Cost-effectiveness analysis of potentially curative and combination treatments for hepatocellular carcinoma with person-level data in a Canadian setting

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Abstract

Patients with early-stage hepatocellular carcinoma (HCC) are potential candidates for curative treatments such as radiofrequency ablation (RFA), surgical resection (SR), or liver transplantation (LT), which have demonstrated a significant survival benefit. We aimed to estimate the cost-effectiveness of curative and combination treatment strategies among patients diagnosed with HCC during 2002–2010. This study used Ontario Cancer Registry-linked administrative data to estimate effectiveness and costs (2013 USD) of the treatment strategies from the healthcare payer’s perspective. Multiple imputation by logistic regression was used to handle missing data. A net benefit regression approach of baseline important covariates and propensity score adjustment were used to calculate incremental net benefit to generate incremental cost-effectiveness ratio (ICER) and uncertainty measures. Among 2,222 patients diagnosed with HCC, 10.5%, 14.1%, and 10.3% received RFA, SR, and LT monotherapy, respectively; 0.5–3.1% dual treatments; and 0.5% triple treatments. Compared with no treatment (53.2%), transarterial chemoembolization (TACE) + RFA (average $2,465, 95% CI: −$20,000–$36,600/quality-adjusted life years [QALY]) or RFA monotherapy ($15,553, 95% CI: $3,500–$28,500/QALY) appears to be the most cost-effective modality with lowest ICER value. The cost-effectiveness acceptability curve showed that if the relevant threshold was $50,000/QALY, RFA monotherapy and TACE + RFA would have a cost-effectiveness probability of 100%. Strategies using LT delivered the most additional QALYs and became cost-effective at a threshold of $77,000/QALY. Our findings found that TACE + RFA dual treatment or RFA monotherapy appears to be the most cost-effective curative treatment for patients with potential early stage of HCC in Ontario. These findings highlight the importance of identifying and measuring differential benefits, costs, and cost-effectiveness of alternative HCC curative treatments in order to evaluate whether they are providing good value for money in the real world.

Keywords
Cost, cost-effectiveness acceptability curve, economic evaluation, effect, intervention, liver cancer

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Introduction

Liver cancer is one of the few cancers that is increasing in incidence and mortality worldwide [1], including Canada [2–4]. This is due to the growing prevalence of underlying chronic liver diseases, mainly chronic viral hepatitis, alcoholic and nonalcoholic liver disease, and the aging of the population that have those diseases [2–4]. Hepatocellular carcinoma (HCC) accounts for the majority (~72%) of liver cancers in both males and females in Canada [4]. For patients with early-stage HCC, potentially curative treatment options include radiofrequency ablation (RFA), surgical resection (SR), and liver transplantation (LT). These treatments provide survival benefits, and outcomes are optimized by identification of appropriate patients [5–7]. Transarterial chemoembolization (TACE) is the standard of care for patients with intermediate-stage HCC [8–10]. However, in clinical practice, TACE has been used as an alternative or combination therapy in patients with early- or advanced-stage HCC [11].

HCC is associated with high costs for treatment, in particular, SR and LT, and management of the cancer presents both clinical and financial challenges [4, 12, 13]. Facilities and resources in Canada to deal with end-stage liver disease are generally inadequate aside from the few pockets of expertise at university-based centers that have limited capacity [14]. As HCC incidence increases, access to potentially curative treatments and the costs associated with them will present new challenges to the Canadian healthcare system. Important considerations include economic evaluation in general, and cost-effectiveness analysis to help guide evidence-based policy decision-making in balancing health gains against the costs of interventions [15].

Net benefit regression framework utilizing person-level data from administrative datasets facilitates the use of regression methods in the economic evaluation [16]. Instead of the usual approach of aggregating cost and effect differences across different intervention strategies, the key advantage of the net benefit framework is the ability to use standard regression techniques, adjusting for explanatory variables to examine the marginal impact on incremental cost-effectiveness [16]. The net benefit calculation determines whether a new treatment meets (or surpasses) the decision-maker’s expectations of “good value for money”. The results can be used to help develop policy, with an aim toward improving efficiency and value in healthcare [17]. We aimed to evaluate the real-world cost-effectiveness of potentially curative and combination (including TACE combined with curative treatment) treatments in patients diagnosed with HCC over a 9-year study time frame from a healthcare payer’s perspective in a Canadian setting.

Materials and Methods

Study design and population

The study included all eligible HCC cases aged 18 years and older in Ontario diagnosed between January 1, 2002 and December 31, 2010 to estimate the effectiveness, cost, and cost-effectiveness of potentially curative treatments compared with no treatment to provide an estimate of the trade-off between extra benefit and extra cost as well as utilizing net benefit regression framework to estimate the incremental net benefit (INB). HCC cases were identified through the Ontario Cancer Registry (OCR). The International Statistical Classification of Disease and Related Health Problems, 9th Revision (ICD-9) site code 155.0, in combination with histology codes 8170–8175 of the International Classification of Diseases for Oncology, Third Edition (ICD-O-3), were used to identify cases of primary liver cancer. Patients who had death dates before or on the HCC diagnosis date during the study period were excluded. Furthermore, patients who received best supportive care within the first year after HCC diagnosis and those who received palliative treatments for advanced-stage of HCC such as sorafenib or chemotherapy over the study period were also excluded.

Data sources and study variables

The OCR is a provincial population-based cancer registry that contains information on all new cases of cancer (except for nonmelanoma skin cancers) in Ontario since 1964 [18]. The OCR includes data regarding date and stage of HCC diagnosis, age, sex, birth location, urban or rural residence, cause of death, and date of death. The OCR cohort was linked to the Discharge Abstract Database maintained by the Canadian Institute for Health Information, the Ontario Health Insurance Plan (OHIP), the Ontario Drug Benefit
Program, and the Canadian census data, to provide person-level information on sociodemographic, screening, treatment, and clinical factors [19]. The OHIP is a publicly funded healthcare program for all Ontario residents; physician billing claims dataset contains service and diagnosis information for outpatient visits in Ontario. The Discharge Abstract Database contains information relating to diagnosis and procedures for all hospitalizations in Ontario, frequency and type of hospital admissions, length of stay, and in-hospital mortality. The Ontario Drug Benefit dataset contains information regarding prescription medications (including sorafenib) dispensed to all adults aged 65 years and older and those receiving social assistance. Although there are some variances in different healthcare services, the system provides free access to hospital and emergency department visits, physician services, homecare, copayments for long-term care placements, and prescription medications for those aged 65 years and older.

Area-level socio-economic status was quantified using median neighborhood household income. Median neighborhood household income was determined through linking of postal codes to Canadian census data; income was categorized into quintiles corresponding to income status of neighborhoods. The least and the most well-off 20% of neighborhoods were included within the first and the fifth quintiles, respectively [20]. Where possible, hospitalization records from the date of diagnosis were used to assign each patient and control subject a baseline Charlson–Deyo comorbidity index. If patients did not have a hospitalization record at their diagnosis date, baseline comorbidity was determined by looking back 2 years into the hospitalization data to find the most recent hospitalization record; the comorbidity score from that hospitalization was then applied [12, 13, 20, 21]. The Charlson–Deyo comorbidity index at baseline was marked as “missing” if the individual had no hospitalization records at diagnosis or during the 2 years before diagnosis. Comorbidity was adjusted for each hospitalization after baseline. The Charlson–Deyo comorbidity index was calculated using methods previously described [22, 23]; an ICD-10 coding algorithm was applied to the diagnostic field codes from the hospitalization data (excluding diagnoses for liver disease, metastatic cancer, diabetes, and HIV). Conditions were weighted and then summed up to provide an overall comorbidity index value for a given episode, which was then categorized into one of five groups (0, 1, 2, ≥3, or no hospitalization record) representing different degrees of comorbidity.

Patients diagnosed with diabetes, HIV, and covariates that denote liver disease stage measured before HCC diagnosis were identified from the Discharge Abstract Database and OHIP using ICD-9 and ICD-10 codes. The study also included viral hepatitis cases identified through OHIP data; defined as subjects having at least two viral hepatitis visits (OHIP diagnostic code “070”) within the 4-year interval before the HCC diagnosis date—to cover as much available OHIP data as possible. Indicators of liver disease stage were categorized exclusively as: (1) viral hepatitis; (2) no cirrhosis; (3) cirrhosis; (4) alcoholic liver disease (ALD) + cirrhosis; (5) viral hepatitis + cirrhosis; (6) ALD + viral hepatitis + cirrhosis; (7) decompensated cirrhosis (i.e., cirrhosis and any recorded ascites, esophageal varices, or hepatic encephalopathy); (8) ALD + decompensated cirrhosis; (9) nonalcoholic fatty liver disease (NAFLD) + decompensated cirrhosis; (10) viral hepatitis + decompensated cirrhosis; and (11) ALD + viral hepatitis + decompensated cirrhosis. Other relevant variables (including ALD alone, NAFLD alone, etc.) which were <20 in total were not considered as covariates.

To identify patients who received screening ultrasonography, we identified all abdominal ultrasonography performed on patients before HCC diagnosis utilizing OHIP fee codes [21]. We obtained exclusive data regarding receipt of abdominal ultrasound screening (at least 4.5 months apart from previous ultrasound), which was defined as receiving one or more ultrasound screening annually for 2 years before HCC diagnosis (i.e., routine screening), at least one screen either within 12 months or between 12 and 24 months before HCC diagnosis (i.e., inconsistent screening), and no screening before HCC diagnosis.

**HCC treatment strategies**

HCC treatments were identified from the Discharge Abstract Database, OHIP, and Ontario Drug Benefit databases determining the timing; for example, if both Discharge Abstract Database and OHIP for SR exits, we considered only the first SR. Mutually exclusive potentially curative monotherapies and combination therapies with palliative treatment (TACE) for HCC considered include: (i) RFA monotherapy; (ii) SR monotherapy; (iii) LT monotherapy; (iv) RFA plus SR; (v) RFA plus LT; (vi) SR plus LT; (vii) TACE plus RFA; (viii) TACE plus SR; (ix) TACE plus LT; (x) RFA plus SR plus LT triple treatment; and (xi) no treatment. Procedure codes used to identify diabetes, HIV, indicators of liver disease stage, HCC screening, and treatments can be found in the Tables S1 and S2 [12, 20, 21].

**Measuring effectiveness**

Life expectancy for each age in this study is the estimated average period that a person may expect to live, according to the age-specific mortality rates for all causes. Potential years of life lost (PYLL, a measure of premature mortality) and quality-adjusted life years lost (QALYL) were used to measure effectiveness. This study followed patients...
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Measuring costs

Full details of data sources and estimation of direct health care costs associated with HCC has been previously published [12, 13]. The total costs of healthcare services included outpatient visits, emergency department visits, acute inpatient hospitalizations, same-day surgery, prescription medications, homecare visits, continuing care, and long-term care. Costs associated with outpatient physician visits and laboratory tests in Ontario were estimated from the Physician Claims History Database of the OHIP. Costs for emergency department visits and same-day surgery were estimated using the National Ambulatory Care Reporting System database [30]. The costs of hospitalization, emergency department visits, and same-day surgery for a particular year were estimated using the Resource Intensity Weight methodology developed by the Canadian Institute for Health Information [30]. Prescription medication costs were obtained from the Ontario Drug Benefit Program [30]. Costs associated with home care, continuing care, and long-term care were estimated from the Ontario Home Care database, Continuing Care Reporting System, and Ontario Drug Benefit Program. Costs were adjusted for inflation to 2013 Canadian dollars using the Statistics Canada Consumer Price Index for healthcare and personal items for Ontario [31]. Purchasing Power Parity for Gross Domestic Product was used to convert 2013 Canadian dollars to 2013 US dollars [32]. Effects and costs were discounted at 3% annually as a base case to capture time preference given somewhat variation in the follow-up time [33].

Statistical analysis

Multiple imputation was used to impute values for variables with a high degree of missing data such as cancer stage at HCC diagnosis, birth country, and Charlson–Deyo comorbidity index. Five independent draws from an imputation model were used to create five completed datasets and results were combined to obtain one imputation inference [34]. Statistically, multiple imputation is an established method to deal with replacing each missing value with a set of plausible values to ensure that the results are unbiased and capture the appropriate degree of precision [34]. Multiple Imputation procedure by logistic regression was used in a sequential process to generate monotone patterns (PROC MI with LOGISTIC in the MONOTONE statement) [35, 36].

Next, we used the net benefit regression framework [16] to evaluate the real-world cost-effectiveness of curative treatments of HCC compared with no treatment among patients diagnosed with HCC from 2002 to 2010. In the first step, the net benefit value for each person (NB) was calculated using the formula: willingness-to-pay (λ) * Ei − Ci, where Ei is the observed incremental effect (i.e., life year [LY] or quality-adjusted life year [QALY] gained) and Ci is the incremental cost, for the ith person. Various values of λ, for an additional effect [16] were explored ranging from $0 to $500,000. NB, differs by various levels of λ; therefore, the person-level net benefit is denoted as NB(λ).
The net benefit regression (i.e., multiple linear regression) involved fitting a linear regression model while adjusting for the relevant covariates (dummy variables), including sociodemographic characteristics: age (<60, 60–69, 70–79, ≥80 years), sex (male, female), income quintile (Q1-lowest to Q5-highest), residence (urban, rural), birth country (Canada, outside of Canada); clinical characteristics: Charlson–Deyo comorbidity index (0, 1, 2, ≥3), diabetes, HIV, liver disease stage, receipt of ultrasound screening 2 years before HCC diagnosis (routine screening, inconsistent screening, no screening), stage at diagnosis (early-stage I, intermediate-stage II, advanced-stage III-IV), and index year of HCC diagnosis (2002–2004, 2005–2007, 2008–2010). Additionally, we adjusted for propensity score to minimize bias related to the nonrandom allocation of potential curative treatment [37, 38]. The propensity score for an individual is the conditional probability of assignment to having a curative treatment of HCC given the observed individual covariates. Here, it was derived by fitting a logistic regression model with HCC curative treatment as the dependent variable and the aforementioned covariates as independent variables. This approach allows for the adjustment of how covariates may affect the estimate of the intervention’s INB (i.e., the marginal impact on incremental cost-effectiveness, ICER) [16]. The regression coefficient δ on the treatment dummy provides the estimate of the INB of treatment versus no treatment corresponding to a certain level of λ adjusted for the covariates. Treatment is defined to be cost-effective, at a certain level of λ, if the corresponding INB is positive (i.e., INB > 0).

Threshold values of the variance inflation factors were evaluated in the context of several other factors that influence the variance of regression coefficients [39]. We eliminated interaction terms if there was no statistical significance or if the variance inflation factor values exceed 10 (i.e., indicating severe multicollinearity), which can reduce the variance of the regression coefficients. All covariates were included in the model because they were considered to be significant correlates of the outcome (theoretical justification) or were significantly different between the treatments (statistical validation).

The final step was assessing uncertainties and constructing cost-effectiveness acceptability curves (CEACs) using the coefficient estimates of the treatment (T) variable and P-values obtained from the net benefit regression model [16, 40]. CEACs have been widely adopted as a method to quantify and incorporate the uncertainty that exists around the estimates of expected costs and expected effects associated with the interventions. A CEAC shows the probability that an intervention is cost-effective compared with the alternative, given the observed data, for a range of λ values that a decision-maker might be willing to pay for a particular unit change in the outcome (i.e., LY and QALY) [16, 40]. The P-values obtained from the net benefit regression are two-sided but only one-sided values are needed to test whether the INB is positive (i.e., treatment is cost-effective) or negative (i.e., treatment is not cost-effective) at the specified λ; therefore, the regression two-sided P-values are divided by two [16]. For negative INB, the probability that the treatment is preferred equals the one-sided P-value, and for positive incremental net-benefits, the probability that the treatment is preferred equals 1 minus the one-sided P-value [16]. A CEAC was created by plotting a graphical representation that the HCC curative treatment is cost-effective compared with no treatment (y-axis), as a function of societal λ threshold per additional LY or QALY for a range of λ between $0 and $100,000 (x-axis).

Analyses were performed using SAS version 9.4 (SAS Institute Inc., Cary, NC) and STATA version 12.0 (Stataco Corporation, College Station, TX) statistical software applications.

**Ethics approval**

Ethics approval for the study was granted by the University of Toronto Health Sciences Research Ethics Board. Informed consent was not obtained because this secondary analysis accessed existing de-identified data; consent was therefore deemed to be neither feasible nor necessary.

**Results**

**Description of cohort**

Overall, 3,857 patients were identified as having a primary diagnosis of HCC from the OCR between 2002 and 2010. Flowchart of study population can be found in Figure S1. The final study cohort comprised 2,222 patients diagnosed with HCC after excluding 1,154 patients who had best supportive care within 1 year after HCC diagnosis, and 481 patients who had palliative treatments (chemotherapy and sorafenib) during the study period. The median and mean (standard deviation) of follow-up time of patients diagnosed with HCC were 489 days and 735 (772) days, respectively. Overall baseline characteristics for this cohort are summarized in Table S4 and those stratified by treatment are summarized in Table 1. The majority (n = 1,182, 53.2%) of patients diagnosed with HCC did not receive curative treatment. Overall, 34.9% (n = 775) received a single curative treatment (10.5% received RFA, 14.1% SR and 10.3% LT monotherapy), 10.2% (n = 227) received dual treatments (3.1% received RFA plus SR, 2.5% RFA plus LT, 1.7% SR plus LT, 1.2% TACE plus RFA, 0.5% TACE plus SR, and 1.2% TACE plus LT), and 0.5%...
| Variable                        | No treatment n (%) | RFA n (%) | SR n (%) | LT n (%) | RFA + SR n (%) | RFA + LT n (%) | SR + LT n (%) | TACE + RFA n (%) | TACE + SR n (%) | TACE + LT n (%) | RFA + SR + LT n (%) |
|--------------------------------|--------------------|-----------|----------|----------|---------------|---------------|--------------|----------------|---------------|----------------|------------------|
| **Overall**                    | 1182 (53.2)        | 234 (10.5)| 313 (14.1)| 228 (10.3)| 69 (3.1)         | 55 (2.5)       | 37 (1.7)     | 27 (1.2)        | 12 (0.5)       | 27 (1.2)       | 12 (0.5)          |
| **Age group (years)**          |                    |           |          |          |               |               |              |                |               |                |                  |
| <60                            | 320 (27.1)         | 66 (28.2)| 114 (36.4)| 170 (74.6)| 20 (29.0)       | 35 (63.6)      | 23 (62.2)   | 6 (22.2)        | 6 (50.0)       | 17 (63.0)      | 10 (83.3)         |
| 60–69                          | 304 (25.7)         | 65 (27.8)| 79 (35.2) | 58 (25.4) | 22 (31.9)       | 19 (34.6)      | 14 (37.8)   | 6 (22.2)        |                | 10 (37.0)      |                  |
| 70–79                          | 367 (31.1)         | 85 (36.3)| 104 (33.2) | 0        | 22 (31.9)       |                |              |                | 14 (51.9)      |                |                  |
| 80+                            | 191 (16.2)         | 18 (7.7 )| 16 (5.1)  | 0        |                |                |              |                |                |                |                  |
| **Sex**                        |                    |           |          |          |               |               |              |                |                |                |                  |
| Female                         | 275 (23.3)         | 60 (25.6)| 72 (23.0)| 33 (14.5)| 12 (17.4)       | 6 (10.9)       | 7 (18.9)    | 10 (37.0)       |                |                |                  |
| Male                           | 907 (76.7)         | 174 (74.4)| 241 (77.0)| 195 (85.5)| 57 (82.6)       | 49 (89.1)      | 30 (81.1)  | 17 (63.0)       | 9 (75.0)       | 26 (96.3)      | 10 (83.3)         |
| **Income quintile**            |                    |           |          |          |               |               |              |                |                |                |                  |
| Q1 (lowest)                    | 322 (27.2)         | 56 (23.9)| 55 (17.6)| 52 (22.8)| 14 (20.3)       | 13 (23.6)      |              |                |                |                | 7 (25.9)          |
| Q2                             | 247 (20.9)         | 52 (22.2)| 70 (22.4)| 42 (18.4)| 17 (24.6)       | 12 (21.8)      | 7 (18.9)    |                |                |                | 9 (33.3)         |
| Q3                             | 244 (20.6)         | 45 (19.2)| 69 (22.0)| 48 (21.1)| 14 (20.3)       | 9 (16.4)       | 9 (24.3)    |                |                |                |                  |
| Q4                             | 173 (14.6)         | 38 (16.2)| 69 (22.0)| 45 (19.7)| 14 (20.3)       | 12 (21.8)      | 10 (27.0)   | 9 (33.3)       |                |                |                  |
| Q5 (highest)                   | 185 (15.7)         | 42 (18.0)| 50 (16.0)| 41 (18.0)| 9 (13.0)        | 9 (16.4)       | 6 (16.2)    | 0              | 6 (22.2)       |                |                  |
| **Residence**                  |                    |           |          |          |               |               |              |                |                |                |                  |
| Urban                          | 1061 (89.8)        | 220 (94.0)| 288 (92.0)| 207 (90.8)| 67 (97.1)       | 52 (94.6)      | 33 (89.2)  | 27 (100)        | 12 (100)       | 26 (96.3)      | 12 (100)         |
| Rural                          | 118 (10.0)         | 14 (6.0 )| 25 (8.0) | 21 (9.2) |                |                |              | 0              | 0              |                |                  |
| Missing                        |                    |           |          |          |               |               |              |                |                |                |                  |
| **Birth country**              |                    |           |          |          |               |               |              |                |                |                |                  |
| Other                          | 537 (45.4)         | 49 (20.9)| 77 (34.4)| 34 (14.9)| 10 (14.5)       |               |              |                |                |                |                  |
| Canada                         | 533 (45.1)         | 66 (28.2)| 66 (21.1)| 32 (14.0)| 13 (18.8)       |               |              |                |                |                |                  |
| Unknown/missing                | 112 (9.5)          | 119 (50.9)| 170 (54.3)| 162 (71.1)| 46 (66.7)       | 48 (87.3)      | 27 (73.0)  | 12 (44.4)       |                |                | 25 (92.6)        |
| **Comorbidity**                |                    |           |          |          |               |               |              |                |                |                |                  |
| 0                              | 404 (34.2)         | 101 (43.2)| 139 (44.4)| 67 (29.4)| 27 (39.1)       | 24 (43.6)      | 6 (16.2)    | 11 (40.7)       | 8 (66.7)       | 8 (29.6)       |                  |
| 1                              | 266 (22.5)         | 74 (31.6)| 89 (34.8)| 100 (43.9)| 22 (31.9)       | 21 (38.2)      | 21 (56.8)  | 8 (29.6)        |                | 13 (48.2)      | 7 (58.3)         |
| 2                              | 138 (11.7)         | 27 (11.5)| 36 (11.5)| 29 (12.7)| 14 (20.3)       |               |              |                |                |                |                  |
| 3 +                            | 100 (8.5)          | 24 (10.3)| 27 (8.6) | 25 (11.0)|                |               |              |                |                |                |                  |
| No hospitalization record      | 274 (23.2)         | 83 (3.4 )| 22 (7.0) | 7 (2.1)   |                |               |              |                |                |                |                  |
| Diabetes                       | 550 (46.5)         | 121 (51.7)| 137 (43.8)| 149 (65.4)| 40 (58.0)       | 31 (56.4)      | 30 (81.1)  | 13 (48.2)       |                | 20 (74.1)      | 7 (58.3)         |
| HIV                           | 22 (1.9)           | 9 (3.9)  | 6 (2.6)  |         |                |               |              |                |                |                |                  |
| **Indicators of liver disease stage** |                |           |          |          |               |               |              |                |                |                |                  |
| Viral hepatitis                | 35 (3.0)           | 10 (3.2) | 0        |         |                |               |              |                |                |                |                  |
| No cirrhosis                   | 277 (23.4)         | 16 (6.8 )| 102 (32.6)| 0      | 17 (24.6)       |               |              |                |                |                |                  |
| Cirrhosis                      | 205 (17.3)         | 45 (19.2)| 81 (25.9)| 24 (10.5)| 18 (26.1)       | 8 (14.6)       | 9 (24.3)    | 10 (37.0)       |                |                | 10 (37.0)        |
| ALD + cirrhosis                | 41 (3.5)           | 13 (5.6)| 6 (1.9)  |         |                |               |              |                |                |                |                  |
| Variable                                      | No treatment | RFA | SR | LT | RFA + SR | RFA + LT | SR + LT | TACE + RFA | TACE + SR | TACE + LT | RFA + SR + LT |
|----------------------------------------------|--------------|-----|----|----|----------|----------|---------|------------|-----------|-----------|---------------|
| Viral hepatitis + cirrhosis                  | 38 (3.2)     | 10 (4.3) | 18 (5.8) | -    | -        | -        | -       | 0          | 0         | 0         | -             |
| ALD + Viral hepatitis + cirrhosis            | 6 (0.5)      | -    | -   | 0   | -        | 0        | 0       | 0          | 0         | 0         | 0             |
| Decompensated cirrhosis                      | 288 (24.4)   | 70 (29.9) | 61 (19.5) | 85 (37.3) | 12 (17.4) | 18 (32.7) | 14 (37.8) | 11 (40.7) | -         | 12 (44.4) | 6 (50.0)       |
| ALD + Decompensated cirrhosis                | 159 (13.5)   | 44 (18.8) | 7 (2.2) | 66 (29.0) | -        | 16 (29.1) | 7 (18.9) | -          | 0         | 0         | 0             |
| NAFLD + Decompensated cirrhosis              | 8 (0.7)      | -    | 0   | -   | 0        | 0        | 0       | 0          | 0         | 0         | 0             |
| Viral hepatitis + Decompensated cirrhosis     | 53 (4.5)     | 14 (6.0) | 12 (3.8) | 19 (8.3) | -        | -        | -       | -          | -         | 0         | 0             |
| ALD + viral hepatitis + Decompensated cirrhosis | 47 (4.0) | 9 (3.9) | - | 15 (6.6) | - | - | 0 | - | 0 | 0 | 0 |

Ultrasound screening 2 years before HCC diagnosis
- No screening: 622 (52.6)
  - RFA: 86 (36.8)
  - SR: 125 (39.9)
  - LT: 113 (49.6)
  - RFA + SR: 24 (34.8)
  - RFA + LT: 24 (43.6)
  - SR + LT: 9 (24.3)
  - TACE + RFA: 9 (33.3)
  - TACE + SR: 6 (22.2)
  - TACE + LT: -
  - RFA + SR + LT: -
- Inconsistent screening: 464 (39.3)
  - RFA: 98 (41.9)
  - SR: 138 (44.1)
  - LT: 93 (40.8)
  - RFA + SR: 34 (49.3)
  - RFA + LT: 21 (38.2)
  - SR + LT: 20 (54.1)
  - TACE + RFA: 12 (44.4)
  - TACE + SR: 19 (70.4)
  - TACE + LT: 6 (50.0)
  - RFA + SR + LT: -
- >1 screens annually: 96 (8.1)
  - RFA: 50 (21.4)
  - SR: 50 (16.0)
  - LT: 22 (9.7)
  - RFA + SR: 11 (15.9)
  - RFA + LT: 10 (18.2)
  - SR + LT: 8 (21.6)
  - TACE + RFA: 6 (22.2)
  - TACE + SR: -
  - TACE + LT: -
  - RFA + SR + LT: -

Stage at HCC diagnosis
- Early (stage I): 66 (5.6)
  - RFA: 62 (26.5)
  - SR: 65 (20.8)
  - LT: 35 (15.4)
  - RFA + SR: 18 (26.1)
  - RFA + LT: 12 (21.8)
  - SR + LT: 6 (16.2)
  - TACE + RFA: 8 (29.6)
  - TACE + SR: -
  - TACE + LT: 6 (22.2)
- Intermediate (stage II): 83 (7.0)
  - RFA: 45 (19.2)
  - SR: 48 (15.3)
  - LT: 55 (24.1)
  - RFA + SR: 16 (23.2)
  - RFA + LT: 24 (43.6)
  - SR + LT: 7 (18.9)
  - TACE + RFA: 11 (40.7)
  - TACE + SR: -
  - TACE + LT: 13 (48.2)
- Advanced (stage III–IV): 274 (23.2)
  - RFA: 10 (4.3)
  - SR: 51 (16.3)
  - LT: 12 (5.3)
  - RFA + SR: 6 (8.7)
  - RFA + LT: -
  - SR + LT: -
  - TACE + RFA: -
  - TACE + SR: -
  - TACE + LT: -
- Unknown: 759 (64.2)
  - RFA: 117 (50.0)
  - SR: 149 (47.6)
  - LT: 126 (55.3)
  - RFA + SR: 29 (42.0)
  - RFA + LT: 17 (30.9)
  - SR + LT: 22 (59.5)
  - TACE + RFA: 7 (25.9)
  - TACE + SR: -
  - TACE + LT: 6 (50.0)

Year of HCC diagnosis
- 2002: 128 (10.8)
  - RFA: -
  - SR: 34 (10.9)
  - LT: 15 (6.6)
  - RFA + SR: -
  - RFA + LT: -
  - SR + LT: 0
  - TACE + RFA: -
  - TACE + SR: -
  - TACE + LT: -
- 2003: 106 (9.0)
  - RFA: 10 (4.3)
  - SR: 33 (10.5)
  - LT: 23 (10.1)
  - RFA + SR: 7 (10.1)
  - RFA + LT: 0
  - SR + LT: -
  - TACE + RFA: -
  - TACE + SR: -
  - TACE + LT: -
- 2004: 142 (12.0)
  - RFA: 6 (2.6)
  - SR: 23 (7.4)
  - LT: 29 (12.7)
  - RFA + SR: 6 (8.7)
  - RFA + LT: -
  - SR + LT: 7 (18.9)
  - TACE + RFA: -
  - TACE + SR: -
  - TACE + LT: -
- 2005: 135 (11.4)
  - RFA: 14 (6.0)
  - SR: 47 (15.0)
  - LT: 28 (12.3)
  - RFA + SR: -
  - RFA + LT: -
  - SR + LT: -
  - TACE + RFA: -
  - TACE + SR: -
  - TACE + LT: -
- 2006: 141 (11.9)
  - RFA: 22 (9.4)
  - SR: 29 (9.3)
  - LT: 38 (16.7)
  - RFA + SR: 9 (13.0)
  - RFA + LT: -
  - SR + LT: 6 (16.2)
  - TACE + RFA: -
  - TACE + SR: -
  - TACE + LT: -
- 2007: 142 (12.0)
  - RFA: 19 (8.1)
  - SR: 34 (10.9)
  - LT: 32 (14.0)
  - RFA + SR: 7 (10.1)
  - RFA + LT: 6 (10.9)
  - SR + LT: -
  - TACE + RFA: -
  - TACE + SR: -
  - TACE + LT: -
- 2008: 109 (9.2)
  - RFA: 46 (19.7)
  - SR: 25 (8.0)
  - LT: 23 (10.1)
  - RFA + SR: 13 (18.8)
  - RFA + LT: 18 (32.7)
  - SR + LT: -
  - TACE + RFA: 6 (22.2)
  - TACE + SR: 0
  - TACE + LT: 0
- 2009: 137 (11.6)
  - RFA: 56 (23.9)
  - SR: 45 (14.4)
  - LT: 21 (9.2)
  - RFA + SR: 10 (14.5)
  - RFA + LT: 10 (18.2)
  - SR + LT: 0
  - TACE + RFA: -
  - TACE + SR: -
  - TACE + LT: -
- 2010: 142 (12.0)
  - RFA: 59 (25.2)
  - SR: 43 (13.7)
  - LT: 19 (8.3)
  - RFA + SR: 9 (13.0)
  - RFA + LT: 12 (21.8)
  - SR + LT: -
  - TACE + RFA: 7 (25.9)
  - TACE + SR: 0
  - TACE + LT: 0

"-", counts less than six have been suppressed.
RFA, radiofrequency ablation; SR, surgical resection; LT, liver transplantation; HCC, hepatocellular carcinoma.
(n = 12) received triple treatments (RFA plus SR plus LT) during the study period. With regard to dual treatments with RFA, 46 (67%) patients underwent primary SR; with LT, 52 (95%) underwent primary RFA, 12 (32%) underwent primary SR, and 27 (100%) underwent primary TACE; with TACE, 17 (63%) underwent primary RFA and 12 (100%) underwent primary SR. Of the 2,222 patients, 13.0% were stage I, 14.2% stage II, 13.0% stage III, 3.7% stage IV, and 56.2% unknown stage at diagnosis. Patients with unknown stage were less likely to have received curative treatments. Age (except RFA + SR and TACE + RFA), Charlson–Deyo comorbidity index (except TACE + RFA, TACE + SR, TACE + LT, and RFA + SR + LT), liver disease stage (except SR + LT, TACE + RFA, TACE + LT, and RFA + SR + LT), and cancer stage (except SR + LT, TACE + SR, and RFA + SR + LT) were associated with receipt of curative treatments (P < 0.05); additionally, sex was associated with receipt of LT, RFA + LT or TACE + LT (P < 0.05), diabetes was associated with receipt of SR, LT, SR + LT or TACE + LT (P < 0.05), ultrasound screening was associated with receipt of RFA, SR, SR + LT or TACE + LT (P < 0.05), and year of HCC diagnosis was associated with receipt of RFA, SR, LT, RFA + LT or RFA + SR + LT (P < 0.05).

Healthcare effects and costs

Effects and costs stratified by treatment strategies are summarized in Table 2. The lowest QALYL was among those who received RFA + SR (9.1, 95% confidence interval [CI]: 8.8–9.5) and the highest QALYL were among those who received LT (11.9, 95% CI: 11.7–12.0) or RFA + LT (11.8, 95% CI: 11.6–12.1). The lowest costs were among those who did not receive treatment ($38,472, 95% CI: $37,255–$39,689) followed by those who received TACE + RFA ($48,485, 95% CI: $43,663–$53,307), and RFA monotherapy ($55,925, 95% CI: $52,123–$59,727); and the highest costs were among those who received SR + LT ($222,275, 95% CI: $205,992–$238,558) or RFA + SR + LT ($208,484, 95% CI: $190,385–$226,582).

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With regard to dual treatments with RFA, 46 (67%) patients underwent primary SR; with LT, 52 (95%) underwent primary RFA, 12 (32%) underwent primary SR, and 27 (100%) underwent primary TACE; with TACE, 17 (63%) underwent primary RFA and 12 (100%) underwent primary SR. Of the 2,222 patients, 13.0% were stage I, 14.2% stage II, 13.0% stage III, 3.7% stage IV, and 56.2% unknown stage at diagnosis. Patients with unknown stage were less likely to have received curative treatments. Age (except RFA + SR and TACE + RFA), Charlson–Deyo comorbidity index (except TACE + RFA, TACE + SR, TACE + LT, and RFA + SR + LT), liver disease stage (except SR + LT, TACE + RFA, TACE + LT, and RFA + SR + LT), and cancer stage (except SR + LT, TACE + SR, and RFA + SR + LT) were associated with receipt of curative treatments (P < 0.05); additionally, sex was associated with receipt of LT, RFA + LT or TACE + LT (P < 0.05), diabetes was associated with receipt of SR, LT, SR + LT or TACE + LT (P < 0.05), ultrasound screening was associated with receipt of RFA, SR, SR + LT or TACE + LT (P < 0.05), and year of HCC diagnosis was associated with receipt of RFA, SR, LT, RFA + LT or RFA + SR + LT (P < 0.05).

Healthcare effects and costs

Effects and costs stratified by treatment strategies are summarized in Table 2. The lowest QALYL was among those who received RFA + SR (9.1, 95% confidence interval [CI]: 8.8–9.5) and the highest QALYL were among those who received LT (11.9, 95% CI: 11.7–12.0) or RFA + LT (11.8, 95% CI: 11.6–12.1). The lowest costs were among those who did not receive treatment ($38,472, 95% CI: $37,255–$39,689) followed by those who received TACE + RFA ($48,485, 95% CI: $43,663–$53,307), and RFA monotherapy ($55,925, 95% CI: $52,123–$59,727); and the highest costs were among those who received SR + LT ($222,275, 95% CI: $205,992–$238,558) or RFA + SR + LT ($208,484, 95% CI: $190,385–$226,582).

Table 2. Healthcare effects and costs after diagnosis of hepatocellular carcinoma by treatment strategies, 2002–2010.

| Treatment strategies | Effects (mean, 95% CI) | Costs1 (mean, 95% CI) |
|----------------------|------------------------|----------------------|
|                      | PYLL                   | QALYL                |                        |
| No treatment (n = 5910) | 11.2251 (11.1105–11.3397) | 10.1149 (10.009–10.2207) | $38,472 ($37,255–$39,689) |
| Monotherapy          |                        |                      |                        |
| RFA (n = 1170)       | 10.2246 (9.9959–10.4534) | 9.8759 (9.6697–10.0821) | $55,925 ($52,123–$59,727) |
| SR (n = 1565)        | 10.0818 (9.8927–10.2709) | 9.9144 (9.7466–10.0822) | $119,032 ($115,799–$122,265) |
| LT (n = 1140)        | 11.9376 (11.7841–12.0911) | 11.8696 (11.7471–11.992) | $211,286 ($203,566–$219,007) |
| Dual treatments      |                        |                      |                        |
| TACE plus RFA (n = 135) | 9.6379 (8.979–10.2968) | 9.3606 (8.7722–9.9489) | $48,485 ($43,663–$53,307) |
| RFA plus SR (n = 345) | 9.0966 (8.6997–9.4934) | 9.1399 (8.844–9.4935) | $109,927 ($103,953–$115,902) |
| TACE plus LT (n = 60) | 11.4624 (10.4676–12.4573) | 10.999 (10.0909–11.907) | $126,514 ($114,451–$138,577) |
| RFA plus LT (n = 275) | 12.0635 (11.7867–12.3402) | 11.8248 (11.5797–12.0699) | $155,898 ($144,119–$167,677) |
| TACE plus LT (n = 135) | 11.4675 (11.0773–11.8578) | 11.4621 (11.359–11.7883) | $178,354 ($163,494–$193,215) |
| TACE plus LT (n = 185) | 10.756 (10.3731–11.1388) | 10.8734 (10.5621–11.1847) | $222,275 ($205,992–$238,558) |
| Triple treatments    |                        |                      |                        |
| RFA plus SR plus LT (n = 60) | 11.1088 (10.3933–11.8242) | 11.2472 (10.6054–11.844) | $208,484 ($190,385–$226,582) |

RFA, radiofrequency ablation; SR, surgical resection; LT, liver transplantation; PYLL, potential years of life lost (a measure of premature mortality); QALYL, quality-adjusted life years lost.

1All costs reflect 2013 US$ per person. Multiple imputation by logistic regression was used to generate missing data (cancer stage at HCC diagnosis, birth country, and Charlson–Deyo comorbidity index) for outcomes.

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Figure S2A–J (LYs) and Figure 2A–J (QALYs) show estimates of INB (i.e., ICER estimate) and its 95% CIs as a function of willingness-to-pay thresholds. The lowest ICER estimate for TACE plus RFA was $2,465/QALY gained (95% CI: $−20,000-$36,600/QALY); this means TACE plus RFA is cost-effective if decision-makers value >$36,600/QALY threshold, but not cost-effective if decision-makers value <$−20,000/QALY threshold.

Alternative ICER estimates in order were: for RFA monotherapy, $15,553/QALY (95% CI: $3,500–$28,500/QALY); RFA + SR, $48,761/QALY (95% CI: $35,000–$67,200/QALY); RFA + LT, $59,642/QALY (95% CI: $46,600–$78,000/QALY); RFA + SR + LT, $60,602/QALY (95% CI: $42,600–$90,000/QALY); SR + LT, $71,972/QALY (95% CI: $58,100–$91,450/QALY); TACE + LT, $72,941/QALY (95% CI: $52,700–$107,600/QALY); LT monotherapy,
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Figure 3A and B show CEACs which plot the probability that each treatment strategy is cost-effective compared with no treatment as a function of willingness-to-pay threshold for an additional quality-adjusted life year (QALY), respectively. The results showed that if a threshold of $50,000/LY gained was to be chosen, RFA monotherapy, SR monotherapy, RFA plus SR, RFA plus LT, and TACE plus RFA would have a cost-effectiveness probability of 99–100% (Table S5); whereas, if $50,000/QALY gained was to be chosen, RFA monotherapy and TACE plus RFA would have a cost-effectiveness probability of 99–100% (Table 4); and if a threshold of $100,000/QALY gained was to be chosen, all treatments would have a cost-effectiveness probability of more than 95% (except TACE plus SR [12%], Table 4).

Sensitivity analysis

Sensitivity analysis according to the pooled lower and upper bound health state utilities of disease stage from published literature showed that TACE plus RFA, RFA monotherapy, and RFA + SR dual treatment appeared to be acceptable compared to no treatment, similar to base case (Fig. S3A and B). A sensitivity analysis of excluding HCC stage IV appeared robust to the base case (stage
IV was lumped with stage III) relating to the incremental effects (LYs and QALYs) and costs, and ICER of curative treatments relative to no treatment (Fig. S4A and B).

**Discussion**

This study evaluated the real-world cost-effectiveness of potentially curative treatments compared with no treatment among patients diagnosed with HCC over a 9-year study time frame from a healthcare payer’s perspective in Ontario, Canada’s most populated province with approximately 13.6 million people as of year 2013. We note that during this time period, no new curative therapies have become available. Compared with no treatment, the adjusted incremental benefit of LT-related treatments are estimated to yield more units of QALYs than RFA or SR treatments, but are more costly. ICERs of TACE plus RFA dual treatment (i.e., major primary RFA; average $2,465, 95% CI: $−20,000–$36,600/QALY gained) and RFA monotherapy (average $15,553, 95% CI: $3,500–$28,500/QALY gained) are below the commonly cited thresholds of $50,000/QALY [41]. Interventions costing less than $50,000/QALY are often considered cost-effective [42], but oncologists commonly endorse higher thresholds [43]. The CEACs show that if a threshold of $50,000/QALY gained is to be chosen, RFA monotherapy and TACE plus RFA would have a cost-effectiveness probability of 99–100%. If a threshold of $100,000/QALY gained is to be chosen, all treatments would have a cost-effectiveness probability of more than 95% (except TACE plus SR).

Historically, percutaneous RFA was widely used for local control of small unresectable HCC including those patients who could not tolerate SR but it is increasingly being used as first-line therapy for patients with amenable lesions [9]. Based on published papers, the combination treatment of TACE and RFA seems to be a safe and effective treatment strategy for patients with early- or intermediate-stage HCC [44, 45]. In patients included in the waiting list for a LT, tumors are often treated as a “bridge” to transplant while waiting for an organ to become available because of the risk of tumor progression. RFA is widely used as a “bridge” to transplantation in order to avoid this progression [46]. This analysis was able to include patients who may have undergone palliative treatment (i.e., TACE) as a bridge to LT.

A few other studies analyzed the cost-effectiveness of two potentially curative treatment strategies in early-stage HCC within the Milan criteria using Markov cohort models. Cucchetti et al. compared SR with RFA and found that in the presence of two or three nodules ≤3 cm, RFA is more cost-effective than resection; for single larger early-stage HCC, SR is cost-effective at a willingness-to-pay largely acceptable in the oncologic setting [27]. Lim et al. compared SR with LT and found that in patients with HCC within the Milan criteria and Child-Pugh A/B cirrhosis, SR is more cost-effective than cadaveric LT across three different costing scenarios: the USA, Switzerland, and Singapore [28]. Our findings highlight the important implications for identification and measurement of differential benefits, costs, and cost-effectiveness of alternative HCC curative treatments in order to evaluate whether particular healthcare technologies are providing good value for money in the real-world within the context of an organized healthcare system.

HCC is a rapidly rising disease and many patients who develop HCC present late which becomes an incurable advanced-stage disease. The biggest limitation to treating HCC is the function of the underlying liver and the stage of first HCC diagnosis. It is evident that patients with HCC receiving treatments might appear fairly different with regard to clinical and tumor features that are known to affect prognosis [47]. Patients might not be considered suitable for surgery because of liver dysfunction and/or portal hypertension, as well as the presence of comorbidities or advanced age contraindicating general anesthesia [47]. Screening and surveillance programs for early detection of HCC and keeping track of the outcomes for quality

![Figure 3. Cost-effectiveness acceptability curves showing the probability that each curative treatment strategy is cost-effective compared with no treatment for a given willingness-to-pay threshold for an additional (A) life year (LY); and (B) quality-adjusted life year (QALY).](image-url)
Table 4. Estimates of incremental net benefit and probability of cost-effectiveness of curative treatment strategies for hepatocellular carcinoma compared with no treatment as a function of willingness-to-pay threshold per additional QALY over the study period 2002–2010.

| λ thresholds | INB estimate (SE) | Probability of cost-effectiveness | INB estimate (SE) | Probability of cost-effectiveness | INB estimate (SE) | Probability of cost-effectiveness |
|--------------|------------------|---------------------------------|------------------|---------------------------------|------------------|---------------------------------|
| Radiofrequency ablation | Surgical resection | Liver transplantation |
| $0 | $0 | $0 |
| $1,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $10,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $20,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $30,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $40,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $50,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $60,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $70,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $80,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $90,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $100,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |

Radiofrequency ablation plus surgical resection

| λ thresholds | INB estimate (SE) | Probability of cost-effectiveness | INB estimate (SE) | Probability of cost-effectiveness | INB estimate (SE) | Probability of cost-effectiveness |
|--------------|------------------|---------------------------------|------------------|---------------------------------|------------------|---------------------------------|
| TACE plus radiofrequency ablation | TACE plus surgical resection | TACE plus liver transplantation |
| $0 | $0 | $0 |
| $1,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $10,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $20,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $30,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $40,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $50,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $60,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $70,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $80,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $90,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| $100,000 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 | $0.001 |
| λ thresholds | INB estimate (SE) | P-value* | Probability of cost-effectiveness | INB estimate (SE) | P-value* | Probability of cost-effectiveness | INB estimate (SE) | P-value* | Probability of cost-effectiveness |
|--------------|-------------------|----------|-----------------------------------|-------------------|----------|-----------------------------------|-------------------|----------|-----------------------------------|
| $30,000      | 25,804 (15,183)   | 0.045    | 0.9554                            | −82,753 (22,420)  | <0.001   | 0.0001                            | −77,909 (15,248)  | <0.001   | 0.0001                            |
| $40,000      | 35,160 (16,628)   | 0.017    | 0.9828                            | −78,360 (24,541)  | <0.001   | 0.0007                            | −59,752 (16,699)  | <0.001   | 0.0002                            |
| $50,000      | 44,515 (18,386)   | 0.008    | 0.9923                            | −73,968 (27,118)  | 0.003    | 0.0032                            | −41,596 (18,463)  | 0.012    | 0.0122                            |
| $60,000      | 53,870 (20,376)   | 0.004    | 0.9959                            | −69,575 (30,032)  | 0.010    | 0.0103                            | −23,439 (20,461)  | 0.126    | 0.1260                            |
| $70,000      | 63,226 (22,536)   | 0.003    | 0.9975                            | −65,182 (33,196)  | 0.025    | 0.0248                            | −5283 (22,629)    | 0.408    | 0.4077                            |
| $80,000      | 72,581 (24,822)   | 0.002    | 0.9983                            | −60,790 (36,544)  | 0.048    | 0.0481                            | 12,874 (24,924)   | 0.303    | 0.6973                            |
| $90,000      | 81,936 (27,203)   | 0.001    | 0.9987                            | −56,397 (40,030)  | 0.079    | 0.0795                            | 31,030 (27,314)   | 0.128    | 0.8721                            |
| $100,000     | 91,292 (29,655)   | 0.001    | 0.9990                            | −52,005 (43,622)  | 0.117    | 0.1166                            | 49,187 (29,775)   | 0.049    | 0.9508                            |

| λ thresholds | Radiofrequency ablation plus surgical resection plus liver transplantation |
|--------------|--------------------------------------------------------------------------------|
| $0           | −164,675 (20,178)                                                             | <0.001 | 0.0001 |
| $1,000       | −161,956 (20,138)                                                             | <0.001 | 0.0001 |
| $10,000      | −137,487 (20,158)                                                             | <0.001 | 0.0001 |
| $20,000      | −110,300 (20,961)                                                             | <0.001 | 0.0001 |
| $30,000      | −83,113 (22,500)                                                              | <0.001 | 0.0001 |
| $40,000      | −55,925 (24,636)                                                              | 0.012  | 0.0116 |
| $50,000      | −28,738 (27,230)                                                              | 0.146  | 0.1457 |
| $60,000      | −1551 (30,164)                                                               | 0.480  | 0.4795 |
| $70,000      | 25,637 (33,348)                                                               | 0.221  | 0.779 |
| $80,000      | 52,824 (36,717)                                                               | 0.075  | 0.925 |
| $90,000      | 80,011 (40,225)                                                               | 0.023  | 0.977 |
| $100,000     | 107,199 (43,838)                                                              | 0.007  | 0.993 |

λ, willingness-to-pay; INB, incremental net benefit; SE, standard error. *One-sided P-value.
assurance are needed [48]. Given the increasing and high mortality rate associated with HCC, investments and healthcare resources at existing regional cancer centers should be enhanced to facilitate the multidisciplinary care to mitigate the impact of the disease [14, 49]. Thus, treating liver disease is a priority and is needed to act to protect the health and well-being of Canadians of all ages. Recommendations to improve the control of HCC in Canada include healthcare providers to identify, offer testing, and counsel people at risk for HCC such as alcohol and non-alcohol-induced liver disease, cirrhosis, diabetes, obesity, smoking, [50–54] and HIV co-infection [55], in particular, marginalized groups of people. Furthermore, patients need easier access to treatments to reduce the chance of progression to liver cancer.

The net benefit framework can clearly demonstrate how different the ICERs are when adjusted or not adjusted for covariates. The advantage of this is that influential covariates can be adjusted for in the regression model to obtain a more accurate INB [16]. This NBR found several covariates associated with INB (P < 0.05), including age group and sex (from λ $10,000 to λ $100,000), income quintile (from λ $0 to λ $25,000), Charlson–Deyo comorbidity index, cirrhosis, and viral hepatitis + cirrhosis (from λ $0 to λ $100,000), ALD + cirrhosis, decompensated cirrhosis, and routine screening (i.e., one or more ultrasound screening annually for 2 years before HCC diagnosis) (from λ $0 to λ $50,000), and HIV diagnosis and year of HCC diagnosis (from λ $25,000 to λ $100,000). The strategy with the largest incremental benefit while still being cost-effective may be the preferred strategy.

This study is meant to be a cost-effectiveness analysis taken from the perspective of the healthcare payer on a population-level (and not an individual-level decision-analysis to aid individual decision-making). This helps payers understand the trade-offs of cost and benefits of the various curative intent therapies. This information can help with priority-setting to allow payers maximize health outcomes if they are faced with finite budget. Given the rising cost of therapies, especially with the rising cost of new expensive drugs, this study can help payers understand the value for money of the curative intent therapies in the context of demands to fund newer but palliative expensive drugs.

There are a number of limitations in this study that should be considered. Using a 9-year observation period may result in an underestimation of the true benefits of HCC curative treatment. If HCC curative treatment does have an impact on the long-term survival, the full impact will not be observed within the 9-year observation period. Although additional work such as building a mathematical model is not within the scope of this study, data collected can be utilized to support future modeling studies. Other limitations include the stage of HCC at treatment; clearly, smaller lesions detected early by regular serial ultrasound surveillance, will be associated with lower costs and better survival, than more advanced lesions requiring multi-modal treatment or LT. From our analysis, it can be inferred that ultrasound surveillance programs may be associated with lower costs and better cost-effectiveness if small lesions amenable to RFA only, can be diagnosed. This may have an effect but overall, there is only 11.8% receiving regular ultrasound screening. Patients who may have undergone RFA at laparotomy (i.e., because of location of the HCC) may have been coded as dual therapy or SR clinically.

Conclusions

From our analysis, compared with no treatment, TACE plus RFA dual treatment or RFA monotherapy appears to be the most cost-effective modality with lowest ICER value if a threshold of $50,000/QALY gained was to be chosen, but this is generalizable only to those who are diagnosed with early HCC lesions. In order to achieve optimal cost-effectiveness of HCC treatment with curative intent, patients at risk of HCC must be diagnosed early (i.e., via regular ultrasound surveillance) and referred for treatment in a timely manner before HCC disease progression requires more advanced curative treatment (i.e., SR, LT, etc.) that is associated with greater healthcare costs and less favorable cost-effectiveness calculations.

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Conflict of Interest

The authors declare that they have no competing interests.

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Supporting Information

Additional supporting information may be found in the online version of this article:

Figure S1. Selection criteria for the study sample.
Figure S2. Estimates of incremental net benefit (i.e., incremental cost-effectiveness ratio, ICER) and its 95% confidence intervals as a function of willingness-to-pay threshold for an additional life year (LY); (S2A) radiofrequency ablation (RFA) monotherapy versus no treatment; (S2B) surgical resection (SR) monotherapy versus no treatment; (S2C) liver transplantation (LT) monotherapy versus no treatment; (S2D) RFA plus SR versus no treatment; (S2E) RFA plus LT versus no treatment; (S2F) SR plus LT versus no treatment; (S2G) TACE plus RFA versus no treatment; (S2H) TACE plus SR versus no treatment; (S2I) TACE plus LT versus no treatment; and (S2J) RFA plus SR plus LT versus no treatment.

Figure S3. Efficiency frontier: plot of incremental quality-adjusted life years (QALYs) and costs of curative treatments relative to lowest cost scenario (no treatment): Sensitivity analysis according to (S3A) lower bound and (S3B) upper bound of pooled mean health state utilities of disease stage from published literature. The dotted diagonal line represents the willingness-to-pay for health effects (maximum acceptable ceiling ratio). If an intervention lies above the line, it will not be acceptable on cost-effectiveness grounds.

Figure S4. Efficiency frontier: Efficiency frontier: plot of incremental (A) life years (LYs) and (B) quality-adjusted life years (QALYs) and costs of curative treatments relative to lowest cost scenario (no treatment): Sensitivity analysis of excluding HCC stage IV.

Tables S1. ICD-9 and ICD-10 codes for patients diagnosed with diabetes, HIV, and liver disease stage.
Table S2. Screening and treatment procedures for patients with hepatocellular carcinoma.
Table S3. Estimation of utilities for noncirrhosis.
Table S4. Baseline characteristics of patients with hepatocellular carcinoma, 2002–2010.
Table S5. Estimates of incremental net benefit and probability of cost-effectiveness of curative treatment strategies for hepatocellular carcinoma compared with no treatment as a function of willingness-to-pay threshold per additional life year over the study period 2002–2010.