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**Outcomes of liver resection in hepatitis C virus-related intrahepatic cholangiocarcinoma:** **A systematic review and meta-analysis**

Cheo FY *et al*. Hepatitis C in ICC

Feng Yi Cheo, Kai Siang Chan, Vishal G Shelat

**Feng Yi Cheo,** Yong Loo Lin School of Medicine, National University of Singapore, Singapore 117597, Singapore

**Kai Siang Chan, Vishal G Shelat,** Department of General Surgery, Tan Tock Seng Hospital, Singapore 308433, Singapore

**Author contributions:** Cheo FY conception and design of the study, acquisition of data, analysis and interpretation of data, drafting the article, revising the article, final approval; Chan KS conception and design of the study, acquisition of data, analysis and interpretation of data, drafting the article, critical revision, final approval; Shelat VG interpretation of data, revising the article, critical revision, final approval.

**Corresponding author: Kai Siang Chan, MBBS, Doctor,** Department of General Surgery, Tan Tock Seng Hospital, No. 11 Jalan Tan Tock Seng, Singapore 308433, Singapore. kchan023@e.ntu.edu.sg

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**Abstract**

BACKGROUND

Cholangiocarcinoma is the second most common primary liver malignancy. Its incidence and mortality rates have been increasing in recent years. Hepatitis C virus (HCV) infection is a risk factor for development of cirrhosis and cholangiocarcinoma. Currently, surgical resection remains the only curative treatment option for cholangiocarcinoma. We aim to study the impact of HCV infection on outcomes of liver resection (LR) in intrahepatic cholangiocarcinoma (ICC).

AIM

To study the outcomes of curative resection of ICC in patients with HCV (*i.e.,* HCV+) compared to patients without HCV (*i.e.,* HCV-).

METHODS

We conducted a systematic review and meta-analysis of randomized controlled trials (RCTs) and observational studies to assess the outcomes of LR in ICC in HCV+ patients compared to HCV- patients in tertiary care hospitals. PubMed, Embase, The Cochrane Library and Scopus were systematically searched from inception till August 2023. Included studies were RCTs and non-RCTs on patients ≥ 18 years old with a diagnosis of ICC who underwent LR, and compared outcomes between patients with HCV+ *vs* HCV-. The primary outcomes were overall survival (OS) and recurrence-free survival. Secondary outcomes include perioperative mortality, operation duration, blood loss, intrahepatic and extrahepatic recurrence.

RESULTS

Seven articles, published between 2004 and 2021, fulfilled the selection criteria. All of the studies were retrospective studies. Age, incidence of male patients, albumin, bilirubin, platelets, tumor size, incidence of multiple tumors, vascular invasion, bile duct invasion, lymph node metastases, and stage 4 disease were comparable between HCV+ and HCV- group. Alanine transaminase [MD 22.20, 95%confidence interval (CI): 13.75, 30.65, *P* < 0.00001] and aspartate transaminase levels (MD 27.27, 95%CI: 20.20, 34.34, *P* < 0.00001) were significantly higher in HCV+ group compared to HCV- group. Incidence of cirrhosis was significantly higher in HCV+ group [odds ratio (OR) 5.78, 95%CI: 1.38, 24.14, *P* = 0.02] compared to HCV- group. Incidence of poorly differentiated disease was significantly higher in HCV+ group (OR 2.55, 95%CI: 1.34, 4.82, *P* = 0.004) compared to HCV- group. Incidence of simultaneous hepatocellular carcinoma lesions was significantly higher in HCV+ group (OR 8.31, 95%CI: 2.36, 29.26, *P* = 0.001) compared to HCV- group. OS was significantly worse in the HCV+ group (hazard ratio 2.05, 95%CI: 1.46, 2.88, *P* < 0.0001) compared to HCV- group.

CONCLUSION

This meta-analysis demonstrated significantly worse OS in HCV+ patients with ICC who underwent curative resection compared to HCV- patients.

**Key Words:** Cholangiocarcinoma; Bile duct cancer; Hepatitis C; Surgical resection; Hepatectomy

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**Core Tip:** Impact of hepatitis C virus (HCV) infection on survival outcomes in patients with intrahepatic cholangiocarcinoma (ICC) undergoing curative resection remains unclear. This is the first systematic review and meta-analysis comparing outcomes of surgical resection of ICC in HCV-positive patients *vs* HCV-negative patients. Our primary outcomes include overall survival (OS) and recurrence-free survival; secondary outcomes include perioperative mortality, operation duration, blood loss and recurrence. Our review and analysis demonstrated worse OS in HCV-positive patients compared to HCV-negative patients.

**INTRODUCTION**

Intrahepatic cholangiocarcinoma (ICC) is the second most common primary liver malignancy accounting for 15% of all primary liver malignancy, after hepatocellular carcinoma (HCC)[1]. Though rare, its incidence and mortality rates have increased in recent years[2,3]. Incidence amongst males increased from 1.51 per 100000 in 1993-1997 to 4.07 per 100000 in 2013-2017 and incidence amongst females increased from 1.73 per 100000 to 2.95 per 100000 respectively[4]. Mortality rates in cholangiocarcinoma have been reported to be up to 2 deaths per 100,000 in the United States, with mortality rates 3 times higher in Asia, and are still increasing[5]. Surgery remains the only potentially curative treatment modality in resectable ICC. However, the presentation for ICC is non-specific and patients may be diagnosed late; a retrospective study on patients with ICC demonstrated that 54% of ICCs were unresectable at diagnosis[6].

Common causes of ICC include cirrhosis, alcohol, hepatotoxins, chronic viral hepatitis, hepatolithiasis and liver fluke infections[7,8]. Patients with hepatitis C virus (HCV) have a 2-fold increase in risk of developing ICC compared to the general population[9]. To add on, a meta-analysis by Wang *et al*[10] in 2016 on 2842 patients with ICC showed that HCV was associated with worse survival [hazard ratio (HR) 2.64, 95% confidence interval (CI): 1.77-3.93] compared to controls. However, their study included patients who received various forms of treatment, ranging from curative surgery to palliative treatment. In 50 patients who received liver resection (LR) for ICC, Hai *et al*[11] however showed that HCV was not a predictor of survival following LR. While HCV is a significant risk factor for ICC, the prognostic significance of HCV remains uncertain for patients with ICC following LR. This study aims to perform a systematic review and meta-analysis to compare the survival between patients with HCV infection (*i.e*. HCV+) *vs* those without HCV (*i.e.* HCV-) in ICC following LR.

**MATERIALS AND METHODS**

***Study selection and search strategy***

This systematic review and meta-analysis was performed according to the Preferred Reporting Items for Systematic reviews and Meta-analysis (PRISMA) guidelines[12]. The protocol for this systematic review and meta-analysis was registered at PROSPERO (Ref no: CRD42023459605). A systematic search of the following databases (PubMed, Embase, The Cochrane Library and Scopus) was conducted for studies published from inception to 19th August 2023. A combination of the following search terms was used: “cholangiocarcinoma” or “bile duct cancer”, and “hepatectomy” or “liver resection”, and “hepatitis C” or “HCV”. The search was restricted to the title, abstract and keywords. The complete search strategy is appended in Supplementary Table 1. Search strategies for other databases were modified from the initial search strategy done on PubMed based on the database requirements.

Included studies were randomised controlled trials (RCTs) and non-RCTs on patients ≥ 18 years old with a diagnosis of ICC who underwent LR, and compared outcomes between patients with HCV+ *vs* HCV-. Exclusion criteria were studies: (1) On other types of liver malignancies (*e.g.*, HCC) or underwent liver transplantation (LT); (2) single-arm studies without comparison; (3) which did not include our outcome of interest; (4) on the same cohort of patients; and (5) based on article type (non-English studies, conference abstracts, case report or series, editorials, expert opinions, and review articles without original data). There were no studies which reported on the same cohort of patients. LR was defined as any form of surgical resection of the liver, including wedge resection, anatomical resection such as minor LR and major LR, and non-anatomical resection. HCV+ was defined as presence of anti-HCV antibodies detected on serology.

All cross-references were screened for potentially relevant studies not identified by the initial literature search. After removing duplicates, two authors screened abstracts for potential inclusion screening independently (Cheo FY and Chan KS). The included studies' full texts were reviewed and selected based on the inclusion and exclusion criteria. All discrepancies were resolved after review by the senior author (Shelat VG).

***Data extraction***

Data extraction was independently performed by two authors (Cheo FY and Chan KS). The following variables were extracted from each study: Publication details (name of first author, publication year and country), study characteristics (sample size, sex, age, Child-Pugh score, presence of cirrhosis, baseline tumor markers (alpha-fetoprotein, carbohydrate antigen 19-9, carcinoembryonic antigen, and tumor size). Our primary outcomes were overall survival (OS) and recurrence free survival (RFS). Our secondary outcomes were perioperative outcomes, including mortality, operation duration, blood loss, and tumor recurrence.

***Assessment of study quality***

Two authors (Cheo FY and Chan KS) independently assessed the included studies' quality. Observational studies were assessed using the modified Newcastle-Ottawa scale (Supplementary Table 2)[13]. No RCTs were included in this study. Only observational studies with sufficient quality (articles with a score >6) were included. Disagreements between authors were resolved by discussion with the senior author (Shelat VG).

***Statistical analysis***

Study variables were extracted to Microsoft Excel 365 (Microsoft®, Washington, United States). Categorical variables were described as *n* (%), and continuous variables were expressed as mean ± SD, or median [interquartile range (IQR)] unless otherwise specified. For continuous variables expressed only in median and range or IQR, mean and SD were estimated from median and range values using methods described by Wan *et al*[14]. Meta-analysis was performed using RevMan 5.4 (Review Manager 5.4, The Nordic Cochrane Centre, Copenhagen, Denmark). For cumulative OS and RFS, HR and standard error (SE) were estimated indirectly according to the methods described by Parmar *et al*[15]. Pooled HR was calculated through the inverse-variance method using the natural logarithm of HR [ln(HR)] and SE[16]. For studies that used univariate and multivariate analysis to assess the impact of HCV infection, the effect size from the multivariate analysis was used in our pooled analysis. Dichotomous outcomes were pooled and calculated using the Mantel-Haenszel method and expressed as odds ratio (OR) with 95%CI. Continuous outcomes were pooled and calculated using the inverse variance method and expressed as mean difference (MD) with 95%CI. Heterogeneity was assessed using Cochrane's Q and quantified by I2. If data was heterogenous (defined as I2 > 50%), a random-effects model using the DerSimonian and Laird approach was used. Statistical significance was defined as *P* < 0.05. Publication bias was investigated using funnel plots. Due to low sample size, quantitative analysis was not performed for short-term intra-operative and post-operative outcomes.

**RESULTS**

The systematic search identified 697 articles from the four databases. There were 492 articles after removal of the duplicates. Titles and abstracts of all the identified articles were screened. The remaining 53 articles underwent full-text review, of which seven articles were included in the final analysis[11,17-22]. The PRISMA diagram for the study selection process is appended in Figure 1. The funnel plots are appended in Supplementary Figure 1.

***Study characteristics***

There were seven studies with 1181 patients (HCV+ *n* = 205, HCV- *n* = 976)[11,17-22]. Kaibori *et al*[19] performed propensity score matching (PSM) analysis to derive their cohorts; only the PSM cohort was analysed in our study. Uenishi *et al*[20] performed univariate and multivariate analyses on the impact of HCV infection on outcomes of surgical resection in cholangiocarcinoma, of which outcomes of the multivariate analysis was included in our study. Yang *et al*[21] reported on OS of HCV+ and HCV- groups in the early relapse subgroup (within 24 mo), which we excluded from our quantitative analysis of OS due to selection bias and misrepresentation of the entire cohort of ICC. While the study by Hai *et al*[11] performed Kaplan-Meier analysis on OS, HR and SE could not be estimated due to the lack of clarity of the Kaplan-Meier curve; clinical demographics and other outcomes were still included. The study by Terakawa *et al*[22] was reported in Japanese; however, as the tables and figures were in English, relevant data such as survival outcomes were included in our study to avoid dilution to power and sample size.

***Clinical demographics***

The study characteristics and patient demographics of individual studies were summarized in Table 1. The overall mean age was 65.0 years, and 17.2% (*n* = 46/267) patients had cirrhosis. There were 17.7% (*n* = 55/311) patients with multiple tumors on diagnosis, and the mean tumor size ranged from 3.6-6.4 cm. 12.5% of patients (*n* = 16/128) had synchronous HCC lesions. Pooled analysis showed that age, incidence of male patients, albumin, bilirubin, platelets, tumor size, incidence of multiple tumors, vascular invasion, bile duct invasion, lymph node metastases, stage 4 disease were comparable between HCV+ and HCV- (Table 2). However, alanine transaminase (ALT) and aspartate transaminase (AST) levels were significantly higher in HCV+ (ALT: MD +22.2, 95%CI: 13.75, 30.65; AST: MD +27.27, 95%CI: 20.20, 34.34) compared with HCV-. There were also more patients with cirrhosis (OR 5.78, 95%CI: 1.38, 24.14, *P* = 0.02), poorly differentiated disease (OR 2.55, 95%CI: 1.34, 4.82, *P* = 0.004), and concomitant HCC (OR 8.31, 95%CI: 2.36, 29.26, *P* = 0.001) in the HCV+ group compared to HCV- group.

***Oncological outcomes***

Table 3 summarizes the survival outcomes reported in individual studies. Four studies involving 841 patients (HCV+ *n* = 145, HCV- *n* = 696) reported on OS[18-20,22]. Pooled HR showed statistically significantly worse OS in the HCV+ group (HR 2.05, 95%CI: 1.46, 2.88, *P* < 0.0001) (Figure 2). Heterogeneity was not significant among the studies (I2 = 0%, *P* = 0.56). The study by Cai *et al*[18] had very few HCV+ patients compared with HCV- patients (HCV+ *n* = 3, HCV- *n* = 527). In view of this, sensitivity analysis was performed to exclude their study; OS remained significantly worse in HCV+ group (HR 2.07, 95%CI: 1.45, 2.96, *P* < 0.0001) compared to HCV- group.

There were 2 studies involving 294 patients (HCV+ *n* = 135, HCV- *n* = 159) which reported on RFS[19,20]. Meta-analysis was not performed for RFS due to the small sample size and limitations in interpretation. Kaibori *et al*[19] reported significantly worse RFS in the HCV+ group (HR 1.61, 95%CI: 1.09, 2.38, *P* = 0.016) compared to the HCV- group. Uenishi *et al*[20] reported comparable RFS between the HCV+ group and HCV- group (HR 1.59, 95%CI: 0.74, 3.41, *P* = 0.24).

***Secondary outcomes***

There was one study which reported on incidence of post-operative mortality. Uenishi *et al*[20] reported in-hospital mortality of 13% in HCV+ group (*n* = 3/33) and 4.8% in HCV- group (*n* = 1/57). However this did not reach statistical significance (*P* = 0.609).

There was one study which reported on operative time and intraoperative blood loss. Terakawa*et al*[22] reported mean operative duration of 359 ± 74 min in the HCV+ group compared to 336 ± 34 min in the HCV- group. They additionally reported mean intraoperative blood loss of 2037 ± 577 mL in HCV+ group compared to 1226 ± 269 mL in HCV- group. However, no comparative statistical analysis was performed to compare between HCV+ and HCV- groups[22].

There were two studies which reported on tumor recurrence post-LR; Yang *et al*[21] reported comparable tumor recurrence (both intrahepatic and extrahepatic) in HCV+ group and HCV- group (HR 3.28, 95%CI: 0.80, 13.51, *P* = 0.098) . Kaibori *et al*[19] showed comparable incidence of intrahepatic recurrence [HCV+: 36% (*n* = 33/92), HCV-: 30% (*n* = 28/94); *P* = 0.467] and extrahepatic recurrence [HCV+: 36% (*n* = 25/69) *vs* HCV-: 27% (*n* = 18/67); *P* = 0.322] between HCV+ and HCV- groups.

**DISCUSSION**

Hepatitis B virus (HBV) and HCV infection are significant risk factors involved in the pathogenesis of cholangiocarcinoma. Interestingly, while HBV infection has been shown to provide favourable prognosis for patients with cholangiocarcinoma, HCV+ is associated with shorter OS compared to HCV- patients[10]. Our study similarly showed that HCV+ is associated with worse OS in ICC following LR. However, there is limited data on peri-operative outcomes.

Risk factors for ICC include biliary tract diseases such as primary sclerosing cholangitis, recurrent pyogenic cholangitis, primary biliary cirrhosis, congenital malformations of the bile duct (*i.e* choledochal cysts), cirrhosis and chemical exposure[23]. Incidence of HCV has been reported to be 13.8%-23.1% in ICC[24,25]. Various mechanisms have been proposed on the role of HCV in the pathogenesis of ICC[26]. One postulation is that cholangiocytes and hepatocytes share the same liver progenitor cell; cholangiocytes express receptors which are susceptible to HCV infection[27]. Another postulation is that the initial HCV infection of hepatocytes result in transdifferentiation of hepatocytes into cholangiocytes[28]. Interaction of cholangiocytes with HCV protein induces chronic biliary inflammation with resulting development of ICC.

HCV is a significant risk factor in the development of cholangiocarcinoma[29]. Globally, HCV is strongly associated with cholangiocarcinoma, especially in the Western populations[9]. Therefore, an understanding of its impact on outcomes helps to guide clinical decisions and development of treatment pathways. A previous meta-analysis by Wang *et al*[10] explored the impact of HCV infection on survival outcomes in patients with ICC, regardless of treatment modality, and showed poorer prognosis in HCV+ patients. However, we wish to understand the implications of HCV on long-term outcomes following curative LR in ICC. Since then, more studies comparing LR outcomes in ICC between HCV+ and HCV- groups have been published. This updated meta-analysis included five new studies with 1053 patients; we showed that HCV+ patients had worse OS compared to HCV- in patients who received curative LR for ICC[17-19,21,22]. We hypothesize potential reasons for these observations which are discussed below.

Several factors prognosticate OS and RFS in ICC following LR, including cirrhosis, positive surgical margins, tumor morphology patterns, tumor size, nodal involvement, and vascular invasion[30,31]. Chronic HCV is recognised as a significant precursor to liver cirrhosis, due to its process of chronic hepatocellular injury leading to chronic inflammation, resulting in scarring and fibrosis[32]. Cirrhosis has been associated with worse short-term and long-term survival; for instance, Zaydfudim *et al*[33] reported higher postoperative mortality (OR = 2.24; 95%CI: 1.16, 4.34, *P* = 0.016) in patients with cirrhosis; Sasaki *et al*[34] reported worse 5-year disease-specific survival (75.4% in patients with normal liver function *vs* 59.1% in patients with cirrhosis, *P* = 0.04) in cirrhotic patients as well. Liver cirrhosis is also a risk factor of tumor recurrence in cholangiocarcinoma; Tsilimigras *et al*[35] reported a significant association between cirrhosis and very early recurrence (within 6 mo after resection) of ICC post-LR (OR 2.06, 95%CI: 1.25, 3.40, *P* = 0.005) and Zhang *et al*[36] reported a significant association between cirrhosis and late intrahepatic recurrence (more than 24 mo after resection) (HR 1.99, 95%CI: 1.11, 3.56, *P* = 0.019). This may be due to the increased carcinogenic potential of remnant cirrhotic liver and biliary system which predisposes to neocarcinogenesis, resulting in de novo recurrence of cholangiocarcinoma[26]. While our meta-analysis showed that HCV+ group had worse OS compared to HCV- group, there was also increased incidence of liver cirrhosis in the HCV+ group (OR 5.78, 95%CI: 1.38, 24.14, *P* = 0.02). Liver cirrhosis may be a confounding factor for worse OS in HCV+ ICC as discussed above, rather than HCV alone.

An important consideration in surgical candidates is the risk of post hepatectomy liver failure (PHLF)[37]; Lei *et al*[38] showed that patients with PHLF diagnosed using the 50-50 criteria was independently associated with higher 90-d mortality (HR 8.63, 95%CI: 3.33-22.35, *P* < 0.001). Clinically relevant PHLF (grade B/C) has been reported to be associated with postoperative 90-d mortality (OR 7.26, 95%CI: 2.90, 18.17) and significantly worse long-term survival outcomes (HR 1.90, 95%CI:1.32, 2.71)[37]. Post-LR, adequate functional liver remnant (FLR) is required to sustain the body’s metabolic, synthetic and detoxifying requirements[39]. Due to chronic hepatocellular injury leading to scarring and fibrosis in cirrhotic livers, these functions are greatly reduced, predisposing to liver failure[40]. Current guidelines recommend FLR of > 30% in patients with liver steatosis and > 40% in patients with cirrhosis to reduce risk of PHLF[41,42]. One of the possible reasons for worse OS in HCV+ ICC may be due to PHLF in the HCV+ group due to higher incidence of cirrhosis. Unfortunately, in our review, none of the included studies described the incidence of PHLF; this remains a postulation to be validated, and correlation cannot be drawn. Nevertheless, other markers have been used to predict risk of PHLF, such as the use of indocyanine green retention rate at 15 minutes (ICGR15). Makuuchi’s criteria serve as a guide to assess the extent of hepatectomy based on ICGR15 to reduce risk of PHLF[43]. In our review, there was a mix of studies reporting either comparable ICGR15 between HCV+ and HCV- groups (such as the study by Hai *et al*[11] with comparable incidence of ICGR15 > 10% in HCV+ group (*n* = 11/17, 64.7%) compared to HCV- group (*n* = 4/21, 19.0%), *P* = 0.0656), or higher ICGR15 in HCV+ compared to HCV- (such as the study by Kaibori *et al*[19] with 71% with ICGR15 ≥10% in HCV+ compared to 48% in HCV-). Whether or not PHLF is a cause of worse OS in HCV+ ICC following LR remains to be answered.

Another possible reason for poorer prognosis in HCV+ patients may be attributed to synchronous or metachronous HCC in HCV+ patients. Chronic HCV infection is the leading cause of HCC in Western countries. HCV is also associated with a large proportion of HCC in certain Asian and African countries[44,45]. Carcinogenesis of HCC and cholangiocarcinoma in the background of chronic HCV-induced cirrhosis share similarities and has been postulated to be associated with the occurrence of synchronous or metachronous HCC and cholangiocarcinoma lesions[46]. A literature review of reported synchronous HCC and cholangiocarcinoma cases by Watanabe *et al*[47] found that 72.7% of cases were positive for HCV. Survival outcomes in patients with synchronous or metachronous HCC and cholangiocarcinoma are generally poorer and may distort survival outcomes in HCV+ group[48]. In our study, incidence of simultaneous HCC lesions found on pathologic studies is significantly higher in HCV+ group compared to HCV- group (OR 8.31, 95%CI: 2.36, 29.26, *P* = 0.001), which may confound and contribute to worse outcomes in the HCV+ group.

Tumor biology is another important consideration in survival. Higher tumor grade and poorly differentiated tumors confer a worse prognosis on survival. A retrospective study by Mao *et al*[49] identified tumor differentiation as an independent predictor of higher postoperative mortality in cholangiocarcinoma (relative risk 1.356, 95%CI: 1.081, 1.699, *P* = 0.008). Nickkholgh *et al*[50] reported that high grade tumor (defined as Grade 3-4) was an independent determinant of recurrence in ICC post-resection (HR 1.63, 95%CI: 1.04, 2.55, *P* = 0.034). HCV-induced development and progression of liver fibrosis involve epithelial-mesenchymal transition (EMT) of cholangiocytes, resulting in reduced expression of E-cadherin, which is associated with poor tumor differentiation in cholangiocarcinoma[26,51,52]. In our study, incidence of poorly differentiated ICC was significantly greater in HCV+ group compared to HCV- group (OR 2.55, 95%CI: 1.34, 4.82, *P* = 0.004). This may have consequently resulted in worse OS in the HCV+ group.

Advanced tumor stage and metastatic disease are poor prognostic factors in cholangiocarcinoma[53]. In advanced tumors, several factors contribute to more aggressive tumor behavior. Notably, presence of vascular invasion increases the risk of haematogenous spread of tumor cells, and tumor multiplicity provide additional nidus for tumor to grow and spread from[54-56]. Expectedly as well, nodal disease has been shown to be associated with worse survival (22.9 mo *vs* 30.1 mo, *P* = 0.03)[57]. The question lies in whether HCV+ increases the risk of more advanced disease or nodal metastases, since HCV infection results in EMT as described above[58]. This question remains unanswered based on our findings, but may be due to the low sample size of the included studies.

The advent of direct-acting antivirals (DAAs) have revolutionized the treatment of HCV, where it is possible to achieve a cure for hepatitis C[59]. The American Association for the Study of Liver Diseases recommends first-line therapy with glecaprevir/pibrentasvir and sofosbuvir/velpatasvir for 8 wk and 12 wk respectively for treatment-naïve adults[60]. However, there are no guidelines on antiviral therapy duration for patients with HCV+ HCC or ICC. In HCV-related HCC, HCV eradication therapy has been proven to improve long-term outcomes of HCC undergoing curative treatment[61-63]. Further meta-analyses suggest benefits of HCV treatment on long-term HCC survival outcomes[64,65]. While the literature on the utility of DAAs in HCV+ ICC is scarce, the oncogenesis of ICC is similar to that of HCC. Hence, we theorize similar benefits of HCV eradication therapy in the ICC population.

Adjuvant chemotherapy is recommended for patients with resected cholangiocarcinoma[66]. The American Society of Clinical Oncology recommends the use of adjuvant capecitabine as first-line therapy for 6 mo[67]. However, certain chemotherapy agents are hepatotoxic and may exacerbate or accelerate fibrosis in HCV+ patients with chronic liver inflammation. Studies have suggested that treatment of HCV infection may also reverse cirrhosis in some group of patients (*e.g.* those without decompensated liver cirrhosis), allowing for the use of adjunct treatment such as chemotherapy[68]. Unfortunately, the use of adjuvant chemotherapy and underlying liver function was not discussed in the included studies and this falls beyond the scope of our study. The combined role of DAAs and adjuvant chemotherapy on underlying liver function and long-term survival should be evaluated.

While our study excluded patients who underwent LT, the use of LT in treating ICC is worth exploring. LT was previously contraindicated in managing ICC due to poor outcomes and high recurrence post-LT. Initial studies reported 3-year OS post-LT ranging from 4.9%-39.0% without receiving pre-transplant treatment and 3-year RFS rate of 28.8-35.0%[69-71]. However, recent studies have reported reasonable outcomes in certain groups of patients with ICC who received LT, with 3-year OS rates post-LT ranging from 47.9%-83.3% and 5-year OS rates ranging from 31.3%-83.3%. 3-year RFS rates also ranged from 41.7%-52.0% in newer studies[72-74]. Transplant outcomes have improved drastically due to improved effectiveness of neoadjuvant therapy such as gemcitabine-based systemic chemotherapy and locoregional therapy including trans-arterial chemoembolization and radiofrequency ablation, in addition to protocols to determine eligibility for LT in patients who demonstrate disease stability or pathological response to these pre-transplant treatment modalities[73-75]. With these improvements in preoperative treatment and a more stringent organ recipient selection process, LT may provide an alternative treatment of cure as standard of care for ICC in the future. Additionally, LT also deals with the problem of cirrhosis and PHLF that comes with LR, which may be a contributing factor to worse survival. With ongoing trials assessing outcomes of LT in ICC currently underway, we anticipate treatment of cholangiocarcinoma to evolve in the future[76-78]. Although not explored in our study, subsequent studies could analyse the impact of HCV infection on outcomes following other treatment modalities in ICC.

There are a few limitations in our study. All the included studies were retrospective observational studies which have inherent selection bias. The absence of high-quality evidence from RCTs and prospective studies may limit interpretation of the outcomes from our analysis. Subsequent studies should employ methods such as PSM and RCTs to reduce bias for more conclusive results. Nevertheless, quality assessment was performed for the included studies and all the included studies had at least moderate quality evidence. The number of studies included in this meta-analysis is relatively small due to our strict inclusion criteria of studies comparing post-hepatectomy outcomes of ICC in HCV+ and HCV- subgroups. All included studies were conducted in Asia, namely Japan and China, despite including ICC globally, hence causing possible limitations in the generalizability of our results. Global incidence of cholangiocarcinoma is highest in Asia, especially Japan[79]. However, incidence of cholangiocarcinoma is rising in Western countries over the past decade, of which their population is underrepresented in our study[80]. Prevalence of chronic HCV infection share a different distribution globally, with middle-low-income countries in the Eastern Mediterranean and European regions suffering the highest burden of disease[81]. Ideally, a more heterogenous sample including these populations would produce results that may be more representative of the global population, hence future studies involving patients from regions of high HCV and cholangiocarcinoma prevalence will provide more insight. We could not perform meta-analysis on RFS and our secondary outcomes due to inadequate data from our included studies. Lastly, this study did not include subgroup analyses of tumors undergoing major hepatectomy. Performing major hepatectomy on a background cirrhotic liver or chronically HCV-infected liver has its additional risks (*e.g*. PHLF and post-operative mortality). Thus, a separate analysis focusing on this subgroup may provide valuable insight and guidance in management.

**CONCLUSION**

Our meta-analysis demonstrated that HCV infection is associated with significantly worse OS in ICC patients undergoing LR with curative intent. Further studies of the underlying mechanisms of oncogenesis of the biliary tree in HCV infection, including genetic and basic science studies are warranted to understand its disease process. More prospective studies with PSM-derived cohorts including analysis of other aspects of treatment such as PHLF and liver augmentation strategies should be conducted to validate our findings.

**ARTICLE HIGHLIGHTS**

***Research background***

Incidence of intrahepatic cholangiocarcinoma (ICC) has been rising over the past decade. Hepatitis C virus (HCV) infection is an important risk factor in the development of ICC. Currently, liver resection (LR) remains the only curative treatment modality for ICC. Our study aims to study the outcomes of LR in ICC patients with HCV-positive (HCV+) compared to HCV-negative (HCV-) ICC patients.

***Research motivation***

Long-term outcomes of curative LR in ICC can be affected by patient and tumor characteristics. The impact of HCV infection on post-LR outcomes should be reviewed and quantitatively concluded.

***Research objectives***

We aim to identify HCV+ patients as a high-risk subgroup amongst ICC patients undergoing curative LR. Our analysis concluded that HCV+ patients had worse overall survival compared to HCV- patients following LR. Our findings act as a stepping stone for future studies to validate our findings, to determine a cause for this outcome, as well as to devise strategies to improve outcomes in HCV+ ICC patients undergoing curative LR.

***Research methods***

Four databases (PubMed, Embase, Scopus and The Cochrane Library) were systematically searched for relevant studies, which were subsequently screened for inclusion in our study based on our inclusion criteria. We assessed the quality of included observational studies using the modified Newcastle-Ottawa Scale. There were no randomised controlled trials included in our study. Our primary outcomes were overall survival (OS) and recurrence-free survival. Secondary outcomes include perioperative mortality, operation duration, blood loss, intrahepatic and extrahepatic recurrence. Study variables, primary and secondary outcomes were extracted from included studies. Pooled hazard ratio (HR) was calculated through the inverse-variance method using the natural logarithm of HR [ln (HR)] and standard error. Dichotomous outcomes were pooled and calculated using the Mantel-Haenszel method and expressed as odds ratio (OR) with 95% confidence interval (CI). Continuous outcomes were pooled and calculated using the inverse variance method and expressed as mean difference with 95%CI.

***Research results***

Our meta-analysis demonstrated significantly worse OS in HCV+ patients with ICC that underwent curative resection compared to HCV- patients (HR 2.05, 95%CI: 1.46, 2.88, *P* < 0.0001). Our analysis also showed increased incidence of cirrhosis (OR 5.78, 95%CI: 1.38, 24.14, *P* = 0.02), poorly differentiated tumors (OR 2.55, 95%CI: 1.34, 4.82, *P* = 0.004), as well as simultaneous hepatocellular carcinoma (HCC) lesions in HCV+ patients (OR 8.31, 95%CI: 2.36, 29.26, *P* = 0.001) compared with HCV- patients. Our findings identify HCV infection as a significant poor prognostic factor in ICC patients undergoing curative LR and as a significant risk factor of liver cirrhosis, poor tumor differentiation and incidence of simultaneous HCC lesions. However, the presence of increased liver cirrhosis and poor tumor differentiation may be confounding factors for worse OS in HCV+ patients. No statistically significant differences were noted between HCV+ and tumor stage, tumor invasion and metastases in our study.

***Research conclusions***

Our study concluded that HCV infection is associated with significantly worse OS outcomes in ICC post-LR. This may be confounded by increased incidence of cirrhosis and poorly differentiated tumors with HCV infection. The exact pathophysiology and confirmation of our findings ought to be explored in future well-designed prospective studies. The role of viral eradication therapy and chemotherapy in this subgroup of patients should also be explored.

***Research perspectives***

Future research should be performed with randomized controlled trials or propensity score matched cohorts to validate our findings. Further studies should also explore the role of adjuncts such as anti-viral therapy and adjuvant chemotherapy in HCV+ ICC patients who underwent curative LR.

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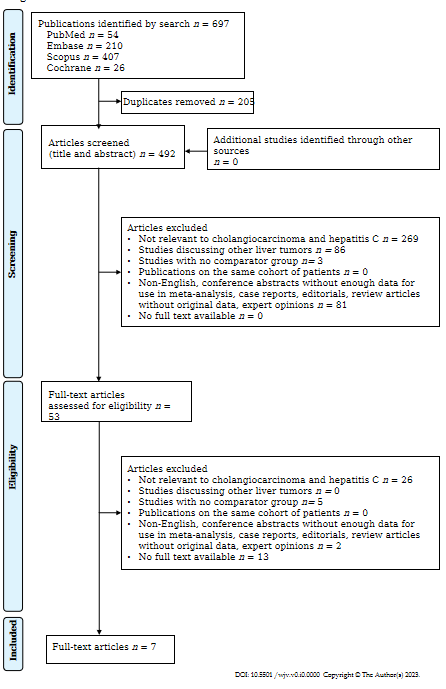
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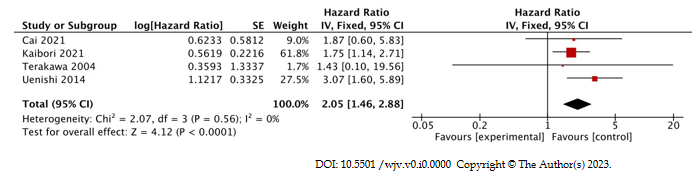
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**Figure Legends**



**Figure 1 Preferred Reporting Items for Systematic reviews and Meta-Analyses flowchart for study selection.**



**Figure 2 Comparison of overall survival between hepatitis C virus-positive group and hepatitis C virus-negative group in patients with intrahepatic cholangiocarcinoma post-liver resection.** CI: Confidence interval.

**Table 1 Baseline characteristics and patient demographics of the included studies (*n* = 7)**

|  |  |  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **No.** | **Ref.** | **Study design** | **Study period** | **Country** | **Sample size, *n* (%)** | **Age, yr** | **Males, *n* (%)** | **Tumor size, cm** | **Cirrhosis, *n*** | **Tumor stage** | **Tumor grade** |
| 1 | Hai*et al*[11], 2005 | Retrospective | Jan 1997-Dec 2002 | Japan | HCV+: 17; HCV-: 21 | HCV+: 69.0 ± 4.9; HCV-: 60.6 ± 12.4 | HCV+: 10; HCV-: 13 | HCV+: 3.6 ± 2.3; HCV-: 6.4 ± 4.5 | NR | HCV+:I: 4, II: 4, III: 3, IV: 6; HCV-: I: 0, II: 6, III: 7, IV: 8 | NR |
| 2 | Kaibori*et al*[19], 20211 | Retrospective | Jan 2000-Dec 2007 | Japan | HCV+: 102; HCV-: 102 | HCV+: ≥ 70: 56/102; HCV-: ≥ 70: 59/102 | HCV+: 64; HCV-: 74 | HCV+: ≥ 3.5cm: 69/102; HCV-: ≥ 3.5cm: 61/102 | HCV+: 24; HCV-: 8 | NR | HCV+:well: 14, moderate: 49, poor: 27; HCV-:well: 21, moderate: 53, poor: 11 |
| 3 | Uenishi*et al*[20], 2014 | Retrospective | Jan 2000-Dec 2011 | Japan | HCV+: 33  HCV-: 57 | HCV+: 66.9 ± 9.0; HCV-: 64.3 ± 11.2 | HCV+: 23;  HCV-: 38 | HCV+: 4.7 ± 1.7; HCV-: 4.8 ± 2.6 | HCV+: 14 HCV-: 3 | HCV+:I: 12, II: 7, III: 5, IV: 9; HCV-: I: 18, II: 9, III: 5, IV: 25 | HCV+: poor: 7; HCV-: poor: 7 |
| 4 | Cai*et al*[18], 2021 | Retrospective | Dec 2008-Dec 2017 | China | HCV+: 3; HCV-: 527 | NR | NR | NR | NR | NR | NR |
| 5 | Ariizumi*et al*[17], 2011 | Retrospective | 1989-2008 | Japan | HCV+: 42; HCV-: 92 | NR | NR | NR | NR | NR | NR |
| 6 | Yang*et al*[21], 2019 | Retrospective | Jan 2005-Dec 2011 | China | HCV+: 1; HCV-: 167 | NR | NR | NR | NR | NR | NR |
| 7 | Terakawa*et al*[22], 2004 | Retrospective | Jan 1992-Dec 2001 | Japan | HCV+: 7; HCV-: 10 | HCV+:64.0 ± 3.0; HCV-: 66.0 ± 3.0 | HCV+: 4; HCV-: 8 | HCV+: 5.0 ± 1.2; HCV-: 5.1 ± 1.0 | NR | HCV+:II: 1, III: 3, IV: 3HCV-: II: 2, III: 4, IV: 4 | HCV+:well: 1, moderate: 4; HCV-: well: 1, moderate: 5 |

1Values included in this study is obtained after propensity score matching. HCV: Hepatitis C Virus; NR: Not reported.

**Table 2 Summary of effect size of different study variables and outcomes between hepatitis C virus-positive group and hepatitis C virus-negative group**

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **No.** | **Study variables and/or outcomes** | **No. of data sets** | **Total number of patients, *n* (HCV+/HCV-)** | **No. of patients (%)** | | **Effect Size, OR (95%CI)/MD (95%CI)/HR (95%CI)1** | ***P* value** | ***I*2, %** | **Model used** |
| **HCV+** | **HCV-** |
| Demographics and histopathological findings | | | | | | | | | |
| 1 | Age, yr | 4 | 349 (159/190) | NA | | 2.55 (-3.09, 8.20) | 0.38 | 82 | RE |
| 2 | Male | 4 | 349 (159/190) | 101 (63.5) | 133 (70.0) | 0.74 (0.47, 1.17) | 0.20 | 0 | FE |
| 3 | ALT, IU/L | 2 | 294 (135/159) | NA | | 22.20 (13.75, 30.65) | < 0.00001a | 0 | FE |
| 4 | AST, IU/L | 2 | 294 (135/159) | NA | | 27.27 (20.20, 34.34) | < 0.00001a | 0 | FE |
| 5 | Albumin, g/L | 2 | 294 (135/159) | NA | | -0.11 (-0.34, 0.12) | 0.34 | 71 | RE |
| 6 | Bilirubin, umol/L | 2 | 294 (135/159) | NA | | -0.07 (-0.33, 0.19) | 0.61 | 84 | RE |
| 7 | Platelets, 104/mm3 | 2 | 294 (135/159) | NA | | -1.96 (-5.88, 1.96) | 0.33 | 71 | RE |
| 8 | Tumor size, cm | 3 | 145 (57/88) | NA | | -0.61 (-1.79, 0.57) | 0.31 | 62 | RE |
| 9 | Multiple tumors | 3 | 311 (142/169) | 25 (17.6) | 30 (17.8) | 1.12 (0.61, 2.06) | 0.70 | 0 | FE |
| 10 | Cirrhosis | 2 | 294 (135/159) | 35 (25.9) | 11 (6.9) | 5.78 (1.38, 24.14) | 0.02**a** | 69 | RE |
| 11 | Vascular invasion | 3 | 145 (57/88) | 20 (35.1) | 36 (40.9) | 0.76 (0.37, 1.54) | 0.45 | 0 | FE |
| 12 | Bile duct invasion | 2 | 294 (135/159) | 53 (39.3) | 64 (40.3) | 0.86 (0.53, 1.39) | 0.53 | 0 | FE |
| 13 | Lymph node metastases | 4 | 349 (159/190) | 41 (25.8) | 54 (28.4) | 0.85 (0.53, 1.37) | 0.51 | 15 | FE |
| 14 | Stage 4 | 3 | 145 (57/88) | 18 (31.6) | 37 (42.0) | 0.64 (0.32, 1.28) | 0.21 | 0 | FE |
| 15 | Poorly differentiated | 2 | 273 (127/146) | 34 (26.8) | 18 (12.3) | 2.55 (1.34, 4.82) | 0.004a | 0 | FE |
| 16 | Simultaneous HCC lesions | 2 | 128 (50/78) | 13 (26.0) | 3 (3.8) | 8.31 (2.36, 29.26) | 0.001a | 0 | FE |
| Outcomes | | | | | | | | | |
| 17 | Overall survival | 4 | 841 (145/696) | NA | | 2.05 (1.46, 2.88) | <0.0001a | 0 | FE |

1Odds ratio and 95% confidence interval (CI) was used for dichotomous outcomes, mean difference and 95%CI was used for continuous outcomes, and hazards ratio and 95%CI was used for time-to-event outcomes.

aData with statistical significance (*P* < 0.05).

ALT: Alanine transaminase; AST: Aspartate transaminase; CI: Confidence interval; FE: Fixed-effects; HCV: Hepatitis C virus; HR: Hazards ratio; I2: Heterogeneity; MD: Mean difference; NA: Not applicable; OR: Odds ratio; RE: Random-effects.

**Table 3 Survival outcomes reported in the included studies (*n* = 7)**

|  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **No.** | **Ref.** | **1-yr OS, %** | **1-yr DFS, %** | **3-yr OS, %** | **3-yr RFS, %** | **3-yr DFS, %** | **5-yr OS, %** | **5-yr RFS, %** |
| 1 | Hai*et al*[11], 2005 | HCV+: 70.9; HCV-: 75.6 | HCV+: 55.7; HCV-: 49.0 | HCV+: 41.4; HCV-: 30.1 | NR | HCV+: 27.9; HCV-: 32.7 | NR | NR |
| 2 | Kaibori*et al*[19], 20211 | NR | NR | NR | NR | NR | HCV+: 32.2; HCV-: 44.7 | HCV+: 25.0; HCV-: 31.3 |
| 3 | Uenishi*et al*[20], 2014 | NR | NR | HCV+: 30.6; HCV-: 65.6 | HCV+: 29.9; HCV-: 31.4 | NR | HCV+: 21.9; HCV-: 32.8 | HCV+: 22.4; HCV-: 20.6 |
| 4 | Cai*et al*[18], 2021 | NR | NR | NR | NR | NR | NR | NR |
| 5 | Ariizumi*et al*[17], 2011 | NR | NR | NR | NR | NR | HCV+: 53; HCV-: 32 | NR |
| 6 | Yang*et al*[21], 2019 | NR | NR | NR | NR | NR | NR | NR |
| 7 | Terakawa*et al*[22], 2004 | NR | NR | NR | NR | NR | NR | NR |

1Values included is this study is obtained after propensity score matching.

DFS: Disease-free survival; HCV: Hepatitis C virus; NR: Not reported; OS: Overall survival; RFS: Recurrence-free survival.



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