Identification of the 33-kDa alkaline protease of Aspergillus fumigatus in vitro and in vivo

M. MOUTAOUAKIL, M. MONOD, M. C. PRÉVOST†, J. P. BOUCHARA‡, S. PARIS and J. P. LATGÉ

Unite de Mycologie, †Station Centrale de Microscopie électronique, Institut Pasteur, 25 rue du Dr Roux, 75015 Paris, France,‡Service de Dermatologie, Centre Hospitalier Universitaire Vaudois, 1011 Lausanne, Switzerland and§Laboratoire de Parasitologie-Mycologie, Centre Hospitalier Universitaire, Angers, France

Summary. Aspergillus fumigatus produced a 33-kDa serine protease (ALP) in vitro and in vivo. In vitro, this alkaline protease was secreted when the fungus was cultivated in the absence of protein, if the pH of the medium remained close to neutrality. Western blotting and immunoelectronmicroscopy studies showed that ALP was localised in the wall of the fungus and was degraded after secretion in the culture medium under conditions of low pH. Although present in the lung during infection, ALP did not appear to be diagnostically useful and was different from the precipitating chymotrypsin antigen used in the diagnosis of aspergilloma.

Introduction

A 32–33-kDa serine protease has been isolated recently from Aspergillus fumigatus by several laboratories.1-4 This alkaline protease (ALP) of the subtilisin family degrades collagen, fibrinogen and elastin. Its secretion is induced in vitro by the presence of a protein (collagen, fibrinogen, elastin, casein or liver extract) in the culture medium.1-4 It has been suggested that such extracellular proteases can play a role during the invasion of lung tissues by A. fumigatus or A. flavus.5-7 However, there is no direct evidence that the protease is secreted in vivo and is required by the fungus to invade host tissues.

The specificity of the antibody response to ALP is questionable. Antibodies were detected with high titres in the sera of only two of six patients with aspergilloma.1 In BALB/c mice, Frosco et al.8 found that ALP was weakly immunogenic whereas other authors produced monospecific rabbit polyclonal antibodies toward this protein.1,2 Western blots and immuno-electronmicroscopy were used to study ALP synthesis in vitro and in vivo. Some of the physicochemical factors involved in the secretion of ALP have been investigated. The low antigenicity of this protein has been demonstrated.

Materials and methods

Strains

A wild-type clinical isolate of A. fumigatus (CBS 144.89) and an alkaline protease-deficient mutant 18 (nia, alp::HPH) derived from the wild strain by genetic engineering8 were maintained on malt agar 2 % slants.

Human sera

Serum samples from patients with aspergilloma, patients with candidiosis and healthy individuals were used throughout this study. Serum reactivity was monitored by both Ouchterlony diffusion assay with somatic and metabolic extracts from A. fumigatus (Diagnostic Pasteur, Marnes, France) and indirect haemagglutination assay (Fumouze, Clichy, France). Candidiosis patients gave negative results in all these assays. All aspergilloma patients were chymotrypsin-positive. A pool of positive human sera was prepared from 75 patients. A control pool of human sera was prepared from 50 healthy donors. Positive and negative serum pools contained equal volumes of all sera.

Rabbit sera

A monospecific rabbit antiserum directed against the 33-kDa protease was prepared as described previously.2 A monospecific rabbit antiserum to the chymotrypsin antigen C or antigen 13 was kindly supplied by Dr J. Longbottom.10 Pre-immune rabbit sera were used as controls.

In-vitro extracts

For the detection of the 33-kDa serine protease, three culture media were used: (a) Sabouraud, i.e., glucose 2 % plus Mycopeptone Biokar (Prolabo, Paris) 1 %; (b) collagen (Serva) 0.2 %; (c) Mycopeptone Biokar 1 %. Conidia of A. fumigatus strain CBS 11489 were inoculated into 150-ml Erlenmeyer flasks containing 50 ml of Sabouraud liquid medium. Flasks were shaken for 2-3 days at 100 rpm and 25°C. Two-litre Biolafitte fermenters, containing 1·2 L of medium, were inoculated with the medium from two shaken flasks.
flasks (inoculum 8% v/v). The culture conditions were as follows: temperature 25°C, aeration 30 L of air/h, agitation 600 rpm, and pH was monitored. Fermentation experiments were also performed in Sabouraud liquid medium in fermenters with pH controlled at 7-2 throughout the duration of the culture. Strain CBS 114-89 was grown for 2-4 days depending on the culture medium. Mycelium was separated from the culture medium by paper filtration. Culture filtrate was precipitated overnight at 4°C by either four volumes of ethanol or ammonium sulphate dissolved in the culture filtrate to 90% saturation. The ethanol precipitate was washed twice with ethanol, resuspended in water and ultrasonicated for 3 min. The undissolved precipitate was removed by centrifugation and the water-soluble material was kept frozen at -20°C. The ammonium sulphate precipitate was collected by centrifugation, resuspended in distilled water and dialysed against water for 24 h and 20 mm Tris-HCl buffer, pH 7.4, overnight at 4°C. Preliminary experiments have shown that both techniques precipitate the 33-kDa protease.

For the preparation of extracts containing the chymotryptic antigen, the wild strain and the ALP-mutant type were grown in unshaken flasks containing Sabouraud medium. After 1 month at 25°C, mycelium was separated from the culture medium by paper filtration. Culture filtrate was concentrated by dialysis against polyethylene glycol 20000. Concentrated culture filtrate was dialysed against distilled water and stored at -20°C.

**Mouse infection**

Male IOPS OF1 mice, 20–25 g (IFFA CREDO, L’arbresle, France) were used for infection experiments. All animals were pre-treated with cortisone: 5 mg of cortisone acetate (200 μl of a NaCl 0.9% aqueous solution containing cortisone acetate 25 mg/ml) were given intraperitonally 2 days before and on the day of infection. Conidia were obtained from subcultures of strain CBS 144-89 grown on malt agar 2% for 4–7 days at 25°C. They were harvested with a NaCl 0.9% aqueous solution and counted with a haemacytometer. Mice were infected intranasally with 50 μl of a conidial suspension of *A. fumigatus* (10⁴ conidia/ml) as described previously. Injection of this dose typically caused 100% mortality within 3 days in the infected mice. Control mice pre-treated with cortisone received intranasal injections of saline solutions. Control mice survived for at least 2 weeks after challenge.

**Infected lung extract**

Five lungs from infected dead mice or control mice were pooled in 0.1 M Tris-HCl buffer, pH 8.4, containing 2 mM PMSF and homogenised in a Sorvall homogeniser for 1 or 2 min at 4°C. Crude tissue homogenates were placed in a bottle with an equivalent volume of 1-mm glass beads and disrupted for 3–4 min under CO₂ cooling in an MSK Braun cell disintegrator. Cell debris was removed by centrifugation (30 min, 15000 rpm). Ammonium sulphate was added to the supernate to achieve 65% saturation. After standing overnight at 4°C, the precipitate was collected by centrifugation (30 min, 15000 rpm), resuspended in the homogenising buffer and extensively dialysed against 2·5 mM Tris-HCl buffer, pH 8.4, containing 2 mM PMSF.

**Protein determination**

Protein concentration of the extracts was estimated by the BioRad method based on Coomassie Blue staining of the proteins with bovine serum albumin (BSA) as a standard.

**Electrophoresis and immunoblotting**

Protein extracts were submitted to SDS-PAGE according to the method of Laemmli with a separating gel of acrylamide 12%, 10–25 μg of protein from in-vitro extracts and 600–700 μg of in-vivo extracts were loaded per well.

Gel contents were transferred electrophoretically on to nitrocellulose membranes (0-2-μm pore size; Schleicher and Schuell, Ceralabo, Paris). Transfer was done in a 25 mM Tris-192 mM glycine buffer, pH 8.3, containing methanol 20% v/v at 30 V overnight. Blots were stained with Ponceau Red S 0.3% in TCA 0.3% to locate mol.-wt markers and to evaluate protein transfer. Blocking was achieved by incubation for 1 h at room temperature in phosphate buffered saline (PBS) containing Tween 20 0.15% v/v (PBST) and non-fat dried milk (Regilait) 5% w/v (PBSTM). The antigenic reactivity was determined by incubation of nitrocellulose strips with transblotted material in PBSTM containing rabbit antiserum at a 1 in 1000 dilution (and 1 in 100 in the case of anti-Ag13 antiserum) or human sera at a 1 in 500 dilution for 1 h at room temperature. In the case of in-vitro extracts, sites of binding of antibodies were determined by incubation of the blot in a peroxidase-conjugated goat anti-rabbit or anti-human IgG (H+L) diluted at 1 in 500 and 1 in 2000, respectively, in the same buffer. The reaction was achieved with an enhanced chemiluminescence Western blotting detection system (ECL kit, Amersham). For this method, blocking was achieved by incubation for 1 h at 37°C in 0.1 M Tris, pH 8.0, containing 15 mM NaCl, 1 mM EDTA, Tween 20 0.05% (TBST) and non-fat milk 10%. Rabbit antibody and peroxidase-conjugated goat anti-rabbit IgG (H+L) were diluted to 1 in 500 and 1 in 20000, respectively, in the same buffer. Final washings were
done with three rinses of TBST and two rinses of 0.1 M Tris, pH 8.0, containing 15 mM NaCl. Development was performed for 30 s.

**Counterimmuno-electrophoresis**

Counterimmuno-electrophoresis was performed on a cellulose acetate membrane (Sartorius, 11200-25-160 IEN) basically as described previously. Serum (15 µl) from a pool of aspergilloma patients was deposited towards the anode whereas the antigen from wild-type and mutant strains (15 µl at 1-1.5 mg protein/ml) was deposited at the cathode. Migration buffer was 50 mM Tris-glycine, pH 8.8. After electrophoresis for 2 h at 100 V, membranes were washed in a NaCl 0.85% solution. Chymotrypsin activity was localised in precipitin arcs by use of N-acetyl-DL phenyl alanine β-naphthyl ester as a substrate.

**Immunofluorescence**

Lungs were fixed overnight at 4°C in formaldehyde 10% aqueous solution (40 vol, Labo Nord, Villeneuve d'Ascq, France) and embedded in paraffin after dehydration with increasing concentrations of ethanol and a final rinse in toluene. After removal of the paraffin, tissue sections (5 µm) were incubated with rabbit anti-ALP serum diluted 1 in 100 in PBS for 30 min at 37°C. The sections were washed extensively in PBS, and then incubated with anti-rabbit immunoglobulin-fluorescein conjugate (Institut Pasteur Production) diluted 100-fold in the same buffer. Before fluorescence microscopy, the tissue sections were stained with Evans blue diluted 1 in 10000 in water. The same experiment was done with a pre-immune rabbit serum as control.

**Immuno-electronmicroscopy**

An infected lung was fixed for 1 h at room temperature, trimmed in 5 x 5-mm blocks and post-fixed overnight at 4°C in a solution containing formaldehyde 2.5% v/v, glutaraldehyde 0.5% v/v in a sodium cacodylate buffer, pH 7.2. After incubation for 12 h in the cacodylate buffer containing 0.01 M ammonium chloride material was dehydrated with increasing concentrations of ethanol and embedded in LR White resin as described by Newman and White at room temperature. Ultra-thin sections recovered on formvar-coated grids were incubated for 1 h at room temperature with anti-ALP anti-serum diluted 1 in 100 and 1 in 500 in PBS, pH 7.2, containing BSA 1% w/v and subsequently washed in the same buffer containing BSA 0.1% w/v. Sections were then incubated with gold-labelled anti-IgG conjugate (10 nm; Janssen, France) diluted 1 in 50 in PBS with BSA 1%. After 1 h at room temperature, sections were washed three times with PBS with BSA 0.1%, twice with PBS and five times with water. A pre-immune rabbit serum was used as control.

**Results**

**Secretion of ALP in vitro**

Immunoblot analysis with the monospecific anti-ALP rabbit antiserum demonstrated that ALP was present in the culture filtrate of three media: collagen 0.2%, mycopeptone 1% and Sabouraud medium with pH controlled at 7.2 (fig. 1). ALP was not detected in the culture filtrate of Sabouraud medium. The change in pH in Sabouraud medium, collagen medium and peptone medium is shown in fig. 2. Conversely, in collagen 0.2% and mycopeptone 1% medium where

---

**Fig. 1.** Immunoblot analysis of culture filtrate extracts from different media: lanes a and e, collagen 0.2%; b, Sabouraud medium without pH control; c, Sabouraud medium with pH controlled at 7.2; d, mycopeptone 1%. Blots were probed with a 1 in 500 dilution of the anti-ALP rabbit antiserum (a-d) and a pre-immune rabbit serum (e).

**Fig. 2.** pH kinetics in three different liquid inoculated with A. fumigatus: Sabouraud ■; collagen 0.2% ▲; mycopeptone 1% ○.
Fig. 3. Immunoblot analysis of culture filtrate extracts of *A. fumigatus* strain CBS 14489 probed with an anti-33 kDa ALP antiserum. Lane 1, collagen medium; 2, collagen medium incubated at pH 4.5 for 24 h at room temperature; 3, collagen medium mixed with Sabouraud medium (2 day growth); 4, Sabouraud medium (2 day growth).

Fig. 4. Reactivity of mycelia in the lung of an immunocompromised mouse infected with *A. fumigatus*. Indirect immunofluorescence was achieved with anti-ALP rabbit antiserum diluted 1 in 100 and fluoresceinylated anti-rabbit IgG antiserum diluted 1 in 100.

Fig. 5. Reactivity of the anti-ALP antiserum with the wall of a hyphal element in the lung of an infected mouse. (a) Anti-ALP and (b) pre-immune sera were diluted 1 in 500 and anti-rabbit IgG immunoglobulin conjugated to colloidal gold (10 nm) was diluted 1 in 40. w, mycelial wall; c, cytoplasm (×51000).

the pH remained slightly basic, a significant decrease in the pH of the medium was observed during mycelial growth in Sabouraud medium. Adjustment of the pH of a culture filtrate from collagen medium, containing ALP, to a value of 4.5 (similar to the pH found in Sabouraud medium at maximal growth) resulted in the disappearance of the 33-kDa protease after 24 h at room temperature. The 33-kDa ALP was still detected in control culture filtrate (pH 7.5) left for 24 h at room temperature. Moreover, co-incubation for 24 h at room temperature of equal volumes of a collagen culture filtrate which contained the 33-kDa protease, and a Sabouraud medium culture filtrate at pH 4.7 (2 days of growth), also induced the disappearance of the 33-kDa protease (fig. 3). Secretion of ALP was also demonstrated by immunofluorescence on germ tubes developed after incubation of conidia overnight at 37°C in Sabouraud medium, during a period of time where no decrease in pH was observed (data not shown). The 33-kDa serine protease was degraded by acid pH and the low molecular mass (18 kDa) immunoreactive band found in Sabouraud medium represented an ALP breakdown product. ALP was also secreted in a yeast extract 1% medium or in a medium containing glucose 2%, asparagine 0.5% and a salt solution (data not shown). The secretion of ALP in the protein hydrolysate media (which do not contain protein as shown by SDS-PAGE) and in the asparagine-based medium showed that the presence of a protein in the culture medium was not necessary to induce the secretion of ALP. The requirement of a
Presence of ALP in vivo

Indirect immunofluorescence studies showed that anti-ALP rabbit antiserum labelled mycelium in the lungs of infected mice. Binding of immunoglobulin was predominantly to the wall of fungal cells. No labelling was seen with pre-immune rabbit serum by either immunofluorescence or immuno-electronmicroscopy (figs. 4 and 5).

The presence of ALP in the lungs of infected mice was confirmed by Western blots of infected lung extracts (fig. 6).

In Western blots, ALP did not discriminate between samples from aspergilloma patients and other patients not suffering from aspergilloma. Amongst 29 sera tested, 25 labelled the 33-kDa ALP at a 1 in 500 dilution. No difference was seen between the A. fumigatus positive patients (17 of 21) and eight negative patients (fig. 7).

The anti-Ag 13 rabbit antiserum did not react with the 33-kDa protein even at a 1 in 100 dilution (although at a 1 in 1000 dilution it bound strongly to a high mol. wt antigen in the collagen extract). Moreover, counterimmuno-electrophoresis of a pool of patients and culture filtrate extracts from the wild strain and a 33-kDa protease-deficient mutant was positive for chymotrypsin antibody in both extracts (fig. 8).

Discussion

Previous work has shown that the alkaline protease from A. fumigatus is secreted in media containing a protein source. Northern blot experiments have shown that the synthesis of ALP was highly stimulated...
if a protein such as collagen was present in the culture medium.\textsuperscript{13} Our results showed that ALP can still be synthesised and secreted in the absence of protein in the culture medium if the pH is held slightly basic, irrespective of the composition of the medium. In Sabouraud medium where the pH becomes acidic, ALP was synthesised and degraded due to acidic conditions. Active synthesis of ALP in Sabouraud medium was confirmed by: (a) the detection of ALP by Western blot using an antiserum from a rabbit immunised with an intracellular extract from a mycelium grown in Sabouraud medium for 2 days, whereas an antiserum from a rabbit immunised with the culture filtrate from the same medium gave negative results (data not shown); and (b) the identification of specific cDNA coding for ALP in a library constructed from mycelium grown for 4 days in a Sabouraud medium.\textsuperscript{17} Battaglino et al. have also demonstrated that the production of the alkaline protease from \textit{A. oryzae}, which is highly related to the one produced by \textit{A. fumigatus} is highly reduced by low pH.\textsuperscript{18} These authors stated that the pH of the culture medium must be maintained over 6-5 for a high protease yield. Although Reichard et al. did not mention the role of pH during secretion of the enzymes, their data showed that the decrease of pH during growth is correlated with a decrease in the proteolytic activity of the culture filtrate.\textsuperscript{1}

The human pathogens \textit{A. flavus} and \textit{A. fumigatus}, like most members of the genus \textit{Aspergillus}, produce high amounts of proteases of different classes \textit{in vitro}: serine protease, aspartyl proteases, metalloproteases.\textsuperscript{2,18-21} A limited number of studies have suggested that in the genus \textit{Aspergillus}, as in the genus \textit{Candida}, proteases may be essential virulence determinants. This study presents immunochromical and cytological evidence that the major 32-33 kDa protein of \textit{A. fumigatus} is indeed secreted \textit{in vivo}. It confirmed previous observations by Reichard et al.\textsuperscript{2} who showed that the 32-kDa protease they studied was also detectable by immunofluorescence in the lung of experimentally infected guineapig.\textsuperscript{1} However, the role of this protease during the infection process remains an open question since ALP\textsuperscript{2} mutants were still pathogenic for mice.\textsuperscript{9}

Reichard et al. have detected high titres of antibodies against the 33-kDa ALP in some aspergilloma patients by ELISA.\textsuperscript{3} With Western blotting assays, we have shown that anti-33-kDa antibodies did not discriminate between aspergilloma and candidiosis patients. The discrepancy between these results could be due to the different immunoassays used. With different purified 33-kDa ALP batches, we have found that trace amounts of high mol.-wt. proteins not detectable by Coomassie Blue staining but very immunogenic and easily seen by Western blotting experiments, can contaminate purified fractions of ALP and give rise to false positive results in ELISA (unpublished observations).

The reactivity of control sera against the 33-kDa protease suggested that this protease may share common epitopes with non-\textit{Aspergillus} proteins. By the combined use of ALP\textsuperscript{2} mutant and a monospecific antiserum directed against the chymotrypsin antigen it was shown that the 33-kDa protease is different from the "chymotrypsin" antigen (= AgC = Ag 13) which is used in the diagnosis of aspergilloma. This antigen is a high mol.-wt, glycosylated (ConA positive) molecule. The absence of cross-reactivity of anti-antigen 13 antiserum with the 33-kDa protease is in agreement with previous results stating that only antigens with molecular masses higher than 43 kDa can precipitate the antibodies of aspergilloma patients.\textsuperscript{10} Moreover, somatic antigens from Diagnostics Pasteur which have a high "chymotrypsin" activity in counterimmunoelectrophoresis have a very low azocoll degradation activity (data not shown) whereas ALP degrades azocoll very efficiently.\textsuperscript{3} The chymotrypsin antigen of \textit{A. fumigatus} has been identified until now only by its ability to degrade N-acetyl-DL-phenylalanin \(\beta\) naphthyl ester. The nature of this chymotrypsin antigen remains to be determined.

This research was supported by INSERM grant 900313.

References

13. Towbin H, Staehelin T, Gordon J. Electrophoretic transfer of
proteins from polyacrylamide gels to nitrocellulose sheets: procedure and some applications. Proc Natl Acad Sci USA 1979; 76: 4350-4354.


ANNOUNCEMENTS

The Royal College of Pathologists
One-day Symposium

Diagnostic Coding and Disease Classification

Thursday 2nd December 1993 to be held at the Royal College of Pathologists, 2 Carlton House Terrace, London SW1Y 5AF

Provisional programme

Chairman: Professor P. G. Toner (Queen's University of Belfast); Introduction; ICD—The original international classification for health care statistics, Speaker from WHO, Geneva (Name to be confirmed); Read codes—The standard coded thesaurus of clinical terms for the NHS, Dr C. Stuart-Buttle (NHS Centre for Coding and Classification, Loughborough); SNOMED International—The most comprehensive system for medical information management, Professor Roger Cote (Chairman, SNOMED International Editorial Board). Lunch (Refreshments provided).

Current Developments in OPCS Mortality Coding, Dr Cleo Rooney (Office of Population Censuses and Surveys, London); Coding, Classification and the National I M & T Strategy, Mr Ian Smith (Director of Strategic Planning, Information Management Group, NHS Management Executive, Leeds); The way forward, General discussion and questions; Demonstrations

The symposium is open to members of the College, to trainee pathologists and to workers in other disciplines with an interest in the subject. The programme is approved by the Thames Postgraduate Deans and hospital doctors may apply to their employing authority for Study Leave under HM(67)27.

The registration fee is £94 and includes coffee, lunch and tea.

Details and application forms may be obtained from: Maureen Russell, Royal College of Pathologists, 2 Carlton House Terrace, London SW1Y 5AF, Tel: 071 930 5862 (Ext: 24/26).