**INTRODUCTION**

In previous papers it was demonstrated that an extract of Triatoma infestans (Ti) – a reduviid belonging to the Triatominae subfamily – was capable to elicit a specific IgG response in rabbits, a specific IgE in atopics suffering perennial rhinitis and asthma and a hypersensitivity pneumonitis in an animal model.

Peptidases are classified in serine, cysteine, aspartyl and metalloproteinases, according to: 1) the reaction that they catalyze; 2) the chemical nature of the catalytic site and 3) the evolutionary relationship revealed by their structures.

In this paper, we present evidence indicating that Ti contain serine-proteinases with gelatinolytic properties which might be involved in their immunogenicity.

**MATERIALS AND METHODS**

**Antigens**

Ti and Periplaneta americana (Pa) were collected and both extracts were prepared as it was previously described following Frugoni-Hansen's method with slight modifications. The Bradford method was applied to establish the protein content of Ti and Pa although this last antigen seems to be similar to that baptized as Per a 7.

**Antisera**

Adult albino rabbits were immunized against the whole Ti extract (13 mg/ml). Thus a rabbit polyclonal anti-Ti serum was obtained. Human sera were collected from atopic patients suffering rhinitis/asthma who exhibited positive immediate skin tests and IgE-RAST ≥ 0.35 PRU/ml to Ti and to Pa separately.

**Polyacrilamide gel electrophoresis**

SDS-PAGE was performed by the method of Laemmli using a 4 % stacking gel and a 15 % running gel in a Mini-Protean II apparatus. Twenty microliters of Ti and Pa were loaded in separated wells with reducing and boiling conditions both for the detection of proteins by Coomasie R 250 brilliant blue and for electrotransference to a nitrocellulose membrane.

**Enzymatic activity assay**

Minigels of 10 × 10 cm each and 1.5 mm thick composed of 12 % acrylamide were made as described by Laemmli (I with gelatin at a final concentration of 0.2 %. They were run at 130 V for 2 hs. When the bromophenol blue used as a marker reached the bottom, the run was stopped and the gels were washed twice in distilled water with Triton-X-100, 0.15 % for 15 min each, then incubated at 37 °C in 0.1 % 2-[N-morpholino] ethane sulfonic acid (MES) buffer at pH 6, in Tris ACH 100 mM pH 3.5 and Tris CH 100 mM pH 8.5 containing 0.5 mM dithiothreitol (DTT).

The reaction was stopped and the remaining protein was stained by incubation at room temperature with 0.2 % Coomassie brilliant blue R-250 in methanol-acetic acid-water 5:1.5:5 (v/v/v).

After destaining in methanol 20 % and acetic acid 10 %, the active bands were observed as colorless over a blue background.

**Inhibitory assays**

The washing and incubation of the gels were done with and without the protease inhibitors. The solutions employed were E 64 (L-trans-epoxy-succinylleucylamido[4-guanidino]-butane) 20 μM; to...
syl-lysyl-chloro-methyl-ketone (TLCK) 100 μM; tosyl-phenyl-allyl-chloro-methyl-ketone (TPCK) 1 mM; phenyl-methyl-sulphonyl-fluoride (PMSF) 10 mM; leupeptin 100 μM; α-phenantrline 1 mM and pepstatin-A 2 μM.

The molecular weight markers were phosphorylase-b (97.4 kDa), bovine serum albumin (BSA) (66.2 kDa), ovalbumin (45 kDa), carbonic anhydrase (31 kDa), trypsin inhibitor (21.5 kDa) and lysozyme (14.4 kDa). For activity gels, the samples were not reduced nor boiled before loading.

**Western blots**

The samples with or without β-mercaptoethanol and boiling were run in 15% standard polyacrylamide gel in the presence of SDS (SDS-PAGE), electro-transferred to nitrocellulose sheets, blocked for 2 hs with a solution containing 2 % fatty acid-free BSA, 0.01 % v/v Tween-20, PBS pH 7.2 and then incubated overnight with rabbit polyclonal anti-Ti serum 1/400, human sera anti-Ti 1/10 and human sera anti-Pa 1/5. After overnight incubation with the rabbit or human antisera, respectively, and repeated washing the sheets were treated with 1/3000 goat anti-rabbit IgG horseradish peroxidase conjugate or 1/500 rabbit anti-human IgE specific for ε-chains peroxidase conjugate at room temperature during 2 hs. The chromagenic detection was developed using N-chloronaphtol and hydrogen peroxide.

**RESULTS**

Ti in SDS-PAGE showed 10 to 12 bands between 14 and 100 kDa meanwhile Pa only showed 2 bands at 28 and 46 kDa. The total protein content of both extracts were 1 mg/ml (fig. 1).

The gelatinolytic activity of Ti in SDS-PAGE with co-polymerized gelatin as substrate was recorded at 56 kDa. The proteolytic activity pattern of Ti was preliminarily analyzed at three different pH levels, 4.5, 6.5 and 8.5. The highest enzyme activity was at pH 8.5 within the band of 56 kDa.

Total activity pattern at pH 8.5 was highly sensitive to PMSF exclusively suggesting that we can tentatively characterized this enzyme as a serine-protease (fig. 3).

When the Ti was separated by SDS-PAGE, transferred to nitrocellulose and incubated with a polyclonal rabbit anti-Ti serum, the bands of 15 kDa, 18 kDa and 36 kDa showed immunoreactivity but only one band of 28 kDa cross-reacted with the Pa extract (fig. 4).
On the other hand, a human anti-Pa serum recognized a band of 28 kDa in Pa and cross-reacted with several bands in Ti (14 kDa, 35 kDa, 36 kDa and 75 kDa) (Fig. 5).

A human anti-Ti serum recognized 2 bands of 17 kDa and 58 kDa with its own antigen meanwhile in the cross-reaction with Pa these 2 bands were located at 28 kDa and 45 kDa (Fig. 6).

### DISCUSSION

Several major allergens in the extracts of insects possess protein hydrolases in their composition. Also, acid phosphatase activities correlate well with allergenic potency in pollen extracts.

Elegant studies showed the identification of a novel serin-protease with allergenic activity from Dermatophagoides pteronyssinus\(^1\).

In previous papers we described the correlation between some proteases with gelatinolytic activity and the allergenicity of house-dust mite, cockroach and bat feces extracts as it was revealed by specific Western blots\(^2\).

We suppose that this is the first report about the serine protease and gelatinolytic activities of the Ti extract although the relationship between these properties and its immunogenicity could not be clearly established.

Considering that many different proteins may be present in each broad band we cannot rule out the fact that the connexion would be possible if we take into account the biological interactions between protease epitopes and the IgE or FcεRI receptors were demonstrated in other models.

It was also confirmed that the extracts of Ti and Pa exhibited common epitopes who were established in the past by our group.

It is very important to define the proteases present in all these extracts and to determine their abili-
ty to degrade other proteins of the same extract or in mixtures of allergen extracts.
Their role in the immunopathological aspects of rhinitis and asthma requires further investigation.

REFERENCES