



Indication of Continuous Renal Replacement Therapy and Nephrology problem with crrt



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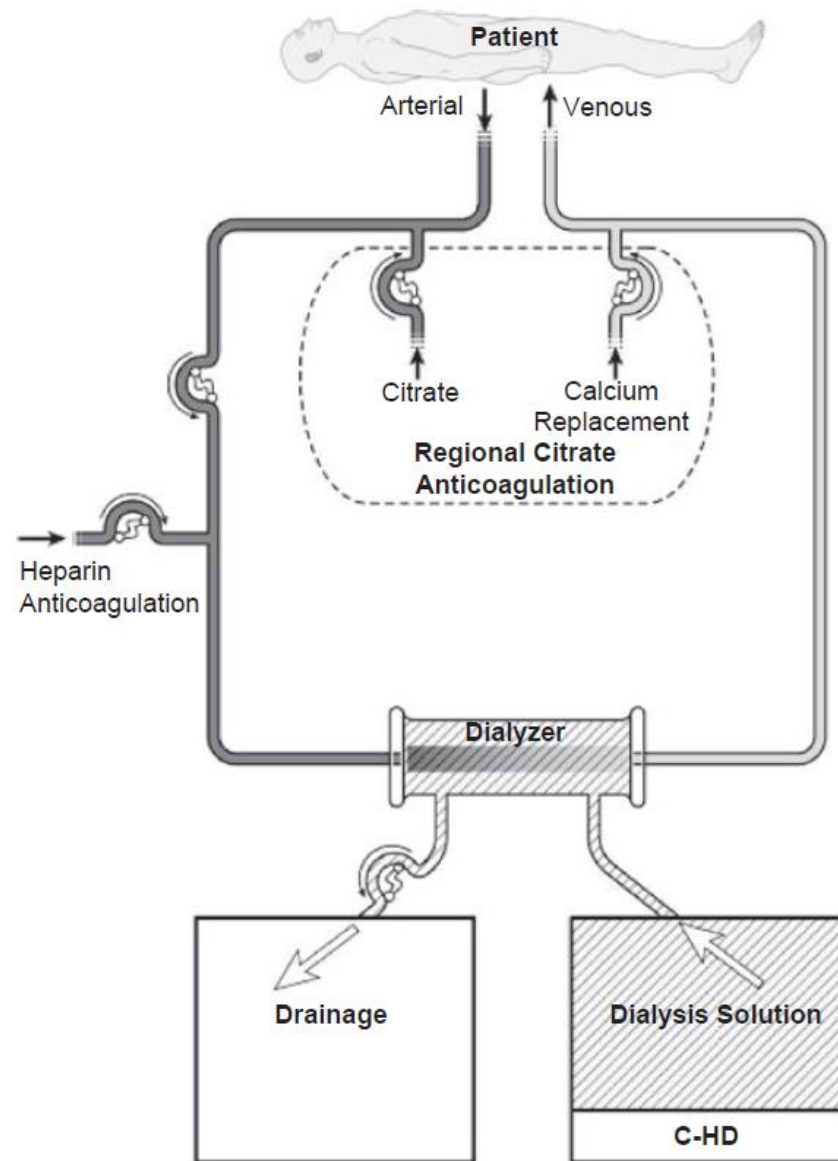
Introduction:

- The most widely used continuous renal replacement therapies (CRRT) for the treatment of critically ill patients in renal failure are continuous hemodialysis and hemodiafiltration.
- Two prolonged intermittent renal replacement therapies (PIRRT), sustained low-efficiency hemodialysis and sustained low-efficiency hemodiafiltration, are also **quite** popular.
- Continuous hemofiltration and slow continuous ultrafiltration are used, but **less** commonly.



Continuous hemodialysis (C-HD).

- Dialysis solution is passed through the dialysate compartment of the filter continuously and at a slow rate.
- In C-HD, **diffusion** is the primary method of solute removal.
- The amount of fluid that is ultrafiltered across the membrane is **low** (usually about 3–6 L per day) and is limited to excess fluid removal.

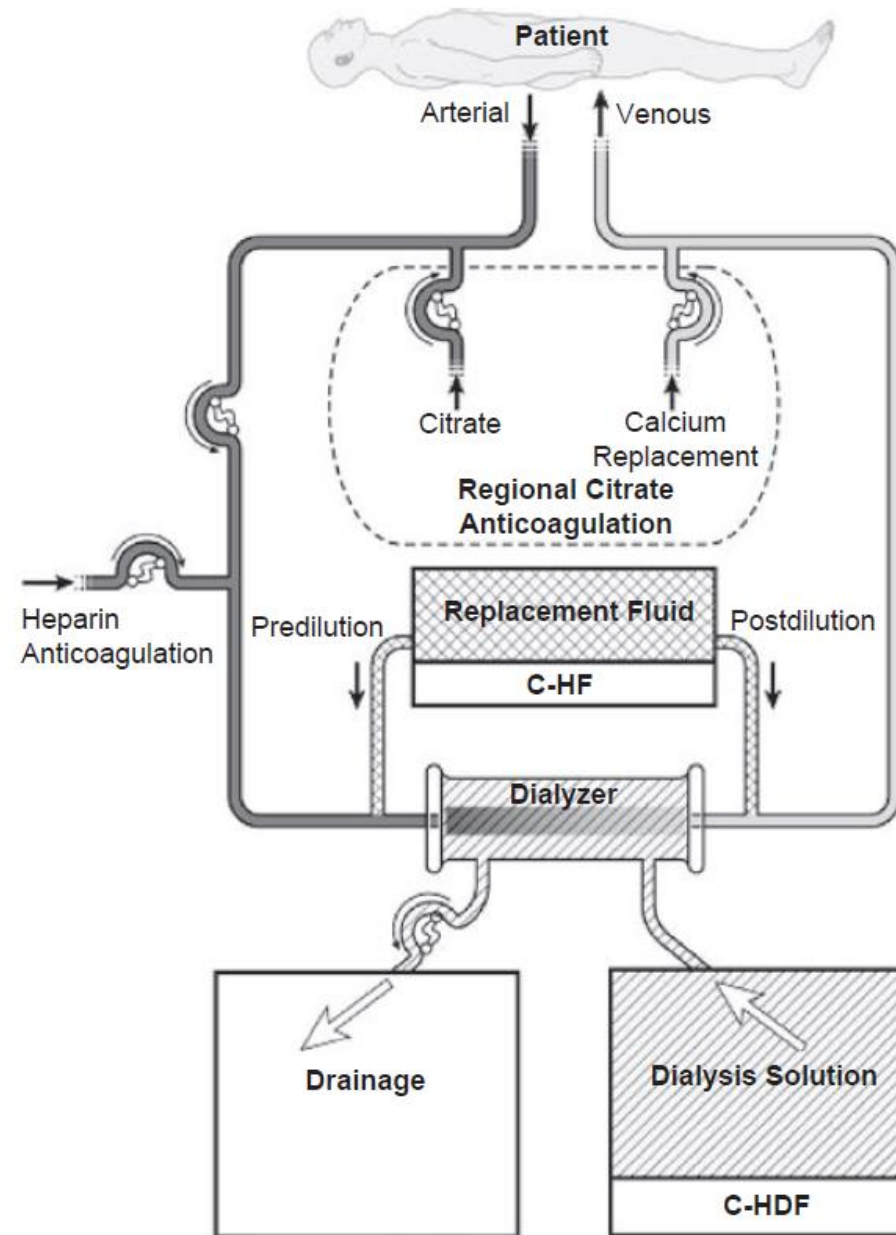


Continuous hemofiltration (C-HF).

- In C-HF , dialysis solution is **not used**.
- Instead, a large volume (about 25–50 L per day) of replacement fluid is infused into either the inflow or the outflow blood line (predilution or post dilution mode, respectively).
- With C-HF, the volume of fluid that is ultrafiltered across the membrane is the sum of replacement fluid and excess fluid removed, and so is much **higher** than with C-HD.

Continuous hemodiafiltration (C-HDF):

- Is simply a combination of C-HD and C-HF.
- Dialysis solution is used, and replacement fluid is also infused into either the inflow or the outflow blood line.
- The daily volume of fluid that is ultrafiltered across the membrane is equal to the replacement fluid infused plus the net volume removed.



Sustained low-efficiency dialysis and hemodiafiltration (SLED)

- SLED is a form of IHD using an extended (6- to 10-hour) session length and reduced blood and dialysate flow rates.

Typically, blood flow rates (BFRs) are about 200 mL/min and dialysate flow rate is 100–300 mL/min.

- The same machine used for IHD during the day often can be used for SLED during the night, and hemodialysis nurses can easily be trained to perform SLED, offering some economy of staff instruction.

CLINICAL INDICATIONS FOR CRRT

- 1. Hemodynamically well tolerated; smaller change in plasma osmolality.
- 2. Better control of azotemia and electrolyte and acid–base balance; correct abnormalities as they evolve; steady-state chemistries.
- 3. Highly effective in removing fluid (post surgery, pulmonary edema, ARDS).
- 4. Facilitates administration of parenteral nutrition and obligatory intravenous medications (i.e., pressor, inotropic drugs)
- 5. Less effect on intracranial pressure.
- 6. New user-friendly machines available.

DIFFERENCES AMONG C-HD, C-HF, AND C-HDF

- **Solute clearance with C-HD:**
 - where the BFR is 150–200 mL/min or more, and dialysate flow rate typically is 25–30 mL/min, clearance of urea and other small molecules is determined primarily by the dialysis solution flow rate.
 - BFR in C-HD should be at least three times the dialysate flow rate.
 - At this slow BFR and high blood-to-dialysate flow ratio, the outflow dialysate is almost 100% saturated with urea and other small-molecular-weight solutes

Continue.....

- The standard dialysis solution inflow rate is now about 20–25 mL/kg per hour.

VASCULAR ACCESS

- Using a dual-lumen cannula inserted into a large (internal jugular or femoral) vein. The subclavian vein can be used but is not the site of first choice.

The KDIGO AKI guidelines recommend using **uncuffed venous catheters for CRRT**).

Uncuffed catheter is easier, that the need for a cuffed catheter might sometimes delay initiation of therapy, and that the average duration of CRRT is only 12–13 days .

- compared use of longer (20–24 cm) soft, versus shorter (15–20 cm) the longer catheters were associated with **longer filter life** and improved dose of therapy.

Arteriovenous blood access:

Use of AV blood access for CRRT is **no** longer widely practiced.

- There is risk of damage to the femoral artery with possible distal limb ischemia, plus AV access will often not deliver high enough blood flows to be able to support the more intensive CRRT therapies in common use today.

TABLE
15.3 Composition of Some Continuous Renal Replacement Therapy Solutions

Component (mM)	Dialysis Machine Generated ^a	Peritoneal Dialysis Fluid ^b	Lactated Ringer Solution	B. Braun Duosol (5-L bag)	Baxter Accusol ^b (2.5-L bag)	Gambro PrismaSol ^c (5-L bag)	Nxstage Pureflow ^d (5-L bag)
Sodium	140	132	130	136 or 140	140	140	140
Potassium	Variable	—	4	0 or 2	0 or 2 or 4	0 or 2 or 4	0 or 2 or 4
Chloride	Variable	96	109	107–111	109.5–116.3	106–113	111–120
Bicarbonate	Variable	—	—	25 or 35	30 or 35	32	25 or 35
Calcium	Variable	1.75 (3.5 mEq/L)	1.35 (2.7 mEq/L)	0 or 1.5 (0 or 3.0 mEq/L)	1.4 or 1.75 (2.8 or 3.5 mEq/L)	0 or 1.25 or 1.75 (0 or 2.5 or 3.5 mEq/L)	0 or 1.25 or 1.5 (0 or 2.5 or 3.0 mEq/L)
Magnesium	0.75 (1.5 mEq/L)	0.25 (0.5 mEq/L)	—	0.5 or 0.75 (1.0 or 1.5 mEq/L)	0.5 or 0.75 (1.0 or 1.5 mEq/L)	0.5 or 0.75 (1.0 or 1.5 mEq/L)	0.5 or 0.75 (1.0 or 1.5 mEq/L)
Lactate	2	40	28	0	0	3	0
Glucose (mg/dL)	100	1,360	—	0 or 100	0 or 100	0 or 100	100
Glucose (mM)	5.5	75.5	—	0 or 5.5	0 or 5.5	0 or 5.5	5.5
Preparation method	6-L bag via membrane filtration	Premix	Premix	Two-compartment bag	Two-compartment bag	Two-compartment bag	Two-compartment bag
Sterility	No	Yes	Yes	Yes	Yes	Yes	Yes

5000 ml	Intravenous use
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1000 ml ready-to-use solution for haemofiltration contain:

Na ⁺	140 mmol/l
K ⁺	2.0 mmol/l
Ca ²⁺	1.5 mmol/l
Mg ²⁺	0.5 mmol/l
Cl ⁻	111 mmol/l
HCO ₃ ⁻	35.0 mmol/l
Glucose anhydrous	5.6 mmol/l (equiv. to 1.0 g)
Theoret. osmolarity [mOsm/l]	296
pH	7.0-8.0

PL 72162/0002

POM

Use as directed by a medical practitioner.

Belief in the existence of
ghosts and spirits, etc.

B | BRAUN

B. Braun Avitum AG
Schwarzenberger Weg 77
34212 Melsungen

Lactate-based solutions:

- Pure lactate-based replacement fluid usually contains 40–46 mM of lactate.
- Lactate based solutions effectively correct metabolic acidosis in most patients.
- Lactate is metabolized on a 1:1 molar basis to bicarbonate, but in practice, the dialysis solution lactate concentration needs to be higher than dialysis solution bicarbonate to effect similar degrees of correction of acidosis.

Bicarbonate-based solutions:

- Bicarbonate-containing bags are sold as two-compartment systems, similar to those used to prepare bicarbonate-containing dialysis solution for peritoneal dialysis.
- Bicarbonate is the buffer of **choice**, and total base concentrations are typically 25–35 mM.
- Lower bicarbonate concentration solutions or bicarbonate-free solutions are also **indicated** when using regional citrate anticoagulation, because citrate is metabolized to bicarbonate by the liver.

When high-lactate solutions should be used with caution:

- Use of solutions using lactate as the primary bicarbonate-generating base has been shown to worsen hyperlactatemia in patients who have severe circulatory instability with tissue hypoperfusion, and in patients with severe liver compromise.

Citrate-based solutions:

- These fluids evolved from attempts to merge the buffering and anticoagulation properties of citrate, and the need to simplify complex regional citrate anticoagulation (RCA) protocols.
- The bulk of citrate-based fluids have to be administered prefilter to allow adequate filter anticoagulation. Forty to 60% of citrate infused in predilution mode is removed in the effluent, and the remainder is mainly metabolized by the liver into bicarbonate (1 mmol citrate yielding 3 mmol bicarbonate).

Continuous Kidney Replacement Therapies: Core Curriculum 2025

J. Pedro Teixeira, Swapnil Hiremath, Abdulghani Omar Kabli, Oleksa G. Rewa, and Edward G. Clark

Critically ill patients that require kidney replacement therapy (KRT) are among the most ill and complex patients routinely encountered in the intensive care unit (ICU). Continuous KRT (CKRT) is used across many ICUs as the therapy of choice for hemodynamically unstable patients with kidney failure. Though existing trials have not shown superior survival or kidney recovery with CKRT relative to intermittent KRT, CKRT has largely become the standard of care in developed nations for the treatment of acute kidney injury (AKI) in patients with shock, acute brain injury, acute liver failure, and other forms of critical illness. As health care systems provide an ever-widening scope of organ-support therapies to increasingly complicated ICU patients, the use of CKRT is likely to expand. In this Core Curriculum, we review the physicochemical principles of CKRT, provide a comprehensive yet practical review of when and how to prescribe CKRT, and summarize seminal trials that serve as the foundations for our approaches to timing of initiation, dosing, vascular access, and anticoagulation for CKRT. We conclude by briefly highlighting a variety of essential, yet often underappreciated, components of the provision of high-value multidisciplinary care to patients receiving CKRT, including drug dosing, nutrition, physical rehabilitation, and CKRT quality assurance programs.

Complete author and article information provided at end of article.

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Case 1

- A 28-year-old woman is admitted to the ICU after a motor vehicle collision with prolonged extrication.
- She is diagnosed with traumatic brain injury with intraparenchymal hemorrhage and multiple fractures with rhabdomyolysis with an initial creatine kinase (CK) level of 40,000 U/L
- After 48 hours, despite aggressive intravenous fluids followed by intravenous (IV) furosemide at 1 mg/kg, she is:
severely oliguric.

She has required several boluses of sodium chloride to control intracranial hypertension with a goal serum sodium of >150 mEq/L

her blood pressure is 142/91 mm Hg without vasopressor support

She is unresponsive and receiving mechanical ventilation

1- 2+ generalized edema.

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- sodium, 154 mEq/L;
- potassium, 6.1 mEq/L;
- creatinine, 3.3 mg/dL (0.9 mg/dL on admission)
- phosphate, 8.2 mg/dL,
- with CK >100,000 U/L (above limit of detection).
- (CT) shows a stable large left frontal hemorrhage with surrounding cerebral edema and persistent 4-mm midline shift.
- Her most recent ICP is 21 mm Hg.

Answer:

- Initiate CKRT with no anticoagulation and a continuous infusion of 3% sodium chloride to generate an effective CKRT bath of 152 mEq/L.



Table 1. Possible Indications for CKRT and Potential Advantages and Disadvantages of CKRT (Relative to IHD or PIKRT) in Critically Ill Patients With AKI

Classic Indications for KRT in the Setting of Hemodynamic Instability	CKRT-specific Indications: Need for KRT in the Setting of Specific Critical Care Scenarios	Advantages of CKRT	Disadvantages of CKRT
<ul style="list-style-type: none"> • Severe hyperkalemia • Severe metabolic acidosis • Diuretic-resistant volume overload • Life-threatening or severe complications of uremia (eg, bleeding in the setting of uremic platelet dysfunction, pericarditis) • Poisoning with dialyzable toxins (eg, toxic alcohols, salicylates, lithium)^a • Persistent oliguria or anuria 	<ul style="list-style-type: none"> • Intracranial hypertension or conditions associated with elevated ICP or requiring maintenance of therapeutic hyponatremia (eg, acute liver failure, acute brain injury) • Gradual correction of severe dysnatremia (eg, serum [Na⁺] < 120 mEq/L or >165 mEq/L) • Cardiopulmonary failure requiring ECMO or other mechanical circulatory support • Organ support in patients with advanced heart or liver disease unable to tolerate IHD, especially when used as a bridge to transplantation or other destination therapy • Conditions requiring continuous solute removal due to high cell turnover or cell lysis (eg, rhabdomyolysis or tumor lysis syndrome) 	<ul style="list-style-type: none"> • Less hypotension • Less effect on ICP in at-risk patients (eg, acute brain injury; acute liver failure) • Superior volume control • Superior solute control (ie, higher total daily or weekly dose) • Usually permits nutrition without restriction in protein, phosphate, or potassium • Less hemodialysis nurse support^b 	<ul style="list-style-type: none"> • Decreased (ie, slower) instantaneous clearance • Increased need for circuit anticoagulation due to extended treatment time • Increased risk of hypophosphatemia • Requires catheter placement^c • Increased risk of immobilization^d • More ICU nurse support^b • Increased overall cost

Classic Indications for KRT in the Setting of Hemodynamic Instability:

- Severe hyperkalemia
- Severe metabolic acidosis
- Diuretic-resistant volume overload
- Life-threatening or severe complications of uremia
(eg, bleeding in the setting of uremic platelet dysfunction, pericarditis)
- Poisoning with dialyzable toxins (eg, toxic alcohols, salicylates, lithium)
- Persistent oliguria or anuria

Need for KRT in the Setting of Specific Critical Scenarios

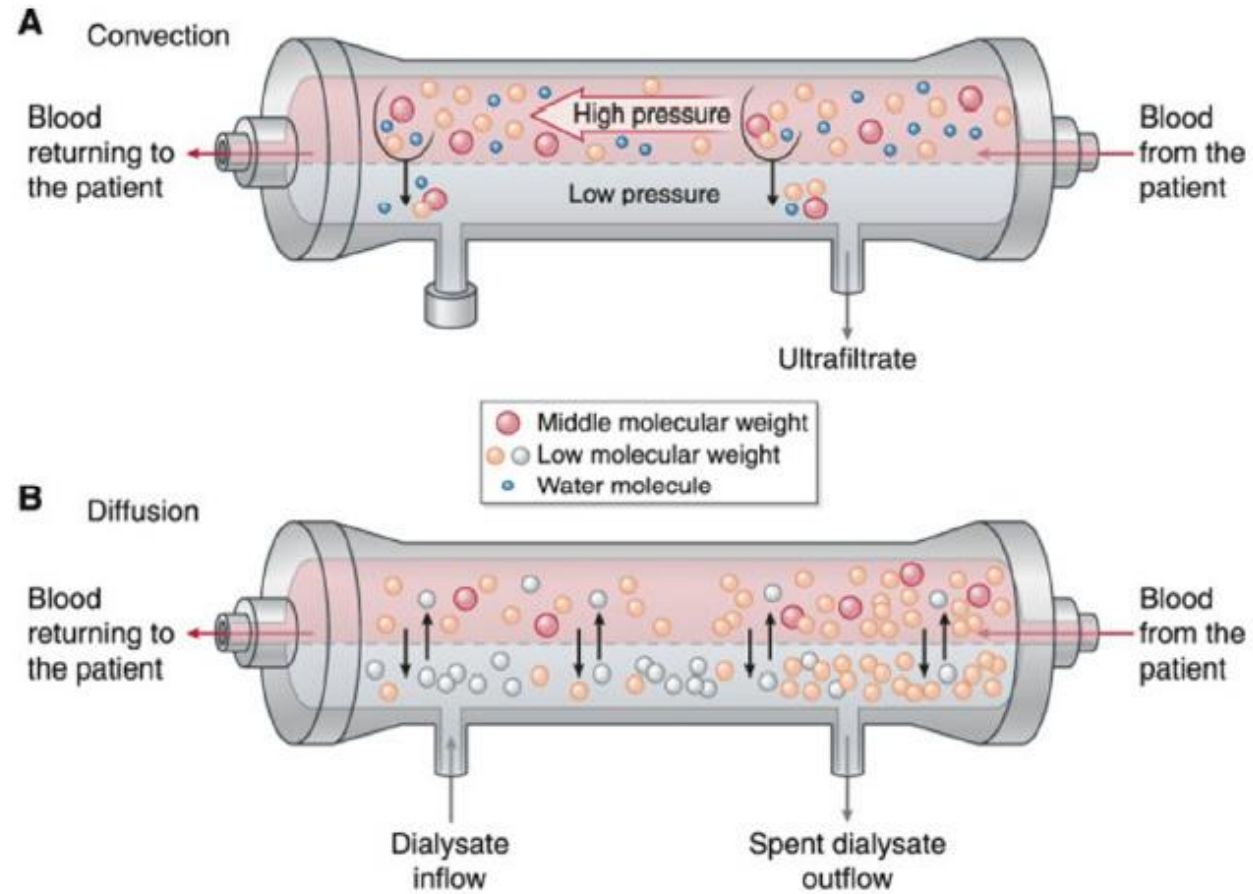
- Intracranial hypertension or conditions associated with elevated ICP or requiring maintenance of therapeutic hyponatremia (eg, acute liver failure, acute brain injury)
- Gradual correction of severe dysnatremia (eg, serum $[\text{Na}^+] < 120$ mEq/L or > 165 mEq/L)
- Cardiopulmonary failure requiring ECMO or other mechanical circulatory support
- Organ support in patients with advanced heart or liver disease unable to tolerate IHD, especially when used as a bridge to transplantation
- High cell turnover or cell lysis (eg, rhabdomyolysis)

Advantages of CKRT

- Less hypotension
- Less effect on ICP in at-risk patients (eg, acute brain injury; acute liver failure)
- Superior volume control
- Superior solute control (ie, higher total daily or weekly dose)
- Usually permits nutrition without restriction in protein, phosphate, or potassium
- Less hemodialysis nurse support

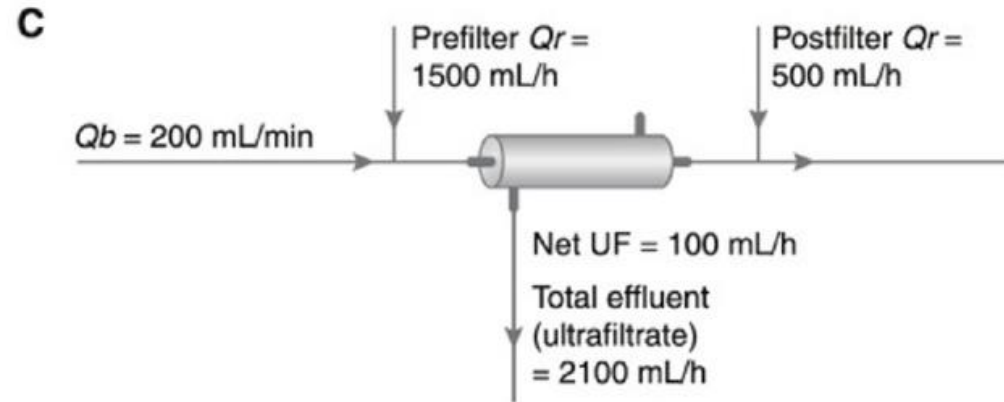
Disadvantages of CKRT

- Decreased (ie, slower) instantaneous clearance
- Increased need for circuit anticoagulation due to extended treatment time
- Increased risk of hypophosphatemia
- Requires catheter
- Increased risk of immobilization
- More ICU nurse support
- Increased overall cost

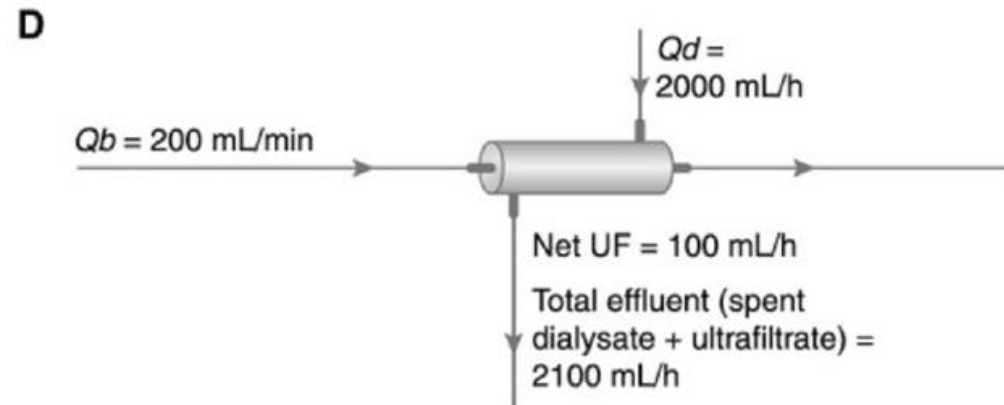


- (A) In hemofiltration, solute clearance occurs primarily by convection.
- In convection, solutes are transported across the hemofilter membrane along with plasma water as a result of a hydrostatic pressure (transmembrane pressure) generated on the blood side of the membrane. Solutes cleared by convection include urea and other small molecules along with larger “middle molecules.”
- (B) In hemodialysis, solute clearance occurs primarily by diffusion, driven by a concentration gradient across the semipermeable membrane. Small solutes in high concentration in the blood diffuse across the membrane into the dialysate, which contains either little (eg, potassium) or none (eg, urea) of the solutes being cleared.

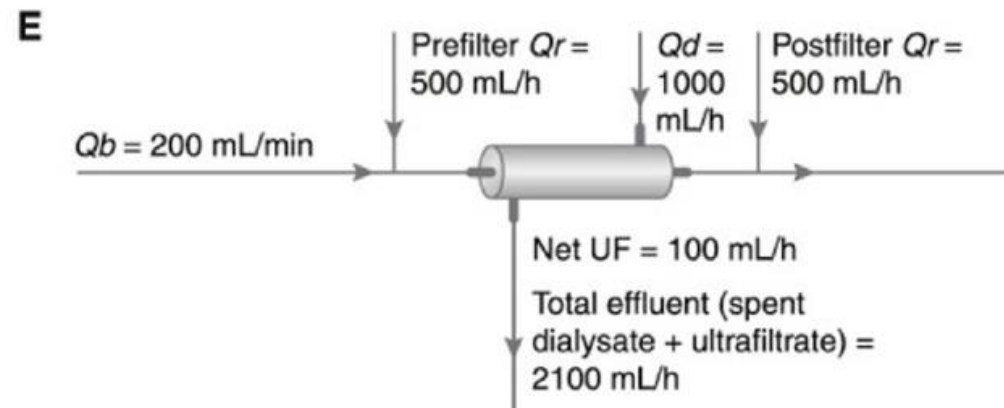
CHF



CHD



CHDF



Parameters	CVVH	CVVHD	CVVHDF	SCUF
Primary solute transport mechanism	Convection ^a	Diffusion	Diffusion + convection ^a	Convection ^a
Blood flow rate (Q_b), mL/min	100-300	100-300	100-300	100-200
Dialysate flow rate (Q_d), mL/h	0	1,000-3,000	1,000-2,000	0
Replacement fluid rate (Q_r), mL/h	1,000-3,000	0	1,000-2,000	0
Net ultrafiltration rate, (UF_{net}), mL/h ^b	0-300	0-300	0-300	50-300
Total ultrafiltration rate (UF_{total}), mL/h	1,000-3,300	0-300	1,000-2,300	50-300
Components of UF_{total}	$Q_r + UF_{net}$	UF_{net}	$Q_r + UF_{net}$	UF_{net}
Total effluent rate (Q_{ef})	1,000-3,300	1,000-3,300	1,000-3,300	50-300
Components of Q_{ef}	$Q_r + UF_{net}$	$Q_d + UF_{net}$	$Q_r + Q_d + UF_{net}$	UF_{net}

Modality	Examples of typical prescriptions for a patient weighing 70kg and with hourly net fluid intake of 50 mL/h
CVVH	Q_b 200 mL/min, $Q_{r,pre}$ 1,200 mL/h, $Q_{r,post}$ 500 mL/h, $UF_{net} = 50$ mL/h, $UF_{total} = Q_{ef} = 1,750$ mL/h (25 mL/kg/min)
CVVHD	Q_b 200 mL/min, Q_d 1,700 mL/h, $UF_{net} = 50$ mL/h, $UF_{total} = 50$ mL/h, $Q_{ef} = 1,750$ mL/h (25 mL/kg/min)
CVVHDF	Q_b 200 mL/min, $Q_{r,pre}$ 400 mL/h, Q_d 900 mL/h, $Q_{r,post}$ 400 mL/h, $UF_{net} = 50$ mL/h, $UF_{total} = 850$ mL/h, $Q_{ef} = 1,750$ mL/h (25 mL/kg/min)
SCUF	Q_b 150 mL/min, $UF_{net} = 100$ mL/h, $UF_{total} = Q_{ef} = 100$ mL/h (1.4 mL/kg/min)

Slow Continuous Ultrafiltration

- consisting of ultrafiltration without fluid replacement.
- Q_b is typically 100-200 mL/min.
- No diffusive clearance occurs because no dialysate fluid is used.
- SCUF is used to treat isolated fluid overload in patients without any need for solute clearance.
- Notably, the trials assessing SCUF performed using peripheral venous access in patients with heart failure showed no mortality benefit compared with protocolized diuretic use.

Adjusting Plasma Composition

- CKRT solutions most commonly vary in their concentrations of potassium, calcium, and bicarbonate.
- Because the total daily dose of solute clearance provided by CKRT is **higher** than that provided by thrice weekly or even daily IHD, The concentration of potassium in CKRT solutions required to control hyperkalemia is typically not as low as is required in IHD (eg, **usually 4 mEq/L** is sufficient unless hyperkalemia is severe).

Acetate or lactate were historically used as primary buffers in KRT solutions, but modern CKRT solutions are almost exclusively **bicarbonate** based, with typical bicarbonate concentrations of 22 to 35 mEq/L.

CKRT solutions:

- CKRT solutions used with RCA are usually free of calcium, which facilitates lowering the intrafilter calcium concentration.
- Additionally, these solutions usually **have lower** concentrations of bicarbonate (typically w25 mEq/L) than other standard CKRT solutions (usually w35 mEq/L)

Importantly, although **commercially** available phosphate-containing CKRT solutions are now available, traditional CKRT solutions are devoid of phosphate.

- With IHD, dialysate sodium concentration can be manipulated in a continuous fashion within a limited range (usually 130-145 mEq/L) by altering the dialysate conductivity, but generating effective sodium dialysate concentrations outside this range is impractical due to the comparatively high Q_d used in IHD.
- In contrast, though the sodium concentrations of premanufactured CKRT solutions are set, the relatively low Q_d and Q_r used in CKRT allow for easier manipulation of the effective sodium concentration, enabling slow and controlled correction of severe hyponatremia or hypernatremia.
- When correcting severe hyponatremia, lowering the effective sodium bath can be achieved by diluting the CKRT solutions by adding sterile water to the bag or replacing some CKRT solution with sterile water.

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- Generally, this approach is impractical if commercially available CKRT solutions are used because large volumes of sterile water are **not routinely stocked** in hospitals due to the safety risk of inadvertent systemic administration, and the addition and removal of fluid from premade sterile bags



Continue...

- Another approach when using commercially available solutions is to provide an additional infusion of dextrose 5% in water (D5W) either into the CKRT circuit, usually in the postfilter position, or via a separate systemic infusion.
- In hyponatremic patients, the additional volume of D5W in liters per hour to administer to achieve a targeted sodium concentration ($[\text{Na}^+]$

[Na⁺]_{CKRT} is the sodium concentration in the dialysate and/or replacement fluid (in mmol/L):

$$V_{D5W} = \left[Q_{ef} \times ([Na^+]_{CKRT} - [Na^+]_T) \right] / [Na^+]_T$$

For example:

- To target a sodium concentration of 125 mmol/L using CKRT with commercially available solutions having a sodium concentration of 140 mmol/L while using 2.5 L/h total effluent rate, V_{D5W} can be calculated as follows:

$$\begin{aligned} V_{D5W} &= (2.5 \text{ L / h} \times (140 \text{ mmol / L} - 125 \text{ mmol / L})) / \\ &\quad 125 \text{ mmol / L} \\ &= (350 \text{ mmol / h} - 312.5 \text{ mmol / h}) / 125 \text{ mmol / L} \\ &= 0.3 \text{ L / h} \end{aligned}$$

- When needing CKRT solutions with a high effective sodium concentration to gradually correct severe hypernatremia or to achieve therapeutic hypernatremia, hypertonic saline can either be added to CKRT solutions or can be infused (usually as 3% sodium chloride) into the CKRT circuit or systemically.

	Maximal Theoretical Clearance (mL/min)	Typical Approximate Total Weekly Dose (ie, Standardized Kt/V _{urea})
IHD, 3 times/wk	280	2
IHD, 7 times/wk	280	5
PD, 7 times/wk	16	2
CKRT, postfilter CVVH, 25 mL/kg/h	25	7
CKRT, prefilter CVVH, 35 mL/kg/h	28	8
CKRT, postfilter CVVH, 35 mL/kg/h	35	10
CKRT, CVVHD, 35 mL/kg/h	35	10
PIKRT	Variable	Variable
Normal kidney	90-140	16

Timing of CKRT Initiation and Discontinuation

- Starting CKRT too late may result in complications from AKI and volume overload.
 - However, starting CKRT too early may expose patients who may not have truly needed KRT to its potential harms.
 - initiation in AKI “accelerated” (earlier) versus “standard” (delayed) initiation.
- The “more delayed” strategy resulted in a trend toward increased mortality (11% higher 60-day mortality, $P = 0.07$).

Thus, no clear benefit to accelerated KRT initiation exists, yet, conversely, the results of AKIKI-2 suggest patients may be harmed by excessive delay beyond standard initiation strategies.

➤ For most patients, CKRT should be initiated in response to concrete clinical indications:

- Most commonly volume overload
- Hyperkalemia
- Metabolic acidemia unresponsive to medical therapy
- Oliguria persisting ≥ 48 -72 hours

➤ Though RCTs are ongoing, no trial data currently exist to guide de-escalation of CKRT

- Observational studies suggest that spontaneous urine output of >500 mL/day
- Diuretic-augmented urine output of >2 L/day

are reasonable criteria for consideration of KRT discontinuation in patients with AKI

Hemodynamic stability

- Ongoing need for vasopressors and higher cumulative fluid balance are associated with intradialytic hypotension in patients who transition to IHD after CKRT.
- In general, **hemodynamic stability** without vasopressor support is a commonly used trigger for consideration of transition to IHD.
- Experts suggest that **volume overload** be corrected before discontinuation or transition.

Risks of CKRT

- Hemodynamic instability from CKRT
- Infection, including catheter-related bloodstream infection
- Other risks associated with vascular access (pneumothorax, procedural bleeding, catheter-associated deep venous thrombosis,
- Clearance of trace elements, water-soluble vitamins, phosphate, amino acids/small peptides, and drugs (especially antibiotics)
- Delayed renal recovery
- Increased risk of immobilization and interference with physical rehabilitation

Patient-centered Factors

- Patient and family wishes and overall goals of care, including willingness to accept risk of long-term dialysis dependence
- Overall prognosis, including likelihood of patient survival

Health Care System Factors

- Availability of machines, disposable supplies, and nursing staff, especially during periods of strain on the health care system (eg, pandemics)
- Health care costs

Dose of Solute Clearance With CKRT

- higher doses of CKRT (35-40 mL/kg/h) had no benefit over lower doses (20-25 mL/kg/h) but were associated with somewhat higher rates of complications including hypophosphatemia and hypotension.

RCA

- In contrast, RCA produces no systemic anticoagulant effect.
- In RCA, citrate is delivered in the pre-filter segment of the circuit, typically targeting a goal blood citrate concentration of 3-6 mmol/L, and chelates calcium to generate a low intra-filter ionized calcium concentration (typically <0.4 mmol/L) to inhibit the clotting cascade.
- To reverse the effect of citrate, to replace the calcium lost as citrate-calcium complexes, and to prevent life-threatening hypocalcemia in the patient, **intravenous calcium** is continuously infused into the return limb of the CKRT circuit or via a separate systemic infusion.

- total calcium should be **monitored at least daily** to monitor for citrate accumulation.
- Protocols have been developed specifically to minimize the risk of citrate accumulation when used in the setting of absent citrate metabolism. A more useful metric than is an organic anion normally metabolized primarily by the liver, and **lactate elevations**, regardless measures of hepatic function to determine the risk of **citrate accumulation** is lactate level. Like citrate, lactate ss of cause (eg, shock or liver disease),
- Imply impaired lactate metabolism and risk of impaired citrate metabolism. As a rough guide, **serum lactate <4 mmol/L, 4-8 mmol/L, and >8 mmol/L** suggest low, intermediate, and high risk, respectively, of citrate accumulation.

RCA OR HEPARIN

The most recent and largest of these was the multicenter RICH trial, which randomized nearly 600 patients to systemic heparin or RCA and found that RCA produced

- Significantly longer filter life by 15 hours
- Fewer bleeding events
- No significant difference in mortality

Complications of CKRT

Complications Related to Catheter Placement

- Hematoma, hemorrhage, or traumatic arteriovenous fistula
- Infection (CRBSI or local soft tissue infection)
- Vein thrombosis or stenosis
- Pneumothorax or hemothorax
- Air embolism
- Visceral injury

Complications Related to Extracorporeal Circuit

- Allergic reaction to dialyzer/hemofilter or circuit tubing (rare)
- Circuit thrombosis
- Hemolysis
- Air embolism
- Hypothermia
- Thrombocytopenia

Metabolic Disturbances

- Complications of regional citrate anticoagulation:
 - ◊ Citrate accumulation, ie, citrate toxicity or citrate lock (see text)
 - ◊ Citrate/buffer excess or citrate/buffer deficit (see text)
 - ◊ Isolated hypo- or hypercalcemia
 - ◊ Hyponatremia (if using formulation containing trisodium citrate)
 - ◊ Hypomagnesemia
- Hypophosphatemia, possibly aggravating respiratory muscle weakness
- Others: hypokalemia, hypocalcemia, hypomagnesemia
- Hypoglycemia (when using dextrose-free CKRT solutions)
- Euglycemic ketoacidosis (when using dextrose-free CKRT solutions)

Others

- Hypotension, especially with initiation or net ultrafiltration
- Inappropriate (excess or inadequate) medication dosing
- Inadequate nutrition due to nonselective clearance of amino acids and other micronutrients

complications of CKRT

- Most complications of CKRT can be categorized into metabolic disturbances, access-related complications, and those related to the extracorporeal circuit . Severe allergic reactions to the hemofilter or circuit tubing and circuit-related hemolysis have been described but are rare.
- Though anecdotally appreciated for years, recent observational data confirm an association between CKRT and thrombocytopenia in patients treated with CKRT.
- However, the degree of decline is usually relatively modest (ie, 33%-50% from baseline), and CKRT as the cause of thrombocytopenia should be considered a diagnosis of exclusion given the many other potential causes of thrombocytopenia in critically ill patients.

Hypophosphatemia

- severe hypophosphatemia can induce a variety of complications, including:
 - Muscle weakness, rhabdomyolysis, and myocardial depression.
 - Prolonged mechanical ventilation or an increased need for tracheostomy.
- Options include use of phosphate-containing CKRT solutions or the pre-emptive initiation of scheduled phosphate replacement as soon as the initial AKI-induced hyperphosphatemia is corrected.

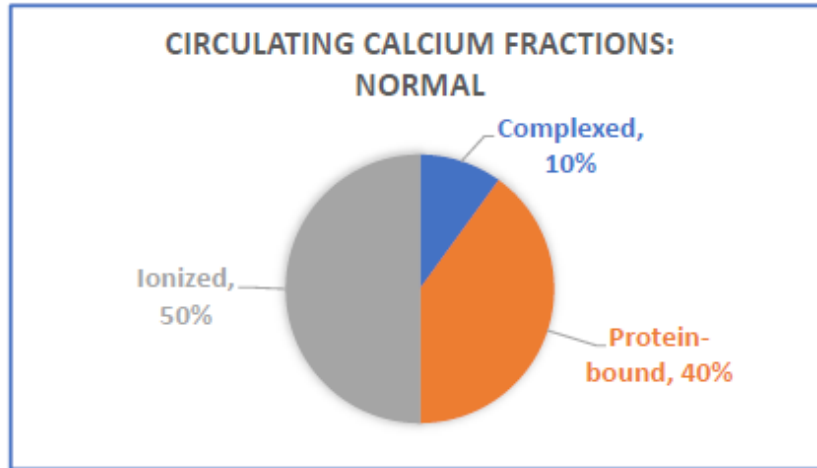
Euglycemic ketoacidosis

- Finally, in contrast to most other CKRT solutions which contain physiologic concentrations of glucose, the commercially available CKRT solutions containing phosphate are **devoid of glucose**, creating the potential for additional complications in patients not receiving nutrition or another glucose source—namely
- Hypoglycemia or euglycemic ketoacidosis.
- The latter manifests with unexplained anion gap metabolic acidosis, normal serum glucose, and ketonemia and requires treatment with infusions of dextrose and insulin.
- Thus, in patients on CKRT who develop **high anion gap acidosis** without elevated lactate or citrate accumulation, euglycemic ketoacidosis must always be considered.

Citrate accumulation

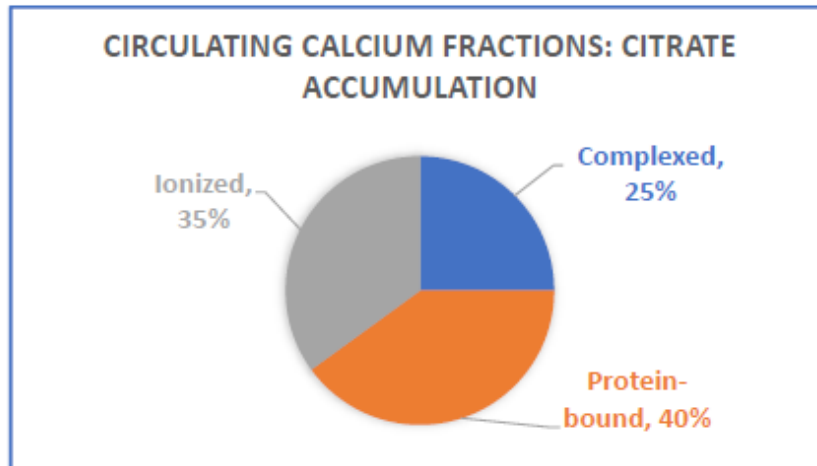
- The most feared complication of RCA is citrate accumulation, which is also referred to as citrate toxicity or citrate lock.
- As most clinical laboratories do not measure plasma citrate levels, the ratio of **total calcium (tCa) to ionized calcium (iCa)** is used as a surrogate measure of citrate levels. Because normally approximately 50% of total calcium is ionized, this ratio is usually 2-1
- Significant citrate accumulation will cause citrate-calcium complexes to accumulate, resulting in an increase in tCa and/or a decrease in iCa, with a tCa/iCa ratio of ≥ 2.5 .
- Notably, correction of the total calcium for hypoalbuminemia is not generally recommended and, based on at least 1 study, likely unnecessary

A



Calcium Levels in Differing Units: Normal				
	%	mmol/L	mEq/L	mg/dL
iCa^{++}	50%	1.25	2.5	5
tCa^{++}	100%	2.5	5	10
tCa^{++}/iCa^{++}	2-to-1			

B



Calcium Levels in Differing Units: Citrate Accumulation				
	%	mmol/L	mEq/L	mg/dL
iCa^{++}	35%	0.96	1.92	3.84
tCa^{++}	100%	2.75	5.5	11
tCa^{++}/iCa^{++}	2.9-to-1			

Nutrition and Physical Rehabilitation

- CKRT may contribute significantly to the negative nitrogen balance that is typically seen with the inflammatory insults and catabolic states characteristic of critical illness. In contrast with intact kidneys, in which amino acids and small peptides are filtered at the glomerulus but fully reabsorbed by the proximal tubule, CKRT can lead to the nonselective loss of 10-20 g of amino acids daily along with other water-soluble micronutrients.
- Though additional data are needed, this amino acid removal by CKRT may plausibly aggravate ICU-acquired weakness.

Continue....

- To overcome this loss, daily nutritional targets of 25- 35 kcal/kg total calories and 1.5-2.5 g/kg of protein are recommended in patients receiving CKRT.
- Likewise, though many perceive CKRT to be a barrier to mobilization, observational studies have shown that cautious physical rehabilitation is feasible and safe in patients undergoing CKRT, and nephrologists should advocate that physical therapy is provided to CKRT patients who are otherwise appropriate candidates for early mobilization.



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Continuous Renal Replacement Therapy

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Continuing Education Activity

Go to:

Managing severe acute kidney injury demands a nuanced understanding of renal replacement therapies. Among the options, including intermittent hemodialysis and prolonged intermittent renal replacement therapy, continuous renal replacement therapy (CRRT) emerges as a preferred modality in various clinical scenarios. A multinational study revealed that CRRT is utilized in 75.2% of intensive care unit visits. CRRT, involving continuous solute removal and fluid balance techniques, employs 3 distinct modalities: continuous venovenous hemofiltration, continuous venovenous hemodialysis, and continuous venovenous hemodiafiltration. CRRT's indications span volume overload, electrolyte disturbances, and uremia complications. This intervention excels in managing hemodynamically unstable patients, allowing controlled fluid management and mitigating risks associated with rapid solute changes.

Collaboration in CRRT implementation involves critical nephrology specialists, a proficient CRRT nursing team, and

Indications

RRT indications in the acute setting often include the same dialysis indications in acute renal failure:

volume overload, acidosis, electrolyte disturbances like hyperkalemia, and uremia complications.

The patient's hemodynamics drive the modality choice. CRRT is employed when signs of hemodynamic instability become evident due to reduced blood flow

This modality allows more controlled fluid management and greater net fluid removal than IHD.

- Additionally, CRRT can achieve better solute control
- in patients with high catabolic rates or have rapid cell breakdown, as in individuals with tumor lysis syndrome or rhabdomyolysis.

Continue.....

- IHD is the modality of choice for life-threatening hyperkalemia and acute intoxications because of its ability to remove toxins and excess potassium through higher blood flow rapidly.
- Ammonia and similar toxins can be difficult to remove from the body because they do not bind well to proteins and are distributed throughout a large volume. Intermittent hemodialysis may be used for initial rapid removal, followed by CRRT, to handle the rebound effect caused by toxin redistribution from the cells to plasma.
- This combination therapy may also be administered in metformin toxicity due to similar redistribution from the erythrocytes to the vascular space.
- Inflammatory mediators such as interleukin 1 (IL-1), IL-6, IL-8, and tumor necrosis factor- α may also be removed via convection, which is presumed to help manage sepsis and its resultant multiorgan failure

Continue.....

- CRRT can easily manage hyponatremia by adjusting the composition of the dialysate and replacement fluids or running a parallel dextrose 5% water infusion to achieve the **desired sodium**.
- **Hyperkalemia** management typically warrants more rapid blood flow, making IHD preferable. However, CRRT may be utilized if the patient is significantly **hemodynamically unstable**

Continue.....

- Patients on extracorporeal membrane oxygenation (ECMO) commonly develop AKI due to hemodynamic insults.

Contraindications

- CRRT is contraindicated when treatment outcomes demand a faster resolution than what this intervention can provide.

Conditions that may render CRRT less favorable include:

- patient directives against dialysis
- vascular access challenges
- Inadequate expertise
- Equipment availability
- Irreversible liver failure in patients ineligible for liver transplantation.

NON-RENAL INDICATIONS FOR CRRT

Non-renal indications for continuous renal replacement therapy

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Non-renal indications for continuous renal replacement therapy. While there is clear support for the use of continuous renal replacement therapy (CRRT) in critically ill acute renal failure patients, there are other illnesses without renal involvement where CRRT might be of value. These include sepsis and other inflammatory syndromes such as acute respiratory distress syndrome (ARDS) and cardiopulmonary bypass where removal of inflammatory mediators by hemofiltration is hypothesized to improve outcome. Adsorption appears to be the predominant mechanism of mediator elimination. However, the observed hemodynamic improvement can, at least partially, be attributed to a reduction of body temperature or to fluid removal, and the evidence for a clinically important removal of proinflammatory cytokines remains limited. Continuous and therefore smooth fluid removal may improve organ function in ARDS, after surgery with cardiopulmonary bypass, and in patients with refractory congestive heart failure. Continuous removal of endogenous toxins, eventually combined with intermittent hemodialysis, is probably beneficial in inborn errors of metabolism, severe lactic acidosis, or tumor lysis syndrome.

Continuous renal replacement therapy (CRRT) is often regarded as one of the more important advances in

SYSTEMIC INFLAMMATORY RESPONSE SYNDROME AND SEPSIS

Sepsis and other inflammatory syndromes represent the most popular non-renal indications for CRRT. The underlying hypothesis is that hemofiltration removes inflammatory mediators (cytokines, complement activation products, contact activation products, arachidonic acid metabolites, and so forth) from the circulation and thereby dampens the systemic inflammatory response while preserving the local effects that are thought to be beneficial. This is indeed an attractive hypothesis. However, at this moment, the evidence for a clinically important elimination of inflammatory mediators, as well as the evidence for a beneficial effect of hemofiltration on the outcome of septic patients, remains limited [2–5].

Removal of inflammatory mediators with hemofiltration

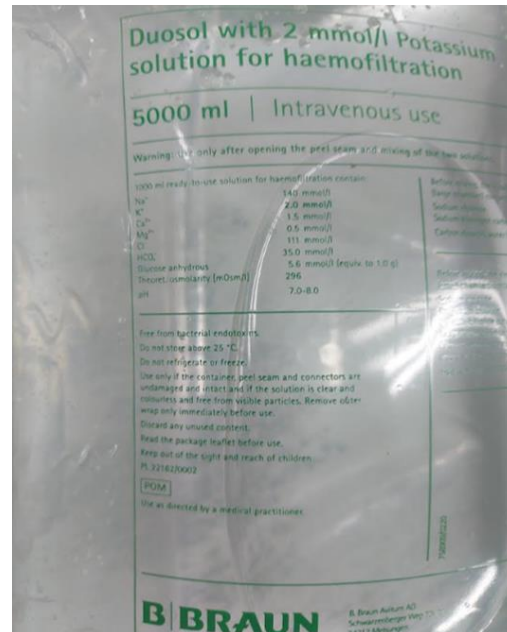
With the exception of endotoxin and the biologically active form of tumor necrosis factor (TNF), which is a trimer with molecular weight of 54,000 Da, the molecular

indications for continuous renal replacement therapy

- SYSTEMIC INFLAMMATORY RESPONSE
- While there is clear support for the use of continuous SYNDROME AND SEPSIS
- Removal of inflammatory mediators with hemofiltration
- ACUTE RESPIRATORY DISTRESS SYNDROME
- CARDIOPULMONARY BYPASS
- CONGESTIVE HEART FAILURE
- INBORN ERRORS OF METABOLISM
- LACTIC ACIDOSIS
- TUMOR LYSIS SYNDROME
- CRUSH INJURY

Nephrologist problem with crrt

- Vascular Access
- Anticoagulation
- Drug Dosing
- Nutritional Management
- Complications
- Timing and Modality





Thank you



