Cost-Effectiveness Analysis of High-Efficiency Hemodiafiltration Versus Low-Flux Hemodialysis Based on the Canadian Arm of the CONTRAST Study

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Abstract

Aim The aim of this study was to assess the cost effectiveness of high-efficiency on-line hemodiafiltration (OL-HDF) compared with low-flux hemodialysis (LF-HD) for patients with end-stage renal disease (ESRD) based on the Canadian (Centre Hospitalier de l’Université de Montréal) arm of a parallel-group randomized controlled trial (RCT), the CONvective TRAnsport STudy.

Methods An economic evaluation was conducted for the period of the RCT (74 months). In addition, a Markov state transition model was constructed to simulate costs and health benefits over lifetime. The primary outcome was costs per quality-adjusted life-year (QALY) gained. The analysis had the perspective of the Quebec public healthcare system.

Results A total of 130 patients were randomly allocated to OL-HDF (n = 67) and LF-HD (n = 63). The cost-utility ratio of OL-HDF versus LF-HD was Can$53,270 per QALY gained over lifetime. This ratio was fairly robust in the sensitivity analysis. The cost-utility ratio was lower than that of LF-HD compared with no treatment (immediate death), which was Can$93,008 per QALY gained.

Conclusions High-efficiency OL-HDF can be considered a cost-effective treatment for ESRD in a Canadian setting. Further research is needed to assess cost effectiveness in other settings and healthcare systems.

Key Points for Decision Makers

The CONTRAST randomized controlled trial evaluated survival of dialysis patients on hemodiafiltration (HDF) compared with low-flux hemodialysis.

A subsequent simulation of costs and health outcomes concluded that HDF is not a cost-effective treatment. However, a major limitation of the CONTRAST study was a failure to deliver high-efficiency HDF.

St. Luc Hospital of the Centre Hospitalier de l’Université de Montréal (CHUM) was part of the CONTRAST study and consistently delivered high-efficiency HDF. Analysis of this data shows that high-efficiency HDF can be considered a cost-effective treatment in a Canadian setting.

This study shows that cost effectiveness of HDF is sensitive to the type of treatment delivered and the prevailing healthcare setting.
End-stage renal disease (ESRD) is the last stage of chronic kidney disease (CKD) and requires renal replacement therapy to keep patients alive, in the form of either dialysis treatment or a transplant. As of 2012, there were 41,252 people in Canada being treated for ESRD, with 58 % (n = 23,814) on dialysis (of whom 82 % were on hemodialysis and 18 % on peritoneal dialysis) [1].

With an increasing number of patients in need of renal replacement therapy, many countries including Canada face the problem of managing ESRD in a sustainable way with limited resources. While, from a patient’s perspective, the effectiveness of the therapy is of major importance, from a payer’s and societal viewpoint, cost effectiveness also plays a critical role. Renal transplantation is clearly the more cost-effective treatment for ESRD [2], but several limitations including the lack of an adequate number of available organs prevent a more widespread application. Since dialysis programs absorb a considerable amount of resources, several studies have evaluated alternative dialysis modalities within the scope of health technology assessment [3, 4].

Recently, an international parallel-group randomized clinical trial (RCT), the CONvective TRANsport STudy (CONTRAST, NCT00205556) [5], was conducted with the aim to evaluate survival on an enhanced convective treatment, post-dilution on-line hemodiafiltration (OL-HDF), compared with low-flux hemodialysis (LF-HD) as standard of care. The trial did not show a significant survival advantage of OL-HDF [5] but suggested (and two other large trials suggested the same) that HDF had a beneficial effect on survival if a higher convection volume was provided [5–7].

From halfway through the CONTRAST trial, data were also collected on quality of life and resource consumption, the latter being multiplied with Dutch unit costs. Based on results of the trial, costs and health outcomes were simulated over a period of 5 years. The resulting cost-utility ratio based on year 2009 euros was €287,679 per quality-adjusted life-year (QALY) [8]. Given this finding, the authors concluded that, at present, OL-HDF cannot be considered a cost-effective treatment for patients with ESRD. However, a major limitation of the CONTRAST study was the failure to achieve the planned volume of post-dilution substitution fluid (19 L instead of the 24 L planned by protocol) in two-thirds of enrolled dialysis centers. As reported by Penne et al. [9], the practice patterns differed considerably between participating centers. According to a recent post hoc analysis [10], these differences result from center-specific practices regarding treatment time and blood flow rate, rather than characteristics of patients. Such differences can lead to varying levels of treatment effectiveness [5–7] and ultimately produce different cost-utility ratios. Moreover, cost differences of OL-HDF depend also on the type of monitor used and the proprietary disposable material (tubing line and filter) for the OL-HDF modality [11].

St. Luc Hospital in Montréal, Canada, which belongs to the Centre hospitalier de l’Université de Montréal (CHUM) and has an annual caseload of approximately 140 hemodialysis patients as well as 20 peritoneal dialysis patients, was part of this international RCT. For the whole duration of the CONTRAST study, the CHUM conducted OL-HDF with the highest convection volume (>24 L/session) of all participating centers. Furthermore, since the introduction of hemodiafiltration in the CHUM in November 2007 (coinciding with the beginning of the CONTRAST study), proprietary cost data were collected in addition to CONTRAST data in the hospital for an independent cost analysis of the two treatments. At the end of the follow-up period of the CONTRAST study (March 2011), the CHUM decided to continue the study and enroll additional patients in the same manner until December 2013. With information on the additional patients, the CHUM aimed to decide whether to continue its OL-HDF program and whether to extend this treatment to all of their hemodialysis patients.

The purpose of this study was to assess the cost effectiveness of high-efficiency OL-HDF defined as OL-HDF performed with an optimal convection fluid volume (that is the sum of substitution fluid volume and net ultrafiltration) compared with LF-HD for patients with ESRD in this Canadian setting.

## 2 Materials and Methods

### 2.1 Clinical Trial

The present study was conducted within the scope of the CONTRAST study [5]. CONTRAST was a non-blinded parallel-group RCT (NCT00205556) that compared post-dilution OL-HDF with LF-HD (both three times a week) on the basis of all-cause mortality. All CHUM CONTRAST study patients (n = 80) were randomized centrally by a computer-based randomization service (Julius Center University Medical Center, Utrecht, The Netherlands) into a 1:1 ratio for treatment with OL-HDF or continuation of LF-HD, stratified per participating center (permuted blocks) [5]. Allocation concealment served to avoid selection bias by concealing what treatment the next patient would receive.

Starting in 2007, cost data were collected at the CHUM. The decision to continue the study after the end of the CONTRAST study was approved by the local medical ethics review board (http://www.clinicaltrials.gov; registered number CE10.253). Written informed consent to
continue the study was requested from all trial participants who were still enrolled by December 31, 2010. Three patients, all on LF-HD, were unable to continue (censored) as their progressive cognitive impairment did not allow giving informed consent. For newly enrolled patients ($n = 50$), written informed consent was also obtained (prior to randomization). The study was conducted in accordance with the declaration of Helsinki and Good Clinical Practice Guidelines. Randomization procedure, monitoring, and data collection were kept the same as in the original trial period. Detailed information on study design and conduct as well as the inclusion and exclusion criteria can be found elsewhere [12].

2.2 Dialysis Procedure

Hemodiafiltration was performed in the online post-dilution mode. The following dialysis machines were used: 4008S with ONLINEplus™ (Fresenius Medical Care, Bad Homburg, Germany) for OL-HDF, and Integra (Gambro AB, Lund, Sweden) for LF-HD. The following synthetic dialyzers were used: Optiflux F200NR (Fresenius Medical Care, Bad Homburg, Germany) for all patients in the OL-HDF arm, and Optiflux F18 (Fresenius Medical Care, Bad Homburg, Germany) for all patients in LF-HD arm. Ultrapure quality of water and dialysis fluids, defined as $<0.1$ CFU/mL and $<0.03$ IU/mL, was used for all treatments. Microbiological tests of water and dialysis fluids were monitored each month in the CHUM, and microbiological assessments were compliant with reference quality levels in 2958 out of 2960 samples between November 2007 and December 2013.

Total duration of follow-up including the study period of CONTRAST and the period after CONTRAST was 74 months. Survival and economic data were assessed based on an ‘intent to treat’ analysis, thus analyzing patients according to the assigned treatment.

2.3 Economic Evaluation

The analysis was conducted from the perspective of the Quebec public healthcare system. Given this perspective, the study did not include productivity costs and copayments. On the other hand, the study included costs resulting from additional survival time as these are relevant from a healthcare system perspective.

We conducted a cost-effectiveness analysis using life-years gained as a measure of health benefits as well as a cost-utility analysis using QALYs. QALYs are the product of life-years and a representation of preference for different health states (preference weight or score). Preference weights are anchored on a scale from zero to one, where zero and one represent death and full health, respectively. The incremental cost-effectiveness ratio (ICER) was determined by dividing the incremental costs (i.e., additional costs) of OL-HDF compared with LF-HD by the incremental (i.e., additional) life-years or QALYs gained. We conducted both an economic evaluation for the period of the RCT (74 months) based on data collected in the trial and an analysis over lifetime, thus following recommended practice [13]. For extrapolation beyond the trial period and in line with other recently published models [8, 14, 15], we used a Markov (state transition) cohort model, thus assuming that transition probabilities do not depend on history. The Markov model used RCT data as well.

2.4 Trial-Based Economic Evaluation

2.4.1 Cost Analysis

The analysis considered costs both related and unrelated to renal disease including costs of dialysis side effects. Given the perspective of the evaluation, only direct costs were considered. Direct medical costs such as those related to dialysis staff and other medical staff, depreciation of dialysis equipment, water installation, disposables (dialyzers, blood lines, needles, etc.), routine diagnostics of patients, lab checks for water and dialysis fluid quality, and direct non-medical costs such as those related to transportation and laundry were obtained from the administration of the hospital. These costs are referred to as treatment costs. Treatment costs were assumed to be constant across time and patients. In reality, some variation may be possible (e.g., due to differences in the number of disposables), but these were considered negligible.

Pharmaceutical costs were calculated on the basis of a quarterly check of all prescriptions active during the month of the check. As a source of unit costs, we used the official price list issued by the healthcare authority in 2007 (list of medications covered by the basic prescription drug insurance plan, la Régie de l’Assurance-maladie du Québec, from June 2007) for all drugs available since the beginning of the study, and the official price at the time of introduction for all those subsequently authorized.

The cost of hospitalization was calculated using Quebec’s diagnosis-related group (DRG) system, which assigns a relative measure of the volume of resources (Niveau d’Intensité Relative des Ressources Utilisées or NIRRU) to each patient at the time of discharge. Hospitalization costs were calculated by multiplying the NIRRU for each admission by the average hospitalization cost, which has a NIRRU of 1.0 (Can$5319 in 2011/2012) [16–18].

We used a partitioned estimator proposed by Bang and Tsiatis [19] to calculate cumulative costs over the study period. This allows use of cost information from both censored and uncensored individuals. Accordingly, we
partitioned the study period into 3-month intervals, estimated costs of all patients who were alive at the beginning of the interval, multiplied interval costs by the inverse probability of not being censored until the end of the interval, and calculated average cumulative costs by summing up these costs over the complete study period and dividing them by the cohort size in the first period. As recommended by Curtis et al. [20], we used a resampling method to estimate the standard error of average cumulative costs. Costs were adjusted to Canadian dollars in 2013 using the Canadian Consumer Price Index for Health and Personal Care [21].

2.4.2 Survival Analysis

Cumulative survival was analyzed using the Kaplan–Meier method, thus accounting for censored data. The Cox proportional-hazards model was used to estimate a hazard ratio of death with 95 % confidence interval. The proportional hazards assumption was tested using the Schoenfeld residuals test.

2.4.3 Quality-of-Life Analysis

Preference weights were estimated using the 5-level version of the EuroQol-5D™ (EQ-5D-5L™) questionnaire. The EQ-5D™ measures health-related quality of life in five dimensions: mobility, self care, usual activities, pain/discomfort, and anxiety/depression [22]. Scores for the five dimensions are converted into preference weights by using country-specific value sets elicited from the general population. As the EQ-5D-5L™ is relatively new, only a few country-specific value sets are available. A Canadian value set only exists for the three-level version of the EQ-5D™ [23]. Therefore, we used a UK value set for the base case and a US value set in a sensitivity analysis.

Quality-of-life data were obtained at the beginning of the trial as well as on an annual basis. Calculation of standard errors of mean OL-HDF and LF-HD preference weights accounted for between- and within-patient variation, thus considering correlation of individual quality-of-life data over time.

\[ p \text{ values} < 0.05 \] were considered statistically significant. We discounted both costs and effects at an annual rate of 3 % [24].

2.5 Model-Based Economic Evaluation

2.5.1 Model Description

We constructed a Markov model to simulate the course of 1000 patients with ESRD (and treatment with OL-HDF or LF-HD). Patients were assumed to continue treatment with OL-HDF or LF-HD over lifetime. A Markov model is an iterative process where patients are assumed to stay in one cycle (i.e., a defined health state) for a certain time and then make a transition to another cycle. Markov models are useful when a decision problem involves risk that is continuous over time, when the timing of events is important, and when important events may happen more than once. Our Markov model contains two health states, ESRD and death. Patients may transit to death at any time. Annual mortality was set to 15 % (standard error 3 %) based on trial data (input data to the model are shown in Table 1). During each cycle, patients accumulate (quality-adjusted) life-years and costs. We chose a cycle length of 1 year for the two health states. The life-table method [25] was applied to both costs and life-years based on the assumption that transition events occur on average halfway through each 12-month cycle. The Markov model included 23 cycles to determine costs and effects over lifetime. After 23 cycles the proportion of patients still alive was less than 0.1 % in both arms.

Annual healthcare costs for each of the two groups were calculated by dividing trial-based cumulative costs by the length of the trial period. For the base-case analysis, we discounted both costs and effects at an annual rate of 3 % [24]. All calculations pertaining to the Markov model were done in Microsoft Excel (Microsoft Corporation, Redmond, WA, USA).

2.5.2 Sensitivity Analysis

To address uncertainty around the mean ICER, univariate sensitivity analyses were conducted where one variable was changed at a time while keeping all other variables constant at their mean or base-case value. We ran the analyses based on the upper and lower boundaries of the 95 % confidence interval of the mean. To assess how a simultaneous change of several variables affects the cost-effectiveness ratio, we performed a Monte-Carlo simulation, a type of multivariate sensitivity analysis. This technique runs a large number of simulations (here 1000) by repeatedly drawing samples from probability distributions of input variables. Thus, it provides a probability distribution for the output variables, that is, incremental costs, incremental effectiveness, and incremental cost-effectiveness ratios.

The annual probability of mortality and preference weights were assumed to follow a beta distribution because they are restricted to take on values between 0 and 1. The hazard ratio of death was assumed to follow normal distribution after logarithmic transformation. Cost data were assumed to follow a gamma distribution (reflecting the long right tail and restriction to positive values).
Given that the interpretation of negative ICERs is ambiguous, ICERs were transformed into net monetary benefits (NMB) [26] using the following equation:

\[ \text{NMB} = \frac{k}{C_1} \cdot \Delta E - \Delta C \]

where \( \lambda \) = maximum willingness-to-pay, \( \Delta C \) = incremental costs, \( \Delta E \) = incremental QALYs.

The decision rule used was to adopt the intervention in question if NMB was greater than zero. Given that the appropriate value of \( \lambda \) is unknown, \( \lambda \) was varied from Can$0 to Can$200,000. We generated a cost-effectiveness acceptability curve based on the distribution of NMB for each \( \lambda \). A cost-effectiveness acceptability curve allows a decision maker to consider if OL-HDF is cost effective in relation to the maximum amount a decision maker is willing to pay for a QALY. At each ceiling value for the willingness to pay for a QALY, the cost-effectiveness acceptability curve shows the probability that OL-HDF is cost effective.

3 Results

3.1 Study Population

Eighty CHUM patients were originally enrolled in the CONTRAST study by using randomized 1:1 variable blocks. Patients were requested to remain in the study at the end of the follow-up. Only three patients, all on LF-HD, were no longer able to give an informed consent (censored) due to cognitive deterioration. An additional 50 patients were enrolled between 2011 and 2013 in the same manner (randomized 1:1, variable blocks). A total of 130 patients constituted the basis for this study and follow-up was maintained until December 31, 2013. Median follow-up was 2.53 years.

The baseline characteristics of the enrolled patients were similar in the two treatment arms and are reported in Table 2. Treatment details during follow-up are provided in Table 3. Fourteen patients dropped out of the OL-HDF group because of transplantation (\( n = 6 \)), transfer to another dialysis facility (\( n = 2 \)), prolonged isolation (\( n = 4 \)), or lack of compliance (\( n = 2 \)). In the LF-HD group, 12 patients dropped out due to transplantation (\( n = 6 \)), pre-dementia (\( n = 3 \)), or transfer to another dialysis facility (\( n = 3 \)). During follow-up, 23 patients in the OL-HDF group and 23 in the LF-HD group died.

3.2 Clinical Trial

OL-HDF led to non-significant improvements both in quality of life and survival. The hazard ratio of death of OL-HDF compared with LF-HD was 0.789 (95% confidence interval, 0.440–1.418). Schoenfeld residuals test did not suggest significant overall nonproportionality (\( p = 0.448 \)). The non-significant improvement in quality of life (Table 4) can be partially attributed to differences which were present at baseline but based on data from <50% of patients (i.e., 30 and 28 patients in the OL-HDF and LF-HD arm, respectively). Below, we also report the result of the extreme-case analysis assuming no improvement in quality of life.

3.3 Trial-Based Economic Evaluation

3.3.1 Base Case

Results of the trial-based analysis by arm are shown in Table 4. Treatment costs over 1 year for OL-HDF and LF-HD were estimated to be Can$40,064 and Can$42,017, respectively (Table 4 reports costs over the complete trial period). Treatment costs per session for OL-HDF and LF-HD were estimated to be Can$268.60 and Can$256.12, respectively, resulting in an additional treatment cost of OL-HDF compared with LF-HD of Can$12.48 per session (Table 5). After treatment costs, hospitalization costs

Table 1  Data used for the model-based economic evaluation

| Variable | Base case estimate | Range tested | References |
|----------|--------------------|--------------|------------|
| Hazard ratio of death on OL-HDF compared with LF-HD | 0.789 | 0.440–1.418 | Trial data |
| Annual probability of death (LF-HD) | 15.2% | 9.5–21.0% | Trial data |
| Annual cost (LF-HD) | €59,403 | Obtained through trial-based Monte Carlo simulation | Trial data |
| Annual cost (OL-HDF) | €59,308 | Obtained through trial-based Monte Carlo simulation | Trial data |
| EQ-5D-5L™ index score for LF-HD (UK value set) | 0.64 | 0.55–0.73 | Trial data |
| EQ-5D-5L™ index score for OL-HDF (UK value set) | 0.72 | 0.65–0.79 | Trial data |
| Annual discount rate | 3% | 0–7% | [24] |

EQ-5D-5L™ 5-level version of the EuroQol-5D™ questionnaire, LF-HD low-flux hemodialysis, OL-HDF on-line hemodiafiltration
represent the second highest cost category (Table 4). Figure 1 depicts the hospitalization length and the number of hospitalizations by cause. Infectious complications were the main reason both for the number of hospitalizations and the number of days spent in the hospital.

Disregarding costs of the additional survival time on OL-HDF, discounted total per-patient costs over the trial period amount to Can$203,082 and Can$203,629 for OL-HDF and LF-HD, respectively. That is, assuming no survival benefit, OL-HDF leads to savings of Can$547. ICERs are shown in Table 6. Based on gains in quality of life, the cost-per-QALY-gained ratio is lower than the cost-per-life-year-gained ratio.

### 3.3.2 Sensitivity Analysis

Using a US value set for EQ-5D™ health states results in a cost-utility ratio of Can$35,250 per QALY gained.

### 3.4 Model-Based Economic Evaluation

#### 3.4.1 Base Case

As shown in Table 6, the model-based ICERs are of comparable relative magnitude to those of the trial-based evaluation. That is, based on gains in quality of life, the cost-per-QALY-gained ratio is lower than the cost-per-life-year-gained ratio.

| Characteristics                          | On-line hemodiafiltration | Low-flux hemodialysis |
|------------------------------------------|---------------------------|-----------------------|
| Patients, n                              | 67                        | 63                    |
| Male gender, n (%)                       | 40 (60)                   | 43 (68)               |
| Age (years) a                            | 64.2 ± 14.1               | 66.3 ± 11.0           |
| Body mass index (kg/m²) a                | 27.3 ± 6.0                | 27.6 ± 6.6            |
| Primary renal disease, n (%)             |                           |                       |
| Glomerulonephritis                       | 17 (25.3)                 | 9 (14.3)              |
| Interstitial nephropathy                 | 6 (9.0)                   | 2 (3.2)               |
| Genetic disease                          | 5 (7.5)                   | 2 (3.2)               |
| Nephroangiosclerosis                     | 4 (6.0)                   | 10 (15.8)             |
| Diabetic nephropathy                     | 28 (41.8)                 | 31 (49.2)             |
| Unknown/others                           | 7 (10.4)                  | 9 (14.3)              |
| Comorbidities, n (%)                     |                           |                       |
| Diabetes                                 | 34 (50.7)                 | 35 (55.6)             |
| Coronary artery disease                  | 32 (47.8)                 | 35 (55.6)             |
| Peripheral vascular disease              | 14 (20.9)                 | 17 (27.0)             |
| ESRD vintage (years) a                   | 3.20 ± 4.04               | 3.36 ± 3.55           |
| Vascular access, n (%)                   |                           |                       |
| AV-fistula                               | 42 (63)                   | 34 (54)               |
| Graft                                    | 5 (7)                     | 4 (6)                 |
| Catheter                                 | 20 (30)                   | 25 (40)               |
| Biochemistry                             |                           |                       |
| B2-Microglobulin (mg/L) a                | 34.0 ± 16                 | 35.4 ± 15.3           |
| Hemoglobin (g/L)a                        | 113.6 ± 15.6              | 112.0 ± 11.2          |
| Phosphate (mM/L)a                        | 1.6 ± 0.4                 | 1.7 ± 0.5             |
| Calcium (mM/L)a                          | 2.2 ± 0.2                 | 2.2 ± 0.2             |
| Parathyroid hormone (pg/mL) a            | 35.4 ± 25.6               | 42.2 ± 35.0           |
| Albumin (g/L) a                          | 35.3 ± 4.4                | 35.6 ± 3.4            |
| Treatment parameters                     |                           |                       |
| Session length (h) a                     | 3.76 ± 0.37               | 3.80 ± 0.31           |
| Dialysis frequency 2/week, n (%)         | 2 (3)                     | None                  |
| Dialysis frequency 3/week, n (%)         | 66 (97)                   | 63 (100)              |
| Blood flow (mL/min) a                    | 368.83 ± 46.23            | 350.7 ± 28.7          |
| Ultrafiltration (L/session) a            | 2.67 ± 0.49               | 2.99 ± 1.18           |
| Single-pool Kt/V urea a                   | 1.64 ± 0.37               | 1.5 ± 0.25            |

**ESRD**: end-stage renal disease

*Mean ± standard deviation*
Furthermore, the cost-utility ratio of LF-HD compared with immediate death (assuming that ESRD patients without LF-HD would die immediately) is Can$93,008 per QALY gained (306,826/3.30).

3.4.2 Sensitivity Analysis

Using a US value set results in a cost-utility ratio of Can$54,978 per QALY gained. Using a UK value set for EQ-5D™ health states and assuming the same level of quality of life in both arms increases the cost-utility ratio to Can$82,072 per QALY gained.

In the univariate sensitivity analysis, the variable with largest impact on the cost-per-QALY-gained ratio of OL-HDF versus LF-HD was the hazard ratio of death on OL-HDF compared with LF-HD. Note that the upper limit leads to a positive ratio which is based on a reduction both in costs and QALYs (Table 7).

Figure 2 shows a scatter plot of incremental costs and QALYs of OL-HDF compared with LF-HD. As displayed, the ratio of incremental QALYs to incremental costs is fairly robust as pairs of incremental costs and QALYs fall on a straight constant. In 28% of replications, OL-HDF was less effective (led to fewer QALYs) than LF-HD. The

### Table 3 Patient treatment details during follow-up (mean ± standard deviation)

| Follow-up (months) | On-line hemodiafiltration | Low-flux hemodialysis | p value for difference of means |
|--------------------|---------------------------|-----------------------|---------------------------------|
| Treatment time (min) | 1–3 225.7 ± 21.9 228.1 ± 18.5 0.498 | 10–12 226.2 ± 19.2 227.9 ± 17.5 0.702 |
|                    | 22–24 229.4 ± 19.8 234.2 ± 12.1 0.278 | 34–36 232.5 ± 13.9 225.0 ± 16.4 0.373 |
| Proportion of patients on 3 sessions/week (%) | 1–3 97.0 100 0.264 | 10–12 97.4 100 0.551 |
|                    | 22–24 96.8 100 0.544 | 34–36 87.5 100 0.571 |
| Kt/V | 1–3 1.76 ± 0.34 1.54 ± 0.44 0.02 | 10–12 1.89 ± 0.32 1.54 ± 0.23 <0.001 |
|                    | 22–24 1.96 ± 0.31 1.57 ± 0.23 <0.001 | 34–36 1.94 ± 0.31 1.59 ± 0.22 <0.001 |
| β2-Microglobulin clearance (mL/min) | 1–3 67.5 ± 13.2 | 10–12 69.1 ± 15.4 |
|                    | 22–24 65.2 ± 13.2 | 34–36 76.5 ± 22.9 |
| Blood flow (mL/min) | 1–3 385.8 ± 46.5 358.0 ± 33.7 <0.001 | 10–12 410.5 ± 52.4 367.3 ± 26.7 <0.001 |
|                    | 22–24 428.5 ± 37.7 383.1 ± 32.3 <0.001 | 34–36 415.2 ± 40.9 378.9 ± 27.0 <0.001 |
| Convective volume (L/sessions) | 1–3 26.6 ± 7.1 | 10–12 27.9 ± 7.7 |
|                    | 22–24 27.3 ± 8.6 | 34–36 22.7 ± 11.9 |
| Darbepoietin alfa dose (µg/week) | 1–3 47.6 ± 36.8 50.3 ± 48.6 0.714 | 10–12 51.0 ± 44.9 60.3 ± 49.4 0.297 |
|                    | 22–24 49.3 ± 43.4 67.1 ± 60.9 0.148 | 34–36 59.2 ± 58.1 64.9 ± 61.8 0.759 |
| Sevelamer dose (g/day) | 1–3 2.34 ± 2.93 3.42 ± 3.37 0.055 | 10–12 2.14 ± 2.88 4.05 ± 3.71 0.003 |
|                    | 22–24 1.99 ± 2.72 4.62 ± 3.69 0.001 | 34–36 2.49 ± 2.98 4.71 ± 4.70 0.062 |
| Hospitalization rate (days/patient-year) | All 16.8 ± 28.7 22.2 ± 46.1 0.416 | All 1.06 ± 1.36 1.36 ± 1.59 0.248 |
reason is that the upper bound of the 95% confidence interval of the hazard ratio of death on OL-HDF compared with LF-HD is larger than one. Figure 3 shows the cost-effectiveness acceptability curve, which considers uncertainty in cost effectiveness. The probability of cost effectiveness is largest at a willingness to pay of approximately Can$90,000 per QALY gained. Increasing willingness to pay further leads to a decrease in the probability of cost effectiveness because the joint density of incremental costs and effects in the southwest quadrant of the cost-effectiveness plane (Fig. 2) is excluded for no longer being cost effective.

3.4.3 Internal Validation

Internal model validation was checked as follows. According to Gandjour and Gafni [27], the ratio of downstream costs to effects of both interventions is the same at a single time point regardless of how effective the intervention is. When also assuming the same annual costs and quality of life, both interventions should yield the same ICER. Following this procedure we indeed obtained this result, thus confirming the internal validity of our model.

4 Discussion

Based on the results of the Canadian arm of the CONTRAST study, this study simulated costs and health benefits of high-efficiency OL-HDF versus LF-HD over lifetime. It shows a cost-utility ratio of Can$53,270 per QALY gained. As shown by the sensitivity analysis, the ratio of incremental QALYs to incremental costs is fairly robust. The cost-utility ratio is lower than that of LF-HD compared with no treatment (immediate death), which is Can$93,008 per QALY gained (Canada does not have an explicit cost-effectiveness threshold [28]). Hence, based on the assumption that ESRD patients without LF-HD would die immediately, OL-HDF weakly dominates LF-HD in the base case. That is, OL-HDF has an ICER that is lower than that of LF-HD and at the same time it is more effective than LF-HD. Based on the principle of weak dominance, OL-HDF can be considered a cost-effective treatment for ESRD in this Canadian setting. This result is also obtained based on the cost-effectiveness acceptability curve presented in Fig. 3. Using Can$93,008 per QALY gained (the ICER of LF-HD) as the willingness to pay for a QALY, the probability of cost effectiveness

### Table 4

| Variable                          | On-line hemodiafiltration | Low-flux hemodialysis | p value |
|----------------------------------|---------------------------|-----------------------|---------|
| Treatment costs, Can$            | 259,453$                  | 247,398$              | <0.001  |
| Hospitalization costs, Can$      | 70,717 (940)              | 70,219 (1013)         | NS      |
| Drug costs, Can$                 | 36,059 (112)              | 49,196 (190)          | <0.001  |
| Total costs, Can$                | 366,229 (957)             | 366,813 (1057)        | NS      |
| EQ-5D-5L\textsuperscript{TM} index score (UK value set) | 0.72 (0.03) | 0.64 (0.05) | NS |
| EQ-5D-5L\textsuperscript{TM} index score (US value set) | 0.79 (0.02) | 0.73 (0.03) | NS |

Standard errors of the mean are given in parentheses

EQ-5D-5L\textsuperscript{TM} 5-level version of the EuroQol-5D\textsuperscript{TM} questionnaire, NS not significant

\textsuperscript{a} No uncertainty in estimates was assumed

### Table 5

| Item                                                 | Cost/session OL HDF, Can$ (%) | Cost/session LF-HD, Can$ (%) |
|------------------------------------------------------|-------------------------------|------------------------------|
| Labour                                               | 155.54 (58.0)                 | 155.54 (60.7)                |
| Disposables                                          | 49.85 (18.6)                  | 42.60 (16.6)                 |
| Intravenous drugs                                    | 19.03 (7.1)                   | 19.03 (7.4)                  |
| Laboratory checks                                    | 5.34 (2.0)                    | 5.34 (2.1)                   |
| Vascular access                                      | 3.46 (1.3)                    | 3.46 (1.4)                   |
| Dialysis machines: supplies, monitoring, depreciation, etc. | 19.39 (7.2) | 15.86 (6.2) |
| Water treatment: supplies, monitoring, depreciation, etc. | 4.03 (1.5)                   | 2.52 (1.0)                   |
| Laundry, hygiene, and safety                         | 5.14 (1.9)                    | 5.14 (2.0)                   |
| Transportation                                       | 3.45 (1.3)                    | 3.45 (1.3)                   |
| Other                                                | 3.17 (1.1)                    | 3.17 (1.2)                   |
| Total                                                | 268.40 (100)                  | 256.12 (100)                 |

OL-HDF on-line hemodiafiltration, LF-HD low-flux hemodialysis
effectiveness is above 90%. A word of caution: it cannot be excluded that OL-HDF is less effective than LF-HD as the upper bound of the 95% confidence interval of the hazard ratio of death on OL-HDF compared with LF-HD is larger than one. Furthermore, our result, strictly speaking, only applies to in-center patients for whom home dialysis is not an option. Otherwise, the analysis would need to include home dialysis as a comparator.

One may consider no treatment as an unrealistic alternative for patients who could be treated either with hemodialysis or extended conservative or supportive therapy. Yet, considering conservative management as a comparator of hemodialysis would likely increase the cost-effectiveness ratio of hemodialysis even further. Also, using no treatment as a comparator of hemodialysis is quite common in the health economics literature. In fact, it is the cost-effectiveness ratio of hemodialysis compared with no treatment which is often used as a benchmark of societal willingness to pay in the academic literature [29]. We followed this practice and thus also used it as a benchmark for the cost-effectiveness ratio of OL-HDF.

Our model has the same structure as the cost-utility analysis based on the original CONTRAST study [8], i.e., a Markov model containing two health states, dialysis and death. Two other recently published models [14, 15] consider in addition transitions between different dialysis
modes, between dialysis and transplantation, as well as between dialysis with and without complications. Yet, in our trial, patients did not switch between OL-HDF and LF-HD. Furthermore, the number of dropouts due to transplantation was the same in both groups and thus cancels out from the calculation of the ICER. Finally, while we included the cost of dialysis side effects, we did not model it through the probability of side effects due to availability of patient-level data.

The trial population was representative for the Canadian ESRD population for the variables age, sex, and primary kidney disease [30] and the two arms were well balanced with respect to baseline characteristics. In this study, non-significant improvements in quality of life and survival may be the result of a lack of power due to small sample size. However, considering even non-significant changes in health for the calculation of ICERs is consistent with the concept of ‘irrelevance of inference’ [31]. Accordingly, rules of inference are arbitrary and entirely irrelevant for decisions based on economic evaluations.

The result of this study differs from that for the Dutch evaluation of the CONTRAST study, which, based on year 2009 euros, reported a cost-utility ratio of €287,679 per QALY gained over a period of 5 years [8] (corresponding to Can$420,357 as of August 7, 2014). A major reason is the larger QALY gain by OL-HDF in our study, resulting from larger gains both in quality of life and survival compared with the original CONTRAST study. Indeed, one can hypothesize that larger improvement of patient outcomes is likely attributable to clinical practices of this center, i.e., achievement of a higher convective volume was able to optimize overall treatment effectiveness.

The estimate for the hazard ratio in this sample (0.79) falls in the range of a recently published estimate (0.84; 95% confidence interval 0.73–0.96) from a meta-analysis [32] of six studies comparing OL-HDF with low- or high-

| Table 6 | Discounted incremental costs, effects, and cost effectiveness of on-line hemodiafiltration (OL-HDF) versus low-flux hemodialysis (LF-HD) |
|---------|---------------------------------------------------------------------------------------------------------------------------------|
|         | Costs | Life-years | QALYs | Incremental costs per life-year gained (OL-HDF vs LF-HD) | Incremental costs per QALY gained (OL-HDF vs LF-HD) |
| Trial-based analysis | OL-HDF | 220,018 | 4.01 | 2.87 | 53,153 | 32,112 |
|         | LF-HD | 203,629 | 3.70 | 2.36 | |
| Model-based analysis | OL-HDF | 368,177 | 6.21 | 4.45 | 58,840 | 53,270 |
|         | LF-HD | 306,826 | 5.17 | 3.30 | |

Costs are in 2013 Canadian dollars and are rounded. QALY: quality-adjusted life-year.

| Table 7 | Univariate sensitivity analysis |
|---------|--------------------------------|
| Variable | Incremental costs per QALY gained over lifetime |
| Annual discount rate | |
| Lower limit (0 %) | 56,034 |
| Upper limit (7 %) | 49,832 |
| Annual probability of death on LF-HD | |
| Lower limit (10 %) | 48,328 |
| Upper limit (21 %) | 53,270 |
| Hazard ratio of death for OL-HDF compared with LF-HD | |
| Lower limit (0.440) | 72,129 |
| Upper limit (1.418) | 145,696 |

Costs are in 2013 Canadian dollars. LF-HD: low-flux dialysis, OL-HDF: on-line hemodiafiltration, QALY: quality-adjusted life-year.

Fig. 2 Scatter plot of incremental costs and quality-adjusted life-years (QALYs) of on-line hemodiafiltration (OL-HDF) compared with low-flux hemodialysis (LF-HD). Plotted are a 95% confidence ellipse (black line) and the mean cost-effect pair (red dot).
flux HD (four studies were on post-dilution [5–7, 33], one on pre-dilution [34], and one on mixed [35] OL-HDF). Furthermore, the hazard ratio in this study is lower than for the original CONTRAST study (0.95) [5], indicating improved survival. The latter result can potentially be explained by the higher mean convective volume of 27.4 ± 7.3 L/session as compared with a mean of 19.7 L/session in the original CONTRAST study [10]. According to Sehgal et al. [36], dialysis dose for low molecular weight substances is inversely correlated to morbidity. Thus, patients with lower dialysis dose have higher hospitalization rates. The same relationship holds for the convective dose [7, 37]. For example, in the study of Maduell et al. [7], a convective dose in the range of 23–24 L/session in patients assigned to on-line OL-HDF was associated with a lower all-cause hospitalization rate.

A recent systematic review [38] including 226 utility estimates from dialysis patients (the majority of whom were receiving hemodialysis) reported a preference weight of 0.70, very close to the estimates in the current study of 0.72 and 0.64 for OL-HDF and LF-HD, respectively (when applying the UK value set).

At CHUM, the additional treatment cost of OL-HDF compared with LF-HD was estimated to be Can$12.48 per session, ensuring ‘ultrapure dialysate’ to patients both on OL-HDF and LF-HD. In 2002, Beerenhout et al. [39] estimated this extra cost to be in the range of €6 to €21 when comparing OL-HDF with high-flux ‘ultrapure’ HD and in the range of €17 to €32 when comparing it with LF-HD. In the meantime, the difference in costs between low- and high-flux membranes has decreased and microbiological testing requirements have been reduced.

Our study shows that spending on pharmaceuticals was 14 % lower for patients on OL-HDF. Lower costs of pharmaceuticals mainly resulted from significantly lower doses of phosphate binders. Doses of rHu-erythropoietin and erythropoiesis-stimulating agents were also lower, in line with the result of prospective observational studies [40, 41] and RCTs [42], but the difference did not reach the level of statistical difference. It should be stressed that the level of hemoglobin correction was comparable for patients on OL-HDF and LF-HD (e.g., 11.4 vs 11.0 g/dL in the third month of follow-up and 11.5 vs 11.0 g/dL in the twelfth month of follow-up for OL-HDF and LF-HD, respectively). In the CONTRAST study [5], costs for rHu-erythropoietin were very similar in both treatment groups.

The following limitations of the study have to be taken into consideration. First, the reference treatment of the CONTRAST study was LF-HD. Actually, according to market survey data, only 5 % of the dialyzers sold in Canada are in the low-flux range. However, since the cost difference between low-flux and high-flux dialyzers is very small today and the hazard ratio estimated by our study is in the range of the estimations of Maduell et al. [7] for high-flux HD as a comparator, this limitation should not have significantly biased the results. Also, less costly dialysis modalities exist in Canada for more autonomous and healthier (or less comorbid) patients for whom home dialysis is an option [43]. As a second limitation, while we included costs of hospitalizations for diseases related and unrelated to kidney disease, we did not consider costs of unplanned outpatient visits. However, these costs may be the same in both arms and thus cancel out. Third, due to the nature of the treatments, both patients enrolled and medical personnel were not blinded to treatment. While quality-of-life improvements by OL-HDF could therefore be attributed to a placebo effect, this effect would also occur outside the trial setting and thus should be accounted for. Fourth, it cannot be excluded that an imbalance in baseline characteristics had an impact on results. Yet, even if the real survival benefit were smaller than shown by the study, results would also be favorable for OL-HDF. That is, when assuming no survival benefit, our discounted cost analysis suggests savings through OL-HDF. Finally, renal units considering the introduction of OL-HDF would face implementation costs in terms of modifying dialysis
delivery to a new system with potential capital, operational, and training costs. As is typical for cost-effectiveness analyses, these implementation costs were not included. Also, renal units may be hesitant to introduce OL-HDF for the reason that the survival benefit of OL-HDF carries some uncertainty or is not strongly recommended by guidelines yet. Therefore, our results primarily support those renal units which have already switched or have made the decision to switch to OL-HDF.

In summary, in the Canadian arm of the CONTRAST study, high efficiency OL-HDF was found to be a cost-effective treatment for ESRD compared with LF-HD. Studies in different healthcare systems and regulatory environments are required to confirm the result.

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The study was approved by the local medical ethics review board (CE10.253). It was conducted in accordance with the declaration of Helsinki and Good Clinical Practice Guidelines. Informed consent was obtained from all individual participants included in the study.

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