Are current clinical guidelines on the use of Peripheral Intravenous Cannula for blood draws supported by evidence? An organizational case study

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INTRODUCTION

Sampling a person's blood is a common and frequently used clinical diagnostic procedure (Thakker et al., 2015). Blood sampling through direct venepuncture is an invasive procedure that is associated with patient discomfort and decreased satisfaction (Buowari, 2013). It is possible to reduce patient discomfort by drawing blood through an existing peripheral intravenous cannula (PIVC) and replacing PIVCs only when clinically indicated (Rickard et al., 2012). However, there is limited consensus across Australian jurisdictions as to whether (and why) the practice of drawing blood through PIVC should be supported. This paper will look at sources of evidence used to inform the creation of government health policy using the specific example of Australian state government policies on the use of PIVC for blood drawing.

BACKGROUND

In academia, the word evidence is used to refer to information that supports or disproves a hypothesis and is produced and validated through the use of specific protocols and measures (Cairney, 2016). Evidence can be taken from a variety of sources and be presented in a variety of forms, each of which will have an impact on the evidence's...
decisions are rarely made on the basis of academic and scientific evidence to produce recommendations (ICN, 2012). Newman et al. (2017) suggest that policy makers are either ignorant of evidence related to their policy field or clearly referenced or identified. It is commonly alleged that policy formation occurs at the government policy level, as the use of evidence in policies is not always clear in their jurisdictions (Dixit & Sambasivan, 2018).

Clinical healthcare workers often seek to incorporate evidence into healthcare through the use of evidence-based practice (EBP). The term EBP is associated with the work of Archie Cochrane and his belief that clinicians should use only the most effective and proven healthcare procedures available (Mackey & Bassendowski, 2017). In clinical situations, EBP incorporates the best available evidence with the needs of health service users and the clinical skills of healthcare staff (ICN, 2012).

There can be significant delays between the discovery of quality evidence and implementation of that evidence into clinical practice (Melnyk, Lynn, English, & Ellen, 2014). The biggest difficulty with EBP may not be creating the evidence but translating the evidence and ensuring that clinical practice is changed in favour of the evidence (ICN, 2012). Healthcare organizations may lack EBP when they are unaware of evidence-based research or when research is not translated into policy (Stavor, Zedreck-Gonzalez, & Hoffmann, 2017). Implementing EBP is a collective and corporate responsibility, as individual practitioners work within broader environments of teams and organizational requirements (Williams, Perillo, & Brown, 2015). It should also be acknowledged that patient care decisions are rarely made on the basis of academic and scientific evidence alone but are required to incorporate individual patient and staff factors, judgements and values (ICN, 2012).

Having healthcare policies and guidelines that are informed by current best evidence are of importance to ensure high-quality and safe patient care. Yet there are several notable practices that continue in Australian health care in the face of high-quality evidence. Examples include changing peripheral intravenous cannulas routinely rather than when clinically indicated and routine oxygenation of patients (Melnyk, 2017; Rickard et al., 2012).

In Australia, state governments are responsible for the operation of healthcare service delivery and the creation of health policy within their jurisdictions (Dixit & Sambasivan, 2018).

It is often unclear how the concept of EBP is understood at a government policy level, as the use of evidence in policies is not always clearly referenced or identified. It is commonly alleged that policymakers are either ignorant of evidence related to their policy field or choose to ignore and fail to act on known evidence (Cairney, 2016). However, there is a specific and specialized skill set associated with evaluating large volumes of evidence and then using the results to produce recommendations (ICN, 2012). Newman et al. (2017) have argued that structural limitations within the Australian public service may have left governments unprepared to engage with research-based and academic evidence.

Peripheral intravenous cannula (PIVC) is a medical device that can easily be inserted into and remain in a person's peripheral vein to allow continued venous access (Wong, Cooper, Brown, Boyd, & Levinson, 2018). It is estimated that up to 80% of hospitalized patients will require intravenous therapy at some point during their inpatient hospital stay (Yagnik, Graves, & Thong, 2017). Peripheral intravenous cannula is the most commonly inserted vascular access device and can be used for administration of medications, blood sampling and management of conditions (Carr et al., 2016). Health professionals understand the importance of obtaining accurate blood sampling; however, there is considerable variability in the methods used (Bentley, Thakore, Muir, Baird, & Lee, 2016). Traditionally, most blood samples have been drawn from peripheral venepuncture, although increasingly PIVCs are being used for blood sampling (Carr et al., 2016; Davies, Coventry, Jacob, Stoneman, & Jacob, 2019).

While blood can be drawn from PIVC, there are several states and territories in Australia where this is avoided, in part, due to the perceived association between PIVC blood draws and increased risk of haemolysis, sample dilution, bloodstream infection and phlebitis (Mulloy, Lee, Gregas, Hoffman, & Ashley, 2018). Yet the evidence around the use of PIVC for blood sampling is unclear. A systematic review by Jeong et al. (2019) found that sampling from PIVC or venepuncture is likely to provide equivalent levels of accuracy for the most commonly used tests. This is in line with a systematic review by Coventry and colleagues (2019) that found limited evidence to suggest that use of PIVC for blood sampling contributed to decreased accuracy, but that PIVC use may lead to increased haemolysis rates. Relying on policies that are made based on poorly informed evidence is one of the main contributors to healthcare service failures (ICN, 2012). There are variances in practice regarding obtaining blood samples from PIVC post-insertion between individual nurses, health services and states in Australia (Davies et al., 2019). These variations and the evidence used to support them are the focus of this study. Therefore, the aim of this study is to examine what evidence is used to support Australian State government guidelines on peripheral intravenous cannula use?

3 METHODS

This research was conducted as an organizational case study, following the Strengthening the Reporting of Observational Studies in Epidemiology reporting guidelines (See File S1). Ethics approval was not required for this study. The authors did not have any conflict of interest to declare, and there was no funding for this project.

In June 2018, policy guidelines relating to PIVC use were sought from the relevant health departments from all Australian States and Territories. In most cases, the policies were freely available online from the health-department websites. Where policies were not readily available, the health department in question was contacted directly and the policies were provided via email.
Seven Australian states/territories were included in the comparison. The jurisdictions were Australian Capital Territory (ACT), New South Wales (NSW), Northern Territory (NT), Queensland (QLD), Tasmania (Tas), Victoria (Vic) and Western Australia (WA). During data collection, it was found that South Australia (SA) did not have an overarching policy for the use of PIVC, but that policies were set by several local health networks. Due to the inability to access several local health network policies, it was decided to exclude SA from this study.

The policies were assessed according to firstly, how well the policy covered the procedure of PIVC blood draws and secondly, PIVC dwell time. Data were extracted from the documents including the year the policy was published, the title of the document and whether the policy included reference to the practice of sampling of blood from a PIVC. Also extracted were details of whether blood could be drawn from a PIVC on insertion, could blood be routinely drawn from a PIVC, could a PIVC be inserted for the purpose of blood collection, was there a procedure on how to draw blood from a PIVC, were there any exceptions and did the policy outline when a PIVC should be removed or replaced.

The evidence and type of evidence that was used to support these guidelines were also extracted. To determine the type of evidence being used to justify positions taken in the guidelines, the reference lists and bibliographies of the guidelines were analysed. Data were collected relating to evidence type, where the evidence was taken from, who created the evidence and how recently it was published.

Using assessment guidelines set out by the Oxford Centre for Evidence Based Medicine (OCEBM) (2009), the choices of evidence used by the guidelines were assessed for quality. The OCEBM guidelines provide a way to categorize evidence into levels, allowing evidence in each level to be assigned a numerical value and a grade to be provided for a reference list based on the proportion or number of higher scoring items. Items are scored from 1 (Systematic Reviews of Randomized Control Trials) through to 5 (expert opinion), the number of sources from each category is identified, and a grade from A–D is applied. Allocation of an A grade indicates a strong level of evidence, while D grade indicates a weak level of evidence. The policies were also compared, to highlight possible cross-jurisdictional inconsistencies. The grades of evidence were compared with the content of the policies to assess whether there were consistent recommendations found in guidelines that contained high levels of evidence. Two researchers independently analysed the data extracted from the policy documents. Where disagreements occurred, these were brought to the research team and discussed until consensus was reached.

4 | FINDINGS

Data extracted from the policy documents identified large variations between policy documents in the information provided. Table 1 lists the information covered by the policies. Policies were all published between 2012–2017. Six of the documents directly referred to blood sampling from PIVCs, with three documents allowing routine collection of blood through a PIVC and three documents prohibiting collection of blood through a PIVC. Procedures for drawing blood through a PIVC differed, from specifying flushing techniques, stating only large veins to be used and whether a vacutainer could be used for withdrawal. One policy provided conflicting information, suggesting PIVCs could be used for routine blood collection and stating that blood may only be collected from PIVCs if it was specifically inserted for that reason (Tas, 2016). Some policies provided strict rules in relation to the use of PIVC for blood draws and set out specific exceptional circumstances as the only times at which they could be used (WA, 2017; QLD, 2015). The guidelines were similar in their approach to PIVC dwell time. Most Australian jurisdictions require PIVC to be replaced at 72 hr, with only Queensland allowing replacement to occur as clinically indicated.

The type of evidence used by the different documents also varied (Table 2. Type of evidence used). Policies were examined to assess currency and the frequencies at which they were reviewed. Documents had publication dates between 2012–2017, all were current within the date stated for review, although one state did not list a review date. All policy documents had references cited but differed as to whether the references were cited in text, at the end of the chapter only, or in a reference list at the end of the document. The evidence used for policy development was taken from a wide variety of different sources including recent and dated academic journals, legislation/government policies, manufacturer guidelines and grey literature. Some of the state guidelines included citations for the relevant guidelines in other Australian jurisdictions, creating a circle of “self-citing.”

The level of evidence provided in each document was mapped against the OCEBM Levels of Evidence (Table 3). Identified variations in evidence are graded from categories A–D. New South Wales and QLD both score A for their OCEBM evidence use, however, comparing the content of their policies from table 1, show significant discrepancies. These discrepancies are most apparent in the application of evidence to cannula dwell time, with QLD allowing changes to occur when clinically indicated, as opposed to NSW where they are required to be changed every 72 hr (Table 1). It is also noteworthy that the guidelines in Vic do not seem to be based on any form of evidence and received an evidence grade of D.

5 | DISCUSSION

Understanding the way that state governments view the evidence that they use when creating clinical guidelines can provide significant insight for those working in evidence creation. This study has found that many Australian state governments have endorsed policies relating to the use of peripheral intravenous cannula, including blood sampling and dwell time that are lacking, or informed to a large degree by, non-academic evidence. The use of peripheral intravenous cannula and evidence used to support guidelines positions is inconsistent and lacks clarity.
<table>
<thead>
<tr>
<th>State</th>
<th>Year</th>
<th>Document title</th>
<th>Specifically refers to sampling blood from PIVC</th>
<th>Blood may be drawn from PIVC on insertion</th>
<th>Blood may be routinely drawn from PIVC</th>
<th>PIVC may be inserted for purpose of blood collection</th>
<th>Procedure for PIVC blood draw specified</th>
<th>Specified exceptions</th>
<th>PIVC Dwell Time</th>
</tr>
</thead>
<tbody>
<tr>
<td>ACT</td>
<td>2017</td>
<td>Canberra Hospital and Health Services Procedure Peripheral Intravenous Cannula, Adults and Children (Not neonates)</td>
<td>NO</td>
<td>Not stated</td>
<td>Not stated</td>
<td>Not stated</td>
<td>PIVC must be flushed pre- and postblood sampling</td>
<td></td>
<td>Must be changed after 72 hr</td>
</tr>
<tr>
<td>NSW</td>
<td>2013</td>
<td>Peripheral Intravenous Cannula (PIVC) Insertion and Post Insertion: Care in Adult Patients</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Aspirate via capless valve and transfer to vacutainer. Flush PIVC with 10 ml of 0.9% saline post sampling.</td>
<td></td>
<td>Must be changed after 72 hr</td>
</tr>
<tr>
<td>NT</td>
<td>2015</td>
<td>Peripheral Intravascular Catheters (PIVC) Insertion and Management (Adult) NT Health Services Procedure</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Not stated</td>
<td>Blood samples may be drawn from large veins only</td>
<td></td>
<td>Must be changed after 72 hr</td>
</tr>
<tr>
<td>QLD</td>
<td>2015</td>
<td>Peripheral intravenous catheter (PIVC) Guideline</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Not stated</td>
<td>Blood samples may be drawn from relatively large veins, immediately following insertion only</td>
<td>Blood cultures may not be taken from a PIVC</td>
<td>Change only when clinically indicated</td>
</tr>
<tr>
<td>Tas</td>
<td>2016</td>
<td>Peripheral Intravenous Cannula (PIVC) Insertion, care and Maintenance Protocol</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>Not stated</td>
<td>Blood samples may be taken using either syringe or vacutainer</td>
<td>Blood samples may be taken using a vacutainer, blue connection and interlink needle</td>
<td></td>
</tr>
<tr>
<td>Vic</td>
<td>2012</td>
<td>Clinical Skills in Hospitals Project Intravenous (IV) therapy (2012) Module 1. IV Cannulation</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes</td>
<td>Not stated</td>
<td>Bloods may be taken using a vacutainer, blue connection and interlink needle</td>
<td></td>
<td>Must be changed after 72 hr</td>
</tr>
<tr>
<td>WA</td>
<td>2017</td>
<td>Insertion and Management of Peripheral Intravenous Cannulae in Western Australian Healthcare Facilities Policy</td>
<td>Yes</td>
<td>Yes</td>
<td>No</td>
<td>No</td>
<td>Blood may only be drawn from a PIVC in an emergency; when the patient has limited vascular access or is at increased risk of bleeding or receiving thrombolytic therapy</td>
<td></td>
<td>Must be changed after 72 hr</td>
</tr>
</tbody>
</table>

Abbreviations: ACT, Australian Capital Territory; NSW, New South Wales; NT, Northern Territory; QLD, Queensland; Tas, Tasmania; Vic, Victoria; WA, Western Australia.
It is intuitively known that when health policies are informed by robust and tested evidence, those policies are likely to result in more equitable and sustainable gains to population health (van de Goor et al., 2017). Organizations devoted to the promotion of evidence-based health care have emerged in the United States and internationally, seeking to increase the production of systematic reviews and evidence syntheses (Malekinejad, Horvath, Snyder, & Brindis, 2018). The World Health Organization has long acknowledged the importance of evidence in health through an advisory committee dedicated to health research and set up a sub-committee specifically to examine the use of research evidence (van de Goor et al., 2017). This study was a reverse of the traditional method of policy investigation, whereby the researchers looked at the policies to find the evidence supporting the use of a particular clinical practice, as opposed to looking at the evidence to ascertain whether it is followed. Because only two of the seven Australian jurisdictions received an evidence rating of A and those policies differed in their interpretation of the evidence that they did use, it is quite clear that there is more to be done in the promotion of research into the use of PIVC for blood draws.

van de Goor et al. (2017) conducted a study of the reasons why European countries struggled to integrate evidence into their health policies. They found that having a lack of concrete evidence in specific areas, having a lack of existing evidence on the cost of specific procedures or programmes and lacking a joint and shared understanding of the barriers to implementation were significant reasons for reduced policy evidence uptake. More rigorous policy creation processes may be required to allow readers to understand the inconsistencies between policies, and the way evidence is used in policy creation and development (Oxman, Fretheim, Schünemann, & SURE, Fretheim & Schünemann, 2006).

It may seem intuitive to suggest that policymakers should make policy based on evidence; however, a strict application of evidence would remove the mandate of decision-making from those who were elected as the policymaker and decision-maker (Cairney, 2016). Healthcare decision-makers are often interested in using evidence to support their decision-making but may find that the scientific evidence available is inadequate for their needs (Tunis, Stryer, & Snyder, 2018). The World Health Organization has emphasized the need for policies to find the evidence supporting the use of a particular method of policy investigation, whereby the researchers looked at the evidence that was specifically cited in their interpretation of the evidence that they did use, it is quite clear that there is more to be done in the promotion of research into the use of PIVC for blood draws.

5.1 Limitations

This paper looked at the evidence that was specifically cited in policies. There may have been additional evidence that was used to support the creation of policies, particularly expert opinions that may not have been referenced in the policies and thus could not be included.

5.2 Recommendations

As current guidelines do not appear to use high levels of evidence, policymakers should give weight to the importance of academic...
rigour in the creation of clinical policies and guidelines. State governments should ensure that searches are conducted for academic evidence in support (or opposition) of specific practices prior to guideline publication. Policymakers may wish to consider providing grades of evidence to support specific policies, allowing for more open discussion over where new evidence is required.

Where there is clear evidence for a clinical policy, such as dwell time for PIVCs, decision-makers appear to be ignoring the evidence. State governments should consider employing people with academic skills and clinical experience in related professions in advisory roles prior to publication of health policy documents to review the evidence.
Involving clinicians and clinically trained academics in policy creation could assist with reducing the divide between policy and practice, as well as providing a conduit for communicating guidelines to clinical professionals. Future research could consider the capacity of clinical staff to engage with varying levels of evidence and the upskilling required to assist clinicians to think critically about knowledge translation.

### 6 | CONCLUSION

The use of high-quality evidence in the creation of health policy is important to ensure that the policy results in positive health outcomes. Health policy guidelines for the use of PIVC in blood sampling in Australia are inconsistent and inconsistently supported by evidence. While it may not always be feasible for decision-makers...
to base policies on evidence, they should be at the least informed by evidence. Barriers to evidence-based policy can include lack of available or consistent evidence and lack the skills of skills by decision-makers to understand academic level clinical information. Governments should be at the forefront for using research-based evidence where available instead of informing themselves through the expert opinions and individual contacts.

CONFLICT OF INTEREST
The authors declare that they have no conflicts of interest associated with this study.

AUTHOR CONTRIBUTIONS
AJ and EJ: Study design. AJ: Data extraction. EJ, AJ, HD and LC: Methodology. AJ and LC: Data analysis. AJ, HD, LC and EJ all contributed towards editing and all authors approved the final manuscript.

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REFERENCES


**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section.

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