Production of mouse monoclonal antibodies to Pasteurella multocida type A and the immunological properties of a protective anti-lipopolysaccharide antibody

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Summary. Eight monoclonal antibodies (MAbs) were produced from mice immunised with whole cells of heat-killed Pasteurella multocida type A which had been cultured under iron-restricted conditions. The MAbs were selected by an enzyme-linked immunosorbent assay (ELISA) in which the antigen consisted of whole bacteria of the immunising strain. Their reactivity was investigated further by immunoblotting, indirect haemagglutination, a complement-mediated bactericidal assay and passive protection of mice. One of the eight MAbs was shown by immunoblotting to react with lipopolysaccharide (LPS), was bactericidal, and completely protected mice against homologous challenge with 10 LD50 of live bacteria. This MAb was selected for further study. Its reaction with LPS of 17 type-A strains and of single strains of types B, D and E was investigated by immunoblotting. Strains that reacted with the anti-LPS MAb in immunoblots were susceptible to its bactericidal activity and gave high ELISA absorbances. Those that did not react were not susceptible to its bactericidal activity and gave low ELISA readings. The relation between bactericidal activity and ELISA absorbance was highly significant (p < 0.001). Five of the strongly reacting heterologous strains and one non-reacting strain were selected as challenge organisms in a passive protection experiment: only the mice receiving the reacting strains were protected.

Introduction

Pasteurella multocida is the cause of various diseases in mammalian and avian species.1 Capsular type-A strains cause fowl cholera, pneumonia in cattle, sheep and pigs, and “snuffles” in rabbits;1,2 strains of types B and E cause haemorrhagic septicaemia in cattle and buffaloes;3 and type-D strains cause pneumonia in cattle and atrophic rhinitis in pigs.1,4 Capsular polysaccharides of P. multocida types B and E are protective against haemorrhagic septicaemia.5 However, non-capsulate and capsulate organisms are equally effective in immunising birds against fowl cholera.6 Lipopolysaccharide (LPS) of P. multocida, which is considered to be responsible for the somatic serotype specificity,7 was poorly immunogenic in mice in its purified form, but protective in chickens.8 A protein-LPS complex protected mice, rabbits and chickens.9,10 LPS has been recognised as the major immunogen in ribosomal vaccines.11,12 The role of outer membrane proteins (OMP) in protection against P. multocida infections is unclear. In mice, protection by whole-cell type-A vaccines was unrelated to their content of OMP;13 in rabbits, however, a 37.5-KDa OMP has been identified as a protective antigen, but only against the strains that produce it.14,15

Materials and methods

Bacterial strains

P. multocida strain W674, a calf pneumonia isolate, was used to immunise mice for MAb production. The National Collection of Type Cultures supplied strains of capsular serotypes A (NCTC 10322), B (NCTC 10323), D (NCTC 10325) and E (NCTC 10326). All other strains

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were of *P. multocida* type A, isolated from calves at Veterinary Investigation Centres and submitted to the Moredun Institute for identification as described previously.16

**Immunisation procedure and production of monoclonal antibodies**

Five 6-week-old BALB/c mice were immunised with 10⁸ heat-killed (60°C, 90 min) cells of strain W674, grown in nutrient broth (NB; Oxoid) containing 2,2'-dipyridyl; the iron-restricted medium probably increased the resemblance of the bacteria to those grown *in vivo*. A suspension of washed bacteria (8.0 × 10⁹/ml in PBS) was emulsified with an equal volume of Bayol/Arlacel adjuvant. One volume of the emulsion was mixed with 3 volumes of a 1 in 10 dilution of alhydrogel. Each mouse received 0-1 ml of this mixture intraperitoneally (i.p.) on days 0 and 14. Blood was taken from the tail vein of each mouse on day 22 and the antibody response was measured by ELISA (see below) with an antigen consisting of whole cells grown in the presence of 2,2' dipryridyl. The mouse with the highest antibody titre was selected for fusion. Three days before fusion (day 30+) the mouse was inoculated intravenously with heat-killed *P. multocida* (10⁶ cells) in 0-1 ml of saline. Fusion with NS-0 cells was made by the method of Kohler and Milstein17 except that polyethylene glycol was used as the fusion agent. Hybridomas producing specific antibody were detected by a whole-cell ELISA (see below) and were cloned and sub-cloned by limiting dilution.

Ascitic fluid was produced in adult BALB/c mice which had been primed i.p. with 0-5 ml of pristane (Sigma) 3 days before an injection by the same route of 10⁷ hybridoma cells. Fluid was collected 7–10 days later.

**Characterisation of MAbs**

**Isotype determination.** The class and subclass of the MAbs in culture supernates were determined by means of a mouse monoclonal-antibody isotyping kit (RPN29: Amersham International plc) according to the manufacturer’s instructions.

**Enzyme-linked immunosorbent assay (ELISA).** An ELISA was used (a) to detect the antibody response of mice to vaccination, (b) to screen hybridomas for the production of specific antibody, and (c) to test the reactivity of MAbs against heterologous strains of bacteria. For purposes (a) and (b), *P. multocida* strain W674 was grown in NB containing 2,2'-dipyridyl for 6 h at 37°C, washed once in phosphate-buffered saline (PBS) and resuspended in carbonate-bicarbonate buffer, pH 9-6, to a concentration (10⁷ cfu/ml) previously shown by chess-board titration to be optimal for coating. For (c), an overnight culture of each strain was grown in NB, washed once in PBS and resuspended in formol saline 0-5% to c. 10⁷ cfu/ml (A420 = 0-1). The wells of the microtitration plates were coated with 100 μl of bacterial suspension by incubating overnight at 4°C. The primary antibody (100 μl) was either undiluted hybridoma culture supernate or doubling dilutions of mouse serum, after an initial 1 in 50 dilution, in PBS containing Tween 20 0-05% (PBST). After incubation for 1 h at 37°C and washing four times in PBST, 100-μl volumes of horse-radish peroxidase- sheep anti-mouse Ig conjugate (Scottish Antibody Production Unit, Carluke) diluted 1 in 200 in PBST were added and incubated for 1 h at 37°C. After washing as previously, 100-μl volumes of substrate solution (ortho-phenylenediamine, Sigma; 0-4 mg/ml in citrate phosphate buffer, pH 5, with 30% hydrogen peroxide 4 μl/10 ml) were added and after 1 h at 37°C the colour was read at 492 nm in a Titertek Multiscan.

**Immunoblotting.** Whole-cell lysates of *P. multocida* strain W674 and Sarkosyl-extracted outer membranes prepared as described by Abdullahi et al.13 were separated on polyacrylamide 10% gels with the buffer system of Laemml.18 LPS was prepared from several strains by the Proteinase K method of Hitchcock and Brown19 and separated on polyacrylamide 14% gels with the same buffer system as that for the 10% gels except that sodium dodecyl sulphate was omitted from the separating and stacking gel buffers. Separated antigens were transferred to nitrocellulose membranes (Schleicher and Schuell; pore size 0-2 μm) and probed with undiluted hybridoma culture supernates for 4 h at room temperature. Immune complexes were detected with anti-mouse Ig-horseradish peroxidase conjugate and the BioRad HRP colour reagent.20

**Indirect haemagglutination (IHA).** This was performed essentially by the method of Carter.21 Doubling dilutions of MAbs (ascitic fluids), from 1 in 2 to 1 in 128, were tested against sheep red cells sensitised with capsular antigen prepared from the homologous strain W674 and several heterologous strains.

**Bactericidal assay.** MAbs were tested for their bactericidal capacity by the method of Wijewardana and Sutherland.22 Briefly, triplicate samples (20 μl) of MAb (ascitic fluid) were incubated with 100 μl of a suspension of strain W674 (2000 cfu/ml) in modified barbitone buffer (Sigma) for 10 min at room temperature in microtitration plates (tissue culture grade; NUNC, Denmark). After the addition of 80 μl of undiluted fresh complement (serum from new-born calves or gnotobiotic lambs) to each well the plates were incubated for a further 30 min at 37°C. From each well, triplicate 10-μl samples were removed and plated on 7% sheep blood agar. The number of colonies (T₃₀) was counted after overnight incubation at 37°C. The initial bacterial inoculum was confirmed retrospectively by plate counts23 and the assay count (T₀) was calculated as 0.5 × the mean cfu/ml. The percentage killing (K%) was then calculated by the formula:

\[
K\% = 100 \times \left( \frac{\text{mean cfu/ml at } T_0}{\text{mean cfu/ml at } T_{30}} \right)
\]

Hyperimmune rabbit serum against strain W674 and fetal bovine serum were included as standard positive and negative sera respectively. The bactericidal capacity of the MAbs against heterologous strains was tested by the same method.

**Passive protection of mice.** The protective capacity of MAbs was investigated in BALB/c mice. Each MAb
(0.2 ml) in the form of ascitic fluid was inoculated into five mice. Ascitic fluid (0.2-ml volumes) of an irrelevant MAb (MAb E7, raised against louping-ill virus and kindly supplied by Mr M. Hussein of the Moredun Research Institute) was inoculated into five control mice. One hour later, all mice were challenged with approximately 10 LD50 of homologous or heterologous bacteria. Mice were observed for 48 h after challenge, deaths were recorded and the percentage protected was calculated.

Results

Production and characterisation of monoclonal antibodies

Of 97 hybridoma supernates tested in ELISA, 20 gave optical densities (OD) at 492 nm of greater than 0.5. However, only eight continued to produce specific antibody after cloning and these were characterised further.

Isotype determination. Four MAbs (1/2.16.8, 1/15.9.1, 1/18.13.5 and 1/20.10.15) were of the IgM isotype, two (1/4.1.6 and 1/19.12.13) were IgG1, one (1/13.6/12) was IgG2, and one (1/8.16.11) was IgG3.

Immunoblotting. By this technique, only two MAbs were shown to react with separated, transferred antigens of the immunising strain W674. When the outer membranes were probed, only MAb 1/4.1.6 reacted (with a 77-Kda band; fig. 1a). However, when LPS was used, another MAb, 1/8.16.11, reacted strongly with the material at the gel front, which corresponded to LPS. This is shown in fig. 1b. Included in this figure are tracks probed with pre-cloned and pre-subcloned supernates.

Indirect haemagglutination. Only one MAb, 1/2.16.8, demonstrated haemagglutinating activity (titre of 64) and was considered to be reactive with a component of the capsule. Subsequently, when this MAb was tested with heterologous strains, it was found to be specific for the immunising strain.

Bactericidal activity. Complement-mediated bactericidal activity was demonstrated only by the anti-LPS MAb (1/8.16.11). There was 80% killing of the homologous strain.

Passive protection of mice. The group of mice that received the anti-LPS MAb (1/8.16.11) resisted challenge with 10 LD50 of strain W674 and remained well during the 48-h observation period.

Fig. 1. Immunoblots of surface antigens of P. multocida strain W674 probed with undiluted hybridoma culture supernates. (a) Sarkosyl-prepared outer-membrane proteins: the only tracks showing positive reactions compared with the negative control (track 16, uninoculated culture medium) are 14 and 17, which correspond to MAb 1/4.1.6 and the positive control (serum from the mouse used for fusion) respectively. (b) LPS: the tracks showing strong, positive reactions were probed with MAb 1/8.16.11 (track 5), and supernates from the uncloned primary hybridoma culture well (track 3) and the first cloning of this culture (track 4); the serum from the mouse chosen for fusion is in track 14, and shows a weak positive reaction.
indicating 100% protection; all the control mice died. In the groups of mice inoculated with MAbs 1/19.12.13 and 1/20.10.15, two and five animals respectively were alive at the end of the observation period but they were severely ill. After killing the surviving mice by cervical dislocation, heart blood was plated on sheep blood agar. *P. multocida* was recovered from the sick mice but not from the mice protected with the anti-LPS MAb.

**Reaction of anti-LPS MAb with heterologous strains**

From the results described above, it was evident that the anti-LPS MAb (1/8.16.11) protected mice against homologous challenge and promoted complement-mediated killing. Therefore, this MAb was tested against heterologous strains in the whole-cell ELISA, in immunoblotting with LPS preparations, in bactericidal assays and by passive protection tests in mice.

The whole-cell ELISA results and the bactericidal capacities are shown in table I. The absorbances of heterologous strains are expressed as percentages of the OD₄₉₂ of the homologous strain. Correlation between percentage killing and percentage absorbance, determined by the Spearman-Jackson rank correlation test, was highly significant (p < 0.001).

ImmunobLOTS of LPS from 20 heterologous strains showed that 10 strains reacted strongly with the anti-LPS MAb and 10 did not (fig. 2). These reactions correlated with the ELISA and the bactericidal results. Fig. 3 shows that silver-stained polyacrylamide gels of the LPS preparations were all similar and resembled rough LPS of the enterobacteria.

The passive protection afforded by the anti-LPS MAb against six heterologous strains—five of which reacted in ELISA and in immunoblotting and one (strain A848) which did not—is shown in table II. The only strain against which no protection was obtained was A848; this strain was not killed by complement-mediated bactericidal activity. Protection was 60–100% against the other strains.

**Discussion**

Of the eight MAbs that reacted in ELISA to whole bacteria, three were characterised further and shown by immunoblotting to be specific for LPS, capsule and an OMP respectively. The other five MAbs did not react in immunoblotting or IHA and were probably directed against protein epitopes which were irreversibly denatured during separation on polyacrylamide gel electrophoresis or on transfer to nitrocellulose. As these MAbs were shown subsequently not to be protective in mice they were not investigated further.

The MAb that reacted only with the homologous capsular antigen in IHA did not mediate complement-dependent killing of bacteria *in vitro*. It did not protect mice against challenge with live organisms and was probably directed against a strain-specific surface antigen rather than to the type-specific hyaluronic acid capsule common to all type-A strains of *P. multocida*.

Similarly, the anti-77 Kda OMP MAb did not protect mice and was not bactericidal. Lu et al. found that hyperimmune rabbit antisera containing a high titre of IgG antibody to a 37.5-Kda OMP was protective in rabbits. However, they used a polyclonal serum containing antibody to several epitopes in a single molecule rather than a MAb defining a single epitope.

The most significant finding of the present study was that anti-LPS IgG₃ MAb protected mice against homologous and heterologous challenge by a mechanism associated with complement-depend-
Fig. 2. Examples of immunoblots of LPS from homologous (track 9) and heterologous strains of _P. multocida_ type A probed with anti-LPS MAb 1/8.16.11 diluted 1 in 2. Track 1, strain A757; 2, A36; 3, X110; 4, X1056; 5, A848; 6, W666; 7, Q110; 8, X1016; 9, W674; 10, X120.

ent antibody-mediated bacterial killing. An investigation of the bactericidal activity of serum from vaccinated mice suggested that a cell-surface antigen was associated with this mechanism of immunity, because the bactericidal capacity of mouse polyclonal serum was closely correlated (p < 0.001) with whole-cell ELISA titres. It is clear that anti-LPS antibodies play a major role in immunity in mice by participating in complement-mediated bacteriolysis. This agrees with earlier views.

In addition, the LPS of _P. multocida_ type A has been identified as an important immunogen when

Table II. The activity of the anti-LPS MAb in passive protection of mice challenged with heterologous strains

<table>
<thead>
<tr>
<th>Challenge strains</th>
<th>Number of survivors in groups of five mice</th>
</tr>
</thead>
<tbody>
<tr>
<td>No.</td>
<td>LD50</td>
</tr>
<tr>
<td>A36</td>
<td>4.8 × 10⁶</td>
</tr>
<tr>
<td>X110</td>
<td>3.6 × 10⁵</td>
</tr>
<tr>
<td>X1053</td>
<td>8.8 × 10⁵</td>
</tr>
<tr>
<td>A341</td>
<td>6.3 × 10⁵</td>
</tr>
<tr>
<td>A1174</td>
<td>6.1 × 10⁵</td>
</tr>
<tr>
<td>A848</td>
<td>1.9 × 10³</td>
</tr>
</tbody>
</table>

Mice were challenged with c. 10 LD50.
conjugated with protein in chickens, mice and rabbits. To induce immunity, ribosomal vaccines should contain LPS. An LPS-ribosome complex gave an enhanced humoral but not cell-mediated immunity in chickens.

The epitope recognised by the anti-LPS MAAb was found to be present on the surface of about 50% of the capsule type-A strains investigated. It will be of interest to define the chemical structure of this epitope and to determine whether it is specific to one or more O serotypes in capsule type-A strains.

REFERENCES