Use of data linkage to improve communicable disease surveillance and control in Australia: existing practices, barriers and enablers

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In Australia, communicable diseases are monitored primarily through the surveillance of scheduled notifiable conditions operating under the governance of state and territory health departments. The system is underpinned by robust public health legislation at the jurisdictional level requiring medical practitioners and/or pathology services to notify a range of communicable diseases of public health importance. A subset of jurisdictional data, excluding identifiers, are transmitted to the National Notifiable Disease Surveillance System (NNDSS) under the auspices of the National Health Security Act 2007,1 operated by the Australian Department of Health.

The notification system has two purposes: to enable a swift public health response to prevent further cases of disease; and to monitor disease epidemiology over time to inform public health policy. Gaps in surveillance data may have an impact on communicable disease prevention and control efforts and may limit optimal development of evidence-based policy. Three important examples are Aboriginal and Torres Strait Islander status (missing in 50% of NNDSS cases), mortality (missing in 56% of cases) and vaccination status (missing in 17% of relevant cases).3 These gaps impact on estimates of disease burden in important populations4–6 and hinder evaluation of vaccination programs.7–9 Traditionally, missing communicable disease surveillance data are addressed through active case follow-up, whereby cases and/or notifying practitioners are interviewed by trained public health practitioners to obtain the missing data. However, this approach is resource intensive and often impracticable, particularly in view of the large volume of cases being notified to jurisdictional health departments.6,7 Data linkage therefore represents another method for closing surveillance gaps.

Data linkage (also known as record linkage, data matching or integration) involves bringing together information about an individual or an event from disparate sources (Box 1).8,9 Use of data linkage in Australia has been steadily increasing as a relatively inexpensive method of conducting epidemiological research or monitoring health at a population level.10 It is often used to improve understanding of chronic diseases including cancer, cardiovascular and respiratory diseases, as well as for injury, and mother and child health research, where long-term monitoring, treatment or care is required.10–12 Most data linkage studies in

Abstract

Objectives: To review the use of data linkage by Australian state and territory communicable disease control units, and to identify barriers to and enablers of data linkage to inform communicable disease surveillance and control activities.

Methods: Semi-structured telephone interviews were carried out with one key informant from communicable disease control units in all eight Australian states and territories between October 2017 and January 2018.

Results: Key informants from all Australian states and territories participated in the interview. A variety of existing practices were identified, with few jurisdictions making systematic use of available data linkage infrastructure. Key barriers identified from the review included: a lack of perceived need; system factors; and resources. Existing regulatory tools enable data linkage to enhance communicable disease surveillance and control.

Conclusions: We identified considerable variation in the use of data linkage to inform communicable disease surveillance and control activities between jurisdictions. We suggest that routinely collected, disparate data are systematically integrated into existing surveillance and response policy cycle to improve communicable disease prevention and control efforts.

Implications for public health: Existing gaps in communicable disease surveillance data may affect prevention and control efforts. Data linkage is recognised as a valuable method to close surveillance gaps and should be used to enhance the value of publicly held health data.

Key words: data linkage, communicable disease, surveillance, infectious diseases

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Australia are conducted in New South Wales (NSW) and Western Australia (WA), where linkage infrastructure is well developed and utilised. In the communicable diseases setting, data linkage can be used to evaluate public health interventions and guide policy development. In Australia, data linkage methodologies have been used to quantify case ascertainment in communicable disease registers, to improve estimates of selected communicable diseases; to better describe the burden of selected communicable diseases among Aboriginal and Torres Strait Islander peoples; utilised. In the communicable diseases linkage infrastructure is well developed and managed or overseeing surveillance systems in communicable disease control units in each Australian state and territory were selected as key informants in this study. Semi-structured telephone interviews were carried out with these key informants between October 2017 and January 2018. Interview questions were grouped into the following themes: availability and use of existing data linkage infrastructure; availability and use of disparate datasets (either linked or unlinked); existing governance arrangements to support data linkage; and existing or planned priorities relating to the use of data linkage. Key informants were also asked to identify barriers to and enablers of the use of data linkage in the communicable disease setting. Notes taken during interviews were reviewed and clarification sought from key informants if required. Responses from each jurisdiction relating to their availability and use of existing data linkage infrastructure, as well as their availability and use of disparate datasets were tabulated. Commonly cited barriers to and enablers of data linkage were synthesised into overarching themes where possible. In this study, we use the term ‘jurisdiction’ interchangeably with the term ‘state and territory’.

Methods
Individuals who were responsible for managing or overseeing surveillance systems in communicable disease control units in each Australian state and territory were selected as key informants in this study. Semi-structured telephone interviews were carried out with these key informants between October 2017 and January 2018. Interview questions were grouped into the following themes: availability and use of existing data linkage infrastructure; availability and use of disparate datasets (either linked or unlinked); existing governance arrangements to support data linkage; and existing or planned priorities relating to the use of data linkage. Key informants were also asked to identify barriers to and enablers of the use of data linkage in the communicable disease setting. Notes taken during interviews were reviewed and clarification sought from key informants if required. Responses from each jurisdiction relating to their availability and use of existing data linkage infrastructure, as well as their availability and use of disparate datasets were tabulated. Commonly cited barriers to and enablers of data linkage were synthesised into overarching themes where possible. In this study, we use the term ‘jurisdiction’ interchangeably with the term ‘state and territory’.

Results
Representatives from all eight Australian states and territories participated in the interview (see Acknowledgements, noting that the key informant for Victoria was author, SR). We identified a variety of existing practices in each jurisdiction’s use of data linkage.

Integration of notifiable communicable disease data into existing linkage infrastructure
Communicable disease notification data were incorporated into a consolidated master linkage system (MLS) operated by Data Linkage Units (Box 1) in five of the eight Australian states and territories: the Australian Capital Territory (ACT), New South Wales (NSW), Queensland (QLD), Victoria (VIC) and Western Australia (WA), see Table 1(A). Additionally, in NSW the MLS – maintained by the Centre for Health Record Linkage (CHeReL) – is used to create a separate Communicable Disease Register (CDR). The CDR is distinct from their local notifiable disease register, Notifiable Conditions Information Management System (NCIMS), and contains de-identified linked data integrating NCIMS and Human Immunodeficiency Virus (HIV) database with three selected disparate datasets containing emergency department presentations, hospital admissions and death registrations. It also contains records from these datasets based on predefined International Classification of Disease (ICD-10-AM) and Systematized Nomenclature of Medicine (SNOMED) codes relating to notifiable communicable diseases even if a corresponding person record is not captured on NCIMS. CDR was created under the Public Health and Diseases Registers provision of the Public Health Act to support communicable disease policy, planning and operations.

Availability of data from disparate datasets
Most commonly, communicable disease data were linked to administrative datasets (those containing hospital-based care such as admitted episodes and emergency department presentations) and vital statistics (deaths). None of the jurisdictions reported having their communicable disease notification data routinely linked to national datasets such as the Medicare Benefits Schedule (MBS), the Pharmaceutical Benefits Scheme (PBS) or the Australian Immunisation Register (AIR), although some jurisdictions (NSW, QLD, VIC, WA) reported current or planned ad hoc linkage and analysis of the MBS and PBS. Several jurisdictions reported having regular access to disparate datasets or databases that could be interrogated on a case-by-case basis (herein referred to as unlinked data) to support communicable disease surveillance and control activities. All jurisdictions reported having access to AIR, and six reported having selected access to local hospital administrative systems (ACT, Northern Territory [NT], QLD, South Australia [SA], Tasmania [TAS]) and WA, Table 1(B)).
Uses of data from disparate datasets

Use of the datasets varied considerably by jurisdiction (Table 1[C]) and none reported using linked data for the purpose of updating missing data in local communicable disease registers (sometimes referred to as ‘data cleansing’ [Box 1]). Among the five jurisdictions that reported having their data incorporated into an MLS, few cited routine applications of the linked data. WA first incorporated communicable disease notification data into an MLS in 2007. However, the data obtained through the linkage process are seldom used operationally nor are the returned, linked data incorporated into their local communicable disease notification register. The ACT and NSW incorporated communicable disease notification data into an MLS in 2014. In the ACT, however, the communicable disease data are not routinely refreshed (updated), so linked content data from disparate datasets are not used. In NSW, linked data from the CDR is routinely used to improve sexually transmissible disease estimates among Aboriginal and Torres Strait Islander people.60 Other planned uses included identification of co-infections and other comorbidities associated with communicable diseases.66 Because CDR contains only de-identified data and is distinct from the NCIMS, data obtained via the linkage process are not incorporated into NCIMS nor transmitted.

Table 1: (A) Data linkage infrastructure and availability of linked communicable diseases data; (B) Linked and unlinked data types available and used in the setting of communicable disease surveillance and control; (C) Uses of linked or unlinked data by communicable disease control units.

(A) Infrastructure

<table>
<thead>
<tr>
<th>State / Territory</th>
<th>ACT</th>
<th>NSW</th>
<th>NT</th>
<th>QLD</th>
<th>SA</th>
<th>TAS</th>
<th>VIC</th>
<th>WA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Data linkage unit</td>
<td>ChReL</td>
<td>ChReL</td>
<td>SA-NT DataLink</td>
<td>DLQ</td>
<td>SA-NT DataLink</td>
<td>TDLU</td>
<td>CVDL</td>
<td>WADLB</td>
</tr>
<tr>
<td>Year communicable disease data incorporated into MLS</td>
<td>2014</td>
<td>2014</td>
<td>n/a</td>
<td>2017</td>
<td>n/a</td>
<td>n/a</td>
<td>2017</td>
<td>2007</td>
</tr>
<tr>
<td>Years for which communicable disease data are available</td>
<td>2000 -</td>
<td>1993 – Dec 2016</td>
<td>n/a</td>
<td>1989 -</td>
<td>n/a</td>
<td>n/a</td>
<td>1990 -</td>
<td>1990 -</td>
</tr>
<tr>
<td>Frequency of data refresh</td>
<td>ad hoc</td>
<td>annual</td>
<td>n/a</td>
<td>n/a</td>
<td>n/a</td>
<td>quarterly</td>
<td>annual</td>
<td></td>
</tr>
</tbody>
</table>

(B) Data types

Linked • Unlinked ○

- ED presentations ○ ○ ○ ○ ○ ○ ○ ○ □ □
- Hospital admissions ○ ○ ○ ○ ○ ○ ○ ○ ○
- Death registrations ○ ○ ○ ○ ○ ○ ○ ○
- Laboratory data* ○ ○ ○ ○ ○ ○ ○ ○
- Immunisation records ○ ○ ○ ○ ○ ○ ○ ○
- PBS / MBS records ○ ○ ○ ○ ○ ○ ○ ○

(C) Use

| | | | | | | | | |
| --- | --- | --- | --- | --- | --- | --- | --- |
| | | | | | | | | |
| Aggregate analyses | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| Data cleansing† | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| Contact details | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| Aboriginal and Torres Strait Islander | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| Alive / Died | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| ED or hospital attendance | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| Risk factors, comorbidities | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| Manifestation, outcome | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| Vaccination status | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| Laboratory results | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| Public health investigations | ○ | ○ | ○ | ○ | ○ | ○ | ○ |
| Case ascertainment | ○ | ○ | ○ | ○ | ○ | ○ | ○ |

Notes:

† Selected death registrations only. ACT receive respiratory deaths using a methodology previously described.66

* Denotes access to one or more databases maintained by a public laboratory.

† “Data cleansing” involves the large-scale comparison or matching of two or more sets of personal data (either held by the same organisation or by different organisations), for the purposes of updating one or both of the sets.67 In this context, data cleansing refers to the updating of missing surveillance data from local notifiable communicable diseases registers with content data gleaned through linkage with or interrogation of disparate datasets. ChReL – Centre for Health Record Linkage; SA-NT DataLink – South Australia and Northern Territory DataLink; DLQ – Data Linkage Queensland; TDLU – Tasmanian Data Linkage Unit; CVDL – Center for Victorian Data Linkage; DLB – Data Linkage Branch; MLS – Master Linkage System; n/a – not applicable; tbd – to be determined; ED – Emergency Department; PBS - Pharmaceutical Benefits Scheme; MBS - Medicare Benefits Scheme.
Most jurisdictions indicated that data linkage methodologies were used for *ad hoc* public health investigations or descriptive epidemiological analyses. These uses involved engagement of a Data Linkage Unit, development of user-built data linkage programs/look-up algorithms, or use of publicly available data linkage software. More commonly, linked data are used to address specific *ad hoc* research questions that are initiated and carried out by research institutions in collaboration with, or distinct from, communicable disease units. In contrast, jurisdictions indicated that data obtained from unlinked databases are regularly used for data cleansing (Box 1) to supplement or improve existing communicable disease surveillance data. The most commonly reported use of data among jurisdictions reporting access to unlinked hospital administrative databases (ACT, NT, QLD, SA, TAS, WA) was to obtain missing demographic and clinical data relating to a notifiable disease, such as Aboriginal and Torres Strait Islander status. Data gleaned in this manner is then manually incorporated into local notifiable disease registers. Other uses included the ability to obtain updated case contact details, and to identify clinical (e.g. manifestation and outcome) or risk factor and comorbidity data. Hospital datasets were sometimes used to ascertain case deaths. Some jurisdictions also reported having access to unlinked mortality data (ACT, SA, NT), whereby selected death registrations (those potentially due to a notifiable communicable disease) are manually cross-checked against the notifiable diseases register, which is then updated accordingly. In SA, unlinked mortality data are also used to identify previously un-notified cases of notifiable communicable disease (i.e. to improve case ascertainment).

**Existing governance arrangements**

Four jurisdictions (ACT, NSW, VIC, and WA) had agreements or protocols in place between communicable disease units and Data Linkage Units, clearly documenting the procedural and governance arrangements surrounding integration of communicable disease data into an MLS. Two states and territories – the ACT and NSW that are incorporated into the MLS maintained by the CHerRel – had data dictionaries describing the availability of linked notifiable disease data and corresponding variables on a publicly available website. Only one state (WA) had detailed specifications relating to identification of relevant linked records with a temporal component. Hospitalisation records are identified and linked to cases of communicable diseases only if a hospitalisation (for any diagnoses) occurred within or on 10 days before or after specimen collection date or death date. Similarly, linked death registrations were only identified and returned if the death was registered up to four days prior or 56 days following the date of onset (or specimen collection date if death date not known). Conversely, in NSW’s CDR, person-specific linkages between cases of communicable diseases and disparate datasets occur for all cases regardless of notification or onset dates.

**Barriers to and enablers of data linkage**

A range of barriers and enablers were identified. These were broadly grouped into the following themes: Perceived need; Regulatory tools; System factors; and Resources.

**Perceived need**

Perceived need was considered both a prohibiting and enabling factor to the use of data linkage in the communicable disease setting. Many of the jurisdictions (ACT, NT, SA, TAS) indicated that pertinent information relevant for communicable disease surveillance and control activities could easily be gleaned on a case-by-case basis via access to unlinked databases, thus precluding the need for data linkage. All of these informants acknowledged that data linkage would in theory be useful for high-volume conditions,
but also indicated that establishing a system to routinely integrate disparate data into local notifiable disease registers was not a current priority.

In contrast, the remaining jurisdictions (WA, NSW, QLD, VIC) – which serve larger population sizes – reported a strong desire for using data linkage to support communicable disease surveillance and control activities in Australia. Most of these jurisdictions also expressed a concomitant desire for the development of a nationally standardised approach for establishing linkages with notifiable communicable diseases, as well as for standardised analytical approaches.

**Regulatory tools**

Few of the informants cited regulatory tools as a prohibiting or enabling factor for the use of data linkage in the communicable disease setting. NSW made reference to the NSW Public Health Act 2010, which allows for the establishment of public health registers for a range of purposes including “to facilitate the care, treatment and the follow up of persons who has or may have an infectious disease, or to facilitate the identification of risk factors or outcomes associated with a disease.” VIC made reference to national and state-based legislation governing data collection and use specifically for public health purposes, noting that the powers of the Chief Health Officer under the Victorian Public Health and Wellbeing Act 2008 were recently used to establish the integration of notifiable disease data (captured on Victoria’s Public Health Events Surveillance System – PHESS) into a local MSL. VIC also noted national and state-based privacy legislation governing the handling of personal information, as well as data sharing legislation supporting sharing and use of public sector data to guide policy making, service planning and design.

Some of the informants (VIC, QLD, TAS) discussed potential privacy concerns as a potential barrier to the use of data linkage if the linkage was to be used for ‘data cleansing’ purposes. Integrating content data – that is, data sourced from disparate datasets through the linkage process – with notifiable disease registries would essentially render the individuals re-identified. However, all of these informants also expressed a view that the public health benefits of data cleansing would outweigh the risks posed by the re-identification process, citing national and jurisdictional privacy legislation and associated privacy principles as enabling factors.

They also noted that jurisdictions already have access to a range of publicly held record-level data to support communicable disease surveillance and control activities, so the use of data linkage methodologies in this context simply improves the efficiency of this established practice.

**System factors**

A range of system factors were identified as potential barriers to the use of data linkage. First, NSW, QLD and VIC specifically made note of Australia’s federated political system as a barrier to the use of data linkage to improve completeness of key data items in the NNDSS. Collection of personally identifiable communicable disease data rests with states and territories, and only de-identified communicable disease data are transmitted to the NNDSS. This precludes national data linkage activities by the Australian Department of Health on behalf of the states and territories for public health purposes. Second, some informants noted that the administrative coding (to ICD-10-AM codes) of hospital-based or registry-based data as a barrier to timely use of linked data for improving morbidity and mortality estimates. Third, and among jurisdictions that incorporated notifiable disease data into an MLS, scheduled delays in sending data to, or receiving data from, Data Linkage Units were also reported. These delays impacted on the acceptance of data linkage as a useful tool to augment surveillance and control practices at an operational level.

**Resources**

All informants identified a lack of resources as a barrier to using data linkage in the communicable disease setting. Many acknowledged that integrating notifiable disease data with disparate data in an enduring manner would be resource intensive at the outset, requiring dedicated staff to work with jurisdictional Data Linkage Units and relevant data custodians to establish data transfer and management protocols. Further, such a system would require staff within communicable disease units with skills to analyse, interpret and report on the resultant linked data. A lack of these skills was identified as a key factor precluding the use of data linkage by communicable disease units. Engagement with external agencies and research institutions with staff who are experienced in the analysis of linked health data was identified as a possible solution to this barrier.

**Discussion**

In this study, we identified limited use of data linkage by Australian state and territory communicable disease control units to inform local surveillance and control activities. None of the jurisdictions reported routinely making use of data linkage methodologies to improve completeness of notifiable disease surveillance data, nor to systematically quantify case ascertainment or reporting bias in local notifiable disease registers. Some jurisdictions reported semi-regular or ad hoc usage of aggregated linked health data to improve burden of disease estimates for selected communicable diseases. A majority of jurisdictions reported manually integrating data from unlinked and disparate datasets on a case-by-case basis. All jurisdictions expressed a desire for a more systematic approach to addressing gaps in communicable disease surveillance. Commonly cited priorities included using data linkage to improve completeness of Aboriginal and Torres Strait Islander, vaccination status and mortality outcomes in local and national communicable disease registers.

The inconsistent use of both linked and unlinked data has implications for the interpretation of national surveillance data, and may partially explain the considerable variability previously identified relating to the completeness of important data items captured nationally in the NNDSS and between jurisdictions. Variations in the manner in which data are sourced were evident from this review, such as for determination of mortality status. Most jurisdictions did not integrate local death registration data with notifiable disease registries: One had a data linkage process set up to do so, but the mortality data were not routinely incorporated into their local notifiable disease register; and another integrated mortality data through a systematic albeit manual process of cross-checking these two data sources. Similarly, variations in the determination of Aboriginal and Torres Strait Islander status by jurisdictions were also identified. Some jurisdictions reported using disparate datasets to improve completeness of this data item, but none did so via data linkage methodologies. Variations in data collection methods for this data item has implications for accurate burden of disease estimates among Aboriginal and Torres Strait Islander people and for consequent delivery of...
targeted prevention and control initiatives to those in greatest need. Data linkage is recognised as a valuable tool for improving Aboriginal and Torres Strait Islander identification, and has been identified to assist in the measurement of the Council of Australian Government Closing the Gap Performance Indicators. 44,45 Gaps in communicable disease surveillance data relating to vaccination status among relevant vaccine preventable disease notifications also remain. 2,7,46 This is despite all jurisdictions having access to AIR. The current challenges faced by communicable disease control units in integrating vaccination data from AIR into notifiable disease registries are largely logistical. Data from AIR must be looked up manually on a case-by-case basis and then transcribed into local notifiable disease registries. If an AIR record for the case is not available, vaccination status may be gleaned from the case’s general practitioner, treating clinician, or from the case directly (via self-report). While this commonly occurs for selected low-incidence vaccine preventable diseases – and with good completeness – it is impracticable to carry out this laborious process among high-incidence vaccine preventable diseases. This may be one explanatory factor for gaps in vaccination status among notifiable disease registers. An alternative reason may also relate to the completeness of data within AIR, which has been shown to underestimate childhood immunisation coverage by up to 5%. 46 The use of data linkage would address the first challenge, but not the second. In our study, none of the jurisdictions reported using data linkage methodologies to improve vaccination status in local notifiable disease registries. An alternative reason may also relate to the completeness of data within AIR, which has been shown to underestimate childhood immunisation coverage by up to 5%. 46 The use of data linkage would address the first challenge, but not the second. In our study, none of the jurisdictions reported using data linkage methodologies to improve vaccination status in local notifiable disease registries. We contend, that while the use of data linkage may not be practicable during an immediate public health response, gaps in surveillance data relating to vaccination status could be addressed retrospectively to improve completeness of this key variable in local and national surveillance systems, thereby also improving cross-jurisdictional evaluation of national immunisation programs. 34 Using the AIR in this manner would accord with the stated purposes of the register as set out in the Australian Immunisation Register Act. 47 Minimising inconsistencies in surveillance practices between jurisdictions could be addressed by adopting nationally consistent linkage and analytical approaches should be prioritised. Barriers associated with cross-jurisdictional linkage identified by others include issues with timely data access, project governance and administration, as well as data security, privacy, infrastructure and capability. 12,14,48-50 We identified additional barriers including a lack of perceived need, system factors and resources. Jurisdictions with small populations reported regularly using unlinked disparate data to fill surveillance gaps, which precluded a need for data linkage methodologies to be used for data cleansing purposes. While these practices serve small jurisdictions well for rare or low-incidence notifiable conditions, this approach is unlikely to be appropriate for jurisdictions serving larger population sizes, or for high-incidence notifiable conditions such as chlamydia and gonococcal infection, campylobacter and salmonellosis, influenza, pertussis, and varicella zoster virus. 7 Improved completeness of surveillance data relating to these and other vaccine-preventable conditions is essential for vaccine program evaluation, vaccine safety and adverse event monitoring, which could be met through the systematic application of data linkage methodologies. An additional barrier identified in this study related to the de-identified form in which communicable diseases data are transmitted to the NNDSS, thus precluding systematic data linkage for data cleansing purposes occurring at a national level. In view of this barrier, many of the key informants noted that data linkage – for the purpose of data cleansing – could in theory be conducted by the jurisdictions, with the resultant data incorporated into local notifiable disease registries for transmission to the NNDSS. The process of incorporating data gleaned from linkage activities back into local notifiable disease registers essentially renders the data ‘re-identified’. Despite this, many of the key informants also noted that data obtained in this manner would accord with the local and national public health legislation, which governs the collection, use and disclosure of information relating to communicable diseases for the public health purposes. 3 Such legislation also sets out a regime for jurisdictions to use data to enhance the understanding of the epidemiology of communicable diseases. These legislative tools were all identified as enabling factors to the use of data linkage in this manner. There is, however, conflict between this use of data linkage for public health and safety purposes, versus the traditional use of linked data for research purposes. Data linkage systems are built around the concept of preserving an individual’s privacy while simultaneously releasing the value of stored data for de-identified health research. 51 This best practice approach is appropriate for the release of data to researchers but does little to address underlying gaps in local and national surveillance systems. Further work will be required to better understand public awareness and attitudes relating to the use of data linkage in this manner. Previous research exploring public attitudes to the linkage and sharing of health and administrative data in the UK identified that, in general, people are supportive of their data being used in this manner if there is a personal or societal benefit to do so. 52-54 The cornerstone of good public health surveillance is the systematic collection, analysis, interpretation and use of data to inform public health policy. 53 Traditional data sources to inform communicable disease surveillance are notifications made by medical practitioners and laboratories, augmented by active case investigation and follow-up. However, as evidenced by this review, a range of disparate and routinely collected data sources are commonly used. We suggest that a more systematic approach be taken to make better use of available data to improve communicable diseases surveillance and control activities in Australia, whereby commonly used disparate data are routinely integrated and used in the surveillance and response policy cycle. 55 Specifically, we suggest that:

• Data from disparate but routinely collected data sources identified in this review are integrated into existing communicable disease surveillance systems at the jurisdictional level.

• Standardised methods for linkage and analyses of the resultant linked data are developed, piloted and implemented to improve data quality and ascertainment.

• Engagement between data custodians, linkage analysts and policy-makers within government, as well as researchers, patients and community groups external to government, is prioritised to improve interpretation, transparency and use of data to support prevention and control efforts.

• Intelligence gleaned from data linkage activities is incorporated into existing communicable disease policy and planning
networks, and national reporting and monitoring frameworks. Such an approach would align with the national framework for communicable disease control, which identified that linkages with existing datasets should form part of a modernised surveillance system. Further, it would improve availability and use of data for policy and program development as demonstrated in other parts of the world. The UK, for example, has committed to integrating a range of disparate datasets with bloodborne virus surveillance systems to inform the care and management of people tested for and diagnosed with these conditions. And several countries in Europe routinely link national communicable disease surveillance systems to inform vaccine safety and effectiveness studies. Finland, for example, has established real-time data linkage between their national immunisation and communicable disease registers whereby data from disparate and routinely collected sources are systematically integrated and used in the surveillance and response policy cycle. The development of standardised linkage and analytical approaches, as well as improved multi-sectoral and community-level engagement, will be important for the successful use of data linkage in the communicable disease setting setting.

Conclusion
Australia has a robust surveillance system in place to prevent, monitor and control a range of communicable diseases of public health importance. Despite this, gaps in surveillance data remain, impacting on the system’s utility to support the development of evidence-based prevention and control efforts. Data linkage is recognised as a valuable method to close surveillance gaps and enhance the value of publicly held health data. To our knowledge, this is the first time that jurisdictional practices relating to the use of data linkage in the communicable disease setting in Australia have been described. We identified a range of barriers to and enablers of the use of data linkage in the communicable disease setting in Australia. In the absence of a nationally integrated data linkage system or interoperable communicable disease surveillance system, we suggest enhancements to state and territory-based communicable disease registers whereby data from disparate and routinely collected sources are systematically integrated and used in the surveillance and response policy cycle. The development of standardised linkage and analytical approaches, as well as improved multi-sectoral and community-level engagement, will be important for the successful use of data linkage in the communicable disease setting.

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