Vaccination with the 7-valent pneumococcal conjugate vaccine (PCV) has been recommended in France since 2003 for children under the age of two years who are at risk due to medical or living conditions. From 2006, the recommendation has been extended to all children under two years. The impact of PCV introduction on the incidence of pneumococcal meningitis and bacteremia and on the serotype distribution in French children and other age-groups was assessed using laboratory surveillance data. The coverage with three doses of PCV was 44% in children aged 6-12 months in 2006. From 2001/2002 to 2006, the incidence of pneumococcal meningitis decreased from 8.0 to 6.0 cases per 100,000, and the incidence of pneumococcal bacteremia decreased from 21.8 to 17.5 cases per 100,000 in children under the age of two years. For the vaccine strains, the incidence of pneumococcal meningitis and bacteremia decreased from 20.4 to 6.0 cases per 100,000, while the incidence of pneumococcal meningitis and bacteremia due to non-vaccine strains increased from 9.4 to 17.5 cases per 100,000 in this time period. The incidence in older children and adults did not decrease.

Further expansion of PCV coverage is expected to increase the impact of the vaccination in both children and adults. However, the fact that cases caused by vaccine serotypes have been partially substituted by cases of non-vaccine serotypes is likely to reduce the overall benefit of PCV in France, should this early observation be confirmed in the future.

**Introduction**

*Streptococcus pneumoniae* causes a wide spectrum of diseases, ranging from upper respiratory tract infections to severe invasive diseases. *S. pneumoniae* is the main cause of bacterial meningitis in France [1,2]. Invasive pneumococcal diseases (IPD) are more frequent in young children and the elderly and are associated with high case fatality ratio. The fatality ratio for pneumococcal meningitis has been estimated at 11% in children under the age of two years in a recent French study [3].

Two pneumococcal vaccines are currently licensed in Europe. The 23-valent pneumococcal polysaccharide vaccine was licensed in the 1980s and, although recommended for high risk individuals and elderly in many European countries, is poorly immunogenic in children under two years of age [4]. The 7-valent pneumococcal conjugate vaccine (PCV) was licensed in Europe in 2001, is immunogenic in young children and covers serotypes 4, 6B, 9V, 14, 18C, 19F and 23F. These serotypes account for between 43% and 75% of IPD in children under the age of 18 years in Western Europe [5].

Introduction of PCV in the United States in 2000 led to a dramatic decrease in those IPD that are due to vaccine serotypes, and an overall decrease of 80% in all IPD in children under the age of five years [6-8]. PCV vaccination of children had also a beneficial impact in older unvaccinated cohorts [6,7]. This herd effect is attributed to the reduction of pneumococcal carriage in the oropharynx of young children after PCV vaccination, reducing the transmission of vaccine-type pneumococcal strains to unvaccinated children and adults [9]. A slight increase in IPD due to non-vaccine serotypes was observed in American children after the introduction of PVC [7,8]. This did not significantly affect the overall reduction in pneumococcal disease incidence in American children [7,8], but has been found to negatively affect the impact of PCV vaccination in high risk populations such as native Alaskan children [10].

In France, PCV has been recommended since 2003 for children under the age of two years who are at risk due to medical or living conditions (children in day care with at least two other children for more than four hours per week, children in families with more than two children, or children breast-fed for less than two months [11]). 79% to 89% of children under two years fall in this category in France [12]. Since June 2006, PCV vaccination has been extended to all children under the age of two years [13]. The French vaccination schedule for PCV contains three doses at the ages of two, three, and four months, administered together with the vaccines against diphtheria, tetanus, poliomyelitis, pertussis and *Haemophilus influenzae* type b, and then a booster at the age of 12-15 months.

The impact of PCV on IPD incidence at the national level has not been assessed in France. Moreover, as the serotype coverage of PCV appears to be lower in Europe than in North America, it is of particular interest to analyse the impact of this vaccine on pneumococcal incidence and serotype distribution in France and in other European countries.

We used surveillance data to evaluate the effect of PCV vaccine recommendations for children at risk, a definition that encompasses the majority of each birth cohort, on the incidence of pneumococcal invasive disease and serotype distribution in 2006, four years after the introduction of PCV.
Methods

Data collection

Pneumococcal surveillance in France relies on two hospital-based laboratory surveillance networks, Epibac and the National Reference Centre for Pneumococci (NRCP) network. Since 1987, Epibac, a national hospital-based laboratory network, has collected data on six severe invasive bacterial diseases including *S. pneumoniae*. Pneumococcal invasive cases are defined as the isolation of *S. pneumoniae* from blood (bacteraemia) or cerebrospinal fluid (meningitis). The participating hospital-laboratories collect information on pneumococcal invasive cases prospectively and report annually to the Institut de Veille Sanitaire. The collected data include age, sex, and site of isolation. In 2006, 307 laboratories participated, covering 79% of the French acute care hospital admissions.

Since 2001, all pneumococcal strains isolated from cerebrospinal fluid and from blood in children under 15 years-old have been collected from hospital-laboratories and sent to the NRCP by 22 regional laboratories organised into a pneumococcal surveillance regional scheme (Observatoires Régionaux des Pneumocoques). NRCP serotyped all collected strains using latex particles sensitized with a panel of antisera that was purchased from the Statens Serum Institut (Copenhagen, Denmark) and allowed to determine 90 serotypes. Pneumococcal strains with known serotypes from the Statens Serum Institut and from the NRCP collection were used as internal quality controls.

Data analysis

The annual incidence of pneumococcal bacteraemia and meningitis cases was calculated using the number of cases reported to the Epibac network as the numerator and the French population covered by Epibac participating hospitals as the denominator. The latter was estimated from the proportion of national public and private acute-care hospital admissions covered by the participating laboratories. This proportion was computed using the National Hospital Annual Activities Database which is an exhaustive source of information regarding inpatient hospital stays, managed by the Directorate for Research, Studies, Evaluation and Statistics (DREES) at the Ministry of Health. French population data is issued each year by the National Institute for Statistics and Economical Studies (INSEE).

Age-specific incidence rates were calculated in the same way using INSEE population data by age. Serotype/age-specific incidence rates for pneumococcal bacteraemia and meningitis were estimated by applying the age distribution of pneumococcal serotypes from the NRCP to age-specific incidence rates. For this analysis, serotypes 4, 6B, 9V, 14, 18C, 19F and 23F were grouped as vaccine serotypes (VT), and other serotypes as non-vaccine serotypes (NVT).

Data from 2001 and 2002, representing the pre-vaccination situation, were aggregated.

Confidence intervals (CI) for incidence rates were estimated using Poisson distribution. Differences in age-specific and serotype/age-specific pneumococcal bacteraemia and meningitis incidence rates between 2001/2002 and 2006 were tested using Fisher’s test for binomial data. Statistical analysis was performed using Stata 9.0 (Stata Corporation, College Station, Texas).

Due to the very recent introduction of the PCV vaccine into the immunisation schedule in France, data are not yet available from the routine infant vaccination coverage monitoring tool, which is based on the health certificates filled in for each child at the age of 24th months. Instead we used data based on PCV sales for the trend analysis. In addition, three specific interview studies, one in 2004, one in 2006 and one in 2007, were conducted in representative samples of French mothers including 1,739, 1,008 and 1,005 mothers, respectively [12,14].

Results

Vaccine coverage

PCV sales increased from 0.6 to 1.6 doses per child under two years between 2003 and 2006. Coverage with three doses of PCV was estimated in three specific surveys at 27% in six month-old children in 2004 [12], at 44% in 6-12 month-old children in 2006 and at 56% in six to 12 month-old children in 2007 [14].

Pneumococcal meningitis and bacteraemia in 2001-2002

In 2001/2002, Epibac hospital-laboratories reported 7,469 cases of pneumococcal bacteraemia and 771 cases of pneumococcal meningitis. 181 (24%) cases of pneumococcal meningitis and 493 (7%) cases of pneumococcal bacteraemia occurred in children under two years; 194 (25%) cases of pneumococcal meningitis and 3,806 (25%) cases of pneumococcal bacteraemia occurred in adults over the age of 64 years. The reported number of cases and the estimated incidence by age-group are presented in Tables 1 and 2. The annual incidence of IPD in France was estimated at 9.4 cases per 100,000 population (95% CI [9.2, 9.6]) in 2001/2002.

Evolution of pneumococcal meningitis and bacteraemia incidence by age from 2001/2002 to 2006

From 2001/2002 to 2006, pneumococcal meningitis in children under two years decreased from 8.0 to 6.0 cases per 100,000 population, a decline of 25% (95% CI [2,43], p=0.04), and pneumococcal bacteraemia decreased from 21.8 to 17.5 cases per 100,000 population, a decline of 20% (95% CI [6,32], p=0.04). In the same period, pneumococcal meningitis incidence showed a not statistically significant increase from 0.69 to 0.73 cases per 100,000 population (+11% 95% CI [-7,21]) and pneumococcal bacteraemia incidence showed a statistically significant increase from 8.2 to 9.0 cases per 100,000 population (+11% 95% CI [6,15]) in older children and adults (Tables 1 and 2).

Incidence by serotype

In children under two years, the overall decrease in pneumococcal meningitis and bacteraemia incidences was associated with a shift in serotype distribution, NVT pneumococcal meningitis and bacteraemia cases partially replacing VT pneumococcal meningitis and bacteraemia cases (Figures 1 and 2). VT pneumococcal meningitis incidence decreased from 5.6 to 1.0 cases per 100,000 population, a 81% decline (95% CI [67,89], p<10-3) and VT pneumococcal bacteraemia decreased from 9.4 cases per 100,000 population, a 56% decline (95% CI [41,191], p<10-3) and pneumococcal bacteraemia increased from 7.0 to 12.2 cases per 100,000 population, a 74% rise (95% CI [39,114], p<10-3).
Evolution of serotype distribution

We determined the serotype for 156 pneumococcal strains isolated from meningitis cases and 246 pneumococcal strains isolated from bacteraemia cases in children under two years, in 2001/2002, as well as 67 strains isolated from meningitis and 99 isolated from bacteraemia in 2006.

In children under the age of two years, VT pneumococcal strains accounted for 68% (274/402) of the serotyped strains isolated from pneumococcal meningitis and bacteraemia cases in 2001/2002 and 25% (42/166) in 2006.

Among NVT pneumococcal meningitis and bacteraemia cases that occurred in children under two years in 2006, serotypes 19A and 7F were the most frequent (Figure 3). Together they accounted for 37% of pneumococcal meningitis and bacteraemia. From 2001/2002 to 2006, the proportion of 19A strains increased from 8% to 19% in meningitis cases (p=0.03) and from 11% to 17% in bacteraemia cases (p=0.007).

Table 1
Reported pneumococcal meningitis cases and estimated pneumococcal meningitis incidence by age in 2001/2002 and 2006, France (source: Epibac)

<table>
<thead>
<tr>
<th>Age group</th>
<th>2001/2002</th>
<th>2006</th>
<th>No. of reported cases</th>
<th>Cases/100,000/year</th>
<th>Incidence rate ratio, CI 95%</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 2 years</td>
<td>181</td>
<td>74</td>
<td>8.0</td>
<td>6.0</td>
<td>0.75 [0.57,0.98]</td>
<td>0.036</td>
</tr>
<tr>
<td>2 - 15 years</td>
<td>74</td>
<td>41</td>
<td>0.5</td>
<td>0.5</td>
<td>1.02 [0.70,1.50]</td>
<td>0.922</td>
</tr>
<tr>
<td>16 - 64 years</td>
<td>322</td>
<td>199</td>
<td>0.6</td>
<td>0.6</td>
<td>1.11 [0.93,1.32]</td>
<td>0.254</td>
</tr>
<tr>
<td>&gt; 64 years</td>
<td>194</td>
<td>106</td>
<td>1.4</td>
<td>1.3</td>
<td>0.97 [0.77,1.23]</td>
<td>0.857</td>
</tr>
<tr>
<td>Total</td>
<td>771</td>
<td>420</td>
<td>0.9</td>
<td>0.9</td>
<td>0.98 [0.87,1.11]</td>
<td>0.785</td>
</tr>
</tbody>
</table>

Table 2
Reported pneumococcal bacteraemia cases and estimated pneumococcal bacteraemia incidence by age in 2001/2002 and 2006, France (source: Epibac)

<table>
<thead>
<tr>
<th>Age group</th>
<th>2001/2002</th>
<th>2006</th>
<th>No. of reported cases</th>
<th>Cases/100,000/year</th>
<th>Incidence rate ratio, CI 95%</th>
<th>p</th>
</tr>
</thead>
<tbody>
<tr>
<td>&lt; 2 years</td>
<td>493</td>
<td>217</td>
<td>21.8</td>
<td>17.5</td>
<td>0.80 [0.68,0.94]</td>
<td>0.007</td>
</tr>
<tr>
<td>2 - 15 years</td>
<td>416</td>
<td>274</td>
<td>2.7</td>
<td>3.3</td>
<td>1.22 [1.04,1.42]</td>
<td>0.013</td>
</tr>
<tr>
<td>16 - 64 years</td>
<td>2,754</td>
<td>1,681</td>
<td>4.9</td>
<td>5.4</td>
<td>1.10 [1.03,1.16]</td>
<td>0.003</td>
</tr>
<tr>
<td>&gt; 64 years</td>
<td>3,806</td>
<td>2,329</td>
<td>26.8</td>
<td>29.0</td>
<td>1.08 [1.03,1.14]</td>
<td>0.003</td>
</tr>
<tr>
<td>Total</td>
<td>7,469</td>
<td>4,501</td>
<td>8.5</td>
<td>9.2</td>
<td>1.08 [1.05,1.13]</td>
<td>0.000</td>
</tr>
</tbody>
</table>

Figure 1
Estimated pneumococcal meningitis incidence by serotype in children under two years of age, France 2001-2006 (source: Epibac-NRCP)

Figure 2
Estimated pneumococcal bacteraemia incidence by serotype in children under two years of age, France 2001-2006 (source: Epibac-NRCP)
27% in bacteraemia cases ($p<10^{-3}$); the proportion of 7F strains increased from 3% to 18% in meningitis cases ($p<10^{-3}$) and from 1% to 10% in bacteraemia cases ($p<10^{-3}$). Other non-vaccine serotypes accounted for less than 8% of pneumococcal meningitis and bacteraemia cases in children under two years in 2006.

From 2001/2002 to 2006, the incidence of pneumococcal meningitis and bacteraemia caused by each of the seven vaccine serotypes decreased. The incidence of pneumococcal meningitis and bacteraemia due to serotypes which are not included in the vaccine but are part of the same serogroup as a vaccine serotype – with the exception of serotype 19A, i.e. serotypes 23B, 6A, 18B, 9N, and 23A – remained unchanged (Figure 3).

**Discussion**

French recommendations for PCV vaccination in 2003 included a large proportion of French children under the age of two years, while other European countries targeted only high risk children [4]. Between 2005 and 2006, vaccination with PCV has been extended to all children under two years in France as well as in other European countries such as Belgium, England, Germany, Luxembourg, the Netherlands, and Norway [15]. The early recommendations and the fact that the French surveillance for invasive pneumococcal diseases allows the analysis of trends in incidence and serotype distribution provided an opportunity to analyse the impact of PCV introduction in France. It is the second analysis of this kind in a European country at the national level following the analysis published this year from Norway [16].

Although PCV introduction in France was associated with a 71% decrease in vaccine-type IPD incidence between 2001/2002 and 2006 in children under two years, the overall decrease of IPD in this age group was only 21% (95% CI [10,31]). The fact that the decline was observed only in children under two years and only for cases due to vaccine serotype strains strongly argues for a role of PCV vaccination in this evolution.

**Impact of PCV vaccination on IPD incidence**

The 21% decline of the disease in French children is far below the 77% reduction observed in children under the age of five years in the regions covered by the United States (US) Centers for Disease Control and Prevention’s ‘Active Bacterial Core surveillance’ in 2005 and the 52% reduction observed in children under two years in Norway between 2004/2005 and 2007 [8,16]. Moreover, an indirect benefit of PCV vaccination in other age-groups has so far not been observed in France. The limited estimated vaccination coverage for PCV, below 60% in 2006, could explain in part this modest impact of PCV vaccination. Although PCV vaccine coverage has improved in the recent years, it remains well below the usual vaccine coverage for infants in France [14]. Expansion of the PCV vaccination coverage should lead to a further reduction in the IPD that are caused by vaccine serotypes in children and, through indirect effects, also in adults.

The overall reduction in IPD decrease that we found in young children is in agreement with the results reported by a French network of paediatricians which indicate a 28% decrease in the number of pneumococcal meningitides in 2-24 month-old children.

**Figure 3**

Estimated incidence of pneumococcal meningitis and bacteraemia by serotype in children under the age of two years, evolution from 2001/2002 to 2006, France (source: Epibac-NRCP)
between 2001/2002 and 2005 [17]. Another survey conducted in 18 hospitals in northern France found a much greater reduction (82%) in pneumococcal meningitis incidence in children under two years between 2002 and 2005 than was found in the above and in our results [18]. A high PCV coverage in the northern region of France and the small number of cases involved in this retrospective survey (n<8 in 2005) are possible explanations for these differences.

Serotype replacement
The 71% reduction in IPD incidence due to VT strains was associated with a 85% rise in cases due to NVT strains in children under two years from 2001/2002 to 2006. The magnitude of this replacement impacted the overall effect of PCV vaccination: IPD incidence in French young children decreased between 2001/2002 and 2005 but did not decrease further from 2005 to 2006 despite a 20% rise in PCV sales. During that later period, the decrease in cases due to VT strains was balanced by an increase of the same magnitude in cases due to NVT strains. Replacement of VT by NVT strains has been observed to a smaller extent also in American children following PCV introduction. The increase of NVT pneumococcal disease in children under five years in the US estimated from ‘Active Bacterial Core surveillance’ data between 1998/1999 and 2005 was only 29% [8]. That this increase was higher in France may be due to the lower PCV serotype coverage in young children in France compared to the US. Before PCV introduction, 68% of IPD in children under two years were caused by PCV serotypes in 2001/2002 in France, compared to 83% in children under five years in 1998/1999 in the US [7].

Among NVT strains, two single serotypes – 19A and 7F – accounted for 37% of pneumococcal strains in 2006 in France. In the US, the 19A serotype has been found to be the predominant serotype in pneumococcal invasive cases in the years following PCV implementation, accounting for 40% of cases in children under the age of five 5 years in 2005, according to the results of the ‘Active Bacterial Core surveillance’ [8].

No decline in IPD was observed in older children and adults; on the contrary we identified a small but significant increase. However, as this trend had already been observed from 1998 to 2002 before the introduction of PCV in France, the possible contribution of vaccination to the increase cannot be conclusively assessed [19].

The evolution of pneumococcal invasive incidence in children in France can be compared with the situation observed in different areas of Spain after PCV introduction. Four regional Spanish studies were performed with the following results: no change in IPD incidence in a Barcelona district between 1999/2001 and 2002/2004 [20], a decrease in the Basque region between 2000/2001 and 2004/2005 [21] and in the Basque and Navarre regions between 1998/2001 and 2003 [22], and even an increase in pneumococcal invasive cases in Barcelona between 1997/2001 and 2002/2006 [23]. The reasons given by the investigators for this limited impact of PCV on IPD incidence refer to the conditions of PCV introduction in Spain: Vaccine coverage was low in Spain, where PCV is not subsidised, and the serotype coverage of PCV was significantly lower than the PCV coverage in the US (43% in the Navarre region) [20,22-24]. An increase in the frequency of pneumococcal invasive cases due to non-vaccine strains after PCV introduction was also found in three of these studies [20,23,24].

Strengths and limitations of the study
We are confident about our incidence estimates because of the high and sustained coverage of the Epibac laboratory network combined with extrapolations made on a reliable source of information (the French national Hospital Annual Statistic database). Furthermore, we regularly monitored the reporting of cases by the participating laboratories through three sources capture-recapture analysis to ensure the exhaustiveness of the reports. The rate of underreporting were estimated at between 10% and 20% in these analyses [25,26].

We cannot completely exclude a change in the rate of pneumococcal case reporting in the last years. However, significant changes in reporting for pneumococcal cases alone are unlikely due to the following reasons: Firstly, the reporting rate of other bacterial diseases included in Epibac surveillance has not changed until 2005 as shown by a recent three sources capture-recapture analysis for invasive meningococcal diseases [26]; secondly, pneumococcal data show opposite trends for VT- and NVT-related cases; and thirdly, the cases are reported, by the vast majority of participating laboratories, through automatic extraction of microbiology isolates registration.

Incidence and serotype data are issued from two networks whose regional coverage is not identical. This may have introduced biases in the estimation of serotype/age-specific incidence evolution. However, each both networks covers more than 300 hospitals localised in all French regions, and the PCV serotype coverage did not vary with the geographical origin of pneumococcal strains (data not shown).

The evolution of individual serotypes should be interpreted with caution given the small number of strains involved in the 2006 analysis. Emergence of serotypes 19A and 7F may not be due to PCV introduction alone, as changes in serotypes distribution can also occur for other reasons than vaccination pressure. The findings of this early analysis must be seen as a preliminary description of the PCV impact in France, IPD evolution and the extent of serotype replacement will be closely monitored in the next years through ongoing epidemiological and bacteriological surveillance.

Conclusion
In conclusion, PCV introduction was followed by a significant decrease in IPD in young children in France. Further improvement of PCV coverage should further increase the positive impact of PCV on vaccine-type pneumococcal invasive diseases in both children and adults in the next years, if a positive herd immunity effect is observed. If, on the other hand, the partial substitution of the cases that are caused by vaccine serotypes with cases caused by non-vaccine serotypes, that was observed in our early analysis in young children, is confirmed in the coming years, this would lead to a reduction of the positive impact of PCV vaccination in France. Theses results emphasise the need for ongoing surveillance of the pneumococcal disease incidence and serotype in countries introducing PCV. The imminent availability of pneumococcal vaccines covering a broader range of the serotypes implicated in IPD in young children could limit the effect of serotype replacement and improve the impact of immunisation on IPD.
Acknowledgments:
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