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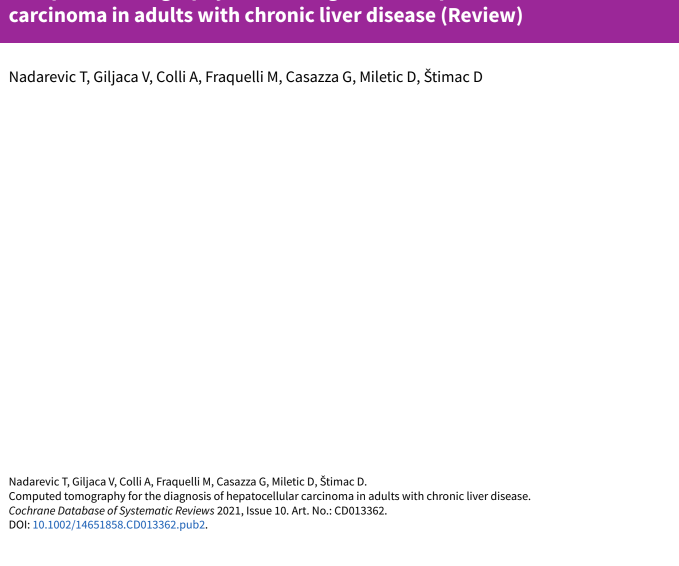






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[Diagnostic Test Accuracy Review]

Computed tomography for the diagnosis of hepatocellular carcinoma in adults with chronic liver disease

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ABSTRACT

Background

Hepatocellular carcinoma occurs mostly in people with chronic liver disease and ranks sixth in terms of global incidence of cancer, and fourth in terms of cancer deaths. In clinical practice, computed tomography (CT) is used as a second-line diagnostic imaging modality to confirm the presence of focal liver lesions suspected as hepatocellular carcinoma on prior diagnostic test such as abdominal ultrasound or alpha-foetoprotein, or both, either in surveillance programmes or in clinical settings. According to current guidelines, a single contrast-enhanced imaging study CT or magnetic resonance imaging (MRI) showing typical hallmarks of hepatocellular carcinoma in people with cirrhosis is valid to diagnose hepatocellular carcinoma. However, a significant number of hepatocellular carcinomas do not show typical hallmarks on imaging modalities, and hepatocellular carcinoma is, therefore, missed. There is no clear evidence of the benefit of surveillance programmes in terms of overall survival: the conflicting results can be a consequence of inaccurate detection, ineffective treatment, or both. Assessing the diagnostic accuracy of CT may clarify whether the absence of benefit could be related to underdiagnosis. Furthermore, an assessment of the accuracy of CT in people with chronic liver disease, who are not included in surveillance programmes is needed for either ruling out or diagnosing hepatocellular carcinoma.

Objectives

Primary: to assess the diagnostic accuracy of multidetector, multiphasic contrast-enhanced CT for the diagnosis of hepatocellular carcinoma of any size and at any stage in adults with chronic liver disease, either in a surveillance programme or in a clinical setting.

Secondary: to assess the diagnostic accuracy of CT for the diagnosis of resectable hepatocellular carcinoma in adults with chronic liver disease.

Search methods

We searched the Cochrane Hepato-Biliary Trials Register, Cochrane Hepato-Biliary Diagnostic-Test-Accuracy Studies Register, the Cochrane Library, MEDLINE, Embase, LILACS, Science Citation Index Expanded, and Conference Proceedings Citation Index – Science until 4 May 2021. We applied no language or document-type restrictions.



Selection criteria

Studies assessing the diagnostic accuracy of CT for the diagnosis of hepatocellular carcinoma in adults with chronic liver disease, with cross-sectional designs, using one of the acceptable reference standards, such as pathology of the explanted liver and histology of resected or biopsied focal liver lesion with at least a six-month follow-up.

Data collection and analysis

At least two review authors independently screened studies, extracted data, and assessed the risk of bias and applicability concerns, using the QUADAS-2 checklist. We presented the results of sensitivity and specificity, using paired forest plots, and tabulated the results. We used a hierarchical meta-analysis model where appropriate. We presented uncertainty of the accuracy estimates using 95% confidence intervals (CIs). We double-checked all data extractions and analyses.

Main results

We included 21 studies, with a total of 3101 participants. We judged all studies to be at high risk of bias in at least one domain because most studies used different reference standards, often inappropriate to exclude the presence of the target condition, and the time-interval between the index test and the reference standard was rarely defined. Regarding applicability in the patient selection domain, we judged 14% (3/21) of studies to be at low concern and 86% (18/21) of studies to be at high concern owing to characteristics of the participants who were on waiting lists for orthotopic liver transplantation.

CT for hepatocellular carcinoma of any size and stage: sensitivity 77.5% (95% CI 70.9% to 82.9%) and specificity 91.3% (95% CI 86.5% to 94.5%) (21 studies, 3101 participants; low-certainty evidence).

CT for resectable hepatocellular carcinoma: sensitivity 71.4% (95% CI 60.3% to 80.4%) and specificity 92.0% (95% CI 86.3% to 95.5%) (10 studies, 1854 participants; low-certainty evidence).

In the three studies at low concern for applicability (861 participants), we found sensitivity 76.9% (95% CI 50.8% to 91.5%) and specificity 89.2% (95% CI 57.0% to 98.1%).

The observed heterogeneity in the results remains mostly unexplained. The sensitivity analyses, which included only studies with clearly prespecified positivity criteria and only studies in which the reference standard results were interpreted without knowledge of the results of the index test, showed no variation in the results.

Authors' conclusions

In the clinical pathway for the diagnosis of hepatocellular carcinoma in adults with chronic liver disease, CT has roles as a confirmatory test for hepatocellular carcinoma lesions, and for staging assessment. We found that using CT in detecting hepatocellular carcinoma of any size and stage, 22.5% of people with hepatocellular carcinoma would be missed, and 8.7% of people without hepatocellular carcinoma would be unnecessarily treated. For resectable hepatocellular carcinoma, we found that 28.6% of people with resectable hepatocellular carcinoma would improperly not be resected, while 8% of people without hepatocellular carcinoma would undergo inappropriate surgery. The uncertainty resulting from the high risk of bias in the included studies and concerns regarding their applicability limit our ability to confidently draw conclusions based on our results.

PLAIN LANGUAGE SUMMARY

How accurate are computerised tomography (CT) scans for detecting liver cancer?

Key messages

In people with chronic liver disease,

- \cdot computerised tomography (CT: cross-sectional scans inside the body) probably misses liver cancer in 22.5% of people who would not receive timely or appropriate treatment, and also, CT incorrectly finds liver cancer in 8.7% of people who would receive unnecessary treatment.
- ·CT probably misses liver cancer in 28.6% of people with liver cancer who could have surgery to remove part of their liver, and CT incorrectly finds liver cancer in 7.7% of people who undergo inappropriate surgery.
- · The studies were too different from each other to allow us to draw firm conclusions based on the evidence.

Why is it important to diagnose liver cancer accurately?

Liver cancer, or 'hepatocellular carcinoma' occurs mostly in people with chronic liver disease, regardless of the cause. It is the sixth most common cancer in the world and the fourth most common cause of death due to cancer. It is difficult to diagnose because early symptoms are similar to those of liver disease. People with blood test or ultrasound results that suggest liver cancer may go on to have further tests, such as scans that produce images of the liver, or biopsy where a small piece of the liver is removed and examined. If liver cancer is detected



early, people may be treated with surgery to remove part of the liver (a liver resection) or with a liver transplant. If the liver cancer is more advanced, people may need chemotherapy. If liver cancer is missed, people will not receive appropriate treatment. However, incorrectly diagnosing liver cancer when it is not present means that people may undergo unnecessary testing or treatment.

What is computed tomography and how might it diagnose liver cancer?

Computed tomography (CT) produces images that show a cross-section or 'slice' of the bones, blood vessels and tissues inside the body. The images consist of a series of X-rays that are directed and combined by a computer. CT scans can detect the presence of abnormalities in the liver that might be cancer. Current guidelines recommend using either CT or another type of imaging, magnetic resonance imaging (MRI), to confirm the presence of liver cancer in people who might have liver cancer, and to judge the size and spread (stage) of the cancer.

What did we want to find out?

We wanted to find out if CT is accurate enough to diagnose liver cancer in adults with chronic liver disease. We were interested firstly, in liver cancers of any size and stage and secondly, in liver cancers that were suitable for resection.

What did we do?

We searched for studies that assessed the accuracy of CT scans compared to the best available tests to confirm liver cancer in adults with chronic liver disease. The best available tests are examination of the liver, or part of the liver under a microscope.

What did we find?

We found a total of 21 studies with 3101 people.

Based on the studies, around 520 (52%) out of 1000 adults with chronic liver disease have confirmed liver cancer. Of these 1000 people, CT may:

- · correctly detect liver cancer in 403 people
- · miss liver cancer in 117 people
- · incorrectly detect liver cancer in 42 cancer-free people
- · correctly detect no liver cancer in 438 people.

Based on the studies, around 350 (35%) out of 1000 adults with chronic liver disease have confirmed resectable liver cancer. Of these 1000 people, CT may:

- · correctly detect resectable liver cancer in 250 people
- · miss resectable liver cancer in 100 people
- · incorrectly detect resectable liver cancer in 50 people; and
- · correctly detect no resectable liver cancer in 600 people.

What are the limitations of the evidence?

Our confidence in the evidence is limited because the studies used different methods to select study participants and used different definitions for the presence of liver disease. This means CT scans could be more or less accurate than suggested by the evidence.

How up to date is this evidence?

The evidence is up to date to 4 May 2021.



Summary of findings 1. Diagnostic accuracy of computed tomography for the diagnosis of hepatocellular carcinoma

Review question: what is the diagnostic accuracy of CT for the diagnosis of HCC in people with chronic liver disease?

Population: adults with chronic liver disease

Setting: clinical setting (secondary or tertiary care setting) or surveillance programmes

Study design: prospective and retrospective cross-sectional studies

Index test: CT

Target condition: HCC of any size, any stage

Reference standards

- Pathology of the explanted liver in case of transplantation
- Histology of resected focal liver lesion(s), or the histology of biopsied focal liver lesion(s) with a follow-up period of at least 6 months to exclude the presence of focal lesions not detected by the index test

Limitations in the evidence: risk of bias and applicability concerns

- Participant selection: high/unclear risk of bias 13 studies (62%); high concern for applicability 18 studies (86%)
- Index tests: high/unclear risk of bias 3 studies (14%); high concern for applicability 0 studies (0%)
- Reference standard: high/unclear risk of bias 12 studies (57%); high concern for applicability 14 studies (67%)
- Flow and timing: high/unclear risk of bias 18 studies (86%)

Findings

Implications in a hypothetical cohort of 1000 people										
Index test	Number of studies (partici- pants)	Sensitivi- ty (95% CI)	Specifici- ty (95% CI)	Preva- lence ^a %	True positives will receive appropriate treatment (surgery or local ablative therapy or systemic chemotherapy)	False negatives will be misdiag- nosed and not re- ceive appropriate treatment	True negatives will not undergo in- appropriate treat- ment or unneces- sary further testing	False pos- itives will undergo in- appropriate treatment	Certainty of the evi- dence	
СТ	21	77.5% (70.9% to	91.3% (86.5% to	20	155	45	730	70	Low ^b	
	(3101)	82.9%)	94.5%)	52	403	117	438	42	_	

35

365

465 135

CI: confidence interval; CT: computed tomography; HCC: hepatocellular carcinoma

^aWe chose for exemplification three values of hepatocellular carcinoma prevalence: 20% for a population with low clinical suspicion, 52% as a median derived from our study analysis, and 60% for population with high clinical suspicion (assessment of nodules detected by ultrasound).

bDowngraded by two levels: for risk of bias, and indirectness.

Summary of findings 2. Diagnostic accuracy of computed tomography for the diagnosis of resectable hepatocellular carcinoma

Review question: what is the diagnostic accuracy of CT for the diagnosis of resectable HCC in people with chronic liver disease?

60

Population: adults with chronic liver disease

Setting: clinical setting (secondary or tertiary care setting) or surveillance programmes

Study design: cross-sectional studies

Index test: computed tomography

Target condition: resectable HCC

Reference standards

- Pathology of the explanted liver in case of transplantation
- Histology of resected focal liver lesion(s), or the histology of resected or biopsied focal liver lesion(s) with a follow-up period of at least 6 months to exclude the presence of focal lesions not detected by the index test

Limitations in the evidence: risk of bias and applicability concerns (total 12 studies which had > 90% of participants with resectable HCC)

- Participant selection: high/unclear risk of bias 5 studies (42%); high concern for applicability 11 studies (92%)
- Index tests: high/unclear risk of bias 3 studies (25%); high concern for applicability 0 studies (0%)
- Reference standard: high/unclear risk of bias 6 studies (50%); high concern for applicability 10 studies (83%)
- Flow and timing: high/unclear risk of bias 11 studies (92%)

Findings

Implications in a hypothetical cohort of 1000 people										
Index test	Number of studies	Sensitiv- ity (95% CI)	Specifici- ty (95% CI)	Preva- lence ^a %	True positives will receive ap- propriate treat-	False negatives will be misdiag- nosed and not	True negatives will not undergo inappro-	False positives will undergo inappro- priate further test-	Certainty of the evi- dence	

	(partici- pants)				ment (surgical resection)	undergo surgical resection	priate further testing or surgical resection	ing or surgical re- section	
СТ	10	71.4% (60.3% to	92.0% (86.3% to	20	143	57	740	60	Low ^b
	(1854)	80.4%)	95.5%)	35	250	100	600	50	
				60	434	166	370	30	

CI: confidence interval; CT: computed tomography; HCC: hepatocellular carcinoma

^aWe chose for exemplification three values of hepatocellular carcinoma prevalence: 20% for a population with low clinical suspicion, 35% as a median derived from our study analysis, and 60% for population with high clinical suspicion (assessment of nodules detected by ultrasound).

^bDowngraded by two levels: for risk of bias, and indirectness.



BACKGROUND

Hepatocellular carcinoma is the most common primary liver neoplasm, usually developing in the setting of a chronic liver disease. It is the sixth most commonly diagnosed cancer type and the fourth leading cause of death from cancer worldwide; 782,000 deaths due to hepatocellular carcinoma were reported in 2018 (Bray 2018). Exceedingly high rates are present in East and Southeast Asia, several areas of Africa and Southern Europe (Bertuccio 2017). In the last decade, hepatocellular carcinoma was one of the few cancers that showed increasing incidence and mortality trends in several areas of the world including Europe, and North and Latin America (Bosetti 2013; Hashim 2016; Ryerson 2016). Mortality rates, even with a recently downward reported trend, are reported to be still two to five times higher in Japan, Hong Kong, and Korea than in most European countries and the Americas (Bertuccio 2017). Most common risk factors include liver cirrhosis, severe liver fibrosis, chronic infections with hepatitis B and C, heavy alcohol intake, tobacco use, diabetes, metabolic syndrome, aflatoxins (poisonous carcinogens produced by Aspergillus flavus and Aspergillus parasiticus, which grow in soil, decaying vegetation, hay, and grains), nonalcoholic fatty liver disease, and being overweight (Yang 2011; Bosetti 2014; Stanaway 2016; Bertuccio 2017). However, people who have developed hepatocellular carcinoma without known risk factors have been reported (Bralet 2000; Young 2012). Hepatocellular carcinoma is rare among adolescents with an incidence of 0.3 to 0.45 people per million per year and accounts for less than 1% of all malignant neoplasms among children younger than 20 years (Mann 1990). The reported hepatocellular carcinomas were associated with hepatitis B virus infection or with inherited metabolic disorders, specifically hereditary tyrosinaemia, a-1-antitrypsin deficiency, and glycogen storage disease type 1. Only approximately 30% of paediatric hepatocellular carcinomas are associated with cirrhosis, and the carcinogenesis and the clinical course are considered peculiar (Ni 2004; Omata 2017; Mogul 2018).

Clinically, hepatocellular carcinoma is frequently diagnosed in the late stages of liver disease because of the absence of specific symptoms, other than those related to chronic liver disease. Less than 20% of people are eligible for curative treatment - such as liver resection, transplantation or ablation - due to advanced tumour stage, liver dysfunction or shortage of liver donors (Davila 2012). Furthermore, curative treatment options are unfeasible in most instances due to severe clinical deterioration at the moment of diagnosis, or due to the inaccuracy of the preoperative clinical evaluation and staging procedure.

Despite the poor initial prognosis (the mortality-to-incidence overall ratio has been reported as 0.95; Ferlay 2019), a five-year survival rate of more than 50% can be achieved if the hepatocellular carcinoma is detected at an early stage (Forner 2012). According to the Barcelona Clinic Liver Cancer (BCLC) staging system, only people with early-stage hepatocellular carcinoma are eligible for curative treatment (Llovet 1999). Therefore, accurate and early diagnosis of hepatocellular carcinoma is of high importance.

Prior to advancements in medical imaging, biopsy and cytologic examination of the liver specimen were used to make a definitive diagnosis of hepatocellular carcinoma (Tao 1984). With the development of advanced imaging techniques, hepatocellular carcinoma has become unique among tumours in that its

characteristics can be accurately detected using imaging, thus reducing the need for invasive

biopsy (Forner 2008; Sangiovanni 2010; Manini 2014). Currently, biopsy is not preferred for the diagnosis of hepatocellular carcinoma due to concerns regarding tumour seeding, bleeding, and rate of false-negative results (Silva 2008; Pomfret 2010). However, it is reserved for lesions with atypical appearance and when imaging results are equivocal (Bruix 2011).

Computer tomography (CT) and contrast-enhanced magnetic resonance imaging (MRI) have been established as the non-invasive imaging modalities for detection and evaluation of liver lesions (Lee 2012a; O'Neill 2015). In comparison with single-detector CT, multidetector computed tomography (MDCT) is superior due to greater speed, thinner slices, and multiphasic scanning; these factors improve spatial and temporal resolution and provide more precise evaluation of liver tumour haemodynamics, and consequently, diagnostic accuracy (O'Neill 2015). The ability of CT to detect hepatocellular carcinoma rests on characterising the enhancement patterns in arterial, portal venous, and subsequent phases relative to the surrounding liver tissue. The differences in blood flow and extracellular volume between hepatocellular carcinoma and normal liver tissue lead to the main radiological hallmarks of hepatocellular carcinoma (Hennedige 2012; Choi 2014; Shah 2014; LI-RADS 2018).

According to the American Association for the Study of Liver Disease (AASLD) and European Association for the Study of the Liver (EASL) guidelines, a single contrast-enhanced imaging study (CT or MRI), performed in high-volume centres with up-to-date radiological equipment showing typical radiological hallmarks in people with cirrhosis, is valid to diagnose hepatocellular carcinoma (Bruix 2011; EASL-EORTC 2012; EASL 2018). However, if a detected lesion presents with some (but not all) of the hallmarks of hepatocellular carcinoma, another imaging study or biopsy is warranted.

According to current relevant guidelines, there are some differences in recommendations for management with regards to the size of a suspected focal liver lesion. In the AASLD guideline, lesions with a diameter less than 1 cm and those with a diameter more than 1 cm without hepatocellular carcinoma hallmarks are labelled as indeterminate lesions and require follow-up (Heimbach 2018). The EASL guideline proposes a diagnostic algorithm for management of suspected focal liver lesions and group lesions in two categories (with a diameter less than 1 cm, and more than 1 cm; EASL 2018). The Asian Pacific Association for the Study of the Liver (APASL) diagnostic pathways focus more on lesion characteristics than on size (Omata 2017).

Previous systematic reviews have assessed the performance of CT in detecting hepatocellular carcinoma, and they have included different studies and yielded different results (Colli 2006; Xie 2011; Chen 2013; Floriani 2013; Chou 2015; Lee 2015; Ye 2015; Guo 2016; Hanna 2016; Roberts 2018; Li 2019). These reviews are comparative reviews that compare two or more tests (ultrasound, CT, MRI) and include studies conducted before 2016, when CT diagnostic criteria were not clearly defined (LI-RADS). Evaluation of risk of bias and definition of inclusion criteria, type of studies, and reference standards are often inconsistent and questionable. Furthermore, these reviews did not put the index tests into context and did not define clearly their role, instead comparing all the available tests as they were used simultaneously. The aim of the present systematic



review and meta-analysis is to determine the accuracy of CT for the diagnosis of hepatocellular carcinoma of any size, as well as the diagnosis of resectable hepatocellular carcinoma in people with chronic liver disease.

Target condition being diagnosed

Hepatocellular carcinoma is the most common primary liver cancer that occurs in people with chronic liver disease. The incidence of hepatocellular carcinoma increases in individuals with chronic hepatitis B and C, alcohol use, and nonalcoholic fatty liver disease, and in those with liver cirrhosis of various aetiologies (Bruix 2011). There is no definite threshold in the definition of lesion size, although literature tends to classify lesions with a diameter equal to or less than 2 cm as 'small' (Hussain 2002; Choi 2014; Park 2017). The histological diagnosis of hepatocellular carcinoma poses many challenges, particularly when dealing with liver biopsy specimens, due to the heterogeneity of hepatocellular carcinoma and occasional difficulties confirming hepatocellular differentiation. Primary liver tumours should be considered as a continuum with typical hepatocellular and cholangiocarcinoma as the two ends of the spectrum. In between, a whole range of tumours showing both hepatocellular and cholangiocellular differentiation with or without an associated progenitor/stem cell component should be differentiated. Characterisation of combined (or mixed) hepatocellular-cholangiocarcinoma can be very challenging. In advanced-stage chronic liver disease, the main challenge for the histopathologist is still to differentiate between hepatocellular carcinoma and its precursors, large regenerative nodule, and a dysplastic nodule, with the potential to progress to hepatocellular carcinoma. The transition from dysplastic nodule to hepatocellular carcinoma is thought to be associated with a change in the lesional vascular supply, from a dual portal-arterial to a predominantly arterial, due to neoangiogenesis (Quaglia 2018). The radiological counterpart of these changes is contrast uptake in the arterial phase and rapid washout in the venous phase, which is considered to be sufficient for a diagnosis of hepatocellular carcinoma (Omata 2017; EASL 2018; Heimbach 2018). An international consensus defined the diagnostic criteria and highlighted the difficulties in histological differentiation between the different stages of hepatocellular carcinoma progression (International Consensus Group for HCN 2009).

In clinical practice, and according to pertinent guidelines, multiphasic CT or MRI with intravascular contrast allow for a highly accurate diagnosis of hepatocellular carcinoma without an invasive liver biopsy. The diagnosis of hepatocellular carcinoma is usually obtained on the basis of cross-sectional CT or MRI features, and liver histology is required only for undefined lesions (Omata 2017; EASL 2018; Heimbach 2018; LI-RADS 2018).

A number of staging systems for hepatocellular carcinoma have been proposed and developed, however, there is no globally applicable staging system (Kinoshita 2015). Among different staging protocols, the BCLC staging system has a notable feature of treatment recommendations for each stage, based on the best treatment options currently available (Llovet 1999; Llovet 2003; Llovet 2008). It is comprised of four elements: tumour extension, liver functional reserve, physical status, and cancerrelated symptoms. According to the BCLC, only people with early-stage hepatocellular carcinoma are eligible for curative treatment such as surgical resection or percutaneous locoregional treatment. Orthotopic liver transplantation is reserved for people

with decompensated cirrhosis, and it is considered a definite curative treatment for hepatocellular carcinoma. When orthotopic liver transplantation for hepatocellular carcinoma was initially introduced in the 1980s, it was associated with poor five-year survival and high recurrences, which led to the treatment being contraindicated for hepatocellular carcinoma (Yokoyama 1990). In 1996, specific criteria, known as the Milan criteria (Mazzaferro 1996), were developed for the selection of people for liver transplantation. These criteria have been repeatedly validated and their value is considerable (EASL 2018). With their implementation, overall five-year survival of people with post-orthotopic liver transplantation exceeded 70% (Mazzaferro 2011). The criteria for people eligible for orthotopic liver transplantation include a single hepatocellular carcinoma lesion with a diameter equal to or less than 5 cm, or up to three hepatocellular carcinoma lesions, each with a diameter equal to or less than 3 cm; no vascular invasion; and no extrahepatic involvement (no metastasis; Mazzaferro 1996).

Index test(s)

Contrast-enhanced multidetector and multiphasic CT is an advanced imaging modality that includes rapid intravenous injection of contrast agent with fast data acquisition using ionising radiation. Minimal CT requirements for the detection of hepatocellular carcinoma include performance on multidetector CT with 8 or more detector rows, acquisition of images in arterial, portal venous, and delayed phase with multiplanar reformations. If people have undergone prior locoregional hepatocellular carcinoma treatment, acquisition of precontrast images is required (LI-RADS 2018).

Although uncommon, physicians should be aware of the acute adverse reactions to iodine contrast which are categorised into mild (nausea, mild vomiting, urticaria, and itching), moderate (severe vomiting, marked urticaria, bronchospasm, facial/laryngeal oedema, and vasovagal attack), and severe (hypotensive shock, respiratory arrest, cardiac arrest, and convulsion). Also, the administration of iodinated contrast agent may lead to contrast-induced nephropathy. However, this entity is more uncommon than the aforementioned adverse reactions (Thomsen 2014). Ionising radiation produced by CT scanners is, by definition, harmful to the molecular structure of human tissue. However, many technological improvements, dose reduction strategies, and radiation effect campaigns have been made for the benefit of reducing radiation risks in people undergoing a CT exam (Kaira 2015; Parakh 2016).

The American College of Radiology established the Liver Reporting and Data System (LI-RADS), with the aim of standardising the terminology, interpretation, and reporting of imaging findings in people with suspected hepatocellular carcinoma. Several versions have been published since the initial release in 2008, most recently in 2018 (LI-RADS 2018). The LI-RADS assign a diagnostic category to each focal liver lesion/observation based on major, ancillary, and other imaging features. Major features include non-rim-like hyperenhancement in arterial phase, non-peripheral washout in portal venous and subsequent phases, enhancing capsule, lesion diameter, and threshold growth (LI-RADS 2018). Based on the presence of major features and morphological suspicion of hepatocellular carcinoma, each lesion is assigned with a category ranging from LR-1 (definitely benign) to LR-5 (definitely hepatocellular carcinoma). Other categories include suspicion for malignancy, but not necessarily hepatocellular carcinoma



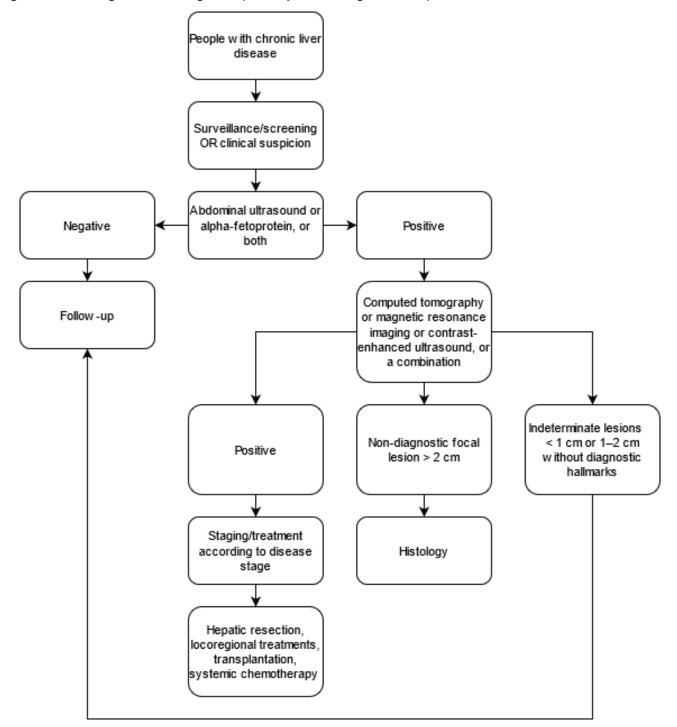
(LRM) and tumour in vein (LR-TIV). If assigning a category is doubtful, many ancillary features have been defined to favour the presence of hepatocellular carcinoma, malignancy other than hepatocellular carcinoma, or benign lesion. Features favouring hepatocellular carcinoma include non-enhancing capsule, nodule in nodule appearance, mosaic architecture, blood products, and fat in the lesion. The main aim of this categorisation is to clearly define the probability that a certain lesion is indeed a hepatocellular carcinoma, and to help guide multidisciplinary clinical management (LI-RADS 2018; Van der Pol 2019).

Clinical pathway

Surveillance of hepatocellular carcinoma (screening performed at regular intervals) in the at-risk population, that is, people with chronic liver disease, regardless of aetiology, is carried out by abdominal ultrasound for detection of nodules. Once a suspected nodule has been detected, other imaging methods are considered according to the size of the nodule and appropriate guidelines. For a flow diagram of the clinical pathway and placement of tests, see Figure 1.



Figure 1. Flow diagram of the diagnostic pathway for the diagnosis of hepatocellular carcinoma



American Association for the Study of Liver Disease (AASLD) diagnostic guidelines

According to the AASLD guidelines, adults with cirrhosis and suspected hepatocellular carcinoma should undergo diagnostic evaluation with either multiphasic CT or multiphasic MRI. Lesions that do not meet the positivity criteria (i.e. arterial phase hyperenhancement in combination with washout appearance and/or capsule appearance), or whose size is less than 1 cm,

are considered indeterminate. For indeterminate lesions, several options are suggested including follow-up imaging, imaging with an alternative imaging modality or alternative contrast agent, or biopsy. No option is preferred and recommended over another. Biopsy may be required in selected instances, but its routine use is not advocated (Bruix 2011; Heimbach 2018).



European Association for the Study of the Liver (EASL) diagnostic guidelines

In cirrhotic liver disease, the diagnostic algorithm proposed by the EASL divides suspected focal liver lesions into two categories: lesions smaller than 1 cm, and those larger than 1 cm. Lesions larger than 1 cm need to be evaluated by CT or MRI straight away. If at least one of these imaging modalities is positive, i.e. proves the existence of hepatocellular carcinoma hallmarks, diagnosis of hepatocellular carcinoma is considered certain. If the results are equivocal, the use of other multiphasic imaging modality is required: multiphasic contrast-enhanced CT or multiphasic contrast-enhanced MRI, gadoxetic-enhanced MRI, or contrast-enhanced ultrasound. If these studies prove the hallmarks of hepatocellular carcinoma, the diagnosis is certain; otherwise, biopsy is warranted. If biopsy appears to be unclear, re-biopsy is to be considered or a repeat ultrasound follow-up every four months is needed. Lesions smaller than 1 cm are to be followed up by ultrasound every four months: if the size of the lesion does not increase, then further ultrasound follow-up is recommended; otherwise, multiphasic contrast-enhanced CT, multiphasic contrast-enhanced MRI, or gadoxetic-enhanced MRI is required (EASL 2018).

Asian Pacific Association for the Study of the Liver (APASL) diagnostic guidelines

Under the APASL guidelines, a single dynamic contrast-enhanced MRI or CT is warranted regardless of the size of suspected liver nodule. If typical hallmarks of hepatocellular carcinoma are shown (presence of arterial hyperenhancement, followed by washout in the portal venous or delayed phases, or both), diagnosis is confirmed. If the lesion is hypervascular but shows no washout, another contrast-enhanced MRI study is needed. If the lesion proves to be hypointense, hepatocellular carcinoma diagnosis is confirmed. However, if the lesion is isointense or hyperintense, biopsy is warranted. If the lesion on the first dynamic MRI or CT study is non-hypervascular, a dynamic MRI study in hepatobiliary phase is needed. If the lesion is isointense or hyperintense, surveillance by ultrasound is recommended every six months, and if the lesion is hypointense, contrast-enhanced ultrasound of the liver nodule is warranted. Depending on lesion features on contrastenhanced ultrasound, biopsy or another dynamic CT or MRI study is recommended every three to six months (Omata 2017).

The expected downstream consequences of the CT results are: people with true-positive results, that is, those with hepatocellular carcinoma and positive test results, will receive the appropriate treatment (surgery, local ablative therapy, or systemic chemotherapy); people with true-negative results, that is, those without hepatocellular carcinoma and negative test results, will not undergo inappropriate treatment or unnecessary further testing; people with false-negative results, that is, those with hepatocellular carcinoma and negative test results, will be misdiagnosed, not receive the appropriate treatment and might be detected later as a more severe case; people with false-positive results, that is, those without hepatocellular carcinoma and positive test results, will undergo further testing and possibly inappropriate treatment. In people on a waiting list for orthotopic liver transplantation for an indication not related to an hepatocellular carcinoma, the consequences of false-negative results of preoperative CT are not completely known and might be less severe: indeed studies report no significant difference in terms of overall survival and tumour recurrence between people with and without previously diagnosed hepatocellular carcinomas (Castillo 2009; Senkerikova 2014; Madaleno 2015; El Moghazy 2016).

Prior test(s)

For surveillance purposes, abdominal ultrasound is recommended as a first-line imaging modality in people with chronic liver disease, regardless of aetiology, who are at risk of developing a hepatocellular carcinoma (Omata 2017; EASL 2018; Heimbach 2018). It is also used as a diagnostic tool in people with clinical suspicion of hepatocellular carcinoma for detecting liver lesions. Alpha-foetoprotein has been used as a diagnostic biomarker even before technological advancements (Kew 1975). However, its role as a screening tool is still a matter of debate. The diagnosis of chronic advanced liver disease is based on clinical judgement derived from history, laboratory testing, physical examination, imaging, liver stiffness measurement, liver histology, or a combination of the aforementioned. Due to the accuracy of non-invasive tests, liver histology is reserved for only a minority of people with unclear diagnosis, and a non-invasive diagnosis of chronic advanced liver disease is considered equivalent to a histological diagnosis of cirrhosis (de Franchis 2015).

Role of index test(s)

Computer tomography is used as an add-on test after ultrasound detection of liver lesions suspected for hepatocellular carcinoma in surveillance programmes or hospital settings in people with clinical suspicion. Based on CT findings, biopsy and other imaging modalities could be avoided, therefore further testing could be reserved for a minority of patients.

Alternative test(s)

An alternative imaging modality in detecting hepatocellular carcinoma is contrast-enhanced dynamic MRI with extracellular and cell-specific gadolinium-based contrast agents. A recent meta-analysis aimed to determine the diagnostic benefit between multiphasic contrast-enhanced CT, extracellular contrast-enhanced MRI, and cell-specific gadoxetate-enhanced MRI for detection of hepatocellular carcinoma in people with cirrhosis (Roberts 2018). No definitive recommendation could be made for the systematic use of gadolinium-enhanced MRI over CT, although other previous meta-analyses reported a preference for MRI (Lee 2015; Ye 2015; Guo 2016).

Contrast-enhanced ultrasound is an advanced form of ultrasound examination in which images are acquired using intravenously injected microbubble contrast agent. Dynamic contrast-enhanced ultrasound images are obtained similarly to contrast-enhanced CT and MRI studies: depending on the time of image acquisition after intravenous contrast injection, the study differentiates arterial and portal venous phases in which sonographic hallmarks for hepatocellular carcinoma, such as arterial hyperenhancement and subsequent washout appearance, are investigated (Chung 2015; LI-RADS). Unlike CT and MRI contrasts, ultrasound contrast agent is a purely intravascular agent; therefore, it is highly accurate in detecting tumour angiogenesis (Schirner 2004).

Lipiodol computerised tomography (Lipiodol-CT) was used in the past as a diagnostic modality for the detection of hepatocellular carcinoma. The method included intra-arterial injection of iodised oil (Lipiodol) through the hepatic arterial supply, following which Lipiodol was deposited within the hepatocellular carcinoma



nodule. The hepatocellular carcinoma was visualised as a hyperattenuating nodule on the subsequent CT, and it showed high sensitivity in detecting small hepatocellular carcinoma (Takayasu 1990). In the context of transarterial chemoembolisation, Lipiodol may be used as an intraprocedural diagnostic modality (C-arm Lipiodol CT) for additional detection of small-size hepatocellular carcinoma (Li 2015).

Rationale

Hepatocellular carcinoma is currently detected by liver ultrasound in people with normal or high alpha-foetoprotein during surveillance programmes of people with chronic liver disease. Following ultrasound, the diagnosis is usually confirmed with high levels of alpha-foetoprotein and contrast-enhanced ultrasound, CT, or MRI. The latter two imaging modalities are also appropriate for staging and allow the choice of the most appropriate treatment. There is no clear evidence of the benefits of surveillance programmes in terms of overall survival: the conflicting results can be a consequence of inaccurate detection, ineffective treatment, or both. Assessing the diagnostic accuracy of CT, the most used confirmatory test after first-line tests, may clarify whether the absence of benefit in surveillance programmes might be related to underdiagnosis or understaging. Furthermore, an assessment of the accuracy of CT for the diagnosis of hepatocellular carcinoma is needed for either ruling out, diagnosing, or supporting further testing in people with chronic liver disease who are not included in surveillance programmes.

This review represents a part of a series of reviews about the diagnostic accuracy of the most commonly used modalities for diagnosing hepatocellular carcinoma in people with chronic liver disease. The first part includes assessment of the diagnostic accuracy of ultrasound and alpha-foetoprotein levels, which are used as triage tests in surveillance (Colli 2021). The second part focuses on the diagnostic accuracy of contrast-enhanced ultrasound in characterising suspected lesions as hepatocellular carcinoma as a second-line diagnostic modality (Fraquelli 2019). The present review focuses on the assessment of CT as a secondline imaging modality in assessing focal liver lesions detected on ultrasound suspected for hepatocellular carcinoma. A comparable review assessing the accuracy of MRI for diagnosing hepatocellular carcinoma is in progress (Nadarevic 2021). We are planning to produce an overview of the reviews that will assess abdominal ultrasound and alpha-foetoprotein, contrast-enhanced ultrasound, CT, and MRI for the diagnosis of hepatocellular carcinoma.

OBJECTIVES

To assess the diagnostic accuracy of multidetector, multiphasic contrast-enhanced CT for the diagnosis of hepatocellular carcinoma (hepatocellular carcinoma) of any size, and at any stage, in adults with chronic liver disease, either in a surveillance programme or in a clinical setting.

Secondary objectives

 To assess the diagnostic accuracy of multidetector, multiphasic contrast-enhanced CT for the diagnosis of resectable hepatocellular carcinoma in adults with chronic liver disease. The definition of resectable hepatocellular carcinoma is a neoplasm amenable to surgical radical resection according to the current guidelines (the Milan criteria): a single lesion with a maximum diameter of less than 5 cm, or fewer than three lesions with a maximum diameter of 3 cm (Mazzaferro 1996).

- To investigate the following sources of heterogeneity:
 - study date (studies published before the year 2005 compared to studies published after the year 2005, due to advancements in technology);
 - study date (studies published before 2016 compared to studies published after 2016, due to changes in diagnostic criteria);
 - inclusion of participants without cirrhosis (studies including more than 10% participants without cirrhosis compared to studies including less than 10% participants without cirrhosis);
 - study location (population differences): studies conducted in North and South America compared to Europe compared to Asia;
 - patient selection (patients recruited from planned surveillance programmes compared to clinical cohorts);
 - different hepatocellular carcinoma stage (studies in which 20% or more of participants have resectable hepatocellular carcinoma compared to studies in which less than 20% of participants have resectable hepatocellular carcinoma);
 - different reference standard (histology of the explanted liver compared to liver biopsy compared to another reference standard);
 - different liver cirrhosis aetiology (hepatitis C or hepatitis B virus-associated cirrhosis compared to all other aetiologies);
 - number of CT detector rows (exams conducted on 64slice or fewer compared with more than 64-slice, due to advancements in technology);
 - o hepatocellular carcinoma mean diameter;
 - prevalence of the target condition (above median compared to below median);
 - prior detection of nodules, studies including study participants with prior tests to detect nodules compared to studies including study participants without prior tests.

We chose the variables listed above for the following reasons. Due to advancements in technology and change in diagnostic criteria, we considered the date of study publication. The proportion of participants without cirrhosis is relevant because hepatocellular carcinoma in absence of cirrhosis has different CT characteristics, prognosis, and treatment. There are differences in epidemiology, and clinical and radiological characteristics of hepatocellular carcinoma in Asia and in Western countries. Selection of patients can induce variability of results: participants recruited from screening or surveillance programmes may be different mainly in severity of the underlying liver disease and consequently in radiological characteristics of the liver. The hepatocellular carcinoma prevalence in included studies can change according to selection and epidemiology. The proportion of resectable hepatocellular carcinoma found in the studies reflects different epidemiology and patient selection. The clinical and radiological characteristics of hepatocellular carcinoma varies according to the aetiology of the underlying liver disease, mainly in the case of chronic infection with hepatitis C or hepatitis B, compared to other aetiologies. The accuracy of CT may vary according to the diameter of the neoplastic lesion and the number of detector rows in the CT equipment. Prior testing and the inclusion of participants with nodules might produce differences in CT accuracy estimates



secondary to this different selection. The investigation of this last possible source of heterogeneity was not planned in the protocol and was added subsequently.

METHODS

Criteria for considering studies for this review

Types of studies

We included studies that, irrespective of publication status and language, have evaluated the diagnostic accuracy of multidetector, multiphasic contrast-enhanced CT for the diagnosis of hepatocellular carcinoma in adults with chronic liver disease. These studies should have used one of the acceptable reference standards (see Reference standards).

We considered studies of cross-sectional design that included participants with clinical suspicion of hepatocellular carcinoma. We excluded studies of case-control design that compared people with known hepatocellular carcinoma to matched control as these are considered to have high risk of bias due to inflated accuracy estimates (Colli 2014). We excluded studies that analysed data only per lesion, rather than per participant, unless study authors made participant data available.

Participants

We included participants aged 18 years and older, of any sex, at risk of developing hepatocellular carcinoma, and with chronic liver disease, irrespective of aetiology, severity of disease, and duration of illness, with or without prior tests, ultrasound, and alphafoetoprotein. The review focused on diagnostic questions related to adults with a first diagnosis of hepatocellular carcinoma. People with previous diagnosis and treatment of hepatocellular carcinoma make up a distinct group for which the diagnosis or natural history of hepatocellular carcinoma has been modified. These people were not the focus of this review; therefore, we excluded studies that included such participants unless they represented less than 5% of all the included participants, or if study authors had presented data in such a way as to allow this group of participants to be isolated from the remaining included participants.

Index tests

We included multiphasic contrast-enhanced CT for the detection of hepatocellular carcinoma in adults with chronic liver disease. Regarding positivity criteria, we accepted any definition of positive/negative test results. This judgment usually, even if implicitly, considers the presence of suspected liver lesion, which shows non-rim-like arterial hyperenhancement and subsequent non-peripheral washout appearance in later phases.

Target conditions

- · Hepatocellular carcinoma of any size and at any stage
- Resectable hepatocellular carcinoma (see Secondary objectives)

Reference standards

We accepted one of the following as a reference standard for the diagnosis of hepatocellular carcinoma.

• The pathology of the explanted liver in case of transplantation

The histology of resected focal liver lesion(s), or the histology
of biopsied focal liver lesion(s) with a follow-up period of at
least three months to exclude the presence of focal lesions not
detected by the index test.

These reference standards, even if commonly used in clinical practice, are not perfect. The pathology of the explanted liver is possible only when all the included participants have undergone liver transplantation; therefore, the setting does not represent the whole spectrum of liver disease severity as only people with advanced and decompensated liver disease are candidates for orthotopic liver transplantation. In the case of histology of resected focal lesion and histology of biopsied liver lesions, the negative result can be confirmed only with an adequate follow-up period. This would introduce an unavoidable differential verification bias.

Search methods for identification of studies

Electronic searches

We searched the Cochrane Hepato-Biliary Group (CHBG) Controlled Trials Register and the CHBG Diagnostic Test of Accuracy Studies Register (both registers are maintained and searched internally by the CHBG Information Specialist via the Cochrane Register of Studies Web), The Cochrane Library, MEDLINE Ovid, Embase Ovid, LILACS (Bireme), Science Citation Index - Expanded (Web of Science), and Conference Proceedings Citation Index - Science (Web of Science) until 04 May 2021. Appendix 1 gives the search strategies with the time spans of the searches.

We did not apply any restrictions on language or document type.

Searching other resources

We tried to identify additional references by manually searching articles retrieved from digital databases and relevant review articles. We sought information on unpublished studies by contacting experts in the field. In addition, we handsearched abstract books from meetings of the AASLD, the EASL, and APASL held during the past 10 years. We also searched for other kinds of grey literature in the System for Information on Grey Literature in Europe 'OpenGrey' (www.opengrey.eu/).

Data collection and analysis

We followed available guidelines as provided in the *Cochrane Handbook for Systematic Reviews of Diagnostic Test Accuracy* (DTA Handbook 2013).

Selection of studies

Two review authors (VG and TN) independently scrutinised titles and abstracts identified by electronic literature searching to identify potentially eligible studies. We selected any citation, identified by either of the two review authors, as potentially eligible for full-text review. The same review authors independently assessed full-text papers for study eligibility, using predefined inclusion and exclusion criteria. We resolved any discrepancies by discussion. After full-text assessment, we recorded all studies and their reasons for exclusion, in the 'Characteristics of excluded studies' table and illustrated the study selection process using a PRISMA diagram (Salameh 2020; Page 2021).



Data extraction and management

We developed a standardised data extraction form and piloted the form on five of the included studies. Based on the pilot, we finalised the form. Then, two review authors (VG and TN) independently completed the data extraction form for each included study. Each review author independently retrieved study data. In cases of disagreement, we reached consensus through discussion with a third review author (AC).

We extracted the following data.

- General information: title, journal, year, publication type, study design and data collection (prospective versus retrospective), surveillance programme or clinical cohorts
- Sample size: number of participants meeting the criteria and total number of participants included and tested
- Baseline characteristics: baseline diagnosis, age, sex, race, and presence of cirrhosis and mean diameter of hepatocellular carcinoma
- · Index test with predefined positivity criteria
- · Target condition
- Time interval between the index test and the reference standard
- · Reference standard tests
- Numbers of true-positive, true-negative, false-positive, and false-negative findings. We extracted these data for the two target conditions (hepatocellular carcinoma of any size and stage and resectable hepatocellular carcinoma)
- Number of uninterpretable results

The unit of analysis was the study participant, and we extracted data per participant. We summarised the data from each study in 2x2 tables (true positive, false positive, false negative, true negative), and we entered the data into Review Manager 5 software (Review Manager 2020).

Missing data

We contacted primary authors of nine primary studies by email to ask for additional information regarding per-patient analyses and data needed to design the 2x2 tables. Two study authors responded but did not provide any additional data. We did not receive a reply from any other study authors. After two weeks we sent a second email but still did not receive a reply. We eventually excluded all the studies in question.

Assessment of methodological quality

Two review authors (VG and TN) independently assessed the risk of bias of included studies and applicability of their results using QUADAS-2 (revised tool for quality assessment of diagnostic accuracy studies; Whiting 2011). In cases of disagreement, we reached a consensus through discussion. We addressed aspects of study quality involving the participant spectrum, index tests, target conditions, reference standards, and flow and timing. Regarding the index test positivity criteria definition, we assessed whether studies reported a clear definition. We recognise that, even if positivity criteria do not present explicit thresholds, they are nevertheless vulnerable to implicit thresholds. We defined a time interval between the index test and the reference standard of three months as appropriate. According to a recent systematic review, the approximate hepatocellular carcinoma volume doubling time is 4 months to 5 months with significant range of 2.2 moths

to 11.3 months (Nathani 2021). In accordance with suggestions from a previous systematic review, which noted the acceptable time interval being from 1 month to 3 months (Kim 2008), we assumed 90 days to be the most acceptable threshold. The visualisation of the liver can sometimes be suboptimal due to patient characteristics; therefore, lack of reporting or exclusion of uninterpretable results from analyses could overestimate the accuracy of CT. We considered the study to be at high risk of bias if uninterpretable results were excluded from the analysis.

We classified a study at a high risk of bias if we judged at least one of the QUADAS-2 study domains as high risk (Appendix 2).

Statistical analysis and data synthesis

We provided a description of the included studies by calculating median values and interquartile ranges (IQR) across studies for some characteristics of our interest, defined at study level. In particular, we considered hepatocellular carcinoma mean diameter and the prevalence of participants with the following characteristics: hepatocellular carcinoma, resectable hepatocellular carcinoma, liver cirrhosis, and viral aetiology of cirrhosis.

We designed 2x2 tables for each primary study for the index test (see Data extraction and management). We planned the following strategy of analyses.

Firstly, we performed a graphical descriptive analysis of the included studies and presented forest plots (sensitivity and specificity separately, with their 95% confidence intervals (CIs)). Secondly, we performed a meta-analysis using the bivariate model and provided estimates of summary sensitivity and specificity (Macaskill 2010). We used the pooled estimates obtained from the fitted models to calculate summary estimates of positive and negative likelihood ratios (LR+ and LR-, respectively).

In case of uninterpretable results, we planned to analyse data according to the intention-to-diagnose principle (Schuetz 2012), also described as worst-case scenario in Cohen 2015. Participants with uninterpretable index test results were classified as false positive if they had a negative reference standard or a false negative for participants with a positive reference standard. If data for the intention-to-diagnose analyses were not retrievable from the text, we contacted publication authors with provided email addresses. If we received no reply, we included the study in the analyses with data retrievable from the published manuscript and we considered it as having a high risk of bias. However, no study reported uninterpretable index test results.

We performed all statistical analyses using SAS statistical software (SAS), and macro METADAS (DTA Handbook 2013).

Investigations of heterogeneity

We investigated the effects of the following sources of heterogeneity.

- Study date (studies published before the year 2005 compared to studies published after the year 2005, due to advancements in technology (categorical)
- Study date (studies published before 2016 compared to studies published after 2016, due to changes in diagnostic criteria (categorical)



- Inclusion of participants without cirrhosis, studies including more than 10% participants without cirrhosis compared to studies including less than 10% participants without cirrhosis (categorical)
- Study location (population differences): studies conducted in North and South America compared to Europe compared to Asia (categorical)
- Participant selection, participants recruited from planned surveillance programmes compared to clinical cohorts (categorical)
- Different hepatocellular carcinoma stage, studies in which 20% or more of participants have resectable hepatocellular carcinoma compared to studies in which less than 20% of participants have resectable hepatocellular carcinoma (categorical)
- Different reference standard, histology of the explanted liver compared to liver biopsy compared to another reference standard)
- Different liver cirrhosis aetiology (hepatitis C or hepatitis B virus-associated cirrhosis compared to all other aetiologies (categorical)
- Number of CT detector rows, exams conducted on 64-slice or fewer compared with more than 64-slice, due to advancements in technology (categorical)
- Hepatocellular carcinoma mean diameter (continuous)
- Prevalence of the target condition, above median compared to below median (categorical)
- Prior detection of nodules, studies including participants with prior tests to detect nodules compared to studies including participants without prior tests (categorical)

We estimated the effects of the predefined sources of heterogeneity by adding covariates to the bivariate model. We assessed the statistical significance of the covariate effect by using the log-likelihood ratio test for comparison of models with and without the covariate term. We considered two-sided P values of less than 0.05 as statistically significant. For interpretation of the results of heterogeneity analysis, we considered the uncertainty of accuracy estimates in the different subgroups, quantified by 95% CIs of the estimated sensitivity and specificity, as an assessment of the degree to which these subgroups could influence diagnostic accuracy.

Sensitivity analyses

We assessed the effects of risk of bias of included studies on diagnostic accuracy by performing a sensitivity analysis in which we excluded studies classified as having high or unclear risk of bias in at least one of the QUADAS- 2 domains (Appendix 2). In addition, we defined the following signalling questions as most relevant, and conducted sensitivity analyses in which we excluded studies with answers of 'no' or 'unclear'.

- Were the positivity criteria defined?
- Were the reference standard results interpreted without the knowledge of the results of the index test?

We also conducted sensitivity analyses in which we excluded studies published only in abstract or letter form, and by limiting the analysis to studies we considered at low concern for applicability.

Assessment of reporting bias

In order to reduce reporting bias, we did not plan to use a filter search strategy nor to implement any language or sample. We did not plan to test for publication bias due to the lack of validated methods for diagnostic test accuracy reviews.

Summary of findings table and assessment of the certainty of evidence

We prepared summary of findings tables to present the main results and key information regarding the certainty of evidence. We assessed the certainty of evidence as recommended using the GRADE approach (Balshem 2011; Schünemann 2008; Schünemann 2016; GRADEpro GDT). We rated the certainty of evidence as either high (when not downgraded), moderate (when downgraded by one level), low (when downgraded by two levels), or very low (when downgraded by more than two levels) based on five domains: risk of bias, indirectness, inconsistency, imprecision, and publication bias. For each outcome, the certainty of evidence started as high when there were high-quality observational studies (cross-sectional or cohort studies) that enrolled participants with diagnostic uncertainty. If we found a reason for downgrading, we used our judgement to classify the reason as either serious (downgraded by one level) or very serious (downgraded by two levels) (Schünemann 2020a; Schünemann 2020b).

Five authors (TN, VG, MF, AC, GC) discussed judgements and applied GRADE in the following way.

- Risk of bias: we used QUADAS-2 to assess risk of bias.
- Indirectness: we assessed indirectness in relation to the population (including disease spectrum), setting, interventions, and outcomes (accuracy measures). We also used prevalence as a guide to whether there was indirectness in the population.
- Inconsistency: we carried out prespecified analyses to investigate potential sources of heterogeneity and downgraded when we could not explain inconsistency in the accuracy estimates based on whether the individual point estimates were similar and if the confidence intervals overlapped sufficiently in the forest plots.
- Imprecision: we looked at the confidence intervals of sensitivity and specificity estimates and at the unexplained heterogeneity of the results.
- Publication bias: we did not evaluate publication bias due to the lack of validated methods for diagnostic test accuracy reviews.

RESULTS

Results of the search

We ran the search on 4 May 2021. We identified 33,282 references by searching the following databases: the Cochrane Hepato-Biliary Group Controlled Trials Register (n = 350), the Cochrane Hepato Biliary Group Diagnostic Test of Accuracy Studies Register (n = 8), The Cochrane Library (n = 1255), MEDLINE Ovid (n = 5785), Embase Ovid (n = 19,833), LILACS (n = 102), and Science Citation Index – Expanded with Conference Proceedings Citation Index – Science (n = 5949). After exclusion of 8055 duplicates, 25,230 references remained for possible eligibility. We retrieved three additional references through handsearching. After reading the title and the abstract of these references, we excluded 25,065 of them, as they did not meet the inclusion criteria. We retrieved full texts of the

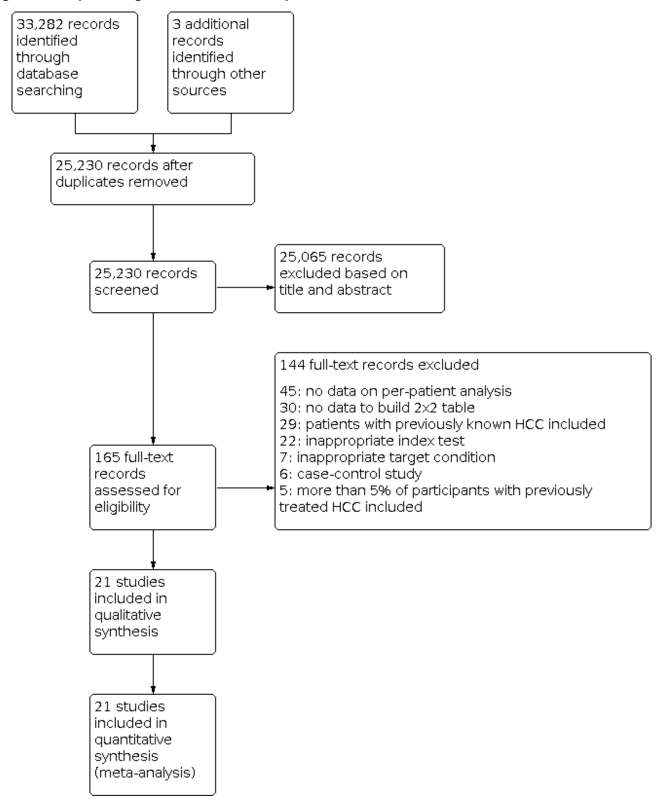


remaining 165 records, and after reading the full texts, we excluded 144 studies for various reasons (see Characteristics of excluded studies). Finally, we included in our review 21 references reporting data on 21 studies (Salameh 2020; Page 2021; Figure 2), including a total of 3101 participants (Pozzato 1997; Chalasani 1999; Gambarin-Gelwan 2000; Mortele 2001; de Ledinghen 2002; Libbrecht 2002; Lim 2002; Freeny 2003; Teefey 2003; Van Thiel 2004; Golfieri 2009; Sangiovanni 2010; Haberman 2011; Kim 2011; Yu 2011; Serste 2012; Maiwald 2014; Lin 2016; Villacastin Ruiz 2016; Hsiao 2019;

Langenbach 2019). Three additional studies, which were retrieved through handsearching, were all included in the analysis (Chalasani 1999; Van Thiel 2004; Maiwald 2014). We applied no language restrictions in the inclusion criteria, which resulted in retrieving full-text articles of 24 studies published in non-English languages of which we included two in the final analysis (Pozzato 1997; Haberman 2011). We requested further information by email for two studies, but did not receive a reply. The studies were conducted from 1997 to 2019.



Figure 2. Study flow diagram. Date of search 4 May 2021



We reported in the Characteristics of included studies tables the main characteristics of the 21 studies. All studies are reported as full-text publications.

Methodological quality of included studies

We have reported in detail results of the quality assessment of included studies in the Characteristics of included studies tables, and we have summarised this information in Figure 3 and Figure 4.



Figure 3. Risk of bias and applicability concerns graph: review authors' judgements about each domain presented as percentages across included studies

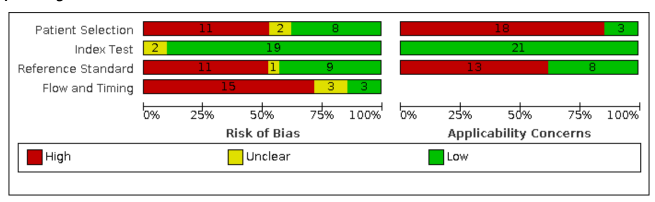
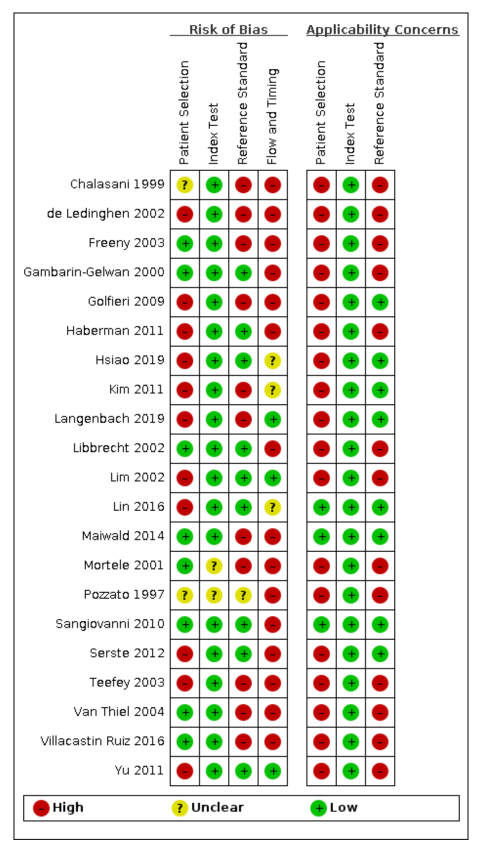




Figure 4. Risk of bias and applicability concerns summary: review authors' judgements about each domain for each included study





Patient selection

We included only studies with a cross-sectional design.

Risk of bias

Eight studies were at low risk of bias in this domain (Gambarin-Gelwan 2000; Mortele 2001; Libbrecht 2002; Freeny 2003; Van Thiel 2004; Sangiovanni 2010; Maiwald 2014; Villacastin Ruiz 2016). We judged two studies unclear for this domain, as they did not provide any data on the presence of exclusion criteria (Pozzato 1997; Chalasani 1999). Eleven studies were at high risk, due to exclusion criteria that we considered inappropriate: missing results of the index test, hepatocellular carcinoma diameter, time interval between index test and reference standards, inconclusive diagnosis, CT performed in institutions outside the study centre, no pathology fibrosis score analysis, absence of liver tumour at the pathology of the explanted liver, or participants removed from the transplant waiting list (de Ledinghen 2002; Lim 2002; Teefey 2003; Golfieri 2009; Haberman 2011; Kim 2011; Yu 2011; Serste 2012; Lin 2016; Hsiao 2019; Langenbach 2019).

Applicability

We judged three studies at low concern (Sangiovanni 2010; Maiwald 2014; Lin 2016). The other 18 studies we judged at high concern because they included only participants with end-stage liver disease on the waiting list for orthotopic liver transplantation (Pozzato 1997; Chalasani 1999; Gambarin-Gelwan 2000; Mortele 2001; de Ledinghen 2002; Libbrecht 2002; Lim 2002; Freeny 2003; Teefey 2003; Van Thiel 2004; Haberman 2011; Yu 2011; Villacastin Ruiz 2016), participants with a defined hepatocellular carcinoma diameter (Golfieri 2009; Kim 2011; Serste 2012; Hsiao 2019), or participants with indeterminate nodules on MRI (Langenbach 2019).

Index test

Risk of bias

We judged 19 studies at low risk, because they clearly predefined the CT positivity criteria (Chalasani 1999; Gambarin-Gelwan 2000; de Ledinghen 2002; Libbrecht 2002; Lim 2002; Freeny 2003; Teefey 2003; Van Thiel 2004; Golfieri 2009; Sangiovanni 2010; Haberman 2011; Kim 2011; Yu 2011; Serste 2012; Maiwald 2014; Lin 2016; Villacastin Ruiz 2016; Hsiao 2019; Langenbach 2019). We judged two studies as unclear for this domain, due to lack of information on CT positivity criteria (Pozzato 1997; Mortele 2001).

Applicability

We judged all studies at low concern.

Reference standard

In 11 studies the reference standard was the pathology of the explanted liver (Pozzato 1997; Gambarin-Gelwan 2000; Mortele 2001; de Ledinghen 2002; Libbrecht 2002; Lim 2002; Freeny 2003; Van Thiel 2004; Haberman 2011; Yu 2011; Villacastin Ruiz 2016), in five studies it was the histology of biopsied focal lesions in all participants (Sangiovanni 2010; Serste 2012; Lin 2016; Hsiao 2019; Langenbach 2019), and in three studies it was the histology of biopsied focal lesions in some participants and follow-up in the others (Chalasani 1999; Kim 2011; Maiwald 2014). Two studies (Golfieri 2009; Teefey 2003), had a mix of pathology of the explanted liver, resection, biopsy, and follow-up.

Risk of bias

We judged nine studies at low risk (Gambarin-Gelwan 2000; Libbrecht 2002; Lim 2002; Sangiovanni 2010; Haberman 2011; Yu 2011; Serste 2012; Lin 2016; Hsiao 2019), 11 at high risk (Chalasani 1999; Mortele 2001; de Ledinghen 2002; Freeny 2003; Teefey 2003; Van Thiel 2004; Golfieri 2009; Kim 2011; Maiwald 2014; Villacastin Ruiz 2016; Langenbach 2019), and one at uncertain risk (Pozzato 1997). The main reasons for judging studies at high risk of bias included statements explaining that reference standard results were interpreted with the knowledge of the results of the index test, and in cases of biopsy, the interventionist had to have knowledge of the presence and location of the lesion in order to perform the procedure. We judged uncertain risk of bias due to lack of detailed information regarding the reference standard.

Applicability

We judged eight studies at low concern (Golfieri 2009; Sangiovanni 2010; Kim 2011; Serste 2012; Maiwald 2014; Lin 2016; Hsiao 2019; Langenbach 2019), and 13 studies at high concern due to orthotopic liver transplantation being the only reference standard (Pozzato 1997; Chalasani 1999; Gambarin-Gelwan 2000; Mortele 2001; de Ledinghen 2002; Libbrecht 2002; Lim 2002; Freeny 2003; Teefey 2003; Haberman 2011; Yu 2011; Van Thiel 2004; Villacastin Ruiz 2016).

Flow and timing

Risk of bias

We judged three studies at low risk of bias (Lim 2002; Yu 2011; Langenbach 2019), 15 studies at high risk (Pozzato 1997; Chalasani 1999; Gambarin-Gelwan 2000; Mortele 2001; de Ledinghen 2002; Libbrecht 2002; Freeny 2003; Teefey 2003; Van Thiel 2004; Golfieri 2009; Sangiovanni 2010; Haberman 2011; Serste 2012; Maiwald 2014; Villacastin Ruiz 2016), and three at unclear risk (Kim 2011; Lin 2016; Hsiao 2019). Reasons for assessing studies at high risk of bias included inappropriate time between index test and reference standard (> 90 days; Pozzato 1997; Gambarin-Gelwan 2000; Mortele 2001; de Ledinghen 2002; Libbrecht 2002; Freeny 2003; Teefey 2003; Van Thiel 2004; Haberman 2011; Villacastin Ruiz 2016)), not all participants underwent the same reference standard (Chalasani 1999; Teefey 2003; Golfieri 2009; Maiwald 2014), and participants missing in the final analysis with no explanations (Freeny 2003; Sangiovanni 2010; Serste 2012; Villacastin Ruiz 2016). We the risk to be unclear due to lack of information on time interval between index test and reference standard. No study reported non-evaluable results.

Overall assessment

We assessed all included studies at high risk of bias. We judged three studies at low concern for applicability for all three QUADAS-2 domains (Sangiovanni 2010; Maiwald 2014; Lin 2016).

Findings

Twenty-one studies with 3101 participants provided data assessing CT for the diagnosis of hepatocellular carcinoma. The median prevalence of the target disease was 52% (IQR 25% to 62%).

Twenty-one studies reported the prevalence of participants with hepatic cirrhosis, and in 16 of them the reported prevalence was 100%. Five studies reported the Child Pugh classification with a median of 54% (IQR 19% to 73%) classified as Child-



Pugh class A. Eighteen studies reported information on liver disease aetiology with a median of 51% (IQR 44% to 73%) having viral aetiology. Sixteen studies reported the proportion of participants with resectable hepatocellular carcinoma, among which 12 reported having more than 90% of participants with resectable hepatocellular carcinoma. Thirteen studies reported the mean diameter of the lesions, with a median of 21 mm (IQR 16 mm to 24 mm).

The studies were conducted from 1997 to 2019. Regarding study location, 10 studies were conducted in Europe, seven in North and South America, and four in Asia. Nineteen studies were conducted in people with clinical suspicion of having a hepatocellular carcinoma, and two studies were conducted in the context of a surveillance programme (Chalasani 1999; Sangiovanni 2010). No study reported uninterpretable index test results.

Among the 11 studies with the pathology of explanted liver as the reference standard, five reported no alternative diagnosis in participants without hepatocellular carcinoma (Pozzato 1997; Gambarin-Gelwan 2000; Lim 2002, Van Thiel 2004; Haberman 2011), Mortele 2001 reported seven macro regenerative nodules in 36 participants without hepatocellular carcinoma, de Ledinghen 2002 reported 16 dysplastic or regenerative nodules in 34 participants without hepatocellular carcinoma, Libbrecht 2002 reported one haemangioma and one focal nodular hyperplasia in 14 participants

without hepatocellular carcinoma, Freeny 2003 reported 296 regenerative nodules in 331 participants without hepatocellular carcinoma, Yu 2011 reported six dysplastic or regenerative macronodules two haemangiomas and one focal infarct in 247 participants without hepatocellular carcinoma, and Villacastin Ruiz 2016 reported six cholangiocarcinomas, two haemangiomas, and six dysplastic nodules in 273 participants without hepatocellular carcinoma.

In the five studies with histology of biopsied focal lesions in all participants, one reported no diagnosis other than hepatocellular carcinoma (Hsiao 2019), one reported 24 out of 60 participants with regenerative nodules (Langenbach 2019), one reported "other liver tumours" without any other specification (Lin 2016), whereas Sangiovanni 2010 reported two out of 69 participants with cholangiocarcinoma, and 21 out of 69 macro regenerative nodules or low-grade dysplastic nodules, and Serste 2012 reported one out of 74 cholangiocarcinoma, one out of 74 epithelioid haemangioendothelioma, nine out of 74 regenerative macro nodule, and nine out of 74 participants with biopsy showing features of chronic liver disease without any features of dysplastic nodule or hepatocellular carcinoma.

Figure 5 shows a forest plot of sensitivity and specificity with 95% CIs. For the 21 studies, the reported sensitivity ranged from 20% to 97% and the specificity ranged from 56% to 100%.

Figure 5. Forest plots of sensitivity and specificity of computed tomography for detection of hepatocellular carcinoma of any size and stage against different reference standards in 21 studies in alphabetical order. Reference standards were: the pathology of the explanted liver in case of transplantation, the histology of resected focal liver lesions, or the histology of biopsied focal liver lesions with a follow-up period of at least six months. Values between square brackets are the 95% confidence intervals (CIs) of sensitivity and specificity. The figure shows the estimated sensitivity and specificity of the study (blue square) and its 95% CI (black horizontal line). CI: confidence interval; FN: false negative; FP: false positive; OLT: orthotopic liver transplantation; TN: true negative; TP: true positive

Study		TP	FP	FΝ	TN	Reference standard	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specifi	icity (95% CI)
Chalasani 19	999	20	11	2	252	Biopsy	0.91 [0.71, 0.99]	0.96 [0.93, 0.98]		•
de Ledinghe	n 2002	17	2	4	11	ÖLŤ	0.81 [0.58, 0.95]	0.85 [0.55, 0.98]		
Freeny 2003	}	15	25	8	296	OLT	0.65 [0.43, 0.84]	0.92 [0.89, 0.95]		-
Gambarin-G	elwan 2000	10	5	9	82	OLT	0.53 [0.29, 0.76]	0.94 [0.87, 0.98]		-
Golfieri 2009)	33	4	21	5	Mixed	0.61 [0.47, 0.74]	0.56 [0.21, 0.86]		-
Haberman 2	:011	28	5	4	25	OLT	0.88 [0.71, 0.96]	0.83 [0.65, 0.94]		-
Hsiao 2019		34	5	7	20	Bi op sy	0.83 [0.68, 0.93]	0.80 [0.59, 0.93]		
Kim 2011		140	4	24	38	Biopsy	0.85 [0.79, 0.90]	0.90 [0.77, 0.97]	-	-
Langenbach	2019	35	1	1	23	Biopsy	0.97 [0.85, 1.00]	0.96 [0.79, 1.00]	-	-
Libbrecht 20	002	1	3	1	11	OLT	0.50 [0.01, 0.99]	0.79 [0.49, 0.95]		
Lim 2002		33	2	8	34	OLT	0.80 [0.65, 0.91]	0.94 [0.81, 0.99]	-	-
Lin 2016		510	41	73	132	OLT	0.87 [0.85, 0.90]	0.76 [0.69, 0.82]	•	-
Maiwald 201	4	22	6	4	18	Bi op sy	0.85 [0.65, 0.96]	0.75 [0.53, 0.90]		
Mortele 200	1	14	0	3	36	OLT	0.82 [0.57, 0.96]	1.00 [0.90, 1.00]		-
Pozzato 199	7	1	2	4	13	OLT	0.20 [0.01, 0.72]	0.87 [0.60, 0.98]	-	
Sangiovanni	2010	15	0	19	21	Bi op sy	0.44 [0.27, 0.62]	1.00 [0.84, 1.00]		_
Serste 2012		35	5	12	22	Biopsy	0.74 [0.60, 0.86]	0.81 [0.62, 0.94]		-
Teefey 2003		6	5	3	11	Mixed	0.67 [0.30, 0.93]	0.69 [0.41, 0.89]		
Van Thiel 20	04	14	0	6	80	OLT	0.70 [0.46, 0.88]	1.00 [0.95, 1.00]		-
Villacastin R	uiz 2016	77	6	18	152	OLT	0.81 [0.72, 0.88]	0.96 [0.92, 0.99]		-
Yu 2011		113	11	36	236	OLT	0.76 [0.68, 0.82]	0.96 [0.92, 0.98]	-	- - - -
									0 0.2 0.4 0.6 0.8 1 0 0.2 (0.40.60.81

We performed a meta-analysis of all 21 included studies using the bivariate model, and we obtained the following pooled estimates: sensitivity 77.5% (95% CI 70.9% to 82.9%), specificity 91.3% (95% CI 86.5% to 94.5%), likelihood ratio: LR+ 8.87 (95% CI 5.67 to 13.86), LR- 0.25 (95% CI 0.19 to 0.32).

Table 1 shows post-test probabilities, calculated using pooled likelihood ratios, according to three different pre-test probabilities.

We assessed the diagnostic accuracy for resectable hepatocellular carcinoma as a secondary objective. We found 10 studies that included participants who all had resectable hepatocellular carcinoma (Pozzato 1997; Gambarin-Gelwan 2000; Mortele 2001; de Ledinghen 2002; Libbrecht 2002; Freeny 2003; Sangiovanni 2010; Yu 2011; Serste 2012; Lin 2016). We performed a meta-analysis and obtained the following estimates: sensitivity 71.4% (95% CI 60.3% to 80.4%) and specificity 92.0% (95% CI 86.3% to 95.5%). Figure 6



shows the forest plot of sensitivity and specificity with their 95% CIs

Figure 6. Forest plots of sensitivity and specificity of computed tomography for detection of resectable hepatocellular carcinoma against different reference standards in 12 studies in alphabetical order. Reference standards were: the pathology of the explanted liver in case of transplantation, the histology of resected focal liver lesions, or the histology of biopsied focal liver lesions with a follow-up period of at least six months. Values between brackets are the 95% confidence intervals (CIs) of sensitivity and specificity. The figure shows the estimated sensitivity and specificity of the study (blue square) and its 95% CI (black horizontal line). CI: confidence interval; FN: false negative; FP: false positive; TN: true negative; TP: true positive

Study	TP	FP	FΝ	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
de Ledinghen 2002	17	2	4	11	0.81 [0.58, 0.95]	0.85 [0.55, 0.98]	
Freeny 2003	15	25	8	296	0.65 [0.43, 0.84]	0.92 [0.89, 0.95]	
Gambarin-Gelwan 2000	10	5	9	82	0.53 [0.29, 0.76]	0.94 [0.87, 0.98]	
Libbrecht 2002	1	3	1	11	0.50 [0.01, 0.99]	0.79 [0.49, 0.95]	
Lin 2016	510	41	73	132	0.87 [0.85, 0.90]	0.76 [0.69, 0.82]	
Mortele 2001	14	0	3	36	0.82 [0.57, 0.96]	1.00 [0.90, 1.00]	
Pozzato 1997	1	2	4	13	0.20 [0.01, 0.72]	0.87 [0.60, 0.98]	
San gio vanni 2010	15	0	19	21	0.44 [0.27, 0.62]	1.00 [0.84, 1.00]	—
Serste 2012	35	5	12	22	0.74 [0.60, 0.86]	0.81 [0.62, 0.94]	
Yu 2011	113	11	36	236	0.76 [0.68, 0.82]	0.96 [0.92, 0.98]	0 0.2 0.4 0.6 0.8 1 0 0.2 0.4 0.6 0.8 1

Heterogeneity analysis

We investigated heterogeneity for all the predefined potential sources (Secondary objectives). Table 2 shows the comparisons of different predefined subgroups. The prevalence of the target disease, reflecting the selection of participants, may in part explain the inconsistency of the overall results. In fact, studies with a prevalence higher than 52% (the median prevalence in the included studies) show a higher sensitivity (81.0%, 95% CI 72.5% to 87.4% compared to 71.1%, 95% CI 60.7% to 79.7%) and a lower specificity (85.5%, 95% CI 78.4% to 90.5% compared to 94.0%, 95% CI 89.8% to 96.5%) than studies with a prevalence lower than 52%. Another possible source of heterogeneity was the inclusion of more than 90% of study participants with resectable hepatocellular carcinoma, that is, the selection of participants with earlier hepatocellular carcinoma. The sensitivity is marginally lower (72.5%, 95% CI 63.2% to 80.1% compared to 76.2%, 95% CI 63.2% to 85.6%) and specificity higher (93.9%, 95% CI 88.7% to 96.9% compared to 82.4%, 95% CI 61.9% to 93.1%) than in studies including less than 90% of resectable hepatocellular carcinoma. The comparison of the other subgroups, assessing the possible role of study date and location, inclusion of participants without cirrhosis, from surveillance programme or clinical cohorts, previous testing with the detection of nodules, and the use of different reference standard did not show differences. Hepatocellular carcinoma mean diameter had no effect on diagnostic accuracy (P = 0.930).

Sensitivity analysis

When considering only the 19 studies that clearly prespecified the positivity criteria, we obtained a pooled sensitivity of 78.3% (95% CI 72.0% to 83.6%) and a specificity of 90.7% (95% CI 85.7 to 94.1%; Table 2).

When considering only the nine studies in which the reference standard results were interpreted without knowledge of the results of the index test, we obtained a pooled sensitivity of 77.5% (95% CI

68.8% to 84.3%) and a specificity of 91.0% (95% CI 83.4% to 95.4%; Table 2).

When considering only the three studies at low concern for applicability (Sangiovanni 2010; Maiwald 2014; Lin 2016), we obtained a pooled sensitivity of 76.9% (95% CI 50.8% to 91.5%) and a specificity of 89.2% (95% CI 57.0% to 98.1%).

We did not perform the planned sensitivity analysis in which studies published only in abstract or letter form were excluded because all included studies were published as full texts.

We did not perform the planned sensitivity analysis in which studies at high risk of bias were excluded as we judged all the included studies to be at high risk of bias.

Summary of findings tables

The main results are shown in Summary of findings 1 and Summary of findings 2.

DISCUSSION

Summary of main results

The aim of this review was to assess the diagnostic accuracy of CT for the diagnosis of hepatocellular carcinoma of any size and at any stage in people with chronic liver disease, either in a surveillance programme or in a clinical setting. We included 21 studies that assessed a total of 3101 participants, 19 were conducted in a clinical setting, and two in a surveillance programme. The main results are presented in Summary of findings 1 and Summary of findings 2.

For the 21 included studies we performed a meta-analysis using the bivariate model, and we obtained the following pooled estimates: sensitivity 77.5% (95% CI 70.9% to 82.9%) and specificity 91.3% (95% CI 86.5% to 94.5%) for the diagnosis of hepatocellular carcinoma at any size and stage (primary outcome). In Table 1 we show the post-test probability of having hepatocellular carcinoma



in the case of positive or negative result of the index test, assuming different values of pretest probability.

Ten studies included only participants with hepatocellular carcinoma amenable for surgical resection, and the pooled estimate of sensitivity was 71.4% (95% CI 60.3% to 80.4%) and specificity 92.0% (95% CI 86.3% to 95.5%) for the diagnosis of resectable hepatocellular carcinoma (secondary outcome).

We judged all included studies to be at high risk of bias in at least one domain, and we assessed the results of 18 out of 21 studies to be at high concern for applicability in the patient selection domain.

Considering only the three studies at low concern for applicability for patient selection (Sangiovanni 2010; Maiwald 2014; Lin 2016), we obtained a pooled sensitivity of 76.9% (95% CI 50.8% to 91.5%) and a specificity of 89.2% (95% CI 57.0% to 98.1%).

We summarised these main results of analyses in Summary of findings 1 and Summary of findings 2, assuming three different prevalence values (20%, 52% and 60%). The prevalence of hepatocellular carcinoma varied widely in all included studies, from 7% to 86%, according to the study design and different settings. For exemplification, we considered three values of hepatocellular carcinoma prevalence: 20% for a population with low clinical suspicion, 52% as a median derived from our study analysis, and 60% for a population with high clinical suspicion (assessment of nodules detected by ultrasound).

For participants with hepatocellular carcinoma at any size and stage, we assumed the following consequences of test results: people with true-positive results, that is, those with hepatocellular carcinoma and positive test results, will receive the appropriate treatment (surgery, local ablative therapy or systemic chemotherapy); people with true-negative results, that is, those without hepatocellular carcinoma and negative test results, will not undergo inappropriate treatment or unnecessary further testing; people with false-negative results, that is, those with hepatocellular carcinoma and negative test results, will be misdiagnosed, not receive the appropriate treatment, and might be detected later as more severe hepatocellular carcinoma patient; people with falsepositive results, that is, those without hepatocellular carcinoma and positive test results, will undergo further testing and possibly an inappropriate treatment. Considering a hypothetical cohort of 1000 people with hepatocellular carcinoma prevalence of 52% (the median value in the included studies), we can expect 117 falsenegative and 42 false-positive results; with a lower prevalence of 20%, we can expect 45 false-negative and 70 false-positive results, and with a higher prevalence of 60%, we can expect 135 falsenegative and 35 false-positive results. We judged the certainty of evidence to be low, downgrading by two levels due to high risk of bias and indirectness.

For participants with resectable hepatocellular carcinoma, considering a hypothetical cohort of 1000 people with hepatocellular carcinoma prevalence of 35%, we can expect 100 false-negative and 50 false-positive results; with a prevalence of 20%, we can expect 57 false-negative and 60 false-positive results; with a prevalence of 60%, we can expect 166 false-negative and 30 false-positive results. We judged the certainty of evidence to be low, downgrading by two levels due to high risk of bias and indirectness.

Strengths and weaknesses of the review

Strengths and weaknesses of included studies

This review included a total of 21 studies, covering a time span of 22 years, from 1997 to 2019 and wide geographical areas, including areas with high and low prevalence of chronic liver disease and hepatocellular carcinoma. Ten studies were conducted in Europe, seven in North and South America, and four in Asia. In terms of number of participants, studies performed in the Americas included 1318 participants, in Asia 1105, and in Europe 678. We found no study from Africa, where hepatocellular carcinoma is highly prevalent (Ferlay 2019).

An overall quality assessment of the studies showed their methodological weaknesses. We assessed all studies at high risk of bias mainly due to inappropriate exclusion criteria, reference standard results interpreted with knowledge of the index test (unavoidable in cases of biopsy), and time interval between index test and reference standard of more than 90 days. The choice of reference standard represents a major concern for all studies, and we recognise none is perfect. The most common reference standard was pathology of the explanted liver (11 studies), the most accurate, allowing the histological evaluation of the whole liver in all participants. However, this almost perfect reference standard is possible only in studies conducted on participants with advanced and decompensated liver diseases on a waiting list for transplantation, that do not represent the intended spectrum of liver disease severity. In fact, we aimed to assess CT accuracy in participants with the whole spectrum of liver disease severity without any exclusion for severity of liver disease or hepatocellular carcinoma volume. Accordingly, correct estimates of CT accuracy can be obtained only at the expense of their applicability. The other reference standards were histology of biopsied focal lesions with adequate follow-up (8 studies) and mix of pathology of the explanted liver, resection, biopsy, and follow-up (2 studies). We judged 10 studies in which the time interval between the index test and reference standard was longer than 90 days to be at high risk of bias. In fact, in diagnostic test accuracy assessment, it is necessary to have the time interval between index test and reference standard as short as possible (Colli 2014). Longer time intervals impair accurate assessment due to possible changes in lesion size and morphologic features during certain periods of time. According to the latest systematic review, the approximate hepatocellular carcinoma volume doubling time is four to five months, with significant range of 2.2 months to 11.3 months (Nathani 2021). In accordance with suggestions from a previous systematic review, which noted the acceptable time interval being from one to three months (Kim 2008), we assumed 90 days to be the most acceptable threshold.

We found no studies that reported on uninterpretable results of the index test. Such a failure of reporting or excluding them from the analysis could have produced an overestimation of the obtained accuracy estimates. In fact, in the process of visual interpretation of CT examinations, sometimes it is impossible for the radiologist to make a definite diagnosis of hepatocellular carcinoma. This is primarily due to unclear visual representation and absence of morphological criteria needed for a definite diagnosis (non-rim like hyperenhancement, non-peripheral washout in portal-venous and subsequent phases, enhancing capsule, etc.; LI-RADS 2018). Technical aspects of a CT examination such as participant movement and breath-hold, scanning protocol, application of



adequate type and amount of contrast, and acquisition of correct phases (arterial, portal-venous, late phase) can impair liver imaging and its correct interpretation.

Using QUADAS 2, we judged 18 out of 21 studies at high concern for applicability mainly due to the selective inclusion of participants with decompensated advanced liver disease or a definite hepatocellular carcinoma diameter, and the use of pathological examination of the whole liver as the reference standard.

Not all studies reported on all covariates that we planned to assess as a possible source of heterogeneity, and this might have impaired the analyses. Most information on MELD (Model for End-Stage Liver Disease), Child A and CT detector number was missing.

Strengths and weaknesses of the review process

Search strategy

Our search strategy provided a significant number of studies that were performed in various geographical areas with high and low prevalence of chronic liver disease and hepatocellular carcinoma. Manually searching the references of the included studies and previous narrative and systematic reviews identified three additional studies, which were ultimately included in the final analysis. We applied no language restrictions in the inclusion criteria, which resulted in retrieving full-text articles of 24 studies published in non-English languages, two of which we included in the final analysis. We requested further information from study authors regarding two studies, but they did not provide any information. We are confident that the search strategy resulted in the detection of most eligible studies, with a low probability of undetected relevant studies.

Quality assessment and data extraction

We consider our attempts to reduce subjectivity in our judgements to minimise errors and miscalculations in data extraction to be the strength of this review. Two review authors independently assessed the risk of bias of the included studies and applicability of their results using the QUADAS-2 tool. We extracted data using a proper form. In case of disagreement, we reached consensus through discussion. Disagreements were most frequent for the two QUADAS-2 domains patient selection (six studies), and reference standard (five studies). All agreements were reached through discussion between two review authors, and the conclusions were discussed and approved by a third review author. For data extraction, most of the discordances were due to miscalculations and typos, which were easily resolved. The same review authors assessed the certainty of evidence using the GRADE approach and the level of agreement was high.

Review analysis

We performed a meta-analysis using the bivariate model, as the results of the index test were reported as dichotomous (positive or negative) with no explicit threshold. Anyway, we recognise that, nevertheless, implicit thresholds cannot be excluded. The pooled estimates of sensitivity ranged from 20% to 97% and those of specificity from 56% to 100%. Two studies included fewer than 20 participants and their results were quite imprecise with a very wide confidence interval (Pozzato 1997; Libbrecht 2002; Figure 5). Studies with a prevalence of the target disease higher than the median shows higher sensitivity and lower specificity,

suggesting that the selection of participants may in part explain the inconsistency of the overall results. The difference in the results between studies that included more than 90% of participants with resectable hepatocellular carcinoma compared to studies that included less than 90% may confirm the possible role of participant selection. Whereas the case mix of the participants in the included studies, at least when adequately reported, seems homogeneous, showing similar numbers and types of alternative diagnosis, encompassing regenerative and dysplastic nodules, and more rarely haemangioma cholangiocarcinoma and focal nodular hyperplasia. Moreover, the study setting (clinical or surveillance programme), different geographic areas, advancements in technology (before and after the year 2005), aetiology of the underlying liver disease, or its severity (prevalence of cirrhosis), hepatocellular carcinoma diameter, and difference in the choice of the reference standards and even prior testing with detection of nodules seem unable to explain the observed inconsistencies.

However, some of our planned investigations were not possible due to lack of data (number of CT detector rows, Child-Pugh classification of severity of cirrhosis), and lack of published studies (before and after 2016 to assess the changes in diagnostic criteria). Furthermore, we were able to investigate only characteristics that could be assessed at study level whereas participants' factors or hepatocellular carcinoma characteristics can only be assessed by aggregate statistics with the inherent risk of ecological bias. Therefore, some important relationships, such as the one with hepatocellular carcinoma volume, could have been missed. In addition, many of the included studies did not report data on the covariates of interest. Finally, other potential sources of heterogeneity were not planned, and might be assessed in future studies, such as CT slice thickness, contrast injection rate, contrast type, number of exam phases, and collimation.

We excluded studies that reported only per-lesion analyses and included only studies with per-patient analyses. Per-patient and per-lesion analyses represent two different approaches to diagnostic accuracy assessment and their choice depends on the type of clinical or scientific question, and requires different and appropriate statistical methodology. In the present review, we aimed to assess the accuracy of CT for the diagnosis hepatocellular carcinoma. Consequently, we chose to include studies that evaluated how CT is able to detect patients with hepatocellular carcinoma at any size and any stage, therefore we applied a perpatient approach. Otherwise, per-lesion analysis is properly used to assess accuracy in detecting multiple lesions on a single image, providing information that is relevant for hepatocellular carcinoma staging. Studies planning per-lesion analysis require a different methodological approach and cannot be pooled together with studies using a per-patient approach (Chang 2006; Zwinderman 2008). Furthermore, the inclusion criteria of studies planning a perlesion analysis are quite different and do not match our review question. In fact, they usually do not include participants with chronic liver disease and suspected hepatocellular carcinoma, but participants with known focal liver lesions, encompassing hepatocellular carcinomas, cholangiocarcinomas, benign liver tumours and even metastases from abdominal or extra-abdominal primary cancers.

We assessed only the impact of the presence of diagnostic criteria on diagnostic accuracy, and we did not assess the differences



in various criteria. In our review, most primary studies used perfusion positivity criteria to assess the lesion as hepatocellular carcinoma: non-rim like hyperenhancement and washout in subsequent phases. A significant number of around 40% of the liver biopsies is reported to be morphologically atypical hepatocellular carcinomas, which surely may impact the accuracy (Kim 2019). The use of perfusion criteria without any additional criteria for non-hyperenhancing hepatocellular carcinoma in arterial phase could explain the high proportion (more than 25%) of false-negative results.

We were unable to assess the effect of uninterpretable results on diagnostic accuracy as no study reported such data. Indeed, it is possible that failures in obtaining adequate images were not reported, with consequent overestimation of CT accuracy.

The sensitivity analysis shows that the obtained results are arguably robust with no variation, after including only studies that clearly prespecified the positivity criteria, and including only those in which the reference standard results were interpreted without the knowledge of the results of the index test.

Comparison with previous research

We found 11 non-Cochrane systematic reviews that assessed the accuracy of CT for detection of hepatocellular carcinomas (Colli 2006; Xie 2011; Chen 2013; Floriani 2013; Chou 2015; Lee 2015; Ye 2015; Guo 2016; Hanna 2016; Roberts 2018; Li 2019). All reviews assessed the accuracy of CT and MRI, and some also assessed ultrasound (Colli 2006; Floriani 2013; Chou 2015; Hanna 2016), contrast-enhanced ultrasound (Xie 2011), and alpha-foetoprotein (Colli 2006). Due to differences in methodological approach, inclusion and exclusion criteria, and in statistical analysis, these results are not comparable to each other nor to our present results. The pooled sensitivity of CT for the diagnosis of hepatocellular carcinoma in these reviews ranged from 61% to 86% and the specificity from 64% to 94% (Table 3). Three systematic reviews performed per-patient analysis (Colli 2006; Floriani 2013; Chou 2015), and the pooled sensitivity and specificity of CT for detection of hepatocellular carcinoma ranged from 67.5% to 83% and 72% to 92.5% (Table 3). These results are in accordance with our present results, despite methodological differences and the number of included studies. We additionally evaluated all the primary studies included in these systematic reviews and assessed them for inclusion in our analysis.

Applicability of findings to the review question

Using the QUADAS-2 tool, we assessed the applicability of the results of the included studies. We judged most studies to be at high concern in the domains of patient selection and reference standard. In the patient selection domain, the main concerns were the inclusion of only patients on the waiting list for orthotopic liver transplantation with decompensated chronic advanced liver disease or the inclusion of participants according to the hepatocellular carcinoma diameter. The choice of the pathology of the explanted liver as the reference standard also impairs the applicability of the results as this reference standard is applied exclusively to transplanted patients.

AUTHORS' CONCLUSIONS

Implications for practice

Hepatocellular carcinoma is a frequent complication of chronic liver disease. The detection of a tumour amenable to surgical resection, thermal ablation, or liver transplantation could improve the prognosis, which in the absence of indications to radical treatment is severe. Being the fourth leading cause of death from cancer worldwide, accurate tests are needed to diagnose hepatocellular carcinoma. In the clinical pathway for the diagnosis of hepatocellular carcinoma in people with chronic liver disease, computed tomography (CT) is currently the second step after ultrasound and alpha-foetoprotein or the combination of the two, and its main role is to confirm the presence of the disease.

As an ideal diagnostic test, CT should ensure a low proportion of false-negative results because people with undetected hepatocellular carcinoma cannot receive proper treatment. People with false-positive results are exposed to unnecessary further diagnostic workup and possible invasive treatment. The estimated pooled sensitivity and specificity derived from our analysis suggest that 22.5% of people with hepatocellular carcinoma would be missed, and 8.7% of people would be unnecessarily treated.

An important piece of clinical information, which is meaningful for further patient workup, is the possibility of surgical resection. Ideally, CT should ensure a low proportion of falsenegative results because people with false-negative results will not undergo surgical resection, and people with false-positive results will undergo inappropriate surgical resection. Based on our results 28.6% of people with hepatocellular carcinoma would be incorrectly classified as without any hepatocellular carcinoma and would improperly not be resected, while 7.7% of people with non-resectable hepatocellular carcinoma will undergo inappropriate surgery. For people on a waiting list for orthotopic liver transplantation for an indication not related to a hepatocellular carcinoma, the consequences of false-negative results of preoperative CT are not completely known and might be less severe: indeed studies report no significant difference in terms of overall survival and tumour recurrence compared to people with previously diagnosed hepatocellular carcinomas (Castillo 2009; Senkerikova 2014; Madaleno 2015; El Moghazy 2016).

The main hallmarks of hepatocellular carcinoma on a CT study are non-rim-like hyperenhancement in arterial phase, and washout in portal-venous and delayed phases. However, around 40% of hepatocellular carcinomas present with atypical morphological features, which pose a significant diagnostic challenge for radiologists. This significant number of atypical hepatocellular carcinomas may influence the sensitivity, and the radiologist should be acquainted to these atypical appearances to correctly interpret CT findings. Another issue is the presence of hepatocellular carcinoma mimickers, such as intrahepatic cholangiocarcinoma, combined hepatocellular carcinoma-cholangiocarcinoma, arterioportal shunt, and haemangioma in cirrhotic liver (Lee 2012b; Shirki 2015; Kim 2019).

Apart from correctly classifying people of having hepatocellular carcinoma and assessing them amenable for resection, another important role of CT is to correctly stage the disease as local, regional, or disseminated disease. However, this issue was not the aim of this review.



Overall, caution is needed in interpreting our review results as we judged all the studies at high risk of bias, and most of them with high concern regarding their applicability, mainly due to patient selection and reference standard domain.

Implications for research

Currently available evidence on the diagnostic accuracy of CT for diagnosis of hepatocellular carcinoma is not conclusive. Therefore, more high-quality primary studies are needed. With introduction of LI-RADS criteria, the results of CT studies are no longer needed to be dichotomised allowing inconclusive and probable results to be assessed also. Apart from typical hepatocellular carcinoma appearances, atypical features of hepatocellular carcinoma need to be taken into consideration, so we hypothesise that further studies using LI-RADS positivity criteria may improve sensitivity. Also, it may be possible that including additional major features such as threshold growth, along with arterial hyperenhancement and subsequent washout may improve sensitivity. Therefore, we welcome future cross-sectional studies using score systems of positivity criteria. In further research, other sources of

heterogeneity may be assessed such as CT slice thickness, contrast injection rate, contrast type, number of exam phases, and collimation.

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CHARACTERISTICS OF STUDIES

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Chalasani 1999			
Study characteristics			
Patient Sampling		r transplantation in	ents with cirrhosis who a tertiary referral center
	166 patients were cons tients were found to ha		ansplantation, and 27 pa-
Patient characteristics and setting	The study included onler transplantation.	y participants with o	cirrhosis who underwent liv
Index tests	chine (CT-Twin; Elscint phased in nature and pionic contrast at 4 mL/ed slice width of 5 mm suspicious lesion on Chancement on arterial venous-phase images. formed all the US exan or abdominal CT were	, Inc; Rockleigh, NJ). performed after IV in s. The scanning was and a reconstructio I was defined as a sc phase and was hypo In general, ultrason ninations. Any lesion biopsied under US g	a a spiral CT scanning ma- The scanning was dual- jection of 150 mL of non- performed with a collimat- n increment of 4 mm. A olid mass that showed en- o-, iso-, or hypervascular on ography technicians per- is suspicious for HCC on US uidance. Only 1 lesion with ose who had multiple le-
Target condition and reference standard(s)	The diagnosis of HCC was based on histology in 26 participants. In 1 participant, the diagnosis of HCC was based on characteristic CT and hepatic angiographic findings. The absence of HCC was based on follow-up with US and CT until liver transplantation or death.		
Flow and timing	No data on interval be	tween index test and	l reference standard
Comparative			
Notes	No data on conflict of i	nterest	
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)		,	
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes	,	



Ch	ala	sani	19	99	(Continued)
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Could the conduct or interpretation of the index test	
have introduced bias?	

Low risk

Are there concerns that the index test, its conduct, or interpretation differ from the review question?

Low concern

DOMAIN 3: Reference Standard

Is the reference standards likely to correctly classify the target condition?

No

No

Were the reference standard results interpreted without knowledge of the results of the index tests?

Could the reference standard, its conduct, or its interpretation have introduced bias?

High risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?

Unclear

Did all patients receive the same reference standard?

Nο

Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?

Yes

Could the patient flow have introduced bias?

High risk

de Ledinghen 2002

Study characteristics

Patient Sampling This prospective study included 34 participants from February 1997-July 1999, who had both MRI and spiral CT performed before OLT. In the participant selection process 20 patients were excluded because they did not have both MRI and spiral CT for medical or economic reasons.

Patient characteristics and setting The study included patients with cirrhosis who underwent liver transplantation.

Index tests On spiral CT, all enhanced nodules during arterial phase were interpreted as HCC.

Target condition and reference standard(s)

Liver histology of the whole explanted liver was considered as the gold standard for HCC. In all cases, the pathologists were aware of the presence or absence of a HCC diagnosed at radiology and, most of the time, the gross location (right or left lobe) of the tumour was known.



le Ledinghen 2002 (Continued)			
Flow and timing	Range of the interval was 1-161 days.	al between index test	and reference standard
Comparative			
Notes	No data on conflict	of interest	
Methodological quality			
Item	Authors' judge- ment	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	No		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	No		
Did all patients receive the same reference standard?	Yes		



de Ledinghen 2002 (Continued)

Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?

Yes

High risk

Freeny 2003

Study characteristics			
Patient Sampling	This study evaluated 354 consecutive patients who had hepatic transplantation between December 1992-March 1999. All participants underwent a dual-phase CT as part of their routine pretrans plantation evaluation. All participants underwent OLT only. 61 hyperattenuating nodules were identified on arterial phase CT in 43 participants.		
Patient characteristics and setting	Only study participa	ants who underwent (OLT were analysed.
Index tests	CT - all arterial phas as potential tumou		odules were considered
Target condition and reference standard(s)	mm intervals and e tion. Nodules suspi by hematoxylin and ipants with suspicion were identified at tl	ach section was evalu cious for HCC, were ev l eosin and reticulin-s ous liver nodules on p ne time of liver section m sections of the exp	were sectioned at 10 lated by gross inspec- valuated histologically tained sections. Partic- retransplant CT scans ning and the scans corre- lanted liver. All included
Flow and timing	Time interval between index test and ref. standard is 11-704 days. In the final count of results, 10 participants were missing.		
Comparative			
Notes	No information on o	conflict of interest	
Methodological quality			
Item	Authors' judge- ment	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			



Yes		
Yes		
	Low risk	
		Low concern
Yes		
No		
	High risk	
		High
No		
Yes		
No		
	High risk	
	Yes Yes No No Yes	Yes Low risk Yes No High risk No Yes No

Study characteristics	
Patient Sampling	This study analysed retrospectively the charts of 106 consecutive adult patients who underwent OLT for treatment of cirrhosis over a 1-year period at Mount Sinai Hospital. All participants had US, CT, and serum AFP measurements within 6 months of OLT. The results were compared to explant histology. Pathological analysis of 106 explants revealed HCC in 19 participants.
Patient characteristics and setting	Only patients who underwent OLT were analysed.
Index tests	CT: All participants underwent conventional CT scans, performed on a GE 9800 CT scanner (General Electric, Milwaukee, WI). Serial transaxial scans were obtained from the diaphragm to the iliac crests by 10 mm collimation sections. Scans were obtained during suspended respiration after administration of oral contrast (E-Z-CAT, E-Z-M, Inc., West-



Gambarin-Gelwan 2000 (Continued)			
	Pharmaceuticals, Inc.,	Barceloneta, Puerto	Omnipaque 240, Sterling Rico). Radiological exami- ecialising in the hepatobil-
		given a semiquantita	results reviewed US and CT ative score of 1–4, based up-
Target condition and reference standard(s)	HCC. A pathologist specialising in the hepatobiliary system reviewed all liver explants. Each liver explant was sectioned every 1 cm. The presence of tumour nodules, their size, and their location were recorded. The underlying liver pathology was evaluated.		ctioned every 1 cm. The d their location were record-
Flow and timing	The time interval is < 180 days, therefore some participants had interval > 90 days.		
Comparative			
Notes	No information on Col		
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		



Could the reference standard, its conduct, or its interpretation have introduced bias?

Low risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?

No

Did all patients receive the same reference standard?

Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?

Yes

Could the patient flow have introduced bias?

High risk

Golfieri 2009

Study characteristics	
Patient Sampling	This prospective study was performed at a tertiary liver care centre. 283 consecutive cirrhotic patients were recruited between July 2003-October 2004. The final study group included 63 participants, out of whom 54 had HCC.
	220 participants were excluded for the following reasons: having no nodules or benign regenerative nodules RN (n = 122), large (> 3 cm) HCC (n = 4), previously treated HCC (n = 94).
Patient characteristics and setting	Patients with HCC > 3 cm were excluded.
Index tests	CT: quadruple-phase MDCT (i.e. unenhanced, hepatic arterial, portal-venous and delayed phases) was performed using a multidetector-row CT scanner (Emotion 6, Siemens Medical Systems, Erhlangen, Germany).
	Positivity criteria: at MDCT and dynamic MRI, nodules showing arterial enhancement plus washout or a coronal enhancement in the portal and/or delayed phase were considered to be typical HCCs, according to the EASL criteria.
Target condition and reference standard(s)	The final diagnosis was established at pathology on the explanted liver (n = 10), resection (n = 6) and biopsy (n = 38) specimens or at 2 years' follow-up (n = 9).
Flow and timing	The mean imaging-resection interval was 3 months (range 12-88 days).
Comparative	
Notes	No information on Col
Methodological quality	



Golfieri 2009 (Continued)

Item	Authors' judge-Risk of bias ment		Applicability con- cerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			
Did the study avoid inappropriate exclusions?	No			
Could the selection of patients have introduced bias?		High risk		
Are there concerns that the included patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (All tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
Were positivity criteria clearly defined?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Low risk		
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern	
DOMAIN 3: Reference Standard				
Is the reference standards likely to correctly classify the target condition?	Yes			
Were the reference standard results interpreted without knowledge of the results of the index tests?	No			
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk		
Are there concerns that the target condition as defined by the reference standard does not match the question?			Low concern	
DOMAIN 4: Flow and Timing				
Was there an appropriate interval between index test and reference standard?	Yes			
Did all patients receive the same reference standard?	No			
Were all patients included in the analysis and analysed accord- ing to intention to diagnose principle (non-evaluable results considered as false)?	Yes			
Could the patient flow have introduced bias?		High risk		



Haberman 2011

Study characteristics				
Patient Sampling	and had liver transp 2007-March 2010. Al histological analysis final reference of th correlated with tom	plantation at a tertiary Il participants include s of the explanted live e diagnosis of HCC. Pa	nts who were evaluated institution from January d were transplanted and r was considered as the athological reports were formed at the time of the ants.	
	Patients with time in dard of > 6 months w		k test and reference stan-	
Patient characteristics and setting	All participants und	erwent OLT.		
Index tests		nade at the time of th	re correlated with the to- e comprehensive evalua-	
	The definition of HCC by MDCT - a solid, hypodense lesion without contrast, with moderate to intense and inhomogeneous enhancement in the arterial phase, with isodensity or decreased enhancement (washout) in the portal phase and with confirmation of the enhancement washout in the late phase.			
Target condition and reference standard(s)	Pathology of explanted liver. The anatomopathological report were correlated with the tomographic reports made at the tim the comprehensive evaluation of the participants.		oorts made at the time of	
Flow and timing	Interval between index test and reference standard is < 180 c		e standard is < 180 days	
Comparative				
Notes	No information on Col			
Methodological quality				
Item	Authors' judge- ment	Risk of bias	Applicability con- cerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Unclear			
Did the study avoid inappropriate exclusions?	No			
Could the selection of patients have introduced bias?		High risk		
Are there concerns that the included patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (All tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
Were positivity criteria clearly defined?	Yes			



Haberman 2011 (Continued)

introduced bias?

	•	
Could the conduct or	interpretation of the index test have	

Are there concerns that the index test, its conduct, or interpretation differ from the review question?

Low concern

DOMAIN 3: Reference Standard

Is the reference standards likely to correctly classify the target condition?

Yes

Yes

Were the reference standard results interpreted without knowledge of the results of the index tests?

.

Low risk

Low risk

Could the reference standard, its conduct, or its interpretation have introduced bias?

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?

No

Did all patients receive the same reference standard?

Yes

Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?

Yes

Could the patient flow have introduced bias?

High risk

Hsiao 2019

Study characteristics	
Patient Sampling	This is a prospective study, which began on 1 December 2017 and finished on 1 December 2018. The first participant was recruited on 14 December 2017 and the last was recruited on 26 November 2018. Participants were recruited from outpatient clinics. Patients with lesions> 3 cm were excluded.
Patient characteristics and setting	The study included solitary liver tumour < 3 cm only, and patients without CLD.
Index tests	CT. Dynamic CT represented quadruple-phase imaging series (precontrast phase, arterial phase, portal-venous phase, and equilibrium phase), in which non-ionic iodine-based contrast medium was used. Physicians involved in this study were blinded from the results of other examinations while formulating their interpretations.
	Positivity criteria: 0—not detected, 1—HCC, 2—metastasis, 3—benign tumour, 4—uncertainty



Hsiao 2019 (Continued) Target condition and reference standard(s) The pathology results were treated as a reference diagnosis, which was originally reported by textual description and later classified using the following schema: 0—no tumour, 1—HCC, 2 metastasis, 3—benign tumour. Physicians involved in pathology analysis were blinded from the results of other examinations while formulating their interpretations. Flow and timing No information on interval between index test and reference stan-Comparative The authors declare no conflict of interest. Notes Methodological quality Item Authors' judge-Risk of bias Applicability conment cerns **DOMAIN 1: Patient Selection** Was a consecutive or random sample of patients enrolled? Yes Did the study avoid inappropriate exclusions? Nο Could the selection of patients have introduced bias? High risk Are there concerns that the included patients and setting do High not match the review question? DOMAIN 2: Index Test (All tests) Were the index test results interpreted without knowledge of Yes the results of the reference standard? Were positivity criteria clearly defined? Yes Could the conduct or interpretation of the index test have Low risk introduced bias? Are there concerns that the index test, its conduct, or inter-Low concern pretation differ from the review question? **DOMAIN 3: Reference Standard** Is the reference standards likely to correctly classify the target Yes condition? Were the reference standard results interpreted without knowl-Yes edge of the results of the index tests? Could the reference standard, its conduct, or its interpreta-Low risk tion have introduced bias?

Are there concerns that the target condition as defined by

the reference standard does not match the question?

Low concern



Hsiao 2019 (Continued)

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?	Unclear
Did all patients receive the same reference standard?	Yes
Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?	Yes
Could the patient flow have introduced bias?	Unclear risk

Kim 2011

Study characteristics	
Patient Sampling	This study prospectively investigated 206 consecutive patients with hepatic masses > 2 cm.
	Patients who had undergone CT as a staging work-up for a known primary extrahepatic malignancy were excluded. Patients who were in the terminal stages of disease and/or who had severe coagulopathy were also excluded because confirmation of the diagnosis would not be helpful for clinical decision and treatment. Patients with intraperitoneal bleeding from spontaneously ruptured tumours were also excluded to perform emergent transarterial chemoembolisation. Patients with inconclusive FNB results were excluded from the analysis and 68 had hepatic nodules between 1-2 cm in diameter and were therefore excluded from this study.
Patient characteristics and setting	Patients with liver mass > 2 cm included only
Index tests	CT examinations were performed using a helical CT (GE Light Speed VC-TXT, General Electric Medical Systems, Milwaukee, WI, USA) with a 4-phase (precontrast, arterial, portal and delayed phases) technique.
	Index test results were interpreted before reference standard (prospective study).
	Positivity criteria: CT enhancement patterns of lesions hypervascular in the arterial phase and washed out in the portal/delayed phase were classi fied as typically vascular. Tumour(s) composed of mixed areas of arterial hyper- and hypovascularity, but > 70% hypervascular area, were arbitrarily considered as showing a typical enhancement pattern to exclude the possibility of hepatocholangiocarcinoma.
Target condition and reference standard(s)	HCC. Biopsy results were considered the gold standard with follow-up 12 months for benign lesions
Flow and timing	No information on interval between index test and reference standard
Comparative	
Notes	The authors declare no conflict of interest.
Methodological quality	



Kim 2011 (Continued)

Item	Authors' judgement	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	No		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as de- fined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis and analysed according to intention to diagnose principle (nonevaluable results considered as false)?	Yes		
Could the patient flow have introduced bias?		Unclear risk	



Langenbach 2019

Study characteristics			
Patient Sampling	This is a retrospective study performed at a single centre. All patients participating in this study underwent a Lipiodol-based angiography, which was followed by percutaneous CT-guided liver biopsy. From March 2016-February 2017, 92 consecutive eligible patients were screened, and 36 had HCC.		
	sies showed a histo generative nodules		er than HCC or benign res s without a contrast-en-
Patient characteristics and setting	odol-based angiogr	aphy which was follow This is according to o	tudy underwent a Lipi- wed by percutaneous CT- our local standard proce-
		all patients whose bid ity other than HCC or	opsies showed a benign regenerative nod-
Index tests	CT. The evaluation of the lesions in angiography and CT was p formed blind by 2 senior radiologists independently, each wit years of experience in diagnostic and interventional radiology agnostic criteria for HCC: combination of 2 independent HCC picious criteria		ependently, each with > 5 erventional radiology. Di-
Target condition and reference standard(s)	Quote: "Biopsy results were considered the gold standard"		he gold standard".
Flow and timing	CT evaluation was performed using the plain CT scan dataset of the upper abdomen used as biopsy planning scan.		
Comparative			
Notes	The authors declare	no conflict of interes	t.
Methodological quality			
Item	Authors' judge- ment	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	No		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)		,	



angenbach 2019 (Continued)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?	Yes		
Could the patient flow have introduced bias?		Low risk	

Libbrecht 2002

Study characteristics	
Patient Sampling	Between January 2000-July 2001, a total of 52 patients with liver cirrhosis underwent liver transplantation. Within the 6 months before transplantation, contrast-enhanced CT was performed in 16 patients (33%). Out of 16 patients, 2 had HCC.
	3 patients without chronic HCV infection for whom it was clear that their tumours exceeded the mentioned number and size limits received a donor liver from a patient with positive serological markers for HCV. These 3 patients were excluded from the study.
Patient characteristics and setting	All participants underwent OLT only.
Index tests	CT examinations were performed in the setting of pretransplantation evaluation and collected after pathological examination



Libbrecht 2002 (Continued) of the explant liver. These reports were made by a fellow and 1 of three different abdominal radiologists who interpreted results of imaging examinations in consensus according to internationally accepted criteria. All cirrhotic explant livers were examined without knowledge of clinical or imaging data. On contrast-enhanced CT, nodular lesions that were hypodense during the arterial phase were interpreted as DNs, and enhanced nodules during the arterial phase were interpreted as HCCs. Target condition and reference standard(s) Reference standard: pathology of the whole explanted liver. All cirrhotic explant livers were examined without knowledge of clinical or imaging data. Flow and timing For participants who underwent CT, time interval range was 22-179 Comparative Notes No information on Col Methodological quality Risk of bias Authors' judge-Applicability con-Item ment cerns **DOMAIN 1: Patient Selection** Was a consecutive or random sample of patients enrolled? Yes Did the study avoid inappropriate exclusions? Yes Could the selection of patients have introduced bias? Low risk Are there concerns that the included patients and setting do High not match the review question? **DOMAIN 2: Index Test (All tests)** Were the index test results interpreted without knowledge of Yes the results of the reference standard? Were positivity criteria clearly defined? Yes Could the conduct or interpretation of the index test have Low risk introduced bias? Are there concerns that the index test, its conduct, or inter-Low concern pretation differ from the review question? **DOMAIN 3: Reference Standard** Is the reference standards likely to correctly classify the target Yes condition? Were the reference standard results interpreted without knowl-Yes edge of the results of the index tests?



Li	bl	orec	ht	200	2 (Continued)
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Could the reference standard, its conduct, or its interpreta- tion have introduced bias?	Low risk
Are there concerns that the target condition as defined by the reference standard does not match the question?	High
DOMAIN 4: Flow and Timing	
Was there an appropriate interval between index test and reference standard?	No
Did all patients receive the same reference standard?	Yes

Yes

Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?

Could the patient flow have introduced bias?

High risk

Lim 2002

Study characteristics	
Patient Sampling	3-phase helical dynamic CT in 77 patients with advanced liver cirrhosis was evaluated prospectively before OLT. From February 1998-April 2002, a total of 108 consecutive adult patients underwent whole liver transplantation. Among them, 77 patients had 3-phase helical dynamic CT for the evaluation of the liver in terms of hepatic anatomy and liver volume as well as detection of tumour before transplantation, and this group formed the basis of this study. A total of 72 HCCs were confirmed histopathologically in 41 patients.
	31 patients who had conventional CT at outside hospitals using third- generation scanners were excluded from the analysis.
Patient characteristics and setting	All participants underwent OLT.
Index tests	3-phase helical CT scanning was performed in all of the 77 participants using HiSpeed Advantage helical scanners (General Electric Medical Systems, Milwaukee, WI).
	The CT criteria for the diagnosis of hepatocellular carcinoma were a nodule showing the enhancement pattern of the hepatic arterial supply and lack of the portal venous supply (e.g. hyperattenuation on the hepatic arterial or portal-venous phases, low attenuation or isoattenuation on the portal-venous and delayed phases compared to the adjacent parenchyma). Low attenuating nodules ≥ 2 cm in diameter showing a distinct margin during all 3 phases or on delayed phase were considered HCC.
Target condition and reference standard(s)	Sectioning of the liver specimens was independently performed without information of CT findings.
Flow and timing	The time interval between CT examination and surgery was 0~76 days (mean, 27.3 days), < 90 days



Lim 2002 (Continued)

Comparative			
Notes	No information on Col		
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	No		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?	Yes		



Lim 2002 (Continued)

Could the patient flow have introduced bias?

Low risk

Lin 2016

Study characteristics			
Patient Sampling	In this retrospective study between January 2006-October 2010, 1016 patients underwent liver tumour resections or liver transplantation in the Chang Gang Memorial Hospital, Kaohsiung, Taiwan. Of these, 841 patients underwent liver CT or MRI examinations or had a pathological fibrosis score analysis, and were therefore enrolled in this study.		
	The exclusion criteria were patients who did not undergo liver CT or MRI examination before surgery, did not have a pathological fibrosis score analysis, or did not have liver tumours in the explanted liver.		
Patient characteristics and setting	Participant characteristics and setting match the scope of this review, patients were included regardless of the stage of CLD. No restrictions in HCC lesion size were applied.		
Index tests	CT examinations were performed using a helical CT (Toshiba, 120KVP) with a 4-phase (non-contrast, arterial, portal and delayed phases) technique.		
	The study defined typical HCC imaging characteristics as early enhancement in the artery phase and early washout in the venous phase.		
	4-phase liver CT or dynamic liver MRI images were read by radiologists with extensive experience in liver and HCC imaging.		
Target condition and reference standard(s)	Reference standard: histological and surgical reports were reviewed to confirm HCC; resection or transplantation.		
	Pathological results were read by pathologists with sufficient experience in the field and who were blinded to the clinical and radiological results.		
Flow and timing	No data on time interval between index test and reference standard		
Comparative			
Notes	No information on Col		
Methodological quality			
Item	Authors' judgement Risk of bias Applicability concerns		
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	No		
Could the selection of patients have introduced bias?	High risk		



Are there concerns that the included patients and set- ting do not match the review question?			Low concern
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?	•		Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the tar get condition?	- Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing	,		
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?	Yes		
results considered as false):		Unclear risk	

Study characteristics	
Patient Sampling	50 patients with suspected or proven HCC were included in this prospective single-centre study to evaluate the diagnostic performance of contrast-enhanced CT and Gd-EOB-DTPA-enhanced MRI in terms of lesion detection. 26 patients had HCC.
	Exclusion criteria: renal failure, allergy to contrast agents, hyperthyreoidism, pregnancy and, especially for the MRI-examination, pace maker or other non-compatible implants and claustrophobia



Maiwald 2014 (Continued)			
Patient characteristics and setting		regardless of the stag	the scope of this review, e of CLD. No restrictions
Index tests	Multiphase-CT was performed using 2 different scanners (Brilliance 64/iCT; Philips Healthcare, Eindhoven, Netherlands) with identical prameters.		
	Positivity criteria: diag in the arterial phase ar layed phase.		d on hypervascularisation al-venous phase or de-
Target condition and reference standard(s)	a surrogate of follow-u technique (US, digital s clinical (loss of weight, pecially AFP) was used	ed as the gold standar p (after 6 months) or c subtraction angiograp general state) and pa in unresected lesions	d for diagnosis, whereas omplementary imaging hy) in combination with raclinical parameters (es-
			psy and liver resection.
Flow and timing	No information on inte	rval between index te	st and reference standard.
Comparative			
Notes	No information on Col		
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern



Maiwald 2014 (Continued)		
Is the reference standards likely to correctly classify the target condition?	Yes	
Were the reference standard results interpreted without knowledge of the results of the index tests?	No	
Could the reference standard, its conduct, or its interpretation have introduced bias?	High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?		Low concern
DOMAIN 4: Flow and Timing		
Was there an appropriate interval between index test and reference standard?	Unclear	
Did all patients receive the same reference standard?	No	
Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?	Yes	
Could the patient flow have introduced bias?	High risk	

Mortele 2001

Study characteristics	
Patient Sampling	In this retrospective study from May 1991-January 1999, 235 patients received hepatic transplants at a tertiary centre, and 53 of them who underwent spiral CT were included in the study. 17 of them had HCC.
	Exclusion criteria: 49 children (up to age 16 years) because of low incidence of malignancies in childhood, 104 patients operated on before December 1995 (no resected specimens were stored), 32 patients because no or inadequate CT scanning
Patient characteristics and setting	Patients underwent OLT only
Index tests	CT scans were performed on a CT unit with available helical CT scanning mode (Somatom Plus 4, Siemens, Erlangen, Germany).
Target condition and reference standard(s)	Reference standard: final diagnosis was established at pathology on the explanted liver. Specimens were re-examined in direct correlation with CT results.
Flow and timing	Mean interval between index test and reference standard was 103 days (range 2-367 days).
Comparative	
Notes	No data on Col



Mortele 2001 (Continued)

Methodological quality

Item	Authors' judge- ment	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	No		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?	Yes		
Could the patient flow have introduced bias?		High risk	



Pozzato 1997

Study characteristics			
Patient Sampling	tal and 20 of them h		nined in a general hospi- 1992-November 1995. 20 HCC.
Patient characteristics and setting	All participants und	erwent OLT only as a	reference standard.
Index tests	carried out by a rad		each examination was unaware of the results of gations.
Target condition and reference standard(s)	The final diagnosis v liver.	was established at pa	thology on the explanted
Flow and timing	Mean time interval I 165 days (range 60-		nd reference standard was
Comparative			
Notes	No data on Col		
Methodological quality			
Item	Authors' judge- ment	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	Unclear		
Could the selection of patients have introduced bias?		Unclear risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Unclear		
Could the conduct or interpretation of the index test have introduced bias?		Unclear risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		



Pozzato 1	.997	(Continued)
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Were the reference standard results interpreted without knowledge of the results of the index tests?

Could the reference standard, its conduct, or its interpretation have introduced bias?

Unclear risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?

No

Did all patients receive the same reference standard?

Yes

Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?

Yes

Could the patient flow have introduced bias?

High risk

Sangiovanni 2010

Study characteristics	
Patient Sampling	In this prospective study, 64 patients with 67 de novo liver nodules (55 with a size of 1-2 cm) were consecutively examined by contrast enhanced-US, CT, MRI, and FNB as diagnostic standard.
	Excluded were patients with a pre-existing nodule, poor liver function (ChildePugh C) indicating liver transplantation independently of HCC, or an echo-coarse US pattern of the liver without a well-defined nodule.
Patient characteristics and setting	Participant characteristics and setting match the scope of this review, patients were included regardless of the stage of CLD. No restrictions in HCC lesion size were applied.
Index tests	CT: CT scan was performed with a 64-multidetector row CT (MDCT; Definition Siemens, Erlangen, Germany.
	The typical radiological pattern of HCC was arterial hypervascularisation followed by portal/venous contrast washout of the nodule.
Target condition and reference standard(s)	Reference standard: the diagnostic gold standard was histology through an FNB performed within the nodule and the surrounding liver parenchyma. All participants underwent biopsy of US-detected nodule even if the nodule was not detected by CT.
Flow and timing	No information on interval between index test and reference standard
	Out of 64 included patients, only 55 were analysed.



Sangiovanni 2010 (Continued)

Notes	The authors declare no CoI.		
Methodological quality			
Item	Authors' judge- ment	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			Low concern
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Unclear		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?	No		
Could the patient flow have introduced bias?		High risk	



Serste 2012

Study characteristics			
Patient Sampling	75 consecutive patients with CLD with US-detected 1-2 cm rules underwent, within 1 month, multiphasic CT, MRI, and b of the nodule.		
	Exclusion criteria: 1 agnosis, and under		I due to inconclusive di-
Patient characteristics and setting	Patients with solitary liver tumour 1-2 cm included only.		
Index tests	CT and MRI results were read by two radiologists in consensus (V.B. and M-P.V.) who were blind to biopsy results.		
	Vascular pattern was qualified as "conclusive" for HCC if contrast washout occurred, defined as the presence of hypervascularity during the arterial phase followed up by a hypodense/hypointense appearance in later phases defining washout. Nodules in which an enhancement was found during the arterial phase without washout were qualified as "suspicious."		
Target condition and reference standard(s)	The diagnostic "gold standard," or the reference method for diagnosis, was the results of biopsy.		
	All biopsies were routinely read by 1 pathologist (P.B.), then independently reviewed in a blinded manner by a second pathologist (V.P.) who was unaware of the previous pathological diagnosis and imaging results.		
Flow and timing	Interval between index test and ref. standard was < 1 month.		
	1 patient was withdrawn from the study because the studied not ule with washout on both examinations, without conclusive diag nosis on biopsy, underwent RFA.		
Comparative			
Notes	No data on Col		
Methodological quality			
Item	Authors' judge- ment	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	No		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			



Serste 2012 (Continued)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			Low concern
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?	No		
Could the patient flow have introduced bias?		High risk	

Teefey 2003

Study characteristics	
Patient Sampling	Between August 1996-December 1998, this prospective study examined 37 patients with end-stage liver disease who had been listed for hepatic transplantation. 10 of the patients either died prior to liver transplantation or their names were removed from the transplant list. 2 patients whose names had been on the transplant list for > 2 years were not included in the study because of an inability to obtain follow-up images. The remaining 25 patients formed the study population. 9 patients were confirmed to have HCC.
Patient characteristics and setting	Only participants with end-stage liver disease on waiting list for OLT were included
Index tests	CT studies were performed on Somatom Plus 4; Siemens Medical Systems, Iselin, NJ.



Teefey 2003 (Continued)	The following criteria we	ere used to evaluate HC	CC.	
	 A well-defined, heterogeneously enhancing, and hyperattenuating lesion on arterial phase images or a lesion associated with vascular invasion was rated 1. A well-defined, homogeneously enhancing, and hyperattenuating area on arterial phase images was rated 2. An ill-defined hyperattenuating area on arterial phase images, a heterogeneously enhancing and hypoattenuating lesion on portal-venous phase images, or a homogeneous hypoattenuating lesion on portal-venous phase images was rated 3 or 4. A well-defined lesion measuring near water attenuation value (features typical of a simple cyst) was rated 5 or 6. 			
Target condition and reference standard(s)	The criterion standard for diagnosis of HCC was histologic examination of the resected liver and liver biopsy. 21 patients underwent liver transplantation, and 4 underwent biopsy of the pertinent liver lesion observed at ≥ 1 of the imaging tests, with unclear follow-up). The presence or absence of all lesions identified with ≥ 1 of the imaging tests (CT, MRI, US, or PET) was determined histologically on a lesion-by-lesion basis.			
Flow and timing	The interval between the last imaging study and the liver transplantation in the 21 participants who had a liver transplant ranged from 1-15 months (mean, 5.3 months).			
Comparative				
Notes	No information on Col			
Methodological quality				
Item	Authors' judgement	Risk of bias	Applicability concerns	
DOMAIN 1: Patient Selection				
Was a consecutive or random sample of patients enrolled?	Yes			
Did the study avoid inappropriate exclusions?	No			
Could the selection of patients have introduced bias?		High risk		
Are there concerns that the included patients and setting do not match the review question?			High	
DOMAIN 2: Index Test (All tests)				
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes			
Were positivity criteria clearly defined?	Yes			
Could the conduct or interpretation of the index test have introduced bias?		Low risk		



Teefe	y 2003	(Continued)
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Are there concerns that the index test, its con-
duct, or interpretation differ from the review
question?

Low concern

DOMAIN	3: Re	ference	Standard
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Is the reference standards likely to correctly classify the target condition?

Yes

No

Were the reference standard results interpreted without knowledge of the results of the index tests?

Could the reference standard, its conduct, or its interpretation have introduced bias?

High risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?

No

Did all patients receive the same reference standard?

No

Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?

Yes

Could the patient flow have introduced bias?

High risk

Van Thiel 2004

Study characteristics

Patient Sampling

Individuals with end-stage liver disease due to any cause, who were evaluated and found to be free of an identifiable HCC and who met United Network for Organ Sharing (UNOS) criteria for listing for liver transplantation, were included in the study.

From October 1998 through July 2003, a total of 300 individuals were evaluated and presented to the liver transplant review board at Loyola University Medical Center. Of these, 282 were listed for transplantation. 15 of these cases were identified as having an HCC at the time of listing and 5 were listed because of fulminant hepatic failure. These cases were eliminated from the subsequent analysis leaving a total of 262 listed liver transplant candidates. Of these, 105 (41%) were transplanted with 4 individuals receiving two and 1 individual receiving three transplants. These later cases receiving multiple transplants were eliminated leaving 100 cases for analysis. Of these, 5 cases (5%) were thought to have

developed an HCC while on the waiting list utilising the US findings and were treated with ethanol injections but were found to have no evidence of HCC when the explanted liver was examined pathologically and included in no HCC group.



Van Thiel 2004 (Continued)			
			nts having an HCC at the time of ure and receiving multiple trans-
Patient characteristics and setting	All participants underwe	nt OLT.	
Index tests			fy a hepatic tumour by CT con- m in diameter with early arterial
Target condition and reference standard(s)	Reference standard: path liver	nologist's identificatio	n of a tumour on the explanted
			ne pathologist to identify lesions or the presence of tumour.
Flow and timing	The time from recognitio was 247.2 +/- 53.8 days	on of the hepatic tumo	ur to the date of liver transplant
Comparative			
Notes	No information on Col		
Methodological quality			
Item	Authors' judgement	Risk of bias	Applicability concerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		



Were the reference standard results interpreted without knowledge of the results of the index tests?

No

Could the reference standard, its conduct, or its interpretation have introduced bias?

High risk

Are there concerns that the target condition as defined by the reference standard does not match the question?

High

DOMAIN 4: Flow and Timing

Was there an appropriate interval between index test and reference standard?

No

Did all patients receive the same reference standard?

Yes

Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?

Yes

Could the patient flow have introduced bias?

High risk

Villacastin Ruiz 2016

Study	chara	ıcteı	ristics
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Patient Sampling

From November 2001-December 2011, 323 OLTs were performed on 313 patients at our centre. Our study is based on the retrospective analysis of data from 273 patients (213 men and 60 women), of an average age of 55 years (31–79), who underwent scheduled transplants because of cirrhosis. A total of 273 consecutive patients with 218 hepatocellular carcinoma (HCC) nodules, who underwent imaging and subsequent transplantation, were examined.

Exclusion criteria were as follows: having undergone urgent nonelective transplants; having undergone retransplantation; and absence of cirrhosis.

Patient characteristics and setting

All participants underwent OLT.

Index tests

CT: abdominal examinations were performed using a 2-slice C Siemens (Siemens Medical Solutions, Forchheim, Germany; November 2001-May 2009) and a 64-slice TC Siemens Somaton Sensation 64 (Siemens Medical Solutions; June 2009-December 2011).

Positivity criteria: lesions suggesting HCC were typically characterised by hypervascularity, especially when accompanied by venous-phase washout.

Target condition and reference standard(s)

The pathological analysis of the explant livers provided the reference standard. Correlation of nodules between the image and pathological results was based primarily on location and secondarily on size.



fillacastin Ruiz 2016 (Continued)			
Flow and timing	Time interval between days.	een index test and ref	erence standard was 105
	Out of 273 patients,	253 patients were ar	nalyzed.
Comparative			
Notes	The authors declare	e no Col.	
Methodological quality			
Item	Authors' judge- ment	Risk of bias	Applicability con- cerns
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	Yes		
Could the selection of patients have introduced bias?		Low risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	No		
Could the reference standard, its conduct, or its interpretation have introduced bias?		High risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			,
Was there an appropriate interval between index test and reference standard?	No		



Villacastin	Ruiz 2016	(Continued)
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Did all patients receive the same reference standard?

Yes

Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?

No

Could the patient flow have introduced bias?

High risk

Yu 2011

Study characteristics	
Patient Sampling	The study analysed data from 638 consecutive adult patients with cirrhosis who received liver transplants within 6 months of imaging at a tertiary care institution.
	Exclusion criteria: patients with studies at outside imaging centres were not included in the study. Previously detected lesions diagnosed as HCC that had undergone locoregional treatment, including thermal or chemical ablation and transarterial chemoembolisation (TACE), before imaging were excluded from analysis.
Patient characteristics and setting	All participants underwent OLT.
Index tests	CT examinations were performed on single-slice (HighSpeed CT/i, GE Medical Systems; Picker PQ 6000, Picker International, Cleveland, OH), 4-slice multidetector (LightSpeed QX/I; GE Medical Systems), or 16 to 64-slice multidetector (Sensation 16, Sensation 64, or Definition 64; Siemens Medical Solutions, Erlangen, Germany) helical scanners with a multiphasic protocol.
	Positivity criteria: lesions suspicious for HCC were typically characterised by ≥ 1 of the following features:
	new or rapidly growing nodule
	 nodule with arterial hypervascularity, especially when accompanied by venous phase washout dominant nodule containing fat
	- dominant noduce containing lat
Target condition and reference standard(s)	Imaging reports and serum AFP levels were compared with results from pathology analysis of explants as the reference standard.
	Pathologists were not specifically provided with the imaging reports regarding number and locations of any suspected lesions.
Flow and timing	Mean imaging-transplantation interval was 2.1 months.
Comparative	
Notes	The authors declare no Col.
Methodological quality	
Item	Authors' judgement Risk of bias Applicability con- cerns



Yu 2011 (Continued)

a zori (continuea)			
DOMAIN 1: Patient Selection			
Was a consecutive or random sample of patients enrolled?	Yes		
Did the study avoid inappropriate exclusions?	No		
Could the selection of patients have introduced bias?		High risk	
Are there concerns that the included patients and setting do not match the review question?			High
DOMAIN 2: Index Test (All tests)			
Were the index test results interpreted without knowledge of the results of the reference standard?	Yes		
Were positivity criteria clearly defined?	Yes		
Could the conduct or interpretation of the index test have introduced bias?		Low risk	
Are there concerns that the index test, its conduct, or interpretation differ from the review question?			Low concern
DOMAIN 3: Reference Standard			
Is the reference standards likely to correctly classify the target condition?	Yes		
Were the reference standard results interpreted without knowledge of the results of the index tests?	Yes		
Could the reference standard, its conduct, or its interpretation have introduced bias?		Low risk	
Are there concerns that the target condition as defined by the reference standard does not match the question?			High
DOMAIN 4: Flow and Timing			
Was there an appropriate interval between index test and reference standard?	Yes		
Did all patients receive the same reference standard?	Yes		
Were all patients included in the analysis and analysed according to intention to diagnose principle (non-evaluable results considered as false)?	Yes		
Could the patient flow have introduced bias?		Low risk	

AFP: alpha-foetoprotein; **CLD:** chronic liver disease; **CoI:** conflict of interest; **CT:** computed tomography; **EASL:** European Association for the Study of the Liver; **FNB:** fine-needle biopsy; **HCC:** hepatocellular carcinoma; **HCV:** hepatitis C virus; **IV:** intravenous; **MDCT:** multidetector computed tomography; **MRI:** magnetic resonance imaging; **OLT:** orthotopic liver transplantation; **PET:** positron emission tomography; **RFA:** radiofrequency ablation; **US:** ultrasound



Characteristics of excluded studies [ordered by study ID]

Study	Reason for exclusion
Abdelfattah 2013	No data on per-patient analysis are available as well as no data for CT.
Addley 2011	This is a case-control study.
Alaboudy 2011	The study did not report the 2x2 table and we could not calculate/extract it based on reported data. The study included a group with previously known HCC, and no data exist on per-patient analysis.
Alhasan 2019	The purpose of this study was to evaluate the diagnostic performance of LI-RADS version 2017 major features, ancillary features, and categories on CT for the diagnosis of HCC. There is no data on per-patient analysis.
Amadei 2008	A group of patients with previously known HCC was included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
An 2019	This is a case-control study.
Aqel 2005	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Ascha 2009	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Aubé 2017	No data on per-patient analysis were reported.
Babar 2019	Data were provided only for the accuracy of CT in detecting metastatic liver lesions, and primary liver tumours were grouped as a single entity.
Baron 1996	The study included participants with previously known HCC, and no data on per-patient analysis was provided. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Basha 2018	No data on per-patient analysis were reported.
Beal 2014	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Bhattacharjya 2004	The study patient group included patients with known HCC. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Born 1998	No data on per-patient analysis were reported. The target condition was not HCC, rather focal liver lesions in general.
Brancatelli 2003	No data on per-patient analysis were reported.
Brehmer 2018	No data on per-patient analysis were reported. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Burrel 2003	This is a case-control study. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Camera 1999	The study did not report the 2x2 table and we could not calculate/extract it based on reported data. No data on per-patient analysis were reported.



Study	Reason for exclusion
Chen 1982	The study included patients with previously known HCC. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Chen 2006	The index test was contrast-enhanced US, not CT. Patients with previously known HCC were included.
Choi 2018	Patients with previously known HCC were included. No data on per-patient analysis were reported.
Clevert 2009	The target condition was malignant or benign hepatic tumours in general, no data specific for HCC were present.
Dai 2008	No data on per-patient analysis were reported.
Denies 2002	No data on per-patient analysis were reported.
De Santis 1992	No data on per-patient analysis were reported.
Di Martino 2010	No data on per-patient analysis were reported.
Di Martino 2013	No data on per-patient analysis were reported.
Fasani 1999	Patients with previously known HCC were included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Fausto 2011	No data on per-patient analysis were reported.
Frey 2015	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Fukunaga 2007	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Furlan 2012	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Gaiani 2004	The index test was US not CT. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Garetti 1988	Patients with previously known HCC were included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Gattoni 1993	Patients with previously known HCC were included.
Giangregorio 2009	Patients with previously treated HCC were included.
Giorgio 2004	Patients with previously known were HCC included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Granito 2013	The index test was MRI, not CT. No data on per-patient analysis were reported.
Grat 2018	Patients with previously known HCC were included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.
Gul 2018	The index test was MRI not CT. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.



Study	Reason for exclusion					
Haberman 2013	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Habermann 2002	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Hafeez 2011	The aim of the study was to assess the diagnostic accuracy of triphasic spiral CT in differentiating benign from malignant focal tumoral liver lesions, not HCC specifically. No data on per-patient analysis were reported.					
Hafeez 2020	Patients with previously known HCC were included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Hasinuzzaman 2018	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Hidaka 2013	No data on per-patient analysis were reported.					
Hirakawa 2011	Patients with treated HCC were included. No data on per-patient analysis were reported.					
Hori 2002	No data on per-patient analysis were reported.					
lavarone 2019	No data on per-patient analysis were reported.					
Ichikawa 2002	The study did not report the 2x2 table and we could not calculate/extract it based on reported data. No data on per-patient analysis were reported.					
Ichikawa 2006	Patients with previously known HCC were included.					
Ichikawa 2010	No data on per-patient analysis were reported.					
Ichikawa 2021	No data on per-patient analysis were reported.					
Imbriaco 2017	No data on per-patient analysis were reported.					
Inoue 1994	No data on per-patient analysis were reported.					
Ismail 1990	No data on per-patient analysis were reported.					
Itai 1981	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
lwamura 1982	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Jia 2017	This is a case-control study.					
Jin 2013	Patients with previously known HCC were included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Jin 2016	Patients with previously known HCC were included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Jonczyk 2017	Patients with previously treated HCC were included.					
Jung 2005	The target condition were focal liver lesions in general, not HCC specifically.					



Study	Reason for exclusion					
Jung 2007	The index test was contrast-enhanced US with quantitative evaluation, not CT.					
Kader 2017	The target condition were focal liver lesions in general, not HCC specifically.					
Kakihara 2014	No data on per-patient analysis were reported.					
Kan 2010	The index test was contrast-enhanced US, not CT. No data on per-patient analysis were reported.					
Kanematsu 1997a	The index test was combined CT hepatic arteriography with CT arterial portography, not the type of CT imaging of interest for this review.					
Kanematsu 1997b	No data on per-patient analysis were reported.					
Kato 2004	The index test is MRI, not CT.					
Kawamori 1991	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Kawata 2002	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Khalili 2011	No data on per-patient analysis were reported.					
Kim 2005	The aim of this study was to assess the value of contrast-enhanced sonography for the characterisation of focal hepatic lesions, not HCC specifically.					
Kim 2006	Patients with previously known HCC were included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Kim 2007a	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Kim 2007b	One of the index tests was the combination of CT hepatic arteriography with CT arterial portography, not the type of CT imaging of interest for this review. Patients with previously known HCC were included.					
Kim 2009	This is a case-control study.					
Kim 2017	This is a case-control study.					
Kim 2018	Patients with previously known HCC included.					
Kim 2019	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Kurucay 2017	Index test was CT perfusion imaging, not the type of CT imaging of interest for this review.					
Laroia 2013	The index test was contrast-enhanced US.					
Laroia 2016	The index test was dual-energy CT, not the type of CT imaging of interest for this review.					
Lee 2003	Patients with previously known HCC were included. No data on per-patient analysis were reported.					
Lee 2009	> 5% of patients with previously treated HCC were included.					



Study	Reason for exclusion					
Lee 2012c	Patients with previously known HCC were included.					
Li 2018	Patients with previously known HCC were included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Lin 2011	Patients with previously known HCC were included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Liu 2012	No data on per-patient analysis were reported. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Liu 2016	No data on per-patient analysis were reported.					
Luca 2010	No data on per-patient analysis were reported.					
Lucatelli 2020	No data on per-patient analysis were reported.					
Maciel 2006	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Maetani 2008	Patients with previously known HCC were included. No data on per-patient analysis were reported The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Malagari 1999	The index test was CT portography and post-Lipiodol CT, not the type of CT imaging of interest for this review.					
Manini 2013	No data on per-patient analysis were reported.					
Marcato 1999	The index test was Lipiodol CT, not the type of CT imaging of interest for this review.					
Marin 2009a	No data on per-patient analysis were reported.					
Marin 2009b	No data on per-patient analysis were reported.					
Masuda 2017	No data on per-patient analysis were reported.					
Mehana 2019	No data on per-patient analysis were reported.					
Miller 1991	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Miller 1994	No data on per-patient analysis were reported.					
Min 2020	No data on per-patient analysis were reported.					
Mok 2004	The index test was Lipiodol CT, not the type of CT imaging of interest for this review.					
Moudgil 2017	No data on per-patient analysis were reported. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Nakamura 2000	Patients with previously known HCC were included. No data on per-patient analysis were reported The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					



Study	Reason for exclusion					
Nusbaum 2015	Patients with previously known HCC were included. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Oberstein 1996	Target conditions were different liver diseases, not HCC specifically.					
Paul 2007	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Peterson 2000	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Pocha 2013	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Putz 2019	The index test was contrast-enhanced US for detection of liver lesions, not HCC specifically.					
Ren 2015	No data on per-patient analysis were reported.					
Ronzoni 2007	> 5% of patients with previously treated HCC were included.					
Ryu 2014	Target condition was focal liver masses, not HCC specifically. No data on per-patient analysis were reported.					
Saada 1994	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Sano 2011	Patients with previously known HCC were included.					
Sekoguchi 1994	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Semaan 2020	Patients previously treated for HCC were included.					
Semelka 1992	The study did not report the 2x2 table and we could not calculate/extract it based on reported data. No data on per-patient analysis were reported.					
Seo 2019	Patients with previously known HCC were included.					
Shapiro 1996	Patients with previously known HCC were included.					
Silberhumer 2004	Patients with previously known HCC were included. No data on per-patient analysis were reported.					
Singh 2007	Target conditions were primary liver tumours, not HCC specifically.					
Sofue 2011	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Soyer 1994	The index test was CT arterial portography for detection of primary malignant liver neoplasms, not HCC specifically.					
Suarez-Munoz 2006	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.					
Sugimoto 2015	Patients with previously known HCC were included, No data on per-patient analysis were reported.					
Takayasu 1990	Patients with previously known HCC were included.					



Study	Reason for exclusion				
Tang 2018	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.				
Valls 2004	No data on per-patient analysis were reported. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.				
Wang 1997	No data on per-patient analysis were reported.				
Wang 2007	The study did not report the 2x2 table and we could not calculate/extract it based on reported data.				
Wang 2018	Patients with previously known HCC were included. No data on per-patient analysis were reported.				
Wang 2019	Patients with previously known HCC were included.				
Watanabe 1986	Patients with previously known HCC were included.				
Yamashita 1996	No data on per-patient analysis were reported. The study did not report the 2x2 table and we could not calculate/extract it based on reported data.				
Yim 2016	No data on per-patient analysis was reported, index test was CT hepatic arteriography and portography, not the type of CT of interest for this review.				
Yukisawa 2007	No data on per-patient analysis were reported.				
Zacherl 2002	Patients with previously known HCC included and only analysis per lesion is presented.				
Zhao 2007	No data on per-patient analysis.				

CT: computed tomography; HCC: hepatocellular carcinoma; MRI: magnetic resonance imaging; US: ultrasound

DATA

Presented below are all the data for all of the tests entered into the review.

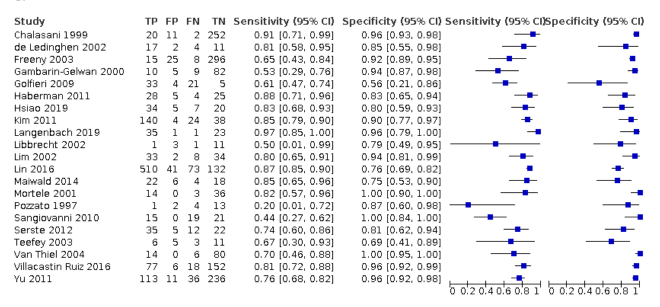
Table Tests. Data tables by test

Test	No. of studies	No. of participants
1 CT	21	3101
2 CT for resectable HCC	10	1854



Test 1. CT

CT



Test 2. CT for resectable HCC

CT for resectable HCC

Study	TP	FP	FN	TN	Sensitivity (95% CI)	Specificity (95% CI)	Sensitivity (95% CI)Specificity (95% CI)
de Ledinghen 2002	17	2	4	11	0.81 [0.58, 0.95]	0.85 [0.55, 0.98]	
Freeny 2003	15	25	8	296	0.65 [0.43, 0.84]	0.92 [0.89, 0.95]	
Gambarin-Gelwan 2000	10	5	9	82	0.53 [0.29, 0.76]	0.94 [0.87, 0.98]	
Libbrecht 2002	1	3	1	11	0.50 [0.01, 0.99]	0.79 [0.49, 0.95]	
Lin 2016	510	41	73	132	0.87 [0.85, 0.90]	0.76 [0.69, 0.82]	
Mortele 2001	14	0	3	36	0.82 [0.57, 0.96]	1.00 [0.90, 1.00]	
Pozzato 1997	1	2	4	13	0.20 [0.01, 0.72]	0.87 [0.60, 0.98]	
San gio vanni 2010	15	0	19	21	0.44 [0.27, 0.62]	1.00 [0.84, 1.00]	→
Serste 2012	35	5	12	22	0.74 [0.60, 0.86]	0.81 [0.62, 0.94]	
Yu 2011	113	11	36	236	0.76 [0.68, 0.82]	0.96 [0.92, 0.98]	0 0 2 0 4 0 6 0 8 1 0 0 2 0 4 0 6 0 8 1

ADDITIONAL TABLES

Table 1. Post-test probabilities

	Likelihood ratio	Post-test probability
if CT positive	8.87 ^a	69%
if CT negative	0.25 ^b	6%
if CT positive	8.87 ^a	91%
if CT negative	0.25 ^b	21%
	if CT negative	if CT positive 8.87a if CT negative 0.25b if CT positive 8.87a



Table 1. Post-test probabilities (Continued)

60%	if CT positive	8.87 ^a	93%		
60%	if CT negative	0.25 ^b	27%		
CT: computed tomography					

^aPositive likelihood ratio.

Table 2. Heterogeneity and sensitivity analyses for computed tomography

Subgroup	No of studies	Sensitivity (95% CI)	Specificity (95% CI)	P value	
All	21	77.5% (70.9% to 82.9%)	91.3% (86.5% to 94.5%)	-	
Positivity criteria clearly defined	19	78.3% (72.0% to 83.6%)	90.7% (85.7% to 94.1%)	-	
Reference standard blinded	9	77.5% (68.8% to 84.3%)	91.0% (83.4% to 95.4%)	-	
Low concern for applic- ability	3	76.9% (50.8% to 91.5%)	89.2% (57.0% to 98.1%)	-	
Before 2005	10	71.4% (60.5% to 80.3%)	93.6% (87.7% to 96.7%)	0.340	
After 2005	11	80.5% (72.3% to 86.7%)	88.7% (81.1% to 93.5%)		
Cirrhosis > 90%	16	75.5% (66.2% to 82.8%)	93.5% (89.0% to 96.2%)	0.225	
Cirrhosis < 90% 4		85.2% (80.8% to 88.7%)	81.5% (73.3% to 87.5%)		
Europe	10	74.3% (59.7% to 85.0%)	90.5% (80.8% to 95.6%)	0.622	
North and South America	7	75.0% (65.7% to 82.4%)	93.7% (87.4% to 96.9%)		
Asia	4	85.5% (81.7% to 88.7%)	85.7% (75.1% to 92.3%)		

^bNegative likelihood ratio.



Table 2. Heterogeneity and sensitivity analyses for computed tomography (Continued)

HCC prevalence ≥ 52%	11	81.0% (72.5% to 87.4%)	85.5% (78.4% to 90.5%)	0.051
HCC prevalence < 52%	10	71.1% (60.7% to 79.7%)	94.0% (89.8% to 96.5%)	
Clinically suspect	19	78.5% (72.7% to 83.3%)	90.2% (84.9% to 93.8%)	0.333
Surveillance	2	72.3% (29.4% to 94.2%)	97.3% (88.7% to 99.4%)	
HCC resectable 100%	10	71.4% (60.3% to 80.4%)	92.0% (86.3% to 95.5%)	
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HCC resectable < 20%	2	75.6% (55.0% to 88.8%)	78.3% (42.6% to 94.6%)	0.116
HCC resectable ≥ 20% 14		73.3% (65.0% to 80.2%)	93.2% (87.8% to 96.3%)	
HCC resectable < 90%	4	76.2% (63.2% to 85.6%)	82.4% (61.9% to 93.1%)	0.081
HCC resectable ≥ 90%	12	72.5% (63.2% to 80.1%)	93.9% (88.7% to 96.9%)	
Biopsy	8	82.9% (69.7% - 91.2%)	90.8% (82.8% - 95.3%)	0.119
OLT	11	78.8% (73.6% - 83.2%)	93.2% (88.7% - 96.0%)	
Mixed	2	61.9% (49.4% - 73.0%)	64.0% (44.0% - 80.1%)	
Viral < 80% ^a	15	81.1% (75.5% to 85.7%)	92.4% (87.5% to 95.4%)	0.332
Viral≥80% ^a	3	62.8% (44.6% to 78.0%)	93.9% (48.2% to 99.6%)	
Prior detection of nod- ules: no	13	79.3% (74.4% to 83.6%)	92.5% (87.6% to 95.6%)	0.797
Prior detection of nod- ules: yes	8	80.9% (67.5% to 89.6%)	88.6% (78.9% to 94.2%)	
CI: confidence interval; I	ICC: hepatocel	lular carcinoma; OLT: orthotopic liver	rtransplantation;	

 $^{\it a}$ Data not reported in three studies.



Table 3. Other systematic reviews on diagnostic accuracy of computed tomography for hepatocellular carcinoma

Systemati- creview	Analysistype	No of in- cluded studies	No of pa- tients analysed	Sensitivity (%) (95% CI)	Specificity (%) (95% CI)	Statistical model
Colli 2006	Per-patient	10	979	67.5 (55 to 80)	92.5 (89 to 96)	Random-ef- fects model
Floriani 2013	Per-patient	2	Not reported	61 (29 to 93)	72 (50 to 94)	Bivariate random-ef- fects model
Chou 2015	Per-patient	17	Not reported	83 (76 to 88)	91 (84 to 95)	Bivariate logistic mixed ran- dom-ef- fects model
Xie 2011	Per-lesion	8	1134	86 (84 to 88)	82 (77 to 86)	
Chen 2013	Per-lesion	15	Not reported	81 (74 to 86)	93 (88 to 96)	_
Floriani 2013	Per-lesion	10	Not reported	68 (56 to 79)	64 (21 to 100)	_
Chou 2015	Per-lesion	80	Not reported	76 (72 to 80)	89 (84 to 93)	_
Ye 2015	Per-lesion	9	469	74 (70 to 77)	93 (91 to 94)	_
Lee 2015	Per-lesion	17	1135	72 (75 to 84)	Not estimated	_
Hanna 2016	Per-lesion	105	Not reported	73.6 (70 to 76)	Not estimated	_
Guo 2016	Per-lesion	12	627	70 (58 to 80)	94 (92 to 96)	_
Roberts 2018	Per-lesion	33	2250	66 (60 to 72)	92 (84 to 96)	_
Li 2019	Per-lesion	8	498	68 (51 to 81)	92 (84 to 96)	_

APPENDICES

Appendix 1. Search strategies

Database Time span Search s		Search strategy	
The Cochrane Hepa- to-Biliary Group Con- trolled Trials Register	May 2021	(computed tomograph* or CT or CECT or MDCT or MSCT or magnetic resonance imaging or MRI or emission tomography or PET) AND (((liver or hepato*) and (carcinom* or cancer* or neoplasm* or malign* or tumo*)) or HCC) AND (advanc* and chronic and (liver* or hepat*))	
The Cochrane Hepa- to-Biliary Group Diag-	May 2021	(computed tomograph* or CT or CECT or MDCT or MSCT or magnetic resonance imaging or MRI or emission tomography or PET) AND (((liver or hepato*)	



(Continued) nostic Test of Accuracy Studies Register		and (carcinom* or cancer* or neoplasm* or malign* or tumo*)) or HCC) AND (advanc* and chronic and (liver* or hepat*))
The Cochrane Library	2021, Issue 5	#1 MeSH descriptor: [Tomography, Emission-Computed] explode all trees
		#2 MeSH descriptor: [Tomography, X-Ray Computed] explode all trees
		#3 MeSH descriptor: [Magnetic Resonance Imaging] explode all trees
		#4 (computed tomograph* or CT or CECT or MDCT or MSCT or magnetic resonance imaging or MRI or emission tomography or PET)
		#5 #1 or #2 or #3 or #4
		#6 MeSH descriptor: [Carcinoma, Hepatocellular] explode all trees
		#7 MeSH descriptor: [Liver Neoplasms] explode all trees
		#8 (((liver or hepato*) and (carcinom* or cancer* or neoplasm* or malign* or tumo*)) or HCC)
		#9 #6 or #7 or #8
		#10 (advanc* and chronic and (liver* or hepat*))
		#11 #5 and #9 and #10
MEDLINE Ovid	1946 to May 2021	1. exp Tomography, Emission-Computed/
		2. exp Tomography, X-Ray Computed/
		3. exp Magnetic Resonance Imaging/
		4. (computed tomograph* or CT or CECT or MDCT or MSCT or magnetic resonance imaging or MRI or emission tomography or PET).mp. [mp=title, abstract, original title, name of substance word, subject heading word, floating subheading word, keyword heading word, protocol supplementary concept word, rare disease supplementary concept word, unique identifier, synonyms]
		5. 1 or 2 or 3 or 4
		6. exp Carcinoma, Hepatocellular/
		7. exp Liver Neoplasms/
		8. (((liver or hepato*) and (carcinom* or cancer* or neoplasm* or malign* or tumo*)) or HCC).mp. [mp=title, abstract, original title, name of substance word, subject heading word, floating sub-heading word, keyword heading word, protocol supplementary concept word, rare disease supplementary concept word, unique identifier, synonyms]
		9. 6 or 7 or 8
		10. (advanc* and chronic and (liver* or hepat*)).mp. [mp=title, abstract, original title, name of substance word, subject heading word, floating sub-heading word, keyword heading word, protocol supplementary concept word, rare disease supplementary concept word, unique identifier, synonyms]
		11. 5 and 9 and 10
Embase Ovid	1974 to May 2021	1. exp computer assisted tomography/
		2. exp positron emission tomography/
		3. exp nuclear magnetic resonance imaging/



(Continued)				
		4. (computed tomograph* or CT or CECT or MDCT or MSCT or magnetic resonance imaging or MRI or emission tomography or PET).mp. [mp=title, abstract, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword, floating subheading word, candidate term word]		
		5. 1 or 2 or 3 or 4		
		6. exp liver cell carcinoma/		
		7. exp liver tumor/		
		8. (((liver or hepato*) and (carcinom* or cancer* or neoplasm* or malign* or tumo*)) or HCC).mp. [mp=title, abstract, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword, floating subheading word, candidate term word]		
		9. 6 or 7 or 8		
		10. (advanc* and chronic and (liver* or hepat*)).mp. [mp=title, abstract, heading word, drug trade name, original title, device manufacturer, drug manufacturer, device trade name, keyword, floating subheading word, candidate term word]		
		11. 5 and 9 and 10		
LILACS (Bireme)	1982 to May 2021	(computed tomograph\$ or CT or CECT or MDCT or MSCT or magnetic resonance imaging or MRI or emission tomography or PET) [Words] and (((liver or hepato\$) and (carcinom\$ or cancer\$ or neoplasm\$ or malign\$ or tumo\$)) or HCC) [Words] and (advanc\$ and chronic and (liver\$ or hepat\$)) [Words]		
Science Citation In-	1900 to May 2021	#4 #3 AND #2 AND #1		
dex Expanded (Web of Science)		#3 TS=(advanc* and chronic and (liver* or hepat*))		
,		#2 TS=(((liver or hepato*) and (carcinom* or cancer* or neoplasm* or malign* or tumo*)) or HCC)		
		#1 TS=(computed tomograph* or CT or CECT or MDCT or MSCT or magnetic resonance imaging or MRI or emission tomography or PET)		
Conference Proceed-	1990 to May 2021	#4 #3 AND #2 AND #1		
ings Citation Index – Science (Web of Science)		#3 TS=(advanc* and chronic and (liver* or hepat*))		
		#2 TS=(((liver or hepato*) and (carcinom* or cancer* or neoplasm* or malign* or tumo*)) or HCC)		
		#1 TS=(computed tomograph* or CT or CECT or MDCT or MSCT or magnetic resonance imaging or MRI or emission tomography or PET)		

Appendix 2. QUADAS 2

Domain	1. Patient selection	2. Index test	3. Reference standard	4. Flow and timing
Signalling ques- tions and crite- ria	Q1: "Was a consecutive or random sample of participants enrolled?"	Q1: "Were the index test results interpreted with- out knowledge of the	Q1: "Is the reference standard likely to correctly classify the target condition?"	Q1: "Was there an appro- priate interval between the index test and the ref- erence standard?"



(Continued)

Yes - If the study reports on a consecutive or a random selection of patients.

No - if the study reports on another form of selection of patients.

Unclear - if the study does not report on how the patients were enrolled.

Q.2: "Did the study avoid inappropriate exclusions?"

Yes - if definitions of exclusion criteria are appropriate (i.e. previous surgery or treatment for hepatocellular carcinoma; patients with cholangiocarcinoma) and all exclusions are reported.

No - if exclusion criteria are inappropriate and exclusions are not reported.

Unclear - if the study does not report causes of exclusions.

results of the reference standard?"

Yes - if the study reports that the results of the index test were interpreted without the knowledge of the results of the reference standard.

No - if the study reports that results of the index test were interpreted with the results of the reference standard.

Unclear - if the study does not report information about blinding of the results of the index test and reference standard.

Q2: "Were positivity criteria clearly defined?"

Yes - if the study clearly reports positivity criteria (i.e. the minimum diameter of a detectable lesion, exclusion of benign criteria).

No - if the study does not report the positivity criteria.

Unclear - if the study does not report information about the definition of positivity criteria Yes - if the reference standard correctly defines the presence/absence of HCC (pathology of explanted liver in a transplant cohort).

No - if other reference tests than pathology of explanted liver were used.

Unclear - if the study does not report on the reference standard used.

Q2: "Were the reference standard results interpreted without the knowledge of the results of the index test?"

Yes - if the study reports that the results of the reference standard were interpreted without the knowledge of the results of the index test.

No - if the study reports that the results of the reference standard were interpreted with the knowledge of the results of the index test.

Unclear - if the study does not report information about blinding of the results of the reference standard and the index test. Yes - if the interval between the index test and the reference standard was less than 3 months.

No - if the interval was longer than 3 months.

Unclear - if the study does not report the interval between the index test and the reference standard.

Q2: "Did all participants receive the same reference standard?"

Yes - if the study has only one reference standard for all the participants

No - if the study has more than one reference standard.(histology of resected focal liver lesion(s), or the histology of focal liver lesion(s) with a follow-up period of at least six months in the participants with a negative result of the index test)

Unclear - if the study information regarding the use of reference standard are unclear

Q3: "Were all participants included in the analysis and analysed according to intention to diagnose principle (uninterpretable results considered as false)?"

Yes - if all enrolled participants were included in the analysis and uninterpretable index test results were analysed according to the intention to diagnose principle).

No - if any participant was excluded from the analysis for any reason or uninterpretable index test results were not analysed according to intention to diagnose principle.



(Continued)

Unclear - if the exclusion of participants from the analysis is unclear.

Risk of bias

Could the selection of patients have introduced bias?

Low risk: "Yes" for all signalling questions.

High risk: "No" for at least one signalling question.

Could the conduct or interpretation of the index test have introduced bias?

Low risk: "Yes" for all signalling questions.

High risk: "No" for at least one signalling question or "Unclear" for the Q2.

Could the reference standard, its conduct, or its interpretation have introduced bias?

Low risk: "Yes" for all signalling questions.

High risk: "No" for at least one signalling question or "Unclear" for the Q 2.

Could the participant flow have introduced bias?

Low risk: "Yes" for all signalling questions.

High risk: "No" for at least one signalling question.

Concerns about applicability

Are there concerns that included participants and setting do not match the review question?

Low concern: the participants included in the review represent the participants in whom the tests is used in clinical practice (i.e. surveillance programme in patients with chronic advanced liver disease; clinical cohort of patients with chronic advanced liver disease).

High concern: the participants included in the review differ from the participants in whom the tests is used in clinical practice

(cohort of patients with advanced and decompensated liver disease, candidates for orthotopic liver transplantation). Are there concerns that the index test, its conduct, or interpretation differ from the review question?

Low concern: the index test, its conduct or its interpretation does not differ from the way it is used in clinical practice.

High concern: the index test, its conduct or its interpretation differs from the way it is used in clinical practice. Are there concerns that the target condition as defined by the reference standard does not match the question?

High concern: the definition of the target condition as defined by the reference standard does not match the question (i.e. pathology of the explanted liver is feasible only in the case of liver transplant; the natural history and prognosis of HCC detected in explanted liver might be different).

Low concern: the definition of the target condition as defined by the reference standard does match the question for all included patients.

HISTORY

Protocol first published: Issue 6, 2019

CONTRIBUTIONS OF AUTHORS

TN wrote the protocol and performed searches for references, evaluated references for obtaining the full reports, evaluated studies for inclusion, extracted data from studies, assessed the risk of bias, and wrote the final review.

VG commented on the protocol and performed searches for references, evaluated references for obtaining the full reports, evaluated studies for inclusion, extracted data from studies, assessed the risk of bias, and wrote the final review.



AC co-ordinated the protocol design and designed and wrote the final review.

MF performed searches for references and critically commented on the review.

GC wrote the protocol, provided statistical expert opinion and critically commented on the final review.

DM commented on the protocol and will critically comment on the final review.

DŠ critically commented on the protocol, will act as arbiter if review authors cannot reach a consensus, and critically commented on the final review.

All authors approved the publication of the review.

DECLARATIONS OF INTEREST

TN: none known VG: none known AC: none known MF: none known GC: none known DM: none known DŠ: none known

SOURCES OF SUPPORT

Internal sources

· None, Other

External sources

· None, Other

DIFFERENCES BETWEEN PROTOCOL AND REVIEW

We did not use Covidence to manage the selection of studies (Covidence).

We added an additional potential source of heterogeneity: prior testing with detection of liver nodules. We recognised that the retesting was different in the included studies, and some studies included participants with previously detected liver nodules.

We performed a sensitivity analysis considering only the studies at low concern for applicability.

INDEX TERMS

Medical Subject Headings (MeSH)

*Carcinoma, Hepatocellular [diagnostic imaging]; Cross-Sectional Studies; *Liver Neoplasms [diagnostic imaging]; Sensitivity and Specificity; Tomography, X-Ray Computed; Ultrasonography

MeSH check words

Adult; Humans