Influenza hospitalizations in Australian children

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SUMMARY

Australia’s National Immunisation Program (NIP) provides free influenza vaccination for children at high risk of severe influenza; a pilot-funded programme for vaccine in all children aged 6 months to <5 years in one of eight states, has seen poor vaccine impact, related to recent vaccine safety concerns. This retrospective review examined influenza hospitalizations in children aged <16 years from three seasons (2011–2013) at two paediatric hospitals on opposite sides of the country. Comparisons of this cohort were made with state-based data on influenza-coded hospitalizations and national immunization register data on population-level immunization coverage. Of 740 hospitalizations, the majority were aged <5 years (476/740, 64%), and a substantial proportion (57%) involved healthy children, not currently funded for influenza vaccine. Intensive care unit admission occurred in 8·5%, and 1·5% of all children developed encephalitis. Use of antiviral therapy was uncommon (20·5%) and decreasing. Of those hospitalized, only 5·0% of at-risk children, who are currently eligible for free vaccine, and 0·7% of healthy children were vaccinated prior to hospitalization. This was consistent with low population-wide estimates of influenza vaccine uptake. It highlights the need to examine alternative strategies, such as universally funded paediatric influenza vaccination, to address disease burden in Australian children.

Key words: Children, epidemiology, hospitalization, Influenza, vaccination.

INTRODUCTION

Influenza is a respiratory infection that is the most frequently reported notifiable disease and a common cause of hospitalizations each year in Australia. Hospitalization rates during 2005–2008 in children aged <5 years (48–65/100,000) exceeded those in the elderly aged >65 years (13·4–20/100,000 annually) [1–3]. The highest hospitalization rates are in those aged <6 months (192–270/100,000), a group potentially amenable to protection through maternal vaccination during pregnancy [3, 4].

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For children and young adults, Australia has ‘targeted’ influenza vaccination funded under the National Immunisation Program (NIP). Since 2010, free NIP-funded vaccine has been available for high-risk individuals with chronic medical conditions aged ≥6 months, pregnant women, and Indigenous children aged ≥15 years [5, 6]. A funded influenza immunization programme for all children aged 6 months to <5 years exists in only one state (Western Australia; WA), introduced in 2008 after three deaths in healthy children in a single season [7]. However, following an unexpected increase in febrile seizures with one brand of trivalent influenza vaccine (TIV) in Australian children aged <5 years during 2010 [8, 9], the majority of which were documented in WA due to higher uptake with its funded vaccination programme (then >40%), public confidence was affected; vaccine uptake fell precipitously to <10% [10] and remains low despite the safety of other brands being confirmed [11, 12]. This contrasts with universal vaccination programmes in the United States, Canada, and UK where coverage rates have increased up to 60% of eligible children [13–18].

Uncertainty remains as to how well Australia’s targeted strategy is working and whether at-risk children are being adequately protected through vaccination. Data on vaccine uptake in those with underlying medical conditions is scarce. One small study conducted in 2012 found that coverage in at-risk children attending outpatient specialist clinics was 41% but relied on unverified parental report in this select population [19]. Previous epidemiological data [1] regarding severe influenza in Australia have relied on administrative hospitalization data, without clinical confirmation; this provides no detailed information on who is affected or vaccine uptake [20]. Other small studies have been limited to examining a single season or centre [21, 22]. To guide future prospective surveillance aimed at assessing annual influenza disease burden and vaccine effectiveness against influenza hospitalization in children and adults [23], we conducted a retrospective examination of influenza-related hospitalizations in two large paediatric hospitals during 2011–2013, to describe the demographic and clinical characteristics and outcomes of severe influenza. The data would also help to better understand the pattern of hospitalization in children and to inform influenza surveillance and vaccination strategies.

MATERIALS AND METHODS

This was a retrospective review of influenza hospitalizations in children aged <16 years, over a 3-year period from January 2011 to December 2013 in two major centres: the Children’s Hospital at Westmead (CHW), Sydney, New South Wales (NSW) and the Princess Margaret Hospital for Children (PMH), Perth, WA, which are part of the Flu Complications Alert Network (FluCAN) [24], a sentinel network of 17 hospitals collecting data on adult and paediatric influenza hospitalizations each season. CHW and PMH function as both general paediatric hospitals for a proportion of each city’s population, as well as tertiary referral centres within their respective states. Cases were obtained from each hospital’s laboratory database. A case was included if they were a hospitalized child who had laboratory-confirmed influenza by polymerase chain reaction (PCR) for A and B viruses (within a multiplex respiratory virus panel), at presentation or during their hospital stay, and an acute respiratory infection (ARI), defined as symptoms or signs of upper or lower respiratory tract infection (e.g. cough, coryza, sore throat, shortness of breath, wheeze, or tachypnoea), with or without fever during the study period. Diagnoses of influenza only by direct fluorescent antibody testing or point-of-care testing were excluded. In both hospitals, it is routine practice, but at the physician’s discretion, for most hospitalized children with ARI to undergo respiratory virological testing for diagnostic purposes and to facilitate infection control.

Demographic data, medical history (including comorbidities), clinical symptoms and signs at presentation (fever, respiratory distress, or other symptoms), investigations performed [chest X-ray, blood culture, lumbar punctures (LP), influenza virus subtyping], and management, including use of antivirals and antibiotics were recorded. Conditions predisposing to severe influenza infection (as listed in the Australian Immunisation Handbook [25]) were documented. Comparisons were made between seasons and according to influenza type. Pneumonia was defined as consolidation recorded on chest X-ray (CXR) report. Bacterial infection included meningitis, bacteraemia, urinary tract infection, or other sterile site infection where a bacterial pathogen was identified on an appropriate specimen by culture and/or antigen testing. We assessed outcomes including length of stay (LOS) in hospital, intensive care unit (ICU) admission, associated bacterial infection, seizures, encephalitis, or in-hospital death.

Numbers of hospitalizations at each site were compared with de-identified state-wide hospitalization data obtained from the WA and NSW Departments
of Health for children aged 0–18 years with ICD-10-AM coded influenza (J09-J10) as a primary or secondary diagnosis.

Influenza vaccination status (including the years prior to and subsequent to hospitalization) was assessed for the included hospitalized children or their mothers (if child aged <6 months) from both the medical record and the Australian Childhood Immunisation Register (ACIR), which records immunizations for all children aged <7 years [26]. In addition, we obtained population-level influenza vaccine coverage during the study period for all children contained within ACIR at the national and state levels.

Differences between groups were assessed using \( \chi^2 \) test for categorical variables and independent-sample \( t \) tests or Mann–Whitney \( U \) test for continuous variables. We used multivariate logistic regression to examine for risk factors associated with ICU admission. Predictors with \( P < 0.20 \) on univariate analysis were incorporated into a stepwise regression model with backwards elimination to determine adjusted odds ratios. For hospital LOS, the distribution was expected to be substantially positively skewed, therefore we used a generalized linear model with negative binomial distribution for multivariate regression analysis. Adjusted ratios of mean LOS were calculated for predictor variables from the final multivariate model. Calculations were performed using SPSS Statistics v. 21 (IBM Corp., USA).

Ethical approval was obtained for this research from the Sydney Children’s Hospitals Network Human Research Ethics Committee (reference HREC/13/SCHN/402).

RESULTS

During the study period, 740 PCR-confirmed influenza hospitalizations occurred (Table 1). The majority were during the July–September winter influenza season, typical for the temperate climate locations. Overall, 65% (n = 480) of cases were admitted to CHW and 35% (n = 260) to PMH. This represented 23% (480/2075) and 41% (260/634) of all ICD-coded influenza hospitalizations during the 3 years in NSW (annual birth cohort: 99,000) and WA (annual birth cohort: 33,000), respectively [27]. Males comprised a larger percentage of hospitalizations (55.5%, \( P = 0.003 \)). Young children were over-represented; 64% (476/740) were aged <5 years; 13.4% (n = 99) of the total were aged 0–<6 months (Fig. 1). Indigenous children accounted for 4.2% of all admissions. Many children (43%, \( n = 317 \)) had one or more medical conditions predisposing to severe influenza infection, most commonly immunosuppression (13.2%), neurological disease (12.7%) and chronic respiratory disease (10.4%). There were no significant differences between CHW and PMH in terms of age distribution or proportion of children with underlying medical conditions; median LOS was slightly shorter at PMH (2 days) than CHW (3 days).

Influenza vaccination

In hospitalized children included in our study, who were aged 6 months–7 years and had data captured by the ACIR, 2.5% (12/474) had at least one influenza vaccination >2 weeks prior to hospitalization and in the same year. The rate of vaccination was greater in children at the WA site compared to NSW (4.3% vs. 1.6%). A higher proportion of children aged 6 months–7 years (10.8%, 51/474) were recorded as receiving an influenza vaccination in the year following their hospitalization. No children had data recorded in their clinical record regarding receipt of vaccination, including no record of maternal vaccination (for those aged <6 months). Population-wide estimates from ACIR revealed similarly low vaccination rates in all children within NSW (0.5–1.0%), WA (3.9–5.1%), and nationally (1.0–1.6%) during 2011–2013 (Table 2).

Influenza types/subtypes

Influenza A was the predominant type in 2011 (64.6%) and 2012 (69%). In 2013, both influenza A and B were equally common (Table 1). The median age was significantly higher for children with influenza B compared to influenza A (5.0 vs. 2.4 years, \( P < 0.001 \)). Limited influenza A subtyping was available, predominantly at PMH (see Table 1). Influenza A or B strains were not associated with any differences in clinical features or outcomes except for myositis which was more common with influenza B infection (\( P = 0.03 \)).

Clinical presentation

Overall 75% of hospitalized patients with influenza presented with influenza-like illness (fever and cough and/or sore throat) [28] (Table 1). Fewer children presented with respiratory distress/pneumonia (19.1%) or fever without source (most common in neonates; 19% of whom presented in this way). Urinary tract infection
was the most common (3·0%) culture-positive bacterial infection (Table 3); bacterial meningitis was rare (one case from *Streptococcus pneumoniae* in an 11-month-old). Bacteraemia was present in 1·8% (*n* = 13) of patients and radiologically confirmed pneumonia in 11·2% (*n* = 83, including three children with associated bacteraemia). Febrile seizures occurred in 5·9% (*n* = 44) of children, and 4·1% (*n* = 30) had an afebrile seizure (23/30 had an underlying seizure disorder). Eleven children had influenza-associated encephalopathy (Table 3).

### Table 1. Demographic information, influenza strain, clinical presentation, and investigations by year in Australian children hospitalized with laboratory-confirmed influenza during 2011–2013 at the Children’s Hospital at Westmead (NSW, Australia) and Princess Margaret Hospital (WA, Australia)

<table>
<thead>
<tr>
<th>Year</th>
<th>2011</th>
<th>2012</th>
<th>2013</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total (<em>n</em>)</td>
<td>Sites</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>199 (65·3)</td>
<td>Children’s Hospital at Westmead</td>
<td>166 (53·2)</td>
<td>184 (80·3)</td>
</tr>
<tr>
<td></td>
<td>69 (34·7)</td>
<td>Princess Margaret Hospital</td>
<td>146 (46·8)</td>
<td>45 (19·7)</td>
</tr>
<tr>
<td>Demographics</td>
<td></td>
<td>Male</td>
<td>116 (58·3)</td>
<td>168 (53·8)</td>
</tr>
<tr>
<td></td>
<td>Median age, years</td>
<td>3·2</td>
<td>3·4</td>
<td>2·8</td>
</tr>
<tr>
<td>Age groups (% of year)</td>
<td></td>
<td>0–28 days</td>
<td>6</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>29 days–&lt;6 months</td>
<td>20</td>
<td>35</td>
<td>28</td>
</tr>
<tr>
<td></td>
<td>6 months–&lt;1 year</td>
<td>24</td>
<td>34</td>
<td>22</td>
</tr>
<tr>
<td></td>
<td>1–&lt;5 years</td>
<td>72</td>
<td>125</td>
<td>100</td>
</tr>
<tr>
<td></td>
<td>5–&lt;10 years</td>
<td>50</td>
<td>82</td>
<td>53</td>
</tr>
<tr>
<td></td>
<td>10–15 years</td>
<td>27</td>
<td>32</td>
<td>20</td>
</tr>
<tr>
<td>Indigenous</td>
<td></td>
<td>12 (6·0)</td>
<td>14 (4·5)</td>
<td>5 (2·2)</td>
</tr>
<tr>
<td>Comorbidity present†</td>
<td></td>
<td>90 (45·2)</td>
<td>122 (39·1)</td>
<td>105 (45·9)</td>
</tr>
<tr>
<td></td>
<td>Respiratory</td>
<td>19</td>
<td>32</td>
<td>26</td>
</tr>
<tr>
<td></td>
<td>Prematurity</td>
<td>19</td>
<td>22</td>
<td>27</td>
</tr>
<tr>
<td></td>
<td>Cardiac</td>
<td>14</td>
<td>22</td>
<td>12</td>
</tr>
<tr>
<td></td>
<td>Diabetes</td>
<td>0</td>
<td>3</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Neurological</td>
<td>29</td>
<td>35</td>
<td>30</td>
</tr>
<tr>
<td></td>
<td>Renal</td>
<td>9</td>
<td>6</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>Immunosuppression</td>
<td>30</td>
<td>38</td>
<td>30</td>
</tr>
<tr>
<td></td>
<td>Hepatic</td>
<td>7</td>
<td>7</td>
<td>4</td>
</tr>
<tr>
<td></td>
<td>Genetic</td>
<td>15</td>
<td>20</td>
<td>16</td>
</tr>
<tr>
<td></td>
<td>Inborn error of metabolism</td>
<td>4</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Influenza strain</td>
<td></td>
<td>A Total</td>
<td>128 (64·6)</td>
<td>216 (69·0)</td>
</tr>
<tr>
<td></td>
<td>A/H1N1 pdm09</td>
<td>83</td>
<td>2</td>
<td>11</td>
</tr>
<tr>
<td></td>
<td>A/H3N2</td>
<td>22</td>
<td>77</td>
<td>13</td>
</tr>
<tr>
<td></td>
<td>A/untyped</td>
<td>23</td>
<td>137</td>
<td>89</td>
</tr>
<tr>
<td></td>
<td>B</td>
<td>70 (35·4)</td>
<td>97 (31·0)</td>
<td>116 (50·7)</td>
</tr>
<tr>
<td>Presentation†</td>
<td></td>
<td>Influenza-like illness</td>
<td>148 (78·8)</td>
<td>247 (79·2)</td>
</tr>
<tr>
<td></td>
<td>Fever without source</td>
<td>17 (8·5)</td>
<td>24 (7·7)</td>
<td>20 (8·7)</td>
</tr>
<tr>
<td></td>
<td>Pneumonia/respiratory distress</td>
<td>38 (19·1)</td>
<td>69 (22·1)</td>
<td>34 (14·8)</td>
</tr>
<tr>
<td>Investigations</td>
<td></td>
<td>Chest X ray</td>
<td>108 (54·3)</td>
<td>179 (57·4)</td>
</tr>
<tr>
<td></td>
<td>Lumbar punctures</td>
<td>31 (15·6)</td>
<td>31 (9·9)</td>
<td>28 (12·2)</td>
</tr>
</tbody>
</table>

Absolute numbers indicated with percentage within stated year in parentheses unless indicated otherwise.

* *P* = 0·003 on binomial testing.

† Individual categories are not mutually exclusive and totals may not equal the sum of all categories.
Investigations, treatment and outcomes are shown in Tables 1 and 3. Antibiotics were used in 72.3% of patients, while antivirals were prescribed to 20.5% of patients, a median of 2 days after admission (Table 3); use of antiviral therapy declined over the study period. Admission to ICU was required in 8.5% (n = 63, including two nosocomial infections), with a median LOS of 3 days [interquartile range (IQR) 2–6 days]. The majority (51/63) of ICU admissions occurred within 24 h of hospital admission. Of 474 children with vaccination status confirmed by ACIR, 36 were admitted to ICU, none of whom were vaccinated. No vaccinated child (n = 12 in total) required ICU admission. Multivariate analysis showed cardiac and neurological comorbidities, and diabetes were significantly associated with ICU admission (Table 4). In addition, bacterial co-infection, CXR abnormality, and encephalopathy were also strong predictors of need for ICU admission. Vaccination status was not incorporated into the multivariate model for ICU admission as it was not available for all participants and was not significant in univariate analysis as a risk factor in subset analysis (children aged <7 years).

The median LOS for all patients admitted with influenza was 2 days (IQR 1–4) (Table 3). Clinical factors significantly associated with LOS are listed in Table 5. Having a febrile seizure was associated with a shorter LOS. Two children, both aged 8 years and with influenza type A, died during their hospitalization.

DISCUSSION

This study provides extensive detail on the characteristics and outcomes of laboratory-confirmed
Table 3. Outcomes and complications in Australian children hospitalized with laboratory-confirmed influenza during 2011–2013 at the Children’s Hospital at Westmead (NSW, Australia) and Princess Margaret Hospital (WA, Australia) by year

<table>
<thead>
<tr>
<th>Year</th>
<th>2011</th>
<th>2012</th>
<th>2013</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>Total (n)</td>
<td>199</td>
<td>312</td>
<td>229</td>
<td>740</td>
</tr>
<tr>
<td>Outcomes</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Median length of stay, days (IQR)</td>
<td>3 (2–6)</td>
<td>2 (1–4)</td>
<td>2 (1·5–4)</td>
<td>2 (1–4)</td>
</tr>
<tr>
<td>ICU admission</td>
<td>15 (7·5)</td>
<td>27 (8·7)</td>
<td>21 (9·2)</td>
<td>63 (8·5)</td>
</tr>
<tr>
<td>Median ICU length of stay, days (IQR)</td>
<td>3 (2–4)</td>
<td>2 (1–4)</td>
<td>4 (2–7)</td>
<td>3 (2–6)</td>
</tr>
<tr>
<td>Antibiotics prescribed</td>
<td>147 (73·9)</td>
<td>221 (70·8)</td>
<td>167 (72·9)</td>
<td>535 (72·3)</td>
</tr>
<tr>
<td>Antivirals prescribed</td>
<td>54 (27·1)*</td>
<td>58 (18·6)</td>
<td>40 (17·5)*</td>
<td>152 (20·5)</td>
</tr>
<tr>
<td>Complications</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Bacterial co-infection</td>
<td>27 (13·6)</td>
<td>31 (9·9)</td>
<td>33 (14·4)</td>
<td>91 (12·3)†</td>
</tr>
<tr>
<td>Meningitis</td>
<td>1 (0·5)</td>
<td>0</td>
<td>0</td>
<td>1 (0·1)</td>
</tr>
<tr>
<td>Bacteraemia</td>
<td>5 (2·5)</td>
<td>2 (0·6)</td>
<td>6 (2·6)</td>
<td>13 (1·8)</td>
</tr>
<tr>
<td>Urinary tract infection</td>
<td>4 (2·0)</td>
<td>8 (2·6)</td>
<td>10 (4·4)</td>
<td>22 (3·0)</td>
</tr>
<tr>
<td>Other</td>
<td>19 (9·5)</td>
<td>22 (7·1)</td>
<td>19 (8·3)</td>
<td>60 (8·1)</td>
</tr>
<tr>
<td>Viral co-infection</td>
<td>43 (21·6)</td>
<td>47 (15·1)</td>
<td>40 (17·5)</td>
<td>130 (17·6)</td>
</tr>
<tr>
<td>Pneumonia/pneumonitis</td>
<td>24 (12·1)</td>
<td>40 (12·8)</td>
<td>19 (8·3)</td>
<td>83 (11·2)</td>
</tr>
<tr>
<td>Seizures</td>
<td>20 (10·1)</td>
<td>31 (9·9)</td>
<td>23 (10·0)</td>
<td>74 (10·0)</td>
</tr>
<tr>
<td>Febrile seizures</td>
<td>10</td>
<td>21</td>
<td>13</td>
<td>44 (5·9)</td>
</tr>
<tr>
<td>Afebrile seizures</td>
<td>10</td>
<td>10</td>
<td>10</td>
<td>30 (4·1)</td>
</tr>
<tr>
<td>Encephalopathy/encephalitis</td>
<td>3 (1·5)</td>
<td>5 (1·6)</td>
<td>3 (1·3)</td>
<td>11 (1·5)</td>
</tr>
<tr>
<td>Myositis</td>
<td>0</td>
<td>4 (1·3)</td>
<td>7 (3·1)</td>
<td>11 (1·5)</td>
</tr>
<tr>
<td>Death</td>
<td>1 (0·5)</td>
<td>0</td>
<td>1 (0·4)</td>
<td>2 (0·3)</td>
</tr>
</tbody>
</table>

IQR, Interquartile range; ICU, intensive care unit. Absolute numbers indicated with percentage within stated year in parentheses unless indicated otherwise. *P = 0·02 using χ² test. †Total bacterial co-infection included individuals with ≥1 co-infection and totals may not equal the sum of all categories.

Table 4. Risk factors associated with intensive care unit admission in Australian children hospitalized with laboratory-confirmed influenza during 2011–2013 at the Children’s Hospital at Westmead (NSW, Australia) and Princess Margaret Hospital (WA, Australia)

<table>
<thead>
<tr>
<th>Predictor variable</th>
<th>Unadjusted OR (95% CI)</th>
<th>P value</th>
<th>Adjusted multivariate OR (95% CI)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Respiratory comorbidity</td>
<td>1·96 (0·97–3·94)</td>
<td>0·06</td>
<td>NI</td>
<td>NI</td>
</tr>
<tr>
<td>Cardiac comorbidity</td>
<td>4·19 (2·05–8·55)</td>
<td>&lt;0·001</td>
<td>3·83 (1·74–8·45)</td>
<td>0·001</td>
</tr>
<tr>
<td>Diabetes</td>
<td>4·41 (0·84–23·2)</td>
<td>0·08</td>
<td>7·44 (1·11–49·7)</td>
<td>0·04</td>
</tr>
<tr>
<td>Neurological comorbidity</td>
<td>3·79 (2·12–6·79)</td>
<td>&lt;0·001</td>
<td>2·15 (1·05–4·40)</td>
<td>0·04</td>
</tr>
<tr>
<td>Immunosuppression</td>
<td>0·20 (0·05–0·83)</td>
<td>0·03</td>
<td>NI</td>
<td>NI</td>
</tr>
<tr>
<td>Genetic comorbidity</td>
<td>2·52 (1·17–5·45)</td>
<td>0·19</td>
<td>NI</td>
<td>NI</td>
</tr>
<tr>
<td>Bacterial co-infection</td>
<td>3·31 (1·82–6·02)</td>
<td>&lt;0·001</td>
<td>2·41 (1·23–4·73)</td>
<td>0·01</td>
</tr>
<tr>
<td>Viral co-infection</td>
<td>1·66 (0·91–3·04)</td>
<td>0·10</td>
<td>NI</td>
<td>NI</td>
</tr>
<tr>
<td>Afebrile seizure</td>
<td>7·32 (3·31–7·33)</td>
<td>&lt;0·001</td>
<td>3·71 (1·34–10·3)</td>
<td>0·01</td>
</tr>
<tr>
<td>CXR abnormality</td>
<td>3·53 (2·05–6·06)</td>
<td>&lt;0·001</td>
<td>3·75 (2·06–6·83)</td>
<td>&lt;0·001</td>
</tr>
<tr>
<td>Encephalitis/encephalopathy</td>
<td>9·64 (2·85–32·5)</td>
<td>&lt;0·001</td>
<td>10·78 (2·43–47·7)</td>
<td>0·002</td>
</tr>
</tbody>
</table>

OR, Odds ratio; CI, confidence interval; NI, variable not included in final model. Unadjusted odds ratios calculated from univariate analysis of predictors, with variables with P < 0·20 listed and incorporated into a backwards stepwise multivariate logistic regression model.
Table 5. Risk factors associated with length of stay in Australian children hospitalized with laboratory-confirmed influenza during 2011–2013 at the Children’s Hospital at Westmead (NSW, Australia) and Princess Margaret Hospital (WA, Australia) excluding nosocomial cases with onset of symptoms ≥7 days after admission

<table>
<thead>
<tr>
<th>Predictor variable</th>
<th>Unadjusted OR of means (95% CI)</th>
<th>P value</th>
<th>Adjusted OR of means (95% CI)</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Female sex</td>
<td>1.27 (1.08–1.49)</td>
<td>0.004</td>
<td>1.22 (1.02–1.44)</td>
<td>0.03</td>
</tr>
<tr>
<td>CXR abnormality</td>
<td>1.65 (1.40–1.95)</td>
<td>&lt;0.001</td>
<td>1.50 (1.25–1.79)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Bacterial co-infection</td>
<td>2.31 (1.83–2.92)</td>
<td>&lt;0.001</td>
<td>2.46 (1.93–3.14)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Viral co-infection</td>
<td>1.64 (1.34–2.91)</td>
<td>&lt;0.001</td>
<td>1.72 (1.38–2.15)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Febrile seizures</td>
<td>0.48 (0.33–0.68)</td>
<td>&lt;0.001</td>
<td>0.62 (0.42–0.89)</td>
<td>0.01</td>
</tr>
<tr>
<td>Afebrile seizure</td>
<td>1.64 (1.10–2.44)</td>
<td>0.02</td>
<td>NI</td>
<td>NI</td>
</tr>
<tr>
<td>Encephalitis/encephalopathy</td>
<td>1.56 (0.80–3.03)</td>
<td>0.19</td>
<td>2.47 (1.24–4.90)</td>
<td>0.01</td>
</tr>
<tr>
<td>Respiratory comorbidity</td>
<td>1.58 (1.22–2.03)</td>
<td>&lt;0.001</td>
<td>1.40 (1.06–1.84)</td>
<td>0.02</td>
</tr>
<tr>
<td>Prematurity</td>
<td>1.54 (1.17–2.02)</td>
<td>0.002</td>
<td>1.45 (1.08–1.94)</td>
<td>0.01</td>
</tr>
<tr>
<td>Cardiac comorbidity</td>
<td>1.38 (1.01–1.89)</td>
<td>0.05</td>
<td>NI</td>
<td>NI</td>
</tr>
<tr>
<td>Diabetes</td>
<td>1.75 (0.80–3.85)</td>
<td>0.16</td>
<td>2.22 (1.00–4.96)</td>
<td>0.05</td>
</tr>
<tr>
<td>Renal comorbidity</td>
<td>1.60 (1.04–2.47)</td>
<td>0.03</td>
<td>2.06 (1.20–2.98)</td>
<td>0.006</td>
</tr>
<tr>
<td>Hepatic comorbidity</td>
<td>2.91 (2.39–6.42)</td>
<td>&lt;0.001</td>
<td>5.60 (3.57–9.31)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Genetic comorbidity</td>
<td>1.63 (1.20–2.23)</td>
<td>0.002</td>
<td>1.45 (1.04–2.03)</td>
<td>0.03</td>
</tr>
<tr>
<td>Metabolic comorbidity</td>
<td>1.48 (0.84–2.60)</td>
<td>0.17</td>
<td>NI</td>
<td>NI</td>
</tr>
</tbody>
</table>

OR, Odds ratio; CI, confidence interval; NI, variable not included in final model after backwards stepwise elimination. Generalized linear regression model with negative binomial distribution. Predictor variables with \( P < 0.20 \) on univariate analysis listed and incorporated into a final multivariate model.

hospitalized influenza in Australian children over a 3-year period in two geographically distinct areas. While the relative over-representation of young children aged <5 years is consistent with administrative data [1, 3], this study provides new insight into the clinical characteristics of children requiring hospitalization. Although 43% of children had underlying medical conditions, the majority were healthy. Infants aged 0–6 months represented the highest proportion (by 6-month age cohorts) of hospitalized children in our study; this age group is too young for influenza vaccination, but may be afforded protection from maternal vaccination [29, 30].

Children hospitalized with influenza had a rate of complications comparable to adults, including pneumonia and seizures in 1/10, with ICU admission required in 1/12 children, similar to reports from seasonal and pandemic years [22, 31, 32]. Despite the median LOS being shorter than in adults, these findings confirm that influenza should not be regarded as a benign paediatric illness. Neurological complications such as acute encephalitis were present in a minority (\( n = 11, 1.5\% \)) but most cases were in previously healthy children and encephalitis was a strong predictor of ICU admission. Influenza has since been confirmed as a significant preventable cause of childhood encephalitis (10%) in a prospective multi-centre Australian study of acute encephalitis [33]. Other risk factors for ICU admission included cardiac, neurological disease, and diabetes, consistent with systematic reviews which found these factors to be predictors of severe influenza [34, 35].

Our study revealed high rates of potentially unnecessary antibiotic use in hospitalized influenza cases and also the infrequent and progressively declining use of antiviral medications (27% in 2011 to 18% in 2013). Prospectively collected FluCAN data from 2014 has shown this to fall further to 15% [23] and demonstrates the consistent decline in utilization rates from a high of around 50% during the 2009 pandemic [36] It is likely that with increasing time since the 2009 pandemic, physician prescribing may be falling back to pre-pandemic levels, when antiviral prescription was relatively uncommon [37].

The cost to the health system of paediatric influenza hospitalization is likely to be considerable but it is difficult to quantify. A recent analysis used weighted costings for the most frequent Australian Refined Diagnosis Related Group (AR-DRG) codes associated with individual national hospitalization data [38] where influenza was coded as the primary diagnosis, and estimated $3473–3560 cost per admission (2016 AU$) for children aged 0–17 years [39]. Applying this cost estimate per episode to our study
(n = 740), the annual direct cost would be AUS 850,000 per year for the two hospitals alone. The cost burden on the health system is likely to be high, particularly given the under-recognition of influenza as a cause of many hospitalizations [20].

Influenza vaccine uptake was low in all children hospitalized with influenza (5% in those with medical conditions and 0–7% in those without). The higher uptake in PMH (4.3%) compared to CHW (1.6%) suggests the availability and promotion of free vaccine for children aged 6 months to <5 years in the state of WA was a factor in vaccine uptake, albeit still only having a modest impact on coverage. Hospitalization of a child increased vaccination uptake in the season following each child’s hospitalization to 10.8%, possibly due to increased awareness by parents of the risk and morbidity of childhood influenza. While this is promising, there remain unanswered questions as to why this figure remains low and how it could be further improved. Increased recommendation of vaccination and addressing of any parental safety concerns at the time of hospitalization by treating physicians may be strategies worthy of consideration.

While our study was limited to directly examining coverage in a cohort of hospitalized children, in whom rates may be lower due to vaccine effectiveness preventing hospitalization, we also found state and national population-level estimates to be very similar. Importantly, our data provides evidence of the low vaccination uptake in children with comorbidities, information which is not collected by the national ACIR. Our study also informed the establishment of prospective active surveillance in specialist paediatric hospitals (annually since the 2014 season) to complement existing FluCAN surveillance in adult and general centres. This prospective surveillance has recently reported on vaccine coverage and effectiveness in the 2014 season [23], and together with a second longitudinal study in WA [40] confirmed low vaccine coverage in hospitalized children, including influenza test-negligible (‘control’) populations, in whom uptake was only 8.5–12.4% overall and 18% for children with comorbidities. The true national population rate is likely to be lower, particularly considering both studies had recruitment of children in WA. These estimates all point to worryingly low influenza vaccination coverage in at-risk children who have been eligible for free vaccination at point-of-care since 2010.

Our data suggests that improving vaccination coverage should be a focus for reduction of influenza morbidity, particularly in those with underlying medical conditions. In addition, the absence of strong recommendations and national funding for vaccination of healthy children means a significant population within the community are unprotected and contribute to the burden of influenza hospitalization nationally. With the knowledge that hospitalization rates in Australia are highest in those aged <5 years [1, 3], the ongoing poor coverage in WA, despite its partially funded (6 months–<5 years) programme for all children is disappointing, but indicates the sustained impact of vaccine safety concerns after elevated post-vaccination febrile seizure rates were seen from one vaccine brand in 2010. Following this, vaccine uptake in WA fell from approximately 45% (2008 and 2009) to 7% during (2011–2015) despite continued funding, as shown in this and other studies [10]; the proportion of parents who felt influenza vaccine to be safe halved [10]. Universally recommended and funded influenza vaccination, as currently provided in numerous countries, has the potential to increase public confidence in influenza vaccine safety and uptake in all children. Coverage of approximately 60% has been achieved in the United States’ [17] and UK’s universal childhood vaccination programmes [18] and has resulted in reductions for a range of influenza-related outcomes both directly in targeted age groups and indirectly via herd protection in unimmunized age groups [18, 41, 42].

Limitations in this retrospective study were that data collection was restricted to information available from case records. Vaccination status was poorly recorded in clinical notes and it is clear that specific medical history taking about influenza vaccination status should become common practice. We confined our analysis to PCR-confirmed influenza only, for consistency across both sites, which together with probable under-ascertainment of cases due to non-specific presentations and likely incomplete testing, suggests the total influenza burden was underestimated. While respiratory virus testing on hospitalized patients with respiratory illness is routinely recommended for infection control purposes, it was ultimately at the physician’s discretion and we could not be certain that all such patients were tested. A strength of our study, is that we have obtained data on influenza vaccine uptake in at-risk populations, albeit influenza cases, but have related that to state and national estimates, and estimates in hospitalized non-influenza patients.

In summary, this study provides detailed vaccination and clinical data on Australian children.
hospitalized with influenza. It demonstrates that influenza has a significant impact on young children, both healthy and with underlying medical conditions, and can be associated with significant morbidity. There is very low vaccination coverage in this hospitalized cohort, which matches low estimates of coverage at a population level and in non-influenza hospitalized children. Funded immunisation programmes for at-risk children, and in WA for some healthy children, are not achieving good vaccine uptake; this strongly suggests the need to examine additional strategies. Accurate local disease modelling to investigate future options, such as universal free influenza vaccination in children, and other studies examining barriers to vaccine uptake, would be valuable.

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DECLARATION OF INTEREST
None.

REFERENCES


