Binding of C-reactive protein to Aspergillus fumigatus fractions

TINE D. BØRGLUM JENSEN, H. SCHØNHEYDER, P. ANDERSEN and A. STENDERUP

Institute of Medical Microbiology, Bartholin Building, University of Aarhus, Department of Medicine, Marselisborg Hospital, DK-8000 Aarhus C, Denmark

Summary. Calcium-dependent binding of C-reactive protein (CRP) to Aspergillus fumigatus was determined by enzyme-linked immunosorbent assay. A homogenate of young hyphae was fractionated by hydrophobic interaction chromatography followed by gel filtration. High CRP-binding activity was found in a fraction of mol. wt c. 500,000 which was characterised by strong binding to the hydrophobic column. Three fractions of less conspicuous CRP-binding activity were identified (c. 500 000, 150 000 and 150 000–50 000 mol. wt respectively). In these four fractions, phosphorylcholine was detected by an anti-phosphorylcholine mouse hybridoma antibody. Some CRP-binding activity in fractions with low affinity for the hydrophobic column did not correspond closely with the presence of phosphorylcholine. It is suggested that C-reactive substance in A. fumigatus is heterogeneous. The C-reactive substances did not correspond with fractions containing major antigens (470 000 and 250 000 mol. wt respectively) which elicit a strong immune response in man.

Introduction

Components of various micro-organisms including bacteria, fungi and parasites, capable of interacting with C-reactive protein (CRP), are referred to as C-reactive substances (Pepys and Longbottom, 1971; Gotschlich et al., 1982). CRP belongs to a group of serum proteins, known as acute-phase reactants, which circulate in increased amounts during infection, inflammation and tissue damage. CRP complexed with calcium has high affinity for polyaniions, galactose residues and phosphorylcholine which have been found in some C-reactive substances (Gotschlich and Edelman, 1967; Volanakis and Kaplan, 1971; Gotschlich et al., 1982). A calcium-independent binding to polycations has also been described (Di Camelli et al., 1980).

Aspergillus fumigatus contains C-reactive substance. Longbottom and Pepys (1964) described a precipitin reaction involving aspergillus C-reactive substance and CRP and pointed to this as a source of error in the serodiagnosis of aspergillosis. Pulmonary aspergillosis occurs in different clinical forms (Pennington, 1980); complement activation may be important in some of them. This activation may be elicited by antigen-antibody complexes but other mechanisms could play a role. The interaction of CRP and C-reactive substances may activate the classical complement pathway (Kaplan and Volanakis, 1974) and thus aspergillus C-reactive substance may participate in the pathogenesis of aspergillus-related diseases through complement activation. In this study we have investigated some of the characteristics of A. fumigatus C-reactive substance.

Materials and methods

Preparation of CRP

CRP was prepared from ascitic fluid from patients with either ovarian or mammary carcinoma, admitted to the Department of Oncology, Radiumstationen, Municipal Hospital, Aarhus. Specimens were tested for CRP by double immunodiffusion (ID) against rabbit anti-human CRP (Dakopatts, Denmark). Only samples with titres above two were used, and CRP was prepared from such samples by anion exchange chromatography essentially as described by Johnson and Prellner (1977). The fractions obtained by chromatography were tested for CRP by ID. Quantitation of purified CRP was done by radial immunodiffusion (RI) in barbital-buffered agarose (HSA, Litex, Denmark), pH 8.2, with 1·5 μl of anti-CRP (Dakopatts) added /cm². The purity of CRP was assessed by crossed immuno-electrophoresis (CIE) in agarose (LSL, Litex) against a homologous antiserum with a Tris-barbital buffer, pH 8.6, according to the method of Weeke (1973) with Na₂EDTA 10 mmol/L added. Plates measured 7 x 7 cm and 10 μl of antiserum were added per/cm². The homologous antiserum was raised in three New Zealand white rabbits by immunisation with 75 mg of
purified CRP every second week for at least 2 months. CIE against the homologous antiserum revealed two faint contaminants together with a CRP arc which cross-reacted with a known CRP antiserum (Dakopatts) by intermediate gel technique. The contaminants did not react with anti-IgG, -IgA or -IgM (Dakopatts) added to the intermediate gel.

The protein content of purified CRP was assayed by the Bio-Rad method (Bio-Rad, USA) with human serum albumin as the reference (KabiVitrum, Sweden).

**A. fumigatus fractionation**

*A. fumigatus* NCPF 2109 was grown in Czapek-Dox broth at 37°C for 2–4 days, harvested and ruptured with a freeze-press technique (Eaton, 1962). The supernate referred to as ruptured mycelium, obtained by centrifugation at 35 000 g for 45 min at 4°C, was submitted to hydrophobic interaction chromatography and gel filtration as previously described (Schönheyder and Andersen, 1984a). In brief, 40 ml of ruptured mycelium with NaCl added to a concentration of 4 M were loaded on a Phenyl-Sepharose CL-4B column equilibrated in Tris buffer, pH 7.4, with 4 M NaCl. The column was eluted successively with Tris buffers of decreasing NaCl molarity (4 M, 2 M, 0.15 M) and with twice-distilled water. At each elution step, fractions of peak absorbance at 280 nm (referred to by their NaCl molarity) were pooled, concentrated by pressure dialysis in collodion bags (Sartorius, FRG) and loaded on a Sephadex G-200 column (Pharmacia Fine Chemicals). The effluent of this column was collected in fractions of 4 ml and assessed for CRP-binding activity. An antigen complex of mol. wt 470 000 and a catalase antigen were identified by rocket immunoprecipitation assays. Immunoprecipitated catalase was visualised by adding hydrogen peroxide to the dried gel (Schönheyder and Andersen, 1984b).

**CRP-binding assay**

An enzyme-linked immunosorbent assay (ELISA) was developed for detection of CRP binding to aspergillus fractions. All determinations were done in duplicate, and in all experiments two identical sets of ELISA microtiter plates (Immunoplate 1, Nunc, Denmark) were processed, i.e., one set saturated with calcium for optimal CRP binding, and one exposed to Na$_2$EDTA for dissociation of calcium-dependent CRP binding.

Undiluted Sephadex G-200 fractions (100 µl) and serial ten-fold dilutions of these fractions in 0.5 M carbonate buffer, pH 9.6 (Voller et al., 1976), were added to the wells and left for 1 h at 37°C. After a wash in TBS-Tw buffer (0.05 M Tris, 0.15 M NaCl, 0.05% Tween 20, pH 7.4) 50 µl of CRP (3.5 g/ml of TBS-Tw) was added to each well and plates were left overnight at 4°C. This concentration of CRP was determined as optimal by using serial dilutions of CRP against one *A. fumigatus* CRP-binding fraction. After a wash in TBS-Tw, 50 µl of peroxidase-labelled rabbit anti-human CRP (Dakopatts) diluted 1 in 300 in TBS-Tw were added to each well and left for 2 h at 37°C. One set of ELISA plates was then rinsed in TBS-Tw and 50 µl of substrate—o-phenylenediamine dihydrochloride (OPD)—in citrate-phosphate buffer, pH 5.6, were added to each well and incubated for 5 min at room temperature. The reaction was stopped by adding 100 µl 1 M H$_2$SO$_4$ to each well.

The replica set of plates was washed in a Na$_2$EDTA containing buffer (0.05 M Tris, 0.15 M NaCl, 20 mM Na$_2$EDTA, Tween 20 0.05%, v/v, pH 7.4) and then the OPD substrate was added and allowed to react as described above. The absorbance of wells was read at 490 nm in absorbance units with a filter photometer (MR 590, Dynatech, USA). The calcium-dependent CRP binding was determined as the difference of means of absorbance of wells with calcium and wells washed with Na$_2$EDTA-containing buffer.

**Anti-phosphorylcholine assay**

A monoclonal antibody against phosphorylcholine from a mouse hybridoma was kindly provided by Dr U. Skov Sørensen, Statens Seruminstitut, Copenhagen (Sørensen et al., 1984). This was used for demonstration of phosphorylcholine ligands in aspergillus fractions as follows. Wells coated with *A. fumigatus* fractions as described above were incubated overnight at 4°C with 50 µl of mouse anti-phosphorylcholine antibody diluted 1 in 1000 in phosphate-buffered saline, pH 7.4, containing Tween 20 0.05%, v/v (PBS-Tw). After a rinse in PBS-Tw, 50 µl of peroxidase-labelled rabbit anti-mouse immunoglobulin (Dakopatts) diluted 1 in 400 in PBS-Tw were added and left for 2 h at 37°C. After a rinse in PBS-Tw, OPD substrate was added and the enzymatic reaction was allowed to proceed for 5 min. All determinations were done in duplicate and wells with PBS-Tw added instead of anti-phosphorylcholine served as blanks. The phosphorylcholine activity was measured by the difference of means of absorbance at 490 nm with anti-phosphorylcholine added and with PBS-Tw added instead of mouse antibody.

**Results**

**CRP binding assay**

The CRP-binding activities of the *A. fumigatus* fractions were determined with and without calcium and the differences were taken to indicate the calcium-dependent binding activities. Investigation of four separate batches of *A. fumigatus* ruptured mycelium exhibited the same pattern of calcium-dependent CRP-binding activity. Results from one representative experiment are shown in the figure. Calcium-dependent CRP-binding activity was detected in all four eluates (4 M, 2 M, 0.15 M and H$_2$O) obtained by hydrophobic interaction chromatography, the highest activity being asso-
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Associated with the H₂O eluate. The absorbance of all fractions in the absence of calcium was below 0.5, which indicated little calcium-independent CRP-binding activity of A. fumigatus fractions.

By gel filtration of the H₂O eluate, the most prominent calcium-dependent CRP binding activity was found in fractions with a mol. wt of c. 500 000. This peak of CRP-binding activity coincided with a distinctive A₂₈₀ peak. Less CRP-binding activity was associated with a fraction of c. 150 000–50 000 mol. wt which did not coincide with an A₂₈₀ peak. In the H₂O eluate the catalase-antigen fraction with mol. wt 250 000 (Schönhheyder and Andersen, 1984b) did not show CRP-binding acti-

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**Fig.** Calcium-dependent CRP binding (---) and anti-phosphorylcholine reactivity (-----) in A. fumigatus fractions as measured by ELISA (A₄₉₀nm). The first chromatographic step was hydrophobic interaction chromatography producing four eluates (4 M, 2 M, 0.15 M and H₂O) of increasing hydrophobicity. Eluates were subsequently processed on a Sephadex G-200 column and effluents were collected in fractions of 4 ml. The A₄₉₀nm profile (-----) and mol.-wt markers—IgM, IgG and human serum albumin (Alb)—are indicated.
vity. The 2 M eluate exhibited two distinct peaks of CRP-binding activity, one with a mol. wt of 500 000 and one of c. 150 000. Only the highest mol. wt peak was associated with high $A_{280}$ activity. A 470 000 mol. wt antigen complex (Schönheyder and Andersen, 1984b) eluted in a prominent $A_{280}$ peak and was not associated with conspicuous CRP-binding activity. In the 4 M eluate all fractions showed some CRP-binding activity whereas the 0-15 M eluate had little activity.

**Anti-phosphorylcholine assay**

A monoclonal antibody against phosphorylcholine was used for demonstration of phosphorylcholine ligands in *A. fumigatus* fractions. The anti-phosphorylcholine assay for all eluates showed an activity profile very similar to that of the calcium-dependent CRP binding (figure). It should especially be noticed that CRP-binding maxima in H$_2$O and 2 M eluates coincided with high phosphorylcholine activity. However, some discrepancies were observed. In the 2 M and 4 M eluates the phosphorylcholine ligand was detectable by anti-phosphorylcholine beyond fraction 75, where no or little CRP binding was seen.

**Discussion**

Combined hydrophobic interaction chromatography and gel filtration has previously been found to resolve several distinct *A. fumigatus* antigens (Schönheyder and Andersen, 1984a and b) and the present study demonstrates that a number of these fractions exhibit calcium-dependent CRP-binding activity. The fraction with the highest binding capacity had a mol. wt of c. 500 000. Phosphorylcholine has been demonstrated to be the determinant group in many microbial C-reactive substances (Volanakis and Kaplan, 1971; Baldo et al., 1977; Gotschlich et al., 1982). The anti-phosphorylcholine reactivity profile corresponded with the CRP-binding activity, indicating that some C-reactive substances from *A. fumigatus* contain phosphorylcholine. It should also be mentioned, however, that phosphorylcholine was detected in some fractions without conspicuous CRP binding.

In the 4 M eluate, fractions with a broad range of mol. wt had CRP-binding activity and this activity did not correspond closely with the presence of phosphorylcholine. Thus it is possible that in these fractions CRP was binding to other ligands, i.e., galactose residues or complexes containing nucleic acids (Robey et al., 1984). The results suggest that *A. fumigatus* C-reactive substance is not a single substance but is composed of different molecules with a broad range of mol. wt and some sharing the phosphorylcholine group.

It should be noticed that the CRP-binding fractions were different from fractions previously found to exhibit the highest antigenic activity in man. Antibodies to the 470 000-mol. wt fraction occur in the majority of healthy subjects and in patients with aspergillus-related disorders. This antigen complex contains protein and carbohydrate (Schönheyder and Andersen, 1984b), but it was found neither to bind CRP nor to contain phosphorylcholine and it seems therefore to be unrelated to C-reactive substance. A catalase-containing fraction of the H$_2$O eluate also induces a strong immune response in patients with aspergillosis (Schönheyder and Andersen, 1984b) and this fraction likewise appeared to be unrelated to aspergillus C-reactive substance.

The high-mol.-wt fraction of the H$_2$O eluate identified as the major CRP-binding fraction may be of particular interest for CRP-mediated reactions towards *A. fumigatus*.

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