Impact of pneumococcal polysaccharide vaccine in people aged 65 years or older

Robert I Menzies
BAppSc, MPH, PhD, Deputy Director, Surveillance, and Senior Lectures
Sanjay H Jayasinghe
MB BS, MPH, FAFPHM, Immunisation Research and Senior Health Physician, 1 MJA 2014; 200: 112–115 Director, 1 and Professor 2 MB BS, PhD, FRACP, Preventable Diseases, 1

Vicki L Krause
MD, DTM&H, FAFPHM, Sydney Medical School, Surveillance of Vaccine

Clayton K Chiu
MB BS, MPH, BAppSc, MPH, PhD, national funded National Immunisation Program from 2005.2

Peter B McIntyre
MB BS, PhD, FRACP, Director, 1 and Professor 2 Public Health Specialist and Director 3

Ecological analysis of trends in invasive pneumococcal disease (IPD) notification rates and vaccine effectiveness estimation using the screening method, using data on Australians aged > 65 years (23vPPV funded) and 50–64 years (23vPPV not funded).

Intervention: National 23vPPV program for people aged > 65 years and national 7vPCV program for infants, both commencing in 2005.


Results: The proportion of people aged > 65 years who were vaccinated within the previous 5 years in jurisdictions excluding Victoria ranged from 41% to 64% over the study period, with no clear trend over time. Incidence rate ratios in the > 65-year age group were 0.11 (95% CI, 0.09–0.14) for 7vPCV serotypes, 1.64 (95% CI, 1.41–1.91) for 23vPPV–non-7vPCV serotypes and 2.07 (95% CI, 1.67–2.57) for non-23vPPV serotypes. The incidence rate ratio for total IPD was 0.65 (95% CI, 0.59–0.71) for people aged > 65 years, and 0.80 (0.71–0.90) for people aged 50–64 years. The estimate of 23vPPV effectiveness was 61.1% (95% CI, 55.1%–66.9%).

Conclusions: The greater reduction in IPD among > 65-year-olds compared with 50–64-year-olds did not reach statistical significance. However, vaccine effectiveness was significant. Greater reductions in IPD in > 65-year-olds would be expected from the indirect effects of using 13-valent pneumococcal conjugate vaccine in infants (introduced for Australian infants in 2011) and an increase in 23vPPV coverage.

Abstract

Objective: To evaluate the impact and effectiveness of the 23-valent pneumococcal polysaccharide vaccine (23vPPV) in > 65-year-old Australians in the context of concurrent 7-valent pneumococcal conjugate vaccine (7vPCV) use in infants.

Design, patients and setting: Ecological analysis of trends in invasive pneumococcal disease (IPD) notification rates and vaccine effectiveness estimation using the screening method, using data on Australians aged > 65 years (23vPPV funded) and 50–64 years (23vPPV not funded).

Methods

Notifications of IPD were obtained from the National Notifiable Diseases Surveillance System (NNDSS), which notionally captures all laboratory-confirmed cases of IPD. To distinguish the impact of the 7vPCV and the 23vPPV, notifications were aggregated by serotype category: serotypes contained in both vaccines (7vPCV serotype), those contained in the 23vPPV but not in the 7vPCV (23vPPV–non-7vPCV serotype) and those not contained in either vaccine (non-23vPPV serotype). Notifications for people in Victoria and people recorded as Indigenous were excluded, as funded programs for these populations commenced earlier than the national program (in 1998 and 1999, respectively). The proportion of notifications for which speci-
moms were serotyped increased from 70% in 2002–2003 to 87% in 2006. To ensure that time trends in serotype categories were not distorted by this change, the serotype distribution among untyped cases was inferred from the serotype distribution among typed cases, by jurisdiction and by year.

This study was exempt from the requirement for ethics approval as it was conducted as a quality assurance exercise pertaining to the National Immunisation Program, under the auspices of the Australian Technical Advisory Group on Immunisation. De-identified data were provided for this purpose by the Communicable Diseases Network Australia.

**Trends in IPD and vaccination coverage**

Population rates of IPD by serotype category in people aged ≥65 years by year from 1 January 2002 to 31 December 2011 were plotted with 95% confidence intervals, which were calculated using the Poisson distribution of notification numbers. The estimated residential populations, minus Indigenous population estimates from the 2006 Australian census, were used as denominators for calculations of rates in the non-Indigenous population.11

Vaccination coverage estimates for people aged ≥65 years, by jurisdiction, were available from adult vaccination surveys conducted by the Australian Institute of Health and Welfare in 2004, 2006 and 2009.12-14 Separate estimates for the 65–74-year and ≥75-year age groups by jurisdiction were available from the 2004 and 2006 surveys.

The proportion of 23vPPV-type IPD cases that were recorded as being in “fully vaccinated” people (ie, those vaccinated within 5 years, according to national recommendations at the time) who were aged ≥65 years was derived from the NNDS for each year that population coverage data were available (2004, 2006 and 2009). For these VE calculations, cases recorded on the NNDS as having occurred in people in Victoria or Indigenous people were not included, as their different dates of 23vPPV funding do not influence VE estimates. In addition, cases that occurred in Indigenous people were included in population coverage data, so the method required their inclusion in the study population.

A logistic regression model was fitted using the GENMOD procedure in SAS 9.1 (SAS Institute) as described previously.16 Data were stratified by year, age group and, for 2004 and 2006, by jurisdiction. A sensitivity analysis of the VE estimates was conducted, recalculating VE using a deviation of ±10% of the total population in coverage estimates. Statistical analysis was carried out in SAS 9.1.3. Statistical significance was established by non-overlapping 95% confidence intervals, of rates and rate ratios.

**Results**

From 2002 to 2011, there were 3978 IPD notifications for Australians aged ≥65 years who were not in Victoria and were not Indigenous.

**1 IPD notification rates by serotype category for ≥65-year-old Australians and pneumococcal vaccination coverage, 2002–2011***

<table>
<thead>
<tr>
<th>Year</th>
<th>7vPCV serotype IPD rate</th>
<th>23vPPV–non-7vPCV serotype IPD rate</th>
<th>Non-23vPPV serotype IPD rate</th>
<th>Vaccination coverage</th>
</tr>
</thead>
<tbody>
<tr>
<td>2002</td>
<td>15%</td>
<td>20%</td>
<td>5%</td>
<td>30%</td>
</tr>
<tr>
<td>2003</td>
<td>20%</td>
<td>25%</td>
<td>10%</td>
<td>40%</td>
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<tr>
<td>2004</td>
<td>25%</td>
<td>30%</td>
<td>15%</td>
<td>50%</td>
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<tr>
<td>2005</td>
<td>30%</td>
<td>35%</td>
<td>20%</td>
<td>60%</td>
</tr>
<tr>
<td>2006</td>
<td>35%</td>
<td>40%</td>
<td>25%</td>
<td>70%</td>
</tr>
</tbody>
</table>

**IPD = invasive pneumococcal disease. 7vPCV = serotypes contained in the 7-valent pneumococcal conjugate vaccine and 23-valent polysaccharide pneumococcal vaccine. 23vPPV = serotypes contained in the 23-valent polysaccharide pneumococcal vaccine but not the 7-valent pneumococcal conjugate vaccine. Non-23vPPV = serotypes not contained in either vaccine. * IPD notification rates do not include people in Victoria or Indigenous people; serotype categories are adjusted for untyped cases; error bars for IPD notification rates are 95% confidence intervals; and error bars for vaccination coverage are ranges of self-reported vaccination coverage for individual jurisdictions excluding Victoria.**
Trends in IPD rates and vaccination coverage

Annual IPD notification rates in people aged ≥ 65 years by serotype category are shown in Box 1. There was a substantial and statistically significant decrease in 7vPCV serotype IPD during the post-7vPCV era (2005–2011), as well as significant increases in 23vPPV–non-7vPCV and non-23vPPV serotypes, based on non-overlapping confidence intervals of annual rates.

The serotype most associated with replacement following 7vPCV introduction internationally — 19A — increased from 3% of isolates in 2002–2004 to 22% in 2010–2011. The range of self-reported vaccination coverage (percentage vaccinated in the previous 5 years) estimates for those aged ≥ 65 years are also shown in Box 1, for all jurisdictions except Victoria. Coverage ranged from 41% to 53% in individual jurisdictions in 2004, increased to 51%–64% in 2006 and decreased to 48%–56% in 2009.

IPD rates in vaccinated and unvaccinated age groups

Pre-7vPCV (2002–2004) to post-7vPCV (2010–2011) changes in IPD rates in the ≥ 65-year age group and the 50–64-year age group are shown in Box 2. In both age groups there were substantial, statistically significant decreases for 7vPCV serotypes and increases for 23vPPV–non-7vPCV and non-23vPPV serotypes, based on IRRs not overlapping 1.0. The magnitude of these changes did not differ significantly between the two age groups. For all serotypes, the IRR point estimate was lower in the ≥ 65-year age group, but confidence intervals for the two age groups overlapped.

Vaccine effectiveness estimates

Numbers of IPD cases and VE estimates for 23vPPV against 23vPPV-type IPD in Australians aged ≥ 65 years are shown in Box 3. All VE estimates were statistically significantly above zero. The point estimate for 2009 was lower than for 2004, but was compatible with it, as confidence intervals overlapped. A sensitivity analysis to evaluate the impact of varying population coverage estimates for 23vPPV yielded an upper VE estimate of 75.8% (95% CI, 72.1%–79.9%) if true population coverage was 10% higher than estimated in the adult vaccination survey and a lower VE estimate of 40.5% (95% CI, 31.1%–49.9%) if true population coverage was 10% lower than estimated.

Discussion

Changes in IPD rates over time by serotype category presented here provide evidence of a substantial herd immunity impact in older people due to 7vPCV use in infants. However, an impact on IPD rates directly resulting from 23vPPV use in older people, by comparing changes in vaccinated and unvaccinated age groups, was not clearly shown. The VE estimate for 23vPPV against 23vPPV-type IPD was 61.1%. An overall decrease of 35% was observed in total IPD rates in ≥ 65-year-olds 6–7 years after the commencement of the nationally funded programs for 7vPCV and 23vPPV (25.2 notifications/100 000 population/year in 2002–2004 v 16.4 in 2010–2011).

Herd immunity impacts in adults from use of the 7vPCV in children have been shown in many countries. Herd immunity impacts on total IPD rates are heavily dependent on the pre-vaccination serotype distribution in adults and the length of time since 7vPCV introduction, as serotype replacement increases over time.
Indigenous people had one of the highest proportions of 7vPCV-type IPD out of total IPD in the world, similar to that in the US, and these are the only two countries with net decreases in IPD reported for older people following 7vPCV introduction.\textsuperscript{13}

Trends in IPD rates by year in Australia in our study did not show clear evidence of a reduction of disease incidence due to use of 23vPPV in people aged $\geq 65$ years. However interpretation of this finding is complicated by two factors: an overall modest level of 23vPPV coverage and a relatively small increase in coverage after national funding began in 2005; and the apparent indirect effects of introducing the 7vPCV for infants at the same time. The comparison of rates in vaccinated and unvaccinated age groups, both subject to herd immunity impacts from infant vaccination, allows the possibility of some impact from the 23vPPV. The absence of impacts on population IPD rates following publicly funded 23vPPV for $\geq 65$-year-olds has also been reported in the US\textsuperscript{18} and United Kingdom.\textsuperscript{19} Gradual increases in coverage also occurred in those settings, and formal VE assessments in adults have consistently shown significant VE.\textsuperscript{20,21}

All observational methods used to estimate VE are subject to bias. For our application of the screening method, different methods were used to ascertain the vaccination status of the general population (telephone survey) and the vaccination status of 23vPPV-type IPD cases (general practitioner and/or patient interview). However, our sensitivity analysis showed that the VE estimate remained statistically significant even if the true population coverage was 10% lower than the adult vaccination survey estimates. A study of 23vPPV vaccination status in older people in Victoria found that patient recall underestimated vaccination status by 6\% compared with medical records.\textsuperscript{23}

During the period of our study, a single revaccination was recommended for people first vaccinated at $\geq 65$ years of age. As of December 2011, this is no longer recommended.\textsuperscript{10} The latest national estimate of the proportion of people aged $\geq 65$ years who have ever received 23vPPV is a modest 59\%.\textsuperscript{13} Given the evidence of the vaccine’s effectiveness, higher coverage would be expected to increase the impact of the vaccine in reducing IPD incidence.

Data are yet to emerge on the herd immunity impact from 13vPCV use in Australian children. However, if similar effects are seen from the additional six serotypes as from the 7vPCV, there would be a further reduction in IPD in older people and, therefore, less potential benefit from the 23vPPV.

The appeal of a conjugate vaccine used in older people includes potential, although unproven, benefits such as a superior response to booster doses and impacts on carriage and non-invasive pneumonia. Herd immunity impacts of 7vPCV on non-invasive pneumonia in older people have been reported as being non-existent, very small or extensive.\textsuperscript{24-26} A randomised controlled trial assessing the impact of 13vPCV use in older people on pneumonia is currently underway.\textsuperscript{27} However, this trial is not being conducted alongside concurrent use of 13vPCV in infants. The incremental benefits of 13vPCV use in older people in addition to an infant program would be more difficult to evaluate.

In conclusion, our data show moderate effectiveness of the 23vPPV against IPD in older Australians, consistent with that shown in comparable populations elsewhere. In combination with herd immunity impacts from 7vPCV in children, this resulted in a 35\% decrease in IPD in those aged $\geq 65$ years. Further benefits could be expected if an increase in 23vPPV coverage in older people could be achieved.

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