CHEMOSAT Hepatic Delivery System

For Melphalan Hydrochloride for Injection

Instructions for Use

EN

EU DOCUMENT ONLY

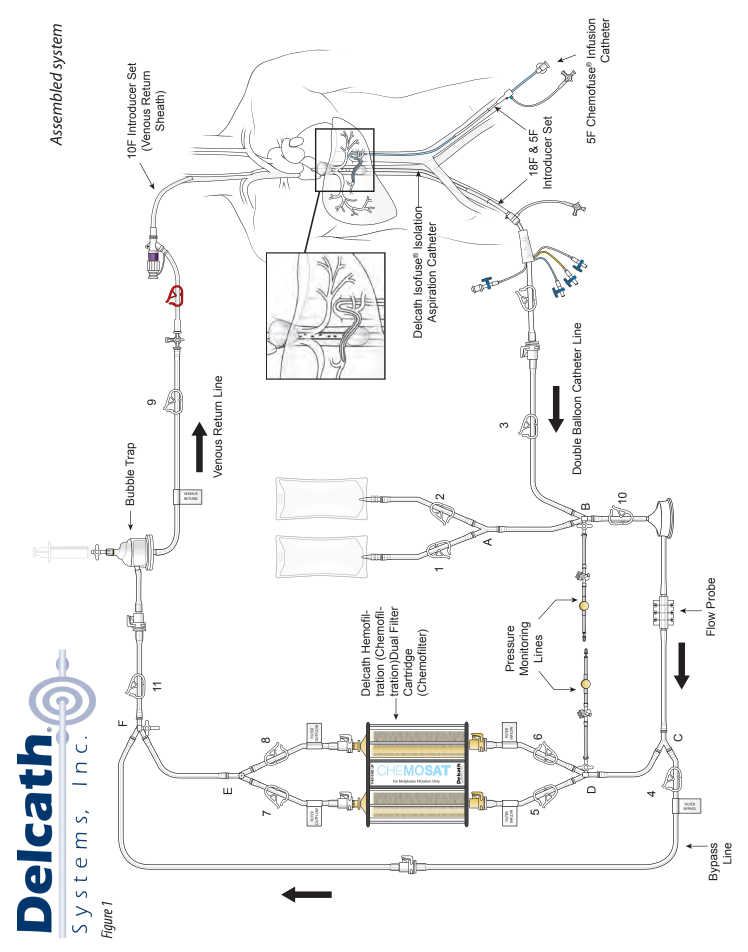


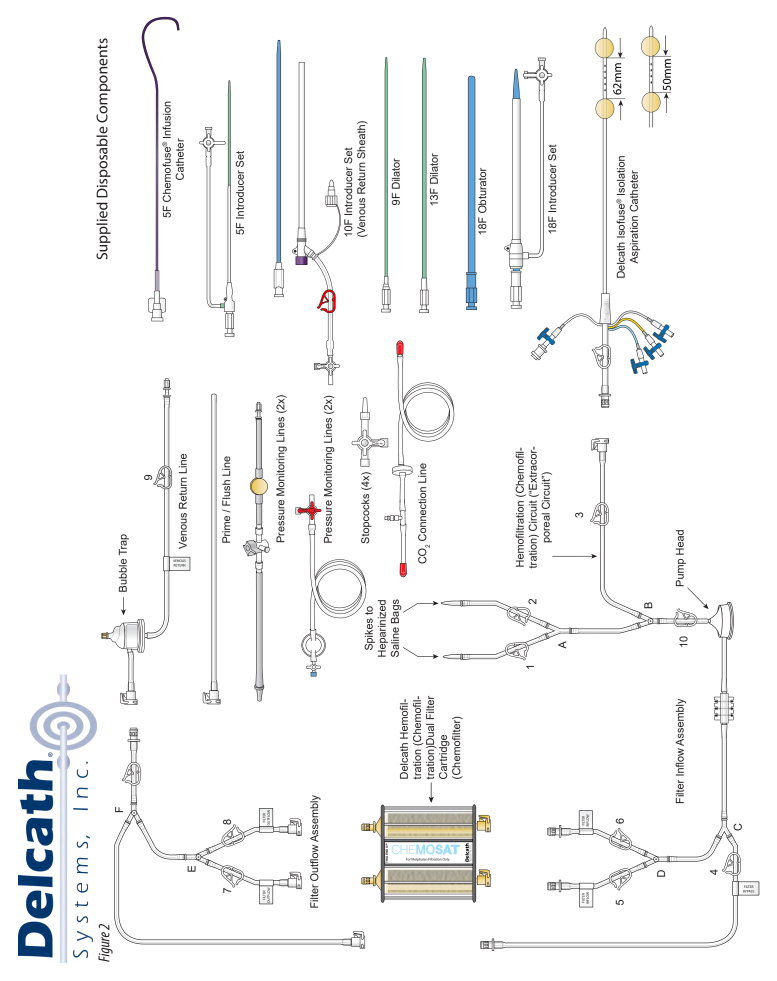
Concentrating the Power of Chemotherapy 120057 C 08-Mar-2021

Systems, Inc.

TABLE OF CONTENTS

ASSEMBLED SYSTEM - FIGURE 1
SUPPLIED DISPOSABLE COMPONENTS - FIGURE 2
DESCRIPTION OF SYSTEM COMPONENTS
INDICATIONS FOR USE
RECOMMENDED DOSE OF MELPHALAN HYDROCHLORIDE
MELPHALAN HYDROCHLORIDE DOSE MANAGEMENT
CONTRAINDICATIONS
WARNINGS
PRECAUTIONS
ADVERSE EVENTS AND COMPLICATIONS
LOCATION OF PROCEDURE
PROCEDURAL TEAM
PREPARATION: PRIOR TO TREATMENT
PREPARING AND PRIMING THE HEMOFILTRATION CIRCUIT
PLACING THE CATHETERS
ESTABLISHING ANTICOAGULATION & PLACING ISOFUSE® ISOLATION ASPIRATION CATHETER
CONNECTING CATHETERS TO HEMOFILTRATION CIRCUIT
ISOLATING THE INFERIOR VENA CAVA
BRINGING HEMOFILTRATION CARTRIDGES ON LINE
SETUP DRUG DELIVERY SYSTEM AND START EXTRACORPOREAL FILTRATION
ENDING EXTRACORPOREAL CIRCULATION
CATHETER REMOVAL
NORMALIZATION OF COAGULATION STATUS FOR SHEATH REMOVAL
PROCEDURE FLOWCHART - FIGURE 29
LIMITED WARRANTY





DELCATH HEPATIC CHEMOSAT® DELIVERY SYSTEM

COMPLETE REQUIRED TRAINING BEFORE USING THIS DEVICE FOR THE FIRST TIME. ENSURE YOU COMPLETELY READ AND UNDERSTAND THE INSTRUCTIONS FOR USE.

DESCRIPTION OF SYSTEM COMPONENTS

The Delcath Hepatic CHEMOSAT® Delivery System (CHEMOSAT® System) consists of a closed circuit of catheters and drug specific filters utilized to deliver a chemotherapeutic agent (melphalan hydrochloride) to the (hepatic) artery of the liver and to lower the concentration of the agent in the blood before it is returned to systemic circulation. A schematic overview of how the Delcath CHEMOSAT® System components work together is presented in Figure 1: Assembled System. The system is designed to be used with a Medtronic Bio-Console® 560 and TX50P Flow Transducer.

 Delcath Isofuse[®] Isolation Aspiration Catheter -- 16F (shaft) polyurethane double balloon catheter that is placed in the retro-hepatic inferior vena cava to isolate the hepatic venous blood and transport it to the extracorporeal Hemofiltration Circuit for filtration. The catheter has one large (central) drainage lumen and four accessory ports. Due to variation in the length of the retro-hepatic segment of the inferior vena cava and relative positions of hepatic and renal veins, the lsofuse[®] Catheter is available in two different balloon configurations: 50 mm or 62 mm between the two balloons.

Using pre-operative computed tomography (CT) imaging, or by performing an inferior vena cavogram prior to placement of the lsofuse[®] Catheter, estimate the length of the retro-hepatic segment of the inferior vena cava and the relative positions of hepatic and renal veins in order to determine the optimum lsofuse[®] Catheter balloon spacing: 50mm or 62mm.

Two (2) of the accessory ports are used to inflate low-pressure occlusion balloons, which are inflated independently to occlude the inferior vena cava above and below the hepatic veins. When inflated, the cephalic (superior – blue port) balloon obstructs the inferior vena cava above the hepatic veins and the caudal (inferior – yellow port) balloon obstructs the inferior vena cava below the hepatic veins, thus isolating hepatic venous blood in the fenestrated segment between the balloons.

The large drainage lumen with a quick connect fitting is a conduit to the fenestrations between the two occlusion-balloons. The fenestrations allow the hepatic venous blood to flow into the drainage lumen and exit the catheter at the proximal end.

The third accessory (translucent) port labeled "CONTRAST" is for injections of iodinated contrast medium through the fenestrations, to check catheter position.

The fourth accessory port (white) is used for over-the-guidewire (OTW) introduction and positioning of the catheter in the retro-hepatic inferior vena cava. This lumen also has a small port opening along the catheter shaft positioned inferior to the caudal balloon and exits at the distal tip, to allow inferior vena cava blood, proximal to the caudal balloon, to bypass the occluded segment of the inferior vena cava and flow into the right atrium.

2. Accessory Pack

- 9F and 13F Dilator Set --These over-the-wire dilators are used to widen the subcutaneous space and venous entry site in preparation for the placement of the 18F Introducer Set.
- 18F Introducer Set (Sheath and Dilator) -- The 18F introducer sheath and coaxial dilator are to be placed over a wire; the dilator is removed and the sheath is available for the insertion of the lsofuse[®] Catheter or the 18F Obturator.

 18F Obturator -- An 18F obturator is used to occlude and support the 18F sheath lumen when it is not in use, and upon removal of the lsofuse[®] Catheter at the end of the procedure.

Delcath

Systems,

- 5F Introducer Set (Sheath & Dilator) -- A 5F hemostasis sheath is used to facilitate the introduction of the 5F Chemofuse[®] Chemo Delivery Catheter through the femoral artery.
- 10F Introducer Set (Venous Return Sheath) -- A 10F sheath used to return the filtered hepatic venous blood through the internal jugular vein. A 3-way high-flow stopcock is included as part of the 10F Introducer Set. The highflow stopcock is attached to the Venous Return Sheath and then to the male connector of the Hemofiltration Circuit, if required. This sheath may also be used for hydration. A 10F obturator is included for insertion through the valve post placement.
- 3. **5F Chemofuse**[®] **Infusion Catheter** -- 5F arterial catheter is used to deliver chemotherapeutic agent (melphalan hydrochloride) into the proper hepatic artery or, it can be used to coaxially introduce a microcatheter (not provided by Delcath), if, at the discretion of the Interventional Radiologist, a microcatheter is preferred for selective catheter tip placement for the drug infusion. The following microcatheters have been qualified for use with the CHEMOSAT[®] System select one from the microcatheters below. See microcathether manufacturer's Instructions for Use. These microcatheters are NOT PROVIDED by Delcath:
 - Merit Maestro (Merit Medical Systems, Inc., So. Jordan, UT, USA)
 - Boston Scientific Renegade Hi-Flo (Boston Scientific Corp., Natick, MA, USA)
 - Terumo Progreat (Terumo Medical Corp., Somerset, NJ, USA)
- 4. **Delcath Hemofiltration (Chemofiltration)Dual Filter Cartridge** (**Chemofilter)** -- One single-use Dual Filter Cartridge designed with the filter cartridges arranged in parallel to lower the concentration of chemotherapeutic agent (melphalan hydrochloride) from the blood. The cartridge frame comes with a built-in pole clamp.
- 5. Hemofiltration (Chemofiltration) Circuit ("Extracorporeal Circuit") -- The Hemofiltration (Chemofiltration) Circuit is used to transport the hepatic venous blood, which has been isolated by the Isofuse® Catheter and aspirated into the fenestration lumen, through the Hemofiltration (Chemofiltration) Cartridges and back to the patient through the Venous Return Sheath. Connections are provided for infusion of normal saline. This circuit includes:
 - Medtronic BP-50 Bio-Pump[®] Centrifugal Pump ("Pump Head"), a disposable pump head to be used with a pump console manufactured by Medtronic, Inc. - see manufacturer's Instructions for Use for Pump Head. (Note: The Medtronic, Inc. Bio-Console 560 extracorporeal blood pumping system is required for use with the CHEMOSAT[®] System: This is NOT PROVIDED by Delcath).
 - Medtronic Bio-Probe® DP-38P blood flow monitoring insert ("Flow Probe"), a disposable flow probe to be used with a blood flow monitoring transducer manufactured by Medtronic, Inc. - see manufacturer's Instructions for Use for Flow Probe. The Flow Probe is used to measure the rate of blood flow during the procedure. (Note: The Medtronic Bio-Probe TX50P blood flow monitoring transducer is required for use with the CHEMOSAT® System: This is NOT PROVIDED by Delcath).
- 6. **Carbon Dioxide (CO₂) Connection Line** -- The CO₂ Connection Line is used to deliver sterile CO₂ gas to the Hemofiltration Cartridges to aid in priming/ debubbling the filter cartridge, prior to the start of the procedure. The CO₂ Line has no patient contact.



WARNING

Only the Components Provided in the CHEMOSAT® System Or Specified By Delcath In The "NOT INCLUDED" Box Below Are to Be Used to Create the Circuit. There Should Be No Substitutions. The Circuit Has Not Been Validated For Use With Other Components.

NOT INCLUDED:

- Chemotherapeutic Agent (Melphalan Hydrochloride)
- Bubble Trap holder
- Medtronic Bio-Console 560 ("Pump")
- Medtronic Bio-Probe TX50P ("Flow Transducer")
- CO, Supply for Priming Dual Filter
- Drug Injector: must be able to inject at a rate of 25 mL/minute
- Drug Delivery Disposables:
 - One (1) Medrad 150mL Syringe (Polypropylene (PP)-Barrel & Polyisoprene-Plunger) or equivalent
 - Two (2) Intravenous Administration Set with spike & drip chamber (Polyvinylchloride (PVC)-tubing, Acrylonitrile butadiene styrene (ABS) & Polyethylene (PE)-Drip Chamber & Polycarbonate (PC)-Luer) or equivalent
 - One (1) 48" injector lines (PVC-Tubing & PC-Luer) or equivalent
 - Five (5) 3-way stopcocks (PC-body, High Density Polyethylene (HDPE) or Acetal-Handles) or equivalent
 - Three (3) 20 mL syringes (PP-Barrel & Polyisoprene-Plunger) or equivalent
- Microcatheters (Maximal Distal End OD = 2.8F) for Selective Drug Infusion (at Interventional Radiologist discretion). Select one from Delcath qualified microcatheters listed below:
 - Merit Maestro (Merit Medical Systems, Inc., So. Jordan, UT, USA)
 - BSC Renegade Hi-Flo (Boston-Scientific Corp.; Natick, MA, USA)
 - Terumo Progreat (Terumo Medical Corp., Somerset, NJ, USA)

INDICATIONS FOR USE

The Delcath Hepatic CHEMOSAT[®] Delivery System is used for percutaneous intra-arterial administration of chemotherapeutic agent (melphalan hydrochloride) for perfusion of the liver with subsequent extracorporeal filtration of the regional (hepatic) venous blood, lowering the concentration of chemotherapeutic agent in the blood before returning it to the systemic venous circulation.

RECOMMENDED DOSE OF MELPHALAN HYDROCHLORIDE

The use of melphalan to treat unresectable primary or metastatic hepatic tumors represents an unlicensed indication.

The recommended dose is 3.0 mg/kg based on ideal body weight, infused over 30 minutes, with a maximum absolute dose of 220 mg during a single treatment. A dose escalation study performed by Delcath supports the safe use of melphalan at 3.0 mg/kg, as does the published literature.

The maximum absolute dose of 220 mg melphalan during a single treatment should not be exceeded.

MELPHALAN HYDROCHLORIDE DOSE MANAGEMENT

Evidence from the literature indicates that patients with unresectable primary or metastatic hepatic tumors receive between 1-8 treatments and possibly more, with CHEMOSAT®/Melphalan (up to 220 mg melphalan/treatment), depending on the patient's clinical and radiologic response.

There should be at least a 5-6 week gap after treatment with CHEMOSAT® before another treatment is considered. Delays in treatment until recovery from toxicities and a dosage reduction to 2.5 mg/kg, or 2.0 mg/kg IBW, are permitted.

Toxicities should have resolved to grade 2 or less, with the exception of hepatic toxicity due to underlying disease which should resolve to baseline, before additional treatments are considered. A dosage reduction to 2.0 mg/kg IBW should be considered for subsequent treatments for the following reasons:

- Grade 4 neutropenia of >5 days duration despite growth factor support or associated with neutropenic fever
- Grade 4 thrombocytopenia of >5 days duration or associated with a hemorrhage that required a transfusion
- Grade 4 anemia of >48 hours duration
- Grade 3 or 4 major non-hematologic organ toxicity not corrected within 24 hours of the
 procedure (excluding fever, nausea, and weight gain); for hepatic toxicity, grade 4 bilirubin
 of any duration, and doubling of liver function test values (AST, ALT, and total bilirubin)
 above the baseline value

CHEMOSAT[®]/Melphalan treatment should be permanently discontinued if patients have persistent toxicity that has not resolved to grade 2, or less, by 8 weeks following treatment.

CONTRAINDICATIONS

- Liver failure or portal hypertension.
- Active intracranial metastases or brain lesions with a propensity to bleed as determined by imaging.
- History of allergies or known hypersensitivity to a component, material, or drug utilized with the CHEMOSAT[®] System including:
 - Heparin
 - Angiographic contrast medium
 - Natural rubber latex component of the product
 - Melphalan hydrochloride

WARNINGS

PLEASE CAREFULLY READ AND UNDERSTAND THE LIST OF WARNINGS BELOW AS SERIOUS INJURY, ILLNESS OR DEATH OF THE PATIENT CAN OCCUR IF THESE WARNINGS ARE NOT PROPERLY FOLLOWED

WARNINGS RELATING TO THE USE OF MELPHALAN HYDROCHLORIDE

- It is the doctor's responsibility to give due consideration to the details of the melphalan marketing authorization in deciding whether treatment with melphalan is appropriate for the patient in his/her care. The melphalan Summary of Product Characteristics (SmPC) must be consulted.
- The dose of melphalan should be assessed according to the individual patient's weight and clinical circumstances.
- Dual cartridge filter efficiency studies demonstrate that the level of melphalan in blood returned to the patient post-filtration is reduced by approximately 95%. Systemic exposure of melphalan can occur. Related side effects that might occur are listed in the melphalan SmPC.



 Due to the known cytotoxic, mutagenic, embryotoxic (and possible teratogenic) properties, melphalan should not be used during pregnancy, women should not breastfeed, and women/men of childbearing potential should use adequate contraceptive measures, as outlined in the SmPC.

WARNINGS RELATING TO THE USE OF CHEMOSAT®

- The CHEMOSAT[®] System must not be used in patients who have undergone surgery or medical treatment of the liver within 4 weeks.
- Screen for prior surgeries that could potentially affect normal hepatic biliary/ vascular anatomy (e.g., Whipples Procedure). If the common bile duct is reimplanted, there is an increased risk of biliary tree infection. If the vascular anatomy (especially hepatic arterial supply) is affected, there could be an increased risk of mis-infusion and reflux of chemotherapeutic agent.
- For patients with 50% or greater tumor burden by medical imaging, a biopsy of the non-involved parenchyma must be performed to show that it is histologically normal. Clinical judgment and biopsy results should be used to guide the decision on whether to perform the procedure.
- Patients on chronic anti-coagulation therapy (e.g., Coumadin), the therapy must be discontinued and switched to short- acting agent for the ease of reversal. Patients should avoid drugs effecting platelet functions such as aspirin, or non steroidal anti-inflammatory drugs (NSAID), starting one week before the procedure. Anti-coagulation therapy may be resumed post-operatively, as indicated, once hemostasis has been restored and no bleeding complication is observed.
- Patients with a history of hypertension receiving angiotensin converting enzyme inhibitors or calcium channel blockers must temporarily discontinue these drugs at least five drug half-lives before the procedure. Short-acting antihypertensive regimen may be used to manage hypertension as needed. After the procedure, previous antihypertensive regimen may be restored under guidance of treating physicians.
- When the catheter is placed intravascularly, it must only be manipulated under fluoroscopic observation. Do not advance or retract the lsofuse[®] Catheter unless both balloons are fully deflated, except when positioning the inflated cephalad balloon at the junction of the inferior vena cava and right atrium (with the caudal balloon fully deflated). If resistance is met during manipulation, determine the cause of the resistance before proceeding.
- When the 3-way high flow stopcock is used to attach the Venous Return Sheath to the male connector of the Hemofiltration Circuit, ensure that the stopcock is fully open to minimize back pressure and maximize flow through the stopcock. ("OFF" handle of stopcock turned 90° to flow path.)
- Use caution to avoid air embolism while performing this procedure. Never use air or any gaseous medium to inflate the lsofuse[®] Catheter balloons.
- Contents are supplied STERILE. Inspect prior to use to verify that no damage has occurred during shipping. Do not use if sterile barrier is damaged. If damage is found, please contact Delcath Customer Service.
- All components are for single patient use only. Do not reuse, reprocess or
 resterilize. Reuse, reprocessing or resterilization may compromise the structural
 integrity of the device and/or lead to device failure which, in turn, may result
 in patient injury, illness, or death. Reuse, reprocessing, or resterilization may
 also create a risk of contamination of the device and/or cause patient infection
 or cross-infection, including, but not limited to, the transmission of infectious
 disease(s) from one patient to another. Contamination of the device may lead to
 injury, illness, or death of the patient.

PRECAUTIONS

PRECAUTIONS RELATING TO THE USE OF MELPHALAN HYDROCHLORIDE

- See melphalan SmPC for contraindications regarding use.
- See melphalan SmPC for side effects related to the drug therapy.
- Melphalan should be used immediately after reconstitution. Refer to section 6.6 of the melphalan SmPC ("Special precautions for disposal and other handling").

PRECAUTIONS RELATING TO THE USE OF CHEMOSAT®

- The CHEMOSAT[®] System is for use only by qualified personnel properly trained and familiar with the procedure in accordance with the Instructions for Use.
- Caution: The CHEMOSAT[®] System contains natural rubber latex which may cause allergic reactions.
- Women who are pre-menopausal (have had a period within the last 12 months) must receive appropriate hormonal suppression to prevent potential bleeding as a result of the procedure.
- The safety and effectiveness of the use of The CHEMOSAT® System on pregnant women and pediatric patients has not been established.
- Patients should be evaluated for gastric acid hypersecretion (e.g., untreated gastrinoma) and clinically managed as indicated.
- In order to avoid mis-infusion of the drug into gastrointestinal branches
 originating from the hepatic artery, careful angiographic examination is required,
 followed by embolization as indicated. During infusion the catheter tip must be
 placed distal to the origins of any gastrointestinal branches that may not have
 been embolized.

WARNING

If The Perfusion Of Melphalan Cannot Be Isolated From The Systemic Circulation, Stop The Drug Infusion Immediately.

- Adequate anticoagulation is required throughout the procedure to prevent intravascular thrombosis and to assure unimpeded flow through the extracorporeal circuit and filters. (See Coagulation Control below).
- During the procedure, the arterial blood pressure must be closely monitored as there will be significant decrease of blood pressure related to:
 - o Initial occlusion of the inferior vena cava by the balloons
 - o The filters coming on line in the extracorporeal circuit

Prompt intervention must be administered as described below under Blood Pressure Control.

- Hepatic artery spasm may occur and potentially cause reflux of the infusate into more proximal non-embolized gastrointestinal branches. In order to avoid such reflux, the hepatic artery must periodically be checked angiographically for spasm. To relieve hepatic artery spasm, which occurs infrequently, use a 50-100 mcg/injection of intra-arterial nitroglycerin locally to relieve spasm. If spasm does not resolve, the procedure must be stopped.
- The following clinical effects have been observed during the procedure:
 - Decreased platelet counts occur in a majority of patients. Transfusion is required as clinically indicated. The CHEMOSAT[®] System must not be administered in patients with platelets <75,000 cells/mm³.
 - o Decreased hemoglobin occurs in a majority of patients. Transfusion is required as clinically indicated. The CHEMOSAT[®] System should not be used on patients with hemoglobin \leq 10 g/dL.



O IR PF AN PH DD IN

- Prolongation of activated partial thromboplastin time is an intended effect of heparin administration during the procedure; however, since removal of platelets and clotting factors by the filters can increase the risk of bleeding, anticoagulation must be reversed promptly by the administration of protamine sulfate, Fresh Frozen Plasma and cryoprecipitate should be administered as clinically indicated to correct remaining coagulopathy. The CHEMOSAT[®] System must not be used on patients with uncorrectable coagulopathies.
- o Decreased serum albumin occurs frequently. Specific corrective intervention is seldom required.
- Decreased blood calcium (hypocalcemia) occurs frequently. As with other less frequent electrolyte disturbances, replacement is required when clinically indicated.
- o Elevated hepatic transaminases (alanine aminotransferase, aspartate aminotransferase), with or without hyperbilirubinemia, occur. They are self-limiting. No intervention is required.

ADVERSE EVENTS AND COMPLICATIONS

In a Phase 3 Ocular and Cutaneous Melanoma Trial, clinically significant (Grade 3-4) adverse events that were observed in >35% of patients in descending order of frequency for the CHEMOSAT® System that occurred within the first 72 hours after the procedure were the following:

Platelet count decreased (69 %), Hemoglobin decreased (60%), Blood Albumin decreased (37.1%), aPTT prolonged (31.0%), Blood Calcium decreased (21.4%), AST increased (20%), INR increased (20%), Lymphocyte decreased (12.9%), ALT increased (8.6%), Blood Bilirubin Increased (10 %), Blood Potassium decreased (10 %), Troponin increased (7.1%).

LOCATION OF PROCEDURE

The procedure must be performed in an appropriately equipped interventional radiology suite with fluoroscopy or an operating room designed and equipped similarly. Resuscitation personnel, equipment, and medications must be immediately available.

PROCEDURAL TEAM

The required procedural team members should be determined by the facility based on institutional experience and clinical judgment. The operating team performing the procedure must include, at a minimum:

- A qualified medical/surgical oncologist experienced in the monitoring of toxicities of chemotherapy and who is responsible for the complete medical management of the patient, including, but not limited to, pre- and postoperative care. The medical/surgical oncologist may also be responsible for monitoring the patient during the immediate post-procedure period.
- O The medical/surgical oncologist will also play a unique role in communicating about chemotherapeutic agent (melphalan hydrochloride) and the CHEMOSAT[®] System risks and coordinating with other oncologists and key health care professionals responsible for patient follow-up care and monitoring for post-procedure toxicities.
- IR A qualified interventional radiologist with the knowledge, skills, experience, and hospital privileges required to perform advanced vascular interventional procedures.

- PF A qualified perfusionist to establish, monitor, and control the extracorporeal pump and veno-venous bypass circuit.
- AN A qualified anesthetist (anesthesiologist) and/or nurse anesthetist responsible for the management of sedation, analgesia, respiratory and cardiovascular support.
- PH A qualified pharmacist, on call during the procedure, to reconstitute the chemotherapeutic agent (melphalan hydrochloride), using national and local safety guidelines. The pharmacist should be aware of the rapid preparation time required for the preparation and administration of melphalan for use with the CHEMOSAT[®] System.
- A qualified chemotherapy healthcare professional certified by the site to deliver chemotherapy, such as Interventional Radiology Technician or Registered Nurse.
- IN A qualified intensivist, or appropriately qualified critical care specialist, responsible for providing medical management of the patient in the immediate post-procedure period during which the patient is in the intensive care unit or step down unit.

The procedure team is required to complete the Delcath training program. Refer to Procedure Flowchart on page 25 which provides an overview of the procedure and how the procedure team and their tasks work together.

To facilitate use of these instructions, the procedural sections include Healthcare User Identifiers to assist each user in identifying procedural steps applicable to them.

PROCEDURE

PREPARATION: PRIOR TO TREATMENT O IR PF AN PH DD IN

All medications and supportive measures must be determined and administered in accordance with each institution's policies, guidelines, procedures, the CHEMOSAT® System Instructions for Use and chemotherapeutic agent (melphalan hydrochloride) prescribing information.

Hepatic Vascular Mapping - Angiography and Embolization

In order to avoid inadvertent infusion of the gastrointestinal or visceral branches, conduct a thorough search for any variant anatomy. In addition, embolization of certain branches supplying the gastro-intestinal tract may be necessary.

- Prior to performing the procedure using the CHEMOSAT® System, complete a visceral angiogram (celiac artery and superior mesenteric artery). The presence of a replaced or accessory hepatic artery should be specifically addressed. Evaluate the portal vein for patency with late imaging during celiac and superior mesenteric arteriography. Completely examine the arterial supply to the liver and assess and understand its impact on chemotherapy infusion. Use of a selective micro-catheter may be beneficial for both embolization and subsequent drug infusion.
- Angiographic mapping of the hepatic arterial circulation with embolization
 of gastrointestinal branches arising from hepatic arteries is required to avoid
 inadvertent infusion of the chemotherapeutic agent into gastrointestinal arterial
 branches. It is recommended to complete embolization at least one week prior to
 treatment to allow the arterial puncture site to heal. Embolization on the same
 day as procedure is discouraged since the intense anti-coagulation required for
 percutaneous hepatic perfusion may not allow stable thrombotic occlusion of the
 coil-embolized vessels.

O IR PF AN PH DD IN



- Assess liver blood supply and formulate a strategy for catheter placement to ensure adequate drug infusion to the entire liver. Dependent on vascular anatomy, this may require repositioning of the catheter during the procedure.
- Embolization of the gastroduodenal artery is frequently necessary, but depends
 on its origin relative to the branches of the distal proper hepatic artery. If the
 infusion catheter tip can be placed sufficiently distally to avoid retrograde reflux
 into the gastroduodenal artery, then the latter may not need to be embolized.
- For patients with supply of the left lobe of the liver from the left gastric artery, selective embolization of left gastric artery branches will be necessary.
- The origin of the right gastric artery and/or supra-duodenal branches, and need for their embolization, must be assessed individually.
- In some patients (~15%), the liver receives arterial blood supply from two (2) different arteries: the right lobe from the superior mesenteric artery (replaced right hepatic artery) and the left lobe of the liver from the celiac artery.

For such patients, the delivery of chemotherapeutic agent to the entire liver may be accomplished by either:

- Temporary balloon occlusion (or permanent coil embolization) of one artery while infusing the remaining patent hepatic artery. Potential risks of this method are ischemia and/or delayed re-perfusion of the occluded lobe. Infusion and filtration periods remain unchanged.
- Sequential volume-adjusted infusion of each branch (60% to right; 40% to left). Infusion times will be different for each lobe, but total infusion and filtration times remain unchanged.
- If the risk assessment is unfavorable or the anatomic variation is too complex to allow selective catheterization for safe delivery of chemotherapy, the procedure must not be performed.
- During the infusion procedure described below, a single optimal angiographic projection of the hepatic artery should be used to periodically document hepatic arterial patency and flow.
- Review computed tomography or magnetic resonance imaging to assess venous anatomy. Select kit (balloon spacing) based on patient anatomy.

Coagulation Studies

- Perform coagulation studies pre-, peri- and post-procedure then repeat daily until normalized. Parameters tested must include:
 - Partial Thromboplastin Time
 - Prothrombin Time / International Normalized Ratio

Blood Products

Type and cross-match for :

- 4 units packed red blood cells
- 4 units Fresh Frozen Plasma
- 6 10 units platelets (depending on institutional guidelines)
- 10 units cryoprecipitate

Hydration

• Place a peripheral large gauge intravenous catheter and begin hydration the night prior to or day of the procedure as per institutional practice, to assist with maintaining mean arterial pressure above 60 mmHg during the procedure.

 A Foley catheter is recommended to closely monitor fluid balance during hydration.

Antibiotics

• Patients with a history of hepatobiliary surgery or ablative procedures must receive antibiotic prophylaxis peri-operatively.

Allopurinol

As a prophylaxis for possible tumor lysis syndrome, patients with more than 25% replacement of normal liver parenchyma with tumor are to be given allopurinol 300 mg/day orally beginning two (2) to three (3) days prior to percutaneous hepatic perfusion (PHP) with the CHEMOSAT® System and continuing two (2) to three (3) days following procedure.

Proton Pump Inhibitors

 To prevent gastritis which may occur as a result of regional melphalan absorption during the procedure, administer prophylactic proton pump inhibitors (for example: omeprazole, one 20 mg delayed release capsule by mouth no later than 8 PM the night before and at 7:30 AM the morning of the procedure followed by pantoprazole 40 mg intravenous every 8 hours during hospitalization).

Anticoagulation

- The patient will be systemically anticoagulated with heparin during the procedure. Proper anticoagulation is required to assure free extracorporeal flow and filtration. Activated clotting time must be closely monitored to ensure adequate anticoagulation.
 - Obtain the baseline activated clotting time value.
 - Administer heparin to the patient only AFTER placement of the 18F (femoral vein), 10F (jugular vein), and 5F (femoral artery) sheaths. Use Ultrasound guidance and single anterior wall puncture technique during sheath placement in order to avoid bleeding complications.
 - The patient must be fully heparinized prior to the insertion of the lsofuse[®] Catheter into the inferior vena cava. Begin with an initial intravenous bolus of heparin at 300 units/kg, dose adjusted to achieve activated clotting time.
 - A minimum activated clotting time of 400 seconds is necessary prior to balloon inflation and initiation of veno-venous bypass.
 - Maintain the activated clotting time above 400 seconds by repeat heparin bolus administration as necessary.
 - Evaluate activated clotting time frequently (approximately every 5 minutes) until adequate anti-coagulation is established (activated clotting time > 400 seconds). Maintain activated clotting time at > 400 seconds throughout the procedure, by checking it every 15 30 minutes depending on the patient's response and by administering intravenous heparin as needed.

Anesthetic Management

 Treatment must be administered with patients being monitored and under general anesthesia. Emergency resuscitation equipment must be available during the procedure.



Blood Pressure Control

- Procedure related decrease of blood pressure occurs when the balloons occlude blood return from the inferior vena cava (decreased cardiac inflow) and when the filters are brought into the extracorporeal bypass circuit. The reasons for filterrelated hypotension are multifactorial, but hypersensitivity to non-physiological surfaces (inflammatory response) and removal of catecholamines by the filters play a role. To aid blood pressure maintenance for extracorporeal bypass, the following actions are recommended per institutional practice:
 - Pre-operative hydration and intra-procedural administration of fluids.
 - Vasopressor use in accordance with institutional practices.
 - Administration of steroids prior to initiating extracorporeal bypass, in accordance with institutional practice.
- Blood pressure must be constantly monitored throughout the procedure and maintained at levels required for adequate perfusion of critical end-organs.

Drug Preparation and Delivery Planning

Prior to set up, provide pre-notification to the hospital pharmacy to be ready to prepare chemotherapeutic agent (Melphalan Hydrochloride for Injection). An actual request for drug preparation and delivery should be timed so that the start of the infusion of the melphalan is within thirty minutes of preparation. Drug administration should be completed within 60 minutes of the start of preparation.

System Components

Confirm that all components of the CHEMOSAT[®] System are available for assembly. Note: Certain components are not supplied by Delcath. Verify that the Medtronic pump is functioning properly (see pump operating manual for instructions on proper functionality).

PREPARING AND PRIMING THE HEMOFILTRATION CIRCUIT

PF

Systems,

Delcat

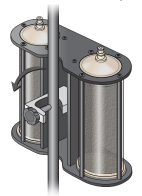
CAUTION: Adherence to strict sterile procedures is mandatory at all times.

1. Assemble Hemofiltration Circuit

See Figure 1 (Assembled System) for reference to a completely assembled circuit.

- (a) Utilizing strict aseptic technique, heparinize nine (9) liters of 0.9% Sodium Chloride Injection (normal saline) by adding 2000 units of heparin per liter.
- (b) Remove Hemofiltration (Chemofiltration) Dual Filter Cartridge from sterile pouch. Attach the filter to the intravenous pole using the built-in pole mount clamp, see Figure 3. Refer to "THIS END UP" label on filter faceplate, see Figure 4. Figure 4

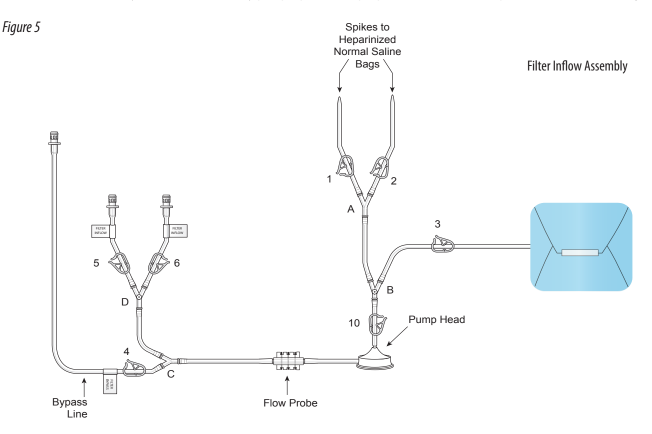
Figure 3





PF

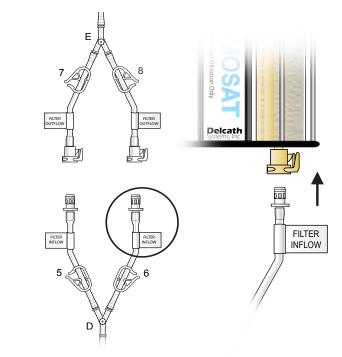
- (c) Open the circuit tray and remove the components that are in the pouches, and set aside for later assembly.
- (d) Remove the "Filter Inflow Assembly" section from the circuit tray, place pump head on the pump drive motor, and insert flow probe into flow transducer, see Figure 5.





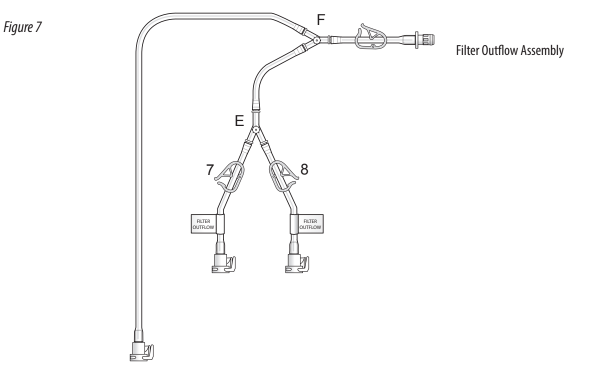
(e) Connect the Inflow lines (labeled and identified as in Figure 6) to the cartridge inlet connectors located on the bottom of the dual filter. To complete the assembly, push the quick connector couplings together, as shown in Figure 6 (male to female) until an audible "click" is heard to verify connection (push and click).





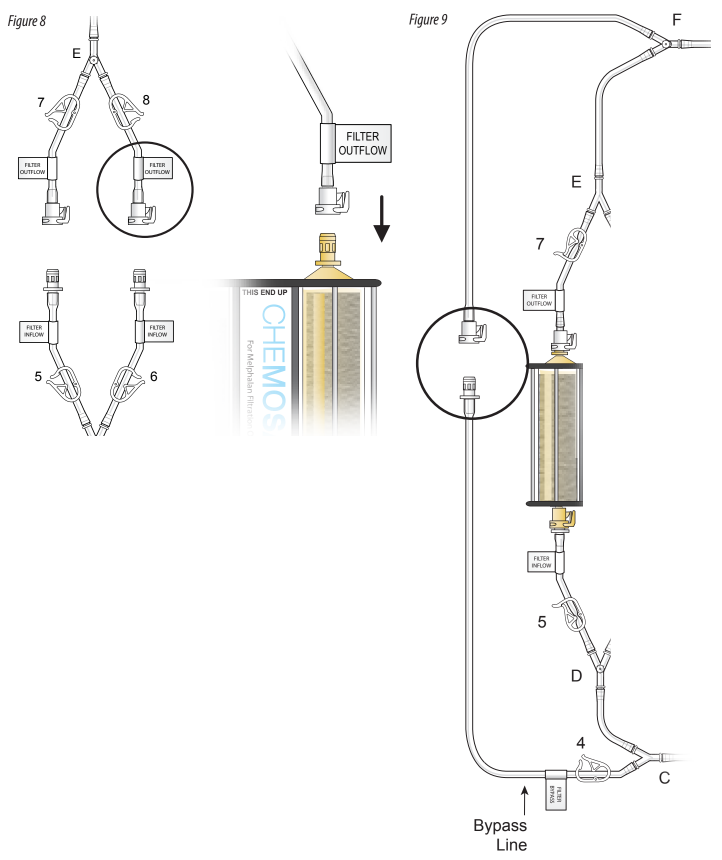
(f) Remove the "Filter Outflow Assembly" section from its sterile pouch, see Figure 7.

PF



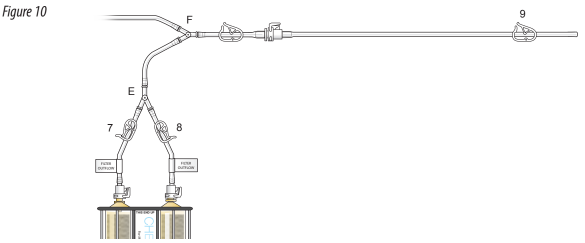


- (g) Connect the outflow lines (labeled and identified as in Figure 8) to the filter cartridge outlet connectors located on the top of the dual filter using the quick connector couplings (push and click), see Figure 8.
- (h) Assemble the two ends of the bypass line by pushing the quick connectors together (push and click), see Figure 9.

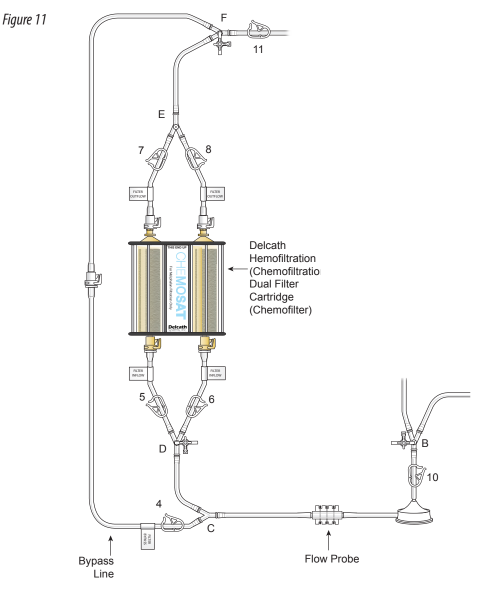




(i) Remove the "Prime/Flush Line" from its sterile pouch and attach it to the quick connect coupling located proximal to Y-connector "F", as shown in Figure 10. Place the open end of the "Prime/Flush Line" into the basin for collecting the flushed effluent during filter hydration.



(j) Attach the supplied stopcocks to the "B" (pre-pump), "D" (pre-filter) and "F" (outlet) Y-connector ports. Verify "Bypass Line" clamp 4 is open, as shown in Figure 11.

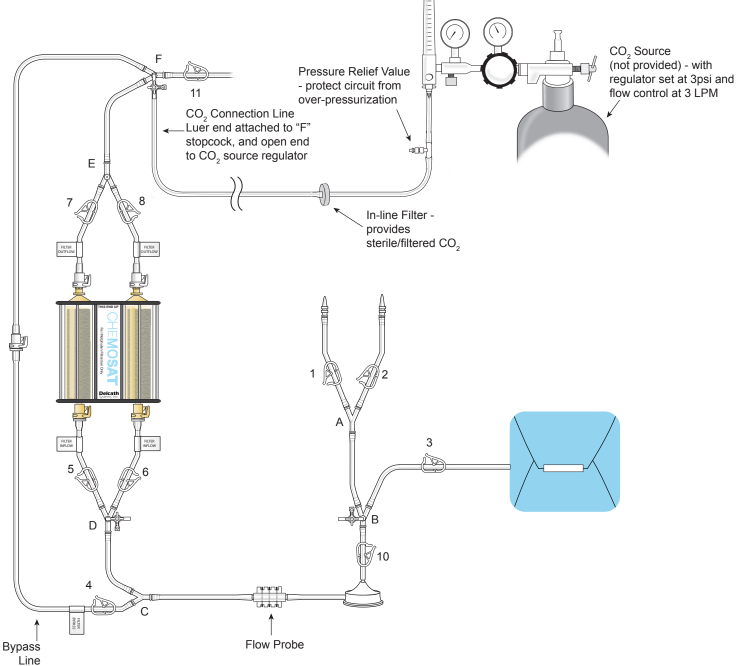




(k) System CO₂ Prime:

• Close outlet clamp (11), attach the CO₂ connection line to the stopcock "F", and open the stopcock, see Figure 12.

Figure 12



- Connect the open end of the CO, connection line to the CO, source, and set the CO, source regulator to 3psi (approximately 3.0 liters per minute (LPM)).
- Start the CO₂ gas flow and allow the CO₂ to flow through the hemofiltration circuit. Adjust CO₂ regulator to maintain 3psi (as necessary). Verify CO₂ flow through the circuit.
- Close bypass clamp (4) after approximately 1 minute to ensure flow through the hemofiltration cartridges. Allow CO₂ to flow through the cartridges (after closing clamp 4) for at least 5 minutes.
- Close saline clamps (1, 2), double balloon catheter line clamp (3), filter inlet clamps (5, 6), and then filter outlet clamps (7, 8) to lock CO₂ within the circuit. Stop the CO₂ flow, and close the stopcock "F" and disconnect the CO₂ connection line and discard.

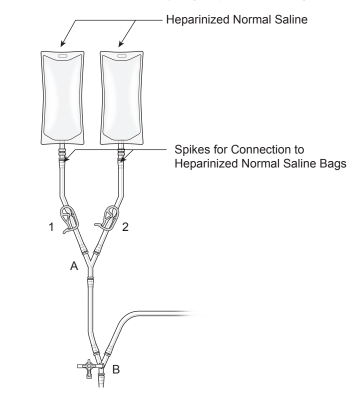


Figure 13

CAUTION: Use strict aseptic technique while spiking the heparinized normal saline bags.

PF

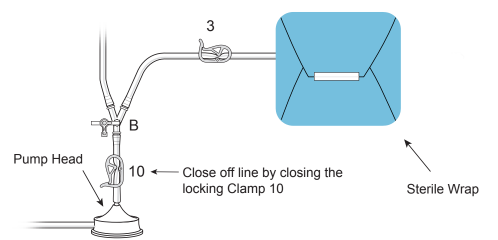
(I) Hang two bags of the heparinized sterile normal saline and connect to circuit by using the spikes, as shown in Figure 13, to allow for gravity priming of circuit components.



2. Prime Delcath Isofuse® Isolation Aspiration Catheter

(a) Close pre-pump clamp (10), open double balloon catheter line clamp (3), open saline line (clamp 1 or 2), and stopcock "B" to allow heparinized normal saline to prime line only up to clamp 3, see Figure 14. Once the stopcock line is primed and the double balloon catheter line is primed up to clamp 3, close clamp 3 and stopcock. **Do not allow excess** heparinized normal saline to fill sterile wrap.



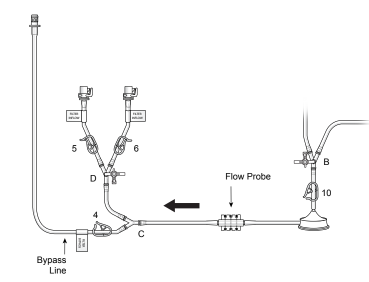




3. Prime Bypass Line

Figure 15

(a) Prime the pump head, filter inflow and bypass lines by opening clamp 10, stopcock "D", and clamp 4, see Figure 15.



<u>4. Prime and Flush Delcath Hemofiltration (Chemofiltration) Dual Filter Cartridge (Chemofilter)</u> <u>CAUTION:</u> Do NOT allow heparinized normal saline bags to run dry or air will enter the system.

(a) Close clamp 4 and open filter inlet clamps (5, 6) then filter outlet clamps (7, 8) and circuit outlet clamp (11) (see Figure 16) and adjust the flow of heparinized normal saline into the filter to a rate of approx. 0.5 liters per minute. Note: Hemostats (forceps) are required to adjust flow rate if using gravity.

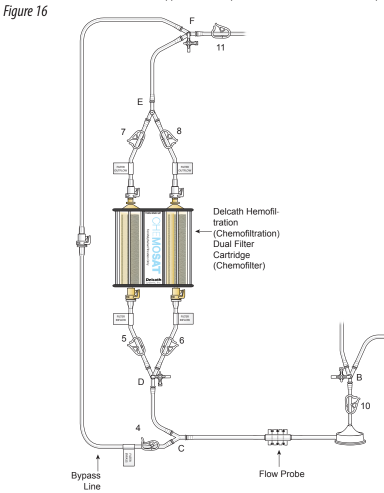




Figure 17

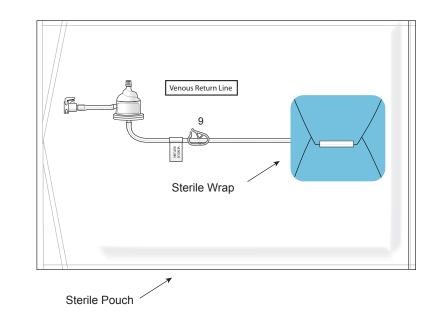
<u>CAUTION:</u> Do not use excessive force when tapping the plastic housing.

PF

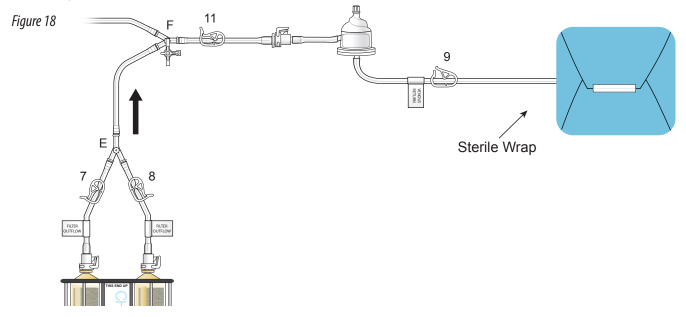
- (b) Allow heparinized normal saline to flow out through the filters and out the "Prime/Flush Line" for approximately six (6) minutes or until the filter appears gas free (solid black). Once all gas appears to have been displaced, tap gently to encourage any trapped gas bubbles to rise, turn cartridge within housing to visualize and access entire filter. Only tap on the sides of the cartridge (Do not tap on the end caps or quick connector couplings).
- (c) When Filter Cartridges are gas free, flush with an additional six (6) liters of heparinized normal saline (3 L/cartridge).
- (d) Clamp all filter lines (5, 6, 7, 8) and outlet clamp 11.

5. Prime Venous Return Line and Bubble Trap <u>CAUTION:</u> Do NOT install the Return Line with built-in Bubble Trap until flushing is complete.

- (a) Disconnect and dispose of the "Prime/Flush Line", by pressing in the latch located on the female quick connect coupling and pulling it a part.
- (b) Open the Venous Return sterile pouch and remove the venous return line and built-in bubble trap, (Figure 17).



(c) Attach the female to the male quick connector (push and click) located by outlet clamp (11), as shown in Figure 18. Position the bubble trap in the bubble trap holder higher than filter cartridges.

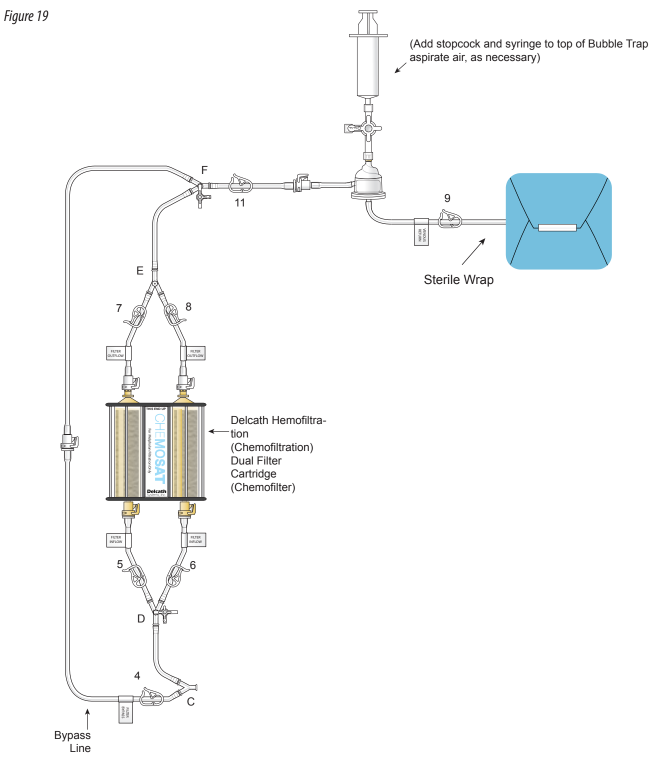




WARNING

Ensure That All Air Is Purged From The System Prior To Use In Order To Avoid An Air Embolism

(d) Prime venous return line and bubble trap by opening clamps 4, 11 and 9; attach stopcock to bubble trap and use syringe to aspirate air, as necessary, see Figure 19.



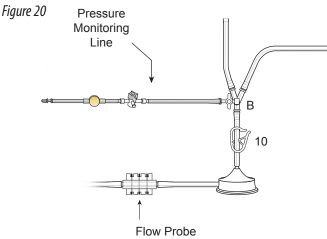
(e) Close clamp 9 once venous return line and bubble trap are primed up to clamp 9.



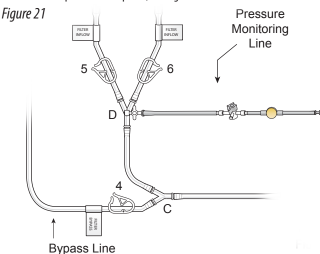


6. Install Pressure Monitoring Lines

(a) Attach pre-pump (to measure negative pressure – pump suction) pressure monitoring line to stopcock "B" and prime, see Figure 20.



(b) Attach pre-filter (to measure positive pressure – pre-filter) pressure monitoring line to stopcock "D" and prime, see Figure 21.



- (c) Attach the pressure monitoring lines to the P1 and P2 ports on the rear of the Medtronic Bio-Console 560.
- (d) Zero the pressure transducers (refer to Bio-Console Manual for details).
- (e) Coiled pressure monitoring lines are included for use with DLP Pressure Display Boxes, as necessary.

7. Pressure Test Circuit

- (a) Pressure test circuit by slowly ramping up the pump head speed (RPM) until a pressure reading of 300 mmHg is achieved on the pressure transducer attached to the line on Y-connector "D" (pre-filter).
- (b) Visually inspect all connections and cartridges to ensure no leaks are present.

<u>CAUTION</u>: If leak is noted, ensure connections are secure before proceeding.

- (c) Turn off pump and close cartridge inlet (5, 6) and outlet (7, 8) clamps. Ensure bypass line clamp (4) is open.
- (d) System is now primed, hydrated, de-bubbled, and ready for use.
- (e) Ensure there are two (2) liters of normal saline available for later use.

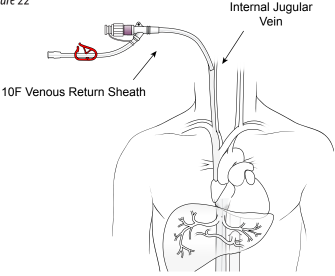
PLACING THE CATHETERS

8. Insertion of the 10F Venous Return Sheath

(a) Attach the stopcock to the sheath side port tube. Using standard Seldinger technique (with ultrasound guidance), insert the venous return sheath, into the internal jugular vein (preferably the right side internal jugular vein, see Figure 22). Flush the sheath with sterile heparinized normal saline. Close the stopcock. Use of sonographic guidance and a single anterior wall puncture of the vein are required to avoid inadvertent carotid artery puncture. If a carotid artery puncture is inadvertently made, the procedure must be aborted and postponed to a later date. Insert the obturator through the valve post placement of the sheath.

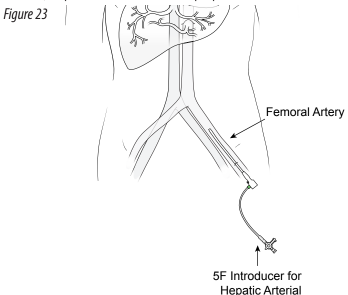
IR





9. Insertion of the 5F Femoral Arterial Sheath

(a) Using Seldinger puncture technique and standard fluoroscopic and arteriographic techniques, place the 5F Introducer Sheath into the femoral artery, see Figure 23. Use of sonographic guidance and a single anterior wall puncture of the femoral artery over the femoral head are required so as to assure compressibility of the artery when the sheath is removed. If a supra-inguinal puncture is inadvertently made, the procedure must be aborted and postponed to a later date.

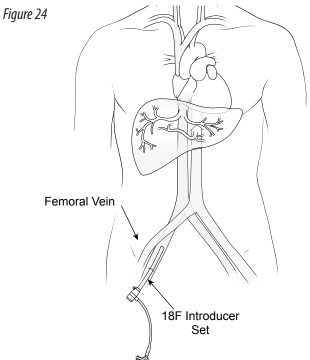






10. Insertion of the 18F Venous Sheath

(a) Using Seldinger technique and standard fluoroscopic and angiographic techniques, place the 18F introducer sheath into the femoral vein after serial dilation with 9F and 13F dilators. The venous sheath may be placed ipsilateral or contralateral to the 5F femoral arterial sheath placement, see Figure 24. Flush the sheath with sterile heparinized normal saline. Use of sonographic guidance and a single anterior wall puncture of the femoral vein over the femoral head are required so as to assure compressibility of the vein when the sheath is removed. If a supra-inguinal puncture is inadvertently made, the procedure must be aborted and postponed to a later date.



11. Insertion of 5F Chemofuse® Infusion Catheter

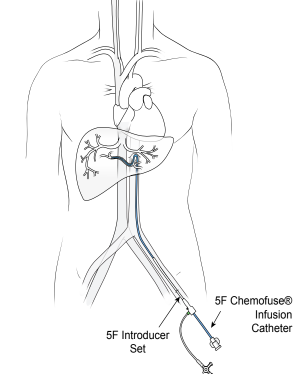
(a) Introduce the 5F Chemofuse[®] catheter through the sheath and manipulate it over a guidewire into the proper hepatic artery, see Figure 25. At the discretion of the Interventional Radiologist, a microcatheter may be coaxially introduced through the 5F catheter for selective catheter tip placement for drug infusion. If a microcatheter is used, attach a rotating hemostatic valve (Tuhoy-Borst type) to the 5F catheter and insert the microcatheter into the 5F catheter through the valve. Three microcatheters have been qualified by Delcath for use with the CHEMOSAT[®] System. Select one of the three microcatheters qualified (see Description of System Components (pg. 5).

WARNING

The Catheter Must Be Positioned As Described Below So That Drug Is Infused ONLY Into The Liver. Perfusion Of Drug Into Any Other Abdominal Organ Or Gastrointestinal Branches Must Be Avoided As This May Result In Serious Injury Or Death.

(b) Position the infusion catheter (5F catheter or microcatheter) in the proper hepatic artery well beyond the origin of the gastroduodenal artery to deliver chemotherapeutic agent (melphalan hydrochloride). Affix the 5F catheter to the skin at the groin. (c) Connect the infusion catheter (5F catheter or microcatheter) to the drug delivery system (see step 18), and maintain catheter patency by hospital catheter infusion protocols (e.g., infuse heparinized normal saline: The concentration of heparin should be 1000 units per 500 mL of normal saline.)

Figure 25



ESTABLISHING ANTICOAGULATION & PLACING ISOFUSE[®] ISOLATION ASPIRATION CATHETER

12. Anticoagulation

- (a) Obtain the baseline activated clotting time value.
- (b) Administer Heparin AFTER percutaneous placement of all introduction sheaths is completed, BUT PRIOR to the insertion of the Isofuse[®] Isolation Aspiration catheter into the inferior vena cava.
- (c) Administer an initial intravenous bolus of 300 units/kg of heparin. Heparin dose should be adjusted to achieve a minimum activated clotting time of 400 seconds prior to initiation of veno-venous bypass and balloon inflation.
- (d) Evaluate activated clotting time frequently (approximately every 5 minutes) until adequate anti-coagulation is established (activated clotting time > 400 seconds). Maintain activated clotting time at > 400 seconds throughout the procedure, by checking it every 15-30 minutes depending on the patient's response, and administering intravenous heparin as needed.

WARNING

The Start Of The Intra-Arterial Infusion Of The Drug Solution Must Be Within 30 Minutes Of Its Preparation In The Pharmacy.



NOTE: TIMING OF CHEMOTHERAPEUTIC AGENT DELIVERY

Time the request for delivery of the chemotherapeutic agent (Melphalan Hydrochloride) so that the start of the intra-arterial infusion of the drug solution is within thirty minutes of its preparation. Since preparation and delivery times vary, depending upon local practices, the timing of the request is critical and should be pre-arranged with the pharmacist. Typically, the time to request chemotherapeutic agent from the pharmacy would be when the lsofuse[®] Isolation Catheter is being positioned in the inferior vena cava.

13. Insertion of the Isofuse® Isolation Aspiration Catheter

- (a) Flush the lsofuse[®] catheter with heparinized normal saline.
- (b) Introduce the Isofuse[®] catheter through the 18F sheath. Under fluoroscopic guidance advance it over a guidewire into the inferior vena cava and position the catheter tip at the level of diaphragmatic hiatus. Do NOT inflate balloons.
- (c) Upon successful placement, remove guidewire and create a heparin lock within the "OTW" lumen to maintain patency.

CONNECTING CATHETERS TO HEMOFILTRATION CIRCUIT

14. Connection of Catheter to Hemofiltration (Chemofiltration) Circuit

- (a) Remove sterile wrap from hemofiltration circuit double balloon catheter line while maintaining sterility, and transfer sterile end to interventional radiologist.
- (b) Open saline line (clamp 1 or 2) clamp and clamp 3 to allow for a "wet connection" of the hemofiltration circuit to the lsofuse[®] catheter. After connection is made, close saline line clamp (clamp1 or 2). Ensure that all air is removed from the lsofuse[®] catheter.
- (c) Remove the sterile wrap from the hemofiltration circuit venous return line while maintaining sterility, and transfer sterile end to the interventional radiologist and flush normal saline to fill the line.
- (d) Connect the hemofiltration circuit venous return line to the stopcock of the 10F venous return sheath placed in the jugular vein (Venous Return Sheath tubing has a red clamp), flush normal saline through the line. When all air is removed and line is completely filled with normal saline turn stopcock to close the side port. Ensure the stopcock (at the venous return line to sheath connection) is fully open to minimize back pressure and maximize flow through the stopcock. ("OFF" handle of stopcock turned 90° to flow path.)

15. Establishing Hemofiltration Circulation

- (a) Start pump and slowly increase RPM control to achieve a maximum allowable flow rate which does not cause flow induced vibration or exceed the 0.80 L/min flow rate or -250 mmHg pre-pump pressure.
 - Flow rates of approximately 0.40 to 0.75 liters/minute are typical; however,
 0.80 L/min is the maximum allowable flow rate for this system.
 - In-line pressure transducers should be used to monitor pressures:
 - Pre pump pressure (suction side) should not be more negative than
 -250 mmHg, as lower pressures indicate possible catheter collapse or kink.
 - o Pre-cartridge pressures (pre-filter) should not exceed **200 mmHg**, as higher pressures indicate increasing filter resistance potentially due to thrombus or a kinked return line. Check filters to assure free flow and return line for kinks.

(b) The hemofiltration circuit is now established. Venous blood is aspirated from the central lumen through the fenestrations in the lsofuse[®] catheter. This blood flows through the lsofuse[®] catheter to the pump, through the bypass line, and returns to the patient through the venous return sheath.

<u>CAUTION</u>: Continously monitor any perfusion related events including:

- Blood flow-rate as displayed by the Medtronic Bio-Console.
- Systolic, diastolic and mean arterial blood pressure.
- Heart rate and vital signs.
- Activated clotting times.
- Bubble-trap for entrapped air.
- Leaks from any part of the circuit.

ISOLATING THE INFERIOR VENA CAVA

16. Inflation of Balloons

WARNING

There Is An Anticipated Significant Decrease Of Blood Pressure Following The Initial Occlusion Of The Inferior Vena Cava By The Balloons. It Is Critical To Maintain Mean Blood Pressure Above 65 mmHg..

Vasoactive Agents Response Testing: Prior to inflation of either balloon (occlusion of inferior vena cava), administer vasoactive agent to assess patient responsiveness to this agent. After inflation of the balloons, assess patient blood pressure for two (2) to five (5) minutes before proceeding. Significant decreases in blood pressure will occur within two (2) to five (5) minutes.

Continue to administer vasoactive agents to maintain mean blood pressures above 65 mmHg. Vasopressor agents are typically not required after the conclusion of the procedure.

(a) Perfusionist must carefully monitor the flow rate during the balloon inflation.

WARNING

Do NOT Overinflate The Balloons. Overinflation Of The Balloons Could Cause The Balloons To Burst Which Could Result in Life-Threatening Injury.

(b) Maximum balloon inflation volumes:

- Cephalad Balloon: **38** mL of dilute contrast medium
- Caudal Balloon: **38** mL of dilute contrast medium
- (c) Under fluoroscopy, partially inflate the cephalad balloon with approximately 15-25 mL of dilute contrast media (e.g., 35% dilution) within the right atrium (the balloon will have a rounded appearance).
- (d) With the caudal balloon still deflated, slowly retract the lsofuse[®] catheter until the cephalad balloon is at the junction of the right atrium and inferior vena cava. If needed, further inflate the cephalad balloon until indentation of the diaphragmatic hiatus is visible at the inferior margin (the balloon will acquire an acorn shaped appearance, see Figure 26). Do not inflate balloons beyond required volume to achieve an adequate seal. Never advance or retract the lsofuse[®] catheter when both balloons are inflated. If resistance is met during manipulation, determine the cause of the resistance before proceeding.





Figure 26



(e) Under fluoroscopy, inflate the caudal balloon with dilute contrast medium until the lateral edges of the inflated balloon start to become effaced by the inferior vena cava wall.

WARNING

Never Stop Blood Flow Through The Hemofiltration Circuit For More Than 30 Seconds.

(f) With balloons inflated, perform a limited (retro-hepatic) inferior vena cavagram (using digital subtraction angiography technique) through the fenestrations. Prior to injection of contrast medium, reduce the pump speed to 1000 RPM and clamp off the circuit. Inject iodinated contrast medium through the CONTRAST port to confirm that the catheter properly isolates hepatic venous flow between the balloons. The cephalad balloon must occlude the inferior vena cava just above the highest (closest to right atrium) hepatic vein, and the caudal balloon must occlude the inferior vena cava just below the lowest hepatic vein (above the renal veins) as shown in the radiographic image in Figure 27.

Figure 27



Re-establish flow through the hemofiltration circuit by unclamping the circuit and returning pump RPM to deliver previous flow rate.

WARNING

Never Adjust the Position of the Double Balloon Catheter Unless Both Balloons are Fully Deflated.

- (g) If the Isofuse[®] catheter is not in the proper position, deflate both balloons (caudal balloon first) and then reposition the catheter, while maintaining flow in the hemofiltration circuit.
- (h) Once satisfactory position is attained (i.e., the isolated segment is well sealed), gently hold the proximal end of the Isofuse[®] catheter to prevent upward migration of the catheter into the right atrium. The catheter must be held and its position checked for the duration of the procedure (approximately 60 minutes).

<u>CAUTION</u>: Check Isofuse[®] catheter balloon positions fluoroscopically every four (4) to five (5) minutes during drug administration and filtration to ensure continued hepatic venous isolation.

BRINGING HEMOFILTRATION CARTRIDGES ON LINE

17. Bringing Hemofiltration Cartridges on Line

- (a) Continuously monitor and check the patient's blood pressure as required (see "Blood Pressure Control").
- (b) Leaving the bypass line open, open clamps on the left cartridge (5 and 7), and allow blood to displace the heparinized normal saline into the patient.
- (c) After the heparinized normal saline in the left cartridge and its lines is fully replaced with blood, wait approximately 30 seconds and open clamps on the right cartridge (clamps 6 and 8), while keeping the bypass line open. Once the heparinized normal saline in the right cartridge and its lines is fully replaced with blood, wait approximately 30 seconds and then close the bypass line by **securely closing clamp 4.** Add a reusable tube clamp as a redundant bypass closure mechanism high on the bypass line in clear view of the team.

WARNING

Close Bypass Line Prior To Infusion of Drug.

SETUP DRUG DELIVERY SYSTEM AND START EXTRACORPOREAL FILTRATION

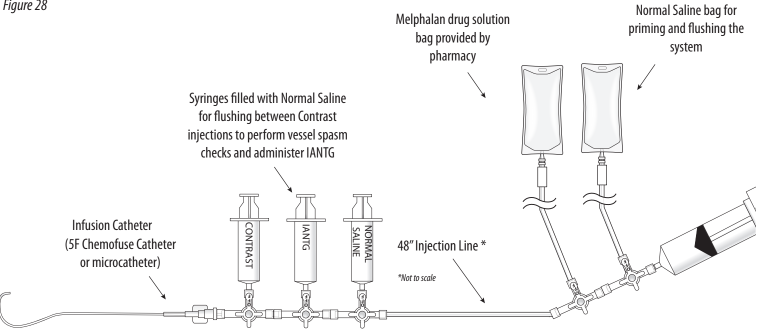
18. Drug Administration and Extracorporeal Filtration

- (a) When the hemofiltration circuit is running satisfactorily and the patient is hemodynamic stable, flush the hepatic arterial infusion line with normal saline to avoid directly mixing heparin with chemotherapeutic agent (melphalan hydrochloride). Connect drug infusion line to Hepatic Artery Infusion catheter (5F Chemofuse® Catheter or microcatheter), as shown in Figure 28 to complete the drug delivery circuit.
- (b) Following normalization of blood pressure, perform an arteriogram to assess patency of the hepatic artery. Use undiluted iodinated contrast agent to check for hepatic artery spasm via CT. The contrast is injected by hand via the syringe for the arteriogram. In circumstances where hepatic arterial spasm is noted, administer nitroglycerin intra-arterially to alleviate the spasm. Always flush the injection line with normal saline after contrast injections.





Figure 28



WARNING

Assess Arterial Patency Approximately Every Four (4) to Five (5) Minutes Via Contrast Administration During Drug Infusion. Administer Intra-Arterial Nitroglycerin if Arterial Spasm is Noted. If Spasm Cannot be Relieved, Terminate the Procedure (see Ending Extracorporeal Circulation below).

(c) Initiate administration of the chemotherapeutic agent (melphalan hydrochloride) through the infusion catheter (5F Chemofuse® or microcatheter) over a period of 30 minutes.

WARNING

Immediately Stop The Procedure If Perfusion Of Drug Is Detected Outside Of The Isolated Region And Cannot Be Corrected. Once the Infusion Of Chemotherapeutic Agent (Melphalan Hydrochloride) Has Started, Do NOT Deflate Balloons Unless Administration of Drug Has Been Stopped And A Full Washout Cycle (30 Minutes) Has Been Completed.

(d) After the prescribed dose has been fully administered, continue extracorporeal filtration for an additional 30 minutes (washout period).

ENDING EXTRACORPOREAL **CIRCULATION**

19. Ending Extracorporeal Circulation

- (a) At the end of the 30-minute wash-out period, deflate the caudal balloon fully.
- (b) Then deflate the cephalad balloon fully.
- (c) Discontinue filtration by reducing the pump RPM to 1000, closing clamps 3 and 9, and then stop flow by turning off the pump.

(d) A portion of blood in the circuit may be returned to the patient by adding sterile heparinized normal saline into the circuit to flush the blood back into the patient. Open clamps 1 and 2 of the circuit to use the heparinized normal saline from the bags used to prime the circuit. Do not open the bypass line clamps.

IR

CATHETER REMOVAL

20. Catheter Removal

- (a) Remove the infusion catheter. The 5F arterial sheath should only be removed when coagulation status has been normalized
- (b) Remove Isofuse[®] catheter carefully and replace with 18F obturator. Place the obturator completely into the sheath so the obturator hub bottoms out onto the sheath hub. The 18F venous sheath should only be removed when coagulation status has been normalized.
- (c) Close the stopcock or red clamp on the 10F venous return sheath sideport and disconnect the venous return line from the sheath. Do not remove the 10F venous return sheath until coagulation status has been normalized.
- (d) Dispose of all components appropriately in accordance with hospital, local, state, and federal biohazard guidelines.

NORMALIZATION OF COAGULATION STATUS FOR SHEATH REMOVAL O IR IN

21. Normalization of Coagulation Status for Sheath Removal

- (a) Administer protamine sulfate by slow intravenous infusion in a dose appropriate to the amount of heparin given and the activated clotting time.
- (b) Administer 10 units of cryoprecipitate and/or Fresh Frozen Plasma based on coagulation profiles to correct remaining abnormalities per institutional guidelines.
- (c) Repeat coagulation profile.

IR PF



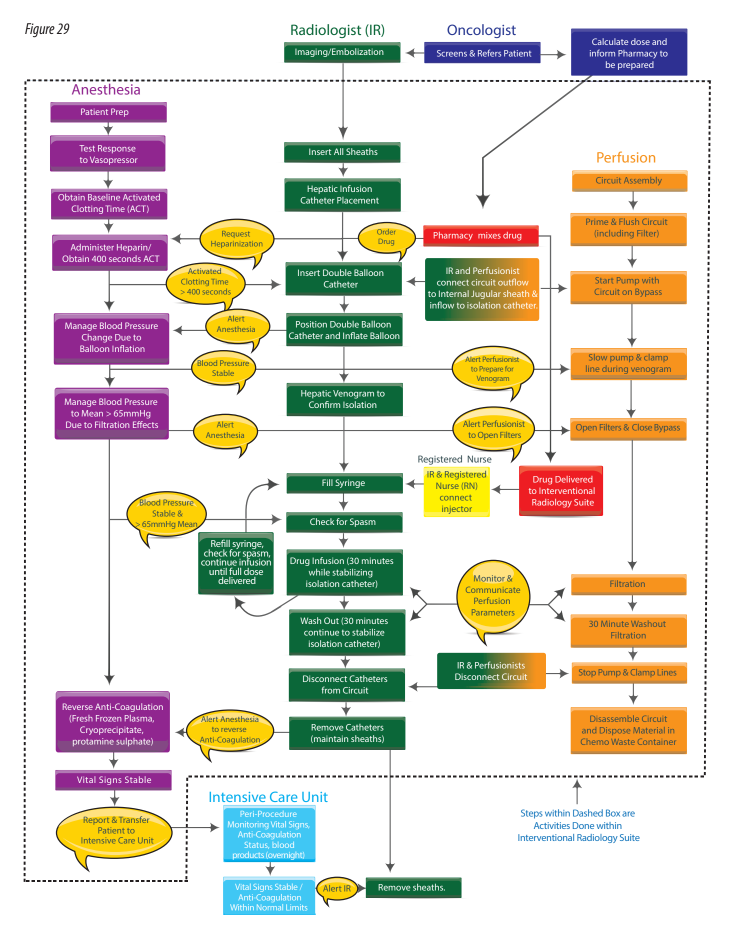
(d) Correct remaining coagulopathy following institutional guidelines. The following recommendations are provided for consideration:

Coagulation Profile	Action
Prothrombin time greater than 2 seconds of normal	Administer Fresh Frozen Plasma
Partial thromboplastin time greater than 5 seconds of normal	Administer protamine

- (e) Plasma albumin and globulin levels are replenished when Fresh Frozen Plasma is given. If replacement is needed, infuse according to institutional guidelines.
- (f) Measure blood platelet levels to determine if replacement is needed. Follow hospital protocols and transfuse accordingly.
- (g) Follow institutional guidelines for administration of packed red blood cells for anemia.
- (h) All sheaths may be safely removed if the platelet count is greater than 50,000/ mm³ and after the patient's coagulation status has normalized. Compress puncture sites until adequate hemostasis is achieved.
- (i) Dispose of all components appropriately in accordance with hospital, local, state, and federal biohazard guidelines.
- (j) Carefully monitor the patient until complete recovery.



PROCEDURE FLOWCHART





LIMITED WARRANTY

Delcath Systems, Ltd. ("Delcath") warrants that the Delcath Hepatic CHEMOSAT® Delivery System ("Product") will be free of defects in material and workmanship at the time of delivery and until the stated expiration date as indicated on the Product. This warranty is available and extended only to the original end-user purchaser of the Product. THE FOREGOING IS DELCATH'S SOLE WARRANTY.

Any Product, or part of the Product, that is judged by Delcath to be defective in material or workmanship during the warranty period will be replaced or repaired by Delcath at its sole option and its expense. Remedies available to the purchaser under this warranty are limited to replacement of the entire Product or malfunctioning parts of Product with the specific remedy subject to determination by Delcath in its sole and reasonable judgement. Application for warranty coverage and remedy must be made in writing to Delcath within ten (10) days of the apparent malfunction. All authorized Product returns will be subject to Delcath's Return Goods Policy.

This warranty is void if the Product has been (a) improperly stored or handled; (b) modified, altered or repaired in any way; (c) reused, reprocessed, or re-sterilized; (d) improperly or negligently used or misused; (e) combined or used with a third party's products; or (f) damaged by negligence, accident, or intentional misconduct.

THE FOREGOING WARRANTIES ARE EXCLUSIVE AND IN LIEU OF ANY OTHER WARRANTIES, EXPRESS OR IMPLIED, WITH RESPECT TO THE GOODS SOLD HEREUNDER. EXCEPT AS EXPRESSLY PROVIDED HEREIN, DELCATH MAKES NO WARRANTY OF ANY KIND, EXPRESS OR IMPLIED, ORAL, WRITTEN OR OTHERWISE, WITH RESPECT TO THE PRODUCT(S) SOLD HEREUNDER, INCLUDING, BUT NOT LIMITED TO, ANY IMPLIED WARRANTIES OF MERCHANTABILITY AND FITNESS FOR A PARTICULAR USE OR PURPOSE. DUE TO BIOLOGICAL DIFFERENCES IN HUMAN PATIENTS AND BECAUSE DELCATH HAS NO CONTROL OVER THE CONDITIONS UNDER WHICH ITS PRODUCTS ARE USED, DIAGNOSIS OF THE PATIENT, TYPE OF TREATMENT, SURGICAL PROCEDURE, THE METHOD OR ADMINISTRATION OF THE PRODUCT OR THE STORAGE AND HANDLING OF THE PRODUCT AFTER IT LEAVES DELCATH'S POSSESSION, DELCATH DOES NOT WARRANT EITHER A GOOD EFFECT OR AGAINST ILL EFFECT FOLLOWING THE USE OF THE DELCATH PRODUCT AND DELCATH MAKES NO WARRANTY AS TO WHETHER OR NOT ANY PARTICULAR OR DESIRED RESULT IS OBTAINABLE BY APPLICATION OR USE OF THE DELCATH PRODUCT.

DELCATH SHALL UNDER NO CIRCUMSTANCES BE LIABLE TO THE PURCHASER OR ANY THIRD PARTY FOR SPECIAL, INDIRECT, INCIDENTAL, PUNITIVE, CONSEQUENTIAL OR EXEMPLARY DAMAGES OF ANY NATURE, WHATSOEVER, INCLUDING, BUT NOT LIMITED TO, COMMERCIAL LOSS FROM ANY CAUSE, BUSINESS INTERRUPTION OF ANY NATURE, LOSS OF PROFITS OR REVENUE, REAL OR PERCEIVED LOSS OF USE, LOSS ARISING FROM A DEFECT IN DESIGN, MATERIAL AND/OR MANUFACTURE OR WORKMANSHIP AND/OR THE FAILURE OF THE PRODUCT(S) TO PERFORM AS SPECIFIED, EVEN IF DELCATH SHALL HAVE BEEN ADVISED TO THE POSSIBILITY OF SUCH DAMAGES.

IT IS REQUIRED THAT THE DELCATH PRODUCT BE OPERATED BY OR UNDER THE IMMEDIATE, DIRECT SUPERVISION OF A LICENSED PHYSICIAN OR OTHER LICENSED HEALTHCARE PROFFESIONAL QUALIFIED TO USE THE PRODUCT AND PERFORM THE PROCEDURE. DECATH DISCLAIMS LIABILITY FOR ALL INJURIES, DEATHS, OR PROPERTY DAMAGE ARISING FROM THE USE OF THE PRODUCT BY ANYONE, OTHER THAN QUALIFIED PERSONNEL DESCRIBED ABOVE, OR THE IMPROPER, NEGLIGENT OR RECKLESS USE OF THE PRODUCT, OR THE USE OF THE PRODUCT FOR ANY UNAPPROVED INDICATION OR FOR ANY USE NOT SPECIFICALLY INDICATED IN THE INSTRUCTIONS FOR USE FOR THE PRODUCT.





Sterilized using STERILE EO ethylene oxide.

Sterilized using STERILE steam or dry heat

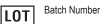


Keep Away From Sunlight.



Do not use if package is damaged.





Keep Dry



Contains or presence of LATEX natural rubber latex.



Manufacturer

Federal Law (USA) restricts R ONLY this device to sale by or on the order of a physician.



PHT

Consult Instructions For Use

Contains or presence of phthalate: bis (2-ethylhexyl) phthalate (DEHP).





 Customer Service

 Phone: +353 91 746200

 Fax: +353 91 746208

Distributed by:



medac GmbH Theaterstraße 6 22880 Wedel Germany EC REP Importer Delcath Systems Limited Unit 19 Mervue Industrial Estate Mervue, Galway IRELAND

Delcath is a registered trademark of Delcath Systems, Inc. CHEMOSAT[®], ISOFUSE[®] and CHEMOFUSE[®] are registered trademarks of Delcath Systems, Inc. © 2021 Delcath Systems, Inc. All rights reserved. Medtronic's Bio-Medicus® System, is a registered trademark of Medtronic Inc.



