

## Paediatric Intensive Care Unit

### Guidelines for Continuous Veno-Venous Haemofiltration (CVVH) and Continuous Veno-Venous Haemodiafiltration (CVVHDF) within the Cardiac Paediatric Intensive Care Unit at Leicester Royal Infirmary.

Staff relevant to:	PICU nursing staff trained in extended role, Consultants and ANP's within PICU.
Approval date:	December 2021
Version:	5
Revision due:	December 2024
Written by:	M McLaughlin & J Whitelaw
Reviewed by:	Kate Peace, Claire Westrope & Fiona Taylor
Trust Ref:	C151/2016



## **1. Introduction and Who Guideline applies to**

The purpose of this guideline is to provide support and direction when caring for a child requiring haemofiltration and haemodiafiltration. The following guidelines refer only to the use of the Prismaflex machine for children on the Cardiac Paediatric Intensive Care Unit (CPICU) at Leicester Royal Infirmary who are receiving Continuous Veno-Venous Haemofiltration (CVVH) or Continuous Veno-Venous Haemodiafiltration (CVVHDF). This guideline is intended to be used by all PICU nursing staff trained in extended role, Consultants and ANP's within PICU.

Clinical guidelines are 'guidelines' only. The interpretation and application of clinical guidelines will remain the responsibility of the individual. If in doubt consult a senior colleague or expert. Caution is advised when using the guidelines after the review date. Deviation from this guideline must be documented in the nursing notes with an explanation of the circumstances.

Please use this guideline in conjunction with the following:

- [Continuous Renal Replacement Therapy UHL Paediatric Intensive Care Guideline C15/2018](#)
- [ECMO Levitronix CentriMag - Neonatal and Infant UHL Childrens Intensive Care Guideline C100/2016](#)

### **Contents**

1. Introduction and Who Guideline applies to .....	2
2. Introduction and description of therapies .....	3
Ultrafiltration: .....	4
Convection: .....	4
Diffusion: .....	4
Continuous Veno Venous Haemofiltration (CVVH) .....	5
Continuous Veno Venous Haemodiafiltration (CVVHDF) .....	5
Prescribing .....	5
Calculating treatment parameters .....	5
'Unintended Pt Fluid Loss or Gain Limit' screen .....	6
Blood Flow Rate .....	6
Replacement rate .....	6
Dialysate flow rate .....	7
High Volume Treatment .....	7
Pharmacokinetics .....	7
Patient fluid removal rate .....	7
Replacement fluids and adding electrolytes .....	8
Adding electrolytes to replacement and dialysate fluid .....	9
Management of hypernatraemia on CRRT .....	9
Considerations for therapy .....	10
Setting up .....	10
Priming .....	10
Blood priming .....	10
Care and clinical monitoring of patient .....	10
Blood sampling guidelines .....	11

Temperature control.....	11
Connecting CVVH/DF circuit to ECMO circuit .....	12
Potential problems whilst undergoing treatment.....	12
Cardiovascular instability .....	12
Hypernatraemia .....	12
Hyponatraemia.....	12
Coagulation of circuit in a patient with Haemolytic Uraemic Syndrome. (H.U.S).....	13
Emergency Disconnection.....	13
Cardiac Arrest.....	13
Anticoagulation .....	13
Contraindications: .....	13
Administration & dosage: .....	14
Anticoagulation whilst on ECMO .....	15
Monitoring.....	15
Epoprostenol.....	15
3. Education and Training .....	18
4. Monitoring Compliance .....	18
5. Supporting References .....	18
6. Key Words .....	19
Contact and review details .....	20

## 2. Introduction and description of therapies

Continuous renal replacement therapy (CRRT) is a supportive therapy used for the management of fluid balance and metabolic derangement, with or without evidence of acute injury kidney injury (AKI), in critically ill patients.

Aims of the therapy include:

- Relieving hypervolaemia and maintaining fluid balance
- Removing excess urea & creatinine
- Correcting and maintaining metabolic and electrolyte balance
- Removing toxins

Indications for commencing treatment:

- Fluid overload
- Electrolyte imbalance
- Acute Kidney Injury (AKI)
- Inborn errors of metabolism
- Sepsis
- Rhabdomyolysis
- Tumour lysis
- Drug intoxications
- Optimising nutrition

In the East England, East Midlands and South Yorkshire (EMEESY) Children’s Kidney Network, CRRT is utilised in the following forms; Peritoneal Dialysis (PD), Continuous Veno Venous Haemofiltration (CVVH) and Continuous Veno Venous Haemodiafiltration (CVVHDF). This guideline focuses on the use of CVVH and CVVHDF; these are both delivered using the Prismaflex (Baxter) machine.

Using a dedicated double lumen vascath and an extracorporeal circuit, blood is continuously removed, and returned, to the patient. This continuous technique allows fluid and waste products to be removed gradually over a 24hr period. This provides more haemodynamic stability for critically ill children who are often cardiovascularly unstable. This treatment can also be incorporated into a patient's Extra Corporeal Membrane Oxygenation (ECMO) circuit and run simultaneously.

The extracorporeal circuit incorporates a blood pump and a filter containing a semi-permeable membrane. A pressure gradient within the circuit forces fluid and solutes across the membrane to form effluent fluid. Dialysis fluid can also be added to the other side of the filter, this causes molecules to move across with a concentration gradient. The processes utilised are ultrafiltration, convection and diffusion.

### **Ultrafiltration:**

This is the movement of fluid through a semi-permeable membrane driven by a hydrostatic pressure gradient. On the Prismaflex a positive pressure is generated by the blood pump on the blood side of the semi-permeable membrane and a negative pressure by the effluent pump on the other side. This results in the movement of fluid from the positive pressure side (blood side) to the negative pressure side.

### **Convection:**

This occurs when large volumes of fluid crossing the semi-permeable membrane down the hydrostatic pressure gradient results in solvent drag across the membrane. Large molecules can be moved efficiently if the flow is fast enough. Pre blood pump and replacement fluid is added to the circuit to increase the flow across the semi-permeable membrane.

### **Diffusion:**

This is the movement of solutes across a semi-permeable membrane caused by a concentration gradient. Solute move from an area of higher concentration to an area of lower concentration. On the Prismaflex this is achieved by the addition of dialysate fluid on the other side of the semi-permeable membrane, this is run counter-currently to the blood flow to ensure the concentration gradient is maintained the whole length of the filter.

The movement of molecules in CRRT are driven by the above processes but are limited by the size of the solute particles. The Prismaflex filters have pores that allow passage of molecules up to 35,000 Daltons. This allows for free movement of small ions and molecules (e.g. urea, creatinine, ammonia) across the membrane as well as some larger molecules such as myoglobin (17,200 Daltons), insulin (active) and interleukins (varies). Small molecules are more efficiently removed by diffusion and larger molecules by convection.

Some molecules are removed in CRRT by adherence to the artificial semi-permeable membrane; this is the process by which inflammatory markers are thought to be removed. High levels of these molecules can cause the filter to clog and become less effective.

On CPICU at UHL using the Prismaflex (Baxter), two modes of CRRT are utilised, these are CVVH and CVVHDF.

## Continuous Venous Haemofiltration (CVVH)

This mode of CRRT utilises the principles of ultrafiltration and convection and is ideal for removing middle and large molecules.

- Biochemistry is controlled by removing large volumes of filtrate and replacing it with electrolyte containing fluid (replacement fluid). The more filtrate you remove and replace the more efficient haemofiltration is in controlling biochemical disturbance.
- As most solutes are distributed within the extracellular and intracellular fluid compartments (total body water), the volume of filtration (replacement) necessary to control biochemistry relates to total body water. Clinical experience has shown that a replacement of approximately 50% of bodyweight (1kg = 1litre) is usually adequate for solute and electrolyte removal.
- Low blood flow rate, high haematocrit and high plasma protein concentration will limit the rate at which filtration can occur and solutes (particularly of higher molecular weight) can be removed.

## Continuous Venous Haemodiafiltration (CVVHDF)

This mode of CRRT utilises the principles of ultrafiltration, convection and diffusion; using both convective and diffusive transport systems will optimise the clearance of molecules of varying sizes.

- The addition of dialysate flow with CVVHDF will improve the efficiency of acid base, waste and control of electrolyte balance. Waste and electrolytes diffuse from the patient's blood into the dialysate fluid surrounding the fibres. The "saturated" dialysate fluid (effluent) is removed from the filter and discarded.
- Diffusion is a two way process, molecules that are at a low concentration in the blood and high in the dialysate fluid will diffuse into the blood and vice versa (e.g. bicarbonate).
- Convective transport systems utilised in CVVH are limited by the blood flow achieved. If higher replacement rates are indicated, but not achievable, adding in dialysate flow will optimise clearance. Reasons for not achieving desired replacement rates could be due to patient stability or issues with access limiting the blood flow rates.
- This is practically important when treating children with inborn errors of metabolism, for overdose of drug or therapeutic agents and for patients with a high lactate.
- Increasing dialysate flow will at least theoretically improve clearance; however this is limited by the relatively low dialysate flow rates generated by CRRT equipment.

## Prescribing

### Calculating treatment parameters

The CVVH/DF prescription is based on a 24hr period. Shorter sessions of CVVH/DF will require adjustment of the flow rates and filters to achieve the same **daily** amount of filtration.

The prescription needs to account for the patient's current situation and the desired management over the next 24 hours. In general it is difficult to achieve a negative balance of more than 5-10% of patient's body weight over 24 hours.

### **All fluid handling rates on the Prismaflex are set in millilitres**

#### **'Unintended Pt Fluid Loss or Gain Limit' screen**

This is the difference between the fluid removal measured on the machine and the set fluid removal rate over the last 3 hours. Once this limit has been reached the machine will stop and no longer run. Therefore if a clamp has been left on a bag the alarm cannot be continuously overridden. The default limit is 60-400mls, but should be set based on patient's weight, clinical condition and haemodynamic stability. This should be assessed and prescribed on the prescription chart.

#### **Blood Flow Rate**

This is the volume of blood passing through the filter in a given time relative to body weight. These are affected and determined by the patient stability and the vascular access. Clotting in the lines or filter is most likely to occur when blood is travelling slowly therefore blood flow rates should be run at the higher end of the range if possible. There is no such thing as too much blood flow, other than maximum pre-programmed limits determined by the circuit size/machine. Blood flow rates should be adjusted in high flow CRRT to compensate for excessive haemoconcentration of the filter

Standard blood flow rates are 6-9ml/kg/min (up to 12-15ml/kg/min in lower weight babies). With a minimum rate of 30ml/min once initiated on treatment. Refer to CVVH/DF prescription chart for guidance on standard flow rates.

In patients >30kgs aim for a blood pump speed of 180-240ml/min. The machine will allow speeds up to maximum of 400ml/min but this is not often achievable or necessary.

#### **Replacement rate**

**Replacement rates = convection + ultrafiltration** medium size molecule clearance (e.g. products of inflammation/cytokines)

Total replacement rate is the volume of fluid taken out of the patient and replaced with replacement fluid every hour in ml/hr. It is also known as the volume of exchange.

Total replacement is calculated on 30ml/kg/hr this is the starting point to provide effective solute waste control. This is calculated based on normal creatinine clearance of 25ml/kg/min allowing for interruptions to treatment. However, replacement can be increased for enhanced clearance. For example septic/metabolic patients may have higher replacement rates, up to 80ml/kg/hr have been used. If a higher replacement is used a proportional higher blood pump speed is needed to stop the filter clotting. This is only possible if the patient's cardiovascular status and vascular access will tolerate it.

The Prismaflex allows for a mix of pre and post dilution replacement rates. As a usual starting rate 10mls/kg/hr pre dilution and 20mls/kg/hr post dilution is recommended. This may be altered dependent on the patient's clinical status and the condition of the circuit e.g if circuit pressures are rising indicating that the filter is clotting then increasing the amount of



pre-dilution may prolong the life of the filter. Maximum pre dilution would be 90%, 10% post dilution minimum is required to prevent clots forming in the bubble trap.

## Dialysate flow rate

**Dialysate rates = diffusion** small molecule clearance (e.g urea/ammonia)

Initial dialysate flow rate should be 20ml/kg/hr. As with replacement, increasing the rate will increase efficiency of therapy. Dialysate will rapidly change electrolyte and waste balance. Dialysate flow should **not** start at high volume rates because of the potential to change patient's osmotic pressure. Dialysate flow should only be increased if treatment is not being effective (up to 50ml/kg/hr). Dialysate flow is not affected by blood flow and increasing it does not cause haemo-concentration in the filter.

## High Volume Treatment

In patients with metabolic diseases ideally total body water should be exchanged in 8 hours, however this is usually not possible to achieve as enough blood flow cannot be achieved to avoid the filter clotting. In the case of metabolic patients the highest practical replacement rate should be used and dialysate flow used from the start. The normalisation of ammonia and metabolic acidosis being the point at which to consider a reduction in flow rates.

High volume replacement or dialysate flow rates have a dramatic effect on electrolyte balance, regular review of potassium, phosphate, calcium, sodium and pH balance are essential. All flow rates should be reviewed every 24 hours and reduced when patient improves or when desired outcome has been achieved. **Medical staff must prescribe any changes in flow rates.** The danger of haemoconcentration of blood in filter can be a problem with high volume replacement.

***In AKI no evidence exists to support one mode over another. Some proponents of CRRT recommend a combination of convection and diffusion to optimise removal of small and medium sized molecules (assuming that SIRS/Sepsis plays a part in most AKI models)***

***A typical starting prescription for fluid overload, oliguria would be 30ml/kg/hr convection, with the addition in SIRS/Sepsis of 20ml/kg/hr diffusion***

## Pharmacokinetics

Drug removal by haemofiltration depends on molecular weight, albumin binding, water solubility and volume of distribution, therefore some drugs handling properties are changed by the therapy. Filtration will remove non-protein bound, water soluble drugs, which are available in the circulation. Adding dialysate flow will cause more efficient drug removal by diffusion. Drug dose changes should be checked with literature (in drug information folder) and a pharmacist.

## Patient fluid removal rate

This is calculated on an hourly basis depending on actual patient fluid balance, desired fluid balance and condition and stability of the patient. Individual prescription depends on input (i.e. nutrition, IV fluids, drugs) and output (i.e. urine, gastric, drain losses). The Prismaflex will continue to run at the set hourly rate until changed by staff. The rate **must** be reviewed every hour. Beware of removing fluid too quickly.

If a negative balance is required this may be calculated on a 12 hourly basis or over a specified period of time. This should be prescribed on the patients prescription chart. Desired negative balance and hourly actual fluid balance both need to be added together to work out fluid loss total to be programmed. Do not remove fluid bolus given to support blood pressure or intravascular loss. Blood products which have not been administered for volume should be added to the fluid loss to be programmed.

If the machine shows a minus number that means that it has gained that amount of fluid. This may occur when you first commence treatment, but will readjust itself after a few minutes.

Deciding exactly how much fluid to remove from the patient is difficult to quantify as it depends on many clinical factors including urine output, insensible loss, hypervolaemia/hypovolaemia and clinical observations. Once fluid removal is started, close observation of the patient's cardiovascular and fluid balance status needs to be undertaken. Continuous evaluation of heart rate, core/peripheral temperature gap, capillary refill, CVP, blood pressure and blood biochemistry are essential.

**EXTREME CAUTION** should be used when setting the patient fluid loss rate, as the haemofiltration machine will try to remove that fluid; it has no way of assessing the effect on the patient. It will continue to remove fluid even when the patient is **HYPOVOLAEMIC**.

**Always treat the patient NOT the machine**

### Replacement fluids and adding electrolytes

Prismasol 4 & Hemosol BO

<b>Prismasol 4</b>									
Sodium	Potassium	Calcium	Magnesium	Chloride	Phosphate	Lactate	Bicarbonate	Glucose	Citrate
140	4	1.75	0.5	113.5	0	3	35	6	0

<b>Hemosol BO</b>									
Sodium	Potassium	Calcium	Magnesium	Chloride	Phosphate	Lactate	Bicarbonate	Glucose	Citrate
140	0	1.75	0.5	109.5	0	3	35	0	0

Prismasol 4 should be used as the standard replacement fluid unless the patient's serum potassium is over 6.0mmol/l. If the patient is receiving a potassium infusion, has potassium chloride in the intravenous maintenance, or as a supplement this should be reviewed and usually stopped at the commencement of therapy.

Prismasol 4 and Hemosol BO (HBO) are bicarbonate based replacement/dialysate fluids and must be mixed before use. If the patients U&E's/ blood sugars drop after starting treatment ensure fluid has been mixed properly.

Prismasol 4 has added potassium at a dose of 4mmol/L and glucose of 6.1mmol/L. HBO and Prismasol 4 also contain a low magnesium concentration (0.5mmol/l). When considering electrolyte balance remember the patients electrolyte levels will become similar to the concentration in the replacement/dialysate fluid and the higher the replacement/dialysate flow rate the faster this will happen.



If the patient has serum potassium **over** 6.0.mmol/L the potassium free HBO solution should be used as the initial replacement/dialysate fluid. Caution should be taken when using HBO as this fluid does not contain any glucose therefore very close monitoring of patients blood glucose is essential. As the serum potassium falls below 6.0mmol/L, the replacement/dialysate fluid should be exchanged for the Prismaol 4 solution.

If the patient has serum potassium **less than** 6.0mmol/L at the commencement of therapy Prismaol 4 solution should be used at the outset. The use of Prismaol 4 solution negates the use of extra potassium infusions. The patient's serum potassium will settle out at about 4.0mmol/L if Prismaol 4 is used.

### Adding electrolytes to replacement and dialysate fluid

With the exception of sodium and glucose if the patient requires other electrolytes the first line would be to administer this directly to the patient. If access is a major problem you may need to consider adding electrolytes to the replacement/dialysis solution and where possible advice should be sought from pharmacy to ensure there are no compatibility problems.

- Prior to additions being made to bags discussion must take place with pharmacy.
- Additives are always prescribed to be added per litre of fluid and as a total dose. Care needs to be taken when making up the fluid as a result.
- When additives are required more frequent blood sampling will be required.

### Management of hypernatraemia on CRRT

In cases of hypernatraemia in patients requiring CRRT it may be necessary to add sodium to replacement/dialysis fluids to ensure sodium correction does not occur too rapidly. The sodium content of both Prismaol 4 and Hemosol BO is 140mmol/L, so without the addition there is a significant risk the patient's serum sodium could drop too quickly.

If the patient's plasma sodium is greater than 160mmol/L it will be necessary to add 30% sodium chloride to bags of Prismaol 4 or Hemosol BO to prevent a rapid fall in plasma sodium. It should never fall >10mmol/L in 24 hours. Sodium chloride 30% contains 5mmol/ml of sodium.

Target Na <sup>+</sup> (mmol/L)	Addition to 5 litre bag (either Prismaol 4/Hemosol BO)	
150	10mls of sodium chloride 30%	Remove same volume from replacement/dialysate bag before addition (i.e. for target 150mmol/L remove 10mls from Prismaol 4/Hemosol BO, then add 10mls sodium chloride 30%)
160	20mls of sodium chloride 30%	
170	30mls of sodium chloride 30%	
180	40mls of sodium chloride 30%	
190	50mls of sodium chloride 30%	

If an addition is required this must be clearly documented on the patients CVVH/CVVHDF Daily Prescription and Record Chart and the patient's serum sodium must be closely monitored.

## Considerations for therapy

### Setting up

- Ensure at least one adult unit of packed cells available.
- Circuit of appropriate size. If patient is on ECMO only use the ST circuits as blood flow from ECMO circuit is too fast for the HF20 circuits.

Circuit	Patient Weight (kg)	Prime Volume (ml)	Machine Maximum Replacement Rate (ml/hr)	Machine Maximum Blood Pump Speed (ml/min)
HF20	0-12	60	780	100
ST60	12-25	93	2250	180
ST100	25-100	150	9000	400

- Once the CVVH/CVVHDF catheter is inserted obtain baseline bloods and a non-heparinised Activated Clotting Time (ACT).
- Obtain most recent haematocrit (Hct), ensure this is programmed into Prismaflex.
- Each lumen should then be locked with the exact volume as stated on the catheter with heparin 100 units/ml and labelled stating 'heparin 100 units/ml insitu'.
- Heparin **MUST** be aspirated and the aspirate discarded, before connecting to patient.
- If patient on ECMO continue with ECMO parameters for heparinisation

### Priming

Circuit	If serum potassium <6mmol/l		If serum potassium >6mmol/l	
	1 <sup>st</sup> prime	2 <sup>nd</sup> prime	1 <sup>st</sup> prime	2 <sup>nd</sup> prime
HF20	1 litre Plasmalyte 148	N/A	1 litre sodium chloride 0.9%	N/A
ST60 & ST100	1 litre Plasmalyte 148 with 2000 units heparin	1 litre Plasmalyte 148	1 litre sodium chloride 0.9% with 2000 units heparin	1 litre sodium chloride 0.9%

### Blood priming

This procedure should no longer be undertaken; if patient needs blood give as transfusion prior to or alongside commencing treatment.

### Care and clinical monitoring of patient

- Continuous monitoring of cardiovascular and respiratory parameters should be undertaken and recorded hourly on the PICU chart.
- All patients who are receiving CVVH/DF must have central/peripheral temperature\*, heart rate, arterial blood pressure and preferably CVP monitoring (\*central temperature does not have to be a rectal temperature).
- Fluid balance needs careful monitoring because of the large volumes of fluid being removed/infused.

- There are inherent errors in all the measures of fluid balance therefore it is prudent to assess the following factors before deciding upon the hydration status of the patient in relation to CVVH/DF:
  - Fluid balance
  - Clinical examination (HR,ABP,CVP,etc..)
  - Biochemistry
  - Haematocrit
  - Weight (if possible)

### Treat the patient NOT the machine

#### Blood sampling guidelines

This specifies the minimum frequency, but will be dictated by patient's clinical condition.

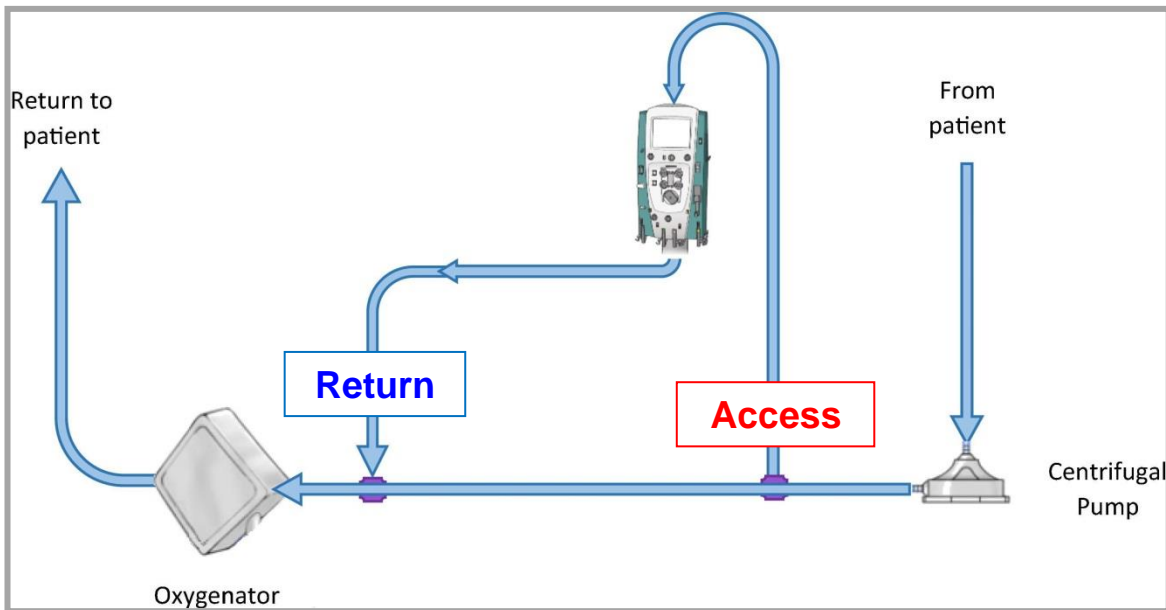
Test	Frequency	Special notes
Arterial blood gas	4 hourly	Carry out initial gas 1 hr after treatment commenced. Particular attention to; pH, lactate, bicarbonate, sodium, potassium & blood sugar.
FBC	Daily	Consider wash back/blood prime.
U&E's	4 hourly on ABG 12 hourly in labs	If using PrismaSol 4 fluid remember to gradually discontinue potassium in maintenance, infusions & supplements.
Clotting	Every 12hrs	Increase frequency if on Protein C or Antithrombin III.
Other electrolytes	Calcium, Magnesium & Phosphate 12 hourly	These are slowly filtered out.

#### Temperature control

- If patient is also on ECMO no further heating of the circuit is required
- The sleeve heater should be set 1-2°C above the desired patient temperature and attached to the **Return** line.
- Beware of the heater masking pyrexia.
- As the heater sleeve covers the **Return** line it is essential that a portion of the line is left exposed next to the patient's catheter so any air will be visible.
- If low weight baby and not on ECMO then consider using heater wires on both the **Return** and **Access** lines
- Additional warming devices may be required (e.g. overhead heater, Bair hugger).

## Connecting CVVH/DF circuit to ECMO circuit

Below diagram shows where to connect **Access** and **Return** lines, these should be connected using the pigtails on the ECMO circuit. This can only be done by ECMO trained staff.



## Potential problems whilst undergoing treatment

### Cardiovascular instability

This normally occurs as patient is being commenced on treatment, and can be more common in patients who are already unstable e.g. septicæmia.

*Action:* Ensure emergency drugs and fluid boluses are drawn up.  
Consider the use of inotropes.

### Hypernatraemia

Often occurs as a result of fluid restriction and excessive sodium intake e.g. in fluids and drugs. Haemofiltration replacement fluid itself contains approximately 140mmol/l.

*Action:* Minimise sodium intake in drugs, infusions and TPN.  
Increase free water intake (water which does not contain electrolytes).

### Hyponatraemia

May be present as part of water overload. When haemofiltration removes large amounts of fluid, the hyponatraemia may be worsened, as the fluid removed is plasma water (therefore has a high sodium content).

*Action:* Increase sodium intake and decrease free water intake.

## Coagulation of circuit in a patient with Haemolytic Uraemic Syndrome. (H.U.S)

Despite a low platelet count and high dose heparin it may be very difficult to keep continuous therapies running.

*Action:* A combination of heparin and Epoprostenol may be a more effective form of anticoagulation. NB Epoprostenol will not affect ACT.

## Emergency Disconnection

In some circumstances emergency disconnection is required such as lines have clotted, technical failure or evacuation of the unit. In these circumstances wash back should be considered if possible. If time allows lines should be disconnected and CVVH/CVVHDF catheter flushed and locked with heparin or at least with sodium chloride 0.9%.

## Cardiac Arrest

In the event of cardiac arrest while the patient is undergoing CVVH/CVVHDF treatment should be stopped and the line aspirated, flushed and locked as time allows. It may be an option to wash back the blood in the circuit, if deemed necessary to the patient. In some circumstances continuing filtration may be appropriate (e.g. if being used for life threatening electrolyte disturbance like HYPERKALEMIA). Haemofiltration catheters are central venous access and can be used instead of standard central lines in an emergency.

## Anticoagulation

As with any extracorporeal circuit, the circuits used for CRRT require the administration of anticoagulation to prevent the development of clots and keep the blood flowing in the circuit. On CPICU unfractionated heparin should be used as the first line anticoagulant. Heparin inhibits coagulation by preventing the conversion of prothrombin to thrombin and fibrinogen to fibrin. Heparin displays almost immediate action following intravenous administration. The clotting time then returns to normal within 2-6 hours after the infusion is discontinued. It binds extensively to plasma proteins and is activated in the liver and excreted in the urine. The action of heparin can be reversed with Protamine Sulphate. The dose is dependent on the heparin rate - see the BNFC for more information. The protamine dose should be based on the cumulative heparin dose given over the previous 2 hours.

## Contraindications:

- Active bleeding
- Imminent or recent surgery
- Coagulopathies such as thrombocytopenia, especially in septic and oncology patients
- Heparin Induced Thrombocytopenia (H.I.T)
- Disseminated Intravascular Coagulation (D.I.C)
- Liver disease
- Haemophilia
- Diabetes
- Pericarditis

CVVH/CVVHDF may be carried out in the above circumstances with minimal or no heparin or with an alternative agent such as Epoprostenol/Prostacyclin (see table below).

**NB.** It is possible to use less heparin if a faster blood flow is obtained, if the circuit is pre-diluted, or if the patient is on warfarin, aspirin or another anti platelet agent.

## Administration & dosage:

If running CVVH/DF as a stand-alone therapy heparin should be drawn up as per the IV Monograph, set up on an external syringe pump and administered into the heparin line on the circuit. The heparin bolus and infusion should be prescribed on the patient's prescription chart. Heparin for line locks is prescribed on the CVVH/DF prescription chart.

The initial bolus (if appropriate) should be given as blood reaches the pre filter port; this is then followed by a continuous infusion. The aim is to raise the patients clotting time sufficiently to prevent clotting in the filter and lines, while not increasing the risk of bleeding to the patient.

<b><u>HEPARIN</u></b>	
<b><u>Care: there are different strengths available – check!</u></b>	
<b><u>Use 1,000units/ml for these infusions</u></b>	
<b>Dose: for treatment or prevention of thrombotic episodes, after complex cardiac surgery procedures and for patients on CVVH</b>	
<b><u>Loading dose:</u></b>	
<b>Neonate &lt;35weeks post-conceptual age:</b>	50units/kg
<b>Neonate &gt;35weeks post-conceptual age:</b>	75units/kg
<b>Paediatric:</b>	75units/kg
<b><u>Initial maintenance dose:</u></b>	
<b>Neonates and children &lt;1year of age:</b>	25units/kg/hour
<b>Paediatric (&gt;1 year of age):</b>	20units/kg/hour
Usual range 10-40 units/kg adjusted according to APTT - Higher doses may be required	
<b>Method of administration:</b>	
<u>Loading dose:</u> IV bolus over 10 minutes (check if required with medical team)	
<u>Maintenance dose:</u> Continuous IV infusion	
Draw up 500 units/kg into a syringe and make up to 50ml with sodium chloride 0.9% or glucose 5%.	
1ml/hour = 10 units/kg/hour	4ml/hour = 40 units/kg/hour
<b>Dilution:</b>	
Sodium chloride 0.9% or glucose 5%	
Change infusion every 24 hours	
<b>Adverse effects:</b>	
Haemorrhage and thrombocytopenia	
Very rarely: hyperkalaemia (via hyperaldosteronism), hypersensitivity reactions, local skin irritation or skin necrosis	
<b>Notes:</b>	
Check clotting screen (baseline FBC, INR & APTT) prior to commencing heparin	
Check APTT regularly, advise 4 hours post loading dose and 4 hours after each alteration in dose	
Check APTT and FBC daily and potassium levels on alternate days	
Consult the pharmacist before use in renal or hepatic impairment	
Heparin has short half-life but if antidote required give protamine	

Above taken from IV Monograph (Sept 2021), please also refer to Medusa for any recent updates



## Anticoagulation whilst on ECMO

If patient is also on ECMO there is no need to separately anticoagulate the CVVH/DF circuit, anticoagulation will be provided from the heparin used for the ECMO circuit. Refer to guidance in ECMO protocol for more information.

## Monitoring

The level of anticoagulation is monitored by measuring the Activated Clotting Time (ACT). This is measured using the Hemochron machine regardless if patient is on ECMO or receiving treatment as a stand-alone therapy.

A clotting time that is taken pre-filtration and is not heparinised, such as at initial insertion of haemofiltration catheter must always be taken. This is essential because each individual will respond differently to the same dose of heparin; therefore, their reaction to it (the increase in the clotting time following heparin) will be different. The initial ACT should be 100-130 seconds. If already on ECMO no need for pre ACT.

Once commenced on treatment you will need to check an ACT. Further ACT's should be taken 1-2 hourly. The aim is to maintain the patients clotting time within the pre-determined limits on the prescription chart, usually between 140-160 or 190-210 seconds however this may change depending on patient situation

If patient already on ECMO take an ACT when first commencing therapy, if stable further monitoring can then be done at the same time the ACT's are taken for ECMO circuit.

If the ACT is low, thrombosis and clotting may occur – in the circuit and patient  
If the ACT is too high, bleeding may occur – in the patient

**The patients clotting must also be monitored closely via laboratory bloods; should the patients clotting become deranged the ACT target range may need to be reviewed and reduced.**

## Epoprostenol

If patient has contraindications to heparin then Epoprostenol can be used as an anticoagulant, can also be used in combination with heparin if heparin alone is proving inadequate. Epoprostenol is used to anticoagulate the patient and therefore the blood that is traveling through the circuit.

Epoprostenol is a prostaglandin and a potent inhibitor of platelet aggregation. This inhibition is dose related. Epoprostenol will not affect the ACT. It may however be of use to check the overall coagulation status regularly, but should not be used to titrate the Epoprostenol rate.

**When administering Epoprostenol it must be delivered directly to the patient, not through the Prismaflex machine. Use a dedicated central line for administration, if struggling with access add a 3-way tap on to return line of circuit.**

## **EPOPROSTENOL**

**(Prostacyclin)**

### **Dose:**

#### **Persistent pulmonary hypertension:**

All ages: 2 - 40 nanograms/kg/minute = 0.12 - 2.4 micrograms/kg/hour

(Remember that 1 microgram = 1000 nanograms)

#### **Platelet aggregation inhibitor / digital ischaemia:**

The dose for these indications in children are unclear

Suggested dose for all ages: 2.5 – 10 nanograms/kg/minute

However higher doses may be used, likely upper dose of 20 nanograms/kg/min, but can go up to 40 nanograms/kg/min

### **Method of administration:**

Continuous IV infusion

### **Dilution: DO NOT MIX WITH GLUCOSE 5%**

Epoprostenol is complicated to prepare, please follow instructions carefully:

1. Reconstitute a 500 microgram powder vial with 10ml of the glycine buffer provided in with the 50ml solvent vial.
2. Return this solution to the 50ml solvent vial and mix well. **\*SAVE THE 50ML VIAL\***

The concentration of this solution is 10 micrograms/ml (10,000 nanograms/ml).

3. **This solution needs filtering using the 0.22 micron filter provided before further dilution or administration**

For older children and higher doses this 10 microgram/ml solution can be given without further dilution via a central line.

**Rates:** 0.012ml/kg/hour = 2 nanograms/kg/minute  
0.24ml/kg/hour = 40 nanograms/kg/minute



### Anticoagulation treatment table

	Heparin	Epoprostenol	Heparin/Epoprostenol combination
Indication	Standard first line therapy	HIT Allergy to heparin  Bleeding (particularly intracranial)  Risk factors for bleeding	Heparin alone inadequate (filter life <24hrs excluding elective changes or access problems)  High risk of filter clotting (e.g. HUS, hypercoagulopathic state including DIC septicaemia)
Dosage range	10-40 unit/kg/hr	4-8nanogram/kg/min <b>MAX</b> <b>10nanogram/kg/min</b>	Heparin: 10-40unit/kg/hr Epoprostenol: 4-8nanogram/kg/min
Starting dose	20 units/kg/hr	4nanogram/kg/min	Heparin: 10units/kg/hr Epoprostenol: 4nanogram/kg/min
Titration point	140-160 seconds	Circuit life <48hrs, increase by 2nanogram/kg/min increments	ACT 140-180 seconds  If ACT >180 seconds ↓heparin by 10%  If ACT<140 seconds ↑ heparin by 10%  Circuit life <36hrs increase epoprostenol by 2nanogram/kg/min increments
Delivery method	Via heparin port on CVVH/DF circuit using an external port	To patient, via a central line. If no spare access add a 3-way tap onto return line of CVVH/DF circuit	Heparin to CVVH/DF circuit Epoprostenol to patient

### 3. Education and Training

All PICU nursing staff caring for patients receiving CVVH & CVVHDF are required to be trained in line with this extended role. Initial training session and ongoing training will be provided.

### 4. Monitoring Compliance

None

### 5. Supporting References

Kidney diseases improving global outcomes (2012) *Clinical Practice Guideline for Acute Kidney Injury* (online) KDIGO Clinical Practice Guideline

National institute for health and care excellence (2019) Acute kidney injury: prevention, detection and management (online) NICE Guidance NG148

Shaheen I, Harvey B, Watson A (2009) Haemofiltration Therapy *Paediatrics and Child Health* **19** (3) 121-126

Braun M. C and Welch T. R (1998) Continuous venovenous hemodiafiltration in the treatment of acute hyperammonemia. *American Journal Nephrology* **18** (6) 531-3

Brophy P, Somers M, Baum M et al. (2005) Multi-centre evaluation of anticoagulation in patients receiving continuous renal replacement therapy (CRRT) *Nephrology Dialysis Transplant.* **20** (7) 1416-21

Brophy P, Mottes T, Kudelka T et al. (2001) AN-69 membrane reactions are pH-dependent and preventable. *American Journal of Kidney Disease* **38** (1) 173-8

Brunet S, Leblanc M, Geadah. D et al. (1999) Diffusive and convective solute clearances during continuous renal replacement therapy at various dialysate and ultrafiltration flow rates. *American Journal of Kidney Disease* **34** 486-92.

Bunchman T, Maxvold N, Kershaw D et al. (1994) Continuous venovenous hemodiafiltration in infants and children. *Pediatric Nephrology* **8** 96-9

Goldstein S. (2003) Overview of pediatric renal replacement therapy in acute renal failure. *Artificial Organs* **27** (9) 781-5

Goldstein S, Somers M, Baum M et al. (2005) Pediatric patients with multi-organ system failure receiving continuous renal replacement therapy. *Kidney Int* **67** (2) 653-8

Harvey B, Watson A, Jepson S (2002) A renal critical care educator: the interface between paediatric intensive care and nephrology. *Intensive Critical Care Nursing* **18** (4) 250-4

Pediatric Continuous Renal Replacement Therapy. [www.pcrct.com](http://www.pcrct.com)

Ronco C et al. (2000) Effects of different doses in continuous veno-venous haemofiltration on outcomes of acute renal failure: a prospective randomised trial. *The Lancet* **356** 26-30

Ronco C, Pohlmeier R, Tetta C (2003) Intermittent or continuous treatment of acute renal failure? *Critical Care Medicine* **31** (9) 2417.

Schaefer F et al. (1999) Dialysis in newborns with inborn errors of metabolism. *Nephrology Dialysis Transplant* **14** 910-918.

Strazdins V, Watson A, Harvey B (2004) European Pediatric Peritoneal Dialysis Working Group: Renal replacement therapy for acute renal failure in children: European guidelines. *Pediatric Nephrology* **19** (2) 199-207

[www.baxterhealthcare.co.uk](http://www.baxterhealthcare.co.uk)

## **6. Key Words**

Acute Kidney Injury (AKI), Anticoagulation, Dialysate, Fluid balance, Electrolyte, Extra Corporeal Membrane Oxygenation (ECMO), Hemosol, Prismaflex, PrismaSol

The Trust recognises the diversity of the local community it serves. Our aim therefore is to provide a safe environment free from discrimination and treat all individuals fairly with dignity and appropriately according to their needs. As part of its development, this policy and its impact on equality have been reviewed and no detriment was identified.

<b>Contact and review details</b>	
<b>Guideline Lead (Name and Title)</b> Kate Peace – Paediatric Renal Critical Care Nurse Claire Westrope – Consultant PICU/ECMO Fiona Taylor – Senior Sister	<b>Executive Lead</b> <b>Chief Nurse</b>
<b>Details of Changes made during review:</b> <b>Major review and update undertaken 2021 – main changes as follows;</b> <ul style="list-style-type: none"> <li>• Added rationale, indications and explanations of CRRT treatments available</li> <li>• Removed step by step 'how to' guides</li> <li>• Removed standard flow rate table</li> <li>• Removed heparinisation section – replaced with IV monograph</li> <li>• Removed air embolism section</li> <li>• Removed wash back/return bloods and recirculation procedure sections</li> <li>• Added diagram showing circuit connections</li> </ul> <b>Prescribing</b> <ul style="list-style-type: none"> <li>• Blood flow minimum rate increased from 20ml/min to 30ml/min once initiated on treatment</li> </ul> <b>Setting up</b> <ul style="list-style-type: none"> <li>• ST60 prime volume reduced from 97ml to 93ml</li> <li>• Added obtain most recent Hct and programme into Prismaflex</li> <li>• Priming fluids composition updated to incl plasmalyte</li> <li>• Added statement highlighting to no longer blood prime</li> </ul> <b>Monitoring</b> <ul style="list-style-type: none"> <li>• Added consider heater wires if low weight baby not on ECMO</li> <li>• Added consider all factors when assessing fluid balance</li> </ul> <b>Emergency disconnection</b> <ul style="list-style-type: none"> <li>• Added wash back should be considered if possible</li> </ul> <b>Anticoagulation whilst on ECMO</b> <ul style="list-style-type: none"> <li>• Removed reference to Actalyke MINI macine</li> <li>• ACT monitoring changed from hrly for 1st 4 hours, to 1-2 hourly, if stable take at the same time for ECMO circuit</li> <li>• Added 190-210 seconds as an alternative clotting time to 140-160 dependent on patient situation</li> </ul> <b>Epoprostenol</b> <ul style="list-style-type: none"> <li>• Amendments made to dosing recommendations</li> <li>• IV Monograph added</li> </ul> References updated Format updated	

<b>REQUIREMENT</b>	<b>ACTION</b>
Where is the Policy available:	Within PICU attached to each Prismaflex machine. Intranet.
Copy to be sent to personnel with a request for inclusion in induction documents	No
Process for monitoring the effectiveness of this document	Will be audited internally.
Patient version.	No
Groups/persons consulted.	PICU Clinical Nurse Manager, PICU Educators, PICU Consultants, PICU Pharmacists, Nephrology Consultants
This Policy is subject to the Freedom of Information Act	