A monoclonal antibody reacting with a determinant on leptospiral lipopolysaccharide protects guinea pigs against leptospirosis

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Summary. An IgA monoclonal antibody (MUM/F1-1/copenhageni) was produced from a mouse immunised with Leptospira interrogans serovar copenhageni. The antibody showed partial serogroup specificity by agglutination and by reaction in enzyme immunoassay, and opsonised homologous leptospires for phagocytosis by cultured mouse macrophages. Immunodiffusion and Western-blotting experiments indicated that MUM/F1-1/copenhageni reacted with a carbohydrate determinant in the leptospiral lipopolysaccharide. Daily administration of purified MUM/F1-1/copenhageni IgA before and after challenge with $2 \times 10^8$ virulent homologous leptospires passively protected newborn guinea pigs against lethal leptospirosis.

Introduction

It is well established that development of a humoral immune response during leptospirosis is important in resistance to infection (Adler and Faine, 1976, 1977, 1978a and b; Adler et al., 1980a). Immunity appears to depend on the production of agglutinating, opsonising antibody; cell-mediated immune mechanisms have no detectable role. Although the immune response to infection with leptospires is well documented, the nature of the antigens important in protection remains unclear. Many preparations of leptospiral antigens have been reported (Faine, 1974) but only a few have been purified for definitive study. One of these antigens, the leptospiral lipopolysaccharide (LPS) structurally resembles endotoxins of other gram-negative bacteria, although it has different chemical and biological properties (Vinh et al., 1986). As leptospiral LPS elicits the production of protective antibody during infection (Adler and Faine, 1978a and b), it is possible the LPS is an important protective antigen. Antibodies against LPS from other gram-negative organisms can be used to passively protect against homologous infection and LPS can also be used as an immunogen against experimental infection (Pennington and Kuchmy, 1980; Colwell et al., 1984; Kirkland and Ziegler, 1984; Sawada et al., 1984).

To determine the nature of the leptospiral antigens that stimulate the production of protective antibody, monoclonal antibodies (MCAs) have been produced. This strategy has been used successfully for other organisms, notably the malaria parasite Plasmodium (Potocnjak et al., 1980; Holder and Freeman, 1981; Boyle et al., 1982). Anti-leptospiral MCAs have been described (Ono et al., 1982, 1984; Adler and Faine, 1983a and b; Kobayashi et al., 1984), but have been mainly of taxonomic significance. If these MCAs can be used to passively protect against infection in a suitable animal model, they should be useful to identify and characterise potential protective antigens whose determinants could be the basis for immunogens suitable for human and veterinary use.

This paper reports the production and characterisation of a protective MCA directed against a determinant on the leptospiral LPS.

Materials and methods

Leptospires

Leptospira interrogans serovar copenhageni strain designated H45 in this laboratory was isolated from a rat (Faine and van der Hoeden, 1964) and serovar copenhageni strain L136 was provided by Professor R. Yanagawa, Hokkaido University, Sapporo, Japan. L. illini serovar illini was obtained from A.D. Alexander, Walter Reed Medical Center, Washington D.C., USA. The other leptospiral serovars used in this study were provided by N. Stallman, W.H.O. Leptospira Reference Laboratory,
Brisbane, Australia. Cultivation and enumeration of leptospires were as described by Adler and Faine (1976).

Antigens, antisera and serological methods

Leptospiral LPS was prepared by the method of Westphal and Jann (1965), with the modification that the LPS was finally purified by centrifugation at 105,000 g for 3 h. LPS polysaccharide fraction was prepared by boiling LPS 1 mg/ml in acetic acid 1.5% v/v for 2 h (Naess and Hofstad, 1984). The precipitated lipid was removed by centrifugation at 400 g and the supernate was used as the polysaccharide fraction of LPS. Protease treatment of LPS was by overnight incubation of LPS 50 μg/ml with Proteinase K (Sigma) 40 μg/ml at 37°C. Leptospiral sonicate antigens were prepared as previously described (Adler et al., 1980b) and a phenol-extracted LPS preparation, designated TM antigen (Shinagawa and Yana-gawa, 1972), was the kind gift of Professor R. Yanagawa, Hokkaido University, Sapporo, Japan. The preparation of mouse antisera was described by Tu et al. (1982), and the methods for performing the microscopic agglutination test (MAT), enzyme immunoassay (EIA) and gel immunodiffusion were described by Adler and Faine (1983a). Subclasses of mouse immunoglobulin were determined by reaction with specific anti-mouse immunoglobulins (Nordic Immunologicals, Tilburg, The Netherlands) in gel immunodiffusion.

Production of monoclonal antibodies

Adult female Balb/c mice were immunised intraperitoneally (ip) with 0.5 ml of a suspension of serovar copenhageni strain H45 2.5 × 10⁶ leptospires/ml grown in low protein EMJH medium (Bey and Johnson, 1978) and washed once in physiological saline. Four weeks later the mice received a similar, booster injection. After 3 days, spleen cells were harvested and 10⁶ spleen cells were fused with 10⁶ NS-1 (P3-X63-Ag8.653) myeloma cells by the method of Kohler and Milstein (1975). The fusion mixture was dispensed into two 24-well tissue culture trays (Costar) which contained 5 × 10⁴ mouse macrophages/well. Resultant hybridomas were grown in Dulbecco Modified Eagle’s Medium (DME, Microbiological Associates, Bethesda, MD, USA) supplemented with fetal calf serum 10% v/v (FCS, Commonwealth Serum Laboratories, Australia). MAT and EIA were used to screen undiluted hybridoma supernatants for specific antibody. Antibody-positive cultures were cloned by limiting dilution in 96-well Costar tissue culture trays with a cloning medium of L929-conditioned medium 25% v/v in DME supplemented with FCS 15% v/v (DME-15% FCS). L929-conditioned medium consisted of DME-10% FCS in which mouse fibroblasts (cell line L929, Common-wealth Serum Laboratories, Australia) had been cultured for 2 days at 37°C. All hybrids were cloned at least twice and were grown to extinction in large volumes of DME-10% FCS. The cells were removed by centrifugation and the culture supernate was treated by precipitation with 50% saturated ammonium sulphate (Göding, 1980) to purify the immunoglobulin fraction. Monoclonal antibody (MCA) levels were determined by absorbance at 280 nm.

Polyacrylamide gel electrophoresis and Western blotting

Leptospiral antigens were separated by electrophoresis according to the method of Laemmli (1970) on a 1 mm thick discontinuous SDS-polyacrylamide 15% w/v gel in a Bio-Rad Protein II gel box for 4 h. Resolved antigens were electrophotorethetically transferred on to nitrocellulose membranes (Bio-Rad, USA) by the method of Towbin et al. (1979) with a 1 in 2 dilution of the described transfer buffer, in a Bio-Rad Transblot cell for 3 h at 60V. Immunological detection of specific antigens was performed with a peroxidase-conjugated second antibody with chloro-1-naphthol as the chromogen by the method of Hawkes et al. (1982). Western-blotted membranes were stained for protein by Penguin India Ink 0-1% w/v in PBS containing tween 20 0.3% v/v (Merck, West Germany). Gels were stained for LPS by the modified silver stain of Tsa and Frasch (1982).

Opsonisation assays

Mouse macrophages for in-vitro opsonisation of leptospires for phagocytosis were prepared as described by Tu et al. (1982). L. interrogans serovar copenhageni strain H45 2 × 10⁶ cells were added to Leighton tubes containing macrophage monolayers. Dilutions of hybridoma culture supernate, antiserum, normal mouse serum (NMS) or DME-10% FCS were also added and the tubes were incubated and fixed as described (Tu et al., 1982), in preparation for indirect immunofluorescence staining. The second antibody used was a 1 in 20 dilution of fluoroisothiocyanate-conjugated anti-mouse immunoglobulin (Wellcome Diagnostics, Australia).

The methods for generation and measurement of the chemiluminescence (CL) response by phagocytic cells were as described by McGrath et al. (1984), but with mouse macrophages. Macrophages were harvested into warm DME-10% FCS by peritoneal lavage and samples containing 1 × 10⁶ macrophages in 1-ml volumes were pipetted into glass vials. The vials were incubated for 1 h at 37°C in a humidified CO₂ incubator before measurement of CL.

Passive protection experiments

Outbred guinea pigs of either sex, weighing less than 100 g and obtained from the Monash University animal house, were given ip injections of 2 × 10⁸ leptospires of a virulent serovar copenhageni strain L136 in EMJH medium. The virulence of the leptospires was maintained by continuous passage in guinea pigs. Infected guinea pigs always died within 3–4 days of challenge. To confirm that death was due to acute leptospirosis, all dead animals were examined post mortem and samples of blood, liver and kidneys removed aseptically, emulsified in 5-ml
volumes of EMJH medium and incubated at 30°C for up to 21 days or until viable leptospires could be detected by dark-ground microscopy. At autopsy, infected animals consistently displayed distinctive pathological changes indicative of acute leptospirosis which included extensive haemorrhaging of the peritoneal and abdominal walls, peri-renal haemorrhages and extensive jaundice. Leptospires could also be cultured from various sites. Guinea pigs given similar volumes of sterile EMJH medium were killed after 4 days and were normal on autopsy. In this model, death of the animal with a characteristic pathology and the presence of live leptospires were considered to be the criteria of lethal leptospiral infection. A group of three guinea pigs each received 175 μg of purified F1-lc IgA MCA ip 1 h before challenge with $2 \times 10^8$ virulent $L. interrogans$ serovar copenhageni strain L136. These animals subsequently received four similar doses of F1-lc IgA at daily intervals. To determine passively-acquired serum antibody levels, guinea pigs were bled by cardio-puncture after pentobarbitone anaesthesia and antibody titres in the separated serum were determined by MAT.

**Results**

**Serological specificity**

MUM/F1-1/copenhageni (designated F1-1c) was an IgA class antibody which reacted in both the microscopic agglutination test (MAT) and enzyme immunoassay (EIA). The antibody agglutinated 11 of the 17 serovars of the Icterohaemorrhagiae serogroup, but did not react significantly with representative serovars of the other serogroups of $L. interrogans$ or other species of *Leptospira*. Representative members of all other $L. interrogans$ serogroups, $L. biflexa$ serovar patoc and $L. illini$ were tested. For all non-reactive serovars the MAT titres were zero and the EIA optical density values were at background level (table). Those serovars agglutinated by F1-1c also reacted by EIA with the exceptions of serovar naam and the cross-reactive canicola serovar (table).

**Nature of the antigen**

F1-1c IgA reacted in immunodiffusion with purified homologous LPS, TM and sonicate antigens to produce lines of identity (fig. 1). An additional precipitin band visible with LPS was not observed when the antigen was boiled before testing. This pattern of reactivity is often seen for leptospiral LPS suggesting that the LPS exists in different aggregation states. The MCA also reacted with these antigens in EIA (results not shown).

Homologous LPS 1 μg and LPS polysaccharide fraction 5 μg were resolved by electrophoresis on polyacrylamide gels 15% w/v and silver-stained (fig.

<table>
<thead>
<tr>
<th>Serogroup of $L. interrogans$</th>
<th>Serovar</th>
<th>MAT titre*</th>
<th>EIA†</th>
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*Expressed as highest dilution showing agglutination.
†Expressed as percentage of OD at 488 nm obtained by heterologous EIA compared with homologous copenhageni EIA (0-80).
- = EIA OD ≤ background level.

Fig. 1. Immunodiffusion reactions (20-μl volumes of serovar copenhageni antigen preparations with F1-1c antibody: (1) Sonicate (equivalent to $1 \times 10^8$ leptospires/ml), (2) TM (400 μg/ml), (3) LPS (400 μg/ml), (4) F1-1c MCA (culture supernate, 20x concentrated); 20-μl volume were added to each well.
Fig. 2. SDS-polyacrylamide gel electrophoresis of serovar *copenhageni* antigen preparations: (a) profile obtained by silver staining; (b) profile obtained by Western-blot analysis with F1-Lc MCA. Lanes: (A) Mol-wt markers, with mol. wts (10³); (B) 1 µg LPS; (C) 5 µg LPS polysaccharide fraction.

2a) or Western-blotted and immuno-stained with purified F1-Lc IgA 20 µg/ml (fig. 2b). Control immunoblots were stained with 1 in 200 dilutions of mouse anti-*copenhageni* serum or NMS as positive and negative controls respectively. The profile observed with the mouse anti-*copenhageni* serum was similar to that with the MCA (not illustrated). F1-Lc reacted with the resolved LPS producing a smear of staining unlike the characteristic pattern of discrete bands often seen with protein antigens, but similar to that described for LPS from other bacteria (Linko-Kettunen *et al.*, 1984). However when LPS polysaccharide fraction was used as the antigen, the smear was resolved into a discrete, characteristically arched band of mol. wt c. 44 000 (fig. 2b). A similar band was detected by silver staining (fig. 2a). Reaction of purified LPS polysaccharide fraction with F1-Lc suggested that the specific epitope was of a carbohydrate nature. This result was confirmed by the reaction of F1-Lc with protease-treated LPS in EIA. Treatment of LPS with Proteinase K did not alter its EIA reactivity, which showed that the determinant was resistant to proteolysis.

**Opsonisation experiments**

Mouse macrophages were used to test the opsonic capacity of F1-Lc culture supernate, both by CL and by indirect immunofluorescent (IF) staining. Vials containing macrophages were incubated with 1 x 10⁸ cells of *L. interrogans* serovar *copenhageni* strain H45 and CL was measured continuously. No significant change in CL response was observed when either NMS or DME-10% FCS were introduced. However, addition of mouse anti-*copenhageni* antiserum or F1-Lc culture supernate resulted in CL responses of 2.00 mV and 2.14 mV respect-
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ively (fig. 3). The responses peaked after 5–7 min and slowly declined to base levels in 30–40 min. Addition of a 1 in 50 dilution of fresh guinea-pig serum as a source of complement did not significantly alter the responses observed. Similarly mouse macrophages co-incubated with leptospires and 1 in 500 NMS or 1 in 20 DME-10% FCS and stained by indirect IF showed no fluorescence, in contrast to macrophages treated with 1 in 500 mouse anti-copenhageni antiserum (MAT titre, 512) as a positive control or 1 in 20 F1-lc culture supernate (MAT titre, 4096), which were strongly fluorescent, with fluorescent-staining leptospires seen attached to and within the macrophages.

Passive protection

The capacity of F1-lc antibody to passively protect against infection was investigated. Initial experiments indicated that passively-acquired serum levels of F1-lc decreased rapidly from titres of 32 to 2-4 in less than 24 h. Animals receiving only a single dose of 175 µg of MCA were not protected against challenge, therefore repeated doses of F1-lc antibody were administered daily to ensure that passive serum MAT titres of 32–64 were maintained. A group of three control animals which received physiological saline in place of antibody, followed by challenge, all died within 3–4 days with typical autopsy findings indicative of acute leptospirosis (see Materials and methods). In contrast the three animals given F1-lc antibody survived, and when killed 14 days after challenge were normal at autopsy. Viable leptospires were not detected in blood, liver or kidney cultures taken from these animals at autopsy.

A passive serum agglutination titre of 16–32 in these animals was sufficient to protect. However, protection was not afforded by passive serum titres of 4 and, therefore, the minimum protective agglutinating titre of this antibody must lie somewhere in the range 4–32.

Discussion

This paper reports the production of an IgA monoclonal antibody directed against a partially serogroup-specific determinant found on leptospiral LPS of some but not all Icterohaemorrhagiae serogroup members. The MCA did not react significantly with serovars outside the Icterohaemorrhagiae serogroup. Those serovars that reacted in MAT were also reactive in EIA with the exception of serovar naam and the cross-reactive canicola serovar. Serovar naam did not react significantly in EIA although it was strongly agglutinated by the MCA (table). This unusual reactivity is commonly seen with other anti-leptospiral LPS MCAs and may reflect reactions with minor determinants on the leptospires (Ono et al., 1982; Farrelly et al., 1985; Jost et al., 1985). The cross-reaction between serovar canicola and the MCA was much lower than reactions between MCA and homologous serogroup members. This Icterohaemorrhagiae-Canicola cross-reaction is commonly observed with specific rabbit antisera, possibly indicating conservation of some minor epitope (Kmety, 1967).

Kobayashi et al. (1984) have also reported the production of MCAs directed against serovars of the Icterohaemorrhagiae serogroup and described the identification of serovar-specific determinants. However the authors only tested their MCAs against two serovars (icterohaemorrhagiae and copenhageni) from this serogroup. The results of the present study and of other workers (Ono et al., 1982; Farrelly et al., 1985) show that MCAs can cross-react widely with different serovars of the same serogroup. Therefore it is difficult to determine the sero-specificity of an epitope by testing only a small number of serovars.

F1-lc reacted with purified LPS and the TM antigen of Shinagawa and Yanagawa (1972) to produce a line of identity. Reaction of F1-lc with both LPS and TM suggests that the specific epitope occurs in both these antigens and that both are LPS preparations as previously described for serovar pomona (Adler and Faine, 1983a).
The determinant to which the MCA was directed was present in the polysaccharide fraction of the leptospiral LPS. Silver staining revealed that only one band was present after electrophoretic resolution of the fraction. This band was also detected immunologically by staining the Western-blotted polysaccharide fraction with MCA. The reactive band was characteristically arched, indicative of polysaccharide fraction with MCA. The reactive immunologically by staining the Western-blotted fraction. This band was also detected against lethal infection in susceptible animals. The one band was present after electrophoretic resolution of the fraction. Silver staining revealed that only one protective epitope. Reaction with the epitope divided the group taxonomically in that the epitope was not present in all members of the Icterohae-morrhagiae serogroup. There is insufficient information to predict cross-immunity between serovars unrelated by agglutination (Kemenes, 1964; Plesko and Lataste-Dorolle, 1970), although the nature of these antigens and their roles in immunity are not understood.

The results suggest an important role for leptospiral LPS in immunity to leptospirosis and provide evidence that anti-LPS antibodies are important in protection from experimental infection. If the polysaccharide fraction of the LPS is important as a protective antigen, then it may be suitable for use in a vaccine. Knowledge of the protective antigens involved in immunity to leptospirosis is clearly necessary for the development of effective vaccines for prophylaxis and control of this disease.

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