Role of the polymorphic region 1 of the *Bordetella pertussis* protein pertactin in immunity

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In several countries pertussis is re-emerging, despite a high vaccination coverage. It is suggested that antigenic divergence between *Bordetella pertussis* vaccine strains and circulating strains, in particular with respect to pertactin, has contributed to pertussis re-emergence. Polymorphism in pertactin is essentially limited to region 1, which is composed of repeats and is located adjacent to an Arg-Gly-Asp motif implicated in adherence. Evidence is provided for the immunological relevance of polymorphism in region 1. Region 1 was found to contain a B-cell epitope recognized in both humans and mice. Furthermore, variation in region 1 affected antibody binding and, in a mouse respiratory infection model, the efficacy of a whole-cell vaccine. Moreover, passive and active immunization indicated that region 1 confers protective immunity. An mAb directed against a linear conserved epitope conferred cross-immunity against isolates with distinct pertactin variants. The results indicate an important role of region 1 of pertactin in immunity.

Keywords: antigenic variation, P.69/pertactin, immunization, protective immunity

INTRODUCTION

*Bordetella pertussis* is the principal aetiological agent of whooping cough or pertussis, a respiratory disease that is most severe in infants and young children. In the pre-vaccination era, nearly every child contracted whooping cough and this infection was a major cause of infant death throughout the world (Willems & Mooi, 1996). For about 40 years, widespread immunization of young children with whole-cell pertussis vaccines (WCVs) has been successful in controlling the disease. However, in recent years the incidence of pertussis has increased in a number of countries, including Australia, Canada, USA and The Netherlands, despite a high vaccination coverage (Andrews et al., 1997; Bass & Stephenson, 1987; Bass & Wittler, 1994; de Melker et al., 1997; DeSerres et al., 1995). We have previously reported that two *B. pertussis* antigens implicated in protective immunity, pertactin and pertussis toxin, are polymorphic (Mooi et al., 1998). Three protein variants of both pertactin and pertussis toxin were found in Dutch clinical isolates collected from 1949 to 1996. Temporal trends in the frequencies of pertactin and pertussis toxin variants indicated a divergence between vaccine strains and clinical isolates (Mooi et al., 1998, 1999).

A number of studies have shown that the *B. pertussis* fimbriae, filamentous haemagglutinin, pertussis toxin and pertactin can induce protection in both animals and humans (Sato & Sato, 1984; Zhang et al., 1985; Kimura et al., 1990; Robinson et al., 1985; Shahin et al., 1990). Therefore, these proteins have been included in acellular pertussis vaccines. The importance of pertactin as a protective antigen is particularly convincing: the presence of antibodies against pertactin has been shown to correlate with clinical protection (Cherry et al., 1998; Storsaeter et al., 1998). Furthermore, acellular vaccines containing pertactin have been reported to give better protection than vaccines without this component (Hewlett, 1997; Plotkin & Cadoz, 1997). Pertactin belongs to a family of secreted proteins designated autotransporter proteins (Henderson et al., 1998). It is produced as a large (910 aa) precursor molecule which is...
proteolytically processed at its N and C termini to produce P.69 and P.30 (Fig. 1), which are located at the cell surface and in the outer membrane, respectively (Charles et al., 1994). P.69/pertactin (henceforth designated as pertactin), but not P.30, is included in acellular vaccines. It contains the amino acid triplet arginine-glycine-aspartic acid (RGD), a sequence motif which functions as a cell-binding site in a number of mammalian proteins. It has been shown that the pertactin RGD sequence is also involved in adherence to host cells (Leininger et al., 1992).

Polymorphism in pertactin is mainly found in two regions composed of proline-rich repeated sequences, designated regions 1 and 2 (Fig. 1) (Mooi et al., 1998). Most variation is found in region 1 which is located proximally to the N terminus, flanking the RGD sequence, and consists of repeats of the Gly-Gly-X-X-Pro (GGxxP) motif. The crystal structure of pertactin predicts that both the RGD sequence and the GGxxP motif are exposed at the surface of pertactin (Emsley et al., 1996). Region 2 is located more towards the C terminus and is composed of repeats of Pro-Glu-Pro (PQP).

Although epidemiological studies suggest that variation in surface antigens increases the fitness of B. pertussis, in terms of epidemiological spread, in immunized populations no direct evidence has yet been presented for its immunological relevance (Mooi et al., 1998; van Loo et al., 1999). Here we studied the immunological importance of the polymorphism in pertactin. Since variation in pertactin is essentially limited to region 1, we focused on this domain.

**METHODS**

**Bacterial strains and plasmids.** Bordetella pertussis, Bordetella parapertussis and Bordetella bronchiseptica strains used in this study are indicated in Table 1. The clinical isolates used for mouse challenge studies were isolated from Dutch patients in the period 1949–1996. Bordetella strains were grown on Bordet–Gengou (BG) agar (Difco Catalogue no. 0048-17-5) supplemented with 1% (v/v) glycerol and 15% (v/v) sheep blood at 35 °C for 3 d. Escherichia coli strains DH5α and BL21(DE3) were used for the propagation of plasmids. E. coli strains were routinely grown in L-broth or on L-agar supplemented with antibiotics. The pMAL-c2 vector was purchased from New England Biolabs.

**DNA sequence determination.** The pertactin genes of isolates used in this study were sequenced previously or in this study (Mooi et al., 1998, 1999; Mastrantonio et al., 1999; Li et al., 1991) (Table 1). Except for the clinical isolates, the prn genes were sequenced completely on both strands. In the case of the clinical isolates, region 1 was sequenced in all isolates, whereas region 2 was sequenced in 65% of the isolates. No polymorphism in region 2 was detected. DNA was isolated using standard procedures. DNA sequence determination was performed by direct sequencing of PCR products, using the ABI Prism Dye Terminator Cycle Sequencing Ready Reaction Kit (Perkin Elmer–Applied Biosystems). The products were analysed on a 373 ABI DNA sequencer (Perkin Elmer).

**Construction of maltose-binding protein (MBP) containing the pertactin polymorphic region.** Part of the ORF of the prn1 gene, encoding the amino acid sequence TIRRGDAP-AGGAVPGGAVPGGAVPGGFGGGPFPV (defined as region 1), was expressed in E. coli as a fusion with the MBP by cloning the corresponding pertactin DNA sequence in the pMAL-c2 vector. The pertactin sequence was obtained by PCR using the primers 5'-CGGGATCCAGATACGCGCGCGGAC-3' and 5'-GGCTTAGAGGGACGGGACC-.
**Table 1. Bordetella strains used in this study**

<table>
<thead>
<tr>
<th>Strain</th>
<th>Original designation</th>
<th>Species</th>
<th>Source</th>
<th>prn type</th>
<th>Accession no.</th>
<th>Reference</th>
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<td>Mooi et al. (1998)</td>
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<td>prn</td>
<td>A0243927</td>
<td>This work</td>
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<tr>
<td>B24†</td>
<td>B. parapertussis</td>
<td>Netherlands</td>
<td>prn</td>
<td>X54547</td>
<td>Li et al. (1991)</td>
<td></td>
</tr>
</tbody>
</table>

* Used for construction of the MBP fusion proteins.
† Used for immunoblotting.
‡ Used for challenge experiments.
§ Accession no. refers to an identical, previously submitted, sequence from another strain.
‖ Isolated from Dutch patients in the period 1949–1996.
¶ In the case of the clinical isolates, prn genes were not sequenced completely. Region 1 was sequenced in all strains, whereas region 2 was sequenced in 20% of the strains. No polymorphism in region 2 was detected.
# Streptomycin-resistant derivative of Tohama.

AGCC-3′ and chromosomal DNA of B. pertussis strain B5 as template. In the primers the BamHI and XbaI restriction sites, introduced to facilitate cloning, are underlined, whereas pertactin-derived sequences are indicated in bold type. PCR products derived from prn2, prn3, prn4 and prn5 were obtained in a similar way using the appropriate strains. Pertactin-derived sequences are indicated in bold type. PCR products were digested with BamHI and XbaI and subsequently ligated in BamHI/XbaI-digested pMAL-c2. All plasmid constructs were checked by DNA sequencing.

**Purification of MBP-pertactin fusion proteins.** E. coli BL21(DE3) cultures (500 ml) carrying pMAL-c2 constructs were grown in 1-l broth medium containing ampicillin (200 µg ml−1) and 1% (w/v) glucose at 37 °C. At an OD600 of 0.5, expression of the MBP fusion protein was induced by the addition of IPTG to a final concentration of 0.3 mM. After induction, growth was allowed to proceed for 2 h and cells were harvested by centrifugation at 13000 g for 20 min. The pellet was resuspended in lysozyme buffer (10 mM NaPO4, pH 7.2, 0.5 M NaCl, 0.25%, v/v, Tween 20, 10 mM EDTA and 0.1 mM PMSF) and stored at −20 °C. After thawing, the cell suspension was sonicated in ice for 4×30 s with 2 min intervals (Branson Sonifier 250, normal tip, 50% output). Extracts were diluted to 100 ml with column buffer A (20 mM Tris, 200 mM NaCl, 1 mM EDTA, pH 7.4) and applied on a 5 ml amylose column. After extensive washing with buffer A, bound proteins were eluted with buffer A, supplemented with 10 mM maltose and fractions of 1 ml were collected. Purity was >99%, as estimated by SDS-PAGE followed by Coomassie brilliant blue (Serva) staining.

**Production of polyclonal and monoclonal antibodies.** Polyclonal antisera against pertactin was prepared by repeated injection (on day 0 and day 28) of BALB/c mice with 5 µg Prn1, kindly provided by Chiron–Biocine, in PBS containing 0.35% (v/v) Alhydrogel. Mice were bled on day 42. mAbs PeM1, PeM2, PeM3, PeM4, PeM5, PeM6 and PeM7 were generated by injection of BALB/c mice subcutaneously three times with purified pertactin mixed with Specol. PeM70, PeM71 and PeM72 were generated similarly using MBP–Prn1 instead of Prn1. For the production of PeM68, mice were injected with MBP–Prn3, PeM80, PeM84 and PeM85 were generated using Prn5. Three days before the fusion, mice were boosted intravenously. Spleens cells were fused with mouse SP2/0 myeloma cells using 50% PEG-1500 (Boehringer Mannheim). Hybridomas secreting antibody to pertactin were selected by ELISA and cloned twice by limiting dilution. mAbs were purified by protein-G affinity chromatography (Pharmacia).

**Synthesis of peptides.** Peptides were assembled by using an automated multiple peptide synthesizer, equipped with a 48-column reaction block (AMS 422; ABIMED Analysentechnik) (Brugghe et al., 1994). Peptides used for epitope mapping were N-terminally acetylated multiple (i.e. octa-meric) antigenic peptides (MAPs) (Tam & Zavala, 1989), prepared as described previously (Roupee van der Voort et al., 1997). The amino acid sequences of the pertactin region 1-specific peptides are listed in Table 2a. Peptides used for immunization contained the same amino acid sequence but were monomeric. These peptides were N-terminally elongated with an S-acetylmethylcaptoacetyl group (Drijfhout et al., 1990) and conjugated to tetanus toxoid (Van der Ley et al., 1991; Brugghe et al., 1994). Control peptide–tetanus toxoid conjugates used for immunizations were derived from meningococcal PorA.

**Immunoblotting.** Cell-free extracts of B. pertussis (isolates B391, B596, B647, B705, B935, B1120, B14 and B24), or purified MBP fusion proteins, were analysed by SDS-PAGE as described by Laemmli (1970). Proteins were transferred to nitrocellulose filters by electrophoretic blotting (Biometra semi-dry blotting) using 5 mA cm⁻² for 30 min. After blocking with...
Table 2. Amino acid sequence of peptides used for epitope-mapping and vaccination

<table>
<thead>
<tr>
<th>Peptide</th>
<th>Amino acid sequence region 1</th>
<th>mAb Minimum epitope recognized</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pep1</td>
<td>ATIRGQAPAGAVP</td>
<td>F8E5</td>
</tr>
<tr>
<td>Pep2</td>
<td>GQAPAGAVPGGAVP</td>
<td>PeM3</td>
</tr>
<tr>
<td>Pep3</td>
<td>GAVPGGAVPGGAVP</td>
<td>PeM4</td>
</tr>
<tr>
<td>Pep4</td>
<td>GQAVPGGAVPGGAVP</td>
<td>PeM68</td>
</tr>
<tr>
<td>Pep5</td>
<td>GAVPGGAVPGGFGPGPVLDGWYGVDV</td>
<td>PeM70</td>
</tr>
<tr>
<td>Pep6</td>
<td>GFFGPGGFGPVLDGWYGVDV</td>
<td>PeM71</td>
</tr>
<tr>
<td>Pep7</td>
<td>GFFGPGGFGPVLDGWYGVDV</td>
<td>PeM72</td>
</tr>
<tr>
<td>Meningococcal</td>
<td>TPAVYKUNNNTLVL</td>
<td>TPAVYKUNNNTLVL</td>
</tr>
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</table>

(a) Epitope mapping of mAbs specific for region 1 of pertactin.

(b) Amino acid sequence of minimum epitopes recognized by mAbs (underlined).

Mouse IgG subclasses were determined with a BioTek ELISA plate reader (EL312e).

Immunization and challenge of mice. B. pertussis isolates, all streptomycin-resistant, were grown on BG agar supplemented with streptomycin (30 µg ml⁻¹) at 35 °C for 3 d. Subsequently the bacteria were plated on BG agar plates without streptomycin. After 3 d, bacteria were harvested and resuspended in Verwey medium (Verwey et al., 1949) to a concentration of 5 x 10⁸ bacteria ml⁻¹. An aliquot of the final suspension was diluted and plated to determine the c.f.u. of the challenge inoculum. Groups of eight female BALB/c mice (RIVM or Harlan; OlaHsd) were used for immunization. For passive immunization, 250 µg of a purified mAb was injected intravenously into 4-week-old mice. Mice were infected 24 h later. For active immunization with (poly)peptides, 3-week-old mice were immunized subcutaneously on day 0, day 14 and day 28 with 0.5 ml PBS containing 20 µg Quil A (Spikoside; Isocote) and 50 µg tetanus-conjugated peptide or 15 µg Prn1. In case of vaccination with the Dutch whole cell vaccine (which contains diphtheria and tetanus toxoids, and inactivated polio, in addition to the pertussis component), mice were immunized twice on day 0 and day 14 with a 1/100 human dose in PBS with 0.325 % aluminium hydroxide (Alu-Gel-S suspension; Boehringer Ingelheim). When the mix of seven peptides was used, equal amounts of each peptide conjugate (i.e. 6.25 µg) were present in the vaccine. Mice were infected 14 d after the last immunization. For infection, mice were lightly anaesthetized with ether and a drop of 20 µl of the inoculum was placed on top of each nostril and allowed to be inhaled by the animal. Mice were infected with a total amount of 2 x 10⁸ B. pertussis cells. Three days after infection mice were sacrificed by intraperitoneal injection of an overdose of Nembutal (Sanofi; Sigma) for 10 s followed by four washings as described above. Bound antibodies were detected using HRP-conjugated anti-mouse total IgG (Cappel; Organon Technica). Mouse IgG subclasses were determined by using a mouse mAb isotyping kit (Isostrip; Boehringer). Extinctions (OD₄₉₂) was measured with a BioTek plate reader (EL312e).
vaccination was determined by ELISA as described above. Plates were coated with tetanus toxoid (2 μg ml⁻¹) or Prn1 (2 μg ml⁻¹). Extinctions were measured with a BioTek plate reader (EL312e) and titres were calculated with Kineticalc (KC4; BioLyse).

**Blocking ELISA.** Blocking ELISA was performed essentially as described by Berbers et al. (1993). Plates were coated with Prn1 (2 μg ml⁻¹) as described above. After washing and blocking, the wells were incubated with 100 μl of serial twofold dilutions of human serum diluted in 0.5% (w/v) BSA in PBST for 2 h at 22 °C. Human sera were from two sources. One serum was from a vaccinated child that was recently infected with pertussis. The serum sample was taken 47 d after the onset of the disease. The second serum preparation comprised a mix of sera from several recently infected pertussis patients who were fully vaccinated. After washing the plates, a pertactin-specific murine mAb, PeM7, PeM2, PeM5 or PeM6, was added and the plates were incubated for 2 h at 22 °C. After the plates were washed thoroughly, they were developed and read as described above.

**Statistical analyses.** P-values (two-sided) were calculated by using Student’s t test.

## RESULTS

### Characterization of mAbs against pertactin

Previously, we identified six different pertactin types in *B. pertussis* strains circulating in Europe, of which three (Prn1–3) are found in The Netherlands (Mooi et al., 1998). Variation between the pertactin types is mainly limited to region 1 and consists of deletions or insertions of the repeat unit GGxxP (Fig. 1) (Mooi et al., 1998, 1999; Mastrantonio et al., 1999). Here we investigated whether polymorphism in region 1 affected antibody binding, i.e. represented antigenic variation. To this purpose several mAbs were raised, using pertactin variants and MBP fusion proteins with region 1 derived from different pertactin variants.

Binding of the mAbs to the six *B. pertussis* pertactin types and the homologous proteins derived from the closely related species *B. bronchiseptica* and *B. parapertussis* was analysed by immunoblotting. To identify polymorphism in pertactin outside region 1, the pertactin genes of all isolates used for immunoblotting were sequenced completely. Compared to prn1, the *B. bronchiseptica* and *B. parapertussis* pertactin genes revealed mutations resulting in amino acid substitutions over the whole length of the encoded protein (not shown). However, within *B. pertussis* only Prn6 showed polymorphism outside region 1; compared to Prn1, Prn6 contained 4 aa substitutions at positions 102, 337, 532, 853 and a deletion of 3 aa in region 2 (Fig. 1). Of the 14 mAbs tested, seven cross-reacted with pertactin derived from *B. bronchiseptica* and *B. parapertussis* (Table 3) and two (PeM5 and PeM7) showed differential binding to *B. pertussis* pertactin (Table 3). PeM5 showed highest and lowest binding to Prn1 and Prn6, respectively. Intermediate binding was observed with Prn2–5. PeM7, although raised with Prn1, showed a much stronger reaction with Prn6 compared to Prn1. Weak binding was observed with Prn2–5 (Table 3). Since, with the exception of Prn6, the different pertactin proteins do not vary outside region 1, the differential binding of mAbs is due to recognition of an epitope encompassing region 1, or an epitope outside region 1 that is affected by changes in region 1 (both of which will be henceforth referred to as a region 1-dependent epitopes). Taken together, these results indicate that variation in region 1 affects antibody binding.

The ability of the mAbs to bind to region 1 was also studied with MBP fusion proteins harbouring region 1 derived from the pertactin variants Prn1–5. Binding was assessed by immunoblotting. Of the 14 mAbs tested, 7 (PeM3, PeM4, PeM68, PeM70, PeM71, PeM72 and F6E5) bound to the fusion proteins and not to MBP (Table 3). No quantitative differences were observed between these seven mAbs with respect to binding to the MBP fusion proteins. The mAbs that did not bind to the MBP fusion proteins may recognize an epitope located outside region 1. It is also possible that these mAbs recognize a conformational region 1-dependent epitope, which folds differently in an MBP fusion protein compared to pertactin. In fact this is probably the case with PeM5 and PeM7, which do not bind to the MBP fusion proteins, although the type specificity of these mAbs suggested that they bound to region 1 of pertactin. Thus 4 out of 10 mAbs raised against intact pertactin were directed against a region 1-dependent epitope, indicating that it is immunogenic in mice.

### Epitope mapping

To delineate the epitopes recognized by the mAbs described above, a set of eight overlapping peptides corresponding to region 1 (Pep1–7 and Pep10) was used (Table 2a). The seven mAbs (F6E5, PeM3, PeM4, PeM68, PeM70, PeM71 and PeM72) which bound to the fusion proteins also bound to the peptides (Table 2a). The mAb F6E5 was previously reported to bind a pertactin peptide of 128 aa encompassing the RGD and the GGxxP repeat (Charles et al., 1991). Here we show that F6E5 bound to peptides 6 and 7 only, defining the sequence GGFGPVLDGW as its epitope. Based on their differential binding to the peptides, the epitopes of the other mAbs were also delineated and are shown in Table 2b. The epitopes comprised of GGFGPGFGER and GGFGP were each recognized by two mAbs, respectively, PeM3 and PeM4, and PeM71 and PeM72. Epitopes for PeM68 and PeM70 were identified as GDAPAGGAVP and ATIRR, respectively. PeM5 and PeM7 did not bind to the peptides, which is consistent with the assumption that they recognize a conformational epitope.

### Antibodies against region 1 are induced in humans

We looked for the presence of region 1-specific antibodies in human sera using a blocking ELISA. In this assay the ability of human antibodies to compete with mAb PeM7, which binds to a region 1-dependent epitope (see above), for binding to immobilized Prn1 was
Table 3. Reactivity of mAbs with pertactin variants or MBP–pertactin fusion proteins containing region 1

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<tr>
<th>mAb</th>
<th>IG sub-type</th>
<th>Antigen used for immunization</th>
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*Identical results were obtained with MBP–Prn1 to MBP–Prn5.

![Fig. 2](image-url)  
**Fig. 2.** Sera from pertussis patients harbour antibodies directed against region 1. ELISA plates were coated with Prn1. Binding of mAbs PeM7 (■; directed against region 1) or PeM2 (▲; directed against an epitope present in all pertactin variants) was allowed after pre-incubation with twofold dilutions of human serum. Human serum was from one pertussis patient; similar results were obtained with a pool of sera from different pertussis patients.

assessed. The serum sample was from a child recently infected with *B. pertussis*. Similar results were obtained with pooled sera derived from several children (not shown). Binding of PeM7 to Prn1 was inhibited in a dose-dependent manner (Fig. 2), indicating the presence of human antibodies directed to the region 1-dependent epitope recognized by PeM7. Binding of PeM2, which recognizes an epitope present in all tested pertactin variants (Table 3), was not inhibited by human antibodies (Fig. 2). The mAb PeM5, which is similar to PeM7 in that it bound to a region 1-dependent epitope (see above), also competed with human antibodies (not shown). Interestingly, mAbs that recognized linear...
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epitopes in region 1 (e.g. PeM4 and F6E5) were not able to compete with human serum for binding to pertactin (not shown). These results indicate that, in its native state, region 1 elicits antibodies in humans.

Variation in region 1 affects the efficacy of the Dutch WCV in the mouse model

To determine if variation in pertactin affected vaccine efficacy, mice were immunized with the Dutch WCV and challenged with *B. pertussis* isolates expressing Prn1, Prn2 or Prn3, the predominant types found in most countries (Mooi *et al.*, 1998, 1999; Mastrantonio *et al.*, 1999). Control mice were immunized with PBS. To correct for the different colonizing abilities of isolates in mice, log(protect) values were calculated (see legend to Fig. 3). We found the Dutch WCV protected against isolates with Prn1 significantly better compared to those carrying non-vaccine-type pertactins (*P* = 0.0081) (Fig. 3). When the Prn2 and Prn3 isolates were analysed separately, the WCV was also found to be less effective; however, significance was only attained with the Prn2 group (*P* = 0.0143).

Region 1 harbours a protective epitope

To confirm that region 1 was able to induce a protective immune response, mice were immunized with a mixture of seven overlapping peptides derived from region 1 (Pep1–7; Table 2a). Mice in the control group were immunized with a meningococcal peptide (Table 2a) or with PBS. Immunization with Prn1 served as a positive control. Peptides were conjugated to tetanus toxoid. Analysis of the serum samples of immunized and infected mice revealed that high antibody titres to tetanus toxoid were observed in both groups immunized with the peptides (Fig. 4a), while only mice immunized with the pertactin peptides or with the intact pertactin had anti-pertactin titres (Fig. 4b). Immunization with pertactin peptides reduced colonization in the lungs and trachea 35- and 16-fold, respectively, compared to mice immunized with the meningococcal peptide (*P* = 0.0009.

**Fig. 4.** Protection against intranasal *B. pertussis* infection by immunization with peptides derived from pertactin region 1. Mice were immunized with PBS, a meningococcal peptide, a mixture of peptides derived from region 1 (Pep1–7; Table 2a) or with pertactin (Prn1). Peptides were conjugated to tetanus toxoid. After immunization, mice were intranasally infected with *B. pertussis* strain B213 expressing Prn1. The amount of bacteria in the lungs and trachea, and antibody titres were determined 3 d post-infection. (a) Anti-tetanus toxoid IgG titres; (b) anti-Prn1 IgG titres; (c) c.f.u. in the lungs; (d) c.f.u. in trachea. The thin line indicates the mean. *P* values are indicated. The experiment was performed three times and a representative result is shown.
Colonization in lungs and trachea (P < 0.0001) (Fig. 4). Immunization with intact pertactin resulted in a significant reduction of colonization in lungs and trachea (P < 0.0001). In conclusion, both in the trachea and lungs, peptides derived from region 1 were able to induce protective immunity.

The role of region 1 in conferring immunity was confirmed by passive immunization with mAb PeM4. PeM4, which binds to the GGFGPGGFGP sequence in region 1, was administered intravenously 24 h prior to challenge. Control mice were injected with PBS. Mice injected with PeM4 showed over 10-fold less colonization of the lungs by *B. pertussis*, compared to mice injected with PBS (P < 0.0001) (Fig. 5). In the trachea the reduction in colonization by PeM4 was less pronounced, in some experiments an approximately twofold drop in colonization was seen, whereas in others no significant drop in colonization was observed (not shown).

**Cross-immunity can be conferred by a linear epitope from region 1**

The sequence recognized by PeM4 was present in all six pertactin variants, suggesting that cross-immunity could be induced. To test this assumption, mice were immunized passively with PeM4 and subsequently challenged with a *B. pertussis* isolate expressing Prn2 or Prn3 (Fig. 5). Mice in the control group were given PBS. As was observed with the isolate expressing Prn1, PeM4 significantly reduced colonization (P ≤ 0.0015) in the lungs of the isolates expressing Prn2 or Prn3. Importantly, no significant differences were observed between the Prn groups with respect to the degree of immunity conferred by PeM4 in the lungs, indicating cross-immunity. As with Prn1, variable results were obtained in the trachea (not shown). The mixture of seven overlapping peptides derived from region 1 was also found to confer cross-immunity (results not shown). The results obtained with passive and active immunization indicate that region 1 contains a cross-protecting linear epitope that protects equally against isolates expressing distinct pertactin variants.

**DISCUSSION**

Studies in animals and humans have indicated that pertactin plays an important role in immunity to pertussis (Novotny *et al.*, 1991; Cherry *et al.*, 1998; Shahin *et al.*, 1990; Storsaeter *et al.*, 1998). Especially relevant are the results of two independent field trials, which revealed a correlation between pertactin antibodies and clinical protection (Cherry & Olin 1999; Cherry *et al.*, 1998; Storsaeter *et al.*, 1998). Our previous studies have provided indirect epidemiological evidence that variation in pertactin is immunologically relevant and may be linked to the re-emergence of pertussis in vaccinated populations (de Melker *et al.*, 1997; Mooi *et al.*, 1998, 1999; Mastrantonio *et al.*, 1999). For example, isolates with vaccine-type pertactin were found in lower frequencies in vaccinated children compared to non-vaccinated children, suggesting that variation in pertactin affects vaccine efficacy (Mooi *et al.*, 1998). Further, temporal trends in the frequencies of the pertactin and pertussis toxin variants indicated a divergence between vaccine strains and clinical isolates (Mooi *et al.*, 1998, 1999). Here we extend our studies on the immunological relevance of the variable region 1 of pertactin.

We observed that a significant number of the mice mAbs were directed against a region 1-dependent epitope after immunization with pertactin, indicating that this region is immunogenic. Moreover, also in serum from humans recently infected with *B. pertussis*, we detected antibodies against region 1. Interestingly, mAbs that recognized a conformational region 1-dependent epitope (Fig. 2), but not those that bound to a linear epitope (not shown), were found to compete with the human antibodies. The reason for this difference is not clear, but it is conceivable that antibodies directed against conformational epitopes bind with higher avidity than antibodies against linear epitopes. It is also possible that infection or vaccination mainly results in the induction of antibodies directed against conformational epitopes of pertactin. In this context it is significant that mAbs directed against conformational region 1-dependent epitopes were type-specific, whereas those that recognized a linear region 1 epitope bound to all *B. pertussis* pertactin variants. Thus, to prevent cross-immunity, the bacteria may have evolved mechanisms to direct the immune response towards conformational epitopes. These data show that region 1 is exposed and able to elicit antibodies both in mice and humans. This is consistent with functional studies, which have implicated this region in adherence (Everest *et al.*, 1996) and
crystallographic data which indicate that region 1 is exposed as a loop (Emsley et al., 1996). The fact that the binding of two mAbs to Prn was affected by variation in region 1, indicated that polymorphism in region 1 represented antigenic variation.

A number of linear epitopes recognized by mAbs were mapped within region 1. One of these mAbs (F6E5) was previously shown to inhibit invasion of HeLa cells by B. pertussis and its binding site was mapped to a region of 128 aa residues overlapping the RGD site (Leininger et al., 1992). We defined the F6E5 epitope more precisely with synthetic peptides and located it in the GGxXP repeat region, 23 aa removed from the RGD motif. The inhibitory effect of this mAb on invasion (Leininger et al., 1991) underlines the importance of antigenic variation in region 1 for bacterial fitness.

Immunization of mice with peptides derived from region 1 significantly reduced colonization of both trachea and lungs, demonstrating the importance of this region for immunity. Further evidence for the importance of region 1 in protective immunity was revealed by the observation that an mAb (PeM4) directed against a linear epitope in region 1, comprising two GGFGP repeats, was able to confer passive protection in the lungs. The efficacy of this mAb was much lower in the trachea, however. Protection probably requires transudation of antibodies from the blood to the site of infection (i.e. the mucosal surfaces). It is conceivable that the mAb used was inefficiently transudated to the trachea. It is not clear on what mechanism the protection by the mAb is based. Effector functions, such as opsonization and complement activation, may be involved. The isotype of antibody used, IgG1, shows relatively weak effector functions in mice (Ravetch & Clynès, 1998). Alternatively, the mAbs may block the function of pertactin or hinder colonization by agglutinating bacteria. It should be noted that a function for pertactin in the mouse model has yet to be established: a pertactin-defective mutant was not affected in its ability to colonize the trachea or lungs of mice (Roberts et al., 1991).

A second protective epitope has been located in the PQP repeat region, designated region 2, which is located in the C-terminal part of pertactin (Fig. 1) (Charles et al., 1991). Like the GGxXP repeat, region 2 was found to be immunogenic in mice (Charles et al., 1991). However, the PQP region is less variable than the GGxXP region and of the hundreds of isolates analysed, only two polymorphisms were observed in the PQP region (Fig. 1). To date, this minor variant, represented by Prn6, was found in two isolates only. Thus the vast majority of isolates analysed do not show variation in the PQP region, suggesting that it plays a less important role in escaping immunity compared to region 1.

Studies using the mouse model indicated that the Dutch WCV was less effective against Prn2 and Prn3 isolates compared to those expressing Prn1 (Fig. 3). Prn1 is produced by the vaccine strains and has been replaced by Prn2 and Prn3 in The Netherlands. Prn2 and Prn3 have also been found to dominate in other countries where the WCV contains Prn1 (Mooi et al., 1999, van Loo et al., 1999). It should be noted that we used clinical isolates which may differ in other properties relevant for immunity than pertactin. However, sequencing of 10 surface-associated proteins revealed that variation was mainly restricted to pertactin (unpublished data). The results obtained in the mouse model are consistent with data obtained from human populations, which indicate that Prn2 and Prn3 are less affected by the current WCVs compared to Prn1. The relevance of the mouse as a model for studying pertussis immunity in human models is further substanti- ated by two independent studies which showed that the results obtained with the intranasal mouse model are congruent with results from humans trials (Mills et al., 1998; Guiso et al., 1999). Boursaux-Eude et al. (1999) used the mouse model to compare the efficacy of acellular vaccines against isolates producing different pertactin and pertussis toxin variants. They concluded that a tri-component vaccine was highly effective in promoting lung clearance of all isolates expressing different pertussis toxin and pertactin variants. The data obtained with different isolates were not compared on a statistical basis, however. Furthermore, in the study of Boursaux-Eude et al. (1999) a much higher vaccine dose was used compared to our study (1/4 and 1/1000 human dose, respectively). In this context it should be noted that we found no significant differences in protection against isolates with different Prn variants when mice were immunized with a relatively high dose of Prn1 (1/10 human dose) (not shown). Nevertheless, the study of Boursaux-Eude et al. (1999) may indicate that acellular vaccines are less affected by antigenic variation than the whole-cell vaccine tested by us.

An mAb recognizing a linear epitope found in all Prn variants was equally effective against strains with Prn1, Prn2 and Prn3 when administered intravenously to mice. Furthermore, a similar observation was made when mice were vaccinated with a mixture of peptides harbouring sequences found in all Prn variants. We hypothesize that linear epitopes derived from region 1 may confer cross-immunity, while conformational epitopes do not. Infection and vaccination may induce mainly antibodies against conformational epitopes.

From the data described in this paper we can conclude that the polymorphic region 1 of pertactin is recognized by the immune system of both mice and humans, that variation in this region affects antibody binding and that this region harbours a protective epitope. Mice experiments suggested that variation in region 1 affects the efficacy of the Dutch WCV. Our results support the assumption that the expansion of isolates which carry non-vaccine type pertactin molecules has contributed to the re-emergence of pertussis in vaccinated populations (Mooi et al., 1998). An important, unresolved, question is whether, and to what extent, variation in pertactin affects the efficacy of acellular vaccines. In addition to pertactin, acellular vaccines contain filamentous haemagglutinin, fimbriae and pertussis toxin, which are also involved in protection against pertussis. Possibly,
the presence of high concentrations of multiple antigens may minimize the effect of variation in pertactin.

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