Longitudinal Hemodiafilter Performance in Modeled Continuous Renal Replacement Therapy

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Abstract

Background/Aims: With advanced anticoagulation, many institutions operate continuous renal replacement therapy (CRRT) circuits longer than manufacturers’ recommendations. This extended use may change hemodiafilter performance and clearance properties. However, hemodiafilter performance over time has not been assessed. We investigated solute clearance over time in modeled CRRT. Methods: In vitro continuous hemofiltration (CH) and continuous hemodialysis (CD) were operated for 48 h using AN69 polyacrylonitrile, cellulose triacetate, F70 polysulfone, and Optiflux F160NR polysulfone hemodiafilters with citrated bovine blood. Urea, creatinine, gentamicin, vancomycin, and albumin clearances were assessed in CH (ultrafiltration rates = 1 and 3 l/h). Clearances of urea, creatinine, gentamicin, and albumin, were assessed in CD with dialysate flow rate of 2 l/h. Results: Solute CH clearances were significantly higher at 3 l/h. Only creatinine and gentamicin clearances were affected by time. Creatinine CD clearance significantly declined at 48 h for all hemodiafilters, especially polysulfone hemodiafilters. Conclusions: CRRT duration affects solute transmembrane clearance. Clinicians should consider hemodiafilter age when assessing hemodialysis dose or drug clearance.

Key Words
Clearance · Continuous renal replacement therapy · Hemodialysis · Hemofiltration

Introduction

Continuous renal replacement therapy (CRRT) is a commonly employed treatment for critically ill patients with acute kidney injury. CRRT results in excellent volume control as well as effective electrolyte and metabolic control [1]. Solutes other than electrolytes and nitrogenous waste products are also removed during CRRT. Drugs, macro- and micronutrients, and cytokines are also cleared resulting in adjustments of drug and nutrition regimens [2–6] and nutrition regimens [7–9]. In fact, the use of CRRT [primarily continuous hemofiltration (CH)] as an adjunctive treatment for septic shock [10–13] is largely a result of cytokine clearance.

When De Vriese et al. [13] studied cytokine removal in patients with septic shock by CH, they noticed that the age...
of the hemodiafilter affected cytokine clearance. Cytokine clearance was highest when it was first used, and was considerably lower after 12 h of CH. While these findings may be attributed to saturation of available cytokine binding sites on the hemodiafilter, they prompted us to consider the influence of hemodiafilter age on the transmembrane clearance of other solutes, namely drugs. Certainly, binding to hemodiafilter membranes does occur to some degree [14–16], but for most drugs, the primary method of drug clearance by CRRT is transmembrane clearance by convection and/or diffusion. As blood is exposed to the hemodiafilter, a proteinaceous secondary membrane is formed, possibly as soon as the first few minutes of CRRT [17]. The formation of a secondary membrane over the existing hemodiafilter membrane could hinder convective solute removal and reduce transmembrane clearance as the secondary membrane thickens.

In the early days of CRRT, heparin was the primary anticoagulant used and CRRT circuit duration was limited, usually because of hemodiafilter clotting. Early CH reports indicated that hemodiafilter life averaged approximately 24 h [18]. As CH technology improved, and citrate anticoagulation use increased, hemodiafilter and circuit life spans increased [19]. Hemodiafilter manufacturers often set a time limit for how long the hemodiafilter should be used before changing, typically 48–72 h. However, with hemodiafilters clotting less frequently, many institutions have extended their routine circuit changing schedule in the spirit of cost containment. Hemodiafilters are now being used for as long as 96 h in many institutions, including our own. This ‘extended’ use of hemodiafilters may affect the convective and diffusive clearance capacity. Solute transmembrane clearance when a CRRT circuit is initiated may be quite different than what it is days later, but this question has not been addressed even within the first 48 h of hemodiafilter use. The purpose of these studies discussed herein was to assess transmembrane clearance at various time points with an in vitro CH and continuous hemodialysis (CD) model using 4 different hemodiafilters.

**Methods**

Our previously published in vitro models of CRRT were used for this trial [3, 4].

**Continuous Hemofiltration**

CH was accomplished using a B. Braun Diapact (Bethlehem, Pa., USA) machine that was primed with heparinized saline according to the manufacturer’s instructions. Formed ultrafiltrate was recirculated back into the blood reservoir so that the volume of blood never changed for the duration of an experiment, and no solutes were lost due to transmembrane clearance. Four different hemodiafilter types were assessed in this study:

1. M60 AN69 hemodiafilter (Gambro, Lakewood, Colo., USA), surface area = 0.6 m²;
2. CT190, cellulose triacetate (Baxter, Deerfield, Ill., USA), surface area = 1.9 m²;
3. F70, polysulfone (Fresenius, Waltham, Mass., USA), surface area = 1.7 m²;
4. O160, Optiflux F160NR, polysulfone, (Fresenius, Waltham, Mass., USA), surface area = 1.6 m².

In a water bath, 1 liter of citrated bovine whole blood (Animal Technologies, Tyler, Tex., USA) was warmed to 37 °C and continually stirred. A ‘cocktail’ of solutes was added to the blood reservoir so that their clearance could be assessed periodically (table 1).

<table>
<thead>
<tr>
<th>Solute</th>
<th>Molecular weight, Da</th>
<th>Initial blood concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Urea</td>
<td>60</td>
<td>75 mg/dl</td>
</tr>
<tr>
<td>Creatinine</td>
<td>112</td>
<td>10 mg/dl</td>
</tr>
<tr>
<td>Gentamicin</td>
<td>470</td>
<td>10 mg/l</td>
</tr>
<tr>
<td>Vancomycin</td>
<td>1,448</td>
<td>50 mg/l</td>
</tr>
<tr>
<td>Bovine albumin</td>
<td>approx. 65,000</td>
<td>3 g/dl (already present in bovine blood at 3 g/dl)</td>
</tr>
</tbody>
</table>

Blood with the added solutes was allowed to flow through the circuit at 200 ml/min in bypass mode (no ultrafiltrate formed) for 20 min prior to starting the experiment to permit initial secondary membrane formation. After 20 min of blood recirculation, ultrafiltration was turned on and the experiment ran for 48 h. Two different ultrafiltrate production rates were to be tested, 1 and 3 l/h; however, an ultrafiltrate rate of 3 l/h could not be accomplished with the smaller M60 hemodiafilter. Consequently, the M60 was only tested at an ultrafiltration rate of 1 l/h. Five experiments were performed for each experimental condition, each with a new, nonreused hemodiafilter. Consequently, 35 total experiments were performed; 5 for each hemodiafilter (4 types) and ultrafiltration rate (1 and 3 l/h; except for the M60 which was only assessed at 1 l/h).

**Continuous Hemodialysis**

CD was performed in a method similar to the CH model described above. The only differences were that calcium-free dialysate was run in a single-pass mode, countercurrently at a flow rate of 2 l/h with a blood flow rate of 200 ml/min. The experiments were run so that the blood volume remained unchanged and no ultrafiltrate was formed. In the CD experiments, the CT190, F70, and Optiflux F160NR polysulfone were tested. The same solutes were tested in the blood with the exception of vancomycin, which was not included.

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**Table 1. Study solutes: molecular weights and initial blood concentrations**

Blood Purif 2011;32:82–88
Blood samples were obtained from the prefilter blood access port. Ultrafiltrate/dialysate samples were taken from the ultrafiltrate/dialysate access port simultaneous to the blood sampling. Samples were taken 1, 24, and 48 h after CH or CD was instituted. Blood samples were allowed to clot and were then centrifuged and the plasma immediately extracted. Plasma and ultrafiltrate/dialysate samples were labeled properly, placed in cryovials, and immediately frozen at –80 °C. All study solutes were assayed in batches using a COBAS Integra 400 Plus Auto analyzer (Roche Diagnostics, Indianapolis, Ind., USA).

**Calculations**

Solute sieving coefficient (SC) was calculated at each time point by dividing the ultrafiltrate/dialysate solute concentration by the prefilter plasma concentration. Solute clearance was calculated by multiplying SC by ultrafiltrate/dialysate production rate.

**Statistical Analysis**

The statistical analysis for the experiments was assessed using a three-step analysis. First we fitted a linear mixed model, with hour as a ‘fixed effect’ and filter as a ‘random effect’. This allows the correlation of observation for the same filter over time to be...
taken into account appropriately, and it allows for and controls for variability between filters. Second, we also used a likelihood ratio $\chi^2$ test to examine the significance of the random variability between filters. Finally a post hoc comparison of the means for each time point was carried out using the Tukey-Kramer adjustment for multiple comparisons. Any measurement with a $p$ value of 0.05 or less was considered to be statistically significant.

**Results**

All CH and CD circuits survived to 48 h at each of the filter and ultrafiltrate/dialysate combinations tested. Mean SC and resultant clearance rates varied by filter and solute, with the exception of albumin which exhibited a SC ranging from 0.02 to below the detection limit of the assay.

**Continuous Hemofiltration**

In general, solute clearances were significantly higher when an ultrafiltration rate of 3 l/h was used compared to 1 l/h. Figures 1–4 illustrate a general trend showing a decline in clearance from initial values at 3 l/h. In no case did any solute show a statistically significant increase in clearance from 1 to 48 h. Creatinine transmembrane clearance across the M60 hemodiafilter declined significantly from 1 to 48 h with a 1 l/h ultrafiltration rate ($p < 0.05$). At an ultrafiltration rate of 3 l/h, the creatinine transmembrane clearance declined significantly from 1 to 48 h with the O160 ($p < 0.001$) and CT190 ($p < 0.05$) hemodiafilters.

All hemodiafilters showed a significant decline in gentamicin transmembrane clearance from 1 to 48 h at 3 l/h of CH ($F70 = 0.05$, O160 $\leq 0.0005$, CT190 $< 0.05$). No significant differences in clearance over time were noted for urea or vancomycin. Differences in solute clearances between filter types were noted for the 4 solutes at all 3 time points assessed. In general, the M60 hemodiafilter tended to have the lowest solute clearance; however, no generalizations beyond this observation could be made.
Continuous Hemodialysis

Figures 5–7 show solute clearances for each of the 3 tested hemodiafilters. No significant changes occurred over time or between filters for urea (fig. 5). Creatinine transmembrane clearance declined significantly for all 3 filters from 1 to 48 h (fig. 6). The decline in creatinine clearance was most profound with the polysulfone hemodiafilters (F70, O160) compared to the cellulose triacetate hemodiafilter (CT190) from 1 to 48 h (p < 0.001). Gentamicin clearances did not differ significantly between hemodiafilters but transmembrane clearance for the polysulfone hemodiafilter O160 increased slightly from 1 to 48 h (p < 0.05).

Discussion

This study assesses the influence of time on solute clearance in CRRT in vitro models. Multiple solutes were added to citrated whole bovine blood and samples were obtained from the prefiter blood access port. We found that solute transmembrane clearance declined with time, particularly for creatinine, and in the case of CH, gentamicin. This was more apparent at the higher CH flow rates of 3 l/h, suggesting that membrane factors such as protein layering and clogging may be more significant at higher ultrafiltration rates.

The issue of hemodiafilter performance over time is one that has not received much scrutiny, possibly because it is difficult to assess and may be dependent on which solute clearance is measured. In practice, urea clearance is used to describe hemodialyzer performance in hemodiafilter manufacturer package inserts and is the gold standard for dialysis adequacy (KT/V urea) calculations. Clinicians frequently use urea clearance as a surrogate marker of hemodiafilter performance to determine if the hemodiafilter needs to be changed. Literature and clinical support suggests the use of a ratio known as FUN/BUN, or the ratio of ultrafiltration fluid urea nitrogen to blood urea nitrogen [20, 21]. Similarly, an analogous dialysate urea nitrogen (DUN)/BUN ratio could be used for CD. Over a decade ago, Mehta and Martin [22] suggested that FUN/BUN be assessed every 12 h to determine hemodiafilter performance. Our own institution measures DUN/BUN daily in patients receiving CRRT. Some have suggested that the hemodiafilter should be changed when FUN/BUN decreases by 20–40% [20–22].

At no point in the present trial was a 20% decline in FUN/BUN or DUN/BUN observed, even after 48 h of therapy. Consequently, if this were a clinical situation using this >20% rule of thumb, we would not have changed our hemodiafilter circuit. However, in many of our circuits with ‘normal’ FUN/BUN ratios, significant decrements in creatinine and gentamicin transmembrane clearances were occurring. Even though FUN/BUN was never affected, the decrease in CH gentamicin transmembrane clearance seen within 48 h could translate to changes in drug dosing at the bedside. In the present study, gentamicin transmembrane clearance declined by 11.7% for the CT190, 16% for the O160, and approximately 28% for the F70 from 1 to 48 h. Clearance change of this magnitude could be clinically significant and may necessitate substantial changes to the gentamicin dosing regimen. An advantage of any CRRT is that continual drug clearance will result in a near-steady-state condition, similar to what is seen in patients with unchanging endogenous renal function. Dependent on when the gentamicin was dosed in the life span of the hemodiafilter, an 11.7–28% change in gentamicin clearance would certainly complicate gentamicin dosing. Gentamicin pharmacokinetic monitoring done when a hemodiafilter is new would result in higher serum concentrations as the hemodiafilter ages. In contrast, gentamicin pharmacokinetic monitoring done when a hemodiafilter is old could result in substantial underdosing when an old filter is exchanged with a new one. Many centers use hemodiafilters for 72–96 h. If the trend in gentamicin clearance reduction continues beyond the 48 h that we studied, the implications of these clearance swings could even be more profound. Similarly, over the 48 h of all the CRRT filter and flow rate combinations, creatinine transmembrane clearance declined from 4.6% (1 l/h, O160 filter, CH) to 66.1% (2 l/h, F70 filter, CD). Obviously, a clearance reduction of this magnitude could result in a substantial increase in serum creatinine. These serum creatinine increases would be unanticipated by caretakers and could further complicate management of these patients, primarily because it is unlikely that physicians would consider reduced hemodiafilter performance as the causative factor.

The concept of using a sensitive, readily available marker to predict when a hemodiafilter needs to be changed is attractive. However, by the time a hemodiafilter is so clogged that the clearance of a 60-dalton nonprotein-bound molecule (urea) is hindered by >20%, the clearance of larger solutes may have already been impaired much earlier in the CRRT circuit life span. Our results suggest that reductions in creatinine and gentamicin transmembrane clearance occur well before any...
change in urea clearance occurs. This calls into question the clinical utility of FUN/BUN (or DUN/BUN in the case of CD). Ultrafiltrate creatinine/serum creatinine might prove to be a better marker than FUN/BUN, as changes in ultrafiltrate creatinine/serum creatinine occurred within 24–48 h in our experiments. Whether ultrafiltrate creatinine/serum creatinine would be a predictive marker of overall hemodiafilter performance has yet to be studied. Creatinine is also a relatively small solute (112 Da) and its clearance may not be representative of overall filter performance.

Because each of these solutes appeared to be affected differently by CRRT duration with each hemodiafilter and CRRT modality, it seems that trying to simplify hemodiafilter performance to a single ratio (i.e. FUN/BUN) may not be as useful as first thought. Indeed, initial recommendations for FUN/BUN monitoring were made in the early to mid-1990s [20, 22] when CRRT machines were rather rudimentary. Contemporary CRRT machines now continually monitor arterial, venous, ultrafiltrate/dialysate and transmembrane pressures within the circuit. It is likely that some of these measures are better predictors of membrane performance and impending circuit failure.

These results may have a bearing when considering issues like actual delivered dose of dialysis, which is often assumed from the time that CRRT is running, or from volume of ultrafiltrate/dialysate produced per unit time. Indeed, a decreased efficiency of urea removal by CRRT has recently been identified as a more significant contributor to a decrease in the delivered dose of CRRT than was CRRT circuit downtime [23]. Actual delivered dose of solute clearance no doubt differs by solute, but, as shown in the present study, may also differ with CRRT duration. If this is true, then these concepts should be considered when interpreting studies that have attempted to compare patient outcome with a particular dose of CRRT as well as pharmacokinetic trials that assess CRRT drug clearance. Changes in actual solute/drug clearance over time may influence interpretations of these studies, and in the case of drugs, may result in the potential for differences in drug dosages.

Limitations

One of the limitations of our in vitro study design is that we did not assess the extent of specific solute adsorption nor did we collect postfilter samples. Prior to the experiment, we expected that the largest solutes would show a decrease in clearance over time earlier and to a greater extent than would urea. Our reasoning was that the larger solute’s transit through hemodiafilter pores would be most sensitive to a narrowing in pore size that might be seen as a hemodiafilter clogs. Consequently, we should have seen the biggest changes in vancomycin transmembrane clearance in this experiment, but we did not. Changes in clearances were far more evident for gentamicin and creatinine than for vancomycin. Whether this has to do with some physicochemical property of vancomycin compared to the other tested solutes or some other reason is unclear. Uchino et al. [24] reported that ‘protein fouling’ in high-volume hemofiltration (6 l/h) might reduce vancomycin SC with CH duration. We did not perform CH at 6 l/h, and did not find evidence of substantial vancomycin SC decline after 48 h of CH conducted at 1 and 3 l/h.

Conclusion

The transmembrane clearance of most solutes in CRRT is likely influenced by duration of therapy. Our data suggest that while urea clearance is maintained for 48 h of modeled therapy, the clearance of other solutes may be altered. More sensitive markers need to be determined to assess changes in hemodiafilter performance. These data need to be validated in clinical trials.

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Disclosure Statement

The authors claim no conflict of interest.
References


