Epitopes of *Bordetella pertussis* lipopolysaccharides as potential markers for typing of isolates with monoclonal antibodies

Karine Le Blay, Martine Caroff, Frédéric Blanchard, Malcolm B. Perry and Richard Chaby

**INTRODUCTION**

*Bordetella pertussis*, the aetiological agent of whooping cough, expresses several antigenic proteins (pertussis toxin, adenylate cyclase, filamentous haemagglutinin) and, like all other Gram-negative bacteria, an ‘endotoxin’ which is actually a family of structurally related lipopolysaccharides (LPSs). We established previously the existence of two types of variations in preparations of *B. pertussis* LPSs: one is due to the presence or absence of a phosphate group on a 3-deoxy-β-manno-2-octulosonic acid (Kdo) unit in the ‘core’ region (Le Dur et al., 1980), and the other is due to the presence or absence of a distal trisaccharide (Caroff et al., 1990). The latter finding explained the presence of two bands (designated A and B) in silver-stained SDS-PAGE of *B. pertussis* cell lysates (Peppler, 1984). The observation that LPSs from *B. pertussis* (Ackers & Dolby, 1972; Mountzouros et al., 1992) and from other bacteria (Terashima et al., 1991) can elicit the production of protective antibodies, and the recent recrudescence of whooping cough in several developed countries, prompted us to produce anti-LPS monoclonal antibodies (mAbs) able to detect antigenic modifications in LPSs from variant *B. pertussis* strains, or from related *Bordetella* species (*B. parapertussis*, *B. bronchiseptica*).

The production of mAbs to *B. pertussis* LPSs has been reported by various authors (Mountzouros et al., 1992; Frank & Parker, 1984; Gustafsson et al., 1988; Li et al., 1988; Archambault et al., 1991; Martin et al., 1992). However, the characterization of the epitopes recognized by these mAbs was hampered by the absence of structural data on the antigens used. Our structural studies (Chaby & Caroff, 1988; Lebar et al., 1994; Di Fabio et al., 1992) of different *Bordetella* LPSs allowed us to re-examine this problem. In this paper we describe the production of mAbs that we used to characterize three antigenic epitopes of *B. pertussis* LPSs. These mAbs may be useful for the rapid screening and typing of clinical isolates.

**METHODS**

**Chemicals and reagents.** Polyoxyethylene-sorbitan monolaurate (Tween 20), 2,6,10,16-tetramethylpentadecane (pristane), horseradish-peroxidase-labelled goat anti-mouse Ig.
anti-mouse H chain of different classes and subclasses, and the peroxidase substrates o-phenylenediamine dihydrochloride and 3,3'-diaminobenzidine were from Sigma. Polyisobutyl methacrylate was from Aldrich, gelatin from Rousselot Kuhlmann, and polyethylene glycol 1500 from Boehringer.

**Lipopolysaccharides.** The *B. pertussis* LPSs from the vaccine strain 1414, and from the variant strain A100, were isolated by the phenol/water extraction procedure, and purified as described previously (Le Dur et al., 1980; Caroff et al., 1990). The LPSs from *B. parapertussis* (strain ATCC 15989 from the American Type Culture Collection) and *B. bronchiseptica* (strain NRCC 4170 from the National Research Council Veterinary Collection) were prepared by extraction of the cells by the modified enzyme/phenol/water method and ultracentrifugation of the concentrated, dialysed, aqueous phases (105 000 g, 4 °C, 12 h) (Johnson & Perry, 1976). The putative structures of these four LPSs (shown in Fig. 1) were obtained in the present study also suggest that the core of LPS BpA100 contains one (or more) HF-labile residue(s).

<table>
<thead>
<tr>
<th>LPS</th>
<th>Structure</th>
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<tbody>
<tr>
<td>Bp1414</td>
<td>GlcNAc→2,3-NAcManA→FucN(Ac)Me</td>
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<tr>
<td></td>
<td>(2,3-NAcGalA)→Hep→Kdo→Lipid A</td>
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<tr>
<td>BpA100</td>
<td>GlcNAc→2,3-NAcManA→FucN(Ac)Me</td>
</tr>
<tr>
<td></td>
<td>(2,3-NAcGalA)→Lipid A</td>
</tr>
<tr>
<td>B. bronchi</td>
<td>GlcNAc→2,3-NAcManA→FucN(Ac)Me</td>
</tr>
<tr>
<td></td>
<td>(2,3-NAcGalA)→Hep→Lipid A</td>
</tr>
<tr>
<td>B. para</td>
<td>GlcNAc→2,3-NAcManA→FucN(Ac)Me</td>
</tr>
<tr>
<td></td>
<td>(2,3-NAcGalA)→Hep→Lipid A</td>
</tr>
<tr>
<td>PeS-BSA</td>
<td>GlcNAc→2,3-NAcManA→FucN(Ac)Me</td>
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<td></td>
<td>2,5-anhydroMan→BPA</td>
</tr>
<tr>
<td>DIS-BSA</td>
<td>2,3-NAcManA→FucN(Ac)Me</td>
</tr>
<tr>
<td></td>
<td>O-CH₂-CHOH-CH₂→BPA</td>
</tr>
<tr>
<td>GlcNAc-BSA</td>
<td>GlcNAc→[cysteamine/glutaredyhyde arm]</td>
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**Fig. 1.** Putative structures of the antigens used. LPSs Bp1414, BpA100, B. bronchi, and B. para, LPSs from *B. pertussis* strain 1414, *B. pertussis* strain A100, *B. bronchiseptica* strain NRCC 4170 and *B. parapertussis* strain ATCC 15989, respectively. NA/LPS Bp1414, nitrous-acid-deaminated LPS Bp1414. GlcNAc-BSA, Dis-BSA, and PeS-BSA, conjugates consisting of BSA covalently coupled to GlcNAc, or to the indicated disaccharide and pentasaccharide, respectively. FucN(Ac)Me, N-acetyl-N-methylamyl glucosaminuronic acid; Glc, glucose; GlcA, glucuronic acid; GlcN, glucosamine; GlcNAc, N-acetylgalactosamine; Hep, heptose; Kdo, 3-deoxy-o-manno-2-octulosonic acid; Man, mannose; 2,3-NAcGalA, 2,3-dideoxy-2,3-di-N-acetylgalacturonic acid; (*+) Phosphate and phosphorylethanolamine groups, present in all LPS preparations, are not represented. Unpublished studies suggest that the core of LPS B. bronchi, is substituted with at least one phosphate group, and that the core of LPS B. para, is shorter than that of LPS Bp1414. The results obtained in the present study also suggest that the core of LPS BpA100 contains one (or more) HF-labile residue(s).

**Cleavage of LPSs with nitrous acid.** The LPSs from the *B. pertussis* strains 1414 (LPS Bp1414) and A100 (LPS BpA100) were incubated (5 g l⁻¹) for 4 h at 20 °C in a freshly prepared mixture of water/5% (w/v) sodium nitrite/30% (v/v) acetic acid (1:1:1, by vol.), as described previously (Caroff et al., 1990; Deprun et al., 1993). The suspensions were ultracentrifuged (200 000 g for 2 h) and the fragments containing the Kdo-lipid A region (NA/LPS Bp1414 and NA/LPS BpA100) were isolated in the pellets. If TLC (solvent A; see below) indicated incomplete deaminative cleavage, the pellet was treated with nitrous acid for a second time. The structure of NA/LPS Bp1414 was directly coupled to GlcNAc-BSA, or as yet unpublished studies (M. Caroff, D. Karibian, H. Zarrour, J. C. Richards and M. B. Perry, indicated in Fig. 1.

**Analyses.** The heptose content of the preparations was determined by quantitative estimation of heptitol acetate by GLC, after acid hydrolysis (2 M HCl, 2 h, 100 °C) of the preparations, further reduction (1 M NaBH₄, 4 h, 20 °C) of the sugars, and peracetylation of their alditols. Methylation, acetylations and GLC analyses (25 m×0.32 mm capillary column coated with a BP 10 bonded phase) were carried out as described previously (Caroff et al., 1990).

**Preparation of the PeS-BSA conjugate.** The PeS isolated by nitrous acid deamination of LPS Bp1414 was directly coupled to BSA by reductive amination, as described by Roy et al. (1984). Briefly, PeS (5 mg) was incubated for 48 h at 37 °C with BSA (10 mg) and sodium cyanoborohydride (20 μmol) in 0.2 M phosphate buffer (1 ml, pH 8). After neutralization with acetic acid, the solution was made up to 0.15 M NaCl and the conjugate was separated from uncoupled polysaccharide by...
chromatography on a Sephadex G-150 column. Elution was monitored by absorption at 220 nm. The purified PeS-BSA conjugate (36 mg) contained 10% (w/w) PeS according to its heptose content. The structure of PeS-BSA is indicated in Fig. 1.

**Diphosphorylation by HF treatment.** LPS Bp1414, LPS BpA100 and NA/LPS BpA100 (10 g l-1) were incubated in aqueous HF (50%, v/v) at 4°C for 48 h in sealed polyethylene tubes (Lipkin et al., 1969). After removal of HF under a stream of nitrogen at room temperature, the dried residues were dissolved in water and dialysed against distilled water. HF/LPS Bp1414, HF/LPS BpA100 and HF-NA/LPS BpA100 were recovered by lyophilization.

**Preparation of the DiS-BSA conjugate.** After hydrolysis of a 1:5% (w/v) solution of HF/LPS Bp1414 in sodium acetate (pH 4.5) for 2 h at 100°C in the presence of 1% (w/v) SDS (Caroff et al., 1988), the polysaccharide fragment (HF/PS) was isolated by chromatography on a Sephadex G-50 column. Smith degradation of HF/PS was carried out by incubations (20°C) of an aqueous solution (4 g l-1, 33 ml) of the polysaccharide with 0.7 mM sodium periodate (48 h), 15 mM ethylene glycol (15 min), 75 mM sodium borohydride (18 h), and 0.1 M acetic acid (1 h), successively. The material was desalted by filtration (Diaflow YC-05 membrane, Amicon) and submitted to chromatography on a Biogel P2 column eluted with water. The fragment consisting of the disaccharide 2,3-NAcManA-Fuc(NAc)Me-erythritol (DiS), identified by NMR analysis (data not shown) was isolated and lyophilized. The erythritol unit of DiS was oxidized by incubation of 3.65 mg DiS for 20 min at 20°C in 0.7 mM sodium periodate (48 h), 15 mM ethylene glycol (15 min), 75 mM sodium borohydride (18 h), and 0.1 M acetic acid (1 h), successively. The material was desalted by filtration (Diaflow YC-05 membrane, Amicon) and submitted to chromatography on a Sephadex G-50 column. The structure of the DiS-BSA conjugate is indicated in Fig. 1.

**Preparation of the GlcNAc-BSA conjugate.** N-Acetylglucosamine-b-allylsaldehyde was synthesized by Dr D. Charon (Châteanay-Malabry, France). Covalent coupling to cysteamine (Roy et al., 1984; Lee & Lee, 1974) was performed by incubation (18 h, 20°C) of a 2 M solution of this compound with a 7.7 M solution of cysteamine hydrochloride. The propylaminoethanethiol derivative of GlcNAc was isolated by chromatography on a column of AG 50 WX8 (H+) resin, eluted with 0.1 M sodium periodate (48 h), 15 mM ethylene glycol (15 min), 75 mM sodium borohydride (18 h), and 0.1 M acetic acid (1 h), successively. The material was desalted by filtration (Diaflow YC-05 membrane, Amicon) and submitted to chromatography on a Sephadex G-50 column. The structure of the GlcNAc-BSA conjugate is indicated in Fig. 1.

**Animal immunizations.** Groups of four BALB/c mice (R. Janvier, Le Genest Saint-Isle, France) were immunized at different intervals with B. pertussis LPS (1 g ml-1, i.p.), or with phenol-killed and lyophilized B. pertussis cells (1 g ml-1, i.v.). Each preparation was suspended in 200 µl nonpyrogenic 0.15 M NaCl solution (Bioseda, Malakoff, France). Sera were collected 7 d after each injection.

**Production of mAbs.** The mouse which produced the highest titre of anti-LPS antibodies (detected after the second immunization) was selected for fusion. Three days after the last injection, the spleen cells were fused at a 10:1 ratio with SP2/0-Ag.14 myeloma cells, in 45% (w/v) polyethylene glycol 1500, as described previously (Girard & Chaby, 1985). Fusion products were cultured in a hypoxanthine/azaserine (50 µM:10 µM) selection medium. Resulting colonies were tested for secretion of anti-LPS antibodies, and cloned by limiting dilution (Nowinsky et al., 1979). Clones were expanded as ascites by injection (5 x 106 cells per mouse, i.p.) in BALB/c mice treated with pristane (0.5 ml, i.p.) two weeks previously.

**ELISA.** Sera and cell culture supernatants were assayed for anti-LPS or anti-BSA-conjugate antibodies by ELISA. The 96-well Immulon-A plates (Greiner Labortecnik) were coated with the antigen (100 µl of 20 µg ml-1 suspensions in 50 mM Tris/HCl buffer plus 20 mM MgCl2, pH 9.6). After overnight incubation at 20°C with gentle stirring, the unbound antigen was washed away with 0.1% Tween 20 in PBS (TWEEN buffer), and the plates were incubated for 1 h at 37°C with 0.25% (w/v) gelatin in the same buffer. The plates were rewashied with this solution and the sample of serum or culture supernatant (diluted in the same solution) added to each well. After incubation for 2 h at 37°C, the plates were washed with the Tween buffer, and the peroxidase-labelled goat anti-mouse Ig reagent added. After incubation at 37°C for 90 min, the plates were washed and developed by adding 100 µl of a freshly prepared solution of o-phenylenediamine (0.5 mg ml-1) in 0.05 M phosphate/citrate buffer (pH 5) containing sodium perborate (0.03%, v/v) as a substitute for hydrogen peroxide. After 15 min at room temperature, the reaction was stopped with a solution of 0.5% (w/v) sodium sulphite in 1 M H2SO4 (50 µl per well). The plates were scanned at 490 nm in a Dynatech MR 5000 spectrophotometer. Antibody isotypes were determined by the same ELISA procedure, by using peroxidase-labelled goat antibodies against mouse Ig of different classes and subclasses.

reactivities with the immunizing LPS (Bp1414), and with LPSs from other Bordetella strains (BpA100, B. bronchiseptica). The results obtained with two representative mice (Fig. 2) show that a marked reactivity with the homologous LPS (Bp1414) was always obtained after the third injection, whereas none of the sera reacted with the variant strain A100 of B. pertussis, even after prolonged immunizations. Reactivities of sera with the B. bronchiseptica LPS were more variable: some sera (like those from mouse 1) reacted with this LPS, whereas others (like those from mouse 2) did not. These results show that although the three LPSs share some common substructures (Fig. 1), the immunogenicities of these substructures in mice are very different.

Production of hybridomas and ascitic fluids

Mice were injected, at 14 d intervals, either with phenol-killed B. parapertussis cells, or with B. pertussis LPS. Fusion experiments were performed 3 d after the second injection with animals immunized with killed cells, and 3 d after the fifth injection with animals immunized with LPS. Hybridomas were first screened for their reactivity with the LPS preparation used for immunization, and then re-screened for the absence of cross-reactivity with a lipid A preparation obtained from the B. pertussis LPS. Among the remaining hybridomas, three clones of high reactivity with LPS were selected for this study (Table 1). One clone (60.5) was produced with a mouse immunized with killed B. pertussis organisms, and two clones (P1P3 and D7) were obtained with mice immunized with the B. pertussis LPS. Analyses of the isotypes of the mAbs prepared from ascitic fluids indicated that P1P3 and 60.5 are IgM, whereas D7 is an IgG3 (Table 1).

Reactivity of the mAbs with unmodified Bordetella LPSs

The specificities of the three mAbs were analysed by their ability to interact with the LPSs from B. pertussis 1414, B. pertussis A100, B. bronchiseptica and B. parapertussis. Results obtained by ELISA (Fig. 3) showed clearcut differences in the specificities of these mAbs. All three mAbs reacted...
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4. Analysis of different LPSs by TLC. LPSs from *S. minnesota* Rd1 (A), *S. typhimurium* Rcl (B), *B. pertussis* 1414 (C), and the nitrous-acid-treated LPS from *B. pertussis* 1414 (D) were analysed by TLC in isobutyric acid/1 M ammonium hydroxide (5:3, v/v). LPS bands were revealed by charring with sulfuric acid (a) or by immunostaining with mAb P1P3 (b). With LPS Bp1414 (Fig. 3a), whereas only two of them (mAbs P1P3 and 60.5) interacted with the *B. bronchiseptica* LPS (Fig. 3c), and only one (mAb P1P3) reacted with the *B. parapertusis* LPS (Fig. 3d); none of the three mAbs could recognize LPS BpAlOO (Fig. 3b). These results clearly show that although the three mAbs interact with similar intensities with the LPS from strain 1414 of *B. pertussis*, they do so by binding to distinct epitopes.

**Reactivity of the mAbs with rough-type LPSs**

We established previously (Caroff *et al.*, 1990) that a rough-type LPS consisting of a pentasaccharide linked to lipid A (NA/LPS Bp1414) can be isolated from the nitrous-acid-treated LPS of *B. pertussis* 1414 (Fig. 1). The reactivity of mAb P1P3 with NA/LPS Bp1414 was compared to its reactivity with untreated LPS Bp1414, and with rough-type LPSs from *Salmonella* (*S. minnesota* Rd1 and *S. typhimurium* Rcl). Reactivities were analysed by immunostaining the LPS bands after migration on TLC. The results (Fig. 4) show that mAb P1P3 reacted exclusively with the untreated LPS Bp1414, and with rough-type LPSs from *Salmonella*. Identical results (not shown) were obtained with the two other mAbs, 60.5 and D7. These results demonstrate that the mAbs interact neither with the Kdo-lipid A region of the LPSs, nor with the four other sugar units which are vicinal to this region in LPS Bp1414 (GlcA-Hep-Hep, Glc).

**Reactivity of mAb P1P3 with HF-treated LPS BpA100**

The absence of reactivity of P1P3 with the short-chain lipopolysaccharides LPS BpA100 (Fig. 3) and NA/LPS Bp1414 (Fig. 4) could suggest that this mAb reacts with distal carbohydrate units present only in *Bordetella* LPSs with longer chains. Another experiment (Fig. 5), which confirmed this observation, provided an additional and unexpected result: after treatment of LPS BpA100 with HF, the isolated material (HF/LPS-BpA100) was recognized by mAb P1P3, whereas the material obtained after sequential treatments with nitrous acid and HF (NA-HF/LPS BpA100) was no longer recognized by P1P3 (Fig. 5). These results suggest (1) that the epitope recognized by P1P3 is released during nitrous acid treatment, and (2) that an HF-labile substituent modifies this epitope in the variant A100 strain of *B. pertussis*.

**Reactivity of the mAbs with carbohydrate substructures coupled to BSA**

The reactivities of the mAbs with distal carbohydrate substructures were analysed with conjugates consisting of the terminal monosaccharide GlcNAc, the disaccharide DiS, and the pentasaccharide PeS, covalently coupled to BSA. The reactivities of the three mAbs with these conjugates, and with unmodified BSA, were determined by ELISA. The results (Fig. 6) show that none of the mAbs reacted with BSA and DiS-BSA. Only one mAb (60.5) reacted with GlcNAc-BSA. The conjugate PeS-BSA, which carries the larger oligosaccharide substructure, was recognized by both P1P3 and 60.5. A significant but much lower reactivity was also observed with mAb D7.

**DISCUSSION**

The aim of the present study was to produce mAbs against distinct epitopes of the *B. pertussis* LPS that can potentially recognize the expression of atypical forms of...
Fig. 6. Reactivity of mAbs P1P3, 60.5 and D7 with different BSA-conjugates. Plates coated with PeS-BSA, GlcNAc-BSA, DIS-BSA, or BSA alone, were incubated with ascitic fluids (dilution 1/100) from clones P1P3, 60.5 and D7. The binding of the mAbs was determined by ELISA. Values represent the mean ± SD of triplicate determinations.

The specificity of mAb 60.5 can be easily established. This mAb reacted with the LPSs from *B. pertussis* strain 1414, three mAbs, termed 60.5, D7 and P1P3 were produced.

The specificity of mAb 60.5 can be easily established. This mAb reacted with the LPSs from *B. pertussis* strain 1414, *B. bronchiseptica*, but not with the LPSs from *B. pertussis* A100 and *B. parapertussis* (Fig. 3). The only structure present in the two former and absent from the two latter LPSs is the distal trisaccharide GlcNac → 2,3-NAcManA → FucN(Ac)Me. The reactivity of 60.5 with the GlcNAc-BSA conjugate (Fig. 6) confirmed this assumption and shows that the epitope recognized by mAb 60.5 consists of the terminal unit(s) of this distal trisaccharide (Fig. 7).

The reactivity of 60.5 with GlcNAc-BSA is, however, markedly lower than that obtained with the intact LPS. This may suggest that the presence of the next carbohydrate unit (2,3-NAcManA) can optimize the reactivity, and thus may be partially involved in the recognized epitope.

The specificity of mAb D7 can also be deduced from the data in Figs 3 and 6. D7 reacts strongly with the *B. pertussis* 1414 LPS, reacts only at high concentrations with the *B. bronchiseptica* LPS, and does not react with the *B. parapertussis* LPS. This means that the presence of the O-side chain, consisting of 2,3-NAcGalA repeating units, blocks the reactivity of D7 and thus modifies the epitope recognized by this mAb. The low reactivity of D7 with the PeS-BSA conjugate and with the *B. bronchiseptica* LPS shows, however, that the presence of the distal trisaccharide slightly enhances the reactivity of D7. Taken together, these results indicate that the epitope recognized by D7 is affected by the addition of the O-side chain and that a portion of the distal trisaccharide could be involved in this epitope. This suggests therefore that this epitope may consist of the FucN(Ac)Me → GlcN disaccharide (Fig. 7).

The third mAb, P1P3, reacted with a substructure shared by the LPSs of three conventional strains of *Bordetella*: *B. pertussis* 1414, *B. bronchiseptica* and *B. parapertussis* (Fig. 3), thus indicating that the O-chain is not involved in this reaction. The reactivity of P1P3 with the *B. parapertussis* LPS, and with the HF-treated LPS from *B. pertussis* A100 (HF/LPS-BpA100), shows that P1P3 does not react with the distal trisaccharide, which is absent from these two LPSs. Therefore, the epitope recognized by P1P3 is apparently located in the common ‘core’ structure (Fig. 1). The absence of reactivity of P1P3 with the rough-type fragment (NA/LPS Bp1414) isolated after nitrous acid cleavage (Fig. 4) indicated that the epitope is localized in the terminal sugar units of the core which are released during nitrous deamination (GlcN, GalNA, or Hep → GlcN).

The interaction of P1P3 with the PeS-BSA conjugate (Fig. 6) demonstrated that the epitope is a part of the Hep → GlcN moiety, which is the only substructure present in both the PeS-BSA HF/LPS-BpA100. However, the epitope recognized by P1P3 cannot be the terminal heptose alone, since P1P3 does not react with the rough-type LPS from *S. minnesota* Rd1 (Fig. 4), which contains a terminal unsubstituted heptose unit. Therefore, we suggest that a region of the glucosamine unit to which the terminal heptose is attached also contributes to the epitope recognized by mAb P1P3. The carbon atoms 3, 4, and 6 of this glucosamine unit could be involved in the epitope since this glucosamine substructure remains unmodified during the nitrous deamination of this residue, leading to 2,5-anhydromannose.

Whereas the inability of mAbs D7 and 60.5 to react with LPS BpA100 can be easily ascribed to the absence of the corresponding epitopes, the inability of mAb P1P3 to react with this LPS (Fig. 3b) was unexpected. A similar absence of reactivity with LPS BpA100 was also systematically observed with sera from mice immunized with LPS Bp1414 (Fig. 2). This may suggest that the ‘core’ of the LPS from the variant strain A100 of *B. pertussis* could be different from that of the more conventional, vaccine strain 1414. However, in a previous study (Caroff et al., 1990), we established that the only detectable difference in the polysaccharide fragments isolated from LPS Bp1414 and LPS BpA100 after HF treatment and mild acid hydrolysis was a loss of the distal trisaccharide in the latter, which suggests that the ‘cores’ of the HF-treated
LPSs are identical. This observation prompted us to examine the influence of HF treatment on the antigenicity of the LPS. We found that HF/LPS-BpA100 was recognized by P1P3 (Fig. 5). This could indicate that the LPS of this variant strain of *B. pertussis* carries an HF-labile substituent in its core region. Structural analyses should soon shed light on this point.

Regarding the immunogenicity of *B. pertussis* LPS substructures, the observation that sera from some mice immunized with this LPS did not react with the *B. bronchiseptica* LPS (Fig. 2) indicates that the epitopes recognized by P1P3 and 60.5 are less immunogenic than that detectable with D7. The observation that none of these sera could interact with the LPS from the *B. pertussis* A100 strain (Fig. 2), and that we were unable to obtain a mAb that recognized the NA/LPS Bp1414, shows that the carbohydrate region proximal to lipid A is very poorly immunogenic.

The anti-*B. pertussis* mAbs produced in this study, in addition to the anti-*B. parapertussis* and anti-*B. bronchiseptica* mAbs prepared previously (Le Blay et al., 1994), represent an efficient tool for routine identification of these organisms in clinical isolates, as well as for the detection of variations in the LPS components of these bacteria. It is noteworthy that these clinical applications of the mAbs should be much easier if the conclusions drawn from the present study, in which experiments were carried out with isolated LPSs, could be extended to LPS in its natural environment, where it is exposed on the surface of intact organisms. It is clear that this situation can modify the accessibility of certain LPS epitopes, and thus limit the use of the corresponding antibodies. Experiments designed to analyze the interactions of the mAbs with intact organisms are now planned.

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**REFERENCES**


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