

Continuous Renal Replacement Therapy

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Definition of Terms

- SCUF - **S**low **C**ontinuous **U**ltrafiltration
- CAVH - **C**ontinuous **A**rterio**V**enous **H**emofiltration
- CAVH-D - **C**ontinuous **A**rterio**V**enous **H**emofiltration with **D**ialysis
- CVVH - **C**ontinuous **V**enovenous **H**emofiltration
- CVVH-D - **C**ontinuous **V**enovenous **H**emofiltration with **D**ialysis

Indications for Continuous Renal Replacement Therapy

- Remove excess fluid because of fluid overload
- Clinical need to administer fluid to someone who is oliguric
 - Nutrition solution
 - Antibiotics
 - Vasoactive substances
 - Blood products
 - Other parenteral medications

Advantages of Continuous Renal Replacement Therapy

- Hemodynamic stability
 - Avoid hypotension complicating hemodialysis
 - Avoid swings in intravascular volume
- Easy to regulate fluid volume
 - Volume removal is continuous
 - Adjust fluid removal rate on an hourly basis
- Customize replacement solutions
- Lack of need of specialized support staff

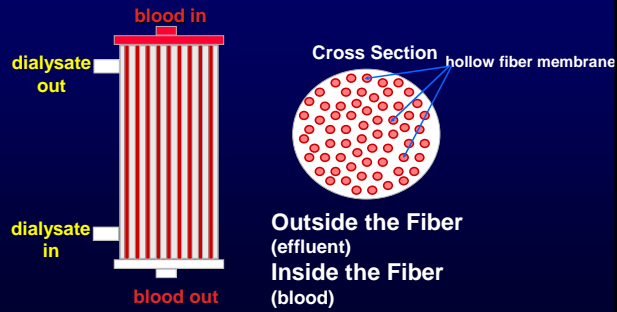
Disadvantages of Continuous Renal Replacement Therapy

- Lack of rapid fluid and solute removal
 - GFR equivalent of 5 - 20 ml/min
 - **Limited role in overdose setting**
- Filter clotting
 - Take down the entire system

Basic Principles

- Blood passes down one side of a highly permeable membrane
- Water and solute pass across the membrane
 - Solutes up to 20,000 daltons
 - Drugs & electrolytes
- Infuse replacement solution with physiologic concentrations of electrolytes

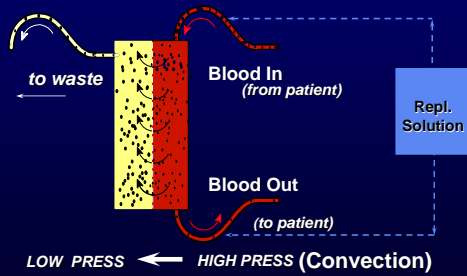
Anatomy of a Hemofilter



Basic Principles

- Hemofiltration
 - **Convection** based on a pressure gradient
 - 'Transmembrane pressure gradient'
 - Difference between plasma oncotic pressure and hydrostatic pressure
- Dialysis
 - Diffusion based on a **concentration gradient**

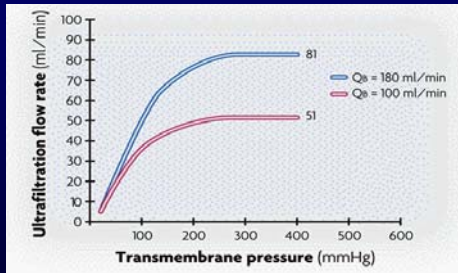
CVVH Continuous Veno-Venous Hemofiltration



CVVH Continuous VV Hemofiltration

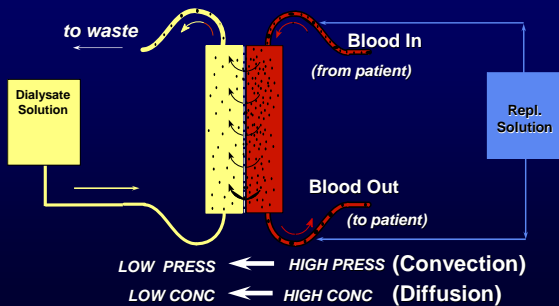
- Primary therapeutic goal:
 - Convective solute removal
 - Management of intravascular volume
- Blood Flow rate = 10 - 180 ml/min
- UF rate ranges 6 - 50 L/24 h (> 500 ml/h)
- Requires replacement solution to drive convection
- No dialysate

CVVH Performance



Continuous venovenous hemofiltration
 "In vitro" ultrafiltration with blood (post-dilution)
 (values \pm 15%) (Bovine blood at 37° C, Hct 32%, Cp 60g/l)

CVVHDF Continuous Veno-Venous Hemodiafiltration



CVVHDF Continuous VV Hemodiafiltration

- Primary therapeutic goal:
 - Solute removal by diffusion and convection
 - Management of intravascular volume
- Blood Flow rate = 10 - 180ml/min
- Combines CVVH and CVVHD therapies
- UF rate ranges 12 - 24 L/24h (> 500 ml/h)
- Dialysate Flow rate = 15 - 45 ml/min (~1 - 3 L/h)
- Uses both dialysate (1 L/h) and replacement fluid (500 ml/h)

Pharmacokinetics of Continuous Renal Replacement Therapy

Basic Principles

- Extracorporeal clearance (Cl_{EC}) is usually considered clinically significant only if its contribution to total body clearance exceeds 25 - 30%

$$Fr_{EC} = Cl_{EC} / Cl_{EC} + Cl_R + Cl_{NR}$$

- Not relevant for drugs with high non-renal clearance
- Only drug not bound to plasma proteins can be removed by extracorporeal procedures

Determinants of Drug Removal by CRRT

- Drug Same as hemodialysis but increased MW range
- Membrane Permeability
Sieving Coefficient
- Renal replacement technique Convection \pm diffusion CI
Flow rates
Blood, Dialysate, UF
Duration of CRRT

Sieving Coefficient (S)

- The capacity of a drug to pass through the hemofilter membrane

$$S = C_{uf} / C_p$$

C_{uf} = drug concentration in the ultrafiltrate

C_p = drug concentration in the plasma

$S = 1$ Solute freely passes through the filter

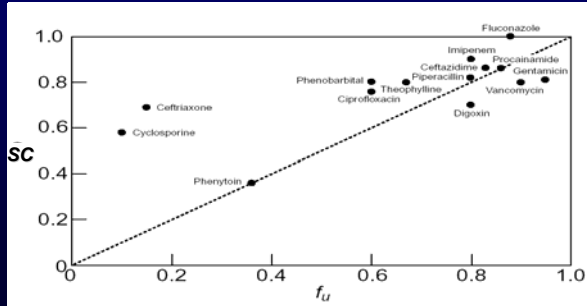
$S = 0$ Solute does not pass through the filter

$$CL_{HF} = Q_f \times S$$

Determinants of Sieving Coefficient

- Protein binding
 - Only unbound drug passes through the filter
 - Protein binding changes in critical illness
- Drug membrane interactions
 - Not clinically relevant
- Adsorption of proteins and blood products onto filter
 - Related to filter age
 - Decreased efficiency of filter

Relationship Between Free Fraction (f_u) and Sieving Coefficient (SC)



Dialysate Saturation (S_d)

- Countercurrent dialysate flow (10 - 30 ml/min) is always less than blood flow (100 - 200 ml/min)
- Allows complete equilibrium between blood serum and dialysate
- Dialysate leaving filter will be 100% saturated with easily diffusible solutes
- Diffusive clearance will equal dialysate flow

Dialysate Saturation (S_d)

$$S_d = C_d / C_p$$

C_d = drug concentration in the dialysate

C_p = drug concentration in the plasma

- Decreasing dialysate saturation
 - Increasing molecular weight
 - Decreases speed of diffusion
 - Increasing dialysate flow rate
 - Decreases time available for diffusion

$$Cl_{HD} = Q_d \times S_d$$

CVVHDF Clearance



Continuous venovenous hemofiltration - post dilution
 QB = 150 ml/min - QD = 2000 ml/h (in vitro saline)

Extracorporeal Clearance

- Hemofiltration clearance ($Cl_{HF} = Q_f \times S$)
 Q_f = Ultrafiltration rate
 S = Seiving coefficient
- Hemodialysis clearance ($Cl_{HD} = Q_d \times S_d$)
 Q_d = Dialysate flow rate
 S_d = Dialysate saturation
- Hemodialfiltration clearance
 $Cl_{HDF} = (Q_f \times S) + (Q_d \times S_d)$

Case History

- AP 36yo HM s/p BMT for aplastic anemia
- Admitted to ICU for management of acute renal failure
- CVVH-D initiated for management of uremia
- ICU course complicated by pulmonary failure failure requiring mechanical ventilation, liver failure secondary to GVHD and VOD, and sepsis

Case History Antibiotic Management on CRRT

- Gentamicin 180 mg IV q24h
- Vancomycin 1 g IV q24h
- Dialysis rate 1000 ml/hour
 - 12 hour post gentamicin levels: 3 - 4 mg/L
 - 12 hour post vancomycin levels: 20 - 23 mg/L
- Dialysis rate increased to 1200 ml/hour
 - 12 hour post gentamicin levels: < 0.4 mg/L
 - 12 hour post vancomycin levels: < 4 mg/L

Dosage Adjustments in CRRT

- Will the drug be removed?
 - Pharmacokinetic parameters
 - Protein binding < 70 - 80%
 - Normal values may not apply to critically ill patients
 - Volume of distribution < 1 L/kg
 - Renal clearance > 35%
- How often do I dose the drug?
 - Hemofiltration: 'GFR' 10 - 20 ml/min
 - Hemofiltration with dialysis: 'GFR' 20 - 50 ml/min

Drug Removal During CRRT

- Recommendations not listed in PDR
- Limited to case reports or series of patients
- Different filter brands, sizes, flow rates
- Limited information in many reports
 - Rarely report % of dose removed
- Many journals will not publish case reports
- Artificial models and predictions have no clinical value

Dosage Adjustments in CRRT

- Loading doses
 - Do not need to be adjusted
 - Loading dose depends solely on volume of distribution
- Maintenance doses
 - Standard reference tables
 - Base on measured loses
 - Calculate maintenance dose multiplication factor (MDMF)

Dosage Adjustments in CRRT

- Frequent blood level determinations
 - Aminoglycosides, vancomycin
- Reference tables
 - Bennett's tables or the PDR recommendations require an approximation of patient's GFR
 - The CVVH 'GFR' is approximated by the ultrafiltrate rate (UFR), plus any residual renal clearance
 - Using Bennett's or the PDR's tables, in most CVVH patients, drug dosing can be adjusted for a 'GFR' in the range of 10 to 50 ml/min

Supplemental Dose Based on Measured Plasma Level

$$\text{Dose}_{\text{Suppl}} = (C_{\text{target}} - C_{\text{measured}}) V_d$$

Adjusted Dose Based on Clearance Estimates

$$MDMF = \frac{CL_{EC} + CL_R + CL_{NR}}{CL_R + CL_{NR}}$$

COMPARISON OF DRUG REMOVAL BY INTERMITTENT HD AND CRRT

DRUG	$CL_R + CL_{NR}$ (mL/min)	MDMF	
		INTERMITTENT HEMODIALYSIS	CONTINUOUS RENAL REPLACEMENT
CEFTAZIDIME	11.2	1.6	2.2
CEFTRIAZONE	7.0	1.0	3.4
CIPROFLOXACIN	188	1.0	2.4
THEOPHYLLINE	57.4	1.1	1.4
VANCOMYCIN	6	3.9	4.9
