Continuous Renal Replacement Therapy

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Definition of Terms
- SCUF - Slow Continuous Ultrafiltration
- CAVH - Continuous Arteriovenous Hemofiltration
- CAVH-D - Continuous Arteriovenous Hemofiltration with Dialysis
- CVVH - Continuous Venovenous Hemofiltration
- CVVH-D - Continuous Venovenous Hemofiltration with Dialysis
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

Illustration of the anatomy of a hemofilter.

Blood flows through hollow fibers and the dialysis fluid is outside.
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

Illustration of Veno-Venous hemofiltration procedure.
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

Chart showing continuous venovenous hemofiltration and the role of transmembrane pressure.
CVVHDF
Continuous Veno-Venous Hemodiafiltration

Illustration of this process.
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 (\(\text{Cl}_{\text{EC}}\)) is usually considered clinically significant only if its contribution to total body clearance exceeds 25 - 30%.

Formula for calculating extracorporeal clearance.

- 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

<table>
<thead>
<tr>
<th>Determinant</th>
<th>Description</th>
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<tbody>
<tr>
<td><strong>Drug</strong></td>
<td>Same as hemodialysis but increased MW range</td>
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<tr>
<td><strong>Membrane</strong></td>
<td>Permeability</td>
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<td></td>
<td>Sieving Coefficient</td>
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<td><strong>Renal replacement technique</strong></td>
<td>Convection + diffusion Cl</td>
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<td></td>
<td>Flow rates</td>
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<tr>
<td></td>
<td>Blood, Dialysate, UF</td>
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<td>Duration of CRRT</td>
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Sieving Coefficient (S)
- The capacity of a drug to pass through the hemofilter membrane

Formula for calculating the sieving coefficient.

\[ C_{uf} = \text{drug concentration in the ultrafiltrate} \]
\[ C_p = \text{drug concentration in the plasma} \]

\[ S = 1 \quad \text{Solute freely passes through the filter} \]
\[ S = 0 \quad \text{Solute does not pass through the filter} \]

**Formula**
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 ($fu$) and Sieving Coefficient ($SC$)

Chart illustrating this relationship.
**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 ($S_d$)

Formula for Dialystate Saturation (SD)

\[ C_d = \text{drug concentration in the dialysate} \]
\[ C_p = \text{drug concentration in the plasma} \]

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

Formula
CVVHDF Clearance

Bar chart showing continuous venovenous hemofiltration – post dilution comparing urea + or-10%. with Vit B12 + or – 20% with Inulin + 20%.

Role of dialysis flows.
Extracorporeal Clearance

- Hemofiltration clearance - formula
  \( Q_f = \text{Ultrafiltration rate} \)
  \( S = \text{Seiving coefficient} \)

- Hemodialysis clearance - formula
  \( Q_d = \text{Dialysate flow rate} \)
  \( S_d = \text{Dialysate saturation} \)

- Hemodialfiltration clearance
  Formula
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 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

Formula for calculating supplemental dose based on measured plasma level.
Adjusted Dose Based on Clearance Estimates

Formulation for calculating adjusted dose based on clearance estimates.
COMPARISON OF DRUG REMOVAL BY INTERMITTENT HD AND CRRT

Chart comparing drug removal by intermittent HD and CRRT.

Examples: Ceftazidime
           Ceftriazone
           Ciprofloxacin
           Theophylline
           Vancomycin