March 2018

*Clostridium difficile* infection

A model to improve the management and control of *Clostridium difficile* in Australia
Improving the prevention and control of *Clostridium difficile* infection in Australia
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Summary

In order to identify the measures that are needed in the future to maintain low rates of *Clostridium difficile* infection (CDI) in Australia, the Australian Commission on Safety and Quality in Health Care (the Commission) established a Community of Practice (CoP) in October 2016 to investigate the variations and gaps associated with the current surveillance and management of *Clostridium difficile* infection. Using the driver diagram methodology, the CoP identified the barriers affecting CDI surveillance in Australia and used this information to identify targeted solutions to address each of these barriers. Each solution was then assessed for ease of implementation and scope of impact.

The CoP identified that improvements to the management and prevention of CDI in Australia were impeded by knowledge gaps, practice variations and practical constraints related to laboratory testing, clinical case management, hospital-based surveillance and uncertainty about the burden of disease in the community. The following actions were identified by the CoP as a priority to improve CDI prevention and control in Australia:

1. Recommend that the Australian Infection Prevention and Control guidelines include a requirement for contact precautions for all patients with diarrhoea until pathology results are known
2. Develop educational resources for clinicians and bed managers on bed placement priorities for patients with diarrhoea or known/suspected CDI
3. Develop a national monitoring mechanism for healthcare associated CDI
4. Update the ASID/ACIPC 2011 infection control position statement
5. Disseminate guidance to promote the requirement for contact precautions, including appropriate bed placement, for all patients with diarrhoea until pathology results are known
6. Review the barriers to appropriate CDI testing
7. Perform a desktop audit of all CDI clinical management and treatment guidelines available in Australia and identify inconsistencies
8. Review utility of other indicators for providing more meaningful outcome data to drive practice change
9. Review utility of current hospital administrative data, hospital laboratory surveillance and identify and review any other mechanisms for monitoring CDI in the community.

Given that work has already been undertaken to address Actions 1, 3 and 4, the CoP recommends that future work is undertaken to address the remaining priority actions:

- Developing national educational resources on bed placement prioritisation (Action 2)
- Reinforcing the need for contact precautions, including appropriate bed placement, at the jurisdictional and health service level (Action 5)
- Expanding the current knowledge base on CDI laboratory testing, case management and surveillance of community exposure through targeted research (Actions 6-9).

The CoP has also scoped which outstanding actions are within the remit of the Commission and which actions are best led by other agencies or organisations.
Background

**Clostridium difficile in Australia**

**Prevalence**

Between 2011 and 2016 the average rate of *Clostridium difficile* infection (CDI)-related diagnoses in Australian public hospitals was 4.0 diagnoses per 10,000 patient days.\(^1\) As seen in Figure 1, the rate of CDI-related diagnoses peaked in early 2012 (5.0 diagnoses per 10,000 patient days) and again in late 2012 (4.9 diagnoses per 10,000 patient days). Previous work suggested that these peaks may have been the result of changes in laboratory testing processes and the emergence of ribotype 027; this latter theory however has been since discounted given that there has been little circulation of ribotype 027 has been observed in Australia to date.\(^3,4\)

**Figure 1. Monthly rate of A047 diagnoses in Australia, 2011-2016\(^1\)**

The rate of CDI diagnosis in Australia is comparable to recent infection rates reported in the United Kingdom (UK). This is despite considerable circulation of ribotype 027 in Europe in the early 2000s.\(^6,7\) In the United Kingdom (UK), strict adoption infection control measures, including mandatory reporting in 2004 and enhanced surveillance and routine ribotyping in 2007, was required to contain this particular strain.\(^8,9\) Prolonged adherence to these measures had a significant effect on the CDI rate in the UK: the CDI rate dropped from 14.9 cases per 10,000 patient days in 2007-08 to 3.67 cases per 10,000 patient days in 2016-17.\(^10\)

The rate of CDI diagnosis varies between states. As seen in Table 1, there is a 10-fold difference in the rate of diagnosis between the states. In 2015 the highest rate was in
the Australian Capital Territory (6.96 CDI diagnoses per 10,000 bed days); the lowest rate observed was in the Northern Territory (0.66 CDI diagnoses per 10,000 bed days).

Table 1. Rate of CDI diagnosis by state, 2015

<table>
<thead>
<tr>
<th>State</th>
<th>Total Bed Days</th>
<th>Rate of CDI diagnosis per 10,000 bed days</th>
</tr>
</thead>
<tbody>
<tr>
<td>Australian Capital Territory</td>
<td>403,765</td>
<td>6.96</td>
</tr>
<tr>
<td>New South Wales</td>
<td>6,388,427</td>
<td>4.35</td>
</tr>
<tr>
<td>Northern Territory</td>
<td>346,379</td>
<td>0.66</td>
</tr>
<tr>
<td>Queensland</td>
<td>3,702,250</td>
<td>2.43</td>
</tr>
<tr>
<td>South Australia</td>
<td>1,479,654</td>
<td>3.75</td>
</tr>
<tr>
<td>Victoria</td>
<td>4,889,193</td>
<td>3.91</td>
</tr>
<tr>
<td>Western Australia</td>
<td>1,938,631</td>
<td>4.63</td>
</tr>
<tr>
<td>Tasmania</td>
<td>396,435</td>
<td>4.96</td>
</tr>
</tbody>
</table>

As the rate of hospital-diagnosed CDI only accounts for symptomatic illness requiring hospital care and treatment and does not account for individuals with asymptomatic CDI or milder cases of CDI where hospitalisation is not required, it is likely that the burden of CDI in the Australian population is larger. Analysis by the Australian Commission on Safety and Quality in Health Care (the Commission) in 2017 indicates that healthcare delivery is the cause of CDI acquisition in fewer than 25% of hospital admissions with a CDI diagnosis. Instead, it is likely that the majority of CDI acquisition occur prior to hospitalisation, either in the community or from a prior episode of care.¹

Severe disease
Currently only Victoria collects surveillance data on severe CDI consistent with the Implementation Guide for Surveillance of Clostridium difficile Infection.¹²⁻¹⁴ Surveillance data regarding severe CDI from 2011 to 2017 were provided by VICNISS. Using this data it is estimated that 2.2% of all CDI cases result in severe disease and 0.7% CDI cases result in death due to severe disease. It is estimated that this equated to 112 cases of severe disease and 45 deaths from severe CDI in Australia in 2015. These estimates are useful given that the true burden of severe disease and mortality in Australia is unknown due to gaps in current CDI surveillance.
Existing infection prevention and control strategies for Australian healthcare settings

There is a suite of existing infection prevention and control strategies that should be used to prevent and contain the spread of CDI in Australian healthcare settings. These strategies include:

Hand hygiene
To minimise the spread of infection, hand hygiene should always be employed by healthcare workers and other people caring for patients with, or suspected of having, CDI. While using soap and water to perform hand hygiene is more effective against *Clostridium difficile* than alcohol-based hand rub (ABHR), hand hygiene with ABHR is effective if gloves have been used during patient care as glove use substantially reduces the microbial load present on hands. If gloves have not been used, hand hygiene should be performed using soap and water and then dried thoroughly with a single-use towel to ensure the removal of the bacteria and its spores.

Standard and transmission-based precautions
Standard precautions should be employed when caring for all patients, regardless of whether CDI is present or suspected. Given that CDI is spread by contact transmission, contact precautions should also be employed when caring for patients with, or suspected of having, CDI in order to minimise the spread of the pathogen. Specific requirements under contact precautions include:

- Use of gloves and gown in the patient area when in direct physical contact with the patient and their immediate environment
- Use of single use equipment or reusable equipment that is dedicated to the affected patient and undergoes appropriate reprocessing between patients
- Ideally, a patient suspected of having or known to have CDI should be placed in a single room with dedicated bathroom facilities
- Minimising unnecessary movement of affected patients around the healthcare facility
- Increased frequency of environmental cleaning to include daily cleaning using detergent and disinfectant, of frequently touched surfaces and objects, and thorough terminal cleaning of the patient room on discharge.

Antimicrobial stewardship
It is critical that strategies are in place to ensure appropriate antimicrobial use given the possible link between antibiotic exposure in hospital and CDI. Appropriate antimicrobial usage refers to appropriate clinical indication, dosage, duration and route of administration.

Surveillance and reporting
Infection surveillance also informs understanding of current and local disease incidence and prevalence and epidemiology and can be used to identify areas for improvement and innovation. Relevant surveillance findings should be reported back to surveillance teams and relevant clinicians in a timely manner to enable an informed interpretation of surveillance findings, as well as engage clinicians to identify opportunities to improve their own clinical practice. Health service organisations are required to undertake infection surveillance as per the National Safety and Quality Health Service Standards. In addition, CDI surveillance is a specific requirement of the national Performance and Accountability Framework (PAF), which is the reporting instrument for the National Health Reform Agreement. The Commission has previously prescribed
a set of standardised data parameters to enable a consistent approach to CDI surveillance across Australia. Laboratory-based surveillance is used in most states and territories to monitor cases of hospital identified CDI as outlined in Table 2.

Consumer engagement

Healthcare consumers who have CDI, or who are at risk of getting CDI, should be provided with information on the cause of infection, disease effects and strategies consumers can use to prevent further disease spread. A consumer factsheet on CDI is available and will be updated in line with the 2018 update of the Australian Guidelines for the Prevention and Control of Infection in Healthcare.

Table 2. Summary of state and territory-based CDI surveillance programs

<table>
<thead>
<tr>
<th>State or Territory</th>
<th>Year that jurisdictional surveillance program commenced</th>
<th>Surveillance elements</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Hospital-identified</td>
</tr>
<tr>
<td>ACT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>NSW</td>
<td>2010</td>
<td>YES</td>
</tr>
<tr>
<td>NT</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Qld</td>
<td>2008†</td>
<td>YES</td>
</tr>
<tr>
<td>SA</td>
<td>2006‡</td>
<td>YES</td>
</tr>
<tr>
<td>Tas</td>
<td>2008†</td>
<td>YES</td>
</tr>
<tr>
<td>Vic</td>
<td>2010§</td>
<td>YES</td>
</tr>
<tr>
<td>WA</td>
<td>2010§</td>
<td>YES</td>
</tr>
</tbody>
</table>

*Only collects healthcare-associated, health facility onset
† The national surveillance definition for CDI has been used since 2009
‡ The national surveillance definition for CDI has been used since the beginning of the jurisdictional surveillance program

Reference material

In addition to the Australian Guidelines for the Prevention and Control of Infection in Healthcare, the Implementation Guide for Surveillance of Clostridium difficile infection, the national definition and calculation of hospital identified Clostridium difficile infection and the Clostridium difficile DSS, a number of other national agencies have developed material that informs on the clinical management of and the infection prevention and control of CDI.

Public Health Laboratory Network laboratory case definition

The Public Health Laboratory Network developed a standard laboratory case definition in 2016 to reduce the variation associated with the laboratory testing and diagnosis of CDI. Key aspects of case definition are:

- Diagnosis requires laboratory detection of C. difficile toxins or toxigenic C. difficile in faeces, rectal swab or bowel contents PLUS relevant clinical manifestations (diarrhoea, ileus, toxic megacolon or pseudomembranous colitis)
- Criteria for diagnostic testing
tests for toxigenic *C. difficile* should only be performed on unformed stool specimens (or gut contents from patients with diarrhoea), unless ileus is suspected.

- all adults and children over 2 years, who have been hospitalized for >48 hours and develop diarrhoea (>3 unformed stools on a 24-hour period) should be tested for CDI.

- all adults and children over 2 years, in whom diarrhoea has persisted for >48 hours and no other enteropathogen has been identified should be tested for CDI.

- repeat testing of faecal specimens during the same episode of diarrhoea is not recommended a) within 4 weeks of a positive test (response to treatment is determined by clinical criteria) or b) following a negative test – unless CDI is strongly suspected and a more sensitive method (e.g. Nucleic Acid Amplification Testing) is used after a negative immunoassay.

- tests for *C. difficile* in children <2 years old should be performed in consultation with a paediatrician.

The laboratory case definition provides a description of the different types of laboratory tests available to identify CDI, but does not indicate a preferred testing strategy.

**Australian Infection Control Association and Australasian Society of Infectious Diseases infection control position statement**

In 2011 the Australian Infection Control Association (now the Australasian College of Infection Prevention and Control (ACIPC)) and the Australasian Society of Infectious Diseases (ASID) published a position statement summarising strategies that should be in place in healthcare facilities to prevent the spread of CDI.17 The position statement reinforces the need for infection surveillance, hand hygiene, contact precautions, environmental cleaning, staff and patient education, and outbreak management as effective mitigation against CDI. The position statement was disseminated as an open-access article in Healthcare Infection (the official journal of the former Australian Infection Control Association).

**Australasian Society of Infectious Diseases guidelines**

In response to there being little clinical experience with severe CDI at the time, the Australasian Society of Infectious Diseases (ASID) published guidelines in 2011, with an update in 2016, to inform clinicians on the best practice approaches for CDI diagnosis and treatment.31, 32 These guidelines provide advice relevant to the principles of diagnosis, patient assessment, use of laboratory methods, disease prevention and management, antibiotic therapy and patient monitoring.
2017 CDI Community of Practice

Introduction

Since 2012, the rate of CDI diagnosis in Australia has been relatively stable. In order to continue to reduce the overall disease burden and to prevent further disease spread, further effort is needed to ensure that the rate of disease in this country remains low.

The purpose of this paper is to report on the work undertaken by a Community of Practice that was established to identify what future measures need to be put in place to further improve CDI prevention and control in Australia.

Methods

CDI Community of Practice

A community of practice (CoP) was established in October 2016 with the objective of examining the practice variations and gaps associated with the monitoring and management of CDI in Australia. Membership of the CoP is included in Appendix 1. The main functions of the Community of Practice were to:

- Identify opportunities to address existing and emerging practice variations and gaps in CDI surveillance
- Examine the predictors of best practice CDI surveillance
- Explore and develop models to integrate CDI surveillance data into quality improvement activity
- Promote and disseminate the learnings obtained from projects undertaken by members.

Process

The CoP met every six weeks via teleconference between October 2016 and October 2017 (9 meetings). The key activities of each teleconference are detailed in Figure 2 and key elements of the process are described below.
Structured discussions focussed on better understanding the cause of a single knowledge gap, practice variation or practical constraint. Discussion firstly centred on identifying all the underlying barriers that contributed to the practice gap, variation or constraint, and the relevant stakeholders that need to be engaged to overcome these barriers. A diagrammatic representation of this discussion is included in Appendix 2. Potential solutions were then workshopped to address each barrier.

The discussion template was used to focus dialogue within the group. This template was also used to record the discussion. Completed templates on laboratory testing, clinical case management, hospital based surveillance and community burden and impacts are included in Appendix 3.

Driver diagrams
A series of driver diagrams were built using the information gathered from the structured discussions. According to the Institute of Healthcare Improvement, a driver diagram illustrates the relationship between a project aim, the primary drivers that contribute to the achievement of the project aim, the secondary drivers that underpin the primary drivers and the specific change ideas that respond directly to the secondary drivers. It was necessary to develop five separate driver diagrams in order to reflect that change and improvement was required at the national level, the jurisdictional and
local health service level and within the broader community and that change also need to occur in the context of disease prevention, clinical management and community

Each driver diagram begun with the same aim - to improve the control and management of CDI in Australia - and the same primary drivers. Primary drivers were determined from the practice variation, knowledge gaps and practical constraints identified in the structured discussions. Secondary drivers and change ideas were populated based on the barriers and solutions identified in the structured discussions and their applicability to the context at hand. Change ideas that were related to at least two secondary drivers were highlighted for priority assessment. The driver diagram series is included Appendix 4.

**Action prioritisation**

Highlighted change ideas were assessed by the CoP on the basis of ease of implementation and potential impact. Ease of implementation was determined as easy or hard. Change ideas that were easy to implement were those that were able to be rapidly actioned, not dependent on other actions being undertaken or were relatively close to the status quo. Potential impact was judged as low impact (i.e. few people would benefit or be impacted upon by the change) or as a high impact (i.e. many people would benefit).

Change ideas that were deemed easy to implement and of high impact were considered as priority actions for future testing and implementation. The completed action prioritisation grid is included in Appendix 5.

**Context mapping**

Change ideas were also mapped back to the variations, constraints and knowledge gaps affecting CDI management and control in Australia in order to visualise the specific contexts in which these actions would likely have greatest effect. Context maps are included in Appendix 6.

**Findings**

**Variation, gaps and constraints**

The group identified that improvements to the management and prevention of CDI in Australia were impeded by a mixture of knowledge gaps, practice variations and practical constraints related to laboratory testing, clinical case management, hospital-based surveillance and uncertainty about the burden of disease in the community.

**Laboratory testing**

Currently there is an inconsistent approach to laboratory testing for CDI across Australia which impairs understanding of how much disease is in circulation. This inconsistency has arisen because there is wide variation in the testing practices of individual laboratories and, more broadly between private and public laboratories. For example, in some settings all diarrhoeal specimens are routinely screened for CDI whereas in other settings, diarrhoeal specimens will only be tested for CDI only if requested by the doctor.

It was highlighted that the practical constraints that lead to inadequate pre-analytic preparation, such as inappropriate storage and transportation, may also impact on the ability to control disease spread. Inadequate pre-analytic preparation results in poor specimen quality which may affect both whether laboratory testing can be performed and
the quality of the results generated. In turn this can limit the usefulness of laboratory testing for clinical management and infection prevention and control purposes. Additionally, there is also potential for inappropriate and unnecessary diagnostic testing to occur which may further question the validity of current laboratory-based surveillance methods.

**Clinical case management**

Considerable unwarranted clinical variation is likely to occur given that there is no national standardised approach used to assess, manage and treat patients with CDI. While the ASID guidelines do exist, these are not necessarily adopted in all clinical settings. As a result, there is still much variation in the clinical management approaches used by individual clinicians as well as variation between health services. The lack of a standardised clinical management approach, combined with diagnostic uncertainty, may result in delayed and inconsistent clinical decision-making and patient placement decisions.

It also remains largely unknown what current clinical management strategies are effective in reducing the disease spread as the ASID guidelines have not been evaluated for effect and there has been very little investigation into which strategies are most effective for mitigating disease spread in the community.

**Hospital-based surveillance**

The utility of existing hospital-based infection surveillance programs for infection prevention and control purposes is limited. Firstly, current hospital-based surveillance methods (i.e. detection of hospital-identified CDI) cannot be used to distinguish the burden of CDI acquired directly from the delivery of health care. Enhanced surveillance classification is needed to provide this level of detail however to do this requires extra resourcing and is cost-prohibitive for many health services. Without this additional information it is difficult to identify where to focus local infection prevention and control efforts.

Secondly, the data generated from hospital-based infection surveillance programs cannot be used to inform national infection control programs because there variation between states regarding denominators (despite the publication of a national CDI definition) and data validation. As such state-level data cannot be aggregated with a high level of confidence. Lastly, existing hospital-based infection surveillance programs are not useful for identifying CDI outbreaks or monitoring disease recurrence, meaning that others methods need to be employed in parallel to track disease spread.

**Uncertainty about the burden of disease in the community**

Control and prevention of CDI in Australia is hampered very much by a lack of knowledge about the transmission of CDI in community. Specifically, there is little understanding about the risk factors for acquisition of CDI in the community and the relationship between hospital-acquired CDI and community-acquired CDI. Developing this knowledge is hard at the moment, given that a wide spectrum of disease exists in the community and there are no surveillance definitions that presently suitable for this setting. As a result of this knowledge gap, it is unclear whether the burden of CDI in community is indeed problematic.
Priorities for future action

Improving CDI prevention and control is currently limited by a number of knowledge gaps, practice variations and practical constraints related to laboratory testing, clinical management and surveillance of CDI. In order to further reduce the overall disease burden and to prevent further disease spread, nine change ideas have been identified to address these gaps, variations and constraints and, in turn, improve CDI prevention and control in Australia. These change ideas, their current status, which context of effect and a proposal for how to progress each of these ideas are described in Table 3. Each change idea has been determined as a priority action because of its ease of implementation (i.e. can be readily actioned) and its anticipated broad impact (i.e. many people will benefit).
Table 3. Proposed change ideas to improve CDI prevention and control in Australia

<table>
<thead>
<tr>
<th>Change ideas identified by the Community of Practice</th>
<th>Current status and proposed action</th>
<th>Context of effect</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>1. Recommend Australian Infection Prevention and Control guidelines include requirement for contact precautions for all patients with diarrhoea until pathology results are known</strong></td>
<td>Updated Australian Infection Prevention and Control Guidelines is due for release in 2018. Recommendation 23 is relevant to this action: &quot;Recommendation 23: It is suggested that contact precautions, in addition to standard precautions, are implemented in the presence of known or suspected infectious agents that are spread by direct or indirect contact with the patient or the patient's environment.&quot; “A single-patient room is recommended for patients who require contact precautions.&quot; <strong>Proposed action:</strong> It is proposed that states, territories and individual health service organisations continue to support the implementation of contact precautions, including appropriate bed placement, in line with the updated Australian Guidelines for the Prevention and Control of Infection in Healthcare.</td>
<td>✓ ✓</td>
</tr>
<tr>
<td><strong>2. Develop educational resources for clinicians and bed managers on bed placement priorities for patients with diarrhoea or known/suspected CDI</strong></td>
<td>No action has been undertaken to date. <strong>Proposed action:</strong> It is proposed that the Commission adapt existing state resources (listed under (5)) as national guidance.</td>
<td>✓</td>
</tr>
<tr>
<td><strong>3. Develop national monitoring mechanism for healthcare acquired CDI</strong></td>
<td>As reported at the October 2017 meeting, the Commission has developed a mechanism to monitor hospital-identified CDI using administrative data available from the Admitted Patient Care National Minimum Data Set. The ongoing use of this approach was supported by the Interjurisdictional Committee in October 2017. Monitoring will commence in 2018. <strong>Proposed action:</strong> A snapshot report on 2016 administrative data will be provided at the April IJC meeting.</td>
<td>✓</td>
</tr>
<tr>
<td><strong>4. Update ASID/ACIPC infection control position statement</strong></td>
<td>This update is currently in progress. A consultation draft has been provided to ASID/ACIPC members. It is anticipated that the updated statement will published by mid 2018. <strong>Proposed action:</strong> A copy of the updated statement will be provided to the IJC for noting when it becomes available.</td>
<td>✓ ✓ ✓</td>
</tr>
</tbody>
</table>
Improving the prevention and control of *Clostridium difficile* infection in Australia

<table>
<thead>
<tr>
<th>Change ideas identified by the Community of Practice</th>
<th>Current status and proposed action</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>5. Disseminate guidance to promote single room placement and contact precautions for all patients with diarrhoea until pathology results are known</strong></td>
<td>SA: Some guidance was produced and disseminated within the context of the <em>management of patients with multi-resistant organisms</em> (2017). NSW: Produced and disseminated <em>clinician factsheet on patient placement</em> (2016). Qld: Some guidance was produced and disseminated within the context of the <em>management of patients with multi-resistant organisms</em> (2014). <strong>Proposed action:</strong> It is proposed that states, territories and individual health service organisations continue to support the implementation of contact precautions, including appropriate bed placement, in line with the updated Australian Guidelines for the Prevention and Control of Infection in Healthcare.</td>
</tr>
<tr>
<td><strong>6. Review barriers to appropriate CDI testing</strong></td>
<td>No action has been undertaken to date. <strong>Proposed action:</strong> It is proposed that the Commission will write to relevant Commonwealth agencies and professional colleges to encourage these groups to undertake targeted and exploratory research on these specific topics.</td>
</tr>
<tr>
<td><strong>7. Perform desktop audit of all CDI clinical management and treatment guidelines in Australia and identify inconsistencies</strong></td>
<td>No action has been undertaken to date. <strong>Proposed action:</strong> It is proposed that the Commission will write to relevant Commonwealth agencies and professional colleges to encourage these groups to undertake targeted and exploratory research on these specific topics.</td>
</tr>
<tr>
<td><strong>8. Review utility of other indicators for providing more meaningful outcome data to drive practice change</strong></td>
<td>No action has been undertaken to date. <strong>Proposed action:</strong> It is proposed that the Commission will write to relevant Commonwealth agencies and professional colleges to encourage these groups to undertake targeted and exploratory research on these specific topics.</td>
</tr>
<tr>
<td><strong>9. Review utility of current hospital data for the monitoring CDI in the community</strong></td>
<td>The Commission has developed a mechanism to monitor hospital-identified CDI using administrative data made available from the Admitted Patient Care National Minimum Data Set. This mechanism allows for the monitoring of the CDI cases which present to hospital, including those cases that have been acquired prior to hospitalisation. It is unable to inform on the prevalence of CDI acquisition where hospitalisation is not required. Other mechanisms for monitoring CDI acquisition when hospitalisation does not occur need to be identified and evaluated. <strong>Proposed action:</strong> It is proposed that the Commission will write to relevant Commonwealth agencies and professional colleges to encourage these groups to undertake targeted and exploratory research on these specific topics.</td>
</tr>
</tbody>
</table>

**Context of effect**

- Improving prevention and control at the national level
- Improving clinical management at the national level
- Improving prevention and control at the jurisdictional or health service level
- Improving prevention and control in community settings
- Developing the evidence base to improve future efforts for CDI control and management

| **5.** | **Yes** | **Yes** |
| **6.** | | **Yes** |
| **7.** | | |
| **8.** | | **Yes** |
| **9.** | **Yes** | **Yes** |
Future action for the Commission

In response to the change ideas proposed by the CoP, the Commission has identified the following three pieces of work are needed to drive further improve the prevention and control of CDI in Australia.

1. National educational resources on bed placement prioritisation needs to be developed

Decision-making around patient bed placement is an area of substantial variation between individual health organisations and between jurisdictions. National guidelines recommend single room placement for patients who require contact precautions. Bed placement decisions, however, are impacted by a number of factors, including:

- Unknown infection status at the time of admission
- Configuration and availability of beds
- Availability of suitable patient equipment
- Competing demands for the limited number of single room available (e.g. other infectious diseases, protective isolation, patient palliation, patient security).

The ability to contain the spread of CDI in a hospital is much easier if affected patients can be placed in single rooms. However, hospital infrastructure and high patient volume affects the availability of single rooms. Jurisdictions and health services need to consider the risk of CDI transmission when making patient placement decisions, particularly when resource prioritisation is necessary. Given the current variation in patient placement decision-making across the country, there is a need to develop national educational resources that supports bed managers, patient flow personnel and other clinicians to make patient placement decisions that consistently minimises transmission risks and adequately manages competing resource demands.

Currently, New South Wales, South Australia and Queensland provide local guidance to health services on resource prioritisation in the context of managing the spread of infectious disease. The development and dissemination of national educational resources addressing this topic would be very useful for health services in other states as well as those in the private sector.

To ensure the utility of educational resources in preventing disease spread, it is important that the resources address the symptomatic presentation of disease, such as diarrhoea and vomiting, rather than just the presence of a CDI diagnosis as this information is not always available on admission. Addressing symptomatic disease rather than disease aetiology will also expand the usefulness of these resources beyond CDI and will enable local decision-makers to use a consistent bed placement approach for all acute gastroenteritis presentations.

2. Support for contact precautions at the jurisdictional and health service level

The updated Australian Guidelines for the Prevention and Control of Infection in Healthcare will provide a recommendation for contact precautions in the presence of known or suspected infectious agents. To ensure the application of this recommendation at the bedside, it is critical that jurisdictional health departments and individual health services continually support clinicians to employ contact precautions, including appropriate bed placement, for known or suspected CDI cases. Clinicians need to also be made aware that suspected CDI cases include those patients who have presented with
diarrhoea but do not yet have definitive laboratory results. This information needs to be disseminated as part of ongoing clinician education and through different teaching modes (e.g. teaching on the run, clinical assessment, in-service training, promotional activity, peer learning).

Enabling additional single room capacity should also be considered by jurisdictions and individual health services as part of routine service planning cycles. Provisions for single room capacity should be in line with the specifications provided in Part D of the Australian Health Facility Guidelines.37

3. More evidence is needed on CDI laboratory testing, case management and community-based surveillance methods

Several of the identified change ideas (Actions 6-9) indicate that further improvement in CDI prevention and control in Australia is contingent on the development of a more detailed evidence base with regards to the mitigation, surveillance and management of CDI. These change ideas are effectively targeted research questions which will provide critical information needed to inform more appropriate and accurate laboratory testing, more appropriate case management and more robust community-based surveillance. In particular, understanding of the extent of variation associated with clinical case management of patients with CDI is still very limited. As a first step to understanding whether this variation is a significant problem and why variation may be occurring, further work needs to be undertaken to establish the extent of variation among the clinical management guidelines currently used in Australia. Findings from this work can then be used to identify the optimal and most efficient practice for managing CDI.

As this work is outside the remit of the Commission, the CoP has suggested a number of key stakeholder groups which may be in a position to undertake these specific research questions (see Appendix 3). It is proposed that the Commission write to the identified stakeholders and encourage these groups to focus on addressing these specific research questions as part of their future work.
References

15. Oughton MT, Loo VG, Dendukuri N, Fenn S and Libman MD. Hand hygiene with soap and water is superior to alcohol rub and antiseptic wipes for removal of Clostridium difficile. Infection Control and Hospital Epidemiology. 2009; 30: 939-44.


Appendix 1 - Community of Practice membership

The CoP brought together a group of individuals who have a shared interest in improving and using surveillance data and infection prevention and control strategies to reduce the incidence of CDI in Australia. The following individuals were members of the CoP:

- Lisa Hall, Queensland University of Technology, Chair
- Ann Bull, VICNISS
- Philip Russo, Deakin University
- Leanne Frazer, Hunter New England Local Health District (NSW)
- Mareeka Gray, Queensland Health/Sydney Local Health District Mental Health services (NSW)
- Rebecca McCann, Healthcare Infection Surveillance Western Australia
- Tom Riley, Pathwest/University of Western Australia
- Brett Mitchell, Avondale College of Higher Education
- Fiona Wilson, Tasmanian Infection Prevention and Control Unit
- Leon Worth, VICNISS
- Allison Peterson, Healthcare Infection Surveillance Western Australia
- Simone Tempone, Healthcare Infection Surveillance Western Australia
- Janet Li, Liverpool Hospital (NSW)
- Jennifer Caldwell, Liverpool Hospital (NSW)
- Christine Cope, SA Health
- John Gerrard, Gold Coast University Hospital (Qld)
Appendix 2 - Variations, gaps and constraints in the current management and prevention of CDI in Australia

**Domain 1: Laboratory testing**
- Variation in the testing effort between public and private settings and jurisdictions
- CDI testing practices vary between labs
- CDI testing is not being routinely done for all diarrhoeal specimens
- CDI testing is dependent on the receipt of good quality specimens
- Structures not in place to support CDI testing outside of hospital settings
- The effect of overdiagnosis and underdiagnosis is unclear

**Domain 2: Clinical case management**
- Clinical management decisions are sometimes delayed because CDI status is unknown
- There is no nationally endorsed standard protocol for CDI case management
- Adequate environmental cleaning processes may not always be used
- Patients with diarrhoea or CDI are not always allocated to a single room
- Compliance with clinical management guidelines has not been evaluated
- There are few strategies in place to address the risk of post-discharge transmission

**Domain 3: Hospital based surveillance**
- Broader impact of the hospital-identified CDI is unclear
- There is a large work burden attached to current enhanced surveillance methods and outcomes are not meaningful
- Current surveillance definitions for recurrent CDI are difficult to apply because there is a lack of reference data sources available
- It is difficult to identify a CDI outbreak using current surveillance methods without typing results
- Uptake of current enhanced surveillance methods is poor
- There is variation in the denominators used by the states and territories
- Uptake and conduct of data validation varies between facilities

**Domain 4: Understanding community burden**
- Understanding of the risk factors community acquired CDI is limited
- The transmission dynamics between community and hospitals are unclear
- It is unclear whether the community burden of CDI is actually problematic
- There are no established surveillance definitions available to enable the surveillance of CDI across the broad community
- It is difficult to conduct surveillance in the community

Improving the prevention and control *Clostridium difficile* infection in Australia
### Appendix 3 - Structured discussions

<table>
<thead>
<tr>
<th>Domain</th>
<th>Issues</th>
<th>Why are these issues occurring? (Barriers)</th>
<th>Identified stakeholders</th>
<th>Estimated time frame</th>
<th>Possible solutions</th>
</tr>
</thead>
</table>
| A. LABORATORY TESTING | 1. There is no standard testing practice for CDI in the country | 1.1 The PHLN case definition provides a review of the literature but does not provide explicit recommendations for diagnostic testing | Public Health Laboratory Network  
Department of Health – MBS (Medical services advisory committee)  
Royal College of Pathologists of Australasian  
Private labs (Pathology Australia)  
Australasian Society for Infectious Diseases  
Australian Commission on Safety and Quality in Health Care  
Australian College of Infection Prevention and Control  
States and territories (IJC) | 1-2 years | • Develop and disseminate easy to read recommendations for clinicians on diagnostic testing  
• Mandate a recommended testing procedure  
• Review financial incentives/disincentives for CDI testing |
| | | 1.2 There is no national diagnostic testing algorithm for CDI | | | |
| | | 1.3 The level of compliance with the approaches described by PHLN is not monitored | Royal College of Pathologists of Australasian (Quality Assurance Program)  
National Association of Testing Authorities, Australia  
Public Health Laboratory Network  
States and territories (IJC) | 1-2 years | • Develop and carry out regular compliance surveys |
| | 2. There is variation in the testing effort between public and private settings | 2.1 Private laboratories do not have any incentive to test unformed stools if a laboratory order has not been placed for CDI testing | Department of Health  
Private labs (Pathology Australia)  
Australian Institute of Health and Welfare  
States and territories (IJC) | 1-2 years | • Develop and disseminate easy to read recommendations for who needs to be testing and when (expand on ASID management guidelines)  
• Create and disseminate factsheet on who needs to be testing and when  
• Make CDI nationally notifiable  
• Report CDI nationally |
| | | 2.2 There is no national diagnostic testing algorithm for CDI | | | |
| | | 2.3 There is a lack of clinician awareness of indications for CDI testing | | | |
| | 3. It is difficult to do micro testing in aged care environments | | REFER TO TABLE D. COMMUNITY | | |
| | 4. Laboratories often receive poor quality specimens for CDI testing | 4.1 The PHLN case definition provides information about specimen quality but this information has not been disseminated to clinicians working at the bedside. | Royal Australian College of General Practitioners  
Acute care physician networks (e.g. Agency of Clinical Innovation, professional societies)  
Therapeutic Guideline: Gastrointestinal | 1-2 years | • Disseminate case definition in Australian Doctor or Australian Family Physician  
• Perform desktop audit of all CDI management/treatment guidelines used in Australia and address inconsistencies between guidelines |
<p>| | | 4.2 Some clinicians are not aware of when and why specimens should be collected for CDI testing | | | |
| | | 4.3 The timeliness of sending specimens to the lab is not prioritised | | | |
| | | 4.4 The geographic distance between some hospitals and laboratories make it difficult to get specimens to the lab in a timely manner | | | |</p>
<table>
<thead>
<tr>
<th>5. Overdiagnosis and underdiagnosis of CDI is likely</th>
</tr>
</thead>
<tbody>
<tr>
<td>5.1 Some clinicians are not aware of when and why specimens should be collected for CDI testing</td>
</tr>
<tr>
<td>5.2 Asymptomatic patients are being tested and diagnosed with infection based on microbiological findings only (not based on overall clinical picture)</td>
</tr>
<tr>
<td>5.3 Symptomatic people with disease are not being tested for CDI and are not being diagnosed with infection</td>
</tr>
<tr>
<td>5.4 Some co-morbidities and medications replicate CDI symptoms and get treated as CDI without proper investigation</td>
</tr>
<tr>
<td>5.5 High risk patient groups are often subjected to overtesting and asymptomatic colonisation in these groups may be misinterpreted as infection</td>
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</table>

<table>
<thead>
<tr>
<th>6. CDI testing is not routinely done for all diarrhoeal specimens</th>
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</thead>
<tbody>
<tr>
<td>6.1 CDI is not included in the usual testing battery for diarrheal specimens</td>
</tr>
<tr>
<td>6.2 Clinicians are not aware that CDI is not included in routine diarrhoeal testing</td>
</tr>
<tr>
<td>6.3 Use of multiplex PCR may prohibit simultaneous CDI testing</td>
</tr>
<tr>
<td>6.4 Medicare and hospital billing influence the decision to order/not order the CDI test</td>
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</tbody>
</table>

**Clinical networks working with high risk groups**
(Haematology Society of Australia and New Zealand, Clinical Oncology Society of Australia, Transplantation Society of Australia and New Zealand, Renal Dialysis Society of Australia, Australian and New Zealand Society for Geriatric Medicine)

**A. LABORATORY TESTING**

- Factsheets for high risk patient populations e.g. elderly, paediatric, cancer patients
- Provide guidance as appendices in Australian Infection Prevention and Control Guidelines.

- Disseminate case definition in Australian Doctor or Australian Family Physician
- Perform desktop audit of all CDI management/treatment guidelines used in Australia and address inconsistencies between guidelines

- Review pros and cons of new technology and disseminate findings through Pathology or MJA

- Review financial incentives/disincentives for CDI testing
<table>
<thead>
<tr>
<th>Domain</th>
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<th>Why are these issues occurring? (Barriers)</th>
<th>Identified stakeholders</th>
<th>Estimated time frame</th>
<th>Possible solutions</th>
</tr>
</thead>
<tbody>
<tr>
<td>1. There is no standard protocol for case management</td>
<td>1.1 Clinical management guidelines from ACIPC are not current</td>
<td>National Health and Medical Research Council Australian Society for Infectious Diseases Australasian College of Infection Prevention and Control States and territories (Safety and Quality departments) Australian Commission on Safety and Quality in Health Care</td>
<td>2-5 years</td>
<td>- Recommend CDI to be included as a topic for review for next update of the Australian Infection Prevention and Control guidelines. - Investigate if the ASID/ACIPC clinical management guidelines for CDI are being updated.</td>
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<tr>
<td>2. Compliance with clinical management guidelines has not been evaluated</td>
<td>2.1 There is no system currently available to monitor compliance with guidelines</td>
<td>States and territories (Safety and Quality departments) Australian Commission on Safety and Quality in Health Care</td>
<td>1-2 years</td>
<td>- Survey healthcare facilities to assess compliance with CDI guidelines (although compliance with various guidelines is within the remit of Standard 3 of the NSQHC Standards). - Include as part of accreditation (inclusion in Std 3 Safety and Quality Improvement Guide or example material to support Safety and Quality Improvement Guide) - Measure compliance with CDI clinical management guideline as part of larger compliance surveys (e.g. compliance with transmission-based precautions)</td>
<td></td>
</tr>
<tr>
<td>3. Patients with CDI are not always allocated a single room</td>
<td>3.1 There is limited availability of suitable decision support tools for resource prioritisation</td>
<td>National Health and Medical Research Council Australian Society for Infectious Diseases (HICSIG) Australasian College of Infection Prevention and Control States and territories (Safety and Quality departments) Australian Commission on Safety and Quality in Health Care</td>
<td>1-2 years</td>
<td>- Develop resource prioritisation tool for the management of known and suspected patients. - Ensure guidelines address the requirement for contact precautions for patients with diarrhoea until pathology results are known (and if transmissible cause of diarrhoea found, then CP should continue).</td>
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<tr>
<td>4. Clinical management decisions are delayed</td>
<td>4.1 CDI status is not always included during clinical handover</td>
<td></td>
<td>1-2 years</td>
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<tr>
<td>5. Clinicians are not always aware of CDI status</td>
<td>5.1 Visibility of flags/alerts in some patient administration systems requires interrogation of the patient record (i.e. the flag is not obvious to clinicians)</td>
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<tr>
<td>6. Inadequate environmental cleaning processes are sometimes used</td>
<td>6.1 Usual cleaning products may not be sufficient for cleaning CDI-contaminated areas</td>
<td>National Health and Medical Research Council Australian Commission on Safety and Quality in Health Care Australasian College of Infection Prevention and Control States and territories (Safety and Quality departments, Environmental Cleaning services)</td>
<td>1-2 years</td>
<td>- Awaiting environmental recommendations from updated Australian Infection Prevention and Control Guidelines - Develop online education resources for environmental cleaning staff</td>
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Improving the prevention and control *Clostridium difficile* infection in Australia
| 7. Limited patient information is available about post-discharge care | 7.1 NHMRC consumer factsheet does not provide sufficient information on post-discharge care | National Health and Medical Research Council
Australian Commission on Safety and Quality in Health Care
States and territories (Safety and Quality departments) | 1 year | • Review and update NHMRC CDI factsheet with additional information with post discharge precautions |
<table>
<thead>
<tr>
<th>Domain</th>
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<th>Why are these issues occurring? (Barriers)</th>
<th>Identified stakeholders</th>
<th>Estimated time frame</th>
<th>Possible solutions</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1. Data validation is not done in all facilities</td>
<td>1.1 Data validation is not standard practice in all facilities</td>
<td>Australian Commission on Safety and Quality in Health Care States and territories (S&amp;Q departments)</td>
<td>2 years</td>
<td>• Develop and publish national specifications for CDI surveillance data validation</td>
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<td></td>
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<td></td>
<td>Private laboratories, Individual hospital services States and territories Office of Health Protection/Communicable Diseases Network, Australia</td>
<td>5 years</td>
<td>• Establish legislation, documentation (including definitions and guidelines) and processes required to make CDI a notifiable condition</td>
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<td></td>
<td></td>
<td>1.2 Jurisdictions have limited access to private laboratory data which makes it difficult to do data validation on cases identified in private hospitals</td>
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<td></td>
<td>2. There are varied processes for data validation</td>
<td>2.1 The incompatibility between some surveillance databases and other electronic data sources means manual workarounds often need to be employed to perform data validation</td>
<td>IT systems vendors State IT services</td>
<td>5 years</td>
<td>• Allocate specific funding to infection control units or health networks for surveillance</td>
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<td></td>
<td></td>
<td></td>
<td>Australian Commission on Safety and Quality in Health Care Hospital networks/districts States and territories (S&amp;Q departments) National Classification in Health group (ICD10) Individual hospital services</td>
<td></td>
<td>• Establish structures that enable existing surveillance databases to easily interface with existing data sources (e.g. patient admissions system, electronic medical record)</td>
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<td></td>
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<td>2.2 Scope of some hospital surveillance programs does not include data validation</td>
<td></td>
<td></td>
<td>• Develop and publish national specifications for CDI surveillance data validation</td>
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<tr>
<td></td>
<td></td>
<td>2.3 There is variation in data validation processes within health networks/health districts</td>
<td></td>
<td></td>
<td>• Explore a range of data sources to develop an algorithm to validate data</td>
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<tr>
<td></td>
<td>3. There is variations in the denominators used across states and territories</td>
<td>3.1 States are currently choosing their own denominators</td>
<td>Australian Commission on Safety and Quality in Health Care Hospital networks/districts States and territories (S&amp;Q departments)</td>
<td>2-5 years</td>
<td>• Harmonise state denominators: – reinforce that patient days be used as the denominator for reporting as per the national CDI definitions</td>
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<td></td>
<td></td>
<td>3.2 Completeness/incompleteness of surveillance data influences the denominators that being used</td>
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<td>• Replicate SABSI requirement for denominators, with relevant exclusions</td>
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<td></td>
<td>3.3 Historical IT specifications in various states are in place to collect patient bed days/OBDs/separations</td>
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<td>4. Poor uptake and a large work burden associated with enhanced surveillance</td>
<td>4.1 Too many elements in the enhanced surveillance data collection (attribution, severity, recurrence, antimicrobial use</td>
<td>Australian Commission on Safety and Quality in Health Care States and territories (S&amp;Q departments)</td>
<td>2-5 years</td>
<td>• Establish a smaller set of core data items for enhanced surveillance.</td>
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<tr>
<td></td>
<td></td>
<td></td>
<td>IT systems vendors State IT services</td>
<td></td>
<td>• Prioritise surveillance in high risk units (e.g. ICU with ANZICS)</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4.2 There is no standardised approach to enhanced surveillance</td>
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<td></td>
<td>• Establish better definitions and mechanisms to identify true burden associated hospital and community</td>
</tr>
<tr>
<td></td>
<td></td>
<td>4.3 Hospitals have limited IT and human resources available to support the data collection required for enhanced surveillance</td>
<td></td>
<td></td>
<td>• Review and update current enhanced surveillance definitions</td>
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<td></td>
<td></td>
<td>4.4 There is little motivation for doing enhanced surveillance for case attribution as it does not provide meaningful hospital performance outcomes</td>
<td></td>
<td></td>
<td>• Identify other indicators that provide more meaningful information for enabling practice change</td>
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<tr>
<td></td>
<td></td>
<td>4.5 Enhanced surveillance definitions are extremely difficult to interpret and apply</td>
<td></td>
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<td></td>
<td></td>
<td>4.6 Current data handling and management is not timely enough to have a positive effect on changing practice</td>
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</table>

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## C. Hospital-Based Surveillance

<table>
<thead>
<tr>
<th>Issue</th>
<th>Recommendation</th>
<th>Timeframe</th>
</tr>
</thead>
</table>
| 5. There is substantial work burden associated with doing enhanced surveillance for severity | - Make CDI severity status a mandatory surveillance data item  
- Prioritise severity surveillance over other enhanced surveillance data collection  
- Prioritise surveillance in high risk units (e.g. ICU with ANZICS)  
- Employ structured lab reporting/standardisation reporting specifications (AURA) to streamline laboratory data collection | 1-2 years |
| 6. It is difficult to do surveillance for recurrence | - Review and update current enhanced surveillance definition for recurrence  
- Explore other mechanisms for monitoring recurrence (e.g. snapshots)  
- De-prioritise recurrence for enhanced surveillance purposes | 1-2 years |
| 7. There is little understanding of the national burden of CDI in hospitals | - Develop a mechanism to monitor true healthcare associated CDI at a national level | 2-5 years |
| 8. It is difficult to identify a CDI outbreak | - Establish satisfactory baseline rate  
- Establish notification/legislation for CDI  
- Set outbreak case definition for CDI  
- Explore other mechanisms for monitoring outbreak potential that do not rely on diagnostic typing (e.g. Liverpool method where 2 cases or more in 1 ward constitutes an outbreak) | 2-5 years |
<table>
<thead>
<tr>
<th>Domain</th>
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<th>Why are these issues occurring?</th>
<th>Identified stakeholders</th>
<th>Estimated time frame</th>
<th>Possible solutions</th>
</tr>
</thead>
<tbody>
<tr>
<td>D. UNDERSTANDING THE COMMUNITY BURDEN</td>
<td>1. It is unclear whether the burden of CDI in the community is problematic</td>
<td>1.1 There are no suitable data collection systems or proxy markers available to determine the burden of CDI in the community 1.2 Have no understanding of the extent of normal disease endemicity in the community 1.3 No thresholds have been established to define CDI outbreaks 1.4 CDI testing (particularly in aged care) is sporadic, therefore even if private laboratory data was readily accessible to jurisdictions, the true burden may not be reflected</td>
<td>Australian Commission on Safety and Quality in Health Care Office of Health Protection/Communicable Diseases Network, Australia Royal Australasian College of General Practitioners Office of Health Protection State and territory Public Health Units Aged care facilities</td>
<td>• Adopt the Victorian continuous surveillance in aged care for 2017/2018 in aged care facilities across the country</td>
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<tr>
<td></td>
<td>2. Understanding of risk factors for community acquisition of CDI is limited</td>
<td>2.1 All reservoirs of infection in the community have not been definitively identified 2.2 The link between antimicrobial usage in the community and the impact on CDI acquisition in the community is not well understood</td>
<td>Academic research</td>
<td></td>
<td>• Increase uptake of acNAPS in across the country</td>
</tr>
<tr>
<td></td>
<td>3. It is difficult to do surveillance of CDI in the community</td>
<td>3.1 Disease may present asymptomatically and may not require medical attention, testing or treatment 3.2 It is difficult, expensive and not clinically appropriate to screen the community for CDI using laboratory methods</td>
<td>Office of Health Protection/Communicable Diseases Network, Australia Public Health Laboratory Network Royal Australasian College of General Practitioners National Prescribing Service</td>
<td></td>
<td>• Identify whether there are alternative non-laboratory mechanisms to monitor for CDI in the community</td>
</tr>
<tr>
<td></td>
<td>4. There are no established surveillance definitions to do comprehensive CDI surveillance in the community</td>
<td>4.1 Current surveillance definitions for aged care settings are not sensitive enough to identify asymptomatic infection 4.2 Besides those that exist for aged care settings, there are no other established CDI surveillance definitions for other community settings. 4.3 Aged care definitions may not be suitable in Australian settings</td>
<td>Australian Commission on Safety and Quality in Health Care Australian Aged Care Quality Agency Office of Health Protection/Communicable Diseases Network, Australia Public Health Laboratory Network</td>
<td></td>
<td>• Update current aged care definitions to enable better identification of asymptomatic infection</td>
</tr>
<tr>
<td></td>
<td>5. The transmission dynamics between community and hospitals are unclear</td>
<td>5.1 CDI status is not always available at time of transfer into and out of hospital</td>
<td>Australian Commission on Safety and Quality in Health Care Australian Aged Care Quality Agency State and territories (S&amp;Q departments) Individual hospital facilities Individual aged care facilities</td>
<td></td>
<td>• Mandate that CDI status be determined on hospital admission</td>
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Improving the prevention and control *Clostridium difficile* infection in Australia

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**Domain**: Understanding the Community Burden

**Issues**:

1. It is unclear whether the burden of CDI in the community is problematic
- There are no suitable data collection systems or proxy markers available to determine the burden of CDI in the community.
- No understanding of the extent of normal disease endemicity in the community.
- No thresholds have been established to define CDI outbreaks.
- CDI testing (particularly in aged care) is sporadic, therefore even if private laboratory data was readily accessible to jurisdictions, the true burden may not be reflected.

2. Understanding of risk factors for community acquisition of CDI is limited
- All reservoirs of infection in the community have not been definitively identified.
- The link between antimicrobial usage in the community and the impact on CDI acquisition in the community is not well understood.

3. It is difficult to do surveillance of CDI in the community
- Disease may present asymptomatically and may not require medical attention, testing or treatment.
- It is difficult, expensive and not clinically appropriate to screen the community for CDI using laboratory methods.

4. There are no established surveillance definitions to do comprehensive CDI surveillance in the community
- Current surveillance definitions for aged care settings are not sensitive enough to identify asymptomatic infection.
- Besides those that exist for aged care settings, there are no other established CDI surveillance definitions for other community settings.
- Aged care definitions may not be suitable in Australian settings.

5. The transmission dynamics between community and hospitals are unclear
- CDI status is not always available at time of transfer into and out of hospital.

**Estimated time frame**

- Adopt the Victorian continuous surveillance in aged care for 2017/2018 in aged care facilities across the country.
- Explore utility of hospital administrative data and laboratory surveillance data to develop a community-based surveillance system.
- Make CDI notifiable.
- Increase uptake of acNAPS in across the country.
- Develop primary care NAPS.
- Identify whether there are alternative non-laboratory mechanisms to monitor for CDI in the community.
- Monitor severe disease on admission separately to HCA inpatient severe disease.
- Update current aged care definitions to enable better identification of asymptomatic infection.
- Develop a community-based surveillance system.
- Mandate that CDI status be determined on hospital admission.
- Include CDI status on discharge/transfer summary.
- Establish a standardised stool testing protocol for all laboratories receiving specimens from residential aged care facilities.
Appendix 4 - Driver diagram series

Driver diagram 1: National actions to improve the control of CDI

**National Actions to Improve CDI Control**

**Problem:** Our ability to control and manage the spread of CDI in Australia is affected by knowledge gaps, practice variation and practical constraints.

**Aim:** To improve the control and management of CDI in Australia.

**Stakeholders:**
- ACSQHC
- OHP/CDNA
- State and Territory SA&Q departments
- ASID
- ACIPC
- NHMRC
- AIHW
- Aus Aged Care Quality Agency
- PHLN
- NCAS

**NATIONAL ACTIONS TO IMPROVE CDI CONTROL**

**Problem:** Our ability to control and manage the spread of CDI in Australia is affected by knowledge gaps, practice variation and practical constraints.

**Aim:** To improve the control and management of CDI in Australia.

**Stakeholders:**
- ACSQHC
- OHP/CDNA
- State and Territory SA&Q departments
- ASID
- ACIPC
- NHMRC
- AIHW
- Aus Aged Care Quality Agency
- PHLN
- NCAS

**Primary Drivers**
- Reduce variation in the testing effort for CDI
- Overcome practical constraints associated with CDI lab testing
- Address the knowledge gaps associated with CDI lab testing
- Reduce variation associated with the validation of hospital-based surveillance
- Overcome practical constraints associated with CDI surveillance
- Address the knowledge gaps associated with CDI case management
- Reduce the practice variation associated with clinical case management
- Overcome practical constraints associated with understanding CDI burden in the community
- Address the knowledge gaps about the CDI burden in the community

**Secondary Drivers**
- Reduce variation in how and when we test for CDI
- Reduce testing variation between public and private settings
- Improve the quality of specimen sent for testing
- Overcome technical limitations associated with using multiplex PCR assays to test diarrhoeal specimens
- Improve understanding of the effect of overtreating and undertreating for high-risk patients
- Ensure that surveillance data is always validated
- Minimise variation in the data validation processes
- Reduce the number of denominators being used
- Improve uptake of exposure classification surveillance
- Reduce work burden associated with exposure classification
- Improve the utility of surveillance for outbreak management
- Mitigate difficulties associated with recurrence surveillance
- Identify and implement risk management processes
- Standardise the clinical management of CDI cases
- Reduce delays associated with clinical management
- Recommend single-room placement where possible
- Improve clinician awareness of CDI status
- Evaluate compliance with clinical management guidelines
- Increase access to patient information on post-discharge transmission risks and care
- Minimise difficulties associated with doing CDI surveillance in the community
- Improve CDI surveillance in community settings
- Improve understanding of the transmission dynamics between community and hospital settings
- Establish an understanding of whether community burden of CDI is problematic
- Improve understanding of the risk factors for community acquired CDI

**Change Ideas**
- Publish PHLN's CDI definition in Australian/Doctor/APR
- Develop spec for surveillance data validation
- Develop data validation algorithm
- Provide national guidance on prioritisation and de-prioritisation of exposure surveillance measures
- Request S&I to use patient bed days as denominator
- Request S&I to require SAABPI denominator requirements for CDI surveillance
- Focus enhanced surveillance in high risk areas
- Promote uptake of severity surveillance
- Review current enhanced surveillance definitions for usability
- Mandate surveillance of CDI severity status
- Establish outbreak case definition for CDI
- Establish baseline rate and outbreak thresholds
- Mandate reporting of CDI rates
- Review surveillance definition for recurrent CDI
- Develop online education resources re CDI for environmental cleaning staff
- Update ASDIA/ACPC infection control position statement
- Recommend Aus AEC guidelines include requirement for single room placement and contact precautions for all patients with diarrhoea and/or new onset abdominal pain
- Develop national prioritisation tool for management of patients requiring single rooms for control of CDI
- Develop educational resources for clinicians and bed managers on patient placement priorities for patients with diarrhoea or new onset abdominal pain
- Review and update NHMRC CDI consumer factsheet
- Develop national monitoring tool for management of patients with diarrhoea or new onset abdominal pain
- Mandate CDI a nationally notifiable disease
- Retine surveillance definitions for hospital and community acquired CDI
- Review quality of current hospital administrative data, hospital laboratory surveillance and identify and review any other mechanisms for monitoring CDI in the community
- Update current surveillance definitions to enable better detection of asymptomatic infection
- Develop primary care NAPs
- Mandate adoption of eNAPs nationally

**Key**
- Practice variation
- Practical constraint
- Knowledge gap
- Addresses 2 or more secondary drivers
Driver diagram 2: National actions to improve the clinical management of CDI

NATIONAL ACTIONS TO IMPROVE THE CLINICAL MANAGEMENT OF CDI

PROBLEM:
Our ability to control and manage the spread of CDI in Australia is affected by knowledge gaps, practice variation and practical constraints.

AIM:
To improve the control and management of CDI in Australia.

STAKEHOLDERS:
- PHLN
- ACSQHC
- NHMRC
- RACP (QAP)
- Pathology Australia
- Private labs
- ASID
- ACIPC
- NATA
- Acute care physician clinical networks
- TG. Gastrointestinal

KEY
- Practice variation
- Practical constraint
- Knowledge gap
- Addresses 2 or more secondary drivers

PRIMARY DRIVERS
- Reduce the variation in the testing effort for CDI
- Overcome practical constraints associated with CDI lab testing
- Answer address the knowledge gaps associated with CDI lab testing
- Reduce variation associated with the validation of hospital-based surveillance
- Overcome practical constraints associated with CDI surveillance
- Answer address the knowledge gaps associated with CDI surveillance
- Reduce the practice variation associated with clinical case management
- Answer address the knowledge gaps associated with CDI case management
- Overcome practical constraints associated with understanding CDI burden in the community
- Answer address the knowledge gaps about the CDI burden in the community

SECONDARY DRIVERS
- Reduce variation in how and when we test for CDI
- Reduce testing variation between public and private settings
- Reduce testing variation between jurisdictions
- Eliminate instances of over-testing and undertesting
- Ensure routine testing of all diarrhoeal specimens
- Improve the quality of specimens sent for testing
- Overcome technical limitations associated with using multiplex PCR assays to test diarrhoeal specimens
- Improve understanding of the effect of over-testing and undertesting for high-risk patients
- Ensure that surveillance data is always validated
- Minimise variation in the data validation processes
- Reduce the number of denominators being used
- Improve uptake of exposure classification surveillance
- Reduce work burden associated with exposure classification
- Improve the utility of surveillance for outbreak management
- Reduce the work burden associated with severity surveillance
- Mitigate difficulties associated with recurrence surveillance
- Improve understanding of the impact of hospital-acquired CDI
- Improve uptake of severity surveillance
- Identify optimal environmental cleaning processes
- Standardise the clinical management of CDI cases
- Reduce delays associated with clinical management
- Recommend single room placement where possible
- Improve clinician awareness of CDI status
- Evaluate compliance with clinical management guidelines
- Increase access to patient information on post-discharge transmission risks and care
- Minimise difficulties associated with doing CDI surveillance in the community
- Establish surveillance definitions for community settings
- Improve understanding of the transmission dynamics between community and hospital settings
- Establish understanding of whether community burden of CDI is problematic
- Improve understanding of the risk factors for community acquired CDI

CHANGE IDEAS
- Mandate a standard diagnostic testing procedure for CDI
- Examine feasibility of mandatory adoption of the UK’s two-step testing procedure
- Include guidance on when to test for CDI in high-risk populations in the AusPIC guidelines
- Develop survey tool to monitor compliance with the standard diagnostic testing procedure
- Create clinician fact sheet on indications for CDI testing
- Develop and implement a standardised stool testing protocol for all laboratories receiving specimens from residential aged-care facilities
- Publish PHLN’s CDI definition in Australian Clinical Guidelines
- Develop national premissum tools for management of patients requiring single rooms for contact precautions
- Update ASDAC/ACPG infection control position statement
- Recommend that compliance with national CDI treatment and management guidelines assessed as part of ISO 3 accreditation
- Develop national survey tools to monitor compliance with CDI treatment and management guidelines
- Perform desktop audit of all CDI clinical management and treatment guidelines in Australia and identify inconsistencies
- Promote use of AUHA’s structured tool reporting space
- Review and update NHMRC CDI consumer fact sheet
- Develop educational resources to clinicians and bed managers on bed placement priorities for patients with diarrhoea or known/suspected CDI
- Disseminate guidance to promote the requirement for single room placement and contact precautions for all patients with diarrhoea until pathology results are known

Improving the prevention and control Clostridium difficile infection in Australia
Driver diagram 3: Jurisdiction and health service actions to improve the control of CDI

**JURISDICTION AND HEALTH SERVICE ACTIONS TO IMPROVE CDI CONTROL**

**Problem:**
Our ability to control and manage the spread of CDI in Australia is affected by knowledge gaps, practice variation and practical constraints.

**Goal:**
To improve the control and management of CDI in Australia.

**Stakeholders:**
State and Territory S&Q departments
State and Territory IT services
State and Territory health education units
State and Territory State and Territory pathology providers
Private pathology providers

**Drivers:**
- **Primary Drivers:**
  - Reduce the variation in the testing effort for CDI
  - Overcome practical constraints associated with CDI lab testing
  - Overcome practical constraints associated with clinical case management
  - Answer the knowledge gaps associated with clinical case management
  - Answer the knowledge gaps associated with the burden of CDI in the community

- **Secondary Drivers:**
  - Reduce the variation in how and when we test for CDI
  - Reduce testing variations between public and private settings
  - Reduce testing variation between jurisdictions
  - Overcome technical limitations associated with using multiplex PCR assays to test diarrhoeal specimens
  - Overcome practical constraints associated with understanding CDI burden in the community
  - Answer the knowledge gaps about the CDI burden in the community

**Change Ideas:**
- Allocate specific funding to hospital networks to resource CDI surveillance
- Develop an online education resource for CDI for environmental cleaning staff
- Promote uptake of severity surveillance
- Develop online education resources/re CDI for environmental cleaning staff
- Disseminate guidance to promote the requirement for single room placement and contact precautions for all patients with diarrhoea until pathology results are known
- Promote use of AURA’s structured lab reporting forms
- Mandate that CDI status needs to be determined on admission
- Include CDI status on discharge/transfer summary templates
- Stricter severe disease by present on admission or not present on admission
- Incorporate measurement of compliance with CDI clinical treatment and management guidelines in large P&Q compliance surveys

Key:
- Practice variation
- Practical constraint
- Knowledge gap
- Addresses 2 or more secondary drivers
Driver diagram 4: Research actions to improve the control of CDI

**COMMUNITY HEALTH ACTIONS TO IMPROVE CDI CONTROL**

**PROBLEM:**
Our ability to control and manage the spread of CDI in Australia is affected by knowledge gaps, practice variation and practical constraints.

**AIM:**
To improve the control and management of CDI in Australia.

**STAKEHOLDERS:**
OHIP/CONA
PHLN
RACGP
RACP
NCAS
Primary Health Networks
Pathology Australia
Private Pathology labs
Individual primary care facilities
Individual residential care facilities

**KEY**
- Practice variation
- Practical constraint
- Knowledge gap
- Addresses 2 or more secondary drivers

**PRIMARY DRIVERS**
- Reduce the variation in the testing effort for CDI
- Overcome practical constraints associated with CDI lab testing
- Overcome practical constraints associated with CDI surveillance
- Reduce the variation associated with the validation of hospital-based surveillance
- Overcome practical constraints associated with clinical case management
- Reduce the practice variation associated with clinical case management
- Overcome practical constraints associated with understanding CDI burden in the community
- Overcome the knowledge gaps associated with CDI case management
- Overcome the knowledge gaps associated with the CDI burden in the community
- Address the knowledge gaps associated with CDI lab testing

**SECONDARY DRIVERS**
- Reduce variation in how and when we test for CDI
- Reduce testing variation between public and private settings
- Reduce testing variation between jurisdictions
- Eliminate instances of overtesting and undertesting
- Ensure routine testing of all diarrhoeal specimens
- Improve the quality of specimens sent for testing
- Improve understanding of the effect of overtesting and undertesting for high risk patients
- Ensure that surveillance data is always validated
- Minimise variation in the data validation processes
- Reduce the number of denominators being used
- Improve uptake of exposure classification surveillance
- Reduce work burden associated with exposure classification
- Improve the utility of surveillance for outbreak management
- Reduce the work burden associated with severity surveillance
- Improve uptake of severity surveillance
- Improve understanding of the impact of hospital-acquired CDI
- Mitigate difficulties associated with recurrence surveillance
- Identify optimal environmental cleaning processes
- Standardise the clinical management of CDI cases
- Reduce delays associated with clinical management
- Recommend single room patient placement where possible
- Improve clinician awareness of CDI status
- Evaluate compliance with clinical management guidelines
- Increase access to patient information on post-discharge transmission risks and care
- Minimise difficulties associated with doing CDI surveillance in the community
- Establish surveillance definitions for community settings
- Improve understanding of the transmission dynamics between community and hospital settings
- Establish understanding of whether community burden of CDI is problematic
- Improve understanding of the risk factors for community-acquired CDI

**CHANGE IDEAS**
- Publish PHLN's CDI definition in Australian Doctors'AFP
- Examine feasibility of national adoption of the UK's two-step testing procedure
- Mandate a standard diagnostic testing procedure for CDI
- Develop a solution to enable easier extraction of surveillance data from electronic data systems
- Stratify severe disease by present on admission or not present on admission
- Include CDI status on discharge/transfer summary templates
- Promote use of AU-RAN's structured lab reporting spec
- Establish better definitions for hospital and community-acquired CDI
- Review utility of other indicators for providing more meaningful outcome data to drive practice change
- Review utility of current hospital administrative data, hospital laboratory surveillance and identify and review any other mechanisms for monitoring CDI in the community
- Make CDI a nationally notifiable disease
- Examine utility of hospital administrative data and hospital lab data to develop a community-based surveillance system
- Mandate adoption of esICUAPs nationally
Driver diagram 5: Actions for further research and exploration

**RESEARCH AND EXPLORATORY ACTIONS TO IMPROVE CDI CONTROL**

**PROBLEM:**
Our ability to control and manage the spread of CDI in Australia is affected by knowledge gaps, practice variation and practical constraints.

**AIM:**
To improve the control and management of CDI in Australia.

**KEY**
- Practice variation
- Practical constraint
- Knowledge gap
- Addresses 2 or more secondary drivers

**PRIMARY DRIVERS**
- Reduce the variation in the testing effort for CDI
- Overcome practical constraints associated with CDI lab testing
- Reduce variation associated with the validation of hospital-based surveillance
- Overcome practical constraints associated with CDI surveillance
- Reduce the practice variation associated with clinical case management
- Overcome practical constraints associated with understanding CDI burden in the community
- Answer/address the knowledge gaps associated with CDI case management
- Answer/address the knowledge gaps about the CDI burden in the community

**SECONDARY DRIVERS**
- Reduce variation in how and when we test for CDI
- Reduce testing variation between public and private settings
- Eliminate instances of overtesting and undertesting
- Ensure routine testing of all diarrhoeal specimens
- Improve the quality of specimens sent for testing
- Overcome technical limitations associated with using multiplex PCR assays to test diarrhoeal specimens
- Improve understanding of the effect of overtesting and undertesting for high-risk patients
- Ensure that surveillance data is always validated
- Minimise variation in the data validation processes
- Reduce the number of denominators being used
- Improve uptake of exposure classification surveillance
- Reduce work burden associated with exposure classification
- Improve uptake of severity surveillance
- Reduce the work burden associated with severity surveillance
- Mitigate difficulties associated with recurrence surveillance
- Improve understanding of the impact of hospital-acquired CDI
- Improve the utility of surveillance for outbreak management
- Identify optimal environmental cleaning processes
- Standardise the clinical management of CDI cases
- Reduce delays associated with clinical management
- Recommend single room placement where possible
- Improve clinician awareness of CDI status
- Evaluate compliance with clinical management guidelines
- Increase access to patient information post-discharge transmission risks and care
- Minimise difficulties associated with doing CDI surveillance in the community
- Establish surveillance definitions for community settings
- Improve understanding of the transmission dynamics between community and hospital settings
- Establish understanding of whether community burden of CDI is problematic
- Improve understanding of other risk factors for community acquired CDI

**CHANGE IDEAS**
- Review of the impact of current $ incentives for CDI testing
- Review pros and cons of new diagnostic technology
- Develop specs for surveillance data validation
- Develop data validation algorithm
- Implement better CDI surveillance definitions
- Review current enhanced surveillance definitions for usability
- Review utility of other indicators for providing more meaningful outcome data to drive practice change
- Review surveillance definition for recurrent CDI
- Review utility of other mechanisms for monitoring recurrent CDI
- Use historical data to establish satisfactory baseline rate and outbreak thresholds
- Use international literature to establish satisfactory baseline rate and outbreak thresholds
- Review utility of other mechanisms for monitoring outbreaks potential
- Perform desktop audit of CDI clinical management and treatment guidelines in Australia and identify inconsistencies
- Review utility of current hospital administrative data, hospital laboratory surveillance and identify and review any other mechanisms for monitoring CDI in the community
- Refine current aged care definitions to enable identification of asymptomatic infection
- Develop primary care NAPS

**Figure 3. Driver diagram 5: Community-level actions to improve the control of CDI.**
### Appendix 5 - Action prioritisation

<table>
<thead>
<tr>
<th>Highlighted change ideas</th>
<th>Ease of Implementation (EASY/HARD)</th>
<th>Anticipated Impact (HIGH/LOW)</th>
<th>Priority for testing</th>
</tr>
</thead>
<tbody>
<tr>
<td>Publish PHLN’s CDI definition in Australian Doctor or AFP</td>
<td>EASY</td>
<td>LOW</td>
<td></td>
</tr>
<tr>
<td>Examine feasibility of national adoption of the UK’s two-step testing procedure</td>
<td>HARD</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Develop specs for surveillance data validation</td>
<td>HARD</td>
<td>LOW</td>
<td></td>
</tr>
<tr>
<td>Recommend Australian IP&amp;C guidelines include requirement for contact precautions for all patients with diarrhoea until pathology results are known</td>
<td>EASY</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Develop national resource prioritisation tool for management of patients with known or suspected CDI</td>
<td>HARD</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Develop educational resources for clinicians and bed managers on bed placement priorities for patients with diarrhoea or known/suspected CDI</td>
<td>EASY</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Develop national monitoring mechanism for healthcare acquired CDI</td>
<td>EASY</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Make CDI a nationally notifiable disease</td>
<td>HARD</td>
<td>LOW</td>
<td></td>
</tr>
<tr>
<td>Mandate a standard diagnostic testing procedure for CDI</td>
<td>HARD</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Develop survey tool to monitor compliance with the standard diagnostic testing procedure</td>
<td>HARD</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Create clinician factsheet on indications for CDI testing</td>
<td>EASY</td>
<td>LOW</td>
<td></td>
</tr>
<tr>
<td>Develop and implement a standardised stool testing protocol for all laboratories receiving specimens from residential aged care facilities</td>
<td>HARD</td>
<td>LOW</td>
<td></td>
</tr>
<tr>
<td>Update ASID/ACIPC infection control position statement</td>
<td>EASY</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Disseminate guidance to promote the requirement for single room placement and contact precautions for all patients with diarrhoea until pathology results are known</td>
<td>EASY</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Review the barriers to appropriate CDI testing</td>
<td>EASY</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Develop e-solution to enable easier extraction of surveillance data from electronic data systems</td>
<td>HARD</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Develop data validation algorithm</td>
<td>HARD</td>
<td>LOW</td>
<td></td>
</tr>
<tr>
<td>Implement better CDI surveillance definitions</td>
<td>HARD</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Review utility of other indicators for providing more meaningful outcome data to drive practice change</td>
<td>EASY</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Perform desktop audit of all CDI clinical management and treatment guidelines in Australia and identify inconsistencies</td>
<td>EASY</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Examine utility of hospital administrative data and hospital lab surveillance to develop a community-based surveillance system</td>
<td>EASY</td>
<td>HIGH</td>
<td></td>
</tr>
<tr>
<td>Develop and implement primary care NAPS</td>
<td>HARD</td>
<td>HIGH</td>
<td></td>
</tr>
</tbody>
</table>
### Appendix 6 - Context mapping

<table>
<thead>
<tr>
<th>Priority change ideas</th>
<th>Topic Domain</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>CDI laboratory testing</td>
</tr>
<tr>
<td></td>
<td>Reduce variation in the testing effort</td>
</tr>
<tr>
<td>1. Recommend Aus IP&amp;C guidelines include requirement for contact precautions for all patients with diarrhoea until pathology results are known</td>
<td></td>
</tr>
<tr>
<td>2. Develop educational resources for clinicians and bed managers on bed placement priorities for patients with diarrhoea or known/suspected CDI</td>
<td></td>
</tr>
<tr>
<td>3. Develop national monitoring mechanism for healthcare acquired CDI</td>
<td></td>
</tr>
<tr>
<td>4. Update ASID/ACIPC infection control position statement</td>
<td></td>
</tr>
<tr>
<td>5. Disseminate guidance to promote the requirement for contact precautions, including appropriate bed placement, for all patients with diarrhoea until pathology results are known</td>
<td></td>
</tr>
<tr>
<td>6. Review the barriers to appropriate CDI testing</td>
<td>X</td>
</tr>
<tr>
<td>7. Perform desktop audit of all CDI clinical management and treatment guidelines in Australia and identify inconsistencies</td>
<td></td>
</tr>
<tr>
<td>8. Review utility of other indicators for providing more meaningful outcome data to drive practice change</td>
<td></td>
</tr>
<tr>
<td>9. Review utility of current hospital administrative data, hospital laboratory surveillance and identify and review any other mechanisms for monitoring CDI in the community</td>
<td>X</td>
</tr>
</tbody>
</table>

Improving the prevention and control *Clostridium difficile* infection in Australia