Differences of circulating *Bordetella pertussis* population in Argentina from the strain used in vaccine production

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**Abstract**

In Argentina, as in other countries, the number of pertussis cases has been increasing, even in highly vaccinated zones. Many reports suggest that the decline of vaccine efficacy due to antigenic shifts in the circulating *Bordetella pertussis* might be among the factors that contribute to pertussis re-emergence in different parts of the world. To evaluate the incidence of this factor in Argentina, we decided to characterize the circulating bacteria of an important demographic area of this country in comparison with the strain used for vaccine production. From 1997 to 2003 we collected nasopharyngeal samples from pediatric patients with signs of *Bordetella* infection hospitalized in the metropolitan area of Buenos Aires and La Plata, Argentina. From these samples we identified 28 *B. pertussis*, which were characterized by biochemical techniques, PCR, DNA fingerprint, prn and ptx genes sequencing, and lipopolysaccharides (LPS) pattern. BOX-PCR from *B. pertussis* isolates yielded one cluster containing 13 isolates and some smaller ones, being all fingerprints different from the vaccine strain. Differences between Argentinean circulating bacteria and the vaccine strain were also observed for the Prn and Ptx variants as well as for the LPS pattern. Moreover, this last pattern seemed to change over the years. In addition, we identified two *B. bronchiseptica*. The presence of this *Bordetella* species together with the observed differences between circulating *B. pertussis* and the strain used in vaccine production should be considered for the development of an improved vaccine.
1. Introduction

The genus *Bordetella* currently contains eight species, of which the three referred to as the classical *Bordetella*, are closely related respiratory tract pathogens [1]. These three species, *B. pertussis*, *B. parapertussis*, and *B. bronchiseptica*, can be viewed as subspecies [2] and express similar virulence factors regulated by the BvgAS two-component system [3]. *B. pertussis* and *B. parapertussis* are human pathogens that cause whooping cough or pertussis in humans, a respiratory disease that is especially severe in young children. *B. parapertussis* has also been isolated from sheep but available data suggest that human and sheep strains are distinct and that transmission from sheep to humans does not occur [4]. *B. bronchiseptica* is an animal pathogen [5] but can also infect humans [6], [7] and [8].

From these three pathogens, *B. pertussis* causes the most severe disease in humans and because of that, attempts to develop vaccines for both treatment and prevention were made soon after *B. pertussis* was first isolated. The initial vaccines consisted of killed whole *B. pertussis* bacteria with which routine immunizations of children were started in the mid-1940s. In some countries this continues up to now and in some others acellular vaccines replaced whole cell vaccines. Although the introduction of both types of vaccines significantly reduced the incidence of pertussis, the disease is still endemic worldwide. Moreover, in some countries with highly vaccinated populations like The Netherlands [9], the United States [10], Canada [11], Sweden [12], and Australia [13] a re-emergence of pertussis was recently reported. Several explanations have been suggested for the re-emergence of pertussis including changes in vaccine potency, decreased vaccine coverage, waning immunity and adaptation of the *B. pertussis* strains to vaccination. The latter hypothesis is mainly based on the finding of antigenic divergence between clinical isolates and the strains used in vaccine production. Genetic polymorphisms were mainly described for two of the main *B. pertussis* protective antigens, namely, pertussis toxin (Ptx) and pertactin (Prn) [14], [15], [16], [17], [18] and [19]. In particular, four Ptx S1 subunit variants (S1A-D) and 10 Prn variants (Prn1-9 and Prn11) were identified in the bacterial circulating populations [16] and [19]. These polymorphisms have been attributed to the ability of the bacterium to resist induced immunity, and cause an increase in the disease incidence [14] and [15].

In Argentina, the Public Health Ministry also registered an increase in pertussis incidence, even before improvements on diagnostic methodology were introduced. In this country whole cell pertussis vaccine was introduced 50 years ago and is still the only kind of pertussis vaccine being used. Up to now there are no data about the local circulating *Bordetella* strains in comparison with the strain used in vaccine production. Hence, it is of epidemiological interest to see whether the allelic frequencies in Argentinean *Bordetella* circulating strains are similar or not to the reported in the northern hemisphere. Therefore, we focused the present study on molecular characterization of *Bordetella* isolates obtained in a region of Argentina during the past 6 years. To our knowledge this is the first epidemiological molecular study on *Bordetella* performed in Latin America.
2. Methods

2.1. Patients, samples and bacterial growth conditions

During the period 1997–2003 we collected nasopharyngeal samples from 400 hospitalized patients from Buenos Aires with signs of *Bordetella* infection. From these samples we searched for the presence of *Bordetella* spp. by specific PCR and culture. *Bordetella* was isolated on Bordet Gengou agar (Difco) supplemented with 15% (v/v) defibrinated fresh sheep blood (BGA medium) at 36 °C for 72 h.

To identify the *Bordetella* species causing the respiratory infection we performed discriminative PCR assays [20] and [21], biochemical typification using API 20 NE system (bioMerieux, Marci l’Etoile, France), and agglutination test with antiserum for *B. pertussis* and *B. parapertussis* (Murex Diagnostic, Dartford England) on the clinical *Bordetella* spp. isolates.

All isolates were stored at −80 °C in 0.9% (w/v) NaCl containing 20% (v/v) glycerol.

We also grew *B. pertussis* strain Tohama, and *B. bronchiseptica* strain CIP 9.73 (Collection de l’Institut Pasteur [CIP] designation) on BGA medium at 36 °C for 48–72 h. In the experiments described below we used the Pasteur–Merieux–Connaught vaccine strain as a control since it is the vaccine strain used in Argentina.

2.2. PCR for detection of *B. pertussis*, *B. parapertussis* and *B. bronchiseptica*

Using the Fla1/Fla2 primers designed by us [20] we could detect the three species of *Bordetella*: *B. pertussis*, *B. parapertussis*, and *B. bronchiseptica*, and distinguish *B. pertussis* (size of amplified product 195 bp) from the other two pathogens (size of the amplicon 164 bp). As the sensitivity of this PCR is low for *B. pertussis* we also employed the oligonucleotide primers derived from the ptx promoter (PT1/PT2) (size of the amplicon, 191 bp) designed by Grimpel et al. [21]. To discriminate *B. bronchiseptica* from *B. parapertussis* we used the primers Fla2/Fla3 [20]. All the oligonucleotides were provided by DNAGency Inc. (Malvern, USA). The PCR assays were performed as described earlier [19]. Positive PCR products were detected by ethidium bromide staining of 2% agarose gels after electrophoresis.

2.3. DNA fingerprinting

We performed total DNA amplification fingerprints using different primers BOXC1, BOXAR1, ERIC, MBOREP1, REP1 and NGREP2, whose sequences were reported by Versalovic et al. [22].

The deoxyoligonucleotide primers were synthesized by DNAGency (Malvern, PA, USA). We carried out PCR reactions in 25 μl containing: 50 mM Tris, pH 8.3; 500 μg/ml BSA; 3 mM MgCl2; 200 μM dNTPs; 1 U Taq polymerase (Promega Corp.); 10 μM primer, and 2 μl of template DNA, previously obtained by phenol extraction. The amplifications were carried out in capillary tubes in an Idaho 1605 Air Thermo Cycler (ATC, Idaho Technology). The cycling conditions were as follows: 94 °C for 2 min, 35 cycles at 94 °C for 10 s, at 52 °C for 60 s, and at 65 °C for 8 min with a final cycle of 65 °C for 16 min. After the reaction, 10 μl of
the PCR products were separated in 1.5% agarose gels containing 0.5–1 μg/ml ethidium bromide, and photographed by using a digital camera Kodak DC290.

Computer-assisted analysis of the BOXAR1 fingerprint patterns was carried out using GelCompar software (Windows NT version 4.1, Applied Maths, Kortrijk, Belgium).

2.4. Sequence analysis of the genes for pertactin and pertussis toxin S1 subunit

We performed DNA sequencing of relevant regions of the prn and ptxS1 genes on PCR fragments as described [16]. We used the primer pair AF/AR (5′-CAATGTCACGGTCCAA-3′/5′-GCAAGGTGATCGACAGGG-3′) for sequencing of prn region 1, and for sequencing of prn region 2 the primer pair BF/BR (5′-AGCTGGGGGTCAAGGT-3′)/5′-CCGGATTCAGGGCAACTC-3′). For pertussis toxin, we used the primers S1-F2 (5′-CCCCCTGCGATGATGATTCA-3′) and S1-R2 (5′-AGGAGTTTCTTGCCTG-3′) to amplify a 930 bp product, which contains the complete ptxS1 gene. We also used the S1-F2/S1-R2 for sequencing [18]. We confirmed mutations by sequencing using two additional primers: S1-MF/S1-MR (5′-AACAATGCCCGGGCCGATCCTCAGGCTGGCGGC-3′).

2.5. Immunoblots

Cells corresponding to 2 × 10^10 colony-forming units of the Bordetella isolates were treated with Laemmli sample buffer and the extracts run on 12.5% (w/v) polyacrylamide SDS-gels. After electrophoresis, we transferred the proteins from the polyacrylamide to a polyvinylidenephosphate membrane (Immobilon P, Millipore) and incubated it with serum obtained from vaccinated patients, or with a 1:5000 dilution of mice polyclonal immune sera directed against AC-Hly (purified and subsequently inoculated into BALB/c mice as previously described [23]) or Ptx (kindly provided by Dr. Nicole Guiso, Laboratoire des Bordetelles, Institut Pasteur, Paris). In all the cases, we used alkaline-phosphatase-labeled sheep anti-mouse immunoglobulins to detect the presence of immune complexes.

2.6. Lipopolysaccharide extraction and sodium dodecylsulfate-polyacrylamide gel electrophoresis (SDS-PAGE)

Cells grown at 36 °C in BGA medium were resuspended in PBS buffer, centrifuged (10,000 × g, 15 min, 4 °C) and washed twice in distilled water. After adjustment of bacterial concentration, we extracted the lipopolysaccharides (LPS) by affinity chromatography as described [24]. We solubilized the isolated LPS by heating at 100 °C for 5 min in Laemmli sample buffer. We then applied the LPS suspensions to SDS-gels.

We performed electrophoresis at room temperature and constant voltage, and then visualized the LPS by the BioRad silver-staining technique.

2.7. Peripheral blood polymorphonuclear leukocyte (PMN)

We isolated peripheral blood PMN from heparinized venous blood of healthy human volunteers (age range: 25–45 years old) using Ficoll-Histopaque (Sigma–Aldrich, St. Louis, MO) gradient centrifugation. We harvested polymorphic leukocytes and removed the remaining erythrocytes by hypotonic lyses. Cell viability was >99% as determined by trypan blue exclusion. Before functional assays, we washed twice the PMN with RPMI 1640 medium supplemented with
10% heat-inactivated fetal calf serum (FCS); the PMN were then resuspended, and used immediately. All experiments described in this study were conducted with freshly isolated PMN lacking FcγRI expression, as monitored by FACS analyses with FITC-conjugated anti-FcγRI mAb 22 [25].

2.8. Phagocytosis

We evaluated phagocytosis of B. pertussis and B. bronchiseptica as previously described [26]. To label bacteria for this study, we introduced into B. bronchiseptica CIP 9.73 and B. pertussis Tohama strains by conjugation the plasmid pGB5P1 that codes for Km resistance and for the green fluorescence protein. This plasmid was kindly provided by Dr. Allison Weiss (Department of Molecular Genetics, Biochemistry and Microbiology, University of Cincinnati). We grew GFP-expressing B. pertussis and GFP-expressing B. bronchiseptica overnight on BG agar plates and resuspended them in RPMI 1640 medium containing 10% FCS. Bacteria were incubated 30 min at 37 °C with human sera (2.5 or 5%) obtained from a 6 years old child either before or 30 days after vaccination with whole cell pertussis vaccine After washing, we incubated the opsonized bacteria with phagocytic cells in a 70:1 ratio for 45 min at 4 °C to allow binding of bacteria to PMN. After extensive washing to remove nonattached bacteria, we further incubated the cells in two aliquots and further incubated them for 30 min, either at 4 °C or 37 °C. Next, we detected remaining cell surface-bound opsonized bacteria by incubation (30 min at 4 °C) with Phycocerythrin (PE)-conjugated goat F(ab')2 of anti-human IgG (from Southern Biotechnology Associates, Birmingham, AL). After washing, we analyzed the samples by flow cytometry. Five thousand cells were analyzed per sample. Green and red fluorescence intensities of cells maintained at 4 °C throughout served as control for bacterial binding (i.e. 0% phagocytosis). The decrease in red fluorescence of green positive cells after incubation at 37 °C reflects bacterial phagocytosis, as confirmed microscopically [26]. We calculated phagocytosis rates from the drop in mean red fluorescence intensity of green-positive cells and expressed them in arbitrary units (AU), as described [27].

3. Results

3.1. Description of samples and isolates

For this study we collected nasopharyngeal samples over a period of 6 years from 400 hospitalized pediatric patients at the metropolitan area of Buenos Aires and La Plata (hereafter referred to as Buenos Aires). This important demographic region where an increase of pertussis cases was registered from 1999, concentrates 30% (11,500,000 people) of the total Argentinean population (37,000,000 people). The patients had signs of Bordetella infection with dry and emetic coughing for approximately 14 days. Apnea and cyanosis were frequent in infants younger than 3 months of age and sometimes necessitated intensive care in hospital. The age distribution of the patients was as follows: 90% younger than 1-year-old, 7% between 19 months and 3 years old, and only one patient 12-year-old. The vaccination statuses of the majority of the patients were completed according to their age. Only for nine hosts the vaccination status was unknown. The vaccine used during 1997–2003 was the whole cell Pasteur–Merieux–Connaught vaccine and the vaccination schedule consisted in five doses given at 2, 4, 6, and 18 months of age, and finally reinforcement at the age of 6 years.

From the collected samples, 25% were shown to be positive for specific Bordetella PCR [20] and [21] (98% positive for B. pertussis and 2% for B. bronchiseptica). However, from the 100 PCR-positive nasopharyngeal samples, only 30 Bordetella isolates could be cultured on BGA plates. PCR assays using the flagellin promoter
sequence as a target DNA region [20] showed that 28 out of the 30 isolates were 
*B. pertussis* (isolated from patients with ages ranging from 1 month to 3 years 
old) and the other two were *B. bronchiseptica* (isolated from vaccinated patients 
of 19 months and 12 years old). To confirm these PCR-based typifications, we 
analyzed oxidase, urease and motility by the API 20 NE system, and agglutination 
with antiserum for *B. pertussis* and *B. parapertussis*. The results obtained were 
coincident with those obtained with PCR assays.

### 3.2. Virulence state

We investigated the hemolytic activity of each clinical bacterial isolate in BGA and 
observed that all isolates were hemolytic and thus most likely virulent. Moreover, 
we evaluated the expression of virulence determinants by Western blot analysis 
using mouse polyclonal immune sera directed against AC-Hly and Ptx (data not 
shown). The 28 *B. pertussis* isolates expressed both toxins and *B. bronchiseptica* 
isolates only expressed AC-Hly. The absence of Ptx expression by *B. bronchiseptica* 
is consistent with previous studies [28].

### 3.3. Lipopolysaccharide variation

Phenotypic changes of surface bacterial components are commonly observed as a 
result of bacterial interaction with the host [29], [30], [31] and [32]. Among 
these surface components are the lipopolysaccharides, the major structural 
component of the outer membrane of Gram-negative bacteria and a highly 
immunogenic molecule. We here compared SDS-PAGE profiles of the LPS from 
clinical isolates and from the vaccine strain (Fig. 1). The LPS were extracted from 
cells grown in the same conditions. Vaccine strain LPS showed the typical two-
band pattern, LPS A and LPS B. In turn, although the LPS from clinical isolates 
also showed a pattern with a preponderance of LPS A, some of them seemed not 
to have LPS B. This shift from the classical two-band pattern (LPS A and LPS B) to 
a single band (LPS A) occurred in the more recent isolates. In order to exclude 
possible artifacts due to growth and/or LPS extraction conditions we repeated the 
whole procedure three times for all samples starting from the glycerol stock, and 
the same results were consistently observed. We also analyzed the possibility that 
the absence of LPSB band could have been caused by a difference in LPS sample 
concentration loaded in the gel. However, similar LPS patterns in the SDS-PAGE 
were observed when four times higher concentration of LPS was loaded in the gel 
(not shown).

![Fig. 1. Electrophoretic LPS profiles from different B. pertussis clinical isolates. LPS samples were separated in SDS-PAGE (15% [w/v]) and visualized via silver staining. The figure is a composite of several gels showing the LPS band patterns of each isolate. As internal standard Tohama and vaccine strain LPS were included in each gel. LPS samples in the gel correspond to the bacterial isolates obtained at the years indicated at the top of the figure. Tohama and B. bronchiseptica CIP 9.73 collection strains, and Pasteur–Merieux–Connaught vaccine strain are also included in lanes 1, 31 and 30, respectively. This LPS analysis was repeated three times with the same result.](image-url)
3.4. Sequencing of the structural genes encoding the S1 subunit of pertussis toxin and the repeated region of pertactin

Polymorphism in the genes for pertactin (prn) and the PtxS1 subunit (ptxS1) was studied in the 28 isolates of *B. pertussis* by DNA sequencing [16]. Our results on prn sequencing showed *B. pertussis* harboring prn2 variant to be prevalent in Buenos Aires (Table 1). In fact, of the 28 isolates studied 26 (93%) contained the prn2 allele and 2 (7%) the prn1/7 allele. Studies on PtxS1 subunit polymorphism showed that all *B. pertussis* isolated by us from 1997 to 2003 contained the ptxS1A allele. By contrast, the vaccine strain used in Argentina contains prn1 and ptxS1D alleles. Thus with respect to these two genes the vaccine strain was different than the prevalent circulating bacteria.

Table 1.

Frequencies of ptxS1 and prn alleles in different periods in Argentina

<table>
<thead>
<tr>
<th>Period</th>
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<th>No. of <em>B. pertussis</em> clinical isolates with allele (%)</th>
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<tr>
<td></td>
<td></td>
<td>ptxS1</td>
</tr>
<tr>
<td>1997–1998</td>
<td>3</td>
<td>0 (0)</td>
</tr>
<tr>
<td>2000–2001</td>
<td>14</td>
<td>14 (100)</td>
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<tr>
<td>2002–2003</td>
<td>11</td>
<td>11 (100)</td>
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Tohama collection strain contained S1B and prn1; Pasteur–Merieux–Connaught vaccine strain contained S1D and prn1.

3.5. DNA fingerprinting

The degree of genotypic similarity among the *Bordetella* clinical isolates was examined by a PCR-fingerprinting method. We tested the reproducibility of fingerprints with primers BOXC1, BOXAR1, ERIC, MBOREP1, REP1 and NGREP2 in PCR assays [22] using purified genomic DNA from four different isolates as templates in three independent experiments. With BOXAR1 primer we obtained the same results in each of the three independent reactions and in addition, the four fingerprint patterns could be easily distinguished from each other thus giving the best discriminatory results (data not shown). Based on these results the BOXAR1–PCR was selected for genotypic typing of the other clinical isolates.

Purified genomic DNA of the clinical isolates, the reference strains, or the vaccine strain, were used as templates in the PCR assays. PCR-amplified fragments were separated on an agarose gel (Fig. 2A) and the fingerprint patterns scanned into a computer. We used the GelCompar software package to derive phylogenetic relationships. The dendrogram derived is shown in Fig. 2B. BOXAR1 PCR of genomic DNA from all *B. pertussis* isolates yielded one major fingerprint profile and a small number of unique profiles. Clear differences in the band patterns between the vaccine strain and the isolates can be observed in the agarose gel. Small differences among the profiles of *B. pertussis* isolates can also be seen.
Fig. 2. Genotypic characterization of *Bordetella* clinical isolates. (A) BOXAR1-PCR fingerprint analysis of genomic DNA obtained from *B. pertussis* and *B. bronchiseptica* clinical isolates. The figure is a composite of several gels showing the characteristic band patterns of each isolate. As internal standard Tohama and vaccine strain DNA were included in each gel. BOXA-PCR products derived from different templates were separated on a 1.5% agarose gel and stained with ethidium bromide. The year of isolation and the number of the isolate are indicated at the top of the figure. *B. pertussis* Tohama and *B. bronchiseptica* CIP 9.73 collection strains and Pasteur–Merieux–Connaught vaccinal strain are also included in lanes 1, 31 and 30, respectively. Lane labeled M contain λ HindIII size marker. (B) Dendrogram of *B. pertussis* clinical isolates BOXA-PCR fingerprint. The year of isolation and the number of the isolate are indicated at the right of the figure. The scale above the dendrogram describes percent relatedness of each branch of the dendrogram as determined by GelCompare software. This fingerprint analysis was repeated at least three times with the same result.
3.6. Cross immune protection between *B. pertussis* and *B. bronchiseptica*

In order to investigate the cross-immune protection between *B. pertussis* and *B. bronchiseptica* we tested the reactivity of sera from a donor vaccinated with whole cell vaccine exhibiting high titers of anti-*B. pertussis* antibodies, as determined by ELISA, against whole cell proteins of both strains by western blot. As expected, this serum recognized polypeptides of both *B. pertussis* and *B. bronchiseptica* (not shown). However, the biological relevance of this cross reactivity has to be established. Opsonophagocytosis was found to be a key mechanism in preventing *Bordetellae* infection [26] and [27]. Determining whether anti-*B. pertussis* antibodies can act as opsonins for *B. bronchiseptica* might give a good indication of the biological activity of these antibodies in relation to *B. bronchiseptica*. Opsonophagocytosis of *B. pertussis* and *B. bronchiseptica* induced by serum obtained before (pre-immune) and after (post-immune) vaccination with pertussis whole cell vaccine was compared. Post-immune serum-induced efficient attachment and internalization of both *Bordetella* strains, as reflected by the high level of both the green and red mean fluorescence of PMN, as well as the drop of mean red fluorescence of PMN after incubation at 37 °C (Fig. 3A). A positive correlation between serum concentration used for opsonization and bacterial phagocytosis was observed both for *B. pertussis* and *B. bronchiseptica* (Fig. 3B). Phagocytosis antibody dependency was further confirmed by the lack of opsonic activity by serum obtained prior vaccination (exhibiting no detectable antibody titers against *B. pertussis* as determined by ELISA). This was demonstrated by the low green and red fluorescence intensities of PMN incubated with either *B. pertussis* or *B. bronchiseptica* opsonized with pre-immune serum even at highest concentration (5%) (Fig. 3).
Fig. 3. Effect of antibodies induced by pertussis vaccination on *B. pertussis* and *B. bronchiseptica* phagocytosis by PMN. (A) PMN were incubated with either *B. pertussis* or *B. bronchiseptica* opsonized with serum (5%) from a donor either before (pre-immune) or after (post-immune) pertussis vaccination at 4 °C during 30 min. Cells were split over two aliquots, and subsequently incubated for 30 min at either 4 °C or 37 °C. Remaining surface-bound opsonized bacteria were detected by addition of PE-conjugated goat F(ab')2 fragments of anti-human IgG antibodies. (B) PMN phagocytosis of *B. pertussis* (dark bars) and *B. bronchiseptica* (white bars) opsonized with either two different concentrations (2.5% and 5%) of post-immune serum or 5% of pre-immune serum from a pertussis vaccinated donor. Data represent the mean ± S.D. of four experiments with PMN from different donors. AU: arbitrary units.
4. Discussion

Many reports have documented the increase of *B. pertussis* incidence even in highly vaccinated populations [9], [11] and [13]. The causes for this increase are still under investigation. However, it has been proposed that, in addition to the waning immunity in adolescents and adults [33] and [34], who could then be reinfected and serve as source of pathogen to young infants, *B. pertussis* may have evolved in a way that is less sensitive to vaccine-induced immunity [35] and [36]. An increasing body of evidence collected in different countries supports this hypothesis [14], [15], [16], [17], [18] and [19]. Here we report the first study on *B. pertussis* molecular epidemiology in Buenos Aires, showing a clear divergence between the strain used in vaccine production and clinical isolates.

Samples from pediatric patients with pertussis symptomatology were collected during 6 years. Twenty-five percent of the total number of samples proved to be *Bordetella* by PCR (98% positive for *B. pertussis* and 2% positive for *B. bronchiseptica*) but only 7.5% (30 isolates) were culture positive. The difference between the initial detection of *Bordetella* positives by PCR and the number of strains that could actually be cultured may be due not only to the difference in the sensitivity of the techniques used but also to the decrease in bacterial viability that may occur after sampling and during transport.

From the 30 *Bordetella* isolates, 28 were identified as *B. pertussis* and two as *B. bronchiseptica*. No differences in the virulent state of the isolates could be detected. However, an interesting difference was observed in the bacterial LPS pattern in the *B. pertussis* isolates. A shift from the classical pattern of two bands (LPS A and LPS B) to only one band (LPS A) was observed (Fig. 1) in some of the isolates. This enrichment in LPS A with respect to LPS B is consistent with the hypothesis that the membrane-distal region of the LPS molecule (i.e. the LPS A), which is in intimate contact with the bacterial environment and might have an important role in host-bacterial interactions, would be expressed at a higher level within the host [37]. Harvill et al. [30] demonstrated that Δwlb mutants of *B. pertussis* which do not produce the LPS A form, were defective in colonization of the respiratory tracts of BALB/c mice. Furthermore, it was reported that *B. pertussis* LPS mutants lacking even one of the sugars in the terminal trisaccharide were bound and aggregated by surfactant protein A (SP-A). Phagocytosis by human monocytes was enhanced on LPS mutants that were able to bind SP-A, but not on wild-type bacteria [38]. The authors concluded that the LPS of wild-type *B. pertussis* shields the bacteria from SP-A-mediated clearance, possibly by sterically limiting access to the lipid A region. Taken together, these results seem to indicate that LPS A is an essential form of LPS for *B. pertussis* infection. If we assume that a selection pressure towards more infective genotypes operates within the bacterial population then the enrichment of LPS A-expressing bacteria can be expected. Thus, it would be of interest to study whether vaccination has any role in this selection.

We also compared isolates and the vaccine strain following the recommended approach of sequencing the genes that code for two proteins that confer protective immunity in animals and humans, pertactin (Prn) and pertussis toxin (Ptx). In addition, we used a more discriminative technique, the PCR fingerprinting, to establish the relatedness among the isolates and in relation to the vaccine strain.

Several reports from Europe and USA [9] and [10] documented in that countries the prevalence of isolates expressing Prn and Ptx variants not present in vaccine. Four PtxS1 types (PtxS1A-D) have been described, from which PtxS1A was most
frequently found as a prevalent variant. By contrast, the acellular and cellular vaccines currently in use contain PtxS1B and PtxS1D but not PtxS1A [39]. Regarding pertactin, many variants have been described, of which Prn2 was mostly observed (vaccines contain Prn1). The alleles prn1 and ptxS1B or D have been termed by Cassiday et al. [40] old alleles and the strains possessing both alleles, old strains. Strains possessing prn2 or 3 and ptxS1A alleles have been assigned to the group of new strains, and those that possess one new and one old prn or ptxS1 to the group of transitional strains. In The Netherlands, Finland, Italy, France, and United States it was observed that the strains possessing both an old prn and an old ptxS1 allele have origins prior to the 1970s. In contrast new strains having prn2, prn3 and ptxS1A were first detected from the mid-1980s till to now in all the countries mentioned above [10], [17] and [41].

In Buenos Aires the majority of circulating bacteria possesses ptxS1A and prn2 allelic variants. These allelic frequencies found in Argentina almost exactly match those found in distant countries at the Northern hemisphere, arguing in favor of vaccine-driven selection pressure. Since vaccine was introduced in Argentina in the 1950s, our observations probably reflect the equilibrium allelic frequencies reached at present, after more than 40 years of selection; thereby we did not observe changes between 1997 and 2003.

The divergences between the vaccine strain (PtxS1D and Prn1) and current isolates with the prevalence of the PtxS1A and Prn2 variants seem to be critical in protection against pertussis [14], [15] and [42]. The results obtained in animal models first by King et al. [15] and recently confirmed independently by Gzyl et al. [14], showed that whole cell vaccines were less effective against strains that carried non-vaccine-type Prn variants compared to strains that carried the vaccine-type. These results, together with the finding that antibodies against both Ptx and Prn are critical in protection against pertussis [36] and [42], also seem to support the hypothesis of vaccine-driven bacterial selection.

In agreement with a previous report, we observed that PCR-fingerprinting employing enterobacterial repetitive consensus primers (ERIC-PCR) was poorly discriminatory. However, when BOXAR1 primers were used the techniques proved to be appropriate since it has a good discriminatory power and reproducibility. In addition to the PtxS1 and Prn differences referred to above, BOXAR1-PCR underscored the genotypic differences between the vaccine and circulating strains. Fingerprints of genomic DNA from all B. pertussis isolates yielded a major cluster containing 13 clinical isolates and a few small ones, all with DNA fragment sizes ranging from 0.2 to 4.0 kb (Fig. 1). Although there were small differences among the profiles of B. pertussis isolates, the vaccine strain profile was significantly distinct from all of them.

We found B. bronchiseptica in two of the patients (19 months and 12 years old) that were investigated for Bordetella infection, none of them known to be in contact with infected animals. This microorganism has been rarely found in humans and only in adult patients with known exposure to infected animals [43]. We wondered whether lack of cross protection by the vaccine was the cause for these atypical infections. Our results show that whole cell pertussis vaccine-induced antibodies that reacted with B. bronchiseptica components, and that these antibodies proved capable of inducing opsonophagocytosis of B. bronchiseptica by professional phagocytes, a potent immune mechanism against these bacteria [26] and [27]. This result is in agreement with those published by Kirimanjeswara et al. [44] who observed that B. bronchiseptica but not B. pertussis is rapidly cleared by adoptively transferred serum antibodies. Kirimanjeswara et al. [45] showed that B. pertussis blocked migration of
neutrophils and inhibit their recruitment to the lung during the first week of infection by a pertussis toxin-dependent mechanism. Consequently PTx allows B. pertussis to avoid rapid antibody-mediated clearance, a mechanism not shared by B. bronchiseptica since it does not express pertussis toxin.

Although more research will be necessary to rule out the possibility of inefficient cross protection between these two Bordetella species, other phenomena might explain these infections with B. bronchiseptica, for instance, waning immunity for the 12-year-old patient. However, we cannot provide a satisfactory explanation for the 19-month-old patient who had the vaccination schedule completed according to his age.

In summary, the results of this study show that B. pertussis circulating bacteria were different from the strain used for vaccine production and similar to those previously found responsible for most pertussis outbreaks in other parts of the world. This result and the fact that B. bronchiseptica was found infecting the pediatric population should be considered for the development of an improved vaccine.

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