Sherlock 3CG® Tip Confirmation System for Placement of Peripherally Inserted Central Catheters: A NICE Medical Technology Guidance

Megan Dale1 · Ailish Higgins2 · Grace Carolan-Rees1

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Abstract In current clinical practice, peripherally inserted central catheters (PICCs) are typically inserted using external anatomical measurements and a confirmatory chest X-ray, or using fluoroscopy. The Sherlock 3CG® Tip Confirmation System (TCS) allows magnetic tracking of the PICC tip during insertion and confirmation of the final location using ECG, meaning that most patients will not require a chest X-ray or fluoroscopy. The Sherlock 3CG® TCS was evaluated in 2014 by the UK National Institute for Health and Care Excellence (NICE) as part of the Medical Technologies Evaluation Programme. The company (C.R. Bard Ltd) identified four abstracts, one paper pending publication and questionnaire data from NHS users of the Sherlock 3CG® TCS. None of the evidence included a comparator arm. Placement accuracy of PICCs using the Sherlock 3CG® TCS where a chest X-ray was also used ranged from 79.5 to 100 %. The company reported that 9 out of 16 NHS centres that used the Sherlock 3CG® TCS were no longer using chest X-rays to routinely confirm PICC tip location. The evidence did not report the need for catheter repositioning, re-insertion, staff time savings, treatment delays, length of stay, quality of life outcomes or complications. The company’s model found that the Sherlock 3CG® TCS was cost saving by GBP9.37 per patient compared to blind bedside PICC insertion. The use of the Sherlock 3CG® TCS in the UK NHS compared to blind PICC insertion using a confirmatory chest X-ray appears to hover around being cost neutral. Staff time and accuracy were key drivers in the model: evidence for these is sparse and the reality will vary in different situations. If evidence became available for outcomes after the initial insertion, such as replacement, complications and adverse events, the cost implications may change. The direction of this potential change is not known. NICE published guidance MTG24 in March 2015 recommending that the case for adoption of Sherlock 3CG® TCS was supported by the evidence.

Key Points for Decision Makers

The Sherlock 3CG® TCS has already been implemented in 16 NHS sites, and routine use of chest X-rays for confirming PICC tip location has been eliminated in some of these.

Use of the Sherlock 3CG® TCS appears to be approximately cost neutral compared to blind PICC insertion, based on the available low quality evidence. It may be cost saving if there are reductions in nurse time, X-ray provision, portering or the number of reinsertions required.

There is some evidence that the Sherlock 3CG® TCS can improve tip positioning accuracy compared to blind placement of PICCs.

There is no evidence on the effect of Sherlock 3CG® TCS on the time to treatment, length of stay, clinical outcomes or patient experience.

1 Cedar, Cardiff and Vale University Health Board, Cardiff, Wales, UK
2 National Institute for Health and Care Excellence, Manchester, UK

Megan Dale megan.dale@wales.nhs.uk

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1 Introduction

This is part of a series summarising guidance produced by the National Institute for Health and Care Excellence (NICE) Medical Technologies Evaluation Programme (MTEP). The process is explained by Campbell and Campbell in the first publication of this series [1]. This article summarises the External Assessment Centre (EAC) report [2] and Medical Technology Guidance (MTG) for the Sherlock 3CG® Tip Confirmation System (TCS) for placement of peripherally inserted central catheters [3]. The EAC that produced the assessment report for the Sherlock 3CG® TCS was Cedar, a collaboration between Cardiff and Vale University Health Board, Cardiff University and Swansea University.

2 Background

Peripherally inserted central catheters (PICCs) use a catheter inserted at a peripheral vein using ultrasound guidance and gently moved through the vein until the tip is in the superior vena cava or right atrium. The insertion site is typically the brachial or cephalic vein in the upper arm, leading to the right subclavian vein [4, 5] (Fig. 1).

PICCs are used in different circumstances, including:

- measuring circulatory or heart functions
- providing long-term access route for infusions and blood tests
- delivering drugs that require rapid dilution, e.g. chemotherapy
- delivering contrast medium for cardiac imaging.

The different clinical uses for PICCs mean that there is variation in the pathways that occur in practice. PICCs may be inserted in intensive care units (ICUs), in outpatient clinics, at the bedside, and in fluoroscopy and X-ray departments. The clinical staff who insert PICCs include specialist nurses, radiologists and doctors [2].

The two methods for placing PICCs which are currently standard are [2]:

- Blind placement—the catheter is pushed into the vein a set distance according to anatomical measurements taken externally on the patient. Once the PICC is placed, the correct position is confirmed using a chest X-ray.
- Fluoroscopy-guided PICC placement—the tip location can be visualised and the final location confirmed during the procedure.

In the UK, the preferred position for the PICC tip position is normally the mid or lower superior vena cava, cavo-atrial junction, or the high right atrium, based on European guidelines [4]. There is no international consensus.

The length of time for which PICCs are in place ranges from a few days to over a year: 1 week to 3 months is most typical. During this time, the location of the tip may move slightly. This would be identified only if it caused a clinical problem, or if there was an X-ray or imaging for another reason.

There are instances of gross malpositioning, for example if the catheter enters the left subclavian vein or the right internal jugular vein. It may be possible to correct such malpositioning with the catheter in place. Otherwise, the PICC is removed and a new PICC inserted. PICC malpositioning may cause complications such as catheter malfunction, cardiac arrhythmia or tamponade [5]. More minor malpositioning has less obvious clinical relevance. The confirmatory chest X-ray may show that the tip is inserted slightly too far, and in this case it can be pulled back to achieve the desired position. Some PICC teams will intend to always place the tip too far, and pull back by a measured amount following the chest X-ray. The PICC can never be moved further in after initial insertion, as the external section is no longer sterile.

2.1 Sherlock 3CG® Tip Confirmation System

The Sherlock 3CG® TCS uses a sensor to track a magnetic tip at the distal end of the catheter and internal ECG measurements to confirm the position of the catheter tip relative to the heart. For most patients, a chest X-ray will not be required to confirm tip placement, and the procedure can be completed at the patient’s bedside, or in an outpatient clinic.

The Sherlock 3CG® TCS sensor is a class I medical device, which obtained a CE mark in December 2011. The sensor must be used with the Sherlock 3CG® magnetic stylet, and Power PICC SOLO catheter—class II medical devices, which obtained CE marking in February 2012. The Sherlock 3CG® TCS cannot be used with other types of catheter.

2.2 National Institute for Health and Care Excellence (NICE) scope

NICE defines the scope of the evaluation prior to the company’s submission of evidence [6].

Population Adult patients undergoing PICC insertion.

Intervention The Sherlock 3CG® TCS, used by a healthcare professional trained in PICC placement. Previous versions of the system that did not include the use of both magnetic tracking and ECG tip confirmation were excluded.
Comparators There were two comparators, based on current practice:
• Blind PICC insertion followed by chest X-ray.
• Fluoroscopy.

Outcomes Included accuracy of catheter tip placement, requirement for confirmatory chest X-ray, catheter malposition rates, catheter re-positioning rates, impact of malposition-related complications, time to treatment, staff time, length of stay, fluoroscopy needed to place the PICC, time for PICC insertion, patient experience, patient quality of life and device-related adverse events.

The benefits to patients claimed by the company are [2, 6, 7]:
• Better accuracy of PICC placement.
• Better outcomes by reducing the incidence of catheter malposition and post-procedural repositioning.
• Removed the need for a chest X-ray or fluoroscopy to confirm tip location after PICC insertion.
• Reduced treatment delays due to intra-procedural verification of tip position.
• Safe method for PICC tip placement with no associated adverse events or complications.
• PICC placement and tip verification are during the same procedure.
• It improves patient experience and increases the patient’s confidence in the PICC placer.

The benefits to the healthcare system claimed by the company are:
• A reduced and more efficient care pathway because no chest X-ray is needed.
• Lower staff requirements (radiologists/radiology nurses/radiographers/radiology healthcare support workers) because the need for X-ray confirmation is reduced. Also reduced need for porters for patient transfer and doctors for X-ray assessment.
• Potential reduction of bed occupancy due to reductions in treatment delays.
• Reduced costs of consequences of malpositioning.
• Reduced costs of using resource intensive departments such as radiology.

3 Review of Clinical and Economic Evidence

The company provided an evidence submission to NICE that summarised the clinical and cost evidence for the Sherlock 3CG\textsuperscript{C}® TCS and presented a cost-consequence model [8]. The aim is to evaluate whether the Sherlock
3CG® TCS carries a clinical advantage and/or a reduced cost in comparison to current NHS standard care.

3.1 Company’s submission of clinical effectiveness evidence

The company identified and submitted 4 relevant studies [9–12]. None of the submitted studies were peer reviewed or published in full. Two studies were from the USA, one from Australia and one from the UK. The company also identified a study that was pending publication [13], and subsequently published. Given the paucity of evidence, the EAC widened the scope to include any previous models of the technology that included both ECG measurement and magnetic tracking. This resulted in one presentation being included [14]. The company also submitted information from questionnaires that it had sent to selected NHS users of the Sherlock 3CG® TCS.

3.2 Critique of clinical evidence available

3.2.1 Peer reviewed evidence

The only available peer reviewed paper reported a retrospective analysis of the first 250 PICC insertions following introduction of the Sherlock 3CG® TCS to an NHS intensive care unit (ICU) [13]. PICCs were placed by the Vascular Access Team at the bedside and, following tip confirmation with the Sherlock 3CG® TCS, a portable X-ray was used to assess placement accuracy. Eleven out of 250 patients (4%) were excluded due to difficulties interpreting the ECG (n = 4), other PICC difficulties (n = 3), or X-ray difficulties (n = 4). The authors reported 49 out of 239 (21%) tip placements were not in the correct position using European guidelines (Pittiruti guideline).

A paper by the same group reported accuracy of blind bedside PICC placement shortly before the introduction of the Sherlock 3CG® TCS [15]. The EAC has presented the results of both papers in Table 1, although the studies do not directly compare identical populations. The malposition rate using the Sherlock 3CG® TCS is significantly lower than the blind placement (p < 0.0001) and the authors conclude that if this definition of tip placement is acceptable, then the Sherlock 3CG® TCS can be used for tip confirmation without routine chest X-ray confirmation.

3.2.2 Other Evidence

Results from non-peer-reviewed evidence such as abstracts, posters and presentations are summarised in Table 2.

The accuracy of the Sherlock 3CG® TCS in Johnston et al. [13] (Table 1) is lower than that reported in five other studies (Table 2); possibly because ICU patients have less clear heart rhythm or are recumbent. It is unclear if the results from this one study are generalisable to other settings (either to other ICU settings or the wider NHS).

Questionnaires completed by six NHS sites using the Sherlock 3CG® TCS highlighted the variation in normal clinical pathways for tip insertion and confirmation, repositioning and reinsertion. At the time of submission, the company reported that nine NHS sites had stopped using routine chest X-rays for tip confirmation, from a total of 16 NHS sites that have introduced the Sherlock 3CG® TCS.

3.2.3 Summary

All studies used the Sherlock 3CG® TCS as the intervention; none were comparative. The studies reported the use of the Sherlock 3CG® TCS for tip confirmation, followed by a chest X-ray to determine if the tip was in the correct place. There were no comparators giving information on the blind PICC placement malposition rates in the same setting. The studies all stop at the point of tip confirmation and assessment by chest X-ray. It is not known if PICCs identified as malpositioned had to be adjusted or reinserted, or what the consequences might be. Similarly, the studies do not give any indication if any patients would have been identified as having misplaced PICCs without the chest X-ray, and what the clinical implications might have been.

Several of the sites from the published abstracts [9–12, 14] report that routine chest X-rays are no longer used to confirm PICC placement. The sponsors report that nine NHS sites using Sherlock 3CG® TCS have had sufficient confidence in the device to remove the requirement for routine chest X-rays to confirm tip location.

3.3 Cost Evidence

Two studies included cost information [9, 10], but contained insufficient information for critical analysis, and had limited relevance to the decision problem. The EAC did not find any additional studies that should have been included. Only two of the six clinical studies identified report the time taken to insert the PICC and complete tip confirmation using Sherlock 3CG® TCS (without chest X-ray), and compare it to the time taken by the previous method. There was no evidence for the effect on staff time, time to deliver treatment, bed days, clinical outcome or patient experience.

The company created an economic model with four branches:

1. Bedside PICC placement with Sherlock 3CG® TCS with X-ray.
2. Bedside PICC placement with Sherlock 3CG® TCS without X-ray.
3. Blind bedside PICC placement, with X-ray.
4. Fluoroscopy guided placement.

Inputs to the model were based on the studies and NHS questionnaires submitted as clinical evidence. Staff costs were derived from the Personal Social Services Research Unit (PSSRU) [16], and detailed pathway information and costs for blind bedside PICC placement and fluoroscopy were obtained from Walker et al. [17, 18].

Using this cost model, the company found that the Sherlock 3CG® TCS without X-ray gave a cost saving of GBP25.67 per patient compared to blind bedside PICC insertion, and a cost saving of GBP510.03 per patient compared to fluoroscopy. All costs are based on 2014 values.

### 3.4 Critique of Economic Submission

There are uncertainties in the model structure and inputs due to the lack of data available and the variations in patient groups and service provision. The company carried out extensive sensitivity analysis, although the EAC disagreed with some of the key inputs. Important assumptions and data for the model submitted by the company, along with the views of the EAC, are listed below.

#### 3.4.1 Time Horizon

The model stops at the point of confirming PICC tip placement. The EAC considered that a preferable time horizon would include adverse events, complications, the time to starting treatment and the clinical consequences. Unfortunately, there is no evidence available to inform such a model and the impact and direction of any effect is unknown.

#### 3.4.2 Nurse Time

The difference in staff time is one of the model’s key drivers. The two studies used in the company’s model were from different countries with different health systems, causing the EAC to consider comparability unlikely. The values used by the company were:

- 62.49 min for blind PICC insertion with X-ray and the Sherlock 3CG® TCS with X-ray, based on Walker and Todd [17] bedside placement time.
- 39.5 min for the Sherlock 3CG® TCS without X-ray [9].

#### 3.4.3 Reinsertion Rates and Method

The difference in reinsertion rates and cost of reinsertion is a key driver when comparing blind bedside PICC insertion to the Sherlock 3CG® TCS. The proportion of accurate placement the model uses is:

- 96 % for the Sherlock 3CG® TCS [10] for trained staff.
- 93.1 % for blind bedside PICC placement [17].
- 100 % for fluoroscopy [17].

The base case of the cost model assumes that all identified tip malpositions will require replacement, and that all reinsertions would be performed using fluoroscopy. Not all malpositions will require reinsertion procedures, and although some services may reinsert PICCs using fluoroscopy, others will use bedside techniques. Because the model includes more malpositions for blind PICC placement than for the Sherlock 3CG® TCS, and fluoroscopy is the most costly option, these assumptions favour the Sherlock 3CG® TCS.

#### 3.4.4 Patient Population

The scope defined by NICE included all adult patients undergoing PICC insertion. The population included in the company’s model was adult patients undergoing PICC insertion who were suitable for ECG tip confirmation using the Sherlock 3CG® TCS. The company assumed 83.5 % of patients were suitable [9], and did not include costs for treatment of the remaining 16.5 % by an alternative method.

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Table 1  Number of tips correctly placed using the Sherlock 3CG® TCS and prior to its introduction [13, 15]

| Definition of adequate tip position | Sherlock 3CG® TCS (Johnston 2014 [13]) | Blind bedside PICC placement prior to the Sherlock 3CG® TCS (Johnston 2013 [15]) |
|-----------------------------------|----------------------------------------|----------------------------------------------------------------------------------|
| ICU (n = 239) % and 95% CI | Mid and low SVC/CAJ/high RA (upper 2 cm) 190 79.5 (74–84) | ICU (n = 246) % and 95% CI | Non-ICU (n = 233) % |
| CAJ cavo-atrial junction, CI confidence interval, ICU intensive care unit, PICC peripherally inserted central catheter, RA right atrium, SVC superior vena cava | | |

CAJ cavo-atrial junction, CI confidence interval, ICU intensive care unit, PICC peripherally inserted central catheter, RA right atrium, SVC superior vena cava
| References | Study design and details | Population | Intervention and previous technology | Accuracy of Sherlock 3CG® TCS (ECG tip placement in agreement with subsequent X-ray\(^a\)) | Comments by the EAC |
|------------|--------------------------|------------|--------------------------------------|----------------------------------------------------------------------------------|---------------------|
| Johnston (peer reviewed paper) [13] | Consecutive case series (UK) | ICU patients needing PICC insertion, excluding those not in sinus rhythm \((n = 250) 11/250 (4.4 \%)\) excluded | Sherlock 3CG® TCS. Previously used chest X-ray confirmation | 79.5 \% of PICCs in acceptable position (mid and lower SVC/CAJ/high RA) | Peer-reviewed publication. Aims, methods, patient population, results and exclusion criteria are fully reported. Chest X-rays reviewed by at least 2 authors |
| Barton [12] | Case series (UK), abstract and additional information from author | Adults with no AF needing PICC insertion \((n = 65\) in original study, \(n = 225\) total, including follow up) \(11/225 (4.9 \%)\) excluded | Sherlock 3CG® TCS. Previously used chest X-ray confirmation | 100 \% of PICCs in acceptable position (lower 1/3 of SVC or CAJ) | Full critical appraisal not possible, not peer reviewed |
| Adams [9] | Case series (USA), abstract and poster | People needing PICC insertion excluding those without identifiable p-wave \((n = 333) 55/333 (16.5 \%)\) excluded | Sherlock 3CG® TCS. Previously used chest X-ray confirmation | 96.4 \% on 1st review 98.2 \% on 2nd review | Full critical appraisal not possible, not peer reviewed |
| Stewart [11] | Case series (Australia), abstract and poster | Unknown, abstract states “over 65” patients treated | Sherlock 3CG® TCS. Previously used chest X-ray confirmation | 96 \% within CAJ 4 \% within RA Number unable to confirm tip location using Sherlock 3CG® TCS not reported | Minimal information available, full critical appraisal not possible, not peer reviewed. Information should be treated cautiously |
| Parikh [10] | Prospective case series (USA), abstract and poster. Two phases, with increased nurse training in 2nd phase. Additional patients after trial also reported | People needing PICC insertion excluding those with AF, atrial flutter, increased risk of bleeding and without identifiable p-wave 1st phase \(n = 65\), 3/65 (4.6 \%) excluded, 2nd phase, \(n = 182\) 35/182 (19.2 \%) excluded | Sherlock 3CG® TCS. Previously used Sherlock II TLS (magnetic tracking) plus chest X-ray confirmation | Phase 1: 83 \% placed within SVC or CAJ Phase 2: 96 \% placed within SVC or CAJ | Poster, not peer reviewed, but with sufficient detail for partial critical appraisal. Clear patient population, tip position criteria, exclusion criteria, assessment by 2 independent operators For follow up phase patients, 437/567 (77 \%) used Sherlock 3CG® TCS, 109/437 (25 \%) still required a chest X-ray |
| Symington [14] | Prospective case series (USA). Presentation | People needing a PICC insertion \((n = 63)\) | Sherlock 3CG® TCS. Previously used Sherlock II TLS (magnetic tracking) plus chest x = ray confirmation | 98.4 \% placed in acceptable position | Minimal information available, full critical appraisal not possible, not peer reviewed |

\(^a\) Accuracy outcomes only include patients where a successful ECG tip placement was thought to have been achieved. If ECG tip confirmation was not achieved, then these patients would be in the excluded category.
method. This has a low impact on the model where the two comparative arms have similar costs. Expert advice indicated that the proportion of patients for whom the Sherlock 3CG® TCS is suitable was likely to be higher than 83.5%. The company’s instructions for use state that use of the technology is limited (but not contraindicated) in patients who do not have an identifiable P wave, e.g. those with atrial fibrillation, tachycardia, etc.

3.4.5 Fluoroscopy

The company’s model finds a large cost saving using the Sherlock 3CG® TCS at the bedside compared to fluoroscopy. This is partially due to the high cost attributed to fluoroscopy—much of the remaining cost saving is due to the move to bedside placement and would be equally true of blind PICC bedside placement.

The cost of theatre for fluoroscopy was assumed by the company to be GBP507.18 [17, 18]. The EAC estimated this cost to be GBP101 [19].

If standard practice was fluoroscopy, and a new bedside PICC placement service had to be set up, there would be costs for nurse training and reorganisation. The model does not include these initial set up costs.

3.5 EAC Alternative Scenarios

There are unavoidable uncertainties due to the lack of evidence, and the variations in patient groups and service provision. There are many inputs and scenarios that can change the model from being cost saving to cost incurring—the most accurate representation will depend on the service being examined. The EAC analysis presents an alternative set of assumptions—there is insufficient evidence to allow absolute certainty over which is most appropriate.

3.5.1 EAC Changes to Company’s Model

The EAC changes to the company’s base model were:

- Nurse time is equal for blind PICC insertion with X-ray, the Sherlock 3CG® TCS with X-ray and the Sherlock 3CG® TCS without X-ray. The EAC used a time of 62.49 min [17]. X-rays are costed separately.
- Where the Sherlock 3CG® TCS is used without X-ray, there is an assumption of zero malpositions being identified within the time frame of the model, since no other tip confirmation system is used.
- Replacements for PICCs placed at the bedside are by the same method as the original placement.
- In patients for whom use of the Sherlock 3CG® TCS is not suitable, treatment costs by an alternative method were included.
- The cost of theatre use for fluoroscopy was changed to GBP101 [19].

Using these inputs, the EAC found that the Sherlock 3CG® TCS without X-ray incurred a cost of GBP9.37 per patient compared to blind bedside PICC insertion, and gave a cost saving of GBP106.12 per patient compared to fluoroscopy. All costs are based on 2014 values.

3.5.2 EAC Cost Neutral Scenario

The EAC explored a scenario where the use of the Sherlock 3CG® TCS is approximately cost neutral when the following assumptions are made (in addition to the previous EAC changes):

- Nurse time for the Sherlock 3CG® TCS is 57.5 min, a 5-min reduction from the original EAC assumption, as there is no requirement for the nurse to interpret the X-ray image.
- The cost of X-ray is assumed to be 10 min of radiologist time (GBP5.67, company’s submission, based on Walker and Todd [17]) with the addition of 15 min for a Band 2 porter (GBP5.25 [16]) to transport patient to and from X-ray. This gives total X-ray cost of GBP10.92.

Using these inputs, the EAC found that the Sherlock 3CG® TCS without X-ray was associated with a cost saving of GBP1.17 per patient compared to blind bedside PICC insertion, and a cost saving of GBP111.27 per patient compared to fluoroscopy.

3.5.3 EAC Scenario Based on ICU Data

The EAC also explored a scenario based on ICU data [13, 15], changing the accuracy rates to those in Table 1. Sherlock 3CG® TCS with X-ray was used in the model, reflecting the local practice at the time of publication [13]. The limitations of this scenario are:

- the data is taken from a single centre and may not be generalisable.
- the comparison is with historical data from the same centre.
- the study only includes intermediate outcomes—the actual number of replacements is not reported.

In this scenario, the Sherlock 3CG® TCS with X-ray confirmation is associated with a cost saving of GBP41.35 per patient compared to blind bedside PICC insertion, because of the significant reduction in the proportion of malpositioned catheters that needed to be re-positioned.
3.6 Impact of Changes

Given current information, use of the Sherlock 3CG® TCS compared to blind PICC insertion using a chest X-ray appears to hover around cost neutral. Staff time and accuracy are key drivers in the model, but evidence for these is sparse and the reality will vary in different situations. If evidence became available for outcomes after the initial insertion, such as replacement, complications and adverse events, the cost implications may change. The direction of this potential change is not known.

4 NICE Guidance

4.1 Preliminary Guidance

The NICE Medical Technologies Advisory Committee (MTAC) met in October 2014 and considered evidence from a range of sources, including the company’s submission, the EAC report and additional economic modelling, and testimony from clinical experts. The committee made provisional recommendations that went to public consultation.

The Committee considered the overall quality and quantity of the evidence to be low, but noted that the general trend of the clinical evidence was in favour of the Sherlock 3CG® TCS, and that the technology seemed likely to be cost neutral. Some additional analysis was carried out by the EAC to further test the impact of staff time on the cost model.

4.2 Consultation Response

During the consultation period, NICE received 14 consultation comments from 7 consultees (3 NHS professionals, 2 manufacturers, the Department of Health, and one EAC representative). The majority of these comments were process-related, and considered if the single-technology assessment methods used by MTEP were appropriate for medical technologies evaluation. At the final guidance meeting (January 2015), the Committee considered these comments with reference to the MTEP’s process and methods guides [20, 21] and concluded that they should not impact on the provisional recommendations. Accordingly, the recommendations did not change substantively as a result of public consultation.

4.3 Final Recommendations

The NICE Medical Technology Guidance on the Sherlock 3CG® TCS for placement of peripherally inserted central catheters was published on 25 March 2015 as MTG 24 [3]. It contains the following recommendations:

1. The case for adopting the Sherlock 3CG® TCS for placement of peripherally inserted central catheters is supported by the evidence. The technology usually avoids the need for a confirmatory chest X-ray in patients who would otherwise have blind insertion, minimising the delay before the catheter can be used for infusion. Using the technology increases staff confidence during catheter insertion.

2. The Sherlock 3CG® TCS should be considered as an option for placement of peripherally inserted central catheters in adults. For patients whose electrocardiogram does not show a P wave (for example, patients with atrial fibrillation), a chest X-ray will still be needed to confirm tip location of the peripherally inserted central catheter.

3. The cost of using the Sherlock 3CG® TCS is similar to that of blind insertion and subsequent chest X-ray in adults who need a peripherally inserted central catheter in a non-intensive care setting. When the Sherlock 3CG® TCS is used instead of fluoroscopy, the estimated cost saving is GBP106 per patient. In an intensive care setting, where the rate of misplacement with blind insertion is generally higher, there is an estimated cost saving of GBP41 per patient per use of the Sherlock 3CG® TCS and a confirmatory chest X-ray compared with using blind insertion and chest X-ray. All these cost savings are subject to some uncertainty and need to be considered in the context of the clinical benefits.

5 Key Challenges and Learning Points

The wide range of uses for PICCs meant that it was difficult to identify a typical patient pathway. Clinical expertise was the key to understanding different pathways and their implications.

The most widely reported outcome was accuracy of the PICC placement judged against chest X-ray. This is an intermediate outcome, rather than the actual clinical or economic information important to a decision.

Sensitivity analysis and alternative scenarios helped to understand the range of possible outcomes within the structure of the model, and to reflect the diverse clinical realities. Where there is no clinical evidence for the actual outcomes needed, there can only be limited confidence in the economic model.

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**Author contributions** MD evaluated the clinical evidence, and both MD and GCR evaluated the economic evidence on which this manuscript is based, as well as the EAC scenarios. MD and GCR wrote the assessment report and AH wrote the assessment report overview both of which were presented to the Medical Technologies Advisory Committee. AH subsequently helped to develop the medical technologies guidance for NICE. GCR reviewed the full EAC report and this article, and can act as a guarantor for the overall content.

**Compliance with Ethical Standards**

Cedar is funded by the NICE Medical Technologies Evaluation Programme to act as an EAC. MD and GCR are NHS employees, and the NHS has a financial interest in the guidance on which this project is based. AH works for the National Institute for Health and Care Excellence, and has no conflicts of interests to declare. This article has not been externally peer reviewed by Applied Health Economics and Health Policy.

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