Drug Dosing During Continuous Renal Replacement Therapy

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ABSTRACT

Continuous renal replacement therapy (CRRT) has given clinicians an important option in the care of critically ill patients. The slow and continuous dialysate and ultrafiltrate flow rates that are employed with CRRT can yield drug clearances similar to an analogous glomerular filtration rate of the native kidneys. Advantages such as superior volume control, excellent metabolic control, and hemodynamic tolerance by critically ill patients are well documented, but an understanding of drug dosing for CRRT is still a bit of a mystery. Although some pharmaceutical companies have dedicated postmarket research in this direction, many pharmaceutical companies have chosen not to pursue this information as it is not mandated and represents a relatively small part of their market. This lack of valuable information has created many challenges in the care of the critically ill patient as intermittent hemodialysis drug dosing recommendations cannot be extrapolated to CRRT. This drug dosing review will highlight factors that clinicians should consider when determining a pharmacotherapy regimen for a patient receiving CRRT.

Unlike intermittent hemodialysis (IHD), continuous renal replacement therapy (CRRT) provides relatively constant and predictable drug clearance. The dialysate and/or ultrafiltrate flow rates that are employed can yield drug clearance similar to an analogous glomerular filtration rate (GFR) of the native kidneys. Obviously, the CRRT system does not account for the secretion and reabsorption of the native kidney, but standard first-order drug clearance equations can be used to calculate dosing regimens. Drug clearance by pumped CRRT is slow, yet constant; consequently determining drug dosing regimens should be “easier” than IHD. However, developing a drug dosing regimen for a patient receiving CRRT is not always as easy as it seems.

Depending on the mode of CRRT, the combined dialysate and ultrafiltrate flow rates have been recommended as a rough estimate of creatinine clearance (CrCl) (1). This method provides a starting point for initial drug dosing estimations. Although an estimate upon which to base empiric dosing, it may not be an appropriate basis for extended durations of drug therapy. Critically ill patients with acute kidney injury (AKI) are dynamic. Renal function, and consequently drug clearance, may be continually changing. Patient-specific, nonrenal factors such as inflammation, hepatic and other organ system function, and changes in volume status all may change profoundly in an AKI patient receiving CRRT. Drug metabolism in patients with AKI can be quite different from what is observed in healthy patients with normal renal function (2,3) or even from patients with chronic kidney disease (4). Further, renal replacement therapy itself can affect drug metabolism (5). All of these factors can have a considerable effect on drug disposition, and consequently, the appropriate drug dosing regimens.

Many CRRT-specific factors have obvious effects on drug removal. In many cases these aspects of CRRT are recognized by clinicians when determining drug dosing, and have been well described in the literature. For example, dialysate/ultrafiltrate flow rates have considerable influence on drug clearance for most drugs (6). The use of higher permeability hemodialyzers will result in considerably higher drug clearance rates than less permeable membranes, especially for drugs of middle molecular weight (MW) size like vancomycin (MW ~1450 Da) (7) or daptomycin (MW ~1620 Da) (8).

Less studied and less recognized by clinicians are some other CRRT technique-specific factors that influence drug clearance. For example, in CRRT systems with a convective component, placement of the replacement solutions has an important impact on solute clearance. Uchino et al. (9) reported that vancomycin sieving coefficient steadily decreased as the proportion of replacement solutions administered prefilter vs. postfilter, decreased.
Prefilter replacement fluid dilutes the blood that is presented to the hemodialyzer and decreases the amount of drug that is cleared. Consequently, in the trial by Uchino where the ratio of prefilter and postfilter fluid replacement was varied, vancomycin clearance was lowest when all fluids were given prefilter and almost 25% higher when the majority of the fluids were given postfilter. The prefilter or postfilter placement of replacement solutions tends to be unit dependent. Where in the CRRT system the replacement solutions are given could account for many of the differences found in published drug sieving coefficients and clearance rates. Brunet et al. (10) suggested that there was a 15–19% clearance reduction for urea and creatinine when ultrafiltrate replacement solutions were administered prefilter.

Another CRRT system-related factor that is usually not considered much by clinicians when assessing CRRT drug clearance is hemodiafilter age. Early data suggest that hemodiafilter membrane performance changes over time (11,12). Drug sieving coefficients are at their highest when the hemodiafilter is first put into use. The sieving coefficient declines with time, ostensibly because of a growing protein layer that builds up on the membrane surface and/or because of a decline in the number of unclotted hollow fibers (11,12). In the earlier days of CRRT, most CRRT centers limited the amount of time that hemodiafilters could be used. Indeed, hemodiafilter package inserts typically stated that they should not be used beyond 48 hours. However, today it is not uncommon for some units to use CRRT circuits for as long as 72–96 hours, in spite of the hemodiafilter manufacturer warnings. Likely the clearance of very small solutes like urea and creatinine are not greatly affected by increasing the lifespan of a CRRT circuit, but clearance of drugs with larger MWs may be reduced as these hemodiafilters become “clogged” over time. This aspect of CRRT drug clearance has not been studied extensively, and most published pharmacokinetic studies and CRRT case reports never mention how long the hemodiafilter was used when the measurements were conducted.

It is interesting to note how “uniform” the practice of IHD has become compared with the practice of CRRT. While dialyzers and machines may differ between units, most dialysis centers and hospitals conduct the hemodialysis procedure itself similarly. They use the same types of vascular access, treatment durations, anticoagulation, and similar blood and dialysate flow rates. This uniformity has made drug dosing relatively predictable and manageable. Many references exist for clinicians to use to make informed decisions on pharmacotherapy. In contrast, the practice of CRRT has tremendous variability. Despite many valiant efforts like the Acute Dialysis Quality Initiative, little agreement has been reached as to the optimal method to deliver CRRT. Indeed, a question as fundamental as whether diffusive or convective modalities should be used is still fiercely debated.

Clinicians often make the assumption that CRRT drug clearance is pretty much the same whatever continuous venovenous hemodialysis (CVVHD), continuous venovenous hemofiltration (CVVH), or the combination, continuous venovenous hemodiafiltration (CVVHDF) is used. With conventional CRRT at low dialysate/ultrafiltrate production rates (< 1 l/hour) this assumption is likely true. However, multiple studies show that dialytic therapies will always have inferior solute clearance to convective therapies at the same dialysate/ultrafiltrate flow rates. The clearance differences become larger as the dialysate/ultrafiltrate flow rates increase and as the MW of the solute increases. This rule of thumb should guide your estimates of drug clearance:

The drug clearance at any given dialysate/ultrafiltrate flow rate is:

CVVH > CVVHDF > CVVHD

The clearance difference might be relatively small for solutes like urea, but for larger molecules like vancomycin or daptomycin, the difference is marked (7,8).

**Patient Case**

An approach toward drug dosing is best conveyed by examining a patient case and discussing how one would determine various drug regimens. Take, for example, an elderly male who developed AKI following abdominal surgery. CVVHDF with prefilter replacement fluid was initiated to remove edema and to correct an electrolyte imbalance. The CRRT flow rates are set at a blood flow of 200 ml/minute, dialysate flow of 2 l/hour (33 ml/minute), and an ultrafiltrate production rate of 500 ml/hour (8 ml/minute). His pertinent dosing information includes his weight 232 lbs (105.5 kg) with an estimated preadmission “dry” weight of 188 lbs (85.5 kg) and the fact that the patient is oliguric (200 ml urine/24 hours). An empiric estimate of creatinine clearance attributed to CVVHDF is ~41 ml/minute (dialysate flow rate + ultrafiltrate production rate). This estimate is likely a slight overestimation by 6–8 ml/minute because the replacement solution is being given prefilter. Obviously, if CRRT were stopped for any period of time, this patient’s estimated clearance would be < 10 ml/minute, as evidenced by his low urine output. At the initiation of CVVHDF, the patient has spiked a fever. The medical team is suspecting a bacteremia and wants to initiate a broad spectrum empiric therapy regimen of gentamicin, daptomycin, and “Drug X.”

Understanding potential drug removal characteristics by these continuous therapies can simplify one’s approach to drug dosing. Knowledge of each drug’s pharmacokinetic and pharmacodynamic properties enables the clinician to empirically drug dose but also adjust the drug’s dose as the patient’s status changes. The clinician must also establish a goal for each therapy and monitor for therapeutic success, failure, and adverse effects.

**Gentamicin**

Gentamicin has a MW ~477 Da and a volume of distribution ($V_d$) of ~0.25 in healthy adults and protein binding rate is only ~10–20% (13). Because of these characteristics, it is easily removed from the blood by CRRT without much regard to hemodialyzer membrane type or size. The so-called “once daily
aminoglycoside regimens" (14,15) take advantage of gentamicin’s concentration-dependent antibacterial activity and post-antibiotic effect, but these regimens are reserved for patients with estimated GFR > 50 ml/minute. Consequently, in this patient, traditional gentamicin dosing methods would be used to achieve peak serum concentrations of 6–8 mg/L. This peak concentration should be sufficient to exceed the minimum inhibitory concentration (MIC90) of the organism.

Gentamicin has serious adverse effects associated with drug accumulation, so it is imperative that an ideal dosing regimen includes a period of time where the drug concentration is below 1–2 μg/L. Based on this patient’s body weight, Vd and estimated drug peak concentration, a loading dose of 2–2.5 mg/kg followed by 1–1.5 mg/kg every 24 hours would be an appropriate initial dose. As edema is removed and the patient approaches his dry weight the gentamicin dose must be adjusted. Whenever possible, serum drug concentrations should be obtained once the drug has had time to distribute. Generally, a peak concentration should be obtained 30 minutes after the end of the infusion and a trough concentration before the next scheduled dose begins. Usually, serum concentrations would not be necessary after the first dose of gentamicin but this patient has a variable concentration-dependent antibacterial "tissue" indicates that unlike gentamicin and daptomycin, other drugs with this pharmacodynamic profile include β-lactam antibiotics, for example.

What information regarding Drug X do you need to identify to dose this drug? Each drug’s distinctive pharmacokinetic and pharmacodynamic property must be considered when dosing to obtain the best possible outcomes. An understanding of the pharmacodynamic goals of the drug and then applying the drug’s pharmacokinetics will assist in developing a dosing strategy for drugs with limited available data. Often, all of the pertinent pharmacokinetic data can be found in the manufacturer’s package insert. Drug characteristics such as Vd, MW, and protein binding rate are important considerations in determining how much of Drug X will be removed by CRRT. A large Vd (> 1 l/kg) decreases the likelihood that a drug will be substantially removed by hemodialysis or CRRT assuming enough time for the drug to distribute. High-flux hemodialyzers allow for the removal of drugs with a MW of ≤2000 Da. Finally, a drug that is highly protein bound is less likely to be removed than one that is mostly unbound as only the nonprotein bound drug can cross the hemodialyzer membrane. The unbound fraction of the drug (1-protein binding rate = unbound fraction) multiplied by the dialysate/ultrafiltrate rate yields the drug’s clearance rate.

With this information, an estimate of the likely removal of Drug X by CRRT can be made. If the clearance of the drug by CRRT is < 25% of the total body clearance, dosing adjustment for CRRT is probably unnecessary. If CRRT looks to be an important source of drug clearance, a typical initial loading dose would be given, and maintenance doses would likely be similar to that given to a patient with a creatinine clearance of 30–50 ml/minute. If serum drug concentrations can be obtained in a timely manner to guide therapy, it should be done. As the patient’s volume status changes, doses should similarly be adjusted.

There are several reliable drug dosing resources that should be consulted including the Renal Drug Dosing Handbook (21) and software packages such as Micromedex (22). Despite these resources, realize that few drugs have been studied in this regard. One of us (Mueller) were involved in the preparation of the CRRT drug dosing recommendations in the Drug Prescribing in Renal Failure book. Our purpose was to make evidence-based CRRT dosing recommendations and guide the evidence upon which we made decisions. Interestingly, of the ~475 drugs that were evaluated for entry into the book, less than 20% of those drugs had any type of published CRRT pharmacokinetic studies conducted on them. Consequently, most of the recommendations had to be made using knowledge of pharmacokinetic and extracorporeal solute removal principles as outlined above.

Daptomycin

Daptomycin has a Vd 0.1 l/kg which is smaller than gentamicin but its MW of 1650 Da is considerably larger. It is highly protein bound (90–92%) in healthy adults (16). Daptomycin removal by CVVHDF poses an interesting problem as most of the drug remains in the vasculature (as evidenced by its small Vd), which increases the probability of its CVVHDF removal despite its high protein binding rate. Daptomycin’s large MW may slow the drugs movement across a hemodialyzer membrane but the nonprotein bound daptomycin is still small enough that it will readily cross highly permeable CRRT membranes (8). The manufacturer’s daptomycin dosing recommendations for bacteremia in patients with a creatinine clearance > 30 ml/minute is 6 mg/kg every 24 hours and every 48 hours with a creatinine clearance < 30 ml/minute (17). CVVHDF is providing an estimated creatinine clearance of ~35 ml/minute for this patient and the clinician may be tempted to select a dose of 6 mg/kg every 24 hours. Based on data from our in vitro continuous dialysis study and early results of an ongoing clinical trial, we would suggest a dose of 8–10 mg/kg every 48 hours (8,18).

Drug X

Drug X represents a drug that has recently been brought to market and for which there are no published studies of this drug’s disposition in CRRT. The “literature” indicates that unlike gentamicin and daptomycin, which have a substantial post-antibiotic effect (19,20), Drug X’s pharmacodynamic properties require that the drug concentration must be continuously maintained above the MIC90 for the drug to be effective and decrease the chance of a resistant organism forming. However, the MIC90 is not always an appropriate target. Consistent with the drug’s high protein binding and low MW, the elimination of Drug X by CRRT can be made. If the clearance rate is > 1 l/min, consider the drug’s dialysate/ultrafiltrate rate and maintenance rate.

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Each drug dosing resource used a different description of renal impairment with little agreement among categories. The authors also found several instances where one resource stated a drug required no dosage adjustment but another resource indicated the same drug as contraindicated in renal impairment. More than one dosing resource should be used when available as dosing evidence remains sparse in this area. In a follow-up letter to this paper, Aronoff (24) described drug dosing in patients with renal disease as well as we have heard it summarized:

"Despite numerous secondary sources of drug dosing information, drug prescribing in renal failure remains imprecise and relies on interpolation, extrapolation, and estimation."

**References**


