Protection against *Streptococcus equi* infection by monoclonal antibodies against an M-like protein

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We have developed an in vivo passive transfer assay using mice to identify monoclonal antibodies (mAbs) which offer protection against *Streptococcus equi* infection. The assay was developed using serum antibodies collected from horses convalescing from strangles. In this study, we show that a preparation of M-like protein, acid-extracted from *S. equi*, affords 80% protection to mice immunized with it. A number of mouse mAbs directed against a preparation of M-like protein were then assessed for their ability to passively protect mice against challenge with a lethal dose of the bacteria. Two mAbs, 1D10 and 2A6, were shown to be highly protective. It was also demonstrated, by means of a competitive enzyme immunoassay, that these mAbs recognized different epitopes in the preparation. Examination of a dose–response curve for mAbs 1D10 and 2A6 revealed that optimal levels of protection were achieved using 1 mg of either 1D10 or 2A6, or 0.5 mg 1D10 and 0.5 mg 2A6 given together. Immunological reactivity of these mAbs with a preparation of M-like protein showed that the antigens they recognized were comparable in size to some of the antigens recognized by convalescent horse serum antibodies. The role of immunoglobulin isotype in conferring protection is discussed.

Introduction

*Streptococcus equi* (a group C streptococcus) causes strangles, a contagious disease of the upper respiratory tract in horses characterized in the initial stages by nasal discharge and fever, and eventually by swelling of the regional lymph nodes due to abscess formation. In severe cases, the infection may become disseminated with internal and external abscess formation and may ultimately result in death [Commonwealth Serum Laboratories (CSL) Veterinary Handbook, 1979]. Mounting evidence in the literature indicates that immunity to *S. equi* is local rather than systemic and that it is mediated by mucosal antibodies (Srivastava & Barnum, 1983, 1985; Timoney & Eggers, 1985; Timoney & Galan, 1985; Galan & Timoney, 1985). The importance of local immunity in conferring protection against group A streptococcal infections is also well documented (Bessen & Fischetti, 1985a, b). Virulence in group A streptococci is mainly associated with the presence of M-protein, a trypsin-labile component with an α-helical coil configuration which spans the bacterial capsule and cell wall (Fox, 1974; Phillips et al., 1981). The M-protein renders the bacterium resistant to non-immune phagocytosis (Bisno, 1979; Peterson et al., 1979). The existence of an M-like protein has also been demonstrated in *S. equi* (Woolcock, 1974). However, the M-like protein has not been conclusively shown to be the major virulence factor of this species, although bacterial extracts containing it confer immunity (Woolcock, 1974; Timoney & Galan, 1985).

Although only one antigenic type of *S. equi* has been observed in the field, vaccines prepared by conventional procedures have proved to be relatively ineffective (Srivastava & Barnum, 1981, 1983, 1985; Timoney & Eggers, 1985; Woolcock, 1974; Jorm, 1990). Our aim is to produce anti-idiotypic antibodies which resemble the protective antigen(s) of *S. equi* and to evaluate their use as potential vaccines. To this end our strategy included the production of protective monoclonal antibodies (mAbs) against *S. equi* (Ab1) followed by the production of anti-idiotypic antibodies (Ab2) using these protective mAbs against *S. equi* (Ab1) as immunogens.
In this study, the ability of a preparation of partially purified M-like protein to protect mice against a challenge with a lethal dose of *S. equi* was evaluated. Subsequently, mAbs raised against a preparation of M-like protein were assessed for their ability to protect mice against challenge with *S. equi* by means of a passive transfer assay which was developed during the course of this investigation.

**Methods**

*S. equi* isolate, growth of bacteria and determination of cell numbers in cultures of *S. equi*. A vaccine isolate of *S. equi*, 572F (CSL), was used throughout this study. The isolate was obtained from a submaxillary lymph node of a horse suffering from strangles. *S. equi* was grown in 'strangles' medium, prepared according to the CSL code of manufacture and containing (g l−1): Neopeptone (Difco), 10; New Zealand Amine B (Sheffield Products, New York, USA), 10; yeast extract (Difco), 20; glucose (AnalarR), 30; NaH₂PO₄, 2H₂O, 4.5; Na₂CO₃, 2.5, and phenol red indicator. After heating the medium to about 50 °C to dissolve the components, the pH was adjusted to 7.6 with 1 M-NaOH.

Lyophilized stocks of the above isolate were reconstituted and used to inoculate pre-cultures. Pre-cultures (50–100 ml) of *S. equi* grown at 37 °C overnight in 50% (v/v) 'strangles' medium and 50% (v/v) veal infusion (CSL) were used as inocula (10%, v/v) for large (500 ml–2 l) cultures which were grown at 37 °C for 4–6 h in 'strangles' medium.

To determine the number of cells in a test culture, the OD₅₇₀ was measured after 4–6 h growth and the number of cells ml⁻¹ was estimated from a standard curve previously constructed by plotting the OD₅₇₀ of samples of a culture taken at half-hourly intervals versus the number of viable cells in the culture at those time points. Viability was determined by plating out serial dilutions of the culture onto horse-blood agar plates (CSL) and counting the number of colonies following incubation at 37 °C for 24 h. Values obtained from the standard curve were in good agreement with the assessed viability count.

**Preparation and partial purification of hot-acid-extracted M-like protein.** The procedure used was essentially that described by Woolcock (1974). *S. equi* cells, washed at least three times with 0.15 M-NaCl, were resuspended in a minimal volume of the same solution. The pH of the suspension was brought down to 2.4 with 1 M-HCl and the suspension was extracted at 95 °C for 15 min. Cells and cell debris were sedimented and removed by centrifugation at 1000 g for 5 min. Protease inhibitors [PMSF (Sigma), final concentration 1 mM; leupeptin (Sigma), final concentration 1 μg ml⁻¹; and pepstatin (Sigma), final concentration 3 μg ml⁻¹] were added to filtered (Whatman fast) extract. The extract was neutralized with 1 M-NaOH.

The extract was then concentrated by salt fractionation with ammonium sulphate at 60% saturation. A precipitate was first obtained at 30% saturation and removed by centrifugation; the supernatant fraction was then made up to 60% salt saturation. Following solubilization and dialysis against 0.05 M-sodium acetate buffer (pH 4.5), the harvested material was loaded onto a carboxymethylcellulose (CM-cellulose) (Whatman) column. The column was washed with the same buffer and the M-like protein was eluted with 0.02 M-HCl. Peak fractions were pooled, the salt concentration adjusted to 0.15 M-NaCl and the pH adjusted to 7.0 with 1 M-NaOH. Protein concentration was determined by the Lowry protein assay.

**SDS-PAGE and Western immunoblotting.** Preparations of M-like protein were analysed by gel electrophoresis on 12.5% SDS-polyacrylamide slab gels (Bio-Rad apparatus). Electrophoresis was carried out at room temperature at a constant current of 20 mA for about 3 h. Proteins were visualized with silver stain (20 μg protein per lane) or after Western immunoblotting (75 μg protein per lane). For immunoblotting, electrophoresed proteins were transferred to nitrocellulose filters at 100 V for 1 h at 4 °C in a Bio-Rad immunoblot apparatus. The filters were blocked overnight at 37 °C with blocking buffer [2% w/v BSA and 0.1% Tween-20 in PBS (10 mM-sodium phosphate, 0.15 M-NaCl, pH 7.4)]. Horse serum or mouse ascites diluted 1:150 in blocking buffer were incubated with the filters for 1 h at 37 °C. The filters were then incubated for 1 h at 37 °C either with goat anti-horse IgG (heavy and light chains) immunoglobulins conjugated to horseradish peroxidase (HRP) (KPL) at a dilution of 1:1000 in blocking buffer, or with goat anti-mouse IgG, IgA and IgM (heavy and light chains) immunoglobulins conjugated to HRP, at the same dilution. The reaction was detected by the addition of 3-chloro-1-naphthol (3 mg ml⁻¹ in methanol) and H₂O₂ (12 μl 30%, w/v) in 20 ml PBS.

**Horse serum antibodies and passive haemagglutination.** Horse serum antibodies were obtained from three horses convalescing from strangles. Blood was collected from horses 1 and 2 early in convalescence following the worst of the clinical manifestations of the disease. These horses had contracted strangles during a natural outbreak of the disease. Horse 3 was experimentally infected with 7 × 10⁶ cfu of *S. equi* strain 572F on each of two occasions with a 9 month interval between infections. Blood was collected 7 weeks after the primary infection and 4 weeks after the secondary infection. Horses 1 and 2 had previously been vaccinated twice at a 5 year interval with an acellular strangles vaccine [hot-acid extract (HAE) of *S. equi* isolate 572F cell walls adsorbed onto aluminium hydroxide adjuvant (CSL)], whereas horse 3 had not been vaccinated and had had no known previous contact with the bacterium.

Anti-*S. equi* antibodies were detected by passive haemagglutination. A packed cell suspension of fresh, normal sheep red blood cells was fixed with pyruvic aldehyde [7.4% (v/v) in PBS, pH 7.2] in a ratio of 1:6 by gentle stirring for 24 h at 4 °C. The cells were centrifuged (2000 g, 10 min), washed three times in PBS and incubated with an equal volume of tannic acid (4% w/v) at room temperature for 10 min. The tanned red blood cells were washed again as above. To sensitize them, the tanned red blood cells were incubated at room temperature for 10 min with 4 vols partially purified M-like protein preparation (3% w/v), centrifuged, washed with normal rabbit serum [0.5% (v/v) in PBS] and finally resuspended in an equal volume of normal rabbit serum.

Passive haemagglutination was carried out on enzyme immunoassay (EIA) plates (96 round-bottomed wells per plate, Dynatech); test sera diluted with normal rabbit sera [1% (v/v) in PBS] were added at serial doubling dilutions to the plates (100 μl per well). Sensitized red blood cells (10 μl) were added and the plates were incubated at room temperature for 3 h. The positive control consisted of horse serum of a known high antibody-titre to *S. equi* and the negative control serum was obtained from a horse with no known previous contact with *S. equi* at the time of blood collection.

**Immunization of mice for mAb production.** Two doses of 50 μg of a partially purified M-like protein preparation in PBS, emulsified with equal volumes of either Freund's complete adjuvant (FCA) or Freund's incomplete adjuvant (FIA) were given intraperitoneally (i.p.) 2 weeks apart to raise anti-M-like protein antibodies in 8–12-week-old BALB/c mice. Mice were eye bled 10 d after secondary immunization. Four days before cell fusion, mice were given an intravenous injection of 50 μg M-like protein in PBS.

**Fusion of mouse spleen cells with mouse myeloma cells.** Spleen cells from mice with the highest serum titres of anti-M-like protein antibody were fused with Sp 2/0 or NS-1 myeloma cells (Stahli et al., 1980). The ratio of spleen cells to myeloma cells was 5:1. After the addition of 0.8 ml 50% (w/v) PEG (gas chromatography grade, Merck)
to the combined spleen/myeloma cell pellet, the fusion mixture was made up to 300 ml in RPMI-1640 medium (CSL) containing 100 μM-hypoxanthine, 0-4 μM-aminopterin and 16 μM-thymidine (HAT), and 1 ml was dispensed into each well of twelve 24-well plates. Hybridomas were screened for the production of M-like protein antibodies by EIA (see below) when they reached 30-40% confluence. Antibody-positive hybridoma supernatant fluids were further characterized by Western immunoblotting. The hybridomas were cloned at least twice by limiting dilution at concentrations of 2 cells per well, 1 cell per well and 0.5 cell per well.

Production of ascites. At least 5 × 10⁶ hybridoma cells in 0.5 ml RPMI-1640 medium were injected (i.p.) into 8–12-week-old BALB/c mice which had been primed with 0.5 ml Pristane [2,6,10,14-tetramethyl pentadecane (Aldrich)] 2–10 d earlier (Galfre & Milstein, 1981).

Immunoglobulin purification, concentration and isotyping. Immunoglobulins were purified from mouse sera, ascites fluid or hybridoma supernatant fluids on a Protein A-Sepharose CL-4B affinity column (Ey et al., 1978). The bound fractions were eluted and concentrated by ultrafiltration on an XM-100 membrane (Amicon). Immunoglobulins from horse sera and IgM from mouse ascites fluid were purified by treatment with ammonium sulphate at 45% saturation. Difficulties were encountered with the purification of mouse IgM immunoglobulins in that they became partially insoluble after salt fractionation. Mouse immunoglobulin classes and isotypes were determined using an EIA kit (CSL) with mouse IgM, IgG1, IgG2a and IgG2b (Miles) as standards.

mAb conjugation to HRP. In order to determine the epitope specificity of mAbs 1D10 and 2A6, HRP-conjugates of these mAbs were prepared. To 5 mg HRP (type VI, Sigma) dissolved in 1 ml freshly prepared 0.3 M-NaHCO₃, 25 μl 0.32% paraformaldehyde was added and the solution was stirred at room temperature for 30 min. NaO₂ (1 ml, 0.04 M) was added and the solution was again stirred at room temperature for 30 min. Ethylene glycol (1 ml, 0.16 M) was then added and the solution stirred at room temperature for 1 h after which it was dialysed against 1 litre 0.01 M-Na₂CO₃ (pH 9.5) overnight at 4°C. About 10 mg IgG in 1 ml sodium carbonate buffer was added and the mixture incubated for 2 h at room temperature after which time 5 mg NaBH₄ was added and the mixture incubated for 2 h at 4°C. The HRP-IgG conjugate was finally dialysed against PBS overnight at 4°C.

Enzyme immunoassays (EIA). Anti-M-like protein antibodies in mouse sera and hybridoma supernatant fluids were determined by EIA. Plates were coated with 2 μg per well of a preparation of partially purified M-like protein in 0.2 M-sodium bicarbonate/sodium carbonate buffer (pH 9.6), incubated for 3 h at 37°C and blocked with 0.1% casein and 2% (w/v) BSA in PBS (blocking buffer) for 1 h at 37°C. The plates were washed three times in 0.2% Tween-20 in PBS. Undiluted hybridoma supernatant fluid (100 μl per well) or serial twofold dilutions of test sera in blocking buffer (100 μl per well) were added. The plates were incubated for 1.5 h at 37°C then washed as above. Anti-mouse IgG, IgM and IgA immunoglobulins conjugated to HRP (100 μl per well) were added and were incubated at a dilution of 1:2000 in blocking buffer. The plates were again incubated for 1.5 h at 37°C and then washed. Equal volumes of 2,2’-azino-di(3-ethylbenzthiazoline-6-sulphonate) (ABTS; KPL) and H₂O₂ substrate (KPL) were thoroughly mixed and 100 μl immediately dispensed into each well. Plates were incubated for 10 min at 20°C and As₀.₅ recorded using a Titertek Multiskan MC (Flow Laboratories). Antibody titres were expressed as the reciprocals of the plates already incubated with 2A6. The plates were incubated for 1.5 h at 37°C, in both instances. The dilutions of conjugated antibodies used had previously been determined by EIA against the M-like protein. The plates were developed using the procedure described above.

Active immunization – challenge assays. Groups of 10 Swiss mice were immunized (i.p.) 2 weeks apart with 50 or 100 μg of either an HAE of S. equi or a partially purified preparation of M-like protein (in PBS) emulsified with an equal volume of FCA or FIA. On day 21, blood was collected from each mouse to determine anti-M-like protein serum antibody titres after which each mouse was challenged (i.p.) with 2 × 10⁶ or 5 × 10⁶ viable S. equi cells in 0.2 ml PBS.

A control group of 10 non-immunized mice was also challenged at the same time. The numbers of mice that died and the times at which they died were recorded over a period of 10 d post-challenge.

Passive antibody transfer assay. Immunoglobulins fractionated from horse serum (30-40 mg total protein per mouse) or mouse ascites fluid (0.5–4 mg IgG or IgM per mouse) were injected (i.p.) into Swiss mice on days 0, 1 and 2. Groups of 10 mice were used to assay each antibody. Two to three hours after the antibody injection on day 0, each mouse was challenged (i.p.) with a lethal dose of viable S. equi cells (2 × 10⁶–3 × 10⁶ cells per mouse). The numbers of mice surviving the challenge were recorded over 4 d post-challenge. Immune responses raised against the injected allogeneic or xenogeneic antibodies could interfere with the interpretation of the results after this time. Control mice received immunoglobulins from the serum of a horse which had not had any known previous contact with S. equi, or from ascites containing mAbs with specificity for a human immunoglobulin epitope. In all experiments at least 80% of the control mice died within 4 d post-challenge.

The ability of an antibody to protect mice was expressed either as the percentage protection or as the cumulative mortality index (CMI) defined as cumulative mortality in test group/cumulative mortality in control group. Thus, a CMI value of 1 indicates no protection whereas a value of 0 indicates full protection. CMI was used for ease of comparison of results in cases where passive transfer experiments were conducted on separate occasions.

Results

Hot-acid-extracted M-like protein: biochemical analysis and induction of protective immunity in mice

Partially purified M-like protein was prepared from S. equi by ion-exchange chromatography on a CM-cellulose column following extraction of the cells for 15 min at 95°C and pH 2.4 (Woolcock, 1974). The protein elution profile and SDS-PAGE polypeptide profile are shown in Fig. 1. The SDS-PAGE polypeptide profile revealed the heterogeneous nature of the preparation, with the presence of an intensely stained band of approximately 55 kDa and several other prominent bands between 10 and 35 kDa. Numerous weakly stained polypeptide bands were also apparent. The protein compositions of M-like protein preparations from different batches of
S. equi cells appeared to be very similar as visualized by SDS-PAGE and silver stain. The presence of a 55 kDa band in a preparation of M-like protein has also been reported by Timoney & Trachman (1985).

The preparation of M-like protein was assessed for its ability to induce protective immunity in mice against a lethal challenge with viable S. equi cells. Mice were immunized (i.p.) with 50 or 100 pg of the HAE of the partially purified M-like protein. The number of mice which survived subsequent challenge with a lethal dose of S. equi was recorded daily over a period of 10 d (Fig. 2). Of the mice immunized with 50 pg HAE, 60% were alive 10 d post-challenge with $5 \times 10^6$ viable S. equi cells. However, only 45% of those immunized with 100 pg HAE survived. Partially purified M-like protein afforded considerably better protection, with 80% of the immunized mice surviving for 10 d post-challenge at both dose levels.

Production and in vitro characterization of mAbs against a preparation of M-like protein

A short-term immunization schedule consisting of two immunizations given i.p. 2 weeks apart was adopted to raise anti-M-like protein antibodies. High serum-antibody titres ranging between 1/51200 and 1/204800 were achieved, suggesting that the M-like protein was highly immunogenic in mice. Three cell-fusions were carried out using the mice that showed the highest antibody titres and 24 hybridomas producing anti-M-like protein antibodies were derived. Approximately 60% of these produced antibodies of the IgM isotype, while the rest produced IgG, the majority of these being of the IgG1 isotype.

Components of the M-like protein preparation which were recognized by the mAbs were identified by Western immunoblotting. The results shown in Fig. 3 are representative of all the mAbs analysed and show an apparently heterogeneous pattern of epitope recognition despite the fact that the hybridomas were cloned at least twice to ensure monoclonality. Fig. 3 also shows that there are two major recognition patterns for these mAbs, one consisting of polypeptide bands predominantly in the 25-55 kDa range (banding pattern 1; lane B) and including the 46 and 41 kDa fragments of the M-like protein described by Galan & Timoney (1987), the other including 2-3 major low molecular mass components (7-15 kDa) as well as a 55 kDa polypeptide band (banding pattern 2; lane D). A comparison of the immunoblot profile of the mAbs with that of convalescent horse serum antibodies (lane F) revealed that mAbs displaying recognition banding pattern 1 recognized several polypeptide bands with $M_r$ values in the 25-55 kDa range which were also recognized by horse serum antibodies.

Development of an in vivo passive transfer assay for the identification of protective mAbs

In order to assess the ability of the mAbs to protect against S. equi infection, an in vivo passive transfer assay was developed in mice using serum antibodies harvested
Passive protection against S. equi

Induction of protective immunity in mice by hot-acid-extracted S. equi antigens. Groups of 10 Swiss mice were immunized (i.p.) 2 weeks apart with 50 or 100 μg of (a) HAE or (b) M-like protein in FCA or FIA. On day 21, each mouse was challenged (i.p.) with (a) 5 × 10⁶ or (b) 2 × 10⁶ c.f.u. viable S. equi cells in 0.2 ml PBS. A control group of non-immunized mice was challenged at the same time. The number of mice that died and the time at which they died was recorded over a period of 10 d post-challenge.

Fig. 3. Immunological reactivity of M-like protein with mouse mAbs and horse serum antibodies. Hot-acid-extracted M-like protein (75 μg) was electrophoresed on 12.5% SDS-polyacrylamide slab gels at 20 mA for 3 h at room temperature and transferred onto nitrocellulose filters at 100 V for 1 h at 4 °C in a Bio-Rad immunoblot apparatus. Pre-stained markers (10 μl, Bio-Rad) were also electrophoresed alongside each sample of M-like protein, transferred onto the same piece of nitrocellulose filter and immunoblotted. The filters were blocked with 2% BSA and 0.1% Tween-20 in PBS overnight at 37 °C. Mouse ascites fluids and horse serum, diluted 1:150 in blocking buffer, were incubated with the filters for 1 h at 37 °C. Goat anti-mouse IgG, IgA and IgM (heavy and light chains) immunoglobulins conjugated to HRP and goat anti-horse IgG (heavy and light chains) immunoglobulins conjugated to HRP were added at a dilution of 1:1000 in blocking buffer. The reaction was detected by the addition of 3-chloro-1-naphthol (3 mg ml⁻¹). Control serum obtained from a horse which had no known previous contact with S. equi and ascites fluids obtained from a murine cell line producing mAbs against human IgG showed no reaction (result not shown). Lanes: A, C, E, molecular mass markers; B, mAb 1D10 (banding pattern 1); D, mAb 7C1 (banding pattern 2); F, serum from a convalescing horse.

Passive protection of mice against S. equi infection by anti-M-like protein mAbs

Eight murine mAbs were assessed for their in vivo ability to protect mice against challenge with S. equi. Four of the mAbs displayed recognition banding pattern 1 on a Western immunoblot, i.e. reactivity with polypeptide bands predominantly in the 25-55 kDa range (Fig. 3), while the other four displayed recognition banding pattern 2, i.e. reactivity with the 55 kDa polypeptide band in addition to two to three polypeptide bands with M, values ranging between 7 and 15 kDa. The results shown in Table 1 demonstrate that the mAbs fall into three categories: those which afforded good protection (CMI, ~0.14-0.25), viz. mAbs 1D10 and 2A6; those which afforded some protection (CMI, 0.77), viz. mAb...
Fig. 4. Passive protection of mice by horse serum antibodies. Immunoglobulins fractionated from horse serum (30-40 mg total protein per mouse) were injected (i.p.) into groups of 10 Swiss mice on days 0, 1 and 2. Two or three hours after the antibody injection on day 0, each mouse was challenged with a lethal dose of viable *S. equi* cells [(a) 3 x 10⁶ and (b) 2 x 10⁶ c.f.u.] The number of mice surviving the challenge was recorded over 4 d post-challenge. Control mice (V) received immunoglobulins from a horse which had not had any known previous contact with *S. equi* (passive haemagglutination titre varying between 1/2 and 1/128). (a) Results obtained with serum from horses 1 (O) and 2 (●), which had previously been vaccinated and were naturally infected with *S. equi* (passive haemagglutination titre: 1/100000). (b) Results obtained with serum from horse 3, which had not been vaccinated previously and was experimentally infected with *S. equi* on each of two occasions. □_serum harvested after the primary infection; ■ serum harvested after the secondary infection (passive haemagglutination titres after the primary and secondary infections were 1/3200 and 1/512, respectively).

Table 1. Some characteristics and properties of the anti-M-like protein mAbs

<table>
<thead>
<tr>
<th>mAb</th>
<th>Isotype/ light chain</th>
<th>Western immunoblot pattern</th>
<th>Passive protection (CMI)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1D10</td>
<td>IgG1 κ</td>
<td>1</td>
<td>0.25</td>
</tr>
<tr>
<td>2A6</td>
<td>IgG1 κ</td>
<td>1</td>
<td>0.14</td>
</tr>
<tr>
<td>2D3</td>
<td>IgM κ</td>
<td>2</td>
<td>0.89</td>
</tr>
<tr>
<td>3D4</td>
<td>IgM κ</td>
<td>1</td>
<td>1.0</td>
</tr>
<tr>
<td>7C1</td>
<td>IgM κ</td>
<td>2</td>
<td>1.0</td>
</tr>
<tr>
<td>7D1</td>
<td>IgG1 κ</td>
<td>2</td>
<td>0.86</td>
</tr>
<tr>
<td>10C5</td>
<td>ND κ</td>
<td>2</td>
<td>0.77</td>
</tr>
<tr>
<td>10D3</td>
<td>IgM κ</td>
<td>1</td>
<td>0.93</td>
</tr>
</tbody>
</table>

ND, Not determined.

Establishment of a dose-response (protection) curve for mAbs 1D10 and 2A6

In order to achieve the maximum level of protection with the minimum amount of antibody, the two protective mAbs, 1D10 and 2A6, were assayed individually and together at various concentrations in an *in vivo* passive transfer assay. Dose–response curves were established using antibody concentrations of 0.5, 1.0, 2.0 and 4.0 mg per mouse (Fig. 5). The highest level of passive protection was achieved using 1 mg of either 1D10 or 2A6, or 0.5 mg 1D10 and 0.5 mg 2A6 given together. The level of protection attained with 2.0 or 4.0 mg 1D10 was less than that attained with 0.5 or 1.0 mg, respectively. When 1D10 and 2A6 were given together, the level of protection was greater than that achieved using 1D10 alone; however, it was not, in most cases, significantly different from the level achieved using 2A6 alone.

mAbs 1D10 and 2A6 recognize different epitopes in a preparation of partially purified M-like protein

The possibility existed that mAbs 1D10 and 2A6 had specificity for the same epitope on a particular protein present in a partially purified preparation of M-like protein. A competitive EIA was developed to investigate this possibility.

Serially diluted ascites fluids containing mAbs 1D10 or 2A6 were added to M-like-protein-coated EIA plates.
HRP-conjugated 2A6 was then added at a fixed dilution to wells which had already been incubated with 1D10, while HRP-conjugated 1D10 was added to wells which had been incubated with 2A6. The results (Fig. 6) showed that the $A_{405}$ values obtained for both mAbs were fairly constant, varying between 1.80 and 2.04 when HRP-conjugated 2A6 was added to wells pre-incubated with serial dilutions of 1D10 (Fig. 6a) and between 0.92 and 1.20 when HRP-conjugated 1D10 was added to wells pre-incubated with serial dilutions of 2A6 (Fig. 6b). The results suggested that, at the concentrations tested, 1D10 and 2A6 did not alter the extent of binding of conjugated 2A6 and 1D10, respectively, to M-like protein. This implied that the epitopes recognized by 1D10 and 2A6 were dissimilar and that at least two mouse-protective epitopes were present in a preparation of M-like protein. As expected, the amount of HRP-conjugated 1D10 (or 2A6) which bound to M-like protein increased as the amount of unconjugated 1D10 (or 2A6) was decreased. The observed increase in $A_{405}$ was greater for the 2A6 conjugate than for the 1D10 conjugate. This may be explained by the fact that the 2A6 conjugate had a titre of 1/2050 against M-like protein as assessed by EIA whereas the 1D10 conjugate had a titre of 1/256.
Discussion

In this study a partially purified preparation of M-like protein, obtained by extracting \textit{S. equi} cells in hot acid (Woolcock, 1974) was shown to afford a high level of protection (80\%) to immunized mice that were subsequently challenged with a lethal dose of \textit{S. equi}. This is in agreement with the findings of Woolcock (1974) and Timoney & Galan (1985). M-protein is believed to be the major virulence factor of group A streptococci (Lancefield, 1928; Beachey, 1985), rendering the streptococcus resistant to non-immune phagocytosis (Bisno, 1979; Peterson \textit{et al.}, 1979). M-protein isolated from group A streptococci by hot-acid extraction (Lancefield, 1928), sonication (Bessin & Pine, 1968), alkaline extraction (Fox & Wittner, 1969), pepsin digestion (Cunningham & Beachey, 1974), streptococcal phage lysis (Fischetti \textit{et al.}, 1974), guanidine. HCl (Russell & Facklam, 1975) and non-ionic detergents (Fischetti \textit{et al.}, 1976) has been shown to be heterogeneous. The hot-acid-extracted M-like protein of group C \textit{S. equi}, which was partially purified in this study, was found to be a heterogeneous preparation, presumably containing M-protein and its smaller breakdown fragments in addition to other protein. Some of these proteins/polypeptides were shown to be mouse-protective. It is interesting that a higher level of protection (60\%) was consistently obtained with 50 \(\mu\)g HAE than with 100 \(\mu\)g of the same preparation (45\%).

In addition mAbs raised against a partially purified preparation of M-like protein isolated from \textit{S. equi} can afford a high level of passive protection to mice challenged with a lethal dose of the bacteria. The passive transfer assay used in this study was adapted from methods described by Woolcock (1974), Erickson & Norcross (1975), Mathews & Roehrig (1982) and Srivastava & Barnum (1983). These authors reported that a single dose of polyclonal antibody was satisfactory for protection. In our experience, effective protection was not achieved with a single dose of either murine monoclonal or equine polyclonal antibody. At least 1.5 mg of antibodies per mouse, administered over 3 d, was deemed necessary for protection from the large challenge dose of bacteria used in the experiments reported here. Both murine monoclonal and equine polyclonal antibodies were equally effective in our assay. Despite the fact that each hybridoma had been cloned at least twice to ensure monoclonality, all the mAbs recognized multiple polypeptide bands on Western immunoblots. This indicates the presence of breakdown products of a single protein in the antigen preparation or cross-reacting epitopes on different proteins/polypeptides, or both. It is interesting to note that the two protective mAbs, 1D10 and 2A6, displayed the same recognition banding pattern, i.e. pattern 1, on a Western immunoblot. In addition, several of the proteins/polypeptides recognized by these mAbs were also recognized by mouse-protective horse antibodies. Determination of epitope specificity of these mAbs revealed the existence of two mouse-protective epitopes in the M-like protein preparation.

In addition to displaying the same immunoblot banding pattern, mAbs 1D10 and 2A6 were both of the IgG isotype. mAbs 3D4 and 10D3, on the other hand, were both of the IgM isotype and non-protective, despite the fact that they appeared identical to 1D10 and 2A6 in their reaction against M-like protein on immunoblots (data not shown). These findings suggest that both the specificity and the isotype of the immunoglobulin in question may be important in conferring protection. The difficulties encountered in purifying the more unstable IgM from ascites fluids containing mAbs 3D4 and 10D3 and the difficulty in producing murine IgM-containing ascites with titres of the same orders of magnitude as IgG-containing ascites may also explain the observed lack of protection with 3D4 and 10D3. Even though it has been shown that the two protective mAbs 1D10 and 2A6 recognized different epitopes in a preparation of M-like protein, their ability to protect mice was not necessarily synergistic. This aspect requires further investigation.

Despite the fact that not all aspects of the passive protection assay were extensively investigated, the assay has been used successfully in this study to identify protective mAbs directed against the M-like protein of \textit{S. equi}. Future work will involve the use of this assay for the identification of other immunogenic epitopes which are required for effective protection against \textit{S. equi} infection.

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References


