Purification of infectious canine parvovirus from cell culture by affinity chromatography with monoclonal antibodies

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Immuno affinity chromatography with virus neutralizing monoclonal antibodies, directed to the haemagglutinating protein of canine parvovirus (CPV) was used to purify and concentrate CPV from infected cell culture. The procedure was monitored by testing the respective fractions in an infectivity titration system, in an ELISA, in a haemagglutination assay and by negative contrast electron microscopy to quantify CPV or CPV antigen. The degree of purification was further estimated by testing the fractions for total protein content in a colorimetric method, for bovine serum albumin content in an ELISA and by SDS-PAGE. Over 99% of the contaminating proteins proved to be removed, and 20% or 70-90% of infectious CPV or CPV antigen, respectively, was recovered.

Canine parvovirus; Purification; Immuno affinity chromatography; Monoclonal antibody

Introduction

Canine parvovirus (CPV) is a nonenveloped single-stranded DNA virus that may cause an occasionally fatal enteritis-panleukopenia syndrome in dogs (Appel et al., 1979; Gagnon and Povey, 1979; Johnson and Spradbrow, 1979; Osterhaus et al., 1980). Purified CPV derived from culture supernatant of CPV-infected cells has been used for biochemical (Paradiso, 1982; Rhode, 1985), ultrastructural (Paradiso, 1982), immunological (Parrish et al., 1982; Burtonboy et al., 1982; Parrish and Carmichael, 1983) and serological (Fiscus et al., 1985) studies. Conventional methods used for the purification of paroviruses from cell culture supernatant.

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including CaCl₂ precipitation (Tattersall et al., 1976), polyethylene glycol precipitation (Paradiso, 1981), banding in isopycnic CsCl gradients (Tattersall et al., 1976; Paradiso, 1981; Parrish et al., 1982) and haemadsorption elution (Osterhaus et al., 1980; Burtonboy et al., 1982), are multi-step and time-consuming procedures which are difficult to apply to large-scale purification. Immuno affinity chromatography (IAC) using monoclonal antibodies (MoAbs) has been shown to be a suitable method for the purification of viruses or viral antigens (Chong et al., 1985; Diaco et al., 1986). Virus purification from tissues infected with Aleutian mink disease virus, a member of the Parvoviridae family, by IAC using a polyclonal antibody preparation has been described (Kenyon et al., 1973).

In the present study, we describe the purification and concentration of CPV by IAC using mouse MoAbs directed against the haemagglutinating (H) protein of CPV, immobilized to Sepharose. The method proved efficient and relatively simple to concentrate and purify CPV from infected cell culture.

Materials and Methods

Virus

The continuous canine cell line A-72 (Binn et al., 1980) was cultivated in Eagle’s medium supplemented with Hanks’ salts, lactalbumine hydrolysate (5.13 g/l), natriumcarbonate 350 mg/l, glutamine 0.002 M, 100 IU penicillin/ml, 100 μg streptomycin/ml, and 10% fetal calf serum in 850 cm² roller bottles (Costar) rolling at 0.4 rpm at 37°C. Complete monolayers were trypsinized, washed and resuspended in phosphate-buffered saline (PBS) (pH 7.4).

The resuspended cells (10⁷ per ml) were infected with CPV (strain C780916; Carmichael et al., 1981) at a m.o.i. of about 10⁻³ TCID₅₀ per cell, by incubating the cells with CPV in a rolling bottle for 90 min at 37°C. After low speed centrifugation, the cells were resuspended in culture medium and cultivated at 37°C in roller bottles. Four to five days postinfection, the cultures exhibiting cytopathic changes were frozen and thawed twice. The suspension was clarified by centrifugation for 15 min at 2000 × g and used as starting material for IAC.

Production and characterization of MoAbs

A panel of hybridomas producing MoAbs reactive with the H protein of CPV was generated essentially as previously described for the generation of MoAbs against poliovirus (Osterhaus et al., 1981). Briefly, spleen cells from BALB/c mice immunized with CPV were fused with a mouse myeloma cell line. Hybridomas producing MoAbs, positive in haemagglutination inhibition (HI) and virus neutralization (VN) tests (see below), were selected and single cell cloned twice. Mouse ascitic fluids were produced in BALB/c mice. Ig was isolated from mouse ascitic fluids by precipitation with ammonium sulphate and affinity chromatography using Protein A-Sepharose (Pharmacia Fine Chemicals, Uppsala, Sweden) (Seppälä et
al., 1981; Manil et al., 1986). Purified antibodies were conjugated to horseradish peroxidase (HRP) according to standard methods (Nakane and Kawaoi, 1974).

The epitope specificity of the MoAbs was determined in a competition ELISA system on CPV-coated microtiter plates (Titertek, Flow Laboratories). For this ELISA, CPV was coated in an optimal dilution in 0.1 M sodium carbonate buffer (pH 9.6). After washing with demineralized water containing 0.05% Tween 80 (Merck, Schuchardt, F.R.G.) (DWT), 100-μl volumes of 10-fold dilutions of unconjugated antibodies (ascitic fluids) were made in PBS containing 1% bovine serum albumin (BSA) (Organon Technika, Oss, The Netherlands) and 0.05% Tween 80 and incubated for 30 min at room temperature. The plates were washed with DWT and homologous or heterologous HRP-conjugated MoAbs were added at the appropriate dilutions and incubated for 30 min. The plates were washed twice and 100 μl of substrate solution (0.1 mg/ml tetramethylbenzidine [TMB]) and 0.003% H₂O₂ in 0.1 M NaAc buffer (pH 5.5) was added to each well. After 10 min at room temperature, 100 μl of 2 M H₂SO₄ was added to stop the reaction. The absorbance at 450 nm was read in a Titertek Multiskan (Titertek, Flow Laboratories).

Isotypes of the MoAbs were determined in an ELISA system. For this ELISA, Protein A-purified anti-CPV MoAbs were diluted in PBS containing 1% BSA (Organon Technika) and 0.05% Tween 80 (Merck) and incubated for 30 min in wells of CPV-coated microtiter plates (Titertek, Flow Laboratories). After washing, 100 μl of rabbit anti-mouse antibody preparations, specific IgG1, IgG2a, IgG2b, IgG3, IGM or IgA (Litton Bionetics, Charleston, U.S.A.), were added in 1:1000 dilutions and incubated for another 30 min. After washing, 100 μl swine anti-rabbit HRP-conjugated antibody (Dakopatts, Denmark) was added in a 1:1000 dilution and incubated for 30 min. The plates were washed and developed with TMB substrate as described above.

IAC chromatography

Protein A-Sepharose IAC-purified MoAbs were dialysed against coupling buffer and coupled to CNBr-activated Sepharose 4-B (20 mg McAb to 10 ml gel volume) as recommended by the manufacturer (Pharmacia Fine Chemicals). The column was equilibrated with TEN buffer (20 mM Tris, 1 mM EDTA, 0.15 M NaCl adjusted to pH 7.8 with HCl). To 90 ml of clarified culture supernatant from CPV-infected cells 10 ml of a 10 times concentrated TEN buffer was added together with solid NaCl to a final concentration of 0.65 M NaCl.

CPV was allowed to bind overnight by running the column at low speed (20 ml/h). After washing the column with 20–30 ml TEN buffer supplemented with 0.5 M NaCl, the virus was eluted with TEN buffer containing 3 M NH₄SCN and dialysed against PBS. All steps were carried out at 4°C.

Effluent (unbound material) and eluate fractions were tested in SDS-PAGE and in the respective biological assays (see below). Protein concentrations were determined by the colorimetric method of Bradford (Bradford, 1976) adapted to a microtiter system.
Double antibody sandwich ELISA

For the detection and quantification of CPV antigen, a double antibody sandwich ELISA was used. Microtiter plates were coated with two Protein A-purified MoAbs (H-1 and H-2) which recognize two different epitopes (see below), 100-μl volumes containing 250 ng of each MoAb were incubated in microtiter plates for 16 h at 4°C. After washing with DWT, 150-μl volumes of a 1% BSA solution containing 0.1% NaN₃ in distilled water were added. The plates were sealed and stored at 4°C until use.

Plates were washed with DWT and 50-μl volumes of 3-fold serial dilutions of CPV suspensions were added simultaneously with 50 μl of HRP-conjugated MoAbs H-1 and H-2. After incubation for 30 min at 37°C, the plates were washed twice and developed with TMB substrate.

HA and HI tests

HA and HI tests were performed in 96-well round bottom microtiter plates as described (Appel et al., 1979; Carmichael et al., 1980; Osterhaus et al., 1980a), using African green monkeys (Cercopithecus aethiops) erythrocytes.

Infectivity and VN tests

For infectivity titrations, 1.8-ml volumes of 10-fold dilutions of CPV preparations in PBS were mixed with 0.9-ml volumes of a suspension containing 8 × 10⁵ A-72 cells per ml in PBS. The mixtures were incubated for 90 min at 37°C and gently agitated every 15 min. Cells were pelleted by low speed centrifugation, resuspended in 5.4-ml culture medium and divided over eight Lab Tek chambers (Miles, Napperville, U.S.A.). After 4 days of cultivation at 37°C, cells were fixed with ethanol at –70°C and examined by immuno fluorescence using one of the Protein A-purified anti-CPV MoAb and a FITC-conjugated swine anti-mouse preparation (Nordic, Tilburg, The Netherlands). For VN tests, the same procedure was followed: after incubation for 1 h at 20°C of 100 TCID₅₀ CPV with dilutions of the respective MoAbs.

Electron microscopy

For negative contrast examination, small drops of the samples to be tested were placed on formvar/carbon-coated grids and contrasted with 2% phosphotungstic acid (adjusted to pH 2.5 with KOH). The preparations were examined in a Philips EM 400T transmission electron microscope.
BSA-ELISA

Cell culture harvests and column fractions were tested for BSA content in an ELISA using an anti-BSA mouse MoAb (IgG1) coupled to HRP (Avest and Osterhaus, submitted for publication). In short, samples were coated to microtiter plates in serial 10-fold dilutions in sodium carbonate buffer (pH 9.6) for 2 h at 37°C. After washing, 100 μl of a 1:3000 dilution of the anti-BSA conjugate in PBS containing 0.05% Tween and 0.25% ovalbumine (Sigma Chemical, St. Louis, MO). Plates were washed and developed with TMB as described above. For the determination of BSA content, ELISA values were compared with those obtained with standard dilutions of a BSA (Boseral, Organon, Oss, The Netherlands) preparation.

SDS-PAGE

Proteins were analysed on 12% polyacrylamide slab gels (Laemmli, 1970). Gels were silverstained (Wray et al., 1981). Proteins of known molecular weights (Bio-Rad, Richmond, U.S.A.) were included to estimate the molecular weights of viral proteins.

Results

MoAbs

A panel of four stable hybridomas producing HI-positive VN MoAbs (H-1, H-2, H-3 and H-4) were selected. These MoAbs also reacted with the haemagglutinin of feline panleukopenia virus (FPV) and recognized at least two different epitopes as was demonstrated in the competition ELISA (Table 1). The isotypes of the four MoAbs proved to be IgG2a, IgG2b, IgG1 and IgG2b, respectively (Table 1). MoAbs H-1 and H-2 were used in further IAC purification studies.

IAC

Two small Sepharose 4-B columns (bed volume 1.0 ml) were prepared with MoAbs H-1 and H-2, respectively, which were shown to recognise different epitopes (Table 1). When 10 ml of CPV-infected culture material (HA titer 1024; double antibody sandwich ELISA titer 30) were passed through either of the columns, no infectious CPV or CPV antigen (ELISA/HA) could be detected in the effluent material of the columns. Upon elution with 3 M NH₄SCN, CPV antigen could be eluted from both columns with about the same efficiency, as judged from ELISA and HA titrations carried out with the respective fractions collected from both columns. The profiles obtained in HA and ELISA coincided well with those obtained with the 10-ml bed volume column described below (not shown). To investigate further the potential of the IAC procedure for the purification and con-
TABLE 1
Properties of anti-CPV MoAbs.

<table>
<thead>
<tr>
<th>MoAb</th>
<th>Log^{10} titer in competition ELISA</th>
<th>isotype</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HRP-conjugate</td>
<td></td>
</tr>
<tr>
<td></td>
<td>H-1</td>
<td>H-2</td>
</tr>
<tr>
<td>H-1</td>
<td>2</td>
<td>-</td>
</tr>
<tr>
<td>H-2</td>
<td>-</td>
<td>4</td>
</tr>
<tr>
<td>H-3</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>H-4</td>
<td>4</td>
<td>-</td>
</tr>
</tbody>
</table>

* -- ≤ 1.

The concentration of CPV from cell culture material. A 10-ml bed volume affinity column was prepared with MoAb H-1. When 100 ml of the CPV-infected culture material were passed through this column again, neither CPV nor CPV antigen could be detected by HA, ELISA or infectivity assays in the effluent (Fig. 1), and no or only few irregular parvovirus-like particles were observed by negative contrast electron microscopy (Fig. 2). The effluent contained relatively high concentrations of contaminating proteins (up to about 4 mg/ml) including BSA as was also visualized by SDS-PAGE in the initial effluent fractions (Fig. 3), which gradually de-

Fig. 1. Purification of CPV by IAC. Arrow indicates change of buffers. Fractions were collected and monitored for ELISA reactivity (○—○), haemagglutination activity (●—●), protein content, (△—△), BSA content, (■—■) and infectivity (△—△).
clined upon washing the column with the starting buffer (Fig. 1). CPV and/or CPV antigen could be readily recovered from the column by elution with 3 M NH₄SCN, as was shown by titration in infectivity assays, HA, and ELISA of the eluate fractions (Fig. 1, Table 2). On the basis of the titers found in the respective assays, it was concluded that a recovery of about 20% of infectious CPV, and of about 70–90% of CPV antigen could be achieved, and that maximal concentration effects of about four and 16 times, respectively, were obtained. Analysis of eluate fractions by negative contrast electron microscopy confirmed the presence of large numbers of predominantly full virus particles, apparently free from contaminating membranous structures, in the fractions with the highest CPV and CPV antigen concentrations (Fig. 2). Total protein and BSA concentrations in these peak fractions measured about 40 and 8 µg/ml, respectively. In eluate peak fractions, only three protein bands could be visualized by SDS-PAGE at molecular weights corresponding to those of parvovirus proteins: 82.5·10^3, 67.5·10^3 and 65·10^3, respectively (Fig. 3).

The column could be used more than 20 times without apparent loss of binding capacity. This capacity proved at least three times higher than the amount of antigen used in the experiments mentioned above.

**Discussion**

In the present paper, we show that IAC, using a MoAb directed against the H protein of CPV, provides a rapid and relatively simple procedure for the purification and concentration of CPV from cell culture material. Analysis of the frac-
Fig. 3. Analysis of effluent (A) and eluate (B) fractions by SDS-PAGE. Gels were silverstained by the method of Wray et al. (1981). Marker proteins were used to estimate molecular weights (MW) of viral proteins (MW in Da × 10^-3).

The possible discrepancy observed between the results obtained with effluent fraction by EM on the one hand and ELISA or HA on the other hand may be explained by the relatively high cutoff values chosen (five times background), and the relative insensitivity of the test, respectively. The possible discrepancy between infectivity and EM results may be explained by the fact that at most only noninfectious or damaged virus particles were present in the effluent. The recovery of CPV antigen, as judged from HA and ELISA titrations was 70-90%, whereas the recovery of infectious CPV, as shown by infectivity titrations, proved to be about 20%. The loss of infectivity probably is a direct effect of the elution conditions used and the procedure of dialysis, followed by freezing and thawing before testing. This was confirmed by testing of CPV culture material.
TABLE 2
Quantitative data of IAC purification and concentration procedure.

<table>
<thead>
<tr>
<th></th>
<th>Culture supernatant</th>
<th>Effluent fractions</th>
<th>Eluate fractions</th>
</tr>
</thead>
<tbody>
<tr>
<td>Volume net (ml)</td>
<td>90</td>
<td>1.8</td>
<td>1.8</td>
</tr>
<tr>
<td>Protein concentration (µg/ml)</td>
<td>3825</td>
<td>&lt;15 - 3925</td>
<td>&lt;15 - 40</td>
</tr>
<tr>
<td>BSA concentration (µg/ml)</td>
<td>3000</td>
<td>2.6 - 3900</td>
<td>0.15 - 8</td>
</tr>
<tr>
<td>HA titer</td>
<td>1024</td>
<td>0</td>
<td>0 - 16.000</td>
</tr>
<tr>
<td>ELISA titer</td>
<td>30</td>
<td>0</td>
<td>0 - 730</td>
</tr>
<tr>
<td>Infectivity (log_{10} TCID_{50}/ml)</td>
<td>4.9</td>
<td>&lt;1</td>
<td>2.2 - 5.5</td>
</tr>
<tr>
<td>EM</td>
<td>+**</td>
<td>-</td>
<td>+++++</td>
</tr>
</tbody>
</table>

* Estimated total recovery.
** Arbitrary units indicating the number of virus particles observed: + = small numbers; +++++ = large numbers (Fig. 3).

directly treated in the same way (not shown). This reduction in infectivity might partly be overcome by using a procedure of NH₄SCN gradient elution in combination with gel filtration. Over 99% of the contaminating proteins were shown to be removed in the IAC procedure, by determination of the reduction of the BSA content in an ELISA, of the total protein content by colorimetry and by comparing the different fractions in SDS-PAGE. Since BSA is the major contaminating protein present in fetal calf serum used in our cell culture system, it may be concluded from our data that the determination of the BSA content may be considered a proper method to validate the purification procedure. The small BSA peak present in the eluate fractions containing most intact CPV particles and infectivity may be explained by a nonspecific binding of BSA to either the affinity column or directly to CPV.

The use of the IAC method described for the purification and concentration of CPV may not only be of interest for the production of purified CPV for biochemical, ultrastructural and immunological studies, but may also be applied to produce infectious CPV or CPV antigen to be used for vaccination purposes or in diagnostic tests. The cross-reactivity of the respective MoAbs with FPV would also offer the opportunity to use these for the same applications with this virus.

A similar approach of IAC using MoAb would also be of interest for the purification of human parvovirus B19, e.g., from viraemic plasma, since no in vitro cultivation systems are available for this virus at present (Siegl, 1984).

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