Production of Murine Monoclonal Antibodies to the Major Axial Filament Polypeptide of Treponema pallidum

By M. J. BAILEY,* A. COCKAYNE AND C. W. PENN
Department of Microbiology, University of Birmingham, Birmingham B15 2TT, UK

(Received 21 November 1986; revised 29 January 1987)

A suitable immunization protocol for the stimulation of a murine antibody response to the axial filament polypeptides of Treponema pallidum was established. A range of monoclonal antibodies (Mabs) specific for different epitopes of the major axial filament polypeptide (37 kDa) was generated which demonstrated diversity in their ability to react with other treponemal species. Immunogold electron microscopy located the 37 kDa antigen on the surface of the axial filament structure. The early appearance of specific antibody to this polypeptide in infected man and rabbit indicates that such Mabs are potentially useful for the diagnosis of early syphilis.

INTRODUCTION

Penn et al. (1985a) recently defined the antigenic components of the axial filament structure of Treponema pallidum (Nichols strain), showing it to be composed of a major polypeptide (37 kDa) and three minor components (34, 33.5 and 31.5 kDa). There has been considerable interest in the antigens of this structure, and diagnostic tests based on the cross-reactive axial filaments of Treponema phagedenis (biotype Reiterii) have been described (Nell & Hardy, 1978; Strandberg-Pedersen et al., 1982). The similarity between the axial filaments of the cultivable Reiter treponeme (T. phagedenis) and T. pallidum is well established (Strandberg-Pedersen et al., 1981; Bharrier & Allis, 1974). Indeed, a small-scale protection study of rabbits immunized with fractionated Reiter treponeme axial filaments has been described (Hindersson et al., 1985) and although the immunized rabbits produced antibody strongly reactive with the minor components of T. pallidum axial filaments, no protection from infection was observed. These findings were predictable as the axial filaments of T. pallidum are located within the periplasmic space and covered by an outer membrane. Therefore, those polypeptides which comprise the endoflagella are not exposed at the surface of the intact organism and would not be expected to elicit neutralizing antibody.

Penn et al. (1985a) demonstrated that antibody to the major axial filament polypeptide (37 kDa) could be detected in the extraction medium used to isolate treponemes from infected testes, suggesting the early recognition of this antigen by the immune system. Further immunological investigations into the development of the humoral response in experimental and human syphilis (Lukehart et al., 1982; Hanff et al., 1982, 1983; Bailey et al., 1987) by Western blotting of sera (Penn et al., 1986) have clearly shown the presence of specific antibody to an antigen of approximately 37 kDa early in infection; comparison of these findings with SDS-PAGE profiles of solubilized T. pallidum generated in our laboratory indicates that one of the main antigens recognized is this major axial filament polypeptide. Such observations suggest the utility of identification of antibody to this polypeptide in syphilitic sera as a basis for the development of an immunodiagnostic assay.

Abbreviations: hRSS, hyperimmune rabbit syphilitic sera; i.p., intraperitoneal(ly); Mab, monoclonal antibody; NCM, nitrocellulose membrane(s); PAT, PBS containing 0-1% BSA and 0-1% Tween 20; PBS, 7-2 mM-Na2HPO4, 2.8 mM-NaH2PO4, 0-15 mMNaCl, pH 7-2.

0001-3842 © 1987 SGM
In order to identify unequivocally the component antigens of the axial filament, monoclonal antibodies (Mabs) to this structure were generated. We have observed that mice respond poorly to the axial filament components after conventional immunization protocols with whole treponemes, and now describe the generation of good responses in mice and the derivation of the requisite Mabs.

**METHODS**

**Bacterial strains.** The Nichols strain of *Treponema pallidum* was propagated in the testes of Dutch rabbits and purified as previously described (Penn, 1981, 1983; Penn & Rhodes, 1982). *Treponema phagedenis* (biotype Reiterii) was cultivated *in vitro* as described by Penn & Rhodes (1982). *Treponema hyodysenteriae* was kindly supplied as a frozen pellet by Dr R. M. Lemcke (IRAD, Compton, Newbury, Berks., UK).

**Antigen preparation.** Axial filaments were extracted from *T. pallidum* by the method of Penn et al. (1985a).

**Immunization.** Balb/c female mice (6–8 weeks old) were immunized with a variety of antigen preparations (Table 1) in order to stimulate a response to the axial filament components. Such mice were used either as a source of spleen cells for the production of hybridoma cell lines, or as a supply of specific mouse anti-*T. pallidum* sera. Hyperimmune mouse sera were generated by the intraperitoneal injection of a 0.1 ml volume containing $10^8$ thrice-PBS-washed azide-killed *T. pallidum* cells included in Freund's complete adjuvant (Sigma) on day 0, subsequently given booster injections on days 14, 28 and 56 with the same quantity of bacteria included in Freund's incomplete adjuvant. Mice were also immunized by the intraperitoneal (i.p.) injection of $10^9$ live, freshly harvested treponemes in PBS on days 0, 14, 28 and 56. Test bleeds were done 10 d after the last injection and sera were analysed by Western blotting for the presence of specific antibody. Selected mice were then given further booster injections or new mice were immunized under the regime described above by injecting 0.1 ml purified axial filaments ($0.1 \text{mg}$) included in an equal volume of adjuvant (complete for new mice, incomplete for boosted mice), of which 0.1 ml was injected i.p. Fusions were done on immunized mice which had been given intravenous booster injections of 50 $\mu$l sonicated purified axial filament in PBS (1 mg ml$^{-1}$) 10–12 d after their last i.p. injection and 4 d before fusion.

Additionally, mice were immunized with a preparation of the 37 kDa polypeptide. After 11% SDS-PAGE electrophoresis (Laemmli, 1970) of purified axial filaments from $10^9$ bacteria, the band corresponding to the 37 kDa polypeptide was located by staining the outside edges of a slab gel with Coomassie blue. The band was cut out, frozen in liquid nitrogen and pulverized with a glass rod. It was resuspended in 500 $\mu$l PBS, mixed with an equal volume of Freund's adjuvant and 100 $\mu$l injected i.p. as above.

Hyperimmune rabbit syphilitic sera (hRSS) were prepared as previously described (Bailey et al., 1985).

**Hybridomas.** Mab-secreting cell lines were produced and isolated essentially as described by Bailey et al. (1987). Washed spleen cells from immunized mice were combined at a ratio of 4:1 with NS-1 myeloma cells (Flow) (Kohler & Milstein, 1975) and fused with polyethylene glycol 1450 (BRL). Cells were maintained in 96-well plates and Mab-producing hybridomas were detected by indirect ELISA or dot-blot ELISA.

ELISA. Antibody-secreting parent hybridomas were initially selected by ELISA. Wells of EIA plates (Nunc I, Gibco) were coated overnight at 4°C with 100 $\mu$l purified, washed and sonicated *T. pallidum* cells diluted (5 $\times$ 10$^{-7}$ ml$^{-1}$) in carbonate/bicarbonate buffer (Voller et al., 1979). Individual supernatants were sampled, and 50 $\mu$l added to 50 $\mu$l double-strength PAT (PAT is 0.1%, w/v, BSA, 0.1%, w/v, Tween 20 in PBS) and allowed to react with the coated solid phase for 1 h at room temperature. After washing five times in PBS, 100 $\mu$l rabbit anti-mouse immunoglobulin conjugated to horseradish peroxidase (Dako) diluted 1 in 1500 in PAT was added to each well and left to react for a further 1.5 h. The substrate, o-phenylenediamine (40 $\mu$g ml$^{-1}$, 0.04% H$_2$O$_2$) in citrate/phosphate buffer pH 5.0 (Voller et al., 1979) was applied to each well of the washed plate and those wells containing antibody were identified colorimetrically. As the intention was to isolate Mabs specific to the axial filament polypeptides, an additional immunooassay was developed which required minute quantities of antigen and allowed for the screening of hundreds of supernatants.

Dot-blot ELISA. Using a nitrocellulose membrane (NCM) as the solid phase, 1 $\mu$l volumes of purified, sonicated axial filament preparations (containing 10 ng protein as estimated by the Lowry method) were spotted evenly onto the membrane at marked positions; duplicate areas were coated with the same amount of sonicated whole *T. pallidum*. After air drying for 60 min the NCM was blocked with 3% BSA, 0.1% Tween 20 in PBS for 1 h, washed twice in PBS and placed on top of a wad of filter paper (Whatman 3MM) soaked in PAT. To each marked area of 1 cm$^2$, containing the antigen preparations, was applied 2–3 $\mu$l ELISA-positive hybridoma supernatant. The filters were left for 1 h at room temperature without shaking and then copiously washed in PBS. Rabbit anti-mouse immunoglobulin conjugated to alkaline phosphatase (Sigma) diluted 1 in 2000 in PAT was allowed to react with the filters for 2 h at room temperature on an orbital shaker, and after further washing, specific anti-axial filament antibody-containing supernatants were identified with the fast red/naphthol ASMX phosphate substrate (Sigma) described by O'Connor & Ashman (1982). Hybridomas were cloned twice by limiting dilution onto splenic feeder
T. pallidum axial-filament Mabs

Table 1. Immunization protocols in female Balb/c for the production of anti-T. pallidum axial filament Mabs

Mice were primed (day 0) with antigen delivered in Freund’s complete adjuvant and boosted with antigen in Freund’s incomplete adjuvant. Antigen preparations: soni. cells, sonicated whole T. pallidum cells; AF, sonicated purified axial filaments; 37 kDa, 60 kDa, 37 and 60 kDa polypeptides eluted from SDS-PAGE gels.

<table>
<thead>
<tr>
<th>Mab code</th>
<th>Type of i.p. injection on day</th>
<th>I.v. boost*</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>0</td>
<td>14</td>
</tr>
<tr>
<td>C</td>
<td>Soni. cells</td>
<td>Soni. cells</td>
</tr>
<tr>
<td>H, I</td>
<td>AF</td>
<td>AF</td>
</tr>
<tr>
<td>J</td>
<td>Live Tp</td>
<td>Live Tp</td>
</tr>
<tr>
<td>K, S</td>
<td>Soni. cells</td>
<td>AF</td>
</tr>
<tr>
<td>L, N</td>
<td>37 kDa</td>
<td>37 kDa</td>
</tr>
<tr>
<td>M</td>
<td>60 kDa</td>
<td>60 kDa</td>
</tr>
</tbody>
</table>

* Intravenous boost with antigen diluted in PBS 4 d before fusion. The time (d) after the first injection is shown in parentheses.

cells (5 × 10^6 per well in 96-well plates) and the specificity of the Mab was determined by Western blotting. Mabs were produced in quantity by cell culture.

**SDS-PAGE and immunoblotting.** Linear gradient acrylamide (8–20%, w/v) or continuous 11% (w/v) gels were run with buffers and conditions essentially as described by Laemmli (1970). Polypeptides were either stained with Coomassie blue (PAGE 83, BDH) or transferred electrophoretically to NCM (0-45 μm, Anderman) according to the method of Towbin et al. (1979). After transfer, the NCMs were blocked as described above for the dot-blot assay and reacted with sera or Mab diluted in PAT. Gels were routinely loaded with bacterial suspensions of 2 × 10^7 organisms in 20 μl sample buffer, the suspensions first solubilized by boiling for 5 min, or with 5 μg axial filament preparation in sample buffer, also boiled as above. The molecular masses of the polypeptides were determined by the method of Weber & Osborn (1981) with molecular mass standards from Bio-Rad. Values presented in the figures refer to the known molecular masses of treponemal polypeptides (Penn et al., 1986). Those polypeptides reactive with antibody were detected by the immunoperoxidase technique. Immunoglobulin class specificity was determined by double immunodiffusion in 1% (w/v) agarose in 0.08 M-barbitone buffer pH 8.2, using anti-mouse sub-class antisera supplied by Dr D. Catty, Department of Immunology, University of Birmingham, UK.

**Immunogold electron microscopy.** After removal of the outer membrane from purified treponemes with Triton X-100, sera (1/50 dilution) or Mabs (1/10 dilution) were allowed to react at 37 °C for 1 h with the released axial filaments in suspension. After washing in PBS, the antibody/treponemal suspension was applied to Formvar-coated copper grids. Non-specific binding sites were blocked with 3% (w/v) BSA in PBS and the samples were then reacted with rabbit anti-mouse antibody gold conjugates (Janssen) as above. After washing, samples were negatively stained with 3% (w/v) phosphotungstic acid pH 7.2 and viewed with a Philips 201 electron microscope.

**T. pallidum immobilization test.** Specific sera and Mabs (the latter concentrated 40-fold by ammonium sulphate precipitation) were assessed for their ability to immobilize T. pallidum in vitro by a modification of the method of Nelson & Mayer (1949) as described by Bailey et al. (1987).

**RESULTS**

**Antibody response**

Mice, when injected with live organisms or immunized with azide-killed bacteria included in Freund’s adjuvant to induce a hyperimmune state, responded poorly to those antigens comprising the axial filament of T. pallidum (Fig. 1) when compared with the response observed with rabbit or human syphilitic sera (Fig. 2). Sonication of the organism prior to injection did not enhance the stimulation of anti-37 kDa polypeptide antibody in Balb/c mice (arrow, Fig. 1, II). Repeated injections of live organisms produced a very weak and restricted range of antibody to T. pallidum. However, purified axial filament (a particulate antigen preparation), either in non-immunized mice or in mice previously immunized with either live or dead whole cells, was capable of producing a good specific response. Further, the 37 kDa polypeptide isolated by SDS-
Fig. 1. Immunoblot analysis of purified *T. pallidum* axial filaments (AF) and whole *T. pallidum* cells (WC) run on an 11% SDS-PAGE gel and reacted with 1 in 100 dilutions of sera prepared as indicated below. Mouse sera were taken 10 d after the final injection. I, hRSS. II, Sera from mice immunized with $10^8$ sonicated dead bacteria included in Freund’s adjuvant (four i.p. injections of 0·1 ml at 14 d intervals). III, Sera from mice immunized as in II and boosted on days 56 and 68, with purified *T. pallidum* axial filaments in adjuvant. IV, Sera from mice immunized with the 37 kDa antigen, extracted from SDS-PAGE gels, included in adjuvant (four i.p. injections of 0·1 ml at 14 d intervals). Va, Sera from mice injected with $10^8$ live treponemes diluted in extraction buffer (four i.p. injections of 0·1 ml at 14 d intervals). Vb, Sera from mice injected as in Va and boosted on days 56 and 68, with purified *T. pallidum* axial filaments in adjuvant. Molecular masses (kDa) of known *T. pallidum* polypeptides are shown.

PAGE was also strongly antigenic in mice. Fig. 1 (III) shows the antibody response of immunized mice given booster injections of purified axial filament preparations. This demonstrates that this structure is composed of at least four clearly distinguishable antigens, of 37, 34, 33·5 and 31·5 kDa. Additional antigens were recognized when anti-axial filament serum was reacted by immunoblotting against whole *T. pallidum*. These additional antigens are highly antigenic membrane-associated components (47, 44 and 32 kDa) (unpublished observations) present as contaminants in purified axial filament preparations in sufficient quantity to elicit an antibody response detectable by immunoblotting with whole *T. pallidum* as the antigen (Fig. 3). Cross-reacting antigens present in both *T. phagedenis* (38·5, 35 and 33 kDa) and *T. hyodysenteriae* (33 kDa) were identified by immunoblotting whole organisms of these species with the antisera to *T. pallidum* axial filaments (Fig. 3), and additional cross-reacting polypeptides were demonstrated with hRSS (Fig. 4).

Identification of Mabs against axial filament polypeptides

By dot-blot assay on ELISA-positive hybridoma supernatants a number of cell lines secreting antibody specific to the axial filament structure were identified. This technique allows for the minimal use of precious antigen from a sub-fraction of *T. pallidum* to isolate desired antibody. By immunoblotting (Fig. 4), a range of Mabs (e.g. Mabs Cc9, HC2, IB8, K3D4) were selected which reacted with the 37 kDa major polypeptide described by Penn et al. (1985a). Mabs which reacted with the other components, the 31·5 kDa polypeptide (JD11), and Mabs that reacted with both the 31·5 and the 33·5 kDa polypeptides (Sc12) were also isolated (data not shown).
Mab (K2F1) which recognized antigenic sites on the 34, 33.5 and 31.5 kDa polypeptides was obtained. Further Mabs derived from the fusions L (LB12), M (ME2) and N (NG4) (Table 1) were ELISA- and dot-blot-positive but reacted only very weakly at 1 in 5 dilutions by Western blotting (data not presented); surprisingly, these Mabs were obtained from mice immunized with denatured polypeptide eluted from SDS-PAGE gels and were exposed to native antigen only in the final boost before fusion. A number of fusions following different immunization protocols were done in order to stimulate the generation of a range of anti-37 kDa Mabs useful both for structural analysis and in diagnosis.

Antigenic relatedness between spirochaetes

Sera and Mabs were assessed for their ability to react with the polypeptide antigens of treponemes other than *T. pallidum* (Nichols). The analysis was done by immunoblotting. hRSS contained cross-reactive antibody to both *T. phagedenis* and *T. hyodysenteriae* (Fig. 4). At least six strongly reactive bands were identified in *T. phagedenis*, three of which are associated with the axial filament of this organism (33, 35 and 38.5 kDa). The Mab IB8 (IgG 1) reacted very strongly not only with the major 37 kDa polypeptide of *T. pallidum* but also with the equivalent polypeptide in *T. phagedenis* (38.5 kDa). Cross-reactivity with *T. hyodysenteriae* was also observed with hRSS and Mabs. Mab Cc9 (IgM against 37 kDa polypeptide) reacted only weakly with a band of 35.5 kDa in *T. hyodysenteriae*. The same polypeptide was recognized by both the hRSS and the anti-axial-filament sera; this polypeptide is considered to be the major axial
filament polypeptide of *T. hyodysenteriae* since Coomassie-blue-stained gels containing purified *T. hyodysenteriae* axial filaments (provided by Dr R. M. Lemcke) contained a predominant band of this size (unpublished observations). The remainder of the Mabs reported here were highly specific for *T. pallidum* and even at high concentrations (1 in 5 dilution) did not cross-react with the other treponemal species investigated. The diversity of antigens recognized by the Mabs possibly reflects the commonality and conserved nature of the axial filament antigens within the genus *Treponema* and demonstrates, especially with K2F1, the presence of shared antibody-binding sites between polypeptides of the *T. pallidum* axial filament.

**Immunogold electron microscopy**

The anti-37kDa Mab Cc9 was strongly reactive with the surface of the axial filament of *T. pallidum*, unequivocally demonstrating the location of this polypeptide. The other Mabs against the 37 kDa antigen showed variability in their reactivity by immunogold techniques,
T. pallidum axial-filament Mabs

Fig. 5. Immunogold electron microscopy of Mabs reacted with the axial filaments of T. pallidum (the outer membrane was removed by detergent treatment). (a) NS-1 supernatant control; (b) Mab Cc9; (c) Mab K3D4. Bars, 0.2 μm.

underlining in the cross-reactivity studies that a diverse range of Mabs against many epitopes had been generated. Cc9 and K3D4 were the most avid immunogold-reactive Mabs (Fig. 5), suggesting that they were directed against external epitopes on the surface of the endoflagella. In a small percentage of the population a degree of heterogeneity of binding with the Cc9 Mab was observed, which has prompted the further investigation of the composition of the axial filament to confirm that the 37 kDa polypeptide is indeed the sheath protein which surrounds the core structure constructed from the remaining polypeptides (Cockayne et al., 1987). None of the other Mabs against the other polypeptides were immunogold-positive when reacted with intact axial filament, and therefore the location of these polypeptides as axial filament components could not be confirmed by the methods employed.

T. pallidum immobilization

All the Mabs tested in the immobilization assay were unable to immobilize T. pallidum in vitro. The periplasmic location of the axial filaments could explain the lack of reactivity, as the Mabs would be excluded from binding in intact organisms. The polyclonal sera against the purified axial filament were not tested as they contained antibody against non-axial-filament polypeptides. One of the contaminating antibody specificities was directed against the 44 kDa antigen, which has been shown to elicit immobilizing antibody (Bailey et al., 1987).

DISCUSSION

A wide variety of anti-T. pallidum antibodies are present in the sera of infected human patients or experimentally infected rabbits (Fig. 2). Certain polypeptides have been investigated and identified, allowing a logical and more accurate assessment of the role of particular antigens in the development of both immunity and humoral responses (Penn et al., 1986). A number of immunodominant antigens have been identified which elicit the stimulation of antibody to specific antigens early in infection. The most dominant of these antigens are the 37 kDa axial filament polypeptide (Penn et al., 1985a) and the 47 kDa outer-membrane-associated polypeptide (Penn et al., 1985b). Mice however, when injected with live T. pallidum do not
develop any clinical signs of disease but do react immunologically following contact with antigen. Saunders & Folds (1986) recently demonstrated the development of a progressive immune response to treponemal antigens in mice injected with live treponemes but not in those injected with dead ones, suggesting that this pathogen is able to survive and multiply in the mouse, but to a lesser extent than in the rabbit. In this work we have shown that mice apparently do not respond as well to T. pallidum antigens as do rabbits, eliciting antibody to only a limited range of antigens. When 10⁸ live treponemes were injected subcutaneously or intraperitoneally into mice only very weak ELISA titres were recorded (unpublished observations) with the predominant antibody detectable being directed against the 47 kDa antigen (Fig. 1, Va). This poor response has been described previously (McLeod & Magnuson, 1951; Ohta, 1972; Robertson et al., 1982), indicating that a large inoculum of live organisms must be repeatedly delivered in order to evoke even a low-titre response. The inclusion of treponemes in adjuvant enhanced the antibody titre, and the sonication of the bacterial suspension increased the range of antibodies stimulated. However, we found that hybridomas generated from mice immunized with unmodified T. pallidum predominantly secreted antibody to the 47 kDa antigen regardless of the route or rate of inoculation. Also, by immunization protocols using unfractionated bacteria as the antigen source we were virtually unable to stimulate specific anti-37 kDa antibody (Fig. 1, 11).

The fractionation of the bacteria and the use of a variety of immunization protocols (Table 1) stimulated antibody specific to axial filament antigens (Fig. 1, III, IV, Vb), indicating that the polypeptides were indeed highly antigenic in mice but were in some way masked when delivered in the form of whole or sonicated T. pallidum cells. Table 1 outlines the immunization schedules adopted from which anti-37 kDa Mabs were selected. The Mabs that reacted most strongly by immunogold electron microscopy were generated from mice immunized only with native antigen (i.e. Cc9 and K3D4), although not all the Mabs produced with these immunization schedules were immunogold-reactive. The use of denatured 37 kDa polypeptide as an antigen was very effective in stimulating mouse antibody (Fig. 1, IV), but the Mabs produced were only very weakly immunoblot-positive, demonstrating that additional epitopes are present on this polypeptide which are not detected by ELISA or the dot-blot ELISA, which only presents native epitopes. This observation is supported by the Mabs generated following fusion M (i.e. ME2), in which mice were immunized with the 60 kDa polypeptide cut from gels and boosted once only with purified axial filaments (Table 1). The 60 kDa polypeptide has been identified as the common bacterial antigen (Hindersson et al., 1984) and is not a component of the axial filament. Thus the Mabs produced that reacted by dot-blot with the axial filament resulted from the final boost and not from the initial immunization protocol. The use of immunization schedules including preparations of purified axial filaments or their constituent polypeptides is considered essential for the production of Mabs to this structure.

From the studies done with the Mabs against the axial filament of T. pallidum we have been able to identify a number of component polypeptides and by immunogold labelling unequivocally confirm the observations of Penn et al. (1985a) that the 37 kDa polypeptide is the major component of this structure. The range of Mabs produced defines a number of different epitopes as determined by the diversity observed in their cross-reactivity with other treponemes, and differential reactivity by gold labelling and immunoblotting. These findings indicate that the denatured polypeptide is strongly antigenic and that there are a number of internalized antibody-binding sites on the native molecule; it is therefore essential that selective immunization protocols be adopted in order to generate Mabs which bind to epitopes on the surface or within the axial filament. The combination of the appearance of strong antibody responses to the 37 kDa polypeptide in both infected rabbits and man, and the demonstration that a variety of epitopes are detectable on this molecule, suggests the usefulness of these reagents in the immunodiagnosis of syphilis.

This research was funded in part by the British Technology Group.
REFERENCES


