# Central Venous Access Device Policy

## Key points
- Types of CVADS
- How to refer for a CVAD
- How to manage CVAD associated complications

## Table of Key Features

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1. ASSOCIATED DOCUMENTS

- Standard operating procedure for referral for CVC insertion (Appendix 17)
- PICC Occlusion Algorithm (Appendix 4)
- Tunnelled CVC Occlusion Algorithm (Appendix 5)
- CVAD Occlusion Algorithm using 3-Way tap (Appendix 9)
- Management of CVC related problems in the community (Appendix 13)
- Urokinase administration policy (Appendix 6)
- CVAD request form (Appendix 18)
- MRSA screening policy
- Anticoagulation policy therapy and management
- Policy for consent to examination and treatment
- Care of your central venous catheter (patient information for tunnelled CV)
- Care of your peripherally inserted central venous catheter (patient information)

2. INTRODUCTION

This policy is to provide information on central venous access devices that are used at The Christie, to ensure patients receive the most appropriate device, the devices are managed correctly and any complications related to them are minimised as much as possible. A central venous access device is defined by the internal portion of the catheter being placed in the central veins, ideally the tip location should be at the lower one-third of the Superior Vena Cava (SVC) at the level of the cavoatrial junction (CAJ) (INS 2016). CVADs provide access to the venous bloodstream of the body and are used for blood sampling, administration of medications, fluids, blood products and pressure monitoring.

2.1 Statement of intent

The intention of this policy is to set out the standards expected by the Christie NHS Foundation Trust, to ensure safe, effective practice in relation to the insertion and removal of CVADs.

2.2 Equality Impact Analysis

As part of its development, this policy was analysed to consider its effect on different groups protected from discrimination by the Equality Act 2010. The requirement is to consider if there are any unintended consequences for some groups, and to consider if the policy will be fully effective for all protected groups. This analysis has been undertaken and recorded using the trust’s e-tool, and appropriate measures taken to remove barriers or advance equality in the delivery of this policy.

2.3 Good Corporate Citizen

As part of its development, this policy was reviewed in line with the Trust’s Corporate Citizen ideals. As a result, the document is designed to be used electronically in order to reduce any associated printing costs.

2.4 The Christie Commitment

We aim to reward our staff who are committed and motivated to do their best for patients every day. The trust’s principles and behaviours describe what our patients and their families or carers can expect from us, and what our staff can expect from each other.

The trust’s behaviours are:

- We always give the best quality care

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Quality & Standards Team Template 2014/15
We treat everybody with compassion, dignity and respect
We listen to our patients and each other
We work together as one Christie team
We share knowledge and learning
We support staff to develop to their full potential
We look for new ideas and better ways of working
We promote a fair culture
We provide a safe, clean and tidy environment

All staff are expected to behave in a way that reflects the trusts principles and behaviours.

2.5 Purpose
This policy will provide staff information and guidance around CVADs (Central venous access devices) used with the Trust.

2.6 Scope
This policy is for all staff involved in any aspect of insertion, care/maintenance and removal of CVADS at The Christie NHS Foundation Trust
- Non-tunnelled (percutaneous) acute CVC
- PICC - Peripherally inserted central venous catheter
- Skin tunnelled central venous catheter (sometimes known as a Hickman line™)
- TIVAD - Totally implantable vascular access device (sometimes known as a Portacath™).

3. DEFINITIONS

<table>
<thead>
<tr>
<th>Term</th>
<th>Meaning</th>
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<tr>
<td>CAJ</td>
<td>Cavoatrial junction; Acceptable tip location for CVAD</td>
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<tr>
<td>CVAD</td>
<td>Central venous access device</td>
</tr>
<tr>
<td>CVC</td>
<td>Central venous catheter</td>
</tr>
<tr>
<td>ECG</td>
<td>Electrocardiograph; ECG based systems are used to confirm catheter tip position during insertion</td>
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<tr>
<td>IV</td>
<td>Intravenous</td>
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<tr>
<td>Normal working hours</td>
<td>Monday to Friday 9am-5pm</td>
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<tr>
<td>Procedure team</td>
<td>Nurse lead team responsible for vascular access and other day case procedures</td>
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<tr>
<td>SVC</td>
<td>Superior Vena Cava; Acceptable tip location for CVAD</td>
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<td>SVCO</td>
<td>Superior Vena Cava Obstruction; A medical condition that contraindicates placement of CVAD in the upper body, femoral approach maybe appropriate</td>
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<td>VHP</td>
<td>Vessel Health and Preservation; evidenced based strategy for vascular access</td>
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4. DUTIES

4.1 Patient Safety Committee
Responsible for:
- Providing the Risk and Quality Governance Committee with assurance that comprehensive approach to improving patient safety is in place.
- Requesting in depth investigation of any clinical concerns.

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• Escalate any procedure related issues to Risk and Quality Governance Committee.
• Medical Devices and Safety Committee must approve any new vascular access devices or medical devices related to vascular access introduced to the Trust.

4.2 Chief Executive
The chief executive has overall responsibility for ensuring that the organisation adheres to the standards set out in this policy. This duty may be delegated to an executive/senior manager but accountability to the Board remains with the chief executive.

4.3 Procedure team
Responsible for setting the standards in relation to best practice to CVAD insertion, removal and the provision of a clinical service.

4.4 Clinical skills training team
Responsible for the dissemination of training for the care and maintenance of CVAD’s in accordance with this policy.

4.5 Matrons and Ward Managers
Ensure that staff are adequately trained in each clinical area to provide routine maintenance, care and trouble-shooting of CVADs during normal working hours, evening and weekends where applicable.

4.7 Chemotherapy Team
To ensure all relevant clinical staff adhere to the standards set out in the central venous access device policy.

4.8 Lead Nurse
Responsible for ensuring that clinical staff in their responsible clinical areas are appropriately trained and maintain standards in accordance with this policy

5. CVADs
CVADs are inserted for a variety of reasons including:
• Access for blood sampling
• Administration of fluids, drugs, blood and blood products
• Parenteral nutrition (PN)
• Safe administration of vesicant drugs
• Ambulatory chemotherapy
• Monitoring of central venous pressure
• Patients with poor peripheral venous access or needle phobia
• Photopheresis.

5.1 Device Selection – Vessel Health Preservation
Site selection for vascular access must include assessment of the patient’s condition, age and diagnosis, vascular condition, infusion device history, and the type and duration of the therapy as well as the potential complications associated with vascular access devices (Wise et al., 2001; Dougherty, 2006; Gabriel, 2008; Scales, 2008a).

Factors in patient assessment:
• Age
- Diagnosis
- Patient’s medical condition
- Medical history
- Vascular access history
- Plan of treatments i.e. intensity and duration
- Type of Central Venous Access Device required
- Minimum number of lumens required
- Patient preference/social factors.

The Vessel Health and Preservation (VHP) protocol, which is used to standardise vascular access practice in the US is being evaluated and adapted for the UK.

It is a comprehensive and inclusive approach promoting the ‘right patient receives the right device at the right time’. This has the intention of minimising all complications of IV access, such as infections, phlebitis and vein loss, and the ensuing interruptions in treatment doses and the risk of subsequent treatment failure.

**Evidence based guidelines that promote principles of Vessel Health Preservation and facilitate device selection**

- Peripheral cannula is chosen for non-irritant, non-vesicant and Hyperosmolar infusion therapy with a relatively short anticipated duration of infusion therapy eg, less than 1 week (INS 2016)
- Episodic chemotherapy for more than 3 months (INS 2016)
- If 4 unsuccessful attempts of cannulation have been made, another plan for vascular access is required (INS 2016)
- Use a catheter with the minimum number of ports or lumens essential for management of the patient. *Class A* (EPIC III, INS 2016)
- CVAD selected is of the smallest outer diameter, with the fewest lumens needed (INS 2016)
- Avoid PICCs and subclavian placement due to risk of stenosis and thrombosis in patients with chronic kidney disease, who may need arteriovenous fistula or graft for hemodialysis (INS 2016)
- In selecting an appropriate intravascular insertion site, assess the risks for infection against the risks of mechanical complications and patient comfort. *Class D* EPIC III (Loveday 2014)
- Replace CVAD with 24-48 hours if the CVAD was placed in a suboptimal aseptic conditions, (INS 2016)

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- PICCs are associated with higher risk of thrombosis in patients who are critically ill and/or have a cancer diagnosis when compared to other CVADs (INS 2016).
- Use the upper extremity for non-tunnelled catheter placement unless medically contraindicated. (EPIC III 2014)
- Modified Seldinger Technique (MST) as the preferred method for PICC insertion due to advantages of decreased vein trauma, decreased insertion complications (INS 2016)
- Site selection for vascular access shall include assessment of the patient’s condition; age; diagnosis; comorbidities; condition of the vasculature at the insertion site and proximal to the intended insertion site; condition of skin at insertion site
- Catheter selection should be based on the prescribed therapy or treatment regimen, length of treatment, duration of dwell, vascular integrity, patient preference, and ability and resources available to care for the device. (INS 2016)
- Promptly remove any intravascular catheter that is no longer essential Category 1A (CDC 2011).

**CVAD device summary**

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<thead>
<tr>
<th>Site</th>
<th>Duration</th>
<th>Advantages</th>
<th>Disadvantages</th>
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| **PICC**              | Upper arm; basilic, brachial veins, Cephalic & enter the superior vena cava | Medium term 4 weeks to 6 months (Loveday et al 2014)                                                                        | 1) Thrombosis is highest in this device than other CVAD’s  
  2) Tip positioning can be difficult.  
  3) Potential for occlusion related to narrow lumens/longer length  
  A portion of the catheter is outside the skin |
| **TIVAD/ Implanted Port** | 1) Internal jugular, axillary & femoral vein. 2) Port chamber can be placed on anterior chest wall, lower lateral ribcage, anterior-lateral thigh or overlying the Trapezius muscle | Long term intermittent access Months/years                                                              | 1) Needles will be required when accessing the device  
  2) A more invasive procedure for insertion but can be done under local anaesthetic  
  3) 3-5cm Scar  
  Small risk of insertion related mechanical complications |
| **Non-tunnelled CVC** | Internal Jugular Axillary or femoral veins | 7-10 Days (Loveday et al 2014)                                                                                          | 1) Proportionally higher risk of infection vs tunnelled catheters and implanted ports (Maki et al 2002)  
  2) Short-term, if IV therapy is continued an alternative device will maybe required.  
  3) Small risk of insertion related mechanical complications |

**5.2 CVAD insertion with ultrasound guidance**

Nice Guidelines (2009) - Two-dimensional (2-D) imaging ultrasound guidance is recommended as the preferred method for insertion of central venous catheters (CVCs) into the internal jugular vein (IJV) in adults and children in elective situations.
The use of two-dimensional (2-D) imaging ultrasound guidance should be considered in most clinical circumstances where CVC insertion is necessary either electively or in an emergency situation.

It is recommended that all those involved in placing CVCs using two-dimensional (2-D) imaging ultrasound guidance should undertake appropriate training to achieve competence.

Audio-guided Doppler ultrasound guidance is not recommended for CVC insertion.

All patients undergoing insertion of a CVAD must have associated veins assessed using ultrasound. The assessment should look at patency, stenosis and thrombosis and general suitability of the vessel.

5.3 Referring for a CVAD

IMPORTANT - Prior to the insertion of any central venous access device it is the responsibility of the referrer to ensure patients are swabbed for MRSA (within 2 weeks of procedure) and take necessary bloods tests. Failure to prepare patient for procedure may lead to cancellation of the procedure.

Process:
1) Phone 3916 to book an insertion time with the Procedure team
2) Complete referral Discover - Documents and guidelines database
3) Obtain Full blood count, coagulation screen (INR) required for Tunneled CVC’s, Port implantation, non-tunneled CVC
4) Referrers please forward results to 0161-446-8265 (fax) or telephone 0161-446-3916
5) Screen all patients for MRSA (within 2 weeks of the procedure)

5.4 Informed consent

Who can seek consent?

It is important that the person who obtains the consent has sufficient knowledge about the procedure, including the potential risks and is trained in line with the Trust Consent to examination or treatment policy. For patients who are unable to consent to investigation or treatment a consent form 4 must be used. Please see consent to examination or treatment policy for further information;

Patient information

Patients should receive information regarding their vascular access device in advance of their procedure, though in urgent cases this might not always be possible but fully informed consent must always be obtained. A contact number will be provided should the patient have any questions in advance and allay any anxieties.

Information sheets are also available electronically;
TIVAD patient information; see Appendix 16

For medical purposes the term 'informed consent' means “permission granted in the knowledge of possible consequences (GMC 1998)

Patients have a fundamental legal and ethical right to determine what happens to their own bodies. If the principle of consent is not respected the individual may be liable to legal action or action by a professional body (DOH 2009)
5.5 Who can insert a central venous access device?
Registered healthcare professionals that have;
• Been assessed as competent in obtaining written informed consent
• Undergone specific training with a structured programme of training including training elements listed in Appendix 1.
• Undertaken a minimum of 25 supervised insertions (depending upon experience, aptitude and learning efficiency) and has been assessed as competent by a Consultant Interventional Radiologist or suitably competent healthcare professional.

Inserters should be familiar with policies, guidelines and relevant documents related to vascular access procedures such as; Epic III, NICE, BCSH, Saving lives etc and relevant Trust policies.

5.6 Insertion bundle
Prior to insertion a checklist should be undertaken including (Example; Appendix 2)
Use a central venous catheter Insertion bundle ensures a structured process for instituting best practice (Loveday et al 2014).
• Hand hygiene
• Maximal sterile barrier precautions during insertion (Hat, mask, gown, gloves, full body drapes)
• Use of 2% Chlorhexidine Gluconate (CHG) in 70% isopropyl alcohol for skin antisepsis at insertion site
• Avoid femoral site
• Remove CVC’s as soon as no longer indicated (EPIC III - Loveday et al 2014)

5.7 Who can access and provide care and maintenance for CVADs
CVADs can only be accessed for blood sampling and flushing by staff Band 3 and above who have received appropriate Trust training. Band 5’s and above who have received appropriate training, can access them to also administer via them.

For staff at The Christie initial training with the clinical skills training team or a standardised practice facilitator must occur before any supervised practice takes place. Healthcare workers must work under the supervision of a member of staff, assessed as competent to access CVADs, until they themselves have been assessed as competent to work without supervision. This assessment must be carried out by member of the clinical skills training team, a standard practice facilitator or a standardised clinical link worker. It is the responsibility of the healthcare worker to ensure this prior to working with central venous catheters. Following the initial assessment, a date will be arranged for update to be undertaken. The healthcare worker must contact the clinical skills training team or their designated link worker to arrange their scheduled update. Staff must also be reassessed following a period of prolonged absence or incident.

5.8 PICC
A PICC is a peripherally inserted central venous catheter. It is a thin flexible tube that is inserted into a vein in the upper arm. The PICC is then threaded along the vein so that the tip lies in the optimum tip position in the should be at the lower one-third of the Superior Vena Cava (SVC) at the level of the cavoatrial junction (CAJ) (INS 2016)

Currently there are two types of PICC used at The Christie;
**Groshong Valved®** - This PICC differs from tunnelled central venous catheters in that they have a pressure sensitive valve at the internal end which allows fluid to be injected into the catheter and blood to be withdrawn. When not in use the valve remains closed, thus preventing blood from flowing back into the catheter and air entering the venous circulation, clamps or switches are not required, this catheter is made of silicone.

**Open ended pressure injectable PICC with clamp** – This PICC is open ended does not have any integral valves. This catheter is made from polyurethane and is compatible with contrast pressure injection (please see manufacture guidelines for details).

**PICC Indications**
- For the delivery of antibiotic therapy, IV fluids, TPN, Blood sampling and the administration of irritants/vesicants, chemotherapy and other anti-cancer intravenous therapies (BCSH 2006)
- Episodic chemotherapy for more than 3 months (INS 2016)
- Patients with limited venous access
- Previous complications or failed with short PIVs
- Patient requires irritating or hyperosmolar medications
- A patient who has bleeding disorder/low platelet levels (<50 · 10⁹/l) where central insertion is contraindicated (BCSH 2006, Hertzog & Waybill 2008)
- Total Parenteral Nutrition (Espen 2009)
- Conditions that contraindicate use of other VAD types
- Need for high pressure injection of contrast media
- Patients consideration
- Intended dwell time – medium term 4 weeks to 6 months (Loveday et al 2014).

**Benefits**
- A PICC can be used to give chemotherapy, anti-cancer drugs, fluids, antibiotics and other drugs directly into the vein.
- It can also be used for taking blood samples. PICC lines are inserted when a central line is needed for short term use
- PICCs are useful when peripheral veins are difficult to find or access, or for people who are very anxious about needles. A PICC is sometimes used temporarily for people who should ideally have a tunnelled central venous catheter or an implantable Port but are not able to because of their clinical condition i.e. anatomy abnormal, blood results or are unable to lie flat. In this situation a PICC means treatment can be started without delay.

**Relative Contraindications**
- Inability to visualise, view suitable vein (Ultrasound)
- Circulatory impairment, peripheral neuropathies
- History of CVA (same side as limb weakness), same side as mastectomy with node dissection surgery, upper extremity fistulas, surgery or neck/head trauma
- Venous thrombosis in vasculature where the PICC is proposed
- Same side as; existing CVAD, cardiac pacemaker, implanted cardiovascular defibrillator, (SVC Vena Cava filter) (Hertzog & Waybill 2008)
- Skin conditions, dermatitis, infection at insertion site (Hertzog & Waybill 2008)
- End stage renal disease, chronic renal insufficiency, to preserve upper limb vasculature for formation of fistula (NKF-DOQI)
- Allergy to material contained in the device.
**PICCs and Ultrasound pre-assessment**

Use Ultrasound pre-assessment to assess optimal vessel size. Ideal vessel should ideally be 3 times greater than the external diameter of the catheter. As a rough guide; 4fr vein size should be no less than 4mm, 5fr no less than 5mm (Nifong & McDevitt 2011). A higher catheter vein ratio may increase thrombosis risk and should be balanced accordingly on the clinical urgency for the PICC CVC vs catheter related and patient thrombotic risk factors. Undertaking comprehensive ultrasound assessment including upper arm veins, axillary, subclavian, internal jugular, brachiocephalic including compression to exclude major abnormalities such as pre-existing thrombosis and select the most appropriate vein using the SIP protocol (Safe insertion of PICCs) advocated by (Pittiruti et al 2014).

**Insertion: general principles.**

In addition to the insertion bundle advocated in EPIC guidelines (Loveday et al 2014) other considerations for PICC insertion are;

- Optimal catheter site selection; Upper arm approach unless contraindicated The Basilic vein is the preferred vein (INS 2016), the Cephalic and Brachial veins may be considered but with caution as the Brachial vein is often in close proximity nerve bundles and artery. The Cephalic vein follows narrower more tortuous pathway and connects to the axillary vein at an acute angle. It is essential ultrasound assessment takes place and the optimal vein is selected on an patient individual basis

- PICCs may be preferred CVAD in patients with coagulopathy disorders or receiving anticoagulants as direct pressure can be applied to control bleeding and in patients with respiratory diagnoses or intracranial bleeding when Trendelenburg position is difficult or contraindicated (INS 2016)

- Cannulation should be ultrasound guided (NICE 2002)

- Modified Seldinger Technique, to reduce vessel trauma (INS 2016)

- Secure with sutureless device (CDC 2011).

Catheter tip position maybe confirmed with ECG tip confirmation system (Nice 2015) and/or fluoroscopy chest X-ray.

If unable to gain satisfactory P-Wave (ECG tip confirmation system) or the patient does not have a discernible P-Wave (Atrial fibrillation or arhythmia). Tip position will be confirmed with chest x-ray and documented.

Patients with arm ports or PICCs should not have additional IV devices placed on the same arm i.e. IV cannula.

**Complications**

- Misplacement of catheter tip following insertion sometimes requires adjustment, this sometimes means includes going to X-ray dept

- Accidental puncture of the artery which may cause bleeding

- Inability to thread catheter into vein/SVC

- Inability to advance sheath/dilator assembly

- Bleeding and some bruising around the insertion site, especially in people whose blood does not clot normally

- Arrhythmias

- Haematoma

- Phlebitis

- Infection, insertion site or systemic
- Nerve injury – patients may feel a shooting pain down their arm if the needle touches a nerve
- Embolism, air or equipment
- Cardiac tamponade, secondary to vessel wall, atrial or ventricular perforation
- Occlusion
- Sepsis
- Phlebitis.

Dressings & Flushing see PICC care and maintenance (Appendix 10)
PICCs must be flushed weekly with 10mls 0.9% sterile saline to maintain patency. This must be done using a firm push-pause technique clamping as the last ml is inserted to maintain positive pressure.

PICCs should be dressed with a securement devices i.e. Griplock®, Statlock® together with steri-strips (12mm width) and transparent semi-permeable dressing. Change dressing, securement device, and steri-strips within 24hrs post procedure and removed gauze if in place. Thereafter, change dressing every 7 days, or sooner, if they are no longer intact or if moisture collects under the dressing (Loveday et al 2014). For more information refer to Care of your peripherally inserted central venous catheter.

PICC dressing including, Statock® and steri-strips

PICC removal, by registered health care professional.
While removing, apply slow, intermittent traction without applying any direct pressure at the insertion site. During the removal process there is possibility that resistance may be present. This could be caused by a venous spasm. Warmth often relieves the spasm and after a brief period of time may easily be removed.

If resistance does occur the catheter must never be forcefully pulled as this could break it and lead to a section embolising in the circulatory system. If still unable to remove it easily, the Procedure team/Radiologists should be contacted.

Treatment – apply warm compress to area for 20 minutes (for vasodilation and to alleviate venous spasm) and re attempt removal. If resistance persists the catheters should not be forcibly removed inform, Procedure team, Radiologists (Drewett 2000).

Following removal of the PICC, pressure should be applied to the site for 2–3 minutes to ensure the bleeding has stopped. Apply a sterile transparent dressing and remove after 24hours.

PICC Complications & Management, post insertion

Phlebitis
- Can be mechanical, chemical or bacterial (RCN 2010, INS 2016). Mechanical phlebitis is caused by irritation of the venous endothelium by the catheter.
Symptoms – Tenderness, erythema, warmth, swelling, induration, pain (INS 2016).

Treatment
- Anti-inflammatory analgesics for inflammation & pain (Higginson 2011)
- Culture and begin antimicrobial therapy if infection present
- Site care and condition/appearance using standardised local assessment scales for phlebitis and/or infiltration/extravasation (RCN 2010)
- Refer to Physician, remove if symptoms don’t improve (Simcock 2001).

Prevention –
- Regular monitoring of site, Aseptic technique during insertion and care/maintenance (INS 2016)
- Stabilisation: Catheter securement to reduce movement (CDC 2011)
- Site: Upper arm U/S rather than ante-cubital (Espen 2009)
- Modified Seldinger Technique reduces trauma (INS 2016)
- Prevention – Upper arm, Modified Seldinger Technique
- Use the Device with ‘Smallest gauge’ (INS 2016).
- ‘The diameter of the catheter should be one third or less of the diameter of the vein, as checked by ultrasound’ (Espen 2009).

Catheter tip Malposition - Ideal Catheter tip position:- Lower one third of the SVC level with the cavoatrial junction (INS 2016)

Symptoms
- Alteration of external catheter length, inability to aspirate blood, “gurgling” sound in ear, discomfort/pain in chest, neck or back arrhythmias (INS 2016)

Treatment
- If the PICC has been pulled out from its original insertion length; please follow PICC Occlusion Algorithm (Appendix 4)
- A chest X-ray may be required to ascertain tip position
- If too short or malpositioned refer to procedure team/Interventional Radiology for repositioning, catheter exchange/replacement
- The nurse should not advance any external portion of the CVAD that has been in contact with skin into the insertion site. (INS 2016)
- A CVAD with a malpositioned catheter tip location that cannot be repositioned to a central vein should be removed (INS 2016)
- Catheters that are malposition have a higher risk of thrombosis if the patient is symptomatic of thrombosis an ultrasound Doppler should be undertaken to assess the associated vessels.

Prevention
- Take meticulous care not to pull PICC out when changing dressings
- Optimal patient positioning during insertion
- Use griplock/statlock securement with Steri-strips
- Thread catheter slowly, do not thread against resistance
- Use of tip navigation/positioning systems/Fluoroscopy.

Insertion related bleeding

Symptoms
- Persistent bleeding, haematoma/swelling from insertion site

Treatment
- Optimal pressure (RCN 2010) should be applied with a sterile dressing following the venipuncture and maintained until bleeding stops (INS 2011)
- Take full blood count & full coagulation screen, is there an underlying cause.
• Use ultrasound guided & Modified Seldinger technique to minimise trauma to
vein, visualise needle tip at all time during insertion
• Assess patient history & current medications prior to insertion
• Re; Use of blood of products; Assess need on an individual basis rather than
routine correction of all minor abnormalities of platelet count and coagulation
studies (BCSH 2006).

5.9 Catheter related Thrombosis
Symptoms;
• Discomfort/pain/Oedema/engorged veins; arm, shoulder, neck, chest
• Leakage of fluid at entry site, discoloration of extremity, unexplained fever
Treatment
• Refer to Respective medical team and to Radiologist to undertake ultrasound
Doppler
• Do not remove the catheter CVAD in the presence of a CVAD-associated vein
thrombosis when the catheter is correctly positioned at the cavo-atrial junction,
the catheter is functioning with a blood return, there is no evidence of any
infection, and there is absence of severe deep vein thrombosis (DVT) - related
symptoms causing pain Prevention (INS 2016).
This is ultimately the medical team's decision based upon, the need to keep the
device in place and balanced individual hypercoagulability factors of the patient.
Monitoring and anticoagulation should be followed in accordance with the
Anticoagulant Therapy Management Policy.

Prevention
• Smallest outer diameter (INS 2016)
• ‘the diameter of the catheter should be one third or less of the diameter of the
vein, as checked by ultrasound’ (Espen 2009)
• Assess patient risk factors i.e. malignancy, critically ill (Chopra et al Lancet 2013)
• Catheter tip position (RCN 2010)
• Modified Seldinger for insertion reduces trauma.

Catheter Occlusion
Treatment
• Eliminate mechanical causes (kinking, pinching of line)
• If chemical, determine which infusates caused precipitates and instill proper
solution (alkaline/acidic) to clear (INS 2016)
• If blood occlusion, instill Urokinase® per protocol.

Prevention
• Flush before and after all infusions and blood draws with an adequate amount of
solution to fully clear device
• Be aware of solution incompatibilities prior to infusion
• Pulsatile flushing technique.

5.10 Non-tunnelled or Percutaneous central venous catheters
Usually found in acute-care settings, critical care, theatre and high
dependency units. A non-tunnelled CVC or percutaneous CVC is a
centrally inserted catheter. It is a polyurethane line, which has
multiple entry ports between 3 and 6 and they are known as lumens.
It is threaded through a large vein and usually sutures are used to
hold it in place with a transparent semi-permeable dressing.
Do not routinely replace CVC access devices to prevent catheter related infection.

**Indications**
- In an acute situation, where the patient deteriorates quickly.
- When the patient urgently needs IV fluids, drugs and anti-biotics
- When patient’s need lots of IV medication at the same time
- If the patient needs specialist monitoring, carried out in Critical Care
- For intended dwell time 7-10 days.

**Benefits**
- Allows patients to receive several amounts of drugs simultaneously
- Allows patients to receive drugs in an emergency
- Allows patients to receive drugs that could irritate smaller vessels
- Allows patients to be ‘centrally monitored’
- Can be inserted quickly and removed quickly.

**Risks**
- Carries high infection risk potential
- Pneumothorax on insertion
- Air emboli on insertion or removal
- Patient discomfort
- Patients often need nursing in specific areas (Morton, 2011, Matching Michigan 2012)
- Infection: as with all surgical procedures there is a risk of infection occurring.
- Accidental puncture of the artery
- Embolus: a small risk of embolus (air/equipment) if the catheter or guide-wire breaks, or air embolus (bubble of air) within the blood stream.
- Cardiac arrhythmia
- Malpositioning.

**Specific points;**
- Axillary vein; insertion can increase risk of pneumothorax secondary to the close proximity of the pleura to the needle insertion site. Damage to the brachial plexus is a risk of this approach (INS 2016)
- Femoral vein; insertion sites present higher risk of infection and should be avoided when possible since catheter stabilisation and occlusive dressing adherence can be difficult to maintain. Femoral insertion does not require Tendelenburg position.
- Subclavian vein; Ultrasound evaluation of the infraclavicular subclavian vein is difficult secondary to the overlying clavicle. Moving outward to the axillary vein make use of the ultrasound more useful. Pinch-off syndrome, where the catheter gets trapped between the first rib and clavicle, can be a risk of subclavian insertions.
- Internal jugular vein insertion sites may be difficult to secure secondary to head movement, and carotid artery puncture is a risk due to its proximity to the vein
- External jugular veins; are more tortuous than internal jugular veins, making catheter threading more difficult, the vessel is superficial which may create difficulty with catheter securement.
5.11 Haematological blood tests for CVADs (Vascular devices other than PICC)

**Insertion Criteria**
Skin tunnelled CVC’s and Implantable Ports, minimum requirements
- Hb 80g/L
- Platelets 50 x10^9/L
- INR less than 1.5 (BCSH 2010)
For further information see referral SOP (Appendix 17).

5.12 Anticoagulants and CVADs (Not relevant for PICCs)

**Intravenous unfractionated (UFH)**
Should be stopped 6 hours before catheter insertion and restarted when haemostasis is secured (Martindale).

**Low molecular weight heparin.**
In patients receiving prophylactic subcutaneous low molecular weight heparin (LMWH) catheter insertion can be undertaken 12 hours after the last injection; for patients receiving therapeutic subcutaneous LMWH, the time to catheter insertion should be extended to 18 hours after the last injection. Heparin can be recommenced once haemostasis is secure, usually within 2 hours of catheter insertion. Substitution with intravenously infused UFH or insertion of an IVC filter should be considered if there is a very high thrombotic risk. Expert haematology advice should be sought.

LMWH may continue >4hrs post removal of the CVAD in patient without abnormal clotting or high risk of bleeding.

5.13 Direct Oral Anticoagulants

**Warfarin**
Patients should be given advice to stop warfarin prior to admission for their vascular device insertion. Provided adequate oral intake and absorption of vitamin-K-containing foods omission of warfarin for 3 days will allow reduction from range 2.0-3.0 down to level safe (eg less than 1.5) for invasive procedures such as central access/biopsies (5 days if in range 3.0-4.0).

Dabigatran (Pradaxa) which is a direct thrombin inhibitor and the direct factor Xa inhibitors Rivaroxaban (Xarelto) and Apixaban (Elquis) and Edoxoban are new drugs which are direct oral anticoagulants and may replace warfarin and other anticoagulants. There is currently no proven method to reverse their anticoagulant effect in bleeding patients and no agreed national guidelines.

Dabigatran **discontinue 2 days prior to the procedure** has a half-life of approximately 12-17 hours.

Rivaroxaban **discontinue 2 days prior to the procedure** has a half-life of approximately 7-11 hours. The effects of these drugs extend in patients with renal impairment. ([https://www.medicinescomplete.com/mc/martindale/current/ms-20786-b.htm?q=dabigatran&t=search&ss=text&p=1#_hit](https://www.medicinescomplete.com/mc/martindale/current/ms-20786-b.htm?q=dabigatran&t=search&ss=text&p=1#_hit))

Apixaban **discontinue 2 days prior to the procedure**

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Half-life 10-15 hours but is only partially renally excreted (30%). For central venous access procedures the drug should be discontinued for 48 hours. For Dabigatran/Rivaroxaban a normal PT and APTT suggest that the anticoagulant effect of the drug has worn off.

**Normal PT and APTT are not reliable indicators to exclude significant concentrations of Apixaban.**

**Edoxaban discontinue 2 days prior to the procedure**
Edoxaban, 50% of the drug is renally excreted and the half-life is 10-14 hours, it should be discontinued preferably at least 24 hours before the procedure (BCSH 2016).

**Stopping DOACs before surgery or invasive procedures for which anticoagulation needs to be stopped** (Table below).

<table>
<thead>
<tr>
<th>Renal Function CrCl ml/min</th>
<th>Estimated half-life (hours)</th>
<th>Low bleeding risk</th>
<th>High bleeding risk</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Dabigatran</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;80</td>
<td>13</td>
<td>24 hours</td>
<td>48 hours</td>
</tr>
<tr>
<td>&gt;50 to &lt;80</td>
<td>15</td>
<td>24-48 hours</td>
<td>48-72 hours</td>
</tr>
<tr>
<td>&gt;30 to &lt;50</td>
<td>18</td>
<td>48-72 hours</td>
<td>96 hours</td>
</tr>
<tr>
<td><strong>Rivaroxaban</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;30</td>
<td>9</td>
<td>24 hours</td>
<td>48 hours</td>
</tr>
<tr>
<td>&lt;30</td>
<td>48 hours</td>
<td></td>
<td>72 hours</td>
</tr>
<tr>
<td><strong>Apixaban</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;30</td>
<td>8</td>
<td>24 hours</td>
<td>48 hours</td>
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<tr>
<td>&lt;30</td>
<td>48 hours</td>
<td></td>
<td>72 hours</td>
</tr>
<tr>
<td><strong>Edoxaban</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>&gt;30</td>
<td>10-14</td>
<td>24 hours</td>
<td>48 hours</td>
</tr>
<tr>
<td>&lt;30</td>
<td>48 hours</td>
<td></td>
<td>72 hours</td>
</tr>
</tbody>
</table>

*All decisions need to consider state of haemostasis and new acquired risk factors for bleeding BCSH (2016). For further information see Anticoagulant therapy Management Policy.

Direct oral anticoagulants can usually be restarted the when complete haemostasis has been secured but if there is concern over bleeding then re-introduction should be delayed, bridging therapy with low molecular weight heparin may be required if there is a >24 hour delay.

If a patient requires emergency central venous catheter then a PT and APTT should be performed, if this is normal there is no residual activity of Rivaroxiban/Dabigatran and the operation can go ahead as normal. If it is prolonged the operation should be delayed as long as possible to allow drug levels to fall or consider a PICC or peripheral IV support. Apixaban and Edoxaban require specialist assay tests, please liaise with a Haematologist.

If an urgent CVC is needed then at least 12 hours is recommended from the last dose until insertion and will be considered an individual basis balancing the risks of the procedure and complications against the benefits for the patients.

**Intravenous unfractionated (UFH)** should be stopped 3 hours before catheter insertion and restarted when haemostostasis is secured.
Anti-platelet therapy
Patients taking anti-platelet medication (e.g. Aspirin, Dipyridamol, Clopidogrel) should discontinue 5 days prior to CVAD insertion.

Emergency/unplanned patients on low dose Aspirin 75mg will be considered for central venous access REMOVAL on an individual basis balancing the risks of the procedure and complications against the benefits for the patients and urgency of device and treatment. The patient should not have problems with their blood clotting and not on concomitant anticoagulants/Antiplatelet therapy.

AVASTIN (Bevacizumab)
Bevacizumab interrupts angiogenesis by inhibiting vascular endothelial growth factor.

Tunneled CVC – Stop Bevacizumab 14 days prior to insertion and it should not be administered until 24hrs after insertion, providing there is no bleeding at the site.

TIVAD – Stop Bevacizumab 4 weeks prior to insertion and not to be used until implantation site has healed, usually 7-10 days.

Complications that occur from patients commenced on Bevacizumab within the above time frame will be the responsibility of the managing Consultant.

5.14 Thrombosis Management and CVADs
Do not remove the catheter CVAD in the presence of a CVAD-associated vein thrombosis when the catheter is correctly positioned at the cavo-atrial junction, the catheter is functioning with a blood return, there is no evidence of any infection, and there is absence of severe deep vein thrombosis (DVT) - related symptoms causing pain (INS 2016).

When a CVAD related thrombus is present and the removal is indicated, anticoagulation should be undertaken prior to the procedure, for thrombosis extending to the Superior Vena Cava or Right Atrium up to 7 days anticoagulation should be considered.

This is ultimately the medical team’s decision based upon, the need to keep the device in place and balanced individual hypercoagulability factors of the patient. Monitoring and anticoagulation should be followed in accordance with the Anticoagulant Therapy Management Policy.

Risk factors;
- History of deep vein thrombosis
- Presence of chronic disease associated with a hypercoaguable state such as cancer, diabetes, irritable bowel syndrome, congenital heart disease and end stage renal failure.
- Surgical and trauma patients
- Critical care patients, hyperglycemia in critical care
- Known presence of genetic coagulation abnormalities (eg Factor V Leiden, prothrombin mutation)
- Pregnancy or the use of oral contraceptives
- Fluid volume deficit
- History of multiple CVAD, especially with difficult or traumatic insertion and the presence of other intravascular devices (eg, pacemakers).
The presence of carcinoma, metastatic disease predisposes oncology patients to hypercoagulability and as a consequence device related thrombosis is high in this group of patients (Chopra et al Lancet 2013).

5.15 Tunnelled central venous catheter

A tunnelled central venous catheter is a long fine hollow tube with an opening at each end. One end is outside (usually anterior chest wall), the other end is situated in a large central vein. There is a Dacron cuff on the CVC which engrafts into the tissue in the skin tunnel, this takes up to three weeks for this to occur, which holds it in position under the skin and prevents it falling out. It can remain in position for several months/years.

A skin tunnelled CVC is usually inserted via the internal jugular, axillary veins and the tip of the catheter is advanced into the lower one-third of the Superior Vena Cava (SVC) at the level of the cavoatrial junction (CAJ) (INS 2016). For femoral placements the tip should lie in the inferior vena cava, with the tip higher than the diaphragm (INS 2016).

The technique of skin tunnelling reduces the risk of sepsis, as the exit site is distant from the site of insertion.

Following insertion there is one suture at the insertion site (usually jugular at the base of the neck) and one or two at the exit site. The top sutures should be removed after 7 days and the exit site sutures should remain in situ for at least 21 days or until the skin appears healed. After this time, the granulation tissue will have grown around the Dacron cuff, securing the line in situ. After insertion the site will be dressed with gauze and a semi-permeable, transparent dressing. After 24 hours the dressing needs to be taken down, the site cleaned with 2% Chlorhexidine in alcohol, single use applicator and allowed to air dry. A large semi-permeable, transparent dressing must be applied and changed weekly, or sooner if soiled.

Following removal of the suture at the exit site (21 days), a dressing is not required providing the exit site remains clean and dry. Tunnelled lines must remain looped and secure at all times with surgical tape when a semi-permeable transparent dressing is not in place.
Tip position may be confirmed with ECG tip confirmation system and/or fluoroscopy, chest X-ray and ultrasound may be used to assess for and identify the “sliding lung sign” to assess for pneumothorax.

If unable to gain satisfactory P-Wave (ECG tip confirmation system) or the patient does not have a discernible P-Wave (Atrial fibrillation or arrhythmia). Tip position will be confirmed with chest x-ray and documented.

**Indications**
- Long term IV therapy Months/years (EPIC III 2014)
- As well as the delivery of antibiotic therapy, IV fluids, TPN, Blood sampling and the administration of irritants/vesicants, chemotherapy and other anti-cancer intravenous therapies.
- Patients with limited venous access
- Previous complications or failed with short PIVs
- Patient requires irritating or hyperosmolar medications
- Patients consideration
- Plasmapheresis – In this instance the catheter must be designed and licensed for Plasmapheresis and be compatible with Plasmapheresis machines.
- Conditions requiring multiple punctures, such as patient’s requiring daily blood withdrawals.
- Total parenteral nutrition.

**Benefits**
- More comfortable and discreet and lower infection rates than the non – tunneled catheters.
- Can remain in situ for long periods
- It allows some therapies to be given at home
- It may also be used for taking blood samples which are needed regularly, avoiding the need for repeated needle stabs in your arm.

**Disadvantages**
Requires a minimally invasive surgical procedure for placement.
Potential complications on insertion
Body image of external catheter
Lifestyle, no swimming.

**Risks**
Insertion related complications;
- Air embolus
- Infection: as with all surgical procedures there is a risk of infection occurring.
- Brachial plexus injury
- Cardiac tamponade
- Endocarditis
- Pneumothorax of the lung: Rate 1 per 13,000 cases
- Accidental puncture of the artery: Rate 1 per 1,500 cases
- Embolus: a small risk of embolus such as air or equipment, if the catheter or guide-wire breaks, or air embolus (bubble of air) within the blood stream.
- Haematoma
- Scarring
- Cardiac arrhythmia
- Malpositioning
- Breakage between first rib/clavicle (Subclavian approach)
- Hydrothorax
- Laceration of the vessel
- Perforation of vessel
- Thoracic duct injury
- Risks normally associated with local anaesthesia

Complications of indwelling catheter;
- Blockage
- Catheter/cuff erosion through skin
- Exit site necrosis/scarring over the catheter site
- Infection, at exit site or systemically
- Fracture of catheter
- Fibrin sheath formation and persistent withdrawal occlusion (Appendix 5)
- Damage, fracture, burst, trauma
- Thrombosis: a blood clot may develop. This is a longer term risk associated with lines. Signs of this include pain, swelling and discomfort in the neck or arm on the side the catheter is placed.
- Extravasation
- Intolerance to the device material
- Spontaneous catheter tip malposition
- Intracardiac thrombosis.

Flushing
Tunneled CVCs must be flushed weekly with 10mls 0.9% normal sterile saline to maintain patency. This must be done using a firm push-pause technique clamping as the last ml is inserted to maintain positive pressure. See tunnelled CVC care and maintenance Appendix 9.

Apheresis catheters – These types of catheters require flushing with 0.9% normal sterile saline and locking with higher does Heparin according to individual patient prescription. This is usually 5000 units Heparin per ml (or as per manufacturer instructions) the volume required is usually printed on each lumen of the catheter.

Tunneled CVC removal
The device should be removed when no longer required or if there is thrombosis or infection related to the catheter or if it is fractured.

The procedure may be undertaken by registered health care professionals who have been trained to undertake the procedure and has a documented assessment of competency, or under the supervision of an experienced operator. Incorrect removal may lead to catheter fracture and embolism, dislodgment of a portion of the catheter into the bloodstream. Removal must be according to SOP for the removal of cuffed tunneled central venous catheter.
Criteria:
- The haemoglobin and white cell count are not critical
- Platelets must be above 30x10^9/litre
- Formal coagulation check is not required.

Patient with history of a known bleeding disorder, history of bleeding should undergo full coagulation screen and take advice from Haematologist if required.

Removal and Anticoagulants
For anticoagulants see Tunnelled CVCs and Anticoagulants.

Aspirin and Aspirin containing products.
- Emergency/unplanned patients on low dose Aspirin 75mg will be considered for central venous access REMOVAL on an individual basis balancing the risks of the procedure and complications against the benefits for the patients and urgency of device and treatment. The patient should not have problems with their blood clotting and not on concomitant anticoagulants/Antiplatelet therapy.

5.16 TIVAD/Implanted Ports
TIVADs (Totally implanted venous access device) are also known as Ports or Portacaths®. The TIVAD is an implantable venous access device consisting of a reservoir with injectable silicone septum, which is called the Port chamber; this is a self-closing membrane. Attached to the Port chamber is a central venous catheter, the tip of the catheter should be at the lower one-third of the Superior Vena Cava (SVC) at the level of the cavoatrial junction (CAJ) (INS 2016). Ports must be flushed monthly with 0.9% normal sterile saline to maintain patency. This must be done using a firm push-pause technique to maintain positive pressure.

These are usually recommended for patients who need intermittent intravenous therapy over a long period of time (Loveday et al 2014) but in recent years the practice as The Christie has shown that this device is also suitable for some types of ambulatory IV therapy. Ports have to be inserted using a sterile procedure which is usually carried out under local anaesthetic and require intraprocedural catheter tip guidance ie. Fluoroscopy/ECG based guidance system.

A TIVAD is implanted by making an incision into the patient’s skin, creating a subcutaneous pocket. Anchor sutures are not routinely used but maybe used to anchor the chamber in place for patients with increased body mass. The overlying tissues and skin are then surgically closed. All the sutures used are all absorbable and but some may require trimming 7 days (at the base of the neck and port incision site) post TIVAD insertion and the wound checked for healing, at this point the skin and tissues should be closed with minimal eschar.
Tip position maybe confirmed with ECG tip confirmation system and/or fluoroscopy and chest X-ray.

If unable to gain satisfactory P-Wave (ECG tip confirmation system) or the patient does not have a discernible P-Wave (Atrial fibrillation or arhythmia). Tip position will be confirmed with chest x-ray and documented.

For inserters with documented competency, ultrasound maybe used to assess for and identifying the “sliding lung sign” to perform point of care assessment for pneumothorax. (Blaivas et al 2005, Lichtenstein et al 2005). If during or post procedure the patient becomes symptomatic of a pneumothorax an urgent X-ray must be performed.

Following implantation the tissues around the chamber may be anaesthetised for 2-4 hours hrs from the time the anaesthetic is administered. The TIVAD should not be used to administer chemotherapy/anti-cancer or any medications that could damage the tissues should extravasation occur, until the local anaesthetic has worn off and the patient has regained sensation to the area. Desensitized tissues carry an increased the risk of extravasation being identified quickly should it occur.

The TIVAD can be used on the day of insertion following the procedure but the access needle should placed intra-procedurally by the Specialist Nurse/Doctor implanting the TIVAD. It may then be used once full sensation is restored to the implantation site. When the TIVAD is not being accessed the device only requires flushing once monthly.

The TIVAD is accessed using a right angled non-coring needle connected to extension tubing with a clamp. A non-coring needle must be used with a TIVAD as hypodermic needles damage the silicone septum by drilling out a ‘core’ and this may lead to extravasation. The length of needle appropriate for the patient will be documented on the patient’s handheld card. This is a guide as the patient’s body mass may change and then an alternative size of non-coring needle maybe required for access, and should be assess each time the device is accessed.

A non-coring needle may stay in situ up to 7 days if required (INS 2016). For outpatients access is usually only required for 48 hours, but can the needle can remain in situ if the patient is undergoing continuous or frequent IV therapy. Strict aseptic non-touch technique must be observed. Any problems or signs of infection must be reported immediately as with any central access device.

Indications
- Long term therapy (Months/years)
- As well as the delivery of antibiotic therapy, IV fluids, TPN, Blood sampling and the administration of irritants/vesicants, chemotherapy and other anti-cancer intravenous therapies.
- Patients with limited venous access
- Previous complications or failed with short PIVs
- Patient requires irritating or hyperosmolar medications
- Patients consideration

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• Prolonged intravenous therapy.

**Benefits (TIVAD)**
Less impact on patient body image
No risk of pulling out the device
Lifestyle advantages, swimming
Lower risk of infection versus other CVADs
Less community care input, devices only need flushing monthly when not in use.

**Disadvantages**
• Still requires a needle to access device
• More invasive to insert than other vascular devices.

**Flushing**
TIVADs must be flushed monthly with 10mls 0.9% normal sterile saline to maintain patency. This must be done using a firm pulsating technique (See Appendix 12).

**Risks**
Bruising or pain at the site of insertion may occur following implantation this usually settles over a few days.

**Infection** - The procedure is carried out using full sterile conditions but there are still risks of the Port getting infected – this may be local infection at the skin or a more general bloodstream infection.

**Port erosion** - There is a small chance that the Port may wear thin the tissue above the Port or even break the skin, patients who lose significant body mass may are at increased risk of this occurring.

**Emboli** – Air or equipment lost into the bloodstream.

**Thrombosis** - Signs of this include pain, swelling in the neck or arm on the side the TIVAD is placed, the arm may show signs of discolouration.

**Arterial puncture** - Puncture of the artery that may cause bleeding, approximately 1 in 1000 patients may have an arterial puncture.

**Lung puncture** - This happens in approximately than 1 every 13,000 (The Christie data) of patients and may need further treatment to avoid breathing complications, further X-rays, a stay in hospital and possible chest drain insertion.

**Stenosis** - The vein may become narrowed or stenosed. If this happens then we may need to put the catheter into a different vein.

**Extravasation** - Damage caused by leakage of solution from the vein to the surrounding tissue spaces during intravenous administration. Extravasation of some Chemotherapy/anti-cancer drugs from Ports can cause tissue necrosis (tissue damage) and may need surgical removal of the Port and the affected tissue. Established fibrin sheath formation have in rare instances lead to extravasation, symptoms of this are pain, erythema, swelling at the site where the catheter enters the vein, i.e. base of neck.
**Mechanical failure.** A break in the catheter or the connection that may lead to extravasation.

**Port rotation/movement.** Refers to the Port moving out of place or turning round within the chest wall in this instance the Port may have to be repositioned or removed.

**Scarring.** There will be a small scar at the base of the neck and a 3-5cm scar where the Port is inserted. The catheter under the skin at the base of the neck may also be visible, scarring may appear worse if the device becomes infected.

**Other complications that may occur include;**
- Haematoma formation
- Brachial plexus injury
- Cardiac arrhythmias
- Endocarditis
- Hydrothorax
- Intolerance reaction to the implanted biomaterial
- Inflammation, necrosis, or scarring of skin over the implanted area
- Thoracic duct injury
- Pericardial tamponade (thrombus formation)
- Spontaneous catheter tip malposition or retraction
- Vessel erosion

PFM medical – Instructions for use (PFMmedical.com).

**Removal**
Patients should be prepared for the removal of the Port using the same haematological requirements for insertion.

**Femoral/Trapezius/tunnelled CVCs/implanted Ports**
On occasion situations arise where placing a central venous device in the upper body is contraindicated, SVCO (Superior vena cava obstruction) or upper body/limb veins are unable to accommodate a CVAD.

**Femoral** - These devices should be placed under fluoroscopy guidance the tip of the catheter should dwell in the inferior vena cava above the level of the diaphragm (INS 2011). The device exit site if inserted in the femoral vein should be on the anterolateral thigh unless contraindicated.

Femoral catheters are at increased risk of thrombosis and persistent withdrawal occlusion. They should be flushed routinely as a normally placed tunnelled CVC but if catheter occlusion problems are on-going, using a lock of higher dose Heparin lock solution may be required i.e Heparin 5000 international units p/ml. Non-tunnelled CVC placed via the femoral vein, consider using biopatch®, as the site is higher risk of infection (Loveday et al 2014).

**Trapezius** – the placement of tunnelled CVC and implanted port overlying the Trapezius muscle maybe appropriate if the usual sites are contraindicated i.e localised skin infection, previous radiotherapy where tissue is at risk of breaking down, subcutaneous disease on the anterior chest wall that may spread over the exit or implantation site.

**Arm TIVADs**
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Arm ports maybe used as an alternative site for patients whom chest placement, and access to internal jugular and axillary veins is not possible. The ports can be placed in the forearm or upper arm (ideally). Accessing, care and maintenance of an arm TIVAD is done in the same way as a TIVAD placed on the chest wall. Patients with arm ports or PICCs should not have additional IV devices placed on the same arm i.e IV cannula.

5.17 CVAD Complications & Management CVAD infection

If a patient develops a pyrexia of unknown origin, blood cultures should be taken from each lumen of the catheter and from a peripheral vein. If attempting to salvage an infected CVAD then intravenous antibiotics should be given via the CVAD.

This should be considered if there is a subcutaneous tunnel or periport infection, septic emboli, hypotension associated with catheter use, or a non-patent catheter.

Specific infections where line removal is recommended include Candida spp and other fungi, and *P. aeruginosa*, and *S. aureus*. Other bacteria that may require removal due to persistent infection/colonisation of the line include *Corynebacterium jeikeium*, *Stenotrophomonas maltophilia*, *Bacillus spp* and *Acinetobacter spp*. Single isolates require confirmation with a repeat blood culture ideally from all lumens of the line and peripherally.

Persistent coagulase negative *staphylococcal* infections may also necessitate line removal.

Where line removal is not possible alcohol 70% injection may be used to try to decontaminate lines. Absolute alcohol injection may be diluted to 70% with water for Injection and used to lock an infected lumen for up to 7 days. Please contact pharmacy and/or microbiologists for advice.

**Not to be used in catheters composed of polyurethane (or check manufacturer guidelines),** i.e. open ended clamped PICC, PFM® implanted Ports (TIVADs).

**Guidelines for the management of sepsis (including Neutropenic Sepsis)**

**Infection**

**Exit site** – Skin/tissue immediately around the catheter exit site has erythema, swelling or is exuding pus. Send swab for MC&S and commence appropriate antibiotics. Monitor for signs of systemic infection.

**Tunnel** - (erythema tracking along the subcutaneous tunnel) Send swab for MC&S and commence appropriate antibiotics. Monitor for signs of systemic infection. If erythema extends to the cuff it becomes very difficult to eradicate infection and in most instances it is appropriate the device is removed.

**Intra-luminal/Systemic** - (resulting in systemic sepsis) If a patient develops a pyrexia of unknown origin, blood cultures should be taken from each lumen of the catheter and from a peripheral vein. Intravenous antibiotics should be given via the CVAD.


**Misplacement of catheter tip**

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Ideal Catheter tip position: should be at the lower one-third of the Superior Vena Cava (SVC) at the level of the cavoatrial junction (INS 2016).

Symptoms
- More of the catheter showing externally, inability to draw back blood, added sounds in ear, arrhythmias (INS 2011).

Treatment
- Chest X-ray
- Seek support from Procedure team/Radiologists
- Misplaced catheters are at increased risk of thrombosis, arrange ultrasound Doppler if the patient symptomatic of thrombosis.

Prevention
- Optimal patient position during placement
- Tip placement lower one third of SVC/CAJ (INS 2016)
- Use catheter securement device
- PICC is are most prone to this - *Take meticulous care not to pull PICC out when changing dressings.*

Tip position
Should be at the lower one-third of the Superior Vena Cava (SVC) at the level of the cavoatrial junction (CAJ) (INS 2016), optimal position reduces risk of thrombosis.

Symptoms – Discomfort, pain, oedema, engorged veins, enhanced venous pattern; extremity, shoulder neck or chest (INS 2011). Leakage of fluid at the insertion site, discolouration of extremity, unexplained pyrexia.

Treatment – Inform medical team/Radiologist to undertake ultrasound Doppler.

Do not remove the catheter CVAD in the presence of a CVAD-associated vein thrombosis when the catheter is correctly positioned at the cavo-atrial junction, the catheter is functioning with a blood return, there is no evidence of any infection, and there is absence of severe deep vein thrombosis (DVT) - related symptoms causing pain Prevention (INS 2016) This is ultimately the medical team’s decision based upon, the need to keep the device in place and balanced individual hypercoagulability factors of the patient. Monitoring and anticoagulation should be followed in accordance with the Anticoagulant Therapy Management Policy.

Use ultrasound pre-assessment to assess optimal vessel size. Ideal vessel should Systemic anticoagulation with or without CVAD removal (INS 2011).

Prevention – Use the device with the smallest gauge (INS 2011)
The diameter of the catheter should be one third or less of the diameter of the vein, as checked by ultrasound (Espen 2009). If the vein is smaller than ideal the risk of thrombosis should be balanced against the urgency of the CVAD. Assess patient risk factors i.e. malignancy, critically ill (Chopra et al Lancet 2013) Catheter tip position, RCN (2010). Modified Seldinger technique reduces trauma.

Extravasation and CVADs
See policy for Management of Extravasation.

Catheter Fracture
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If the catheter is damaged, clamp above the damaged area to avoid blood loss and air emboli and contact the procedure team (08:00-17:00) immediately, or the hotline out of hours.

**Catheter Occlusion**
Catheter Occlusion, causes include;
- **Mechanical causes** (kinking, pinching of line)
- **Chemical**, infusates caused precipitate
- **Blood occlusion** (Proteins and Platelets)

In all cases the occlusion maybe partial or total. **Partial;** flush but not aspirate or **Total;** cannot flush or withdraw.

**Treatment** –
- Eliminate mechanical causes (kinking, pinching of line)
- If chemical, determine which infusates caused precipitates and instill proper solution (alkaline/acidic) to clear (INS 2011)
- Most commonly blood related occlusion, instill Urokinase per protocol Appendix 6.

**Prevention** -
- Use a firm pulsatile flush before and after all infusions and devices use.
- Use 20mls flush following blood transfusion, contrast media, TPN or any viscous infusants
- Be aware of solution incompatibilities prior to infusion

For persistent withdrawal occlusion see Trouble shooting algorithms for the device being use (Appendix 3, 4, 5, 9, Urokinase protocol Appendix 6).

**Effective pulsatile flushing of vascular devices is one of the most important factors in preventing occlusion related problems and must be adopted by all staff caring for CVADs.**

**For further information please see;**
- TIVAD occlusion algorithm, Appendix 3
- PICC occlusion algorithm, Appendix 4
- Tunneled CVC occlusion algorithm, Appendix 5
- Urokinase protocol Appendix 6

**5.18 Anti-infective strategies for CVADs**
Strategies maybe considered for patients;
- at high risk of infection
- or have experienced a CVAD related infection and require a replacement vascular device
- or when treating an infected device.

**Alcohol lock** – For patient who have had recurrent infection, who are high risk of infection or when line removal is not possible alcohol 70% injection may be used to try to decontaminate lines. Absolute alcohol injection may be diluted to 70% with water for Injection and used to lock an infected lumen for up to 7 days. Please contact pharmacy and/or microbiologists for advice. (See Appendix 7).
Alcohol lock is not compatible with many polyurethane catheters and may degrade the composition catheter i.e. open ended clamped PICCs, PFM® implanted Ports (TIVADs), please check with manufacturer guidance or seek advice from the procedure team 0161-446-3916. Vygon Broviac Lifecath catheters are compatible with alcohol lock.

Other anti-infective strategies currently being considered (subject to Trust & business case approvals).

**Taurolock®** - To be administered as a lock solution, this solution is compatible with all CVADs (Subject to Safe Medicines Committee/Medical device committee approval).

**Coated catheters** - may be considered if a patient is at higher risk of infection or who have had CVAD related infection and require a replacement vascular device. Examples of impregnated, coated catheters are those with Endexo technology® (Angiodynamics, Queensbury, N.Y, U.S) Chloragard® (Teleflex, Morrisville, NC 27560). Please note all medical device are subject to proper approvals via the medical devices committee and must follow Trust procurement process.

"Use an antimicrobial-impregnated central venous access device for adult patients those central venous catheter is expected to remain in place for >5 days if catheter-related bloodstream infection rates remain above the locally agreed benchmark, despite the implementation of a comprehensive strategy to reduce catheter-related bloodstream infection".

EPIC III, Class A, Recommendation (Loveday 2014).

**Chlorhexidine dressing** –
"Consider the use of a chlorhexidine-impregnated sponge dressing in adult patients with a central venous catheter as a strategy to reduce catheter-related bloodstream infection".

Class B Revised Recommendation(Loveday 2014).

**Biopatch®** - may be considered if a patient is at higher risk of infection or who have had CVAD related infection and require a replacement vascular device.

"Consider the use of a chlorhexidine impregnated sponge dressing in adult patients with a central venous catheter as a strategy to reduce catheter related bloodstream infection".

EPIC III, Class B, Recommendation (Loveday 2014).

**Chlorhexidine Gluconate – Sensitivity**
Chlorhexidine is a potential allergenic antiseptic that is present in many products and is widely used in health care for skin antisepsis, insertion of urinary catheters or coating CVCs.406 In susceptible individuals, initial contact will cause a minor hypersensitivity reaction that, although not severe, should not go undocumented as subsequent exposures to Chlorhexidine may lead to anaphylaxis.423,424 The Medicines and Healthcare Products Regulatory Agency has alerted all healthcare providers in the UK to the risk of Chlorhexidine allergy425 and requires them to have systems in place that ensure:

- Awareness of the potential for an anaphylactic reaction to Chlorhexidine;
- Known allergies are recorded in patient notes;
- Labels and instructions for use are checked to establish
- if products contain chlorhexidine prior to use on patients
- with a known allergy;
- If a patient experiences an unexplained reaction, checks
- are carried out to identify whether chlorhexidine was used

- or was impregnated in a medical device that was used; and
- Reporting of allergic reactions to products containing
  chlorhexidine to the Medicines and Healthcare Products
  Regulatory Agency. EPIC III.

**Saline vs Heparin CVAD flush/locking.**
Central venous access devices should only be flushed routinely with 0.9% sterile saline. Heparinised saline and higher strength heparin locks maybe beneficial for devices (Appendix 15).

6. CONSULTATION PROCESS
The policy has been developed in collaboration with the Clinical Skills Team, Chemotherapy Team, Lead Nurses and clinical colleagues specialising in the vascular access. It has been updated into the standardised policy template. Overall approval of clinical content has been given by the Patient Safety Committee and ratified by the document ratification committee.

7. DISSEMINATION, IMPLEMENTATION & TRAINING

7.1 Dissemination
- Once ratified the procedural document will be sent to the web team who will replace the historical version which will subsequently be archived. It will be place the document intranet database and on The Christie external website.
- Dissemination of the policy will be via the intranet, nursing structure Matrons, ward managers and via relevant clinical committees with in The Christie and clinical skills team. Peripheral sites will be emailed directly, including chemotherapy teams and AONS, the policy will be forwarded to community nurse liaison to assist with informing community nursing colleagues
- The governance team will be notified of the policy ratification which will enable update of the procedural document register.
- Awareness will be raised clinically via an email to all clinicians, departmental managers and at on-going essential training sessions to The Christie and peripheral site.

7.2 Implementation
This policy will be effective from the date of ratification.

7.3 Training/Awareness
All permanent clinical staff training to use central venous access devices must complete the relevant workbook utilising the Clinical Skills team or Link Staff for advice. All staff must be signed off as competent. For further information refer to the Clinical skills web page.

| Insertion of CVC | Procedure team only, appropriately trained registered healthcare professionals | According to training schedule and assessment by competent, experienced practitioner i.e Consultant |

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8. PROCESS FOR MONITORING EFFECTIVE IMPLEMENTATION

The effectiveness of this policy will be monitored by the Clinical Skills Team relating to care and maintenance through IV training and updates.

<table>
<thead>
<tr>
<th>Standard to be monitored</th>
<th>Process for monitoring</th>
<th>Frequency</th>
<th>Person responsible for:</th>
<th>Committee accountable for:</th>
<th>Frequency of monitoring</th>
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<tr>
<td>Training of care and maintenance of all CVADs</td>
<td>On-going</td>
<td>Yearly (CVAD skills should be assessed 3 yearly and annually for implanted ports)</td>
<td>Clinical skills Training Team</td>
<td>Patient Safety Committee</td>
<td>Annual</td>
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<tr>
<td>Compliance with care and maintenance standards set out in CVAD and Christie policies</td>
<td>Audit Datix</td>
<td>Annual</td>
<td>Procedure team Manager</td>
<td>Patient Safety Committee</td>
<td>Annual</td>
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9. REFERENCES (IF APPLICABLE)

2) Aseptic non-touch technique, (ANTT 2010), The Christie Hospital NHS foundation trust.
3) Bard Limited Forest House, Tilgate, Forest Business Park, Brighton Rd, Crawley, West Sussex, England RH11-9BP WWW.Bardaccess.com
4) BCSH Guidelines for the Insertion and Management of Central Venous Access Devices (2006 page 8).
12) Dawson RB (2011) PICC zone insertional method: a systematic approach to determine the ideal insertion site for PICCs in the upper arm. JAVA 16 (3) 156-165
17) G. A. Goossens,M. Jérôme1, C. Janssens1, W. E. Peetermans, S. Fieuws,


33) Lichtenstein, Daniel A. MD; Mezière, Gilbert MD; Lascols, Nathalie MD; Biderman, Philippe MD; Courret, Jean-Paul MD; Gepner, Agnès MD; Goldstein, Ivan MD; Tenoudji-Cohen, Marc MD. Ultrasound diagnosis of occult pneumothorax. Critical Care Medicine: June 2005 - Volume 33 - Issue 6 - pp 1231-1238 doi: 10.1097/01.CCM.0000164542.86954.B4


40) ‘Matching Michigan’, www.BMJQua Safe.com, September 2012 (last viewed 26.08.14)


51) PFM medical Uk Limited, Poynton, Cheshire, England


57) Safe use and disposal of sharps policy (2009). The Christie NHS foundation trust


60) Mark Gorman, Jennifer Ligibel, Paul Mansfield, and Mark Levine


64) Stephens LC (1997) et al Normal saline versus heparin flush for maintaining central venous catheter patency during apheresis collection of peripheral blood stem cells (PBSC).


70) UKCC (1992) The scope of professional Practice. UKCC. London
The previous policy that pertained to central venous access devices was the; Greater Manchester and Cheshire Cancer Network Policy for the Care of long term Central Venous Catheters. The CVAD policy covers the content of the older Network policy and provides additional information relating to CVADs.

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<td>07/11/2016</td>
<td>Steve Hill</td>
<td>Live</td>
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11. APPENDICES

Appendix 1

Training for nurse led insertion of vascular access device must include:
- Anatomy
- Vein selection
- Potential insertion site selection
- Relevant guidelines
- Vein health preservation; Right device, right time, right patient
- Types of infusants; Osmolarity, pH and which types require a central venous access device.
- Types of devices; associated advantages and disadvantages
- Indications and contraindications
- Informed consent
- Hand hygiene max barrier precautions
- About and how to used ultrasound
- Modified Seldinger technique
- Insertion techniques
- Hands on training with simulation
- Tip position
- Care and maintenance

Complication;
- Signs, symptoms and management i.e infection (exit site or intraluminal/systemic), thrombosis, fracture, emboli, erosion, persistent withdrawal occlusion

Evidence based practice
IRMER; safe radiation practice, working safely whilst in fluoroscopy
Complete comprehensive training programme, assessed by experienced competent practitioner.

Appendix 2

<table>
<thead>
<tr>
<th>Name</th>
<th>DOB</th>
<th>Hosp No.</th>
</tr>
</thead>
<tbody>
<tr>
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<td>(Attach addressograph)</td>
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Affix catheter ID sticker here

<table>
<thead>
<tr>
<th>Has the patient confirmed his/her identity?</th>
<th>YES/NO</th>
</tr>
</thead>
<tbody>
<tr>
<td>Has the patient any allergies</td>
<td>YES/NO</td>
</tr>
<tr>
<td>Allergy to contrast dye</td>
<td>YES/NO</td>
</tr>
<tr>
<td>Risk, Benefits, alternatives discussed Re; CVAD, consent obtained</td>
<td>YES/NO</td>
</tr>
<tr>
<td>Is the patient taking anticoagulants</td>
<td>YES/NO</td>
</tr>
<tr>
<td>Full blood count and clotting screen within protocol</td>
<td>YES/NO</td>
</tr>
<tr>
<td>MRSA Screen within 2 weeks of procedure</td>
<td>YES/NO</td>
</tr>
<tr>
<td>If applicable, has the most recent CT scan (if applicable) been reviewed</td>
<td>YES/NO</td>
</tr>
<tr>
<td>Do you bleed abnormally or have a bleeding disorder</td>
<td>YES/NO</td>
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</table>
Appendix 3

TIVAD Occlusion Algorithm 2016.doc

Appendix 4

PICC Algorithm (Hospital) 2016.doc

Appendix 5

Tunneled CVC algorithm (Hospital) 2

Appendix 6

Urokinase Administration protocol.doc

Appendix 7

Alcohol lock 2016.doc

Appendix 8
Guidelines for the management of sepsis (including Neutropenic sepsis)
Discover - Documents and guidelines database

Appendix 9

Procedure for unblocking a CVAD (3-Way tap method)

Procedure for unblocking CVAD.doc

Appendix 10

Blood sampling from a picc.doc

Appendix 11

Blood sampling from a CVC.doc
Appendix 12
TIVAD care, community guide

Appendix 13
PICC occlusion algorithm - Community

Appendix 14
Tunneled CVC occlusion algorithm - Community

Appendix 15
Heparin vs Saline Lit 2016.doc

Appendix 16
Patient information implanted port 2016.doc

Appendix 17
Referral SOP CVAD 2016.docx

Appendix 18
CVAD REQUEST 2016.doc