Purity and Stability of Online-Prepared Hemodiafiltration Fluid after Storage

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Abstract

Background/Aims: Previous studies have suggested that online hemodiafiltration (OL-HDF) fluid can be used as dialysate for continuous renal replacement therapies, and thus HDF costs can be reduced. The aims of this study were to determine the purity of OL-HDF fluid and to verify the stability of the electrolyte composition and acid-base balance during its storage. Methods: OL-HDF fluid was collected in 70 individual bags and stored for up to 7 days. The following tests were performed daily in 10 bags: natural visible precipitation (macrocrystallization), sample collection for chemical analysis and fluid culture, limulus amebocyte lysate endotoxin test, standard culture of NALGENE\textsuperscript{®} filters after passing of the fluid, and molecular analysis of bacterial DNA. Results: The values of pH and pCO\textsubscript{2} showed a significant change starting at 24 h (p < 0.001); after 72 h, their values were beyond the measurable range. Coefficient of variation for pCO\textsubscript{2} was as high as 25.7%. Electrolyte composition (Na\textsuperscript{+}, K\textsuperscript{+}, Cl\textsuperscript{−}, Ca\textsuperscript{2+} and glucose) showed a statistically significant difference over time (p < 0.05); however, their coefficients of variation were low (1.7, 1.4, 0.6, 2.3 and 0.9%, respectively), which might not be considered clinically significant. Negative results were obtained at all points by fluid and filter cultures, endotoxin test and molecular analysis. No macrocrystallization was observed at any time point. Conclusions: We demonstrate the microbiological purity of OL-HDF fluid stored for up to 7 days. The electrolyte composition was stable, except for a relevant change in pCO\textsubscript{2} and consequently in pH (first noted at 24 h), emphasizing the need to reassess the acid-base balance in multilayer plastic bags in future studies.

Introduction

Acute kidney injury (AKI) is a common complication in hospitalized patients, with an incidence around 5% \cite{1–4}. Mortality in patients with AKI remains high, reaching up to 50% in severely ill patients \cite{1–4}. Approximately 5–6% of critically ill patients require extracorporeal renal support during their intensive care unit stay \cite{4}. Continuous renal replacement therapies (CRRT) have been increasingly used in critically ill patients with AKI mainly under the statement of a better hemodynamic stability, although they result in higher costs, basically due to the extracorporeal circuit, the fil-
Purity and Stability of OL-HDF Fluid

Materials and Methods

OL-HDF Fluid Preparation
OL-HDF solution was prepared with two different dialysis machines, AK 200 ULTRA (Gambro AB, Lund, Sweden) and 4008H (Fresenius, Homburg, Germany). The solution produced was a replication of the standard fluid used during OL-HDF. These dialysis machines use acid and bicarbonate concentrate mixed with heated water (treated by reverse osmosis in our dialysis unit). The predefined standard composition consisted of Na⁺ 138 mmol/l, K⁺ 2 mmol/l, HCO₃⁻ 32 mmol/l and Ca²⁺ 1.5 mmol/l.

Fluid Collection
The whole fluid was passed through the NALGENE® in-line microorganism filter system designed for removal of particles or microbial contaminants from liquids (Nalgene Nunc International, Rochester, N.Y., USA). The membrane of the filter (cellulose acetate) is characterized by a pore size of 0.2 μm and a diameter of 47 mm. All components of the system were sterilized by autoclave (120°C for 20 min). The filter was introduced between the dialysate fluid line and the collecting bag with a sterile technique (fig. 1). Sterile 3-liter ethylvinylacetate bags (SIFEVA, Fresenius Kabi, Isola della Scala, Italy), which were normally used for total parenteral nutrition, were hermetically closed and clamped, and stored at room temperature for up to 7 days. All bags were collected at the same time and under the same conditions.

OL-HDF Fluid Analysis
A total of 70 bags were planned to be stored, using 10 bags per day for microbiological and chemical analysis (time 0, 24, 48, 72, 96, 120, 144 and 168 h), plus 3 extra measurements at 3, 6 and 9 h after its collection only for chemical analysis.

The protocol for fluid analysis consisted of the following order, of (1) assessment of daily natural visible precipitation (microrystallization), (2) sample collection for chemical analysis and fluid culture, and (3) passage of the total fluid remaining through the NALGENE in-line microorganism filter system to complete the microbiological (standard cultures and endotoxin test) and molecular analysis.

Chemical Analysis
The analysis for glucose, calcium, sodium, potassium and chloride were obtained by the SYNCHRON CX3® System (Beckman Instruments, Fullerton, Calif., USA) using the Jaffé method. Acid-base analysis was obtained through the Rapid point 405® System (Bayer Diagnostics, Sudbury, UK), which directly determines pH and pCO₂ using a potentiometric method, and HCO₃⁻ is calculated according to the equation derived from Henderson-Hasselbalch as recommended by the Clinical Laboratory Standard Institute (formerly the National Committee for Clinical Laboratory Standards) [10]. For pH, the detectable range is 6.5–7.8 and for pCO₂ 10–150 mm Hg.

Microbiological Analysis
Fluid cultures for aerobic and anaerobic bacteria were obtained using the BacT/ALERT® System (SA-SN; BioMérieux, Marcy l’Etoile, France) with a colorimetric sensor, which detects microorganisms by tracking CO₂ production. The BacT/ALERT is a fully automated, dual temperature system, which may be used to conduct sterility testing at 35°C. The membranes of the filter (at T0 and for each day of storage) were placed in tryptone glucose extract agar plates (Agromics, Italy) and incubated at 22°C for 7 days [11–13]. Endotoxins from Gram-negative bacteria were detected and quantitated using the kinetic limulus amebocyte lysate endotoxin test (LAL Pyrotell® Single Test Vial, Associates of CAPECOD, East Falmouth, Mass., USA) [14, 15]. The Pyrotell sensitivity used for this study was 0.125 EU/ml.

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Molecular Biology Analysis

The molecular method developed for the detection and identification of bacterial DNA is based on characteristics of the bacterial 16S rRNA gene. This gene consists of highly conserved regions and mixes with highly variable regions, which allow phylogenetic analysis [16–18].

Nucleic Acid Preparation

Genomic DNAs were prepared by standard methods. Filter membrane was treated overnight in a lysis buffer containing 10 mmol/l Tris-HCl (pH 8.3), 50 mmol/l KCl and proteinase K to a final concentration of 0.5 g/l, and Nonidet P-40 (Roche Applied Science, Mannheim, Germany) at 55 °C. After boiling and centrifugation of the solution, the supernatant was used as template.

PCR Amplification and Purification

DNA was amplified by PCR. In this study, the primers used for amplification of 16S rRNA were 355F (5'-CCTACGGGAGGCAGCAG-3') and 910R (5'-CCCGTCCATTCCTTTGAGTT-3') [19]. Approximately 0.2–1 μg of genomic DNA were amplified in a 50-μl reaction. Cycler conditions consisted of 5 min at 95 °C, and then 35 cycles at 95 °C for 45 s, 53 °C for 45 s and 72 °C for 45 s. The final extension step was for 7 min at 72 °C. The amplification products were checked in 3% NuSieve 3:1 agarose (Cambrex Bio Science, Rockland, Me., USA) with 5% GelStar staining (Cambrex) using standard techniques. The PCR products were excised from gel and purified with Wizard SV Gel and PCR Clean-Up System (Promega Corp., Madison, Wisc., USA).

Sequencing

Direct Sanger sequencing was performed using the BigDye terminator v1.1 cycle sequencing kit (Applied BioSystems, Foster City, Calif., USA). The reaction products were purified with Centri-Sep Columns (Princeton Separation, Adelphia, N.J., USA) to remove unincorporated dye terminators. The sequences were determined by fluorescent capillary electrophoresis (ABI PRISM 310 Genetic Analyzer, Applied BioSystems) and then compared with a database library on the GenBank web site (http://www.ncbi.nlm.nih.gov/blast).

Positive Control for Microbiology and Molecular Analysis

The same microbiological and molecular analysis was performed in 4 positive control commercial bags, by adding 0.5 McFarland (0.125 optic density at 550 nm; bacterial concentration 150 × 10^6/ml) of 4 different microorganisms (Escherichia coli, Staphylococcus aureus, Pseudomonas aeruginosa and Candida albicans).

Statistical Analysis

The continuous variables were expressed as medians (25–75th percentiles) or means ± SE. Categorical variables were described as proportions. Kolmogorov-Smirnov and Shapiro-Wilk normality tests were used. Continuous variables were analyzed with ANOVA for repeated measurements or the Friedman test. Contrast analysis was performed to identify the changing point along time. Coefficients of variation were reported to compare changes among all chemical parameters. Values of p < 0.05 were regarded as statistically significant.

Results

Visible precipitation of all bags was not present at any time (T0–T168).

OL-HDF Fluid: Chemical Analysis

At the time of fluid production (T0), the mean values of pH, pCO₂ and HCO₃⁻ were 7.46 ± 0.01, 46.92 ± 1.14 mm Hg and 32.54 ± 0.32 mmol/l, respectively. The median values of the electrolyte composition of the
OL-HDF fluid were Na⁺ 137.4 (137.4–137.7) mmol/l, K⁺ 2.01 (2.01–2.02) mmol/l, Cl⁻ 118.4 (118.1–118.8) mmol/l, Ca²⁺ 6.0 (6.0–6.0) mg/dl and glucose 107.0 (107.0–107.0) mg/dl.

The values of pH and pCO₂ were outside the range starting from T96. Statistical analysis of T0–T72 showed a significant change over time (p < 0.001). Contrast analysis showed that values differed starting from T24 (fig. 2).

There was a statistically significant difference in electrolyte composition over time (T0–T168) for all electrolytes (Na⁺, K⁺, Cl⁻, Ca²⁺ and glucose; p < 0.05). Nevertheless, their coefficients of variation were 1.7, 1.4, 0.6, 2.3 and 0.9%, respectively, compared to a coefficient of vari-

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**Fig. 2.** Acid-base composition of OL-HDF fluid (pH, pCO₂ and HCO₃⁻) from T0 to T96. All values of pH and pCO₂ were out of range (box plot with means, SE and ranges).

**Fig. 3.** Electrolyte composition of OL-HDF fluid (Na⁺, K⁺, Cl⁻, Ca²⁺ and glucose) from T0 to T96 (box plot with medians, 25th–75th percentiles and ranges).
OL-HDF Fluid: Microbiological and Molecular Biological Analysis

The BacT/Alert® system did not detect any positive signal (colorimetric) in anaerobic and aerobic OL-HDF fluid cultures at any time (T0–T168). Standard cultures after 7 days of incubation for every filter membrane were free from microorganisms. Figure 4 shows a comparison with a positive control. The analysis of the endotoxin level was negative at all times (T0–T168).

In addition, bacterial DNA was never detected from the digestion of the filter membranes (T0–T168). The sequencing of the positive controls confirmed the species of inoculated microorganisms (E. coli, S. aureus, P. aeruginosa and C. albicans).

Discussion

Our study has shown that OL-HDF fluid stored for up to 7 days can maintain its purity, by recommended standards for its use as a dialysate solution. In the literature, the microbiological purity of stored OL-HDF fluid was demonstrated at the time of its production and at random measurements up to 72 h [8].

The dialysate required for CRRT must meet the definition of a fluid with no detectable levels of bacteria or bacterial-derived products. Their reference values for dialysate have been established in <10 CFU and <0.125 EU/ml, respectively [11, 20]. These standards might be even more important if OL-HDF fluid is meant to be stored instead of immediately used. Moreover, it is extremely important to monitor its purity during the time of storage in order to exclude unnecessary complications. As a complement to the standard culture methods to determine the microbiological purity of the OL-HDF fluid, we added a specific test. A qualitative molecular biology test based on the detection of bacterial DNA confirmed the negative microbial result in every filter membrane.

It has to be mentioned that storage conditions might influence the preservation of the microbiological quality of OL-HDF fluid. This should be considered if it is planned for use as emergency treatment outside hospital facilities, e.g. during natural disasters.

Although the level of purity tested in our study does not fit the definition of ultrapurity to confirm its use as replacement fluid, it can reduce the need for dialysate bags and therefore the costs of CRRT treatment. We plan to increase the sensitivity of the microbiology analysis to determine if the OL-HDF fluid could be used as ultrapure dialysate for replacement in CRRT, too.

Dialysate solutions may vary in composition and include a buffer and electrolytes to manage acid-base disorders and ion imbalances. Teo et al. [9] demonstrated that a large number of AKI patients in the intensive care unit can be safely, effectively and economically supported with continuous hemodialysis using machine-generated bicarbonate-based dialysate. However, they did not report the electrolyte or acid-base balance stability of the OL-HDF fluid during storage.
Our results demonstrated statistically significant differences among the concentrations of Na⁺, K⁺, Cl⁻, Ca²⁺ and glucose over time (fig. 3), but their coefficients of variation were considerably low, so that these differences might not be clinically significant, supporting the concept of the stability of the electrolyte composition of our OL-HDF fluid stored for up to 7 days.

In contrast, the differences among the values of pH and pCO₂ starting from 24 h pointed towards a relevant aspect in the storage of the fluid. The one-layer ethylvinylacetate bags do not represent a protective barrier for O₂, CO₂ and ultraviolet radiation. In theory, the dispersion of CO₂ and its decrement affects the balance described by the Henderson-Hasselbalch equation, and consequently reduces H⁺ concentration and increases pH. The values reported for HCO₃⁻ should ideally be verified by a quantitative technique, as ours are calculated and not directly measured from the samples. Repeating this study with next-generation multilayer ethylvinylacetate bags could prevent the loss of CO₂ and therefore contribute to acid-base stability.

Finally, in previous years, the possibility of calcium carbonate precipitation prevented the widespread use of premixed fluid bags [8, 21, 22]. In our study, visual inspection was negative at all times, and no clinically significant changes in calcium or bicarbonate concentrations were noticed. Nevertheless, different predefined prescriptions for calcium levels in OL-HDF fluid (1.5 vs. 2.5 or 3.5 mmol/l) might produce different results.

In summary, using advanced qualitative molecular biology analysis to detect bacterial DNA in addition to the standard microbiology analysis, we had demonstrated the purity of OL-HDF fluid stored in plastic bags for up to 7 days.

We verified the clinical stability of the electrolyte composition of prepared OL-HDF fluid, although a relevant change in pCO₂ and consequently in pH was noticed starting from 24 h, which underlines the need to reassess the acid-base balance in multilayer plastic bags.

This study represents an important step to establish a time limit for storing OL-HDF fluid for its use in CRRT and consequently paves the way for a sustainable alternative to the more expensive pharmacy-made solutions.

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Disclosure Statement

All the authors declare that they have no conflicts of interest or relationships with industry relevant to this work.

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