Communicable Diseases Intelligence

Summary of National Surveillance Data on Vaccine Preventable Diseases in Australia, 2016-2018

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Multi-year report

Summary of National Surveillance Data on Vaccine Preventable Diseases in Australia, 2016-2018 Final Report

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Overview

This summary report on vaccine preventable diseases (VPDs) in Australia brings together the three most important national sources of routinely-collected data on VPDs (notifications, hospitalisations and deaths) for all age groups for the three-year period 1 January 2016 to 31 December 2018. Information about each VPD is provided in 16 chapters using a standard structure.

Changes in the National Immunisation Program (NIP) between 2016 and 2018 are listed below:

1. Addition of a booster dose of diphtheria-tetanus-acellular pertussis vaccine (DTPa) at 18 months (March 2016).

2. Replacement of trivalent influenza vaccine with quadrivalent influenza vaccines (from 2016), and availability of two enhanced trivalent vaccines (high-dose and adjuvanted vaccines), for people aged 65 years and older (in 2018).

3. Addition of a single dose of live attenuated herpes zoster vaccine for people aged 70 years from November 2016, with a five-year catch-up program for 71- to 79-year olds.

4. Replacement of the quadrivalent human papillomavirus (HPV) vaccine (4vHPV) for adolescents with a nonavalent HPV vaccine (9vHPV) in January 2018.

5. Replacement of the 12-month dose of combination *Haemophilus influenzae* type b–meningococcal serogroup C vaccine (Hib-MenC) with a single dose of quadrivalent meningococcal conjugate (MenACWY) vaccine, with monovalent Hib vaccine given at 18 months of age (July 2018).

6. Change in the timing of routine infant doses of 13-valent pneumococcal conjugate vaccine (PCV13) from 2, 4, and 6 months (3+0 schedule) to 2, 4, and 12 months (2+1 schedule), commencing July 2018. The schedule remained as 2, 4, 6, and 12 months (3+1) for Aboriginal and Torres Strait Islander children in the Northern Territory, South Australia, Queensland and Western Australia, and for children with specified underlying medical conditions.

7. Addition of a dose of diphtheria-tetanus-acellular pertussis vaccine (dTpa) for all women in the third trimester of pregnancy, from July 2018 (replacing funding by states and territories).
Notifications, hospitalisations and deaths in this three-year reporting period and the previous four-year reporting period (2016–2018 and 2012–2015) are summarised in Table 1. Influenza, pertussis and rotavirus were the most commonly notified conditions whereas the most common causes of hospitalisation were influenza, zoster, rotavirus and pneumococcal disease. There were no notifications or hospitalisations due to polio and a continuing low incidence of diphtheria, rubella and tetanus.

**Important changes in VPDs in the 2016–2018 reporting period**

Influenza was the most commonly reported VPD, with the incidence of notifications and hospitalisations during this period approximately twice that in the previous period. The increase in influenza notifications and hospitalisations was driven by a high-activity influenza season in 2017 that was the largest season since the 2009 pandemic year, although increased laboratory testing, especially greater use of influenza polymerase chain reaction (PCR) tests, also contributed. The highest numbers and rates of influenza-related deaths were reported in those aged ≥ 65 years.

Pertussis was the second most commonly notified VPD, although declines in both notification and hospitalisation annual rates were observed compared with the previous four-year period (2012–2015). Notification rates were highest in the 5–14 year age group, while hospitalisation rates were highest among infants aged < 1 year.

Diphtheria notifications increased from seven in the previous four-year period to 27 in the three years 2016–2018, with the highest number (14) and incidence among persons older than 50 years. The reasons for this increase in notifications should be explored further. The only notified diphtheria death occurred in an unvaccinated male 25–49 years of age; and, of 20 cases with known vaccination status, only two were fully vaccinated.

Meningococcal disease notifications and hospitalisations increased substantially in this reporting period, due to the emergence of serogroups W and Y. The accelerating increase in meningococcal disease notifications between 2015 and 2017 prompted several jurisdictionally-funded meningococcal (MenACWY) vaccination programs, replaced by a nationally-funded vaccination program in 2018.

Acute hepatitis B notifications progressively declined over this reporting period; a trend evident since 2007, but which has accelerated. While hepatitis A infections among Aboriginal and Torres Strait Islander people continued to decline, an increase in overall notifications of hepatitis A was observed due to a number of outbreaks, most notably among men who have sex with men.

Notifications and hospitalisations due to pneumococcal disease increased in this reporting period. The highest numbers of pneumococcal notifications, hospitalisations and deaths occurred in older adults aged ≥ 65 years. Rates of pneumococcal notifications and deaths were also highest in adults aged ≥ 65 years; however, the highest rates of pneumococcal hospitalisations occurred among infants aged < 1 year. Aboriginal and Torres Strait Islanders continue to experience substantially higher rates of pneumococcal disease relative to the rest of the population.

Small overall reductions in *Haemophilus influenzae* type b disease notifications were seen in this reporting period, following on from larger earlier reductions.

Reductions in the overall average annual rate of hospitalisations due to rotavirus were observed during this reporting period, even though a spike was observed in 2017 due to disease outbreaks.
<table>
<thead>
<tr>
<th>Disease</th>
<th>Notifications</th>
<th>Hospitalisations Principal diagnosis</th>
<th>Deaths Underlying cause</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Average annual rate&lt;sup&gt;a&lt;/sup&gt;</td>
<td>Average annual rate&lt;sup&gt;b&lt;/sup&gt;</td>
<td>Rate ratio (95% CI)&lt;sup&gt;c&lt;/sup&gt;</td>
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<tr>
<td>Hepatitis B</td>
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<tr>
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<tr>
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<tr>
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<td>na&lt;sup&gt;f&lt;/sup&gt;</td>
<td>–</td>
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<tr>
<td>Zoster</td>
<td>na&lt;sup&gt;f&lt;/sup&gt;</td>
<td>na&lt;sup&gt;f&lt;/sup&gt;</td>
<td>–</td>
</tr>
</tbody>
</table>

**Notes:**

- Data from 2012–2015 are in the previous report.<sup>l</sup>
- Rate per 100,000 population.
- Rate ratio = (average annual rate in 2016–2018) / (average annual rate in 2012–2015). Please note the following when interpreting rate ratios:
  - Rate ratio > 1: the average annual rate ratio was higher in the current reporting period (2016–2018) than in the previous reporting period (2012–2015)
  - Rate ratio < 1: the average annual rate ratio was lower in the current reporting period than in the previous reporting period
  - Rate ratio = 1: the average annual rate ratio was the same in the current reporting period as in the previous reporting period
  - A rate ratio with a 95% confidence interval (95% CI) that overlaps with 1 is considered not statistically significant.
- Minimum estimates as notifications, hospitalisations and deaths grossly underestimate influenza-related cases.
- Pneumococcal hospitalisations and deaths septicemia and meningitis only.
- na: not applicable, because data are unavailable for one or both periods.
- Rotavirus notifications for the period 2016–2018 are reported for notifications where the month of diagnosis was between 1 July 2018 and 31 December 2018 only.

<sup>a</sup> Data from 2012–2015 are in the previous report.<sup>l</sup>
<sup>b</sup> Rate per 100,000 population.
<sup>c</sup> Rate ratio = (average annual rate in 2016–2018) / (average annual rate in 2012–2015). Please note the following when interpreting rate ratios:
  - Rate ratio > 1: the average annual rate ratio was higher in the current reporting period (2016–2018) than in the previous reporting period (2012–2015)
  - Rate ratio < 1: the average annual rate ratio was lower in the current reporting period than in the previous reporting period
  - Rate ratio = 1: the average annual rate ratio was the same in the current reporting period as in the previous reporting period
  - A rate ratio with a 95% confidence interval (95% CI) that overlaps with 1 is considered not statistically significant.
<sup>d</sup> Minimum estimates as notifications, hospitalisations and deaths grossly underestimate influenza-related cases.
<sup>e</sup> Pneumococcal hospitalisations and deaths septicemia and meningitis only.
<sup>f</sup> na: not applicable, because data are unavailable for one or both periods.
<sup>g</sup> Rotavirus notifications for the period 2016–2018 are reported for notifications where the month of diagnosis was between 1 July 2018 and 31 December 2018 only.
The rate of hospitalisations coded as varicella (chickenpox) in this reporting period was similar to that in the previous four years. Following the introduction of the national zoster vaccination program in November 2016, the increasing trend in zoster hospitalisation rates stabilised for the first time since 1993.

Measles notifications and hospitalisations continued to remain low, but outbreaks linked to imported cases in travellers from high endemicity regions occurred sporadically, with 73% of cases reported to be unvaccinated. The available evidence supports continued elimination of endemic measles, first documented in 2005, with verification by the World Health Organization (WHO) in 2014.3,4

Mumps occurred in localised outbreaks, commencing in central Australia in 2015 and seen through most of the reporting period before a decline in notifications in 2018. Cases were predominantly two-dose vaccinated adolescents and young adults in Aboriginal communities, as previously documented in a 2007–2008 outbreak in the Kimberley, and also seen in close contact settings in other developed countries.6,7 This was in contrast to previous mumps outbreaks in adolescents and young adults in Australia, which had primarily been in persons who had received only one vaccine dose or were unvaccinated.5–9

Rubella notifications and hospitalisations continued to decline in this reporting period and have remained consistently low since 2004, following marked declines in the late 1990s and early 2000s. In 2018, the WHO officially declared that Australia had eliminated rubella.10,11

Q fever notifications and hospitalisations remained stable in 2016–2018 compared with 2012–2015, with rates remaining highest in males aged 50–64 years.

Limitations

The data sources used in this report have a number of limitations, discussed in detail in the body of the report. The datasets for notifications, hospitalisations and deaths differ in their purpose for data collection, reporting mechanisms and accuracy, so comparisons between them should be made with caution. We found that, for some diseases, the number of deaths recorded in the National Notifiable Diseases Surveillance System (NNDSS) differ substantially from cause of death data from the Australian Coordinating Registry and the AIHW National Hospital Morbidity Database. Notification data on vaccination status (vaccine type and vaccination date) in the NNDSS had low levels of completeness for age groups not targeted for vaccination programs. Interpretation of hospitalisation data derived from codes in any diagnosis field and of death data derived from all coded associated causes of death should take into account the lack of specificity compared with principal diagnosis and underlying cause of death. Interpretation of trends in notification rates over time needs to take account of changing patterns of diagnostic testing, particularly for influenza and pertussis, and changes in case definitions.

1. Introduction

The National Centre for Immunisation Research and Surveillance (NCIRS) has published six comprehensive reports on vaccine preventable diseases (VPDs) and vaccination coverage in Australia since 2000. These reports (the ‘VPD reports’) have been published as supplements in Communicable Diseases Intelligence. They serve as a national resource to support surveillance and control of VPDs in Australia, particularly VPDs targeted by the NIP.

In addition to these summary VPD reports, which use a standard structure across all VPDs, a rolling series of epidemiologic reviews provides more detailed data for individual VPDs. Epidemiologic reviews including data relevant to this report include Haemophilus influenzae type b (Hib) and pertussis.13
2. Methods

In keeping with previous practice, three main sources of routinely collected data on VPDs in Australia were used for this report. Disease notification data were obtained from the Office of Health Protection's National Notifiable Diseases Surveillance System (NNDSS), supplied by states and territories; data on coded hospitalisations were sourced from the Australian Institute of Health and Welfare (AIHW) National Hospital Morbidity Database; and causes of death unit record file data were obtained from the Australian Coordinating Registry (ACR).

A comprehensive listing of significant events in vaccination practice in Australia is available from the National Centre for Immunisation Research and Surveillance.\(^1\)

Notifications

The NNDSS was established in its current form in 1991 and includes de-identified information on cases of notifiable disease reported by state and territory authorities in Australia. Each of the eight state and territory health departments collects data on notifiable diseases under its respective public health legislation. Data quality of the NNDSS is continually monitored by the Office of Health Protection within the Australian Government Department of Health, and by the National Surveillance Committee, a committee which includes jurisdictional surveillance and data managers. Historically, state and territory notification criteria were based on the 1994 National Health and Medical Research Council surveillance case definitions.\(^15\) In September 2003, a new set of national case definitions for notifiable diseases reported to the NNDS was endorsed by the Communicable Diseases Network Australia,\(^16\) with all jurisdictions implementing in 2004.\(^17\)

Information on case definitions currently in use for vaccine preventable diseases is available on the Australian Government Department of Health website.\(^i\) For this report, data for a ten-year period from 1 January 2009 to 31 December 2018 were extracted from the NNDSS in November 2019, with the exception of Q fever for which data were extracted in December 2020 for the period from 1 January 2010 to 31 December 2019. Data were checked and cleaned. Disease notification data for cases with a date of diagnosis between 1 January 2016 and 31 December 2018 are included in this report, with the exception of rotavirus (cases reported for 1 July to 31 December 2018) which became nationally notifiable in July 2018. It should be noted that historical notification data (i.e. data for the period 2009 to 2015 for all diseases except Q fever, which was updated for the period 2010 to 2015) included in this report have been updated from previous reports and the updated data used for trend analysis. For data prior to 2009 (2010 for Q fever), existing published data were used for trend analysis. The variables extracted for each VPD in this report were: date of diagnosis, age at onset, sex, Aboriginal and Torres Strait Islander status and state or territory of residence. Date of diagnosis is derived from the date of onset, or, where not supplied, the earliest date recorded among the three fields of date of specimen, date of notification, or date notification received (the only mandatory date field).

Aboriginal and Torres Strait Islander status in the notification data provided by the Department of Health to NCIRS includes three categories: ‘Aboriginal and/or Torres Strait Islander origin’, ‘not Aboriginal or Torres Strait Islander origin’ and ‘not stated’. For the purposes of calculating rates, we used two categories: ‘Aboriginal and Torres Strait Islander’ (individuals identified as Aboriginal and/or Torres Strait Islander) and ‘other’ (individuals recorded as not Aboriginal or Torres Strait Islander and where not stated).

Completeness of Aboriginal and Torres Strait Islander status was assessed for each disease by calculating the proportions of notifications


where Aboriginal and Torres Strait Islander status was ‘not stated’ and/or where the data field was left blank. Incomplete identification may result in the rates reported underestimating the true incidence of disease in Aboriginal and Torres Strait Islander people. Notification data are not reported for influenza due to poor overall completeness of Aboriginal and Torres Strait Islander status, and for pertussis only reported for the < 5 years age group due to poor completeness in other age groups, as per previous practice.18

Vaccination status was assessed using the new vaccine type and vaccination date data fields, which included information on the vaccine type (or brand), date of vaccination and age at time of vaccination for the five most recent vaccine doses received. In jurisdictions where new vaccination data fields were not used for the specified years, we used the old vaccination data fields, which included data on vaccination status (i.e. fully vaccinated, partially vaccinated, unvaccinated or unknown/missing data), whether previous doses were validated using official records or patient/parent recall only, how many previous vaccine doses were received, and the date of last vaccination. Data provided via the new vaccination data fields allows for collection of data for up to five previous vaccinations relevant to the disease; data on additional vaccine doses received (where six or more previous doses of a vaccine were received) was not available for analysis. We also used a cut-off of 14 days between vaccination date and disease onset date for estimating validity of vaccine doses. Vaccine doses given < 14 days prior to disease onset date or given after the disease onset date were considered invalid. We provide details of only valid vaccine doses in this report. Data provided via the old vaccination data fields only reported the date of the most recent vaccination; therefore assessment of the validity of additional previous doses of vaccine (where two or more previous doses were received) was not possible. Where information on the date of vaccination was missing for a case who had received a vaccine dose, the dose was assumed to be valid (i.e. received 14 days or more prior to disease onset date).

Hospitalisations

Hospitalisation data from the AIHW National Hospital Morbidity Database were analysed by calendar year of hospital admission for the three years 2016–2018. Historical data on hospitalisations since July 2001 were updated for trend analysis. For data prior to July 2001, the report uses selected historical data from previous reports.12 Hospitalisations where the date of admission fell within the reporting period were included, with analysis by variables such as age and sex grouped by the calendar year of hospital admission. Data for each VPD were extracted based on the International Statistical Classification of Diseases and Related Health Problems, 10th Revision, Australian Modification (ICD-10-AM). Records of hospitalisations included in analysis were those with the code(s) of interest listed as the principal diagnosis (the diagnosis recorded as chiefly responsible for the hospitalisation) or as any other diagnosis for that episode of hospitalisation. The proportion of hospitalisations where the disease of interest was coded as the principal diagnosis is reported for each disease. For hepatitis B, only hospitalisations with acute hepatitis B coded as the principal diagnosis are included, consistent with the approach taken in previous reports.

The variables extracted for analysis included: month of admission, age on admission, sex, Aboriginal and Torres Strait Islander status, state or territory of residence, length of stay, and diagnosis (principal and other diagnoses) coded using the relevant edition of ICD-10-AM for the collection period.

Aboriginal and Torres Strait Islander status in the hospitalisation data provided by AIHW to NCIRS includes the two categories ‘Aboriginal and Torres Strait Islander Australians’ and ‘Other Australians’. ‘Other Australians’ includes those with a status of ‘not stated’. Completeness
and accuracy of Aboriginal and Torres Strait Islander status in hospitalisation data is high, with an estimated 88% of Aboriginal and Torres Strait Islander patients correctly identified in public hospital admission records in 2011–2012. Incomplete identification may result in the rates reported underestimating the true incidence of disease in Aboriginal and Torres Strait Islander people.

Deaths

Death data were obtained from the Cause of Death Unit Record File (COD URF) data from the Australian Coordinating Registry (ACR). The Queensland Registry of Births, Deaths and Marriages is the ACR for COD URF data. The ACR coordinates the approval and release of COD URF files on behalf of the data custodians – Australian Registrars of Births, Deaths and Marriages, State/Chief Coroners and the National Coronial Information System (NCIS).

Since 1997, ICD-10 has been used to identify the cause of death. Deaths analysed in this report included those recorded as occurring in the calendar years 2016–2018. The variables included were cause of death, age, year of death, sex, and state or territory in which the death was recorded. Both underlying and associated causes of death were analysed for this reporting period. Deaths recorded in the NNDSS are also reported in relevant disease chapters.

Calculations

All rates were calculated using the mid-year estimated resident populations released by the Australian Bureau of Statistics (ABS) as the population denominator and hospital admissions (data from AIHW)/ notifications (data from NNDSS) /deaths (from ACR) as the numerator. Rates are presented as annual rates or average annual rates per 100,000 total population, or population in age, sex or geographical subgroups, as appropriate. The reported rate estimates for the populations were not stratified by age groups (i.e. all ages together) and are crude rates that have not been age-standardised. For notification, hospitalisation and death data, the mid-year population estimates for the corresponding calendar year were used as the denominator population. Averages were calculated for rates of notifications and hospitalisations and for bed-days of hospitalisation episodes per year. Rate ratios were calculated for average annual rates in this reporting period (2016–2018) compared with the previous period (2012–2015). The median (rather than average) and range were used to describe the distribution of notifications and hospitalisations per month, and the length of stay per hospitalisation episode, as these data are not normally distributed.

Current case definitions and significant events in vaccination practice in Australia were taken into consideration for this report. Documentation of changes in case definitions is no longer readily available and hence could not be considered in this report.

Notes on interpreting data

Comparison between the notification, hospitalisation and death data should be made with caution since these datasets differ in their purpose for data collection, reporting mechanisms and accuracy. The rates presented in this report are crude rates, not adjusted for differences in population structure between jurisdictions.

Rate ratios with wide confidence intervals should be interpreted with caution, as they indicate very small numbers of cases with underlying rates typically less than 0.1 per 100,000 population. These rate ratios are subject to wide fluctuations with very small changes in the numbers of cases.

Notification data

A major limitation of notification data is that they represent only a proportion of all the cases occurring in the community, due to under-reporting and/or testing practices. This proportion may vary between diseases, over time, and across jurisdictions. An infectious disease diagnosed by a laboratory test should be notified by
the testing laboratory or treating clinician, as required by jurisdictional legislation, whereas diagnoses without laboratory confirmation rely on notification by clinicians, known to be substantially less complete. Changes in screening programs—including the preferential testing of high-risk populations; the availability and use of less invasive and more sensitive diagnostic tests; and periodic awareness campaigns—may influence the number of notifications over time. Data accuracy may also vary among jurisdictions due to the use of different case definitions for surveillance prior to adoption of the national case definitions in 2004 (a consideration that is particularly relevant to trend graphs which present pre-2004 data) and varying reporting requirements and clinician practice with respect to laboratory testing.

Aboriginal and Torres Strait Islander status completeness in notification data was assessed for the period 2016–2018 and reported in each chapter.

Interpretation of data on vaccination status is limited due to missing data on a substantial proportion of notifications, although the quality of data is improving compared with previous reports. The Vaccine Preventable Diseases Surveillance Program Project Agreement, finalised in late 2017, supports states and territories to improve reporting of vaccination information for children aged less than 10 years, specifically to report these data for at least 95% of notified cases in this age group. In many cases, vaccination history was self-reported and not validated, and should be interpreted with caution. As we assumed vaccine doses received were valid for notifications who had received a vaccine but for whom the date of vaccination was missing, some doses included may have been invalid (i.e. received < 14 days prior to the disease onset date), resulting in an overestimation of the proportion of notifications previously vaccinated.

Hospitalisation data

The AIHW publishes regular overviews of Australian hospitalisation statistics, including details of the number of hospitals reporting and any documented data problems. The AIHW performs logical validations on the ICD-10-AM coded data; for example, for sex- and age-specific diagnoses. Coding audits and coding quality improvement activities are variously performed at hospital level and/or at state or territory level. Some variation in hospital access, admission practices, and record coding may occur between regions and over time and this may impact upon the use of hospitalisation data for monitoring disease trends over time and between jurisdictions.

There are also limitations associated with the use of ICD-10-AM codes to identify cases. Errors that cause the ICD-10-AM code to differ from the true disease may be either random or systematic measurement errors. For rare diseases, such as acute poliomyelitis, tetanus and diphtheria, hospitalisation episodes or deaths so coded are much more likely to be miscoding of other conditions. The ICD-10-AM codes for diagnosis chosen for analysis of a disease should accurately reflect the specific condition which is vaccine preventable. For some diseases, such as *Haemophilus influenzae* type b (Hib) infection, both the previously used ICD-9-CM and current ICD-10-AM codes lack specificity for serotype b, and may not be limited to sterile site isolates, in contrast to more stringent case definitions used for notification data. For each disease in this report, the ICD-10-AM code(s) selected to best represent hospitalisation for the disease of interest are listed on the first page of each disease chapter.

In the AIHW hospitalisation database, there is one record for each hospital admission episode. This means that there will be separate records for each re-admission or inter-hospital transfer. This is unlikely to have a major impact on the numbers reported for most of the diseases reviewed in this report, as they are mostly acute diseases, but severe cases in regional and remote areas may be transferred for specialist care. However, there may be greater impact on numbers reported for diseases where re-admission due to long-term complications is more likely
e.g. meningococcal disease and tetanus. It is also more difficult to gauge the relevance where the coded disease was not the principal diagnosis but was recorded as an additional or secondary diagnosis for that hospitalisation episode.

AIHW has restrictions on release of data that may potentially identify cases. Reporting of fewer than five hospitalisations in the period of interest is expressed as a range, 1–4, rather than as the actual number. The exact date of admission was not provided by AIHW to NCIRS (month of admission was provided instead). Length of stay was provided as '30 days' for hospitalised cases where the length of stay was 30 days or longer.

Aboriginal and Torres Strait Islander status in the hospitalisation data provided by AIHW to NCIRS includes the categories ‘Aboriginal and Torres Strait Islander Australians’ and ‘Other Australians’. As the ‘Other Australians’ category includes both non-Aboriginal and Torres Strait Islander people and those with unknown Aboriginal and Torres Strait Islander status, it is not possible to report the exact proportion with ‘unknown’ Aboriginal and Torres Strait Islander status. However, Aboriginal and Torres Strait Islander status completeness for hospitalisation data has been greater than 80% in all jurisdictions since 2010–2011.\(^{19}\)

Death data

Mortality data are analysed by year of death and cause of death. The ACR has restrictions, which apply to this report, on release of data that may potentially identify cases. Where there are fewer than six (but not zero) deaths in the period of interest, this is reported as a range from 1–5 rather than the actual number. In addition, death data are not provided for some age groups, where back-calculations could compromise confidentiality. Many of the issues for accuracy of ICD coding in hospital separations, such as propensity for laboratory testing, are also relevant for mortality data.

Aboriginal and Torres Strait Islander status in the death data provided by the ACR has a high degree of completeness (> 99%) for all deaths, but there are concerns regarding the accuracy of the data, with the number of deaths among Aboriginal and Torres Strait Islanders possibly underestimated.\(^{25}\) Efforts have been made to improve ascertainment of Aboriginal and Torres Strait Islander status. Year-to-year comparisons should be interpreted with caution.
3. Vaccine preventable diseases

3.1 Diphtheria

Highlights

In the three years 2016–2018, there were 27 notifications of toxigenic diphtheria (0.04 per 100,000 population per year), compared with 7 in the four years 2012–2015 (0.01 per 100,000 population per year).

Diphtheria is caused by toxigenic strains of the bacteria Corynebacterium diphtheriae and Corynebacterium ulcerans. Corynebacterium diphtheriae is transmitted from person to person by intimate respiratory and direct contact. Corynebacterium ulcerans, a bacterium found in cattle and more recently in cats, can also express diphtheria toxin and can cause zoonotic infection in humans. Infection can present as respiratory or cutaneous diphtheria, with cutaneous diphtheria being more common in warmer climates and in settings with poor hygiene and overcrowding. Infection typically remains localised to the throat or skin, but disease manifestations arise from both local inflammation and/or systemic toxemia. Pharyngeal diphtheria presents with a membranous inflammation of the upper respiratory tract, which can be extensive and cause laryngeal obstruction. Damage to other organs including the myocardium, nervous system and kidneys, caused by the organism’s exotoxin, may complicate pharyngeal or cutaneous diphtheria. Non-toxigenic C. diphtheriae usually causes mild throat or skin infection, which is occasionally complicated by invasive disease including endocarditis or arthritis.

Case definition

Notifications

Diphtheria case definition:


Note that this case definition was updated in January 2017.

Hospitalisations and deaths

The ICD-10-AM/ICD-10 code A36 (Diphtheria) was used to identify hospitalisations and deaths.

Secular trends

There were 27 notified cases of toxigenic diphtheria during 2016–2018 (Table 3.1.1): 8 in 2016 (3 cutaneous, 1 pharyngeal, 4 unknown); 8 in 2017 (7 cutaneous, 1 pharyngeal); and 11 in 2018 (9 cutaneous, 2 pharyngeal), a median of 1 case (range 0 – 3) notified per month. This represents an average annual notification rate of 0.04 per 100,000 population, compared with 0.01 per 100,000 population per year during 2012–2015 (7 notified cases). The majority of the 2016–2018 cases were cutaneous (n = 19; 70%), with 4/27 cases (15%) pharyngeal. Twenty-three cases (85%) were due to infection with C. diphtheriae, with four (15%) due to C. ulcerans. The majority of cases (17/27; 63%) were acquired overseas, all from South East Asian or Pacific Island countries, with 9/27 (33%) reported as acquired in Australia. Prior to 2011, there had been no notified cases since 2002. However, even this recent increase corresponds to very low notification rates compared with historical data (Appendix A, Figure A.1).

In contrast to notifications, the number of hospitalisations (any diagnosis) in the current three-year reporting period 2016–2018 was lower than the previous four-year period 2012–2015 (38
versus 61 admissions; average annual rate 0.05 vs 0.07 per 100,000 population) with 13, 15 and 10 hospitalisations recorded in 2016, 2017 and 2018, a median of 1 (range 0–4) per month.

Severe morbidity and mortality

Of the 38 hospital admissions coded as due to diphtheria from 1 January 2016 to 31 December 2018, 23 (59%) were cutaneous and 4 were pharyngeal diphtheria (10%). Diphtheria was the principal diagnosis in 7 (18%) of the hospitalisations, 4 cutaneous and 1 pharyngeal. A total of 264 bed-days were recorded between 2016 and 2018, with a median length of stay of five days, highest in patients aged 65 years and above (12.5 days) and lowest in those aged 15–24 years (1.5 days).

In the causes of death data, there were 1–5 deaths in this reporting period with diphtheria recorded as the underlying or associated cause. One death attributable to diphtheria was recorded in the NNDSS, in a young (aged 25–49 years) unvaccinated male.

Age and sex distribution

There were more notifications and hospitalisations for diphtheria in males than in females, with an average male:female rate ratio of 1.7:1 for notifications and 1.3:1 for hospitalisations. Between 2016 and 2018, there were no notifications of diphtheria in cases aged younger than 15 years, although six hospital admissions in patients aged < 15 years were recorded (16%). Slightly more than half of the notified cases were aged 50 years or older (n = 14/27; 52%). Half of the hospitalisations, also, were recorded among persons 50 years or older (19/38; 50%), with approximately one-quarter among adults aged 25 to 49 years (n = 9/38; 24%).

Geographical distribution

Two-thirds (n = 18/27) of diphtheria notifications between 1 January 2016 and 31 December 2018 occurred in Queensland, where 16 hospitalisations also occurred (42%). Five hospitalisations (71%) where diphtheria was the principal diagnosis occurred in Queensland, with the remaining two (29%) in Victoria. No notifications but three hospitalisations (8%) were recorded in the Northern Territory (NT), in contrast to the four years between 1 January 2012 and 31 December 2015 when 28% of all hospitalisations occurred in the NT and the NT rate of hospitalisations was more than four times as high (1.7 per 100,000 population per year versus 0.41 per 100,000 population per year in the current reporting period). Despite this decrease, the highest rate of diphtheria hospitalisations in this reporting period was still in the Northern Territory.

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for 25 of the 27 notifications (93%) of diphtheria during 2016–2018, with two not stated (7%). Aboriginal and Torres Strait Islander people accounted for 11% (3/27) of notifications and 13% (5/38) of hospitalisations for diphtheria, representing higher rates than those seen in other Australians, with notification rates of 0.12 vs 0.03 per 100,000 population per year, respectively, and a rate ratio of 3.7 over the 3-year period. Of the three notifications among Aboriginal and Torres Strait Islanders, one was cutaneous, one was pharyngeal, and one was unknown.

Vaccination status

Of the 27 notifications, vaccination data were available for 74% (20/27) of cases. Of the seven cases where vaccination data were not available, two were recorded as having been followed up but with no information available, and five as not followed up. Of the 20 cases that had vaccination data available, 15 were previously vaccinated; five were unvaccinated. Of the 15 vaccinated cases, ten cases had received only 1 dose and three had received 2 doses, with one case receiving 4 and one receiving 5 doses of diphtheria-containing vaccine. Among vaccinated cases, 50% (7/14) had received their most recent dose within five years of disease onset,
Table 3.1.1: Diphtheria notifications, hospitalisations and deaths, Australia, 2016 to 2018,\textsuperscript{a} by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Diphtheria notifications</th>
<th>Diphtheria hospitalisations</th>
<th>LOS\textsuperscript{b} per admission</th>
<th>Diphtheria deaths\textsuperscript{c}</th>
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<tr>
<td></td>
<td>n</td>
<td>Rate\textsuperscript{d}</td>
<td>Any diagnosis</td>
<td>Principal diagnosis</td>
</tr>
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<td></td>
<td></td>
<td></td>
<td>n</td>
<td>Rate\textsuperscript{d}</td>
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<td></td>
<td>n\textsuperscript{*}</td>
<td>Rate\textsuperscript{d,e}</td>
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<td>All ages</td>
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<td>1–5</td>
<td>0.00</td>
</tr>
</tbody>
</table>

\textsuperscript{a} Notifications where the month of diagnosis was between January 2016 and December 2018; hospitalisations where the month of admission was between January 2016 and December 2018.

\textsuperscript{b} LOS: length of stay in hospital.

\textsuperscript{c} Deaths include underlying and associated causes of deaths.

\textsuperscript{d} Average annual age-specific rate per 100,000 population.

\textsuperscript{e} Principal diagnosis (hospitalisations).

43% (6/14) within 5–10 years prior to disease onset and 7% (1/14) more than 20 years prior to onset. The notified case who died was recorded as unvaccinated.

Comment

Although diphtheria remains a rare disease in Australia, the fourfold increase in notifications of confirmed toxigenic disease in the 2016–2018 period, compared to the 2012–2015 period (27 vs 7), after no cases were notified between 2002 and 2011, was notable. While diphtheria hospital admissions were lower in 2016–2018 than in 2012–2015, hospitalisation data lack specificity and are unreliable for understanding the burden of diphtheria in Australia. Population-based serosurveys in Australia have found low levels of immunity in middle-aged and older adults.\textsuperscript{37} The continued occurrence of cases, particularly with respect to travel-associated exposure in adults, and fatalities caused by this preventable disease, with one known to occur in an unvaccinated individual, is cause for vigilance in ensuring high vaccination rates in all age groups. Travellers may benefit from booster doses of vaccine prior to travel, particularly given recent outbreaks in the Asia-Pacific region.\textsuperscript{26,38–41}
3.2 *Haemophilus influenzae* type b disease

**Highlights**

Notifications for invasive *Haemophilus influenzae* type b (Hib) disease remained low for the period 1 January 2016 to 31 December 2018. Infants aged < 1 year accounted for 27% and children aged 1–5 years for 20% of invasive Hib notifications for the reporting period.

*Haemophilus influenzae* is a gram-negative bacterium, which occurs in both encapsulated and unencapsulated forms. It is predominantly a commensal organism in the nasopharynx, especially in young children. The *H. influenzae* bacterium can be further characterised, by its capsular polysaccharide, into six serotypes designated by the letters a to f. The serotype which has most often been associated with invasive disease is *H. influenzae* type b, or Hib. Before Hib vaccines became available, Hib was recognised as the most serious bacterial infection in young children in Australia.\(^{42}\) Hib caused at least 95% of invasive disease due to *H. influenzae* in children, and up to 70% of bacterial meningitis in children in Australia was estimated to be attributable to Hib.\(^{43,44}\) Worldwide, 90% of invasive Hib disease occurs in children < 5 years of age.\(^{45}\) Before Hib vaccine was introduced in Australia, infants < 18 months of age had the highest incidence,\(^{43,46}\) and Aboriginal and Torres Strait Islander children had rates among the highest recorded in the world and a significantly younger age of onset.\(^{44}\) While population-level vaccination programs in Australia have successfully reduced the incidence of Hib disease, the decline has been greater among non-Aboriginal and Torres Strait Islander children than among Aboriginal and Torres Strait Islander children and a gap in the rates of disease remains.\(^{47,48}\) The most common manifestations of invasive Hib disease are meningitis and epiglottitis, with meningitis more common in infants than in older children.\(^{46,49,50}\) Epiglottitis has been most often seen in children > 18 months of age and is rare in Aboriginal and Torres Strait Islander children.\(^{43,51}\) Survivors of Hib meningitis commonly have neurological sequelae such as deafness and intellectual impairment. Other manifestations of Hib disease include cellulitis, septic arthritis, pneumonia, pericarditis, osteomyelitis and septicaemia.\(^{46}\)

This report only presents notification data for Hib disease. No hospitalisation or mortality data are presented for invasive Hib disease because the available ICD-10-AM/ICD-10 codes are not type-specific.

**Case definition**

Notifications\(^{52}\)


**Secular trends**

Notifications for invasive Hib disease declined steeply in Australia between 1993 and 1995 following the introduction of Hib vaccine in 1993. There were 51 notifications between 1 January 2016 and 31 December 2018 (Figure 3.2.1), an average annual notification rate of 0.07 per 100,000 population (Table 3.2.1), which is similar to the rate seen in the previous review period (1 January 2012 to 31 December 2015). A median of 1 case per month (range 0–3) was notified (Figure 3.2.1).

**Severe morbidity and mortality**

There were four notified deaths among the 51 Hib notifications to the NNDSS in the three years from 1 January 2016 to 31 December 2018 (3 in 2017 and 1 in 2018). Of these four deaths, two occurred in infants aged < 1 year, one in a child aged 10 years, and one in an adult aged 61 years. Details of some of these deaths have been published elsewhere.\(^{13,53}\)
Age and sex distribution

There were more notifications for invasive Hib disease for females than males, with a male:female ratio of 1:1.3 (0.06 vs 0.08 per 100,000 population per year, respectively) over the three years from 1 January 2016 to 31 December 2018.

Notifications of invasive Hib disease were highest among children aged < 5 years, with infants < 1 year of age (Table 3.2.1) accounting for 14/51 (27%) and children aged 1–4 years accounting for 10/51 (20%) of invasive Hib notifications, incidences of 1.51 and 0.26 per 100,000 population per year, respectively. The annual average notification rate in this reporting period for children 0–4 years of age was 0.5 per 100,000 population, identical to the previous four-year reporting period (1 January 2012 to 31 December 2015). Notifications were rare among children and young adults aged 5–24 years, with one each among those aged 5–14 years and 15–24 years (Table 3.2.1).

Geographical distribution

Eighty percent of notifications occurred across New South Wales (39%), Queensland (29%), and Victoria (12%). Relative to its small population, the Northern Territory accounted for 8% of notifications for invasive Hib disease, with the highest rate of notification (0.50 per 100,000 population per year) rate, sevenfold higher than the national average of 0.07 per 100,000 population per year (Appendix B). All four notified cases in the Northern Territory occurred among Aboriginal and Torres Strait Islander people. No cases of Hib were notified in the Australian Capital Territory or Tasmania.

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for all 51 notifications of invasive Hib disease over the 2016–2018 period. Notification of Hib recorded among Aboriginal and Torres Strait Islander people accounted for 29% of all notifications (15/51) with a rate over 12 times as high as that seen in other Australians (0.61 versus 0.05 per 100,000 population per year respectively). The age distribution was younger for notifications among Aboriginal and Torres Strait Islanders, with 67% of notifications aged 0–4 years compared with 39% among other Australians.

Vaccination status

Vaccination status in the NNDSS was evaluated for all notified cases born after 31 December 1987, i.e. the cohort eligible to receive the Hib vaccine. Of the total 51 notifications, there were 31 vaccine-eligible cases. Of the 20 cases not eligible for nationally-funded vaccine, 7 (35%) were unvaccinated and no information was available for the remaining 13 cases. Of the 31 cases in vaccine-eligible birth cohorts, vaccination data were available for 29 cases (94%), of which 3 (10%) had received no vaccine doses, with one too young to be vaccinated. Among vaccinated cases (n = 26), four had received 1 dose, four had received 2 doses, nine had received 3 doses and nine had received 4 doses.

Comment

Notification rates for invasive Hib disease remained low for all ages and in children < 5 years of age between 2016 and 2018, and were similar to the previous reporting period (2012–2015). Our report is consistent with a recent review of the epidemiology of Hib disease in Australia, which provides further evidence of the persistent gap in rates of disease between Aboriginal and Torres Strait Islanders and other Australians.
Figure 3.2.1: *Haemophilus influenzae* type b notifications for all ages, Australia, January 1993 to December 2018, by month of diagnosis

![Graph showing *Haemophilus influenzae* type b notifications for all ages, Australia, January 1993 to December 2018, by month of diagnosis.]

a Notifications where the month of diagnosis was between 1 January 1993 and 31 December 2018.

Table 3.2.1: *Haemophilus influenzae* type b notifications, Australia, 2016 to 2018, by age group

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<thead>
<tr>
<th>Age group (years)</th>
<th>n</th>
<th>Rate&lt;sup&gt;b&lt;/sup&gt;</th>
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<tr>
<td><strong>All ages</strong></td>
<td><strong>51</strong></td>
<td><strong>0.07</strong></td>
</tr>
</tbody>
</table>

a Notifications where the month of diagnosis was between January 2016 and December 2018.

b Average annual age-specific rate per 100,000 population.
3.3 Hepatitis A

**Highlights**

The average annual notification rate for hepatitis A in the three-year period from 1 January 2016 to 31 December 2018 was 1.1 per 100,000 population, higher than in the previous four years from 1 January 2012 to 31 December 2015 (0.8 per 100,000 population per year), largely due to outbreaks among men who have sex with men (MSM). The notification rate in Aboriginal and Torres Strait Islander people (0.6 per 100,000 population per year) was lower than in other Australians (1.1 per 100,000 population per year). The average annual rate of hepatitis A hospitalisations was higher in the current reporting period than in the previous period (1.5 vs 1.1 per 100,000 population).

Hepatitis A is caused by the hepatitis A virus (HAV), a RNA virus classified within the genus hepatovirus of the picornavirus family. There is only one human HAV serotype. Hepatitis A is an acute inflammatory disease of the liver and can produce either asymptomatic or symptomatic infection. Clinical manifestations of symptomatic infection vary from mild anicteric illness to fulminant hepatic failure. Infection with HAV typically has a sudden onset of symptoms that can include fever, anorexia, malaise, nausea, abdominal discomfort, jaundice and dark urine. The likelihood of having symptoms with HAV infection is related to age. In young children, hepatitis A is usually asymptomatic or associated with mild illness without jaundice. In adults, symptomatic infection is characteristic and 70–95% of infected adults show clinical symptoms. Complications are uncommon, but in rare cases infection can develop into fulminant hepatitis for which mortality can be as high as 60%. Transmission typically occurs via the faecal-oral route or through close physical contact with an infected person. Hepatitis A infection was prevalent among Aboriginal and Torres Strait Islander children in the late 1990s, but a national targeted program in affected jurisdictions since 2005 has been highly effective in decreasing disease incidence in both children and adults. Increasingly, a large proportion of cases in Australia are related to overseas travel to countries where hepatitis A is endemic.

**Case definition**

**Notifications**

Hepatitis A case definition:


**Hospitalisations and deaths**

The ICD-10-AM/ICD-10 codes B15.0 (hepatitis A with hepatic coma) and B15.9 (hepatitis A without hepatic coma) were used to identify hospitalisations and deaths.

**Secular trends**

There were 796 notifications of hepatitis A over the three-year period from 1 January 2016 to 31 December 2018, with a median of 20 (ranging from 5 to 59) per month (Figure 3.3.1). The average annual notification rate was 1.08 per 100,000 population (Table 3.3.1), with the annual rate increasing over the three-year period from 0.60 per 100,000 population in 2016 (n = 145) to 0.88 per 100,000 population in 2017 (n = 217) and 1.74 per 100,000 population in 2018 (n = 434) (Appendix B). Of the 796 notifications, 263 were noted as acquired in Australia (33%), 351 from overseas (44%) and 182 (23%) unknown. The total number of coded hospitalisations (n = 1,129) exceeded notifications, but only 451 (40%) were coded as principal diagnosis (average annual rate of 0.61 per 100,000 population). The rate of hepatitis A notifications and hospitalisations was higher in this reporting period (2016–2018) than in the previous four-year reporting period from
Severe morbidity and mortality

In the three years from 1 January 2016 to 31 December 2018, hospitalisations where hepatitis A was the principal diagnosis accounted for 1,459 bed-days, with a median length of stay of two days, increasing with age (Table 3.3.1). A total of nine deaths were recorded in the causes of death data, although hepatitis A was recorded as an associated rather than underlying cause of death in all of these. All deaths occurred among adults aged 25 years or older. Two deaths were reported in the NNDSS (aged between 55 and 70 years) and two in-hospital deaths where hepatitis A was the principal diagnosis (both aged 60 years or older) were recorded in AIHW hospitalisation data.

Age and sex distribution

The largest number of notifications of hepatitis A (48%) was in the 25-49 year age group (383/796, 1.49 per 100,000 population per year) but notification rates were similar for those aged 1-49 at 1.4 to 1.5 per 100,000 population per year (Table 3.3.1). When restricted to principal diagnosis, hospitalisation rates were highest in those 25-49 years (Table 3.3.1), although this varied by year. Notifications were almost twice as high in males as in females (523 vs 273, ratio 1.95:1).

Geographical distribution

The highest numbers of notified cases during this reporting period were from Victoria (n = 394; a notification rate of 2.08 per 100,000 population per year) and New South Wales (n = 200; a notification rate of 0.85 per 100,000 population per year), where outbreaks of hepatitis A were observed. Hospitalisation rates were similarly high in these states (2.30 per 100,000 population per year in Victoria [n = 436] and 1.46 per 100,000 population per year in New South Wales [n = 344]). Tasmania, the Northern Territory and Queensland had the lowest average annual rates of notifications (0.19, 0.27 and 0.67 per 100,000 population, respectively) and the lowest hospitalisation rates where hepatitis A was the principal diagnosis (0.26, 0.41 and 0.31 per 100,000 population, respectively) (Appendix C).

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for 741 of the 796 notifications (93%) of Hepatitis A during 2016–2018, with 55 not stated (7%). Of the 796 hepatitis A notifications, 15 were in Aboriginal and Torres Strait Islander people (2% of all notifications) and 781 in other people (notification rate 0.6 versus 1.1 per 100,000 population per year, respectively).

Vaccination status

Of the 796 notifications, vaccination data was available for 57% (454/796) with 434 of those (96%) reported to have received no vaccine doses. Of the 20 vaccinated cases, only seven had received 2 or more doses.

Comment

Hepatitis A vaccine has been available on the National Immunisation Program (NIP) for Aboriginal and Torres Strait Islander children in the Northern Territory, Queensland, South Australia and Western Australia since 2005, with a resultant marked decline in disease rates despite relatively modest vaccination coverage. Aboriginal and Torres Strait Islander people remained lower than in other Australians in this reporting period. However an overall increase in notifications of hepatitis A was observed, largely due to outbreaks, primarily among men who have sex with men (MSM). Other key risk groups include people who inject drugs, homeless persons and adult prisoners. State-funded 2-dose hepatitis A vaccination programs targeting
MSM and injection drug users were introduced in Victoria and Tasmania from mid-2018 until June 2019 to control the outbreak, with the program successfully increasing immunity against hepatitis A and likely preventing disease transmission and further outbreaks. Additional food-borne outbreaks resulting in substantial numbers of cases have also been reported, particularly in Victoria and New South Wales. However, a large proportion of cases continue to be acquired overseas (351/796), accounting for 44% of notifications in this reporting period, with data missing for 23% of cases.

The Australian Immunisation Handbook recommends hepatitis A vaccination for people with an increased risk of acquiring hepatitis A, due to occupational or other factors, but coverage of the vaccine in such groups is unknown.
Table 3.3.1: Hepatitis A notifications, hospitalisations and deaths, Australia, 2016 to 2018, by age group

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<th>Age group (years)</th>
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<td>Rate(^d)</td>
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<td>Rate(^d)</td>
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</tbody>
</table>

| All ages         | 796 | 1.08 | 1,129 | 1.53 | 451 | 0.61 | 2 | 2 | 9 | 0.01 |

\(a\) Notifications where the month of diagnosis was between 1 January 2016 and 31 December 2018; hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.

\(b\) LOS: length of stay in hospital.

\(c\) Deaths include underlying and associated causes of deaths.

\(d\) Average annual age-specific rate per 100,000 population.

\(e\) Principal diagnosis (hospitalisations).

\(f\) np: not provided.
3.4 Hepatitis B

Highlights

The declining trend in the number of notifications of newly-acquired hepatitis B observed since 2007 continued over the three-year reporting period. There were 461 notifications of newly-acquired hepatitis B from 1 January 2016 to 31 December 2018, and the majority (58%) were among adults aged 25 to 49 years.

The acute hepatitis B hospitalisation rate was low at 0.37 per 100,000 population per year from 1 January 2016 to 31 December 2018.

The focus of this chapter is acute infection with hepatitis B virus (HBV), a hepadnavirus. Humans are the only known reservoir for human HBV genotypes, but closely related HBV genotypes exist in higher primates. The outcomes of HBV infection depend on host factors including age, sex, genetic background, co-infections, other co-existing diseases and concomitant medications, and on viral factors including the HBV genotype and viral DNA levels. It produces a range of conditions from subclinical infection to acute and, rarely (in < 1% of cases), fulminant hepatitis. The majority of HBV infections are not clinically recognised, with < 10% of children and 30–50% of adults experiencing jaundice. When illness occurs, it is usually insidious, with anorexia, vague abdominal discomfort, nausea and vomiting, sometimes arthralgia and rash, often progressing to jaundice. The main burden of disease is related to chronic HBV infection, which is the primary cause of cirrhosis deaths in the Asia-Pacific region. The risk of an acute infection becoming chronic varies inversely with age. Chronic HBV infection occurs in about 90% of infants infected at birth, in 20–50% of children infected at 1–5 years of age, and with much lower but highly variable risk (1–10%) in people infected as older children and adults.

HBV transmission occurs by percutaneous or permucosal exposure to infective body fluids such as blood, semen, vaginal secretions and any other body fluid containing blood. Major modes of transmission include sexual or close household contact with an infected person; perinatal transmission from mother to infant; injecting drug use; and nosocomial exposure. Screening of blood and organ donors has nearly eliminated the risk of transmission through blood transfusion and organ transplants.

The summary below is restricted to newly-acquired hepatitis B notifications and acute hepatitis B hospitalisations.

Case definition

Notifications

Hepatitis B (newly acquired) case definition:

Hospitalisations and deaths

The ICD-10-AM/ICD-10 code B16 (acute hepatitis B) was used to identify hospital admissions and deaths. As in previous reports, only those hospitalisations with B16 as the principal diagnosis were included.

Secular trends

In the three years from 1 January 2016 to 31 December 2018, there were 461 notifications of newly acquired hepatitis B (an average annual rate of 0.6 per 100,000 population) (Table 3.4.1), decreasing from a median of 14.0 per month in 2016 to 11.5 in 2018 (Figure 3.4.1). There were 274 hospitalisations with a principal diagnosis of acute hepatitis B over the three years, an...
average annual rate of 0.37 per 100,000 population (Table 3.4.1). The number and rate of hospitalisations were lower in this reporting period than in the previous four-year period (2012–2015), but rates have remained fairly stable since 2015 (0.35 per 100,000 population in 2016; 0.36 per 100,000 population in 2017; and 0.41 per 100,000 population in 2018).

Severe morbidity and mortality

In the three-year reporting period, the 274 hospitalisations for acute hepatitis B infection accounted for 1,453 bed days. The median length of stay in hospital was four days for people aged 15–64 years, and slightly higher (median: five days) for people aged ≥ 65 years (Table 3.4.1). Seven hospitalisations occurred among children 5–14 years, all in 2018, of which six were Aboriginal and Torres Strait Islander children. Between 2016 and 2018, there were no hospitalisations among children aged less than 5 years.

There were 533 deaths recorded in the causes of death data over the three-year period where acute hepatitis B infection (ICD-10 code B16) was an underlying or associated cause, with only 52 (9.8%) coded as the underlying cause of death (eight deaths in those aged 25–49 years, 19 in 50–64 years and 25 in ≥ 65 years) (Table 3.4.1). The number of deaths where hepatitis B was the underlying cause of death decreased from 24 in 2016 to 21 in 2017 and 7 in 2018. No deaths were recorded among notifications to NNDSS during this reporting period, and 1–4 in-hospital deaths were recorded in the AIHW dataset among cases with a principal diagnosis of acute hepatitis B, all aged between 45 and 55 years.

Age and sex distribution

From 1 January 2016 to 31 December 2018, the highest rates for notification of newly-acquired hepatitis B infection and hospitalisations for acute hepatitis B were among adults aged 25–49 years, followed by adults aged 50–64 years (Table 3.4.1). In children aged 0–4 years, there were six notifications of newly-acquired hepatitis B and no hospitalisations (Table 3.4.1).

Overall, the number of notifications and hospitalisations was higher for males (male:female rate ratio for notifications 2.6:1 and for hospitalisations 2.1:1), especially in adults aged 50–64 years (male:female rate ratio for notifications 6.4:1 and for hospitalisations 3.3:1).

Geographical distribution

Similar numbers of notifications were reported in Queensland (146/461, 32%) and Victoria (144/461, 31%), with notable numbers reported in Western Australia (73/461, 16%) and New South Wales (48/461, 10%) over the 3 years from 1 January 2016 to 31 December 2018 (Appendix B). The largest numbers of hospitalisations occurred in Victoria (31%), New South Wales (25%) and Queensland (20%). However, the highest rates of notifications and hospitalisations occurred in the Northern Territory (average annual rate of 1.89 per 100,000 population for both notifications and hospitalisations) (Appendices B and C).

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for 430 of the 461 notifications (93%) of hepatitis B during 2016–2018, with 31 not stated (7%). Of the 461 notifications, 37 (8%) were in Aboriginal and Torres Strait Islander people. Of the 274 hospitalisations with hepatitis B recorded as the principal diagnosis, 23 (8%) were in Aboriginal or Torres Strait Islander people. Notification rates in Aboriginal and Torres Strait Islander people were 2.5 times as high (1.51 per 100,000 population per year) as in other Australians (0.59 per 100,000 population per year), and hospitalisation rates were 2.7 times as high (0.94 per 100,000 population per year versus 0.35 per 100,000 population per year). Aboriginal and Torres Strait Islanders accounted for 55 of the 533 deaths (10.3%) where hepatitis B was recorded as underlying or associated cause, and for five of the 52 deaths (9.6%)
Figure 3.4.1: Hepatitis B notifications and hospitalisations with a principal diagnosis of acute hepatitis B, Australia, 1997 to 2018, by month of diagnosis or admission

Notifications with month of diagnosis between 1 January 1997 and 31 December 2018; hospitalisations where the month of admission was between 1 January 1997 and 31 December 2018. This figure includes data from 1997 onwards since it was not until 1996 that acute hepatitis B became notifiable in all states and territories; prior to 1994, hospitalisations for acute hepatitis B were not distinguished from chronic hepatitis B.

where hepatitis B was recorded as the underlying cause, with rates more than three times as high as those that seen in other Australians.

Vaccination status

Of the 12 notifications between January 2016 and December 2018 in people born in 2000 or later who were eligible to receive hepatitis B vaccine as part of the universal infant vaccination program, vaccination status was not available for five cases (42%). Of the seven cases (58%) with vaccination data available, two were reported to have received no vaccine doses (unvaccinated), of whom one was too young to receive the vaccinations scheduled at 2 months of age. Of the five cases who had received a hepatitis B-containing vaccine, one had received 2 doses and four had received 4 doses.

Comment

Universal infant hepatitis B immunisation was introduced in May 2000, although targeted programs had been in place since the 1980s in all jurisdictions. A progressive downward trend in total notifications of acute hepatitis B has been evident since 2007, with notifications continuing to decline between 2016 and 2018. However, the longstanding disparity in rates of hepatitis B infections among Aboriginal and Torres Strait Islander people persists. Modelling has suggested that a nationally-funded hepatitis B immunisation program for Aboriginal and Torres Strait Islander adults could have a considerable impact on the prevention of acute and subsequent chronic hepatitis B infections.
Table 3.4.1: Hepatitis B notifications, hospitalisations and deaths, Australia, 2016 to 2018, \( ^a \) by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications</th>
<th>Hospitalisations( ^b )</th>
<th>LOS( ^c ) per admission ( ^b )</th>
<th>Deaths( ^d )</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Rate( ^e )</td>
<td>n</td>
<td>Rate( ^e )</td>
</tr>
<tr>
<td>&lt; 1</td>
<td>2</td>
<td>0.22</td>
<td>0</td>
<td>0.00</td>
</tr>
<tr>
<td>1–4</td>
<td>4</td>
<td>0.11</td>
<td>0</td>
<td>0.00</td>
</tr>
<tr>
<td>5–14</td>
<td>3</td>
<td>0.03</td>
<td>7</td>
<td>0.08</td>
</tr>
<tr>
<td>15–24</td>
<td>35</td>
<td>0.36</td>
<td>24</td>
<td>0.25</td>
</tr>
<tr>
<td>25–49</td>
<td>269</td>
<td>1.04</td>
<td>158</td>
<td>0.61</td>
</tr>
<tr>
<td>50–64</td>
<td>115</td>
<td>0.88</td>
<td>62</td>
<td>0.47</td>
</tr>
<tr>
<td>≥ 65</td>
<td>33</td>
<td>0.29</td>
<td>23</td>
<td>0.20</td>
</tr>
<tr>
<td>All ages</td>
<td>461</td>
<td>0.62</td>
<td>274</td>
<td>0.37</td>
</tr>
</tbody>
</table>

\( ^a \) Notifications where the month of diagnosis was between 1 January 2016 and 31 December 2018; hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.

\( ^b \) Principal diagnosis (hospitalisations).

\( ^c \) LOS: length of stay in hospital.

\( ^d \) Deaths include underlying and associated causes of deaths.

\( ^e \) Average annual age-specific rate per 100,000 population.

\( ^f \) np: not provided.
3.5 Influenza

**Highlights**

There was a marked increase in influenza notifications, hospitalisations and deaths in 2017, which was the largest season observed since the 2009 pandemic, with both A/H3N2 and type B influenza strains circulating. Increased use of PCR rapid tests and changing testing and reporting practices partially contributed to the increase in notifications observed over the reporting period 2016 to 2018.

Influenza viruses, predominantly influenza type A (H1N1 and H3N2) and type B (Yamagata and Victoria lineages), cause annual epidemics of respiratory disease, which is mainly spread by droplet transmission. The disease is often indistinguishable clinically from that caused by other respiratory viruses. Typical symptoms include abrupt onset of fever, cough, malaise, myalgia, sore throat and headache. Complications of influenza infection include pneumonia, cardiovascular events, stroke, otitis media and exacerbation of chronic medical conditions. Significant antigenic changes can lead to pandemics with higher rates of illness and death. Seasonal epidemics occur in Australia mainly between June and September, with year-to-year variability in terms of activity, severity and age groups affected, based on the viruses predominantly circulating. Influenza can cause severe illness or death, particularly in high-risk groups including children younger than 5 years; people 65 years and older; Aboriginal and Torres Strait Islander people; pregnant women; and people with chronic medical conditions.

**Case definition**

**Notifications**

Influenza laboratory-confirmed case definition:


**Hospitalisations and deaths**

The ICD-10-AM/ICD-10 codes J09 (influenza due to certain identified influenza virus, including avian influenza and the influenza A/H1N1 pandemic strain), J10 (influenza due to identified virus) and J11 (influenza, virus not identified) were used to identify influenza hospitalisations and deaths. As no avian influenza cases have been reported in Australia, J09 in this report refers to the influenza A/H1N1 pandemic strain.

**Secular trends**

In the three years from 1 January 2016 to 31 December 2018, there were 400,907 notifications of influenza with an average annual rate of 543.4 per 100,000 population (Table 3.5.1). Notifications varied substantially by year, with the highest levels of activity since the 2009 pandemic season seen in 2017 (251,163 notifications, 1020.9 per 100,000 population, comprising 63% of the influenza notifications across 2018–2018). Notifications in 2016 were lower at 90,879 (375.7 per 100,000 population), and lower still in the mild 2018 influenza season (58,865 notifications, 235.6 per 100,000 population) (Appendix B). Similar to previous reporting periods, seasonal peaks in influenza notifications were observed between July and September, although the 2017 season appeared to continue into October (Figure 3.5.1). Although notifications were substantially lower in 2018 compared with previous years, a higher number of inter-seasonal notifications were observed following the
regular 2018 season, with three times as many notifications observed in December 2018 as in December 2017 (6,269 vs 2,207, respectively).

From 1 January 2016 to 31 December 2018, there were 87,075 hospitalisations with ICD-10-AM influenza codes J09, J10 or J11. A trend similar to that for notifications was observed, with the highest number of hospitalisations observed in 2017 (50,688 hospitalisations, 206.0 per 100,000 population) (Appendix C). Influenza was the principal diagnosis in 63% of hospitalisations, with 89% of these coded as J10. Influenza coded as J11 and J09 made up only a small proportion of cases (9% and 3%, respectively).

Severe morbidity and mortality

In the three-year reporting period, there were 469,172 bed days for hospitalisations due to any influenza (J09, J10 or J11). The median length of hospital stay was three days, increasing to five days for patients aged ≥ 65 years (Table 3.5.2). In the cause of death data, influenza (ICD-10 codes J09, J10, J11) was recorded in 2,368 cases between 1 January 2016 and 31 December 2018, an average annual rate of 3.2 deaths per 100,000 population (Table 3.5.1). This varied substantially from year to year, being highest in 2017 (1,629 deaths; 6.62 per 100,000 population) followed by 2016 (585 deaths; 2.42 per 100,000) and lowest in 2018 (154 deaths; 0.62 per 100,000 population).

Age and sex distribution

The highest rate of notifications for influenza was among children aged 1–4 years for the overall three-year reporting period (924.2 per 100,000 population per year) (Table 3.5.1), and for each individual year included in the reporting period (575.1, 1722.5 and 469.0 per 100,000 population).
population in 2016, 2017 and 2018 respectively). The second highest rate of influenza notifications was among adults aged 65 years and older (746.0 per 100,000 population per year) (Table 3.5.1), but there was some variation by year, dependent on the season characteristics. For example, in 2018 there were higher notification rates observed among infants aged < 1 years (298.0 per 100,000 population) and children aged 5–14 years (307.9 per 100,000 population) than among adults aged 65 years and older (249.2 per 100,000 population). Hospitalisation rates for influenza coded as J09, J10 or J11 were highest for adults aged 65 years or older, followed by infants aged < 1 year (Table 3.5.1). Adults aged 65 years and older accounted for 91% of recorded influenza deaths.

There were slightly fewer influenza notifications and hospitalisations for males than for females (male:female rate ratio 0.8:1 for notifications and 0.9:1 for hospitalisations).

Geographical distribution

Notification rates were highest in South Australia, followed by New South Wales and Queensland (Appendix B). Notification rates in 2017 were more than twofold higher across every state and territory, relative to 2016, with the exception of Western Australia where rates remained relatively constant across the three-year period. Hospitalisation rates were highest in the Northern Territory and Queensland (196.0 and 151.3 per 100,000 population per year, respectively) (Appendix C). As with notifications, hospitalisation rates were markedly higher in 2017 across all jurisdictions except Western Australia.

Aboriginal and Torres Strait Islander status

Of the 400,907 notifications of influenza over the 2016–2018 period, Aboriginal and Torres Strait Islander status was reported for only 133,604 (33%). Reporting of Aboriginal and Torres Strait Islander status varied across jurisdictions, ranging from the Northern Territory (98% complete) to Tasmania (6% complete), New South Wales (8% complete) and Victoria (17% complete). The Australian Capital Territory (87%), South Australia (70%) and Queensland (56%) also had relatively high completeness. Due to low completeness of reporting of Aboriginal and Torres Strait Islander status in several jurisdictions, notification rates of influenza for Aboriginal and Torres Strait Islander people are not presented in this report.

Of the 87,075 hospitalisations for influenza, approximately 4.7% (4,053) were recorded as occurring in Aboriginal and Torres Strait Islander people, a hospitalisation rate of 165.9 per 100,000 population per year compared with 116.4 per 100,000 population per year in other Australians (rate ratio 1.43). Of the 2,368 deaths recorded in the causes of death data, 40 (2%) occurred among Aboriginal and Torres Strait Islander people.

Vaccination status

Vaccination status data were not provided to NCIRS due to data quality issues. As influenza notifications are reported to the NNDSS by laboratories, generally without further public health follow up, vaccination status is not routinely reported for influenza notifications.

Influenza type and subtype

Influenza A comprised the majority of notifications in each year, decreasing from 2016 (89%) to 2018 (80%) and 2017 (62%), with influenza B comprising the remainder. Of those influenza A viruses that were subtyped, A/H3N2 predominated in 2016 (64.5% vs 35.5% A/H1N1) and 2017 (76% vs 24% A/H1N1), while A/H1N1 was more common in 2018 (66.5% vs 33.5% A/H3N2).

Over the three reporting years, influenza A was responsible for the highest proportion of notifications in infants aged < 1 year (80%) and older adults aged 65 years and over (77%), while
influenza B occurred most frequently among children aged 5–14 years (41%) and adolescents and young adults aged 15–24 years (32%).

Comment

Influenza remains the most commonly notified VPD in Australia. Consistent with a detailed review of the epidemiology of influenza in Australia for the period 2006 to 2015, high rates of disease were observed among young infants and children and older adults aged 65 years and older. Notification rates in Aboriginal and Torres Strait Islander people were not calculated as they would substantially underestimate the burden of influenza due to low completeness of reporting of Aboriginal and Torres Strait Islander status in many states, as has been reported previously.

This reporting period was marked by a longer influenza season of heightened activity in 2017, with excess mortality due to influenza and pneumonia observed above what would normally be expected, and in which influenza subtype A/H3N2 predominated. Influenza A/H3N2-predominant seasons tend to be associated with higher morbidity and mortality, particularly among adults aged 65 years and over. Influenza A/H3N2 also tends to be associated with lower vaccine effectiveness. Influenza vaccine effectiveness in 2017 was lower than in previous seasons, particularly for the A/H3N2 strain. The severity of the 2017 season led to the introduction, in 2018, of two new enhanced influenza vaccines (a high-dose and an adjuvanted influenza vaccine) onto the National Immunisation Program for Australians aged 65 years and older.

In contrast, influenza activity in 2018 was low and comparatively mild, with A/H1N1 the dominant circulating strain. Vaccine effectiveness was also higher in 2018.

Comparisons made with rates of influenza in earlier years, particularly the 2009 A/H1N1 pandemic year, can be misleading, due to improved reporting of cases of influenza and the impact of increased influenza testing in recent years, an effect that is difficult to quantify. Increased uptake of polymerase chain reaction (PCR) testing in primary care, and of rapid PCR testing in hospitals, have contributed to increases in laboratory-confirmed influenza notifications in the last decade. Although geographic variations are expected, differences in notifications, hospitalisations and deaths across jurisdictions are likely due in part to testing and reporting practices. Despite improvements in reporting, notifications of influenza to NNDSS still likely substantially underestimate the true rate of influenza disease. Similarly, coded influenza death data are known to significantly underestimate the true influenza-associated hospitalisation and death rates.
Table 3.5.1: Influenza notifications, hospitalisations and deaths, Australia, 2016 to 2018,\(^a\) by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications</th>
<th></th>
<th></th>
<th>Hospitalisations</th>
<th></th>
<th></th>
<th></th>
<th>LOS(^b) per admission</th>
<th>Deaths(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Rate(^d)</td>
<td>Any diagnosis</td>
<td></td>
<td>Principal</td>
<td></td>
<td>Any</td>
<td>Principal</td>
<td></td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>Rate(^d)</td>
<td></td>
<td>diagnosis</td>
<td></td>
<td>diagnosis</td>
<td></td>
<td>diagnosis</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>n</td>
<td>Rate(^d)</td>
<td></td>
<td>n(^e)</td>
<td>Rate(^d)</td>
<td></td>
<td>Median days</td>
</tr>
<tr>
<td>&lt; 1</td>
<td>5,998</td>
<td>645.6</td>
<td>2,634</td>
<td>283.5</td>
<td>2,001</td>
<td>215.4</td>
<td>2</td>
<td>2</td>
<td>1–5</td>
</tr>
<tr>
<td>1–4</td>
<td>35,081</td>
<td>924.2</td>
<td>4,711</td>
<td>124.1</td>
<td>3,656</td>
<td>96.3</td>
<td>2</td>
<td>2</td>
<td>7</td>
</tr>
<tr>
<td>5–14</td>
<td>65,425</td>
<td>712.7</td>
<td>3,653</td>
<td>39.8</td>
<td>2,709</td>
<td>29.5</td>
<td>1</td>
<td>1</td>
<td>np(^f)</td>
</tr>
<tr>
<td>15–24</td>
<td>38,499</td>
<td>401.0</td>
<td>3,528</td>
<td>36.7</td>
<td>2,443</td>
<td>25.4</td>
<td>1</td>
<td>1</td>
<td>10</td>
</tr>
<tr>
<td>25–49</td>
<td>107,473</td>
<td>417.0</td>
<td>12,488</td>
<td>48.5</td>
<td>8,101</td>
<td>31.4</td>
<td>2</td>
<td>1</td>
<td>55</td>
</tr>
<tr>
<td>50–64</td>
<td>63,508</td>
<td>484.0</td>
<td>13,018</td>
<td>99.2</td>
<td>7,950</td>
<td>60.6</td>
<td>3</td>
<td>3</td>
<td>143</td>
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<td>≥ 65</td>
<td>84,869</td>
<td>746.0</td>
<td>47,043</td>
<td>413.5</td>
<td>27,675</td>
<td>243.3</td>
<td>5</td>
<td>4</td>
<td>2,145</td>
</tr>
<tr>
<td>All ages</td>
<td>400,907(^g)</td>
<td>543.4</td>
<td>87,075</td>
<td>118.0</td>
<td>54,535</td>
<td>73.9</td>
<td>3</td>
<td>3</td>
<td>2,368</td>
</tr>
</tbody>
</table>

\(^a\) Notifications where the month of diagnosis was between 1 January 2016 and 31 December 2018; hospitalisations (ICD-10-AM codes J09, J10 and J11) where the month of admission was between 1 January 2016 and 31 December 2018.

\(^b\) LOS: length of stay in hospital.

\(^c\) Deaths include underlying and associated causes of deaths.

\(^d\) Average annual age-specific rate per 100,000 population.

\(^e\) Principal diagnosis (hospitalisations).

\(^f\) np: not provided.

\(^g\) Total notifications for all ages: includes 54 notifications without data on age group.
3.6 Measles

Measles is an acute and highly communicable disease caused by a member of the genus Morbillivirus. Before the introduction of a vaccine, measles caused more than two million deaths annually worldwide. The virus is transmitted directly from person to person, by respiratory droplets, and is contagious from approximately two weeks before symptoms develop for up to four days after the rash appears. The clinical picture includes a prodromal fever, cough, coryza, conjunctivitis, and Koplik spots on the buccal mucosa, before the onset of rash. Complications include otitis media, pneumonia and encephalitis. Subacute sclerosing panencephalitis (SSPE) occurs very rarely as a late sequel of wild infection but not of vaccination. Complications and deaths occur more commonly in developing countries in children aged < 5 years and adults, and in persons with malnutrition or immune deficiencies. Improved rates of childhood vaccination have resulted in the burden of disease shifting from young children to older age groups. Achieving herd immunity requires 2-dose vaccine coverage rates in excess of 95%. In 2014, the World Health Organization declared that Australia had achieved elimination of measles.

Case definition

Notifications

Measles case definition:


Hospitalisations and deaths

The ICD-10-AM/ICD-10 code B05 (measles) was used to identify hospitalisations and deaths. SSPE, a very rare late sequel of measles infection, was not included in this analysis.

Secular trends

In the three years from 1 January 2016 to 31 December 2018, there were 283 notifications of measles recorded nationally, an average annual notification rate of 0.4 per 100,000 population (Table 3.6.1), a 50% reduction on the notification rate recorded in the four-year period 2012 to 2015 (0.8 per 100,000 population per year). Notification rates were similar in 2016 (n = 99, 0.4 per 100,000 population), 2017 (n = 81, 0.3 per 100,000 population), and 2018 (n = 103, 0.4 per 100,000 population) (Figure 3.6.1 insert and Appendix B). About half of notified cases (151/283, 53%) were acquired within Australia, and 126 (45%) acquired overseas, with only six cases (2%) of unknown origin.

Hospitalisations followed the same general trend as notifications (Figure 3.6.1). From 1 January 2016 to 31 December 2018, there were 163 hospitalisations with the ICD-10-AM code B05 (measles) at an average annual rate of 0.22 per 100,000 population (Table 3.6.1), lowest in 2017 (n = 41, 0.17 per 100,000 population) and similar in 2016 (n = 60) and 2018 (n = 62) (rate=0.25 per 100,000 population in both years) (Appendix C).
Severe morbidity and mortality

In the three-year reporting period, hospital admissions coded as measles accounted for 546 hospital bed days. The median length of stay was three days; of the 163 hospitalisations, 140 (86%) had measles recorded as the principal diagnosis (Table 3.6.1). In the causes of death data, 1–5 deaths were recorded with measles as the underlying or associated cause of death for 2016–2018, all in people aged 65 years and older. There were no deaths recorded in the NNDSS for this reporting period.

Age and sex distribution

From 1 January 2016 to 31 December 2018, the highest age-specific notification rate (3.6 per 100,000 population per year) was in infants (< 1 year) (Table 3.6.1). Notification rates among adolescents and adults in the 2016–2018 period were lower than in the 2012–2015 period (15–24 year olds: 0.49 vs 1.63 per 100,000 population per year; 25–49 year olds: 0.55 vs 0.83 per 100,000 population per year). A greater proportion of cases among infants and young children aged 1–4 years were acquired overseas (52% and 68%) than was evident among older children and adults (5–14 years: 40%; 15–24 years: 34%; 25–49 years: 46%; 50–64 years: 50%).

Age-specific hospitalisation rates reflected notification rates; the highest rate was in infants < 1 year of age (1.61 per 100,000 population per year), with hospitalisation rates below 1 per 100,000 population per year for all other age groups (Table 3.6.1). There were slightly more notifications and hospitalisations for females than males (male:female rate ratio 0.9:1 for both).

Figure 3.6.1: Measles notifications and hospitalisations, Australia, 1993 to 2018, by month of diagnosis or admission

Notifications where the month of diagnosis was between 1 January 1993 and 31 December 2018; hospitalisations where the month of admission was between 1 January 1993 and 31 December 2018.
Geographical distribution

Measles notifications rates over the three-year reporting period were highest in Western Australia (0.85 per 100,000 population per year) (Appendix B). Consistent with population size, cases in Victoria and New South Wales accounted for the majority of notifications (30% and 24%, respectively).

Aboriginal and Torres Strait Islander status

Of 283 notifications of measles 2016–2018, Aboriginal and Torres Strait Islander status was reported for 277 (98% complete), with only seven cases (2.5%) reported in Aboriginal and Torres Strait Islander people, all in New South Wales. The measles notification rate in Aboriginal and Torres Strait Islander people was lower than in other Australians (0.29 vs 0.39 per 100,000 population per year).

Vaccination status

Of the 283 notifications, vaccination data was available for 187 cases (66%), of which 139 (74%) were reported to have received no vaccine doses; 25/139 (18%) were aged < 1 year and hence not eligible for vaccination. Of the 48 cases who had received at least one dose of measles-containing vaccine more than 14 days before disease onset, 26 had received 1 dose and 21 had received 2 doses (data on the number of doses received for one case was missing).

Comment

Measles notifications and hospitalisations continue to remain low in Australia, with the majority of cases in Australia travel-related. Outbreaks linked to imported cases related to travel to or from high endemicity regions continue to sporadically occur, most notably during this reporting period in Western Australia. Evidence continues to support the maintenance of elimination of endemic measles from Australia, which was verified by the World Health Organization in 2014.

A substantial proportion of cases were unvaccinated, highlighting the need to achieve high vaccine coverage across all age groups and to ensure vaccination among travellers to endemic countries. Although vaccinated cases typically have a milder clinical presentation, evidence of declining measles-specific immunoglobulin G (IgG) antibody levels among individuals more than 10 years since vaccination emphasizes the importance of continued vigilance in monitoring rates of disease, particularly in light of the shift of the second dose of measles-containing vaccine from age 4 years to 18 months in 2013.
Table 3.6.1: Measles notifications, hospitalisations and deaths, Australia, 2016 to 2018,\(^a\) by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications</th>
<th>Hospitalisations</th>
<th>LOS(^b) per admission</th>
<th>Deaths(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Rate(^d)</td>
<td>n</td>
<td>Rate(^d)</td>
</tr>
<tr>
<td>&lt; 1</td>
<td>33</td>
<td>3.55</td>
<td>15</td>
<td>1.61</td>
</tr>
<tr>
<td>1–4</td>
<td>19</td>
<td>0.50</td>
<td>24</td>
<td>0.63</td>
</tr>
<tr>
<td>5–14</td>
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<td>0.27</td>
<td>np</td>
<td>np</td>
</tr>
<tr>
<td>15–24</td>
<td>74</td>
<td>0.77</td>
<td>36</td>
<td>0.37</td>
</tr>
<tr>
<td>25–49</td>
<td>126</td>
<td>0.49</td>
<td>69</td>
<td>0.27</td>
</tr>
<tr>
<td>50–64</td>
<td>6</td>
<td>0.05</td>
<td>9</td>
<td>0.07</td>
</tr>
<tr>
<td>≥65</td>
<td>0</td>
<td>0.00</td>
<td>1–4</td>
<td>0.04</td>
</tr>
<tr>
<td>All ages</td>
<td>283</td>
<td>0.38</td>
<td>163</td>
<td>0.22</td>
</tr>
</tbody>
</table>

\(^a\) Notifications where the month of onset was between 1 January 2016 and 31 December 2018; hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.

\(^b\) LOS: length of stay in hospital.

\(^c\) Deaths sourced from the Causes of Death database from the Australian Coordinating Registry. Deaths include underlying and associated causes of deaths.

\(^d\) Average annual age-specific rate per 100,000 population.

\(^e\) Principal diagnosis (hospitalisations).

\(^f\) np: not provided.
3.7 Invasive meningococcal disease

**Highlights**

The reporting period 2016–2018 saw the reversal of a declining trend (evident since the commencement of the national meningococcal C vaccination program in 2003) in notifications and hospitalisations for invasive meningococcal disease. Increases in notifications and hospitalisations commenced in 2015, peaking in 2017 at 380 notifications (1.54 per 100,000 population per year). This increase was largely due to the increase in disease associated with serogroups W and Y (38% and 17% of all notified cases, respectively), with serogroup B cases making up 38%.

Invasive meningococcal disease is defined as the isolation of *Neisseria meningitidis* from cerebrospinal fluid (CSF), blood or other normally-sterile sites including skin lesions. There are 13 *N. meningitidis* serogroups, of which the most common globally are A, B, C, W and Y. Clinical manifestations include meningitis; septicemia with or without meningitis, and septic arthritis. In culture-negative cases with a compatible clinical picture (such as fever, haemorrhagic rash and shock), a diagnosis of meningococcal disease can be supported by a range of laboratory evidence. This includes the identification of gram-negative intracellular diplococci or meningococcal antigen in blood or CSF, the identification of nucleic acid from *N. meningitidis* in body fluids, or demonstration of a serological response to *N. meningitidis*. Between 10% and 30% of survivors of invasive meningococcal disease develop permanent physical, psychological and/or cognitive sequelae.

The summary below includes data on invasive meningococcal disease by serogroup. Serogroup data are only available for notifications reported to the NNDSS, as hospitalisation and death data do not include information on serogroup. Given the high ascertainment of invasive meningococcal disease cases by the NNDSS and high completeness of recording of deaths, we report data on deaths recorded in the NNDSS complemented by data recorded in the causes of death data.

**Case definitions**

**Notifications**

Meningococcal disease (invasive) surveillance case definition:


**Hospitalisations and deaths**

The ICD-10-AM/ICD-10 code A39 (meningococcal infection) was used to identify hospitalisations and deaths.

**Secular trends**

There were 913 notifications of invasive meningococcal disease during the three years from 1 January 2016 to 31 December 2018. This equated to an average annual notification rate of 1.2 per 100,000 population, 55% higher than in the four-year period from 2012 to 2015 (0.8 to per 100,000 population per year, 720 notifications; Table 3.7.1). The number of notifications increased from 252 in 2016, to 380 in 2017 (Appendix B), before declining to 281 in 2018. The monthly median number of notifications over the 2016–2018 period was 23 (range: 9–67 notifications per month).

There were marked increases, compared with the previous 2012–2015 reporting period, in notification rates caused by serogroups W (0.47 per 100,000 population per year [347 cases; 38% of all cases] in 2016–2018, vs 0.07 per 100,000 population per year [69 cases; 10% of all cases] in 2012–2015) and Y (0.22 per 100,000 population per year [159 cases; 17% of all cases] in 2016–2018, vs 0.07 per 100,000 population per year [69 cases; 10% of all cases] in 2012–2015)
year [65 cases; 9% of all cases] in 2012–2015) reported over the three years. Serogroup B accounted for 38% of all notifications (348 cases; 0.47 per 100,000 population per year) in this reporting period, compared with 71% (512 cases; 0.55 per 100,000 population per year) in the previous reporting period. For all three serogroups, notifications increased between 2016 (B: 92; W: 108; Y: 40) and 2017 (B: 137; W: 139; Y: 74), and then declined in 2018 (B: 119; W: 100; Y: 45). Serogroup C accounted for only 2% of cases (21/913; 0.03 per 100,000 population per year) during the reporting period, with these mainly due to an outbreak of serogroup C in 2017 in Victoria (14 cases). In 2018, there were two cases of serogroup E disease, a rare serogroup that has never previously been reported in Australia. There were no notifications for serogroup A and X during this reporting period.

Severe morbidity and mortality

In the three-year reporting period, hospitalisations with an ICD-10 code of meningococcal infection were associated with a total of 8,659 hospital bed days. The median length of stay was six days, varying from three days in 5–14 year olds to eight days in those aged 65 years and older (Table 3.7.1). There were 51 deaths due to meningococcal infection (of which 46, or 90%, were recorded as the underlying cause) recorded in the causes of death data over the three-year reporting period, an average annual rate of 0.07 deaths per 100,000 population (Table 3.7.1), almost double that reported for the 2012–2015 period (0.04 per 100,000 population per year), but lower than the number of deaths recorded in NNDSS (60, 0.08 per 100,000 population per year). The overall case fatality rate from NNDSS data was 6.6% (60/913), higher for serogroup W (10.1%, 35/347) than serogroup B (4.3%, 15/348) and Y (4.4%, 7/159).

Age and sex distribution

Infants aged < 1 year had the highest notification rate for invasive meningococcal disease (11.0 per 100,000 population per year), decreasing to 3.0 per 100,000 population per year in children aged 1–4 years (Table 3.7.1) and with a smaller peak in adolescents and young adults (aged 15–24 years) of 2.2 per 100,000 population per year. Rates of serogroup B meningococcal diseases were higher than those of serogroup W among infants < 1 year old (5.81 vs 3.55 per 100,000 population per year), children aged 1–4 years (1.40 vs 1.21 per 100,000 population per year), and adolescents and young adults aged 15–24 years (1.15 vs 0.56 per 100,000 population per year). In contrast, among adults aged 65 years and older, the rate of serogroup B cases was lower than those of serogroups W and Y (0.23, 0.69 and 0.45 per 100,000 population per year for serogroups B, W and Y, respectively). Of the small number of serogroup C cases, 13/21 (62%, 0.05 per 100,000 population per year) were aged 25–49 years. Overall, numbers of notifications for 2016–2018 were similar in males (n = 455) and females (n = 457).

The case fatality rate, calculated from NNDSS data, was highest for adults aged 65 years and older (22%, 12/60), lowest for infants aged < 1 year (8%, 5/102) and children 1–4 years (8%, 5/112) and intermediate for adolescents and young adults aged 15–24 years (13%, 14/211).

Geographical distribution

Notification rates of invasive meningococcal disease increased in every state and territory from 2016 to 2017 and then declined in 2018, with the exception of the Australian Capital Territory (where notification rates remained stable throughout this period) (Appendix B). Rates were highest in the Northern Territory (Appendices B and C), largely due to an outbreak of serogroup W disease in 2017 among Aboriginal communities in central Australia. Serogroup W accounted for 82% of cases in the Northern Territory (33/45), an average annual rate of 4.46 per 100,000 compared with the national rate of 0.47 per 100,000. Serogroup B accounted for 73% of all notifications in South Australia (71/97), an average annual rate of 1.37 per 100,000 compared with the national rate of 0.47 per 100,000.
Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for 901 of the 913 notifications (99%) of invasive meningococcal disease during 2016–2018, with 12 not stated (1%). Of the 913 notifications, 133 (15%) were in Aboriginal and Torres Strait Islander people, with 82/133 (62%) in children aged < 5 years and 27/133 (20%) among children aged 5–14 years, whereas children aged < 5 years comprised only 17% (133/780) of notifications among other Australians. A greater proportion of cases among other Australians were aged 15–24 years (26%, 205/780) than among Aboriginal and Torres Strait Islander people (4.5%, 6/133), but all-age Aboriginal and Torres Strait Islander notification rates were fivefold higher than rates in other Australians (5.4 vs 1.1 per 100,000 population per year). Aboriginal and Torres Strait Islander people accounted for 14% of all meningococcal deaths recorded in the causes of death data, 4.6 times that of other Australians (0.29 vs 0.06 per 100,000 population per year).

Vaccination status

The completeness of vaccination data improved to 71% (650/913 cases) in the 2016–2018 reporting period, compared to 42% in 2012–2015. Of the 650 cases with vaccination data available, 445 (68%) had received no meningococcal vaccine doses and 205 (32%) had 1 or more doses. Of the 205 previously vaccinated, 189 had received 1 dose, 12 had received 2 doses, and four had received 3 or more.

Comments

Notifications for invasive meningococcal disease increased in 2016 and 2017, driven by serogroup W and Y disease, before declining in 2018. In recent years, cases of meningococcal serogroup W disease caused by strains that are close variants of the virulent ST-11 clone have risen in several regions of the world (namely Australia, Europe, South America and parts of sub-Saharan Africa), and have been associated with atypical clinical presentations (particularly pneumonia and gastrointestinal symptoms), greater disease severity and higher case fatality rates. An outbreak of invasive meningococcal disease caused by serogroup W in the latter half of 2017, primarily among young Aboriginal children (< 10 years of age) in remote communities highlighted the previously-documented disparity in rates of meningococcal disease among Aboriginal and Torres Strait Islander people. A small decline in disease rates in 2018 followed the introduction of several jurisdictionally-funded quadrivalent conjugate meningococcal (MenACWY) vaccination programs, which varied in their target age group, duration and mode of delivery. A nationally-funded MenACWY vaccination program commenced for young children aged 12 months in July 2018, and subsequently for adolescents in 2019. The continued predominance of serogroup B disease in South Australia prompted the introduction of a jurisdictionally-funded meningococcal B vaccination program for young children in October 2018 and for adolescents in February 2019.
Figure 3.7.1: Meningococcal disease notifications and hospitalisations, Australia, 1993 to 2018,\(^a\) by month of diagnosis or admission

Table 3.7.1: Meningococcal disease notifications, hospitalisations and deaths, Australia, 2016 to 2018,\(^a\) by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications</th>
<th>Hospitalisations</th>
<th>LOS(^b) per admission</th>
<th>Deaths(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Rate(^d)</td>
<td>n</td>
<td>Rate(^d)</td>
</tr>
<tr>
<td>&lt; 1</td>
<td>102</td>
<td>10.98</td>
<td>150 16.15</td>
<td>130 13.99</td>
</tr>
<tr>
<td>1–4</td>
<td>112</td>
<td>2.95</td>
<td>188 4.95</td>
<td>160 4.22</td>
</tr>
<tr>
<td>5–14</td>
<td>49</td>
<td>0.53</td>
<td>97 1.06</td>
<td>83 0.90</td>
</tr>
<tr>
<td>15–24</td>
<td>271</td>
<td>2.20</td>
<td>257 2.68</td>
<td>225 2.34</td>
</tr>
<tr>
<td>25–49</td>
<td>144</td>
<td>0.56</td>
<td>194 0.75</td>
<td>148 0.57</td>
</tr>
<tr>
<td>50–64</td>
<td>135</td>
<td>1.03</td>
<td>188 1.43</td>
<td>135 1.03</td>
</tr>
<tr>
<td>≥65</td>
<td>160</td>
<td>1.41</td>
<td>213 1.87</td>
<td>133 1.17</td>
</tr>
<tr>
<td>All ages</td>
<td>913</td>
<td>1.24</td>
<td>1,287 1.74</td>
<td>1,014 1.37</td>
</tr>
</tbody>
</table>

\(^a\) Notifications where the month of diagnosis was between 1 January 1993 and 31 December 2018; hospitalisations where the month of admission was between 1 July 1993 and 31 December 2018.

\(^b\) LOS: length of stay in hospital.

\(^c\) Deaths include underlying and associated causes of deaths.

\(^d\) Average annual age-specific rate per 100,000 population.

\(^e\) Principal diagnosis (hospitalisations).
3.8 Mumps

**Highlights**

During the review period from 1 January 2016 to 31 December 2018, there was a major outbreak of mumps, predominantly affecting Aboriginal communities in Western Australia, the Northern Territory and Queensland. More than 64% of notifications (1,451/2,250) were in Aboriginal and Torres Strait Islander people, with at least 57% known to have received one or more doses of mumps-containing vaccine.

Mumps is an acute viral disease caused by a paramyxovirus. In the pre-vaccine era it was a well-known and common childhood viral disease. Mumps infection is systemic with variable pathology and symptomatology. The classical disease is characterised by fever and painful swelling and inflammation of one or more salivary glands, most commonly the parotid glands. Between 20% and 40% of cases, however, are subclinical. Aseptic meningitis can develop in up to 10% of cases, though long term sequelae are rare. Between 15% and 30% of post-pubertal males experienced epididymoorchitis in the pre-vaccination era, though this is less common (< 10%) in the post-vaccine era. Vaccination programs have resulted in substantial decreases in mumps incidence among young children, but a rise in incidence has been seen among older adolescents and young adults. Increasingly, outbreaks of mumps have been observed among adolescent and young adult populations with high coverage of 2 doses of mumps-containing vaccine and living in close contact.

**Case definition**

**Notifications**

Mumps case definition:


**Hospitalisations and deaths**

The ICD-10-AM/ICD-10 code B26 (mumps) was used to identify hospitalisations and deaths.

**Secular trends**

This reporting period saw the growth of a major outbreak of mumps that started in 2015 (Figure 3.8.1). In the three years from 1 January 2016 to 31 December 2018, there were 2,250 notification of mumps, an average annual notification rate of 3.1 per 100,000 population (Table 3.8.1), more than double that observed between 2012 and 2015 (1.3 per 100,000 population per year). The rate was 3.3 per 100,000 population per year in 2016 and 2017 and then declined slightly to 2.5 per 100,000 population in 2018 (Appendix B and Figure 3.8.1). From 1 January 2016 to 31 December 2018, there were 411 hospitalisations for ICD-10-AM/ICD-10 code B26 (mumps), an average annual rate of 0.6 per 100,000 population (Table 3.8.1), stable over the 3 years (0.53, 0.56 and 0.57 per 100,000 population in 2016, 2017 and 2018, respectively) (Appendix C).

**Severe morbidity and mortality**

Between January 2016 and December 2018, mumps hospitalisations accounted for 1,243 hospital bed days. The median length of stay was 1 day, but was higher among people aged over 65 years (5 days) and infants < 1 years (2.5 days) (Table 3.8.1). There were 1–5 deaths recorded.
in the causes of death data with mumps as any (underlying or associated) cause, all occurring among people aged ≥ 65 years (Table 3.8.1).

Age and sex distribution

Over the three-year reporting period, 88% of mumps notifications were in persons aged 5 to 49 years (23% among 5–14 year olds; 28% among 15–24 year olds; and 38% among 25–49 year olds). Notification rates were highest among 15–24 year olds (6.4 per 100,000 population per year) (Table 3.8.1). A similar trend was observed for hospitalisations, though a greater proportion of hospitalised cases were aged 65 years or older (who accounted for 14% of hospitalisations vs 3% of notifications) (Table 3.8.1). Since 2002, the notification rate of mumps for children aged < 5 years has remained low, mostly below 1 per 100,000 population per year. Over the three-year reporting period, notifications were evenly distributed among males and females (49% vs 51%); hospitalisations were slightly more common among females (53% vs 47% in males, male:female rate ratio of 0.9:1).

Geographical distribution

Notification and hospitalisation rates were highest in the Northern Territory, at more than 14 times and 7 times the national rates, respectively (Appendices B and C). Notification rates were also higher than the national average in Queensland and Western Australia, with all three jurisdictions affected by an outbreak predominantly affecting Aboriginal communities.

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for 2,150 of the 2,250 notifications (96%) of mumps during 2016–2018, with 100 not stated (4%). Almost two-thirds of notifications (64.5%, 1,451/2,250) were in Aboriginal and Torres Strait Islander people, with nearly all of these from Queensland (47%), Western Australia (29%) and the Northern Territory (21%). The notification rate among Aboriginal and Torres Strait Islander people was 53 times higher than that among other Australians (59.4 vs 1.1 per 100,000 population per year). Notified cases among Aboriginal and Torres Strait Islander people tended to be younger than cases among other Australians (29% of cases among Aboriginal and Torres Strait Islander people, vs 12% of cases among other Australians, were aged 5–14 years; 32% vs 20% were aged 15–24 years, 34% vs 44% were aged 25–49 years).

Vaccination status

Of the 2,250 notifications, vaccination data was available for 65% (1,462/2,250) of cases, with completeness higher for cases born from 1983 onwards (and hence eligible for vaccination at 12 months of age under the NIP – 1,281/1,621; 79%) than for those born up to 1982 (181/629, 29%). Of 1,462 notifications with vaccination data available, 170 (12%) had received no vaccine doses, including seven aged < 1 year and so not yet eligible for vaccination. Among the 1,292 cases (88%) recorded as vaccinated, 276 had received 1 dose, 933 had received 2 doses and 72 had received 3 or more doses (data on the number of doses received was missing for 11 vaccinated cases).

Comment

Previous mumps outbreaks in young adults in Australia have largely been concentrated in those who had received only one vaccine dose or were unvaccinated. This reporting period was marked by a series of large outbreaks of mumps, predominantly affecting young adults in Aboriginal communities in northern Australia of whom most (89% of cases aged 1–19 years) had at least two documented doses of mumps vaccine. A similar mumps outbreak was previously documented in 2007–2008 in the Kimberley. Other outbreaks among communities in close contact, where the force of infection is high, have been reported internationally, particularly among people who completed the vaccination series against mumps more than 10 years previously. The Advisory Committee on Immunization Practices in the USA has recommended a third dose of mumps
Figure 3.8.1. Mumps notifications and hospitalisations, Australia,\(^a\) 1993 to 2018,\(^b\) by month of diagnosis or admissions

- Note that the number of jurisdictions notifying mumps increased over the review period until July 1996 when mumps became notifiable in all states and territories. From July 1999 until June 2001, mumps was not notifiable in Queensland. Only the Australian Capital Territory, New South Wales and Victoria notified for the entire review period.

- Notifications where the month of diagnosis was between 1 January 1993 and 31 December 2018; hospitalisations where the month of admission was between 1 July 1993 and 31 December 2018.

Outbreak control strategies in Australia should take into consideration that vaccine effectiveness against the mumps component of measles-mumps-rubella (MMR)-containing vaccines is lower relative to the other antigens,\(^{154,169}\) seroprevalence is below the threshold for herd immunity,\(^{175}\) and waning may occur earlier following the shift of the second dose of mumps-containing vaccine from 4 years to 18 months of age.\(^{122}\)
Table 3.8.1: Mumps notifications, hospitalisations and deaths, Australia, 2016 to 2018,\(^a\) by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications</th>
<th>Hospitalisations</th>
<th>LOS(^b) per admission</th>
<th>Deaths(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Rate(^d)</td>
<td>n</td>
<td>Rate(^d)</td>
</tr>
<tr>
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</tr>
<tr>
<td>&lt; 1</td>
<td>8</td>
<td>0.86</td>
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<tr>
<td>1–4</td>
<td>77</td>
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<td>26</td>
<td>0.68</td>
</tr>
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<td>5–14</td>
<td>516</td>
<td>5.62</td>
<td>69</td>
<td>0.75</td>
</tr>
<tr>
<td>15–24</td>
<td>619</td>
<td>6.45</td>
<td>74</td>
<td>0.77</td>
</tr>
<tr>
<td>25–49</td>
<td>849</td>
<td>3.29</td>
<td>148</td>
<td>0.57</td>
</tr>
<tr>
<td>50–64</td>
<td>112</td>
<td>0.85</td>
<td>30</td>
<td>0.23</td>
</tr>
<tr>
<td>≥65</td>
<td>69</td>
<td>0.61</td>
<td>58</td>
<td>0.51</td>
</tr>
<tr>
<td>All ages</td>
<td>2,250</td>
<td>3.05</td>
<td>411</td>
<td>0.56</td>
</tr>
</tbody>
</table>

\(^a\) Notifications where the month of diagnosis was between 1 January 2016 and 31 December 2018; hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.

\(^b\) LOS: length of stay in hospital.

\(^c\) Deaths include underlying and associated causes of deaths.

\(^d\) Average annual age-specific rate per 100,000 population.

\(^e\) Principal diagnosis (hospitalisations).

\(^f\) np: not provided.
3.9 Pertussis

Highlights

Pertussis notification rates were lower over the three-year period from 1 January 2016 to 31 December 2018 than in the January 2012 to December 2015 period. Notification rates were highest in the 5–14 year age group (183.3 per 100,000 population per year) and hospitalisation rates were highest in infants aged < 1 year (31% of all pertussis hospital admissions; average annual rate 55.4 per 100,000 population).

Pertussis (whooping cough) is an acute respiratory illness, caused by the *Bordetella pertussis* bacterium. The illness begins with an irritating cough that gradually becomes paroxysmal and lasts for 1–2 months or longer. Paroxysms are characterised by repeated violent coughs and are followed by a characteristic crowing or high-pitched inspiratory whoop. The characteristic paroxysmal cough is more common in young children than in infants < 6 months of age, adolescents and adults, and is less common in people who are partially protected due to either vaccine-derived or national immunity.\(^{176,177}\)

The most common complication and cause of pertussis-related deaths is secondary bacterial pneumonia.\(^{177}\) Complications can occur in up to 6% of young children, and up to 24% of infants aged < 6 months.\(^{178}\) The greatest morbidity and mortality occur at the extremes of age, with the case fatality rate in unvaccinated infants aged < 6 months reported at 0.8%.\(^{176,179}\)

Case definition

Notifications\(^{180}\)

Pertussis case definition:


Hospitalisations and deaths

The ICD-10-AM/ICD-10 code A37 (whooping cough) was used to identify hospitalisations and deaths.

Secular trends

In the three years from 1 January 2016 to 31 December 2018, there were 44,918 cases of pertussis notified to the NNDSS. The average annual notification rate over the three-year period was 60.9 per 100,000 population, 19% lower than that in the four-year period 2012–2015 (76.0 per 100,000 population per year).\(^{12}\)

The notification rate was 83.1 per 100,000 population in 2016, which marked a decline from 2015 when a peak in notifications was observed (Figure 3.9.1). The rate declined by 40% to 49.7 per 100,000 population in 2017 and was similar in 2018 (50.4 per 100,000 population; Appendix B). Seasonal peaks were seen from approximately July to December. (Figure 3.9.1).

There were 1,807 hospital admissions coded as pertussis between 2016 and 2018. Hospitalisation rates in this three-year reporting period were lower than the previous four-year reporting period between 2012 and 2015 (average annual rate 2.4 vs 3.2 per 100,000 population, respectively). The annual hospitalisation rate declined from 3.1 per 100,000 population in 2016 to 1.7 per 100,000 population in 2018 (Figure 3.9.1 and Appendix C).

Severe morbidity and mortality

Between 1 January 2016 and 31 December 2018, hospital admissions coded as pertussis accounted for a total of 7,775 hospital bed days with a median length of stay of two days, higher in people aged 65 years and older (Table 3.9.1). In the three-year reporting period, eight deaths were recorded in the causes of death data (of which five were recorded with pertussis as the
underlying cause), all of which were in either infants aged < 1 year or adults aged 65 years or older (Table 3.9.1).

**Age and sex distribution**

Just over half of all notifications (51%) in the three-year reporting period occurred among children < 15 years of age, with the highest rate among 5–14 year olds (183.3 per 100,000 population per year) followed by infants aged < 1 year (142.2 per 100,000 population per year). Among adult age groups, notifications declined with increasing age, and were lowest among people aged 65 years and older (32.3 per 100,000 population per year) (Table 3.9.1). Hospitalisation rates were highest among infants aged < 1 year (55.4 per 100,000 population per year), with infants accounting for 31% of all hospitalisations. Adults aged 65 years and older comprised 26% of hospitalisations, but the hospitalisation rate (4.2 per 100,000 population per year) was tenfold lower than in infants. A greater proportion of notifications were female (55%) than male (45%) (male:female rate ratio of 0.85:1), while hospitalisations were more evenly distributed (male:female rate ratio 0.9:1).

**Geographical distribution**

Pertussis notifications declined across all states and territories from 2016 to 2018, but notifications in some states and territories rose slightly between 2017 and 2018 (Appendix B). Over the three-year period, notification rates were highest in New South Wales (95.2 per 100,000 population per year), followed by South Australia (85.2 per 100,000 population per year) and the Australian Capital Territory (83.7 per 100,000 population per year) (Appendix B). The highest
hospitalisation rate for pertussis occurred in the Northern Territory (4.5 per 100,000 population per year) followed by South Australia (4.0 per 100,000 population per year) (Appendix C).

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for 25,853 of the 44,938 notifications (58%) of pertussis during 2016–2018, with 19,085 not stated (42%). There was a high level of completeness in Western Australia (96%), the Northern Territory (96%), the Australian Capital Territory (95%) and South Australia (93%), with completeness lowest in New South Wales (40%) and Victoria (51%) and intermediate in other jurisdictions (Queensland 66%, Tasmania 74%). However, overall completeness of Aboriginal and Torres Strait Islander status was high for young children (96% for those aged < 1 year and 92% for those aged 1–4 years). Notification rates in Aboriginal and Torres Strait Islander children were higher in infants (310.4 vs 131.2 per 100,000 population per year [rate ratio 2.4:1]) and children aged 1–4 years (152.0 vs 128.9 per 100,000 population per year [rate ratio 1.2:1]).

Of the 1,807 hospitalisations, 9.5% (172 cases) were in Aboriginal and Torres Strait Islander people. The hospitalisation rate in Aboriginal and Torres Strait Islander people was higher than in other Australians (7.0 vs 2.3 per 100,000 population per year, a ratio of 3.1:1).

Vaccination status

Of the 44,938 notifications, vaccination data was available for only 40% (17,754/44,938), with 2,643 of these (15%) having no recorded vaccine doses. In comparison, the completeness of vaccination data was far improved for children aged < 5 years, with vaccination data available for 95% (5,979) of the 6,267 cases. Among all notifications, 15,111 had received a pertussis-containing vaccine, of whom 8,758 had received 1 dose; 267 had received 2 doses; 1,585 had received 3 doses; 3,711 cases had received 4 doses; 487 had received 5 doses; and 59 had received 6 or more doses of pertussis-containing vaccine. Data on the number of doses received was unavailable for 244 vaccinated cases. Among the 5,979 cases aged < 5 years, 786 (13%) had not received any doses of pertussis-containing vaccine; 3,312 had received 1 dose; 154 had received 2 doses; 1,246 had received 3 doses; 431 had received 4 doses; and 3 each received 5 and 6 doses (data on the number of doses received was unavailable for 44 vaccinated cases).

Comments

Rates of pertussis notifications and hospitalisations declined between 2016 and 2018 following a spike in cases in 2015. Reintroduction of the 18-month booster dose of pertussis-containing vaccine in 2016,181,182 and rising coverage of pertussis-containing vaccine in pregnancy (funded by jurisdictions from 2015 and then under the NIP in 2018),183–185 may have contributed to this decline. Although severe disease predominantly occurs in unvaccinated young infants, high healthcare utilisation has also been reported among older adults with pertussis, with burden of disease in older adults believed to be underestimated.186,187
Table 3.9.1: Pertussis notifications, hospitalisations and deaths, Australia, 2016 to 2018, by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications</th>
<th>Hospitalisations</th>
<th>LOS per admission</th>
<th>Deaths^c</th>
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<td>Any diagnosis</td>
<td>Principal diagnosis</td>
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<td>50–64</td>
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<td>214 1.9</td>
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<td>All ages</td>
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<td>60.9</td>
<td>1,807 2.4</td>
<td>1,036 1.4</td>
</tr>
</tbody>
</table>

a Notifications where the month of diagnosis was between 1 January 2016 and 31 December 2018; hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.
b LOS: length of stay in hospital.
c Deaths include underlying and associated causes of deaths.
d Average annual age-specific rate per 100,000 population.
e Principal diagnosis (hospitalisations).
f np: not provided.
g Total for notifications includes 22 cases without data on age of onset.
### 3.10 Pneumococcal disease

#### Highlights

Notification rates for invasive pneumococcal disease were slightly higher in the three-year period 2016–2018 than in the previous four years (2012–2015), as were hospitalisations for pneumococcal disease. Adults aged ≥ 65 years accounted for a large proportion of notifications and hospitalisations and the majority of deaths.

Pneumococcal disease is caused by the bacterium *Streptococcus pneumoniae* (pneumococcus). There are over 90 different serotypes of pneumococcus. Pneumococci can be isolated from the upper respiratory tract in adults and, more often, in children, and can spread directly from the nasopharynx to the respiratory tract, which may cause otitis media, sinusitis or pneumonia. Pneumococci are also able to enter the bloodstream to cause invasive disease, which may manifest as meningitis, pneumonia, septicaemia without focal infection or, less commonly, infection of other sites such as pleural, peritoneal or joint fluid. Invasive pneumococcal disease (IPD) is diagnosed by detecting *S. pneumoniae* from blood, CSF or other sterile site. In the absence of a sterile site isolate, a presumptive diagnosis of pneumococcal pneumonia may be based on a sputum isolate of *S. pneumoniae* and/or or clinical features such as chest X-ray appearance or prompt response to antibiotic therapy. Children with comorbid conditions have higher rates of pneumococcal disease and higher case fatality rates than do otherwise healthy children. While pneumococcal conjugate vaccines have substantially reduced the incidence of pneumococcal disease caused by vaccine serotypes, there is evidence of an increase in disease incidence due to non-vaccine serotypes in some settings.

#### Case definition

**Notifications**

Pneumococcal disease (invasive) case definition:


**Hospitalisations and deaths**

The ICD-10-AM/ICD-10 codes G00.1 (pneumococcal meningitis) and A40.3 (pneumococcal septicaemia) are each considered to be a proxy for IPD; these codes, and J13 (pneumococcal pneumonia), which is less strongly associated with IPD, were used to identify pneumococcal disease hospitalisations and deaths.

#### Secular trends

In the three years from 1 January 2016 to 31 December 2018, notification patterns of IPD remained relatively steady, with marked seasonality noted for notifications and hospitalisations (Figure 3.10.1). Overall, there were 5,742 notifications of IPD over the three-year reporting period at an average annual rate of 7.8 per 100,000 population (Table 3.10.1), an increase of 13% on the average notification rate for the previous four-year reporting period (6.9 per 100,000 population). The annual number of notifications increased from 1,498 in 2015 to 1,664 in 2016, to 2,050 in 2017, and decreased slightly to 2,028 in 2018 (Appendix B).

There has been an increasing trend in hospital admissions with ICD codes of interest since 2006 (Figure 3.10.1). Between 1 January 2016 and 31 December 2018, there were 13,297 hospitalisations coded as pneumococcal meningitis, septicaemia or pneumonia, an average annual rate of 18.0 per 100,000 population (Table
3.10.1), which was 21% higher than the average annual hospitalisation rate in the 2012–2015 period (14.9 per 100,000 population). Of the 13,297 hospitalisations, 3,081 (23.2%) were for presumed IPD (pneumococcal meningitis and/or septicaemia) and 10,216 (76.8%) for pneumococcal pneumonia (Table 3.10.1). The seasonal peak in admissions for pneumococcal disease was highest in 2017, reaching 671 hospitalisations in September 2017 (Figure 3.10.1).

Severe morbidity and mortality

Between 1 January 2016 and 31 December 2018, admissions with pneumococcal disease codes accounted for 97,021 hospital bed days (an average of 32,340 bed days per year), higher than the 29,774 average bed days per year between 2012 and 2015. The median length of stay for hospitalisations coded as pneumococcal meningitis or septicaemia (7 days) was higher than for hospitalisations coded as pneumonia (5 days) (Table 3.10.1).

For the three years from 1 January 2016 to 31 December 2018, there were 159 deaths (average annual rate 0.2 per 100,000 population) recorded in the causes of death data with pneumococcal disease-related ICD-10 codes (A40.3, G00.1 and J13) as the underlying or associated cause of death (Table 3.10.1), of which 82 (52%) were recorded as deaths with pneumococcal disease being the underlying cause. The 159 deaths recorded in the causes of death data was less than half the number of deaths recorded in the NNDSS database (452; average annual rate 0.6 per 100,000 population). The recorded in-hospital death rate among hospitalisations recorded in the AIHW data was almost three times as high for pneumococcal meningitis and septicaemia (8.6%) than for pneumococcal pneumonia (3.0%).

Age and sex distribution

For the three-year period from 1 January 2016 to 31 December 2018, the IPD notification rate was highest in infants aged < 1 year, followed by adults aged ≥ 65 years and children aged 1–4 years (Table 3.10.1). differing from the previous four-year period when the highest incidence was in older adults followed by infants.

Rates of IPD notifications among children aged < 5 years increased between 2016 and 2017, and then stayed steady in 2018. Notification rates in infants aged < 1 year increased from 18.8 per 100,000 population (n = 60) in 2016 to 26.4 per 100,000 population (n = 81) in 2017, and in children aged 1–4 years from 13.9 (n = 174) to 17.1 (n = 217) per 100,000 population per year. The rate in adults aged ≥ 65 years increased from 16.6 per 100,000 population (n = 611) in 2016 to 20.9 per 100,000 population (n = 793) in 2017, with smaller increases for other age groups.

The rate of hospitalisation for pneumococcal disease was highest in adults aged ≥ 65 years (55.0 per 100,000 population per year), comprising almost half (47.1%) of all hospital admissions for pneumococcal disease. For pneumococcal meningitis or septicaemia, the highest rate of hospitalisation was in infants aged < 1 year (12.3 per 100,000 population per year), whereas for pneumococcal pneumonia the highest rate was in adults aged ≥ 65 years (43.4 per 100,000 population per year). Hospitalisations for pneumococcal disease were slightly more common among males than females (18.9 vs 17.2 per 100,000 population per year respectively, rate ratio 1.1:1).

Death rates based on the causes of death data were highest among adults aged ≥ 65 years, followed by adults aged 50–64 years (Table 3.10.1). The recorded in-hospital death rate among hospital admissions for pneumococcal disease was highest among adults aged ≥ 65 years (6.0%).

Geographical distribution

The Northern Territory had the highest rates of notifications of IPD (average annual rate 22.4 per 100,000 population), and hospitalisations coded as pneumococcal disease (average annual rate 66.1 per 100,000 population), across the three-year period 2016–2018. Notification
Figure 3.10.1: Pneumococcal disease notifications and hospitalisations, Australia, 1998 to 2018, \(^a\) by month of diagnosis or admission

\(^a\) Notifications where the month of diagnosis was between 1 January 2001 and 31 December 2018; hospitalisations where the month of admission was between 1 January 1998 and 31 December 2018. Hospitalisations include pneumonia, meningitis and septicaemia.

and hospitalisation rates in the other states and territories did not differ appreciably from the national average (Appendices B and C).

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for 5,260 of the 5,742 notifications of IPD (92%) during 2016–2018, with 482 not stated (8%). Of the total notified cases (5,742), there were 644 cases (11%) in Aboriginal and Torres Strait Islander people, with 36% of these aged between 25–49 years. Notification rates were almost four times higher in Aboriginal and Torres Strait Islander people than in other Australians (26.4 versus 7.1 per 100,000 population per year); among those aged 15–24 and 25–49 years this disparity was more than ninefold. Aboriginal and Torres Strait Islander people accounted for 1,548/13,297 (11.6%) of total hospitalisations for this reporting period, with 40.2% of these (622) in adults aged 25–49 years; 31.5% (487) in adults aged 50–64 years; and 15.1% (233) in adults aged \(\geq\) 65 years. Aboriginal and Torres Strait Islander hospitalisation rates were almost four times higher than those for other Australians (63.4 vs 16.5 per 100,000 population per year; rate ratio 3.8).

Mortality attributed to pneumococcal disease (as either the underlying or associated cause of death) in the causes of death data was 1.6 times greater among Aboriginal and Torres Strait Islander people than in others (0.3 vs 0.2 per 100,000 population per year). The death rate among Aboriginal and Torres Strait Islander people was higher when calculated based on
NNDSS data (1.4 vs 0.6 per 100,000 population per year) and AIHW hospitalisation data (1.8 vs 0.7 per 100,000 population per year).

Vaccination status

Of the total 5,742 notifications, vaccination data was available for 75% of cases (4,294). Of 840 notifications in children aged < 5 years, 818 (97%) had vaccination data available, of whom 79 (10%) had received no vaccine doses and 739 (90%) had received at least one dose of pneumococcal vaccine. Among the 739 vaccinated cases, 103 had received 1 dose; 47 had received 2 doses; 554 had received 3 doses; 32 had received 4 doses; and three had received 5 doses of pneumococcal vaccine.

Of the 2,188 notifications in adults aged ≥ 65 years, vaccination data was not available for 19% (426). Of the 1,762 cases aged ≥ 65 years for whom vaccination data was available, 951 cases (54%) were unvaccinated and 811 (46%) had received at least one dose of pneumococcal vaccine. Of the vaccinated cases, 554 had received 1 dose; 222 had received 2 doses; 33 had received 3 doses; and two had received 4 doses of pneumococcal vaccine.

Of the 243 cases in Aboriginal and Torres Strait Islander adults aged ≥ 50 years, 92% (224) had vaccination data available, of whom 83 cases (37%) had no recorded vaccine doses and 141 (63%) had received at least one dose of pneumococcal vaccine. Of the 141 vaccinated cases, 45 had received 1 dose; 46 had received 2 doses; 37 had received 3 doses; and 13 had received 4 or more doses of pneumococcal vaccine.

Serotype of notifications

Among the 5,742 IPD notifications, 32% (1,852) were due to a serotype contained in the 13-valent pneumococcal conjugate vaccine (PCV13) and 62% (3,552) were due to a serotype contained in the 23-conjugate pneumococcal polysaccharide vaccine (PPV23); 1,712 (30% of total cases) were due to a serotype contained in PPV23 but not in PCV13. Another 30% of all cases (1,716) were due to serotypes not included in PPV23 and 8% (462) were of unknown serotype. Among cases aged < 5 years who had received at least one prior dose of pneumococcal vaccine, approximately 34% had a PCV13 serotype, compared with 41% of unvaccinated cases.

Comment

From 1 January 2005, all children born from January 2003 were eligible for a full course of pneumococcal conjugate vaccine (then 7-valent) funded on the NIP. Since 2005, there have been large reductions in notifications of vaccine type IPD in Australia. For the three-year reporting period 2016–2018, rates of IPD increased slightly, with marked seasonality observed in notified and hospitalised cases. In-depth examination of the epidemiology of pneumococcal disease in Australia has found that reductions in vaccine-type IPD post-PCV13 introduction were inferior in Australia than the reductions seen in countries providing a booster dose at 12 months of age, with more breakthrough cases among fully-vaccinated children. Subsequently, the NIP schedule for PCV13 for infants was updated from a 3+0 schedule to a 2+1 schedule, i.e. 2 doses in infancy and a booster dose at 12 months of age, with the expectation that this will induce longer-lasting protection against pneumococcal disease. Despite the presence of a nationally-funded pneumococcal vaccination program, uptake of pneumococcal vaccine slowly declined between 2011 and 2016 among adults aged ≥ 65 years, and a substantial proportion (54%) of cases in this age group in this reporting period were unvaccinated. The disparity in pneumococcal disease between Aboriginal and Torres Strait Islanders and others continues to persist and warrants consideration of differential vaccination policies to reduce this gap.
Table 3.10.1: Pneumococcal disease notifications, hospitalisations and deaths, Australia, 2016 to 2018,\(^a\) by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications</th>
<th>Hospitalisations</th>
<th>LOSb per admission</th>
<th>Deaths(^c)</th>
</tr>
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<td></td>
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<td>Rate(^d)</td>
<td>n* Rate(^d,e)</td>
<td>n Rate(^d)</td>
</tr>
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<td>301 7.93</td>
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<td>67 0.73</td>
<td>147 1.60</td>
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<td>1.55</td>
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<td>174 1.81</td>
</tr>
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<td>3.98</td>
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<td>417 3.18</td>
<td>17,44 13.29</td>
</tr>
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<td>19.23</td>
<td>663 5.83</td>
<td>3,485 30.63</td>
</tr>
<tr>
<td>All ages</td>
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<td>7.78</td>
<td>1,648 2.23</td>
<td>7,316 9.92</td>
</tr>
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</table>

\(^a\) Notifications where the month of diagnosis was between 1 January 2016 and 31 December 2018; hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.

\(^b\) LOS: length of stay in hospital.

\(^c\) Deaths sourced from the Causes of Death database from the Australian Coordinating Registry. Deaths include underlying and associated causes of deaths.

\(^d\) Average annual age-specific rate per 100,000 population.

\(^e\) Pneumococcal meningitis or septicaemia (proxy for invasive pneumococcal disease).

\(^f\) Pneumococcal pneumonia.
### 3.11 Poliomyelitis

#### Highlights

There were no notifications or deaths due to poliomyelitis between January 2016 and December 2018.

Poliomyelitis (polio) is caused by an enterovirus, poliovirus. Infection involves the gastrointestinal tract, and may progress to the nervous system, resulting in paralysis. Acute flaccid paralysis (AFP) occurs in fewer than 1% of infections. More than 90% of infections are asymptomatic, with a minor illness characterised by fever, headache, malaise and nausea/vomiting occurring in about 10%. The maximum extent of paralysis is usually reached within 3–4 days of disease onset. Any paralysis still present after 60 days is likely to be permanent. In the pre-vaccine era, poliovirus was the leading cause of permanent disability in children. Efforts to eradicate polio have resulted in the eradication of two of three wild poliovirus (WPV) serotypes, with type 2 certified as eradicated in 2015 and type 3 in October 2019. Transmission of WPV type 1 remains uninterrupted only in Afghanistan and Pakistan, with 175 cases reported in 2019.

Vaccine-associated paralytic poliomyelitis (VAPP) is acute flaccid paralysis due to a Sabin-like poliovirus (i.e. a virus similar to that used in the Sabin live attenuated oral poliovirus vaccine [OPV]). A vaccine-derived poliovirus (VDPV) is defined as having 1–15% nucleic acid sequence variation from the prototype Sabin strain. The variation is due to long-term (more than 1 year) virus replication after the administration of OPV. Virus replication may occur in an individual with an immunodeficiency (iVDPV) or through sustained person-to-person transmission in areas with low OPV coverage (circulating or cVDPV). Any VDPVs not clearly assigned to either of these categories are known as ambiguous VDPVs (aVDPV).

Low vaccination coverage is a major risk factor for cVDPV outbreaks and has been observed in under-vaccinated communities in developed countries.

#### Case definition

**Notifications**

Poliovirus infection:


**Hospitalisations and deaths**

The ICD-10-AM/ICD-10 code A80 (acute poliomyelitis) was used to identify hospitalisations and deaths. Note: This code includes VAPP and specific codes for Aboriginal and Torres Strait Islander and imported wild-type poliovirus infection. Sequelae of poliomyelitis (ICD-10 code B91) were not included in these analyses.

There were no acute poliomyelitis notifications or deaths reported during the 3-year period 1 January 2016 to 31 December 2018. There were 1–4 hospitalisations with a principal diagnosis of A80.9 (acute poliomyelitis, unspecified); however, hospitalisations for rare diseases such as poliomyelitis should be interpreted with caution due to possible misclassification, coding and data related issues.

Australia was declared polio-free in 2000, with the only case of polio since 1987 reported in 2007 in an overseas-born student who acquired the disease during a visit to a country with ongoing polio transmission. Regular case review is undertaken by the Polio Expert Panel (PEP) and existing polio surveillance strategies are considered appropriate for Australia. In the annual PEP reports for the three years 2016 to 2018, no cases of poliomyelitis were identified from clinical surveillance, with between 1.24 and 1.38 non-polio AFP cases per 100,000 children less than 15 years of age, exceeding the
WHO AFP surveillance performance criterion of 1 case of non-polio AFP per 100,000 children. While Australia has regularly met national AFP surveillance performance indicators, a recent study identified local areas with AFP surveillance detection rates lower than expected. Serosurveillance in 2012 and 2013 found that over 80% of the Australian population were seropositive for poliovirus type 1 and type 2; lower levels of seropositivity against poliovirus type 3 (67%) are not a major concern given that wildtype poliovirus type 3 has not been detected worldwide since November 2012.
3.12 Q fever

Highlights

The rate of Q fever notifications and hospitalisations observed between 2016 and 2018 was largely unchanged from the previous reporting period between 2012 and 2015. Notification and hospitalisation rates continue to be more than threefold higher among males compared to females, and remain highest for adults aged 50–64 years.

Q fever is a zoonotic disease caused by *Coxiella burnetii*. It has been identified in a wide range of wild and domestic animal hosts including arthropods, birds, rodents, marsupials and livestock, but the most important reservoirs as a source for human infections are cattle, sheep and goats. Humans become infected primarily by inhaling aerosols contaminated by *C. burnetii*. Occupations with higher exposure risks include abattoir and farm workers and veterinarians. Windborne spread and indirect exposures in a contaminated environment account for non-occupational infections.

High proportions, up to 60%, of persons infected with *C. burnetii* are asymptomatic during primary infection or only experience a self-limiting febrile illness. Q fever may present with acute or chronic clinical manifestations. There is increasing acceptance of an association with long-term sequelae, in particular the post Q fever fatigue syndrome which develops in up to 20% of people with acute Q fever. The most common complication in adults is subacute endocarditis. Infection in pregnancy can result in fetal death, intrauterine growth restriction and premature delivery. Although men and women are similarly exposed to the pathogen, epidemiological studies have shown that Q fever symptoms are more than twice as frequent in men than in women.

Australia has a highly effective licensed Q fever vaccine (Q-VAX; CSL Limited) that requires pre-vaccination screening tests. A time-limited nationally funded vaccination program for at-risk populations between 2001 and 2006 resulted in a decline in Q fever notifications by more than 50% to the lowest levels on record in 2005 and 2006. This vaccine is currently not recommended for children aged less than 15 years due to lack of data on safety and efficacy; however, a clinical trial of Q fever vaccination in children aged 10–15 years is underway.

Case definition

Notifications

Q fever case definition:


Hospitalisations and deaths

The ICD-10-AM/ICD-10 code A78 (Q fever) was used to identify hospitalisations and deaths.

Secular trends

From 1 January 2016 to 31 December 2018, there were 1,552 notifications of Q fever (average annual rate 2.1 per 100,000 population) (Table 3.12.1), and 766 hospital admissions (average annual rate 1.0 per 100,000 population) (Table 3.12.1). The average annual rate of notifications and hospitalisations was largely unchanged in this reporting period compared with the previous 4-year reporting period between 2012 and 2015 (2.1 per 100,000 notifications and 0.9 per 100,000 hospitalisations).

Severe morbidity and mortality

There were 4,502 bed days associated with hospital admissions coded as due to Q fever over the period 1 January 2016 to 31 December 2018. The median length of stay was 4 days, but...
increased with increasing age (Table 3.12.1). Q fever was the principal diagnosis in 71.5% of these admissions (Table 3.12.1). Over the 3-year reporting period, 1–5 deaths were recorded as due to Q fever.

Age and sex distribution

The highest age-specific notification and hospitalisation rates for Q fever were in adults aged 50–64 years (4.2 per 100,000 population per year for notifications and 1.9 per 100,000 population per year for hospitalisations) (Table 3.12.1). Notification rates were also relatively high among adults aged 25–49 years (2.3 per 100,000 population per year) and those aged ≥65 years (2.1 per 100,000 population per year). There were more notifications and hospitalisations in males than females (male:female rate ratio 3.3:1 for notifications and 3.1:1 for hospitalisations). The highest gender-specific rates of notifications and hospitalisations were in males aged 50–64 years (6.5 and 3.0 per 100,000 population per year, respectively).

There were no notifications among infants aged <1 year. Among children and young adults, notification rates were low in the 1–4 years (0.2 per 100,000 population per year) and 5–14 years (0.4 per 100,000 population per year) age groups, but were higher among 15–24 year olds (1.4 per 100,000 population per year).

Geographical distribution

The largest numbers of notifications for Q fever occurred in Queensland (43%; 4.5 per 100,000 population per year) and New South Wales (43%; 2.8 per 100,000 population per year) (Appendix B). Trends for hospitalisations were similar to notifications, with the majority of hospitalisations occurring in Queensland (50%) and New South Wales (33%).

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for 1,484 of the 1,552 notifications of Q fever (96%) during 2016–2018, with 68 not stated (4%). Of the 1,552 notifications, 4.3% (n = 66) were in Aboriginal and Torres Strait Islander people, with notification rates slightly higher than other people (2.7 vs 2.1 per 100,000 population per year; rate ratio 1.3:1). The age distribution of Q fever notifications among Aboriginal and Torres Strait Islander people was younger, with a greater proportion aged 15–24 years relative to other Australians (15–24 years: 24.2% vs 7.7%; 25–49 years: 47.0% vs 37.3%; 50–64 years: 24.2% vs 36.0%; ≥65 years: 1.5% vs 16.2%).

Vaccination status

Vaccination status data were not available.

Comment

Despite some variation over the three years between 1 January 2016 and 31 December 2018, notifications and hospitalisations of Q fever were largely similar to rates during the previous four-year period (2012–2015). The highest notification rates were among adults aged 50–64 years, followed by those aged 25–49 years, consistent with occupational exposure and similar to the previous reporting period, although notification rates for 2016–2018 were also relatively high among adults aged ≥65 years. Seroprevalence studies in New South Wales and Queensland have estimated seroprevalence to be between 3% and 5%. There is evidence of variance between metropolitan and non-metropolitan areas, likely linked to occupational exposure. While studies have noted the higher risk of Q fever among occupations involving contact with livestock, the lack of risk factor data limits the ability of routinely-collected notification data in informing policies regarding Q fever prevention and vaccination.
Figure 3.12.1: Q fever notifications and hospitalisations, Australia, 1993 to 2018,\(^a\) by month of diagnosis or admission

Table 3.12.1: Q fever notifications, hospitalisations and deaths, Australia, 2016 to 2018,\(^a\) by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications</th>
<th>Hospitalisations</th>
<th>LOS(^b) per admission</th>
<th>Deaths(^c)</th>
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<td>Rate(^d)</td>
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<td>1–4</td>
<td>8</td>
<td>0.21</td>
<td>10</td>
<td>0.26</td>
</tr>
<tr>
<td>5–14</td>
<td>35</td>
<td>0.38</td>
<td>19</td>
<td>0.21</td>
</tr>
<tr>
<td>15–24</td>
<td>131</td>
<td>1.36</td>
<td>50</td>
<td>0.52</td>
</tr>
<tr>
<td>25–49</td>
<td>585</td>
<td>2.27</td>
<td>249</td>
<td>0.97</td>
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<tr>
<td>50–64</td>
<td>551</td>
<td>4.20</td>
<td>251</td>
<td>1.91</td>
</tr>
<tr>
<td>≥ 65</td>
<td>242</td>
<td>2.13</td>
<td>187</td>
<td>1.64</td>
</tr>
<tr>
<td>All ages</td>
<td>1,552</td>
<td>2.10</td>
<td>766</td>
<td>1.04</td>
</tr>
</tbody>
</table>

\(^a\) Notifications where the month of diagnosis was between 1 January 1993 and 31 December 2018; hospitalisations where the month of admission was between 1 July 1993 and 31 December 2018.

\(^b\) LOS: length of stay in hospital.

\(^c\) Deaths include underlying and associated causes of deaths.

\(^d\) Average annual age-specific rate per 100,000 population.

\(^e\) Principal diagnosis (hospitalisations).

\(^f\) np: not provided.
3.13 Rotavirus

**Highlights**

Since the funding of rotavirus vaccine on the National Immunisation Program in 2007, there has been a substantial decrease in hospitalisations due to rotavirus. The average annual hospitalisation rate in the three-year period 2016–2018 was 5.0 per 100,000 population, lower than the 2012–2015 average annual rate of 5.4 per 100,000 population. In 2017, an outbreak of rotavirus resulted in a higher rotavirus hospitalisation rate of 8.6 per 100,000 population.

Rotavirus is a non-enveloped virus that is the major cause of acute gastroenteritis in young children and infants, and globally the leading cause of severe dehydrating diarrhoea in children aged < 5 years. Infection can be asymptomatic; can cause mild to moderate gastroenteritis; or can cause severe gastroenteritis with dehydration requiring hospitalisation. Virtually all children worldwide are infected with rotavirus by 5 years of age, but severe disease occurs most commonly in those aged 6 months to 2 years. However, disease does occur in all age groups. Rotaviruses are primarily spread by faecal–oral transmission. Infection with rotavirus confers some protection against subsequent serious disease. Rotaviruses are typed based on two surface proteins, VP7 (G protein) and VP4 (P protein). G1P[8] was the dominant genotype in Australia in the pre-vaccine era, with other common strains being G2, G3, G4 and G9. In recent years, following the introduction of the national rotavirus vaccination program, G12P[8] has become the dominant genotype nationally, but other genotypes (specifically G2P[4] and equine-like G3P[8]) have been more common since 2016, although genotype distribution has varied based on the vaccine implemented.

A national case definition for rotavirus was defined by the Communicable Diseases Network Australia (CDNA) in 2017. Rotavirus became a nationally notifiable disease from 1 July 2018. This report includes rotavirus notifications from January 2017, but focuses on notifications from July 2018 to December 2018.

**Case definition**

**Notifications**

Rotavirus case definition:


**Hospitalisations and deaths**

The ICD-10-AM/ICD-10 code A08.0 (rotavirus enteritis) was used to identify hospitalisations and deaths.

**Secular trends**

Rotavirus notifications have only been recorded as per the national definition since 1 January 2017; rotavirus became a nationally notifiable disease in July 2018. There was an increase in rotavirus notifications in mid-2017, with notifications peaking in September 2017, declining again by the end of 2017. Notifications remained low in 2018. A total of 7,267 and 2,881 notifications were recorded in 2017 and 2018, respectively, with 1,453 notifications reported between July and December 2018.

In the three years from 1 January 2016 to 31 December 2018, the number of monthly hospitalisations with an ICD-10-AM code for rotavirus enteritis was largely similar to the monthly average number in the previous reporting period (2012–2015) (Figure 3.13.1). However, a larger peak and higher median monthly cases were observed in 2017 due to an outbreak in the latter half of the year (median monthly cases: 65 in 2016, 160 in 2017 and 55 in 2018). As in previous years, seasonal peaks were seen during the winter months.
There were 3,673 hospital admissions over the three years with an ICD-10-AM code of rotavirus enteritis. The average annual hospitalisation rate was 5.0 per 100,000 population, slightly lower than in the 2012–2015 period (5.4 per 100,000 population per year) (Table 3.13.1). The majority of hospitalisations (2,112/3,673; 57.5%) occurred in 2017, peaking in September 2017 (n = 392).

Severe morbidity and mortality

During the three-year reporting period, rotavirus was recorded as the principal diagnosis in 60.9% of hospitalisations with rotavirus enteritis (2,236/3,673), accounting for 6,625 bed days (average 2,208 bed days per year) with a median length of stay of two days (Table 3.13.1). The length of stay for hospitalisations with rotavirus as any diagnosis was slightly higher at three days (Table 3.13.1).

There were 1–5 deaths recorded in the causes of death data for the three years from 1 January 2016 to 31 December 2018 with rotavirus enteritis (ICD-10 code, A08) as the underlying or associated cause of death; all were in adults aged ≥ 50 years and all occurred in 2017.

Age and sex distribution

Between July and December 2018, young children aged < 5 years comprised half of all rotavirus notifications (726/1,453), though rates were by far the highest among infants aged < 1 year (303.6 per 100,000 population per year) (Table 3.13.1). Notifications were relatively evenly distributed between males and females (male:female rate ratio of 0.95:1), although there were slightly more notifications observed among males in children aged < 5 years (male:female rate ratio of 1.1:1). During the three-year reporting period, rates of hospitalisations with an ICD-10-AM code for rotavirus enteritis as a principal diagnosis were highest among infants aged < 1 year (58.6 per 100,000 population per year), followed by children aged 1–4 years (17.7 per 100,000 population per year) (Table 3.13.1), with these age groups accounting for 54% of hospitalisations with a principal diagnosis of rotavirus enteritis. There were slightly fewer hospitalisations in males than females (male to female ratio 0.9:1). Trends were similar for hospitalisations with rotavirus as any diagnosis (Table 3.13.1).

Geographical distribution

Between July and December 2018, the rotavirus notification rate was highest in the Northern Territory (36.7 per 100,000 population per year), followed by South Australia (32.8 per 100,000 population per year), with Victoria recording the lowest notification rate (0.5 per 100,000 population per year) (Appendix B).

Over the three-year period 1 January 2016 to 31 December 2018, the Northern Territory recorded the highest rate of hospitalisations with rotavirus enteritis (29.7 per 100,000 population per year), representing nearly six times the Australian average (Appendix C), while South Australia recorded the second-highest rate (10.5 per 100,000 population per year). The Australian Capital Territory (1.3 per 100,000 population per year) and Victoria (1.5 per 100,000 population per year) recorded the lowest rates of hospitalisations.

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for 1,050 of the 1,453 rotavirus notifications (72%) between July and December 2018, with 403 not stated (28%). Of the 1,453 cases, 10.7% (156) were among Aboriginal and Torres Strait Islander people. The rate of rotavirus notifications was 3.5 times higher among Aboriginal and Torres Strait Islander people than among others. The majority of notifications among Aboriginal and Torres Strait Islander people (123/156, 79%) were in children aged < 5 years (57 [36.5%] in infants aged < 1 year and 66 [42%] in children aged 1–4 years). In comparison, 46.5% of notifications among
other Australians were aged < 5 years (374/1,297 [29%] aged < 1 year and 229/1,297 [18%] aged 1-4 years).

For the reporting period between 2016 and 2018, 13.1% (481) of the total 3,673 hospitalisations were recorded as being in Aboriginal and Torres Strait Islander people. The proportion of rotavirus hospitalisations that were reported to be in Aboriginal and Torres Strait Islander people increased over the reporting period, from 9.0% in 2016 to 13.4% in 2017 and 17.0% in 2018. Of these 481 hospitalisations, 85% (409) were in children aged < 5 years. The Aboriginal and Torres Strait Islander hospitalisation rate was more than four-fold higher than other Australians (19.7 per 100,000 population versus 4.8 per 100,000 population, respectively).

**Vaccination status**

Of the total 1,453 rotavirus notifications between July and December 2018, vaccination data was available for 830 cases (57%). Of these, 284/830 (34%) had received no vaccine doses and 546 (66%) had received at least one dose of rotavirus vaccine. Among the 546 vaccinated cases, 254 had received 1 dose; 148 had received 2 doses; 121 had received 3 doses; and one had received 4 doses of rotavirus vaccine. Data on the number of vaccine doses received was unavailable for 22 vaccinated cases.

**Comment**

Prior to the national rotavirus vaccination program which commenced in July 2007, thousands of hospitalisations due to rotavirus were reported annually. Since then, there has been a decline in hospitalisations, particularly in children aged < 5 years, with a continued small decline during this reporting period. Seasonality continued to be evident but was substantially dampened compared to the pre-vaccine era. An increase in hospitalisations in 2017 was observed due to an outbreak of rotavirus, caused by novel strains G2P[4], equine-like G3P[8] and G8P[8]. Examination of cases in New South Wales found high vaccine effectiveness over an 8-year period against vaccine-type strains, even in 2017 in the context of the outbreak. However, lower vaccine effectiveness among Aboriginal and Torres Strait Islander people, more akin to that seen in developing countries, has been reported, consistent with the higher rates of disease observed in this high-risk population in the current reporting period.
Figure 3.13.1: Rotavirus hospitalisations for all ages, Australia, 1994 to 2018, \( ^a\), \( ^b\) by month of admission

Table 3.13.1. Rotavirus hospitalisations and deaths, Australia, 2016 to 2018, \( ^c\) by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications (July–December 2018)</th>
<th>Hospitalisations</th>
<th>LOS(^d) per admission (days)</th>
<th>Deaths(^e)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Rate(^d)</td>
<td>Any diagnosis</td>
<td>Principal diagnosis</td>
</tr>
<tr>
<td>&lt; 1</td>
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<td>1–4</td>
<td>295</td>
<td>46.5</td>
<td>891</td>
<td>23.47</td>
</tr>
<tr>
<td>5–14</td>
<td>112</td>
<td>7.2</td>
<td>375</td>
<td>4.09</td>
</tr>
<tr>
<td>15–24</td>
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<td>114</td>
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<td>309</td>
<td>1.20</td>
</tr>
<tr>
<td>50–64</td>
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<tr>
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<td>186</td>
<td>9.5</td>
<td>768</td>
<td>6.75</td>
</tr>
<tr>
<td>All ages</td>
<td>1,453</td>
<td>11.6</td>
<td>3,673</td>
<td>4.98</td>
</tr>
</tbody>
</table>

\( ^{a}\) Notifications where the month of diagnosis was between 1 July 2018 and 31 December 2018; hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.

\( ^{b}\) LOS: length of stay in hospital.

\( ^{c}\) Deaths include underlying and associated causes of deaths.

\( ^{d}\) Average annual age-specific rate per 100,000 population.

\( ^{e}\) Principal diagnosis (hospitalisations).
3.14 Rubella

Highlights

Notification and hospitalisation rates for rubella remained low over the reporting period 1 January 2016 to 31 December 2018. No new cases of congenital rubella syndrome were notified during this period. The highest notification rates were among adults aged 25–49 years.

Rubella is caused by the rubella virus (family Togaviridae), which only infects humans and is transmitted by aerosol droplets. Rubella is usually a mild febrile viral disease characterised by a discrete maculopapular rash; conjunctivitis; sore throat; headache; nausea; and postauricular, suboccipital and cervical lymphadenopathy. However, subclinical infection occurs in up to 50% of cases. Arthralgia and arthritis may occur in up to 70% of infected adult females, but is uncommon in younger females and males. More severe complications, such as encephalitis, are rare. Rubella is important because up to 90% of infants born to women with a primary infection during the first trimester of pregnancy develop abnormalities characteristic of congenital rubella syndrome (CRS). The latter syndrome involves multiple serious defects, including cataract, retinopathy, deafness, heart defects and neurological deficits. Rubella infection during the first 20 weeks of pregnancy also increases the risk of spontaneous abortion, fetal death and infant mortality. Rubella vaccination has successfully decreased the burden of rubella worldwide, with 81 countries verified as having eliminated rubella as of 2018, inclusive of Australia. The risk of rubella is highest in countries that have not yet introduced rubella-containing vaccines in their national program, with women of reproductive age remaining at risk.

Case definition

Notifications

Rubella case definition:

Rubella (congenital) case definition:

Hospitalisations and deaths

The ICD-10-AM/ICD-10 code B06 (rubella [German measles]) was used to identify hospitalisations and deaths for rubella and P35.0 (congenital rubella syndrome) for hospitalisations and deaths from CRS.

Secular trends

In the three years from 1 January 2016 to 31 December 2018, there were 36 notified cases of rubella, an average annual notification rate of 0.05 per 100,000 population (Table 3.14.1). Rubella notifications have remained consistently low since 2004, following a marked decline in the late 1990s and early 2000s (Figure 3.14.1). There were no notifications of congenital rubella syndrome (CRS) between 2016 and 2018. This is consistent with surveillance reports by the Australian Paediatric Surveillance Unit which also reported no cases of CRS over this period.

The average annual hospitalisation rate for rubella remained low in this reporting period at 0.02 per 100,000 population, with 16 hospitalisations (Table 3.14.1 and Appendix C). There were 32 hospitalisations coded as due to congenital rubella syndrome (CRS) between 2016 and 2018 (0.04 per 100,000 population per year).
Severe morbidity and mortality

From 1 January 2016 to 31 December 2018, hospitalisations with an ICD-10-AM code for rubella accounted for 45 bed days with a median length of stay of 1.5 days (Table 3.14.1). The median length of stay was longer for hospitalisations recorded as due to CRS (4 days), which accounted for 191 bed days. There were 26 deaths (0.4 per 100,000 population per year) recorded over the three years from 1 January 2016 to 31 December 2018 in the causes of death data with rubella (ICD-10 code B06) as the underlying or associated cause of death, with rubella recorded as the underlying cause of death in six of these deaths.

Age and sex distribution

Adults aged 25–49 years accounted for 56% (20/36) of rubella notifications for the three-year reporting period and had the highest rate of notifications (0.08 per 100,000 population per year) (Table 3.14.1).

Rates of rubella notifications for children < 5 years of age have remained very low since 2003 and remained low over the three years from 1 January 2016 to 31 December 2018, with three cases reported among children aged 1–4 years (0.08 per 100,000 population per year). All 32 CRS hospitalisations were aged 25 years or older (13 were people aged 25–49 years; 11 were aged 50–64 years; and eight were aged ≥ 65 years).

While hospitalisation rates for rubella for the three-year period were highest for children aged 1–4 years, they were very low in all age groups at 0.05 per 100,000 population per year or lower (Table 3.14.1). Almost all deaths recorded in the causes of death data were in adults aged 50 years and older (85%), with none occurring among people aged < 25 years (Table 3.14.1).

The overall male:female rate ratio for the three-year reporting period was 0.8:1 for notifications and 0.2:1 for hospitalisations.

Geographical distribution

Notification rates for rubella remained low for all states and territories for each year from 2016 to 2018 (all < 0.1 per 100,000 population per year) (Appendix B). Similarly, hospitalisation rates for rubella in each state and territory were also very low between 2016 and 2018 (range 0–0.06 per 100,000 population per year) (Appendix C).

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for 32 of the 36 notifications (89%) of rubella during 2016–2018, with 4 not stated (11%). There were no rubella or CRS notifications or hospitalisations recorded as being in Aboriginal or Torres Strait Islander people.

Vaccination status

Of the 36 notifications, vaccination data were available for 14/36 cases (39%). Among these, six (43%) had not received any vaccine doses and eight had received 1 dose of rubella-containing vaccine.

Vaccination status should be complete in the NNDSS for all notifications of rubella in women of child-bearing age (15–45 years). There were fewer than half as many notifications (14; 0.09 per 100,000 population per year) of rubella among women aged 15–45 years between 1 January 2016 and 31 December 2018 compared to the 2012–2015 period (37; 0.18 per 100,000 population per year).254 Of these 14 cases in women aged 15–45 years, vaccination data were available for seven (50%), of whom two had received no vaccine doses and five had received 1 dose of rubella-containing vaccine.

Comment

The low and declining numbers of rubella notifications indicate that vaccination has been highly effective in Australia. A serosurvey of a nationally representative sample of the Australian population in 2012–2013 showed
Figure 3.14.1. Rubella notifications and hospitalisations,\textsuperscript{a} Australia, 1993 to 2018,\textsuperscript{b} by month of diagnosis or admission

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure3_14_1}
\caption{Rubella notifications and hospitalisations,\textsuperscript{a} Australia, 1993 to 2018,\textsuperscript{b} by month of diagnosis or admission}
\end{figure}

\textsuperscript{a} Note: varying scales between notifications and hospitalisations.

\textsuperscript{b} Notifications where the month of diagnosis was between 1 January 1993 and 31 December 2018; hospitalisations where the month of admission was between 1 July 1993 and 31 December 2018.

High levels of rubella-specific IgG seropositivity, with the effective reproductive number (R) estimated at 0.33 (95\%CI: 0.28–0.39),\textsuperscript{255} consistent with previous serosurveys reporting R less than 0.5.\textsuperscript{256} In October 2018, the WHO certified that Australia had eliminated rubella.\textsuperscript{10,11} Deaths with rubella coded as an underlying or associated cause of death should be interpreted with caution, given the generally mild illness associated with the disease. The low incidence of rubella in Australia means that the majority of IgM-positive tests are likely false positive, and may occur due to the presence of rheumatoid factors (including rheumatologic disease), cross-reacting IgM, or infection with other viruses.\textsuperscript{257,258}
Table 3.14.1: Rubella notifications, hospitalisations and deaths, Australia, 2016 to 2018, by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications</th>
<th>Hospitalisations</th>
<th>LOS(^b) per admission</th>
<th>Deaths(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Rate(^d)</td>
<td>Any diagnosis</td>
<td>Principal diagnosis</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>n</td>
<td>Rate(^d)</td>
</tr>
<tr>
<td>&lt; 1</td>
<td>0</td>
<td>0.00</td>
<td>0</td>
<td>0.00</td>
</tr>
<tr>
<td>1–4</td>
<td>3</td>
<td>0.08</td>
<td>1-4</td>
<td>0.05</td>
</tr>
<tr>
<td>5–14</td>
<td>3</td>
<td>0.03</td>
<td>1-4</td>
<td>0.01</td>
</tr>
<tr>
<td>15–24</td>
<td>7</td>
<td>0.07</td>
<td>1-4</td>
<td>0.04</td>
</tr>
<tr>
<td>25–49</td>
<td>20</td>
<td>0.08</td>
<td>7</td>
<td>0.03</td>
</tr>
<tr>
<td>50–64</td>
<td>1</td>
<td>0.01</td>
<td>1-4</td>
<td>0.01</td>
</tr>
<tr>
<td>≥ 65</td>
<td>2</td>
<td>0.02</td>
<td>1-4</td>
<td>0.01</td>
</tr>
<tr>
<td><strong>All ages</strong></td>
<td><strong>36</strong></td>
<td><strong>0.05</strong></td>
<td><strong>16</strong></td>
<td><strong>0.02</strong></td>
</tr>
</tbody>
</table>

\(^a\) Notifications where the month of diagnosis was between 1 January 2016 and 31 December 2018; hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.

\(^b\) LOS: length of stay in hospital.

\(^c\) Deaths include underlying and associated causes of deaths.

\(^d\) Average annual age-specific rate per 100,000 population.

\(^e\) Principal diagnosis (hospitalisations).

\(^f\) np: not provided.
3.15 Tetanus

Highlights

The number of notifications (14) and hospitalisations (52) for tetanus remained low between 2016 and 2018.

Tetanus is a disease caused by an exotoxin of the Clostridium tetani bacterium, which grows anaerobically at the site of an injury and is not transmissible from human to human.259 The disease is characterised by painful muscle contractions, primarily of the masseter and neck muscles, secondarily of the trunk muscles. The case-fatality rate ranges from 10% to 24% in settings with intensive care facilities, but approaches 100% in their absence, with the highest case-fatality rates in infants and the elderly.260,261

Case definition

Notifications262

Tetanus case definition:


Hospitalisations and deaths

The ICD-10-AM/ICD-10 codes A33 (tetanus neonatorum), A34 (obstetrical tetanus) and A35 (other tetanus) were used to identify hospitalisations and deaths.

Secular trends

In the three years from 1 January 2016 to 31 December 2018, there were 14 notifications of tetanus (average annual notification rate 0.02 per 100,000 population) (Table 3.15.1). The annual number of notifications for tetanus has remained stable since 2005, though a slight increase was seen during this reporting period in 2016 (Figure 3.15.1). There were 52 hospital admissions for tetanus from 1 January 2016 to 31 December 2018 (average annual hospitalisation rate 0.07 per 100,000 population) (Table 3.15.1). Hospitalisations have remained relatively stable since 1998, including during this three-year reporting period (Figure 3.15.1).

Severe morbidity and mortality

Hospitalisations with an ICD-10-AM code for tetanus accounted for 351 bed days. No hospitalisations were coded as tetanus neonatorum (A33) or obstetrical tetanus (A34). The median length of stay in hospital was three days. Adults aged 50–64 years and those aged ≥ 65 years had a longer median stay of 6.5 and 6 days, respectively. A substantially longer median stay was reported for children aged 1–4 years (25 days) (Table 3.15.1).

Causes of death data recorded 1–5 deaths with tetanus as the underlying or associated cause of death for the three years from 1 January 2016 to 31 December 2018, all in people aged ≥ 65 years (Table 3.15.1).

Age and sex distribution

During the three-year reporting period, five notifications (36%) were in adults aged ≥ 65 years (Table 3.15.1). Hospitalisations were most common among adults aged 25–49 years (21/52; 40%), but the highest hospitalisation rate was among adolescents and young adults aged 15–24 years (13/52; 0.14 per 100,000 population per year) (Table 3.15.1). There were more notifications and hospitalisations among males (male:female rate ratio 1.4:1 for notifications and 1.3:1 for hospitalisations).

Geographical distribution

The greatest number of tetanus notifications was in Queensland (n = 6) over the three years 2016–2018. The greatest number of tetanus hospitalisations was in New South Wales (n = 16) followed by Queensland (n = 14) and Victoria (n = 10) (Appendices B and C).
Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander status was reported for all 14 notifications of tetanus over the 2016–2018 period, with none recorded as being in Aboriginal and Torres Strait Islander people. There were 1–4 hospitalisations recorded as being in Aboriginal and Torres Strait Islander people during this reporting period.

Vaccination status

Of the 14 notifications, vaccination data was available for eight (57%); four of these had received no vaccine doses, two had received 1 dose, one had received 4 doses and one had received 5 doses of tetanus-containing vaccine.

Comment

Tetanus immunisation (complete) induces protective levels of antitoxin throughout childhood and into adulthood.4 However, by middle age, approximately 50% of vaccinated persons have low or undetectable levels of antitoxin.5,6 In Australia, tetanus remains largely a disease of older adults,6 but also occurred in adolescents and young adults in this reporting period. Adolescents aged 11–13 years are scheduled to receive a dTpa booster dose under the NIP, and adults aged 50 years or older are recommended to receive a booster dose of tetanus-containing vaccine if their last dose of dTpa was more than 10 years previously. Notification and hospitalisation rates in this reporting period were largely similar to those in the 2012–2015 period, with hospitalisation rates considerably higher than notification rates, as in previous reports.12,13,65 Under-notification by hospital staff, multiple hospital admissions or inter-hospital transfers for true cases, and coding errors are all possible contributing factors.66
Figure 3.15.1: Tetanus notifications and hospitalisations, Australia, 1998 to 2018, by year of diagnosis or admission

![Chart showing tetanus notifications and hospitalisations from 1998 to 2018.](chart)

Table 3.15.1: Tetanus notifications, hospitalisations and deaths, Australia, 2016 to 2018, by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Notifications</th>
<th>Hospitalisations</th>
<th>LOS(^c) per admission</th>
<th>Deaths(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>n</td>
<td>Rate(^a)</td>
<td>n</td>
<td>Rate(^a)</td>
</tr>
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<td>1–4</td>
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<td>5–14</td>
<td>2</td>
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<td>6</td>
<td>0.07</td>
</tr>
<tr>
<td>15–24</td>
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<td>0.03</td>
<td>13</td>
<td>0.14</td>
</tr>
<tr>
<td>25–49</td>
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<td>0.01</td>
<td>21</td>
<td>0.08</td>
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<td>0.03</td>
</tr>
<tr>
<td>≥ 65</td>
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<td>0.04</td>
<td>7</td>
<td>0.06</td>
</tr>
<tr>
<td>All ages</td>
<td>14</td>
<td>0.02</td>
<td>52</td>
<td>0.07</td>
</tr>
</tbody>
</table>

\(a\) Notifications where the month of diagnosis was between 1 January 2016 and 31 December 2018; hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.

\(b\) LOS: length of stay in hospital.

\(c\) Deaths include underlying and associated causes of deaths.

\(d\) Average annual age-specific rate per 100,000 population.

\(e\) Principal diagnosis (hospitalisations).
3.16 Varicella-zoster virus infection

Highlights

The number of varicella (chickenpox) hospitalisations remained low during the three-year period 1 January 2016 to 31 December 2018. The increasing trend in the numbers of herpes zoster hospitalisations, observed since 1993, has stabilised following the introduction of the national zoster vaccination program in 2016, with hospitalisation rates similar to the previous reporting period.

The varicella-zoster virus (VZV) is a DNA virus in the herpesvirus family that causes two distinct illnesses: varicella (chickenpox) following primary infection and herpes zoster (shingles) following reactivation of latent virus. Varicella is a highly contagious infection with an incubation period of 10–21 days, after which a characteristic rash appears. While typically a mild childhood disease, complications occur in about 1% of cases. Acute varicella may be complicated by secondary bacterial skin infections, haemorrhagic complications, encephalitis and pneumonia. Encephalitis occurs in 1 in 100,000 cases. Varicella is more severe in adults, particularly pregnant women and people of any age who are immunocompromised who can develop disseminated disease and are more likely to die from varicella complications. Varicella infection during pregnancy can result in congenital varicella syndrome in the infant, causing skin scarring, limb defects, ocular abnormalities and neurological malformations in the infant.

Herpes zoster (HZ) or shingles is a sporadic disease, caused by reactivation of latent VZV in sensory nerve ganglia. VZV reactivation occurs, in part, due to a decline in cellular immunity to the virus, most commonly due to ageing or with immunocompromising medical conditions or immunosuppressive treatment, and affects half of people who live to 80 years. Herpes zoster infections are characterised by severe pain with dermatomal distribution, sometimes followed by post-herpetic neuralgia, which can be chronic and debilitating, particularly in the elderly, and can last for months to years. Other neurological complications may occur, including ophthalmicus (which can lead to monocular blindness); Ramsay-Hunt syndrome; aseptic meningitis; transverse myelitis; and neurosensory hearing impairment. Approximately 13–30% of patients with herpes zoster develop complications, occurring more often in patients who are older and/or immunocompromised. Treatment with antivirals aims to shorten the duration of disease, alleviate the severity of acute pain and limit viral shedding. Atypical manifestations of HZ, such as glioma, zoster sine herpete and bilateral HZ, are rare, with limited guidelines on how to manage these patients. Given Australia’s ageing population, the burden of HZ is substantial and, in the absence of vaccination, would have been expected to increase over time.

Case definition

Notifications

Varicella-zoster (chickenpox) case definition:

Varicella-zoster (shingles) case definition:

Varicella-zoster (unspecified) case definition:

Hospitalisations and deaths

The ICD-10-AM/ICD-10 code B01 (varicella [chickenpox]) was used to identify varicella
hospitalisations and deaths and B02 (zoster [shingles]) was used to identify herpes zoster hospitalisations and deaths.

Notifications of varicella and herpes zoster

Detailed notification data are not reported due to the large proportion that are unspecified as to whether varicella or zoster. Varicella-zoster virus infection is also not notifiable in New South Wales. For the current reporting period (2016–2018), of the total 85,870 notifications in NNDSS, 12.6% (10,800) were recorded as varicella, 36.0% (30,882) as zoster and 51.5% (44,188) unspecified.

Secular trends, varicella and herpes zoster hospitalisations

In the three years from 1 January 2016 to 31 December 2018, there were 2,459 hospital admissions with an ICD-10-AM code for varicella, an average annual rate of 3.3 per 100,000 population (Table 3.16.1), similar to the 2012–2015 period (3.4 per 100,000 population per year, Figure 3.16.1).

Monthly hospitalisations with an ICD-10-AM code for herpes zoster were also stable, with an average annual rate between 2016 and 2018 of 30.8 per 100,000 population compared to 30.6 per 100,000 population per year between 2012 and 2015. This represents a stabilisation in the previously steadily-rising trend in the number of zoster hospitalisations (Figure 3.16.1).

Severe morbidity and mortality (varicella)

From 1 January 2016 to 31 December 2018, hospitalisations with an ICD-10-AM code for varicella accounted for 15,543 bed days, with a median length of stay of four days (Table 3.16.1). The median length of stay was higher in adults aged ≥ 50 years (five days for 50–64 years, seven days for ≥ 65 years) compared with younger people (two or three days, see Table 3.16.1). For the three-year period, the causes of death data recorded 41 deaths with varicella as underlying or associated cause of death (of which 14 [34%] were recorded as the underlying cause), 88% (36/41) of which occurred in adults aged ≥ 65 years (Table 3.16.1).

Age and sex distribution (varicella)

During the three-year reporting period, the highest hospitalisation rate for varicella was among infants aged < 1 year (11.1 per 100,000 population per year), followed by adults aged ≥ 65 years (8.1 per 100,000 population per year) (Table 3.16.1). In comparison to the previous four-year reporting period, the rate of varicella hospitalisations among children aged 1–4 years declined from 4.5 to 3.4 per 100,000 population per year, and among 5–14 year olds declined from 2.2 to 1.5 per 100,000 population per year. Hospitalisation rates among other age groups remained similar between the two reporting periods.

There were slightly more hospitalisations for males than females over the three years (465 vs 453, male:female rate ratio 1.1:1).

Geographical distribution (varicella)

The Northern Territory had the highest hospitalisation rate for chickenpox (4.6 per 100,000 population per year), but accounted for only 1.4% of hospitalisations for chickenpox (Appendix C). New South Wales (29%), Queensland (24%) and Victoria (24%) accounted for just over three-quarters of hospitalisations with an ICD-10-AM code for chickenpox.

Severe morbidity and mortality (herpes zoster)

Hospitalisations with an ICD-10-AM code for herpes zoster accounted for 151,495 bed days in the three years from 1 January 2016 to 31 December 2018. This is an average of 50,498 bed days per year, a decrease from the average annual number reported in the 2012–2015 period (61,212). The median length of stay was four days for hospitalisations with any herpes
zoster diagnosis and three days for those with a principal diagnosis of herpes zoster (Table 3.16.2).

For the three-year reporting period, the causes of death data recorded 322 deaths with zoster (herpes zoster, shingles) as the underlying or associated cause of death (of which 91 [28%] were recorded as the underlying cause); nearly all (309/322, 96%) were aged ≥ 65 years (Table 3.16.2).

Age and sex distribution (herpes zoster)

During the three-year reporting period the highest hospitalisation rate for herpes zoster was among adults aged ≥ 65 years (145.6 per 100,000 population per year), followed by 50–64 year olds (27.3 per 100,000 population per year) (Table 3.16.2). The rate of hospitalisation among adults ≥ 65 years was slightly lower in this reporting period than in the 2012–2015 period (145.6 vs 155.3 per 100,000 population per year). The median length of stay for zoster hospitalisations was greatest among those aged ≥ 65 years (five days). There were more zoster hospitalisations for females than males (9,615 vs 13,120, male:female rate ratio 1:1.3).

Geographical distribution (herpes zoster)

The highest zoster hospitalisation rate was in Queensland (34.5 per 100,000 population per year). New South Wales (32%), Victoria (25%) and Queensland (22%) accounted for over three-quarters of hospitalisations (Appendix C).

Aboriginal and Torres Strait Islander status

Aboriginal and Torres Strait Islander people accounted for 3.8% of varicella hospitalisations (94/2,459), with the highest rate observed in infants aged < 1 year (21.1 per 100,000 population per year). Aboriginal and Torres Strait Islander varicella hospitalisation rates were slightly higher than those of other Australians (3.9 vs 3.3 per 100,000 population per year; rate ratio 1.2:1). Aboriginal and Torres Strait Islander people accounted for 2% of zoster hospitalisations (425/22,735), with the highest rate in those aged ≥ 65 years (106.1 per 100,000 population per year). The overall Aboriginal and Torres Strait Islander zoster hospitalisation rate was lower than in other Australians (17.4 per 100,000 population per year and 31.3 per 100,000 population per year, respectively). While rates of hospitalisation among adults aged ≥ 65 years were lower among Aboriginal and Torres Strait Islander people than among non-Aboriginal and Torres Strait Islander adults (106.1 vs 146.0 per 100,000 population per year, respectively), rates were higher for Aboriginal and Torres Strait Islander people in younger age groups (adults aged 50–64 years: 52.8 vs 26.8 per 100,000 population per year; adults aged 25–49 years: 14.6 vs 7.9 per 100,000 population per year; 15–24 years: 6.8 vs 3.5 per 100,000 population per year).

Comment

A nationally-funded zoster vaccination program commenced in Australia in November 2016, with people aged 70 years eligible for a single dose of live attenuated zoster vaccine, and a five-year catch-up program for people aged 71 to 79 years. Coverage in the first 17 months of the program (1 November 2016 to 31 March 2018) was less than optimal (33.9% for adults aged 70 years and 25.8% for the catch-up program for adults aged 71–79 years), although there has likely been underreporting of administered vaccines to the Australian Immunisation Register; another study using an electronic primary care dataset has reported higher coverage estimates (46.9% up to 31 December 2018). However, despite this less than optimal coverage, hospitalisation rates for herpes zoster stabilised in this reporting period, breaking the steadily increasing trend observed since 1993, and the rate of zoster hospitalisations was slightly lower among adults aged ≥ 65 years than during the 2012–2015 period. An evaluation of antiviral prescriptions for herpes zoster found that, despite increases in the rate of antiviral prescriptions in the decade prior to the vaccination program, prescriptions decreased markedly in the following two years, providing...
Figure 3.16.1: Varicella and herpes zoster hospitalisations, Australia, 1993 to 2018,\(^a\)\(^b\) by month of admission

Further evidence of impact on disease incidence.\(^{287}\) Rates of zoster hospitalisations among Aboriginal and Torres Strait Islander adults aged < 70 years have been previously reported to be higher than those among non-Indigenous Australians,\(^{288}\) a trend that continued in this reporting period.

Although the vaccine effectiveness of a single dose against varicella hospitalisation has been moderate, reported at 64.7%,\(^{289}\) varicella vaccination has substantially reduced the burden of varicella disease in Australia.\(^{290}\) Varicella hospitalisation rates remained stable in this reporting period compared with the previous 2012–2015 period, indicating the sustained impact of varicella vaccination since its introduction onto the NIP in 2005.\(^{290,291}\) Varicella hospitalisation rates continued to decline among the age groups eligible to receive varicella vaccination under the NIP (1–14 years). The vaccination program has also been successful in reducing the disparity in varicella hospitalisation rates between Aboriginal and Torres Strait Islander children and non-Aboriginal and Torres Strait Islander children.\(^{291}\)

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\(^a\) Hospitalisations where the month of admission was between 1 July 1993 and 31 December 2018.

\(^b\) Inset graph shows hospitalisations, where the month of admission was between 1 July 2001 and 31 December 2018, for vaccine-eligible age groups.
### Table 3.16.1: Varicella hospitalisations and deaths, Australia, 2016 to 2018,\(^a\) by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Hospitalisations</th>
<th>LOS(^b) per admission</th>
<th>Deaths(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Any diagnosis</td>
<td>Principal diagnosis</td>
<td>Any diagnosis</td>
</tr>
<tr>
<td></td>
<td>n</td>
<td>Rate(^d)</td>
<td>n</td>
</tr>
<tr>
<td>&lt; 1</td>
<td>103</td>
<td>11.09</td>
<td>79</td>
</tr>
<tr>
<td>1–4</td>
<td>128</td>
<td>3.37</td>
<td>85</td>
</tr>
<tr>
<td>5–14</td>
<td>140</td>
<td>1.53</td>
<td>93</td>
</tr>
<tr>
<td>15–24</td>
<td>144</td>
<td>1.50</td>
<td>105</td>
</tr>
<tr>
<td>25–49</td>
<td>649</td>
<td>2.52</td>
<td>433</td>
</tr>
<tr>
<td>50–64</td>
<td>377</td>
<td>2.87</td>
<td>173</td>
</tr>
<tr>
<td>≥ 65</td>
<td>918</td>
<td>8.07</td>
<td>375</td>
</tr>
<tr>
<td><strong>All ages</strong></td>
<td>2,459</td>
<td>3.33</td>
<td>1,343</td>
</tr>
</tbody>
</table>

\(^a\) Hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.
\(^b\) LOS: length of stay in hospital.
\(^c\) Deaths include underlying and associated causes of deaths.
\(^d\) Average annual age-specific rate per 100,000 population.
\(^e\) Principal diagnosis (hospitalisations and deaths).

### Table 3.16.2: Zoster hospitalisations and deaths, Australia, 2016 to 2018,\(^a\) by age group

<table>
<thead>
<tr>
<th>Age group (years)</th>
<th>Hospitalisations</th>
<th>LOS(^b) per admission</th>
<th>Deaths(^c)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Any diagnosis</td>
<td>Principal diagnosis</td>
<td>Any diagnosis</td>
</tr>
<tr>
<td></td>
<td>n</td>
<td>Rate(^d)</td>
<td>n</td>
</tr>
<tr>
<td>&lt; 1</td>
<td>16</td>
<td>1.72</td>
<td>1-4</td>
</tr>
<tr>
<td>1–4</td>
<td>33</td>
<td>0.87</td>
<td>np(^f)</td>
</tr>
<tr>
<td>5–14</td>
<td>92</td>
<td>1.00</td>
<td>61</td>
</tr>
<tr>
<td>15–24</td>
<td>354</td>
<td>3.69</td>
<td>179</td>
</tr>
<tr>
<td>25–49</td>
<td>2,091</td>
<td>8.11</td>
<td>1,031</td>
</tr>
<tr>
<td>50–64</td>
<td>3,588</td>
<td>27.35</td>
<td>1,767</td>
</tr>
<tr>
<td>≥ 65</td>
<td>16,561</td>
<td>145.58</td>
<td>6,637</td>
</tr>
<tr>
<td><strong>All ages</strong></td>
<td>22,735</td>
<td>30.82</td>
<td>9,694</td>
</tr>
</tbody>
</table>

\(^a\) Hospitalisations where the month of admission was between 1 January 2016 and 31 December 2018.
\(^b\) LOS: length of stay in hospital.
\(^c\) Deaths include underlying and associated causes of deaths.
\(^d\) Average annual age-specific rate per 100,000 population.
\(^e\) Principal diagnosis (hospitalisations and deaths).
\(^f\) np: not provided
Acknowledgements

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References


Appendix A. Charts of historical national notification data

Selected vaccine preventable diseases

Figure A.1: Notifications of diphtheria, 1917 to 2018, Australia

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Figure A.2: Notifications of *Haemophilus influenzae* type b, 1991 to 2018, a Australia

- 1992 - First Hib vaccines approved
- 1993 - National Hib vaccination program commenced

Notifications per 100,000 population

<table>
<thead>
<tr>
<th></th>
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<tbody>
<tr>
<td>1991</td>
<td>3.5</td>
<td>3.0</td>
<td>2.5</td>
<td>2.0</td>
<td>1.5</td>
<td>1.0</td>
<td>0.5</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
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<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
<td>0.0</td>
</tr>
</tbody>
</table>

Figure A.3: Notifications of hepatitis A, 1952 to 2018, Australia

- 1994 - HAV vaccine approved
- 2005 - HAV vaccination program commenced for Aboriginal and Torres Strait Islander children in NT, Qld, WA and SA


Figure A.4: Notifications of measles, 1917 to 2018. Australia

- 1970 - Measles vaccine became widely available
- 1993 - Second dose of MMR vaccine introduced for 10-16
- 1998 - Second dose of MMR vaccine lowered to 4-5 years; Measles Control Campaign
- 2000 - Second dose of MMR vaccine lowered to 4 years
- 2001 - Second dose of MMR vaccine lowered to 4-5 years
- 2013 - Second dose of MMR vaccine lowered to 18 months and delivered as a combination MMRV vaccine


Figure A.5: Notifications of meningococcal disease (invasive), 1949 to 2018, Australia

- 2003 - National Meningococcal C vaccination program commenced. Meningococcal C conjugate vaccine added to vaccination schedule for 12 month olds.
- 2018 - MenC vaccine replaced with quadrivalent meningococcal conjugate (MenACWY) vaccine.


Figure A.6: Notifications of mumps, 1932 to 2018, Australia

- 1982 - Single dose of mumps vaccine (given as measles-mumps vaccine) funded on the national schedule for 12 month olds
- 1993 - Second dose of MMR vaccine introduced for 10-14 year olds
- 1998 - Second dose of MMR vaccine lowered to 4-5 years; Measles Control Campaign targets children 5-12 years
- 2000 - Second dose of MMR vaccine lowered to 4 years
- 2013 - Second dose of MMR vaccine lowered to 18 months and delivered as a combination MMRV vaccine
- 2015


Figure A.7: Notifications of pertussis, 1917 to 2018, Australia

- 1942 - Mass vaccination with pertussis vaccine commenced
- 1953 - DTP vaccination introduced
- 1994 - Fifth dose of DTPa at 4-5 years added to the vaccination schedule (replacing CDT vaccine)
- 2003 - Fourth dose of DTPa at 18 months no longer recommended
- 2004 - dTpa funded for 15-17 years, replacing the dT dose
- 2004 - dTpa funded for 15-17 years, replacing the dT dose
- 2015 - Booster dose of DTPa added to the schedule at 18 months
- 2018 - Antenatal dose of dTpa added to the schedule

Figure A.8: Notifications of poliomyelitis, 1917 to 2018, Australia

- 1956 - Mass vaccination with IPV commenced
- 1966 - OPV introduced
- 1994 - Reinforcing OPV to 15 year olds
- 2005 - IPV funded to replace OPV, in combination vaccines
- 1998 - OPV booster dose to 4 year olds before starting school


Figure A.9: Notifications of rubella, 1942 to 2018, Australia

- 1971 - Schoolgirl rubella program commenced
- 1989 - MMR replaced MM vaccine for infants
- 1993 - Two-dose schedule introduced
- 2000 - MMR rather than rubella vaccine recommended for non-immune women of childbearing age
- 2013 - Second dose of MMR vaccine delivered as a combination MMRV vaccine


Figure A.10: Notifications of tetanus, 1921 to 2018, a Australia

- 1953 - DTP vaccination introduced

Notifications per 100,000 population

Year

- 1921
- 1924
- 1927
- 1930
- 1933
- 1936
- 1939
- 1942
- 1945
- 1948
- 1951
- 1954
- 1957
- 1960
- 1963
- 1965-66
- 1968-69
- 1971
- 1974
- 1977
- 1980
- 1983
- 1986
- 1989
- 1992
- 1995
- 1998
- 2001
- 2004
- 2007
- 2010
- 2013
- 2016


### Table A.1: Notifications and notification rates for vaccine preventable diseases, Australia, 1 January 2016 to 31 December 2018, by state or territory and year.

<table>
<thead>
<tr>
<th>Disease</th>
<th>Year</th>
<th>Number of notifications</th>
<th>Notification rate per 100,000 population</th>
</tr>
</thead>
<tbody>
<tr>
<td>Diphtheria</td>
<td>2016</td>
<td>0</td>
<td>0.00</td>
</tr>
<tr>
<td></td>
<td>2017</td>
<td>0</td>
<td>0.00</td>
</tr>
<tr>
<td></td>
<td>2018</td>
<td>0</td>
<td>0.00</td>
</tr>
<tr>
<td>Haemophilus influenzae type b</td>
<td>2016</td>
<td>0</td>
<td>0.00</td>
</tr>
<tr>
<td></td>
<td>2017</td>
<td>0</td>
<td>0.00</td>
</tr>
<tr>
<td></td>
<td>2018</td>
<td>0</td>
<td>0.00</td>
</tr>
<tr>
<td>Hepatitis A</td>
<td>2016</td>
<td>2</td>
<td>0.50</td>
</tr>
<tr>
<td></td>
<td>2017</td>
<td>2</td>
<td>0.49</td>
</tr>
<tr>
<td></td>
<td>2018</td>
<td>2</td>
<td>0.48</td>
</tr>
<tr>
<td>Hepatitis B (acute)</td>
<td>2016</td>
<td>3</td>
<td>0.74</td>
</tr>
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<td>2017</td>
<td>1</td>
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<tr>
<td></td>
<td>2018</td>
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<tr>
<td>Influenza</td>
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<td></td>
<td>2017</td>
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<tr>
<td>Measles</td>
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<td>18</td>
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<td></td>
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<td>Total</td>
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<td>Meningococcal disease (invasive)</td>
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<td></td>
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<td>Mumps</td>
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<td></td>
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<td>4</td>
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<td></td>
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<td>Total</td>
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<td>Pertussis</td>
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<td></td>
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<tr>
<td>Total</td>
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<td>22453</td>
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<td>Disease</td>
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<td>Notification rate per 100,000 population</td>
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<td>----------------------------------------</td>
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<tr>
<td></td>
<td></td>
<td>ACT</td>
<td>NSW</td>
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<tr>
<td>Q fever</td>
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<tr>
<td></td>
<td>2017</td>
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<td>210</td>
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<tr>
<td></td>
<td>2018</td>
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<td>223</td>
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<tr>
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- **b** Total cases for 3-year period and average annual rate per 100,000 population.
- **c** Total cases for 6-month period July to December 2018 only (i.e. since rotavirus became a nationally notifiable disease); rates have been annualised.
### Appendix C. Hospitalisations by state or territory

Table A.2: Hospitalisation rates for vaccine preventable diseases, Australia, 1 January 2016 to 31 December 2018, by state or territory and year

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<th>Disease</th>
<th>Year</th>
<th>Hospitalisation rate per 100,000 population</th>
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<tr>
<td></td>
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<tr>
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<tr>
<td>Haemophilus influenzae meningitis</td>
<td>2016</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>2017</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>2018</td>
<td>–</td>
</tr>
<tr>
<td></td>
<td>Total</td>
<td>–</td>
</tr>
<tr>
<td>Hepatitis A</td>
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<td></td>
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<td>Hepatitis B (acute) (principal diagnosis only)</td>
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<tr>
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<tr>
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<tr>
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<tr>
<td></td>
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</tr>
<tr>
<td></td>
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</tr>
<tr>
<td>Tetanus*</td>
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<tr>
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<td>---------</td>
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a: Hospitalisations for rare diseases such as diphtheria, rubella and tetanus should be interpreted with caution due to possible misclassification, coding and data related issues. Diphtheria hospitalisations coded here include non-respiratory and non-toxigenic diphtheria infections.

b: Average annual rate per 100,000 population. The rates are based on hospitalisations (in public and private hospitals) by date of admission and state of residence.

c: Retained for consistency with previous reports in this series. Data for Haemophilus influenzae (Hib) hospitalisations are not presented because the available ICD-10 codes are not type specific.

d: Pneumococcal meningitis, sepsicaemia or pneumonia.
### Appendix D. Abbreviations

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<tr>
<th>Abbreviations</th>
<th>Description</th>
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<tr>
<td>4vHPV</td>
<td>Quadrivalent human papillomavirus vaccine</td>
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<tr>
<td>9vHPV</td>
<td>Nonavalent human papillomavirus vaccine</td>
</tr>
<tr>
<td>ABS</td>
<td>Australian Bureau of Statistics</td>
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<tr>
<td>ACR</td>
<td>Australian Coordinating Registry</td>
</tr>
<tr>
<td>AFP</td>
<td>Acute flaccid paralysis</td>
</tr>
<tr>
<td>AIHW</td>
<td>Australian Institute of Health and Welfare</td>
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<td>AIR</td>
<td>Australian Immunisation Register</td>
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<tr>
<td>COD URF</td>
<td>Cause of Death Unit Record File</td>
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<tr>
<td>CRS</td>
<td>Congenital rubella syndrome</td>
</tr>
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<td>DTPa</td>
<td>diphtheria-tetanus-pertussis vaccine</td>
</tr>
<tr>
<td>dTpa</td>
<td>diphtheria-tetanus-pertussis vaccine (reduced antigen formulation)</td>
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<td><em>Haemophilus influenzae</em> type b</td>
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<td>Hib-MenC</td>
<td><em>Haemophilus influenzae</em> type b- meningococcal serogroup C vaccine</td>
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<tr>
<td>HZ</td>
<td>Herpes zoster</td>
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<td>ICD</td>
<td>International Classification of Diseases</td>
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<tr>
<td>ICD-9-CM</td>
<td>International Classification of Diseases, Ninth Revision, Clinical Modification</td>
</tr>
<tr>
<td>ICD-10-AM</td>
<td>International Statistical Classification of Diseases and Related Health Problems, 10th Revision, Australian Modification</td>
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<tr>
<td>IMD</td>
<td>Invasive meningococcal disease</td>
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<tr>
<td>IPD</td>
<td>Invasive pneumococcal disease</td>
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<tr>
<td>LOS</td>
<td>Length of stay (hospitalisations)</td>
</tr>
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<td>MenACWY</td>
<td>Quadrivalent meningococcal conjugate vaccine</td>
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<td>NCIRS</td>
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<td>NCIS</td>
<td>National Coronial Information System</td>
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<td>National Immunisation Program</td>
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<td>NNDSS</td>
<td>National Notifiable Diseases Surveillance System</td>
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<td>PCR</td>
<td>Polymerase chain reaction</td>
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<td>13-valent pneumococcal conjugate vaccine</td>
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<td>Polio Expert Panel</td>
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<td>PPV23</td>
<td>23-valent pneumococcal polysaccharide vaccine</td>
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<td>Vaccine-derived poliovirus</td>
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<td>Vaccine preventable disease</td>
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<td>Varicella-zoster virus</td>
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<td>World Health Organization</td>
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