Artificial intelligence in the detection, characterisation and prediction of hepatocellular carcinoma: a narrative review

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Abstract: Hepatocellular carcinoma (HCC) is a significant cause of morbidity and mortality worldwide. Despite significant advancements in detection and treatment of HCC, its management remains a challenge. Artificial intelligence (AI) has played a role in medicine for several decades, however, clinically applicable AI-driven solutions have only started to emerge, due to gradual improvement in sensitivity and specificity of AI, and implementation of convoluted neural networks. A review of the existing literature has been conducted to determine the role of AI in HCC, and three main domains were identified in the search: detection, characterisation and prediction. Implementation of AI models into detection of HCC has immense potential, as AI excels at analysis and integration of large datasets. The use of biomarkers, with the rise of ‘omics’, can revolutionise the detection of HCC. Tumour characterisation (differentiation between benign masses, HCC, and other malignant tumours, as well as staging and grading) using AI was shown to be superior to classical statistical methods, based on radiological and pathological images. Finally, AI solutions for predicting treatment outcomes and survival emerged in recent years with the potential to shape future HCC guidelines. These AI algorithms based on a combination of clinical data and imaging-extracted features can also support clinical decision making, especially treatment choice. However, AI research on HCC has several limitations, hindering its clinical adoption; small sample size, single-centre data collection, lack of collaboration and transparency, lack of external validation, and model overfitting all results in low generalisability of the results that currently exist. AI has potential to revolutionise detection, characterisation and prediction of HCC, however, for AI solutions to reach widespread clinical adoption, interdisciplinary collaboration is needed, to foster an environment in which AI solutions can be further improved, validated and included in treatment algorithms. In conclusion, AI has a multifaceted role in HCC across all aspects of the disease and its importance can increase in the near future, as more sophisticated technologies emerge.

Keywords: Hepatocellular carcinoma (HCC); artificial intelligence (AI); machine learning (ML); computer-aided diagnosis; neural networks

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Introduction

Hepatocellular carcinoma (HCC) is the most common adult primary liver malignancy causing the fourth most cancer-related deaths worldwide (1). The highest burden of HCC is found in East-Asia and Sub-Saharan Africa, however, it is also an emerging major cause of morbidity and mortality globally (2). There were significant improvements in our understanding of the aetiology and pathogenesis of HCC throughout the years, which have contributed to novel surveillance, diagnosis and management strategies. However, despite these advancements, HCC still poses challenges for clinicians worldwide. HCC is characterised by presenting at advanced stages, often on a background of pre-existing liver disease, which poses diagnostic challenges. These challenges are both histological, due to heterogeneity of HCC and continuum of its malignant progression and radiological due to HCC mimickers such as i-CCA or c-CCA and prevalence of atypical radiological features (approx. 40%) (3,4). Moreover, extensive background liver disease can impede potentially curative therapies (5). Due to heterogeneity in pathogenesis and multiple recognised risk factors, the biological behaviour of HCC is mostly unknown, which translates into discrepancies in staging systems and prognosis (6). Moreover, HCC has a high recurrence rate following surgical resection, and even orthotopic liver transplantation has a five year survival of only 65–81% despite using specific criteria (Milan, USFC, Kyoto) aimed at selecting patients thought to have better long-term outcomes (6,7).

Development of artificial intelligence (AI) provides a unique set of novel tools to aimed at solving the aforementioned issues and improve HCC detection, characterisation, prediction of survival and treatment outcomes. Although AI has been applied in medicine for over 60 years (8), only the recent research has provided a plethora of successful studies with potential for clinical impact, including classification of skin cancers (9), breast cancer screening (10) or retinal examination (11). AI-driven solutions can help in early detection of HCC, more accurate diagnosis and classification of the tumour, as well as predicting disease course and outcomes. In order to understand the potential future role of AI in various aspects of HCC management, and in-depth review of currently available evidence of the role of AI in HCC was conducted. By providing an overview of the strengths and limitations of AI in HCC, the authors aimed to understand the factors limiting widespread clinical adoption of AI-driven solutions and provide recommendations on future AI research.

This review explores AI solutions applied to HCC that can be classified into three main domains: detection, characterisation and prediction. In the context of this review, detection encompasses technologies that highlight an abnormality, both in imaging and clinical data, which allow for further follow-up. Characterisation includes techniques that differentiate between various hepatic abnormalities, as well as, stratify and stage previously diagnosed HCC. Prediction comprises methods, which use AI for evaluating long-term outcomes including overall survival, tumour progression or assessment of response to treatment. We present the following article in accordance with the narrative review reporting checklist (available at http://dx.doi.org/10.21037/tgh-20-242).

Definitions

In order to discuss AI in context of HCC, its definition needs to be established. AI describes the use of computers and related technology to emulate intelligent behaviour and critical thinking of human beings (12). Within AI, one of the most commonly used terms is machine learning (ML), which can be defined as a discipline in which machines (computers) learn from data, with emphasis on computational algorithms, which are able to analyse billions of data points (13). Of the most commonly used AI techniques are artificial neural networks (ANN); statistical systems, which mimic the complex architecture of biological networks of neurons, to derive outputs, based on interactions of weighted inputs and outputs (14). Convolutional neural networks (CNN) can be especially useful in the medical context, as they have the ability to process data with a grid pattern (e.g., radiological images) using multiple layers, including convolution and pooling layers performing feature extraction to produce final output (15). Additional terms related to AI that were used throughout this review are defined in Table 1.

Methods

A comprehensive search of Embase, MEDLINE and Cochrane Library databases was conducted. The databases were searched from their conception to 14th March 2020. The search was conducted using MeSH terms and keywords for hepatocellular carcinoma (hepatocellular carcinoma, HCC) artificial intelligence (AI, artificial intelligence, machine intelligence), machine learning (machine learning,
ML, deep learning, supervised learning, unsupervised learning) artificial neural network (artificial neural networks, neural network, ANN, NN) and convolutional neural network (convolutional neural network, CNN) combined with Boolean operators. Relevant keywords were identified using recent related publications on AI in HCC (20,21).

In total, 702 records were identified, and after removal of duplicates, 470 records have remained. The titles and abstracts were screened by two authors independently (MK and AD), and a third author resolved any conflicts (TMHG). Inclusion criteria included: original research studies, English language, use of AI-driven solutions and HCC as primary pathology of interest. Exclusion criteria included: malignancies other than HCC as primary pathology of interest, age <18, non-original research studies (e.g., commentaries, letters to editors, reviews). Relevant original research studies were included in the analysis, however, conference abstracts and review articles were also screened. Manual screening of reference lists of included full texts was also performed by two authors (MK and AD) independently to look for any missing studies. After the list of accepted texts was finalised, full texts were classified into one of three domains that have been described in the introduction (detection, characterisation and prediction). Following classification into domains, data extraction was conducted; variables of interest were the number of participants/pathological slides/radiological images, AI algorithm used, type of validation, AUC (or if not available, alternative diagnostic accuracy measures such as sensitivity and specificity, c-index or F-score).

### Detection

The summary of literature on HCC detection, based on pre-HCC disease models, imaging and biomarkers is presented in Table 2.

### Pre-HCC disease models

HCC pathogenesis is strongly linked to chronic inflammatory disease of the liver, which allows for HCC detection based on the range of pre-malignant changes.

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**Table 1** Glossary of terms related to the use of artificial intelligence in medicine and performance of AI-driven solutions

<table>
<thead>
<tr>
<th>Term</th>
<th>Definition</th>
</tr>
</thead>
<tbody>
<tr>
<td>Artificial intelligence (AI)</td>
<td>The use of computers and related technology to emulate intelligent behaviour and critical thinking of human beings (12)</td>
</tr>
<tr>
<td>Machine learning (ML)</td>
<td>A discipline in which machines (computers) learn from data, with emphasis on computational algorithms, which are able to analyse billions of data points (13)</td>
</tr>
<tr>
<td>Supervised learning</td>
<td>Type of ML, which deals with predicting a known outcome, based on inputs, in the presence of an expert ‘supervisor’ (16)</td>
</tr>
<tr>
<td>Unsupervised learning</td>
<td>Type of ML, which deals with finding naturally occurring patterns without a pre-defined outcome, without the presence of an expert ‘supervisor’ (16)</td>
</tr>
<tr>
<td>Artificial neural networks (ANN)</td>
<td>Statistical systems, which mimic the complex architecture of biological networks of neurons, to derive outputs, based on interactions of weighted inputs and outputs (14)</td>
</tr>
<tr>
<td>Convolutional neural network (CNN)</td>
<td>Type of deep learning ANN, for processing data with grid pattern (e.g., radiological images) using multiple layers, including convolution and pooling layers performing feature extraction to produce final output (15)</td>
</tr>
<tr>
<td>Deep learning</td>
<td>A subset of ML, that uses representation learning (automatic discovery of representations from raw data for classification or detection) (17)</td>
</tr>
<tr>
<td>Area under the curve (AUC)</td>
<td>Algorithm performance measure, which can be established based on receiver operator characteristics (ROC) curve. AUC takes values between 0 and 1, depending on average sensitivity and specificity for all analysed values of the, with values approaching 1 indicating higher performance (18)</td>
</tr>
<tr>
<td>Accuracy</td>
<td>Algorithm performance measure, taking values between 0% and 100%, based on the number of true positive and true negative results, compared to the overall size of the population (18)</td>
</tr>
<tr>
<td>C-index (c-statistic)</td>
<td>Algorithm performance measure, describing the goodness of fit of the model, taking values between 0 and 1 (19)</td>
</tr>
</tbody>
</table>
Table 2 Summary of literature on HCC detection, based on pre-HCC disease models, imaging and biomarkers

<table>
<thead>
<tr>
<th>Domain</th>
<th>Sub-category</th>
<th>Notes</th>
<th>N</th>
<th>AI algorithm</th>
<th>Type of validation</th>
<th>AUC (95% CI)</th>
<th>Limitations</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-HCC disease models</td>
<td>NAFLD/NASH</td>
<td>Serum and liver lipids in murine models</td>
<td>15 (5 intervention, 10 controls)</td>
<td>Random forest</td>
<td>Development only (no validation)</td>
<td>N/A</td>
<td>Animal (murine) model not replicated on humans</td>
<td>Chiappini et al. 2016 (22)</td>
</tr>
<tr>
<td></td>
<td>Population screening for NAFLD</td>
<td>500 (146 case, 354 controls)</td>
<td>Logistic regression</td>
<td>Internal validation (cross-validation)</td>
<td>0.87 (0.83–0.90)</td>
<td>Retrospective study design and lack of external validation</td>
<td>Yip et al. 2017 (23)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Cirrhosis/fibrosis/</td>
<td>HCC</td>
<td>442 patients</td>
<td>Random forest</td>
<td>Internal validation</td>
<td>c-index 0.64 (0.60–0.90)</td>
<td>Study performed in a tertiary centre. Attrition bias. Low clinical adaptability potential</td>
<td>Singal et al. 2012 (24)</td>
</tr>
<tr>
<td></td>
<td>hepatitis B/hepatitis C</td>
<td>Progression of chronic HepC infection into fibrosis</td>
<td>533 (349 normal, 184 fibrosis)</td>
<td>Random forest</td>
<td>Internal validation</td>
<td>0.84 (0.82–0.86)</td>
<td>Narrow enrolment criteria, reducing generalisability of conclusions</td>
<td>Konerman et al. 2015 (25)</td>
</tr>
<tr>
<td></td>
<td>Progression of HepB/C into</td>
<td>6,561 (Reddy et al.; 146,218 (Ioannou et al.)</td>
<td>ANN</td>
<td>Internal validation (random sample split)</td>
<td>0.911–0.962 (range, Reddy et al.); 0.89 (Ioannou et al.)</td>
<td>Retrospective and cross-sectional character of the studies and lack of external validation</td>
<td>Reddy et al. 2017 (26); Ioannou et al. 2019 (27)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>HCC</td>
<td>HCC development based on viral status and clinical data</td>
<td>165 patients</td>
<td>Support vector machine</td>
<td>Internal validation (cross-validation)</td>
<td>0.88</td>
<td>Small sample size</td>
<td>Kissapa et al. 2019 (28)</td>
</tr>
<tr>
<td>Imaging</td>
<td>CT</td>
<td>HCC detection on multi-phase CT scans</td>
<td>25 (Lee et al.; 21 (Okumura et al.)</td>
<td>Temporal subtraction, 3D global matching</td>
<td>Development only (no validation)</td>
<td>N/A</td>
<td>Small sample size, proof-of-concept character of both studies</td>
<td>Lee et al. 2015 (29); Okumura et al. 2011 (30)</td>
</tr>
<tr>
<td></td>
<td>MRI</td>
<td>HCC nodule detection using SPIO-MRI in rat models</td>
<td>40 images</td>
<td>ANN</td>
<td>Development only (no validation)</td>
<td>Classification accuracy 91.76%</td>
<td>Animal (rat) models used. No test-retest reproducibility</td>
<td>Guo et al. 2009 (31)</td>
</tr>
<tr>
<td>Biomarkers</td>
<td>N/A</td>
<td>Serum</td>
<td>167 (Poon et al.; 1,582 (Sato et al.)</td>
<td>Multiple techniques combined</td>
<td>Development only (no validation)</td>
<td>Accuracy 75.9% (Poon et al.); 0.844–0.940 (range, Sato et al.)</td>
<td>Single centre, internally validated studies with small sample size</td>
<td>Poon et al. 2001 (32); Sato et al. 2019 (33)</td>
</tr>
<tr>
<td></td>
<td>Transcriptome</td>
<td>3,981 (2,316 HCC, 1,665 non-tumorous tissue)</td>
<td>Multiple techniques combined</td>
<td>External validation</td>
<td>0.91–0.96 (range)</td>
<td>Heterogeneity of data due to pooled analysis of 30 studies</td>
<td>Kaur et al. 2020 (34)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Gene co-expression</td>
<td>57 (38 HCC, 19 normal samples)</td>
<td>PCA</td>
<td>Development only (no validation)</td>
<td>N/A</td>
<td>Use of retrospective databases and no clinical applicability</td>
<td>Zhang et al. 2017 (35)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>mRNA</td>
<td>95 (43 tumour, 52 non-tumour)</td>
<td>Deep belief nets</td>
<td>Internal validation (cross-validation)</td>
<td>F-score 75.48%</td>
<td>Lack of external validation</td>
<td>Ibrahim et al. 2014 (36)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Urine</td>
<td>15 samples</td>
<td>PCA, random forest</td>
<td>Internal validation</td>
<td>0.903</td>
<td>Small sample size</td>
<td>Liang et al. 2016 (38)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Biomarker identification from literature</td>
<td>N/A</td>
<td>Data mining</td>
<td>Internal validation (cross-validation)</td>
<td>F-score 0.89</td>
<td>Use of impact factor as scoring tool, introducing systematic bias into the results</td>
<td>Chang et al. 2017 (39)</td>
<td></td>
</tr>
</tbody>
</table>

HCC, hepatocellular carcinoma; AI, artificial intelligence; AUC, area under the curve; 95% CI, 95% confidence intervals; NAFLD, non-alcoholic fatty liver disease; NASH, non-alcoholic steatohepatitis; HepB, viral hepatitis type B; HepC, viral hepatitis type C; ANN, artificial neural network; PCA, principal component analysis; miRNA, microRNA; N/A, non-applicable; CT, computed tomography; SPIO-MRI, superparamagnetic iron oxide magnetic resonance imaging.
Chronic viral hepatitis B and C infections are both associated with the development of HCC (40,41). Similarly, non-viral causes, including non-alcoholic fatty liver disease (NAFLD), non-alcoholic steatohepatitis (NASH), cirrhosis and fibrosis of the liver are also risk factors for developing HCC (42). The stepwise progression of these pathologies creates the potential for a screening window, during which high-risk individuals can be identified.

The use of AI-driven solutions in detection of NAFLD and NASH has not been comprehensively researched. In a 2016 study, Chiappini and colleagues investigated serum and liver lipids in NAFLD and NASH murine models, using supervised ML (random forest analysis). They identified unique signatures of NASH, opening new possibilities of pre-HCC change detection (22). Population screening for NAFLD has also been suggested, with ML-algorithm, based on 23 clinical parameters, achieving an AUC of 0.88 (95% CI, 0.84–0.91) in detecting NAFLD (23).

Substantially more research has been made into the association of chronic HepB and HepC infections and development of cirrhosis, fibrosis and eventually HCC. Progression of cirrhosis into HCC was studied as early as 2012; it was found that a supervised ML (random forest) model outperformed conventional regression analysis, achieving a c-index of 0.64 (95% CI, 0.6–0.69) (24). Konerman and colleagues used a very similar model to predict progression of chronic hepatitis C infection into fibrosis, with AUC of 0.86 (95% CI, 0.85–0.87) (25).

Throughout the years, as ML models have become more sophisticated, their diagnostic performance has improved. An ANN model by Reddy and Imler, achieved an AUC as high as 0.962 in the prediction of malignant transformation from hepatitis B or C chronic infection (26). Similarly, another recurrent neural network model achieved an AUC of 0.89 (27). It is worth highlighting that in all cases, ML models have proven to be superior to the classical statistical regression model when analysing big data sets.

Moreover, most recent model by Książek et al., used patient characteristics, such as viral status, presence of comorbidities and laboratory results to predict the development of HCC based on 23 quantitative and 26 qualitative features, has achieved 88.5% accuracy using this approach (28).

### Imaging

Detection is the most basic way of utilising AI in imaging, while more novel approaches focus on characterisation and stratification of suspicious imaging regions. While more advanced applications of AI in imaging will be discussed later in this review, it is important to consider the detection of HCC lesions on imaging, which were the precursors to current AI applications.

Lee et al. used computer-aided diagnosis (CAD) on multi-phase CT scans to detect HCC in a set of 15 moderate HCC (mean ø 3.1 cm) and 10 small HCC (mean ø 1.04 cm). Using a non-rigid registration model, which accounted for deformation between phases due to respiratory movements and heartbeats, they have achieved 100% detection accuracy, when compared with radiological diagnosis (29). Similar results were obtained using 3D non-linear image wrapping (30). Moreover, in 2009 Guo et al. have looked into HCC nodule detection on rat models, using SPIO-enhanced MRI, achieving 91.67% classification accuracy (31).

### Biomarkers

A variety of biomarkers have been researched, to equip clinicians with a reliable tool for HCC detection. Recently advances in bioinformatics and technology, resulting in have revolutionised the way large biological datasets (-omics’ datasets) can be generated and analysed, allowing for the integration of multiple datasets (genome, proteome, transcriptome, etc.) (43). Combination of ‘omics with AI algorithms has led to the identification of suitable biomarkers with the potential to translate data into therapeutics (44). The use of biomarkers, identify with the aid of neural networks, combined with the classically used serological marker for HCC (alpha-fetoprotein), was shown to increase diagnostic sensitivity from 60% to 73.8% while maintaining the sensitivity of 88.2% (32).

Harnessing the power of ML has also allowed for new types of clinical data to be analysed in the hope of detecting HCC. Large-scale (2,316 HCC tumour samples and 1,665 non-tumorous tissue samples) transcriptomic profiling by Kaur et al. derived a 3-gene signature (FCN3, CLEC1B, and PRC1), which has achieved an AUC of 0.91–0.96 on its validation set, which used peripheral blood samples containing mononuclear cells of both HCC and healthy patients (34). Further, each one of the three genes, showed prognostic character, as their expression levels, when stratified to greater than mean and lesser than mean groups, correlated with overall survival, progression-free survival (PFS) and disease-free survival (DFS) on univariate analysis.

A wide array of HCC biomarkers have been investigated...
using AI-driven techniques: gene co-expression patterns (35), miRNA (36), HCC-related genes (37) and serum biomarkers (33) have all been suggested as potential diagnostic signatures. Urine biomarkers have also been identified using ML approaches—Liang and colleagues identified five differential metabolites, from 37 urine samples (25 early-HCC and 12 controls), showing a sensitivity of 96.5% and specificity of 83% in differentiating between healthy and HCC patients, on an independent validation set (n=25) (38).

Another application of AI is data mining, which can be used to automatically screen existing literature to identify biomarker candidates, making it a useful tool for bioinformaticians (39).

**Characterisation**

The summary of literature on HCC characterisation, based on biomarker, imaging and pathology is presented in Table 3.

**Biomarkers**

Although the primary use of biomarkers is the detection of HCC, ML approaches also allow for stratification of HCC patients based on their biomarker profile, which can have therapeutic implications, as the response to treatment can be dependent on HCC subtype.

Genomics and epigenomics (DNA methylation patterns) analysis using ML has allowed for accurate differentiation between early-stage (stage I) and late-stage (stages II-IV) HCC (45). Moreover, Estevez et al. have established a biomarker-based classification into HepB-HCC, HepC-HCC and non-viral HCC, using cytokine profile from serum samples (46). Such stratification can have clinical implications for management, as well as understanding differences in disease pathogenesis.

**Imaging**

AI can help analyse radiological features from ultrasound, computed tomography (CT) and magnetic resonance imaging (MRI), all of which are routinely used in the diagnosis and differentiation of liver pathology.

The first use of AI in HCC characterisation based on imaging focused on the region of interest (ROI) analysis and computer-aided diagnosis systems using US and (47), CT (51) or MRI (55). These simple models employed analysis of features, such as lesion border or texture, to differentiate between benign and malignant liver tumour and thus identify HCC. Nowadays, more sophisticated AI-driven solutions are used in lesion differentiation, with greater accuracy and better differentiation, staging and stratification abilities.

For ultrasound scan, differentiation between cirrhotic liver and HCC, using neural networks can be made with 94.5% classification accuracy (48). Differentiation between atypical HCC and focal nodular hyperplasia (FNH) on the contrast-enhanced US, have also been reported, achieving 94.4% classification accuracy, when compared against pathology report analysis (biopsy or resection) and subsequent clinical follow-up (49). High differentiation accuracy has also been found for CT and MRI scans (52,64). What is more, Yamashita et al. have proven the feasibility of CNN assigning Li-RADS grades [an HCC CT/MRI scan probability classification used by American College of Radiologists (65)], to guide treatment decision-making (66).

AI models can also assist in imaging-based grading of HCC. Using contrast-enhanced ultrasound (CEUS), Sugimoto et al. established classification into well-differentiated, moderately differentiated and poorly differentiated HCC, with an AUC of 0.863–0.872 (50). Similar differentiation based on MRI scans was shown by Zhou et al. (56).

Furthermore, tumour segmentation algorithms have been employed to aid in management planning. Visualisation of the tumour can dictate decisions regarding tumour extent and resection. These algorithms, based on contrast-enhanced CT scans, can provide clinically useful 3D projections with a high degree of accuracy, as shown by Li et al. (53).

Most recently, radiomics approach to imaging analysis has been proposed, involving a multi-step process to derive large datasets of radiological features, via image acquisition, segmentation, feature extraction and automated analysis of patterns by using high throughput computing (67). Studies utilising this methodology shed light on the future directions of AI in imaging for HCC, highlighting its potential for high accuracy tumour characterisation and classification on MRI (57) and multi-phase contrast-enhanced CT (54) by analysing of textural features.

**Pathology**

AI has been used in pathology in order to precisely analyse the results of biopsies and resections to help with lesion characterisation and differentiation, using image-analysis
Table 3 Summary of literature on HCC characterisation, based on biomarkers, imaging and pathology

<table>
<thead>
<tr>
<th>Domain</th>
<th>Sub-category</th>
<th>Notes</th>
<th>N</th>
<th>AI algorithm</th>
<th>Type of validation</th>
<th>AUC (95% CI)</th>
<th>Limitations</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>Biomarkers</td>
<td>N/A</td>
<td>Genomics and epigenomics for HCC staging</td>
<td>400 (173 early stage, 177 late stage, 50 normal)</td>
<td>Support machine vector</td>
<td>Internal validation (cross-validation)</td>
<td>0.99 (0.98–0.99)</td>
<td>Biomarkers derived from tissue, which requires invasive approaches for sample isolation</td>
<td>Kaur et al. 2019 (45)</td>
</tr>
<tr>
<td>Imaging</td>
<td>US</td>
<td>Differentiation between focal hepatic lesions (benign and malignant)</td>
<td>51 images</td>
<td>ROI analysis</td>
<td>Development only (no validation)</td>
<td>N/A</td>
<td>Only two data validators (radiologist); data possibly not generalizable</td>
<td>Kim et al. 2009 (47)</td>
</tr>
<tr>
<td>CT</td>
<td>Differentiation between cirrhosis and HCC</td>
<td>188 images</td>
<td>ANN</td>
<td>Internal validation (cross-validation)</td>
<td>Accuracy 94.5%</td>
<td>Small sample size</td>
<td>Bharti et al. 2018 (48)</td>
<td></td>
</tr>
<tr>
<td>CT</td>
<td>Differentiation between atypical HCC and focal nodular hyperplasia</td>
<td>257 images</td>
<td>ANN</td>
<td>Internal validation (cross-validation)</td>
<td>F1-score 94.62%</td>
<td>Small sample size leading to lack of generalisability and network which is difficult to interpret</td>
<td>Huang et al. 2020 (49)</td>
<td></td>
</tr>
<tr>
<td>CT</td>
<td>Grading based on tumour differentiation</td>
<td>232 (76 well-differentiated HCC, 133 moderately differentiated HCC, 23 poorly differentiated HCC)</td>
<td>ANN</td>
<td>Development only (no validation)</td>
<td>Accuracy 87.5%</td>
<td>Use of 2D ultrasound and fine-needle biopsy specimen for establishing differentiation instead of surgical specimen</td>
<td>Sugimoto et al. 2016 (50)</td>
<td></td>
</tr>
<tr>
<td>MRI</td>
<td>Differentiation between focal hepatic lesions (benign and malignant)</td>
<td>147 images</td>
<td>Region of interest (ROI) analysis</td>
<td>Internal validation (cross-validation)</td>
<td>Accuracy 84.96%</td>
<td>Small sample size</td>
<td>Mougiakakou et al. 2007 (51)</td>
<td></td>
</tr>
<tr>
<td>MRI</td>
<td>HCC diagnosis from nodular, diffuse and massive tumours</td>
<td>168 (46 diffuse tumour, 43 nodular tumours, 76 massive tumours)</td>
<td>Convolutional neural networks (CNN)</td>
<td>Internal validation (random sample split)</td>
<td>Accuracy 98.4–99.7% (range)</td>
<td>Segmentation performance for diffuse tumour is not as good as other types, creating noise in the data</td>
<td>Li et al. 2020 (52)</td>
<td></td>
</tr>
<tr>
<td>MRI</td>
<td>Tumour segmentation on contrast-enhanced enhanced CT</td>
<td>201 images</td>
<td>Fully convolutional neural networks</td>
<td>External validation</td>
<td>Accuracy 93.7%</td>
<td>Network which is difficult to interpret and restricted by GPU memory</td>
<td>Li et al. 2018 (53)</td>
<td></td>
</tr>
<tr>
<td>MRI</td>
<td>Differentiation between five phases of CT</td>
<td>502 images</td>
<td>Random forest</td>
<td>External validation</td>
<td>Accuracy 94–98% (range)</td>
<td>Overlap between five phases on CT scan (no clear guidelines on start and end of each phase); decision based on expertise of principal investigators</td>
<td>Derce et al. 2020 (54)</td>
<td></td>
</tr>
<tr>
<td>Pathology</td>
<td>N/A</td>
<td>Differentiation between HCC and cholangiocarcinoma</td>
<td>106 whole-slide images</td>
<td>Deep learning</td>
<td>Internal validation (random sample split)</td>
<td>0.842</td>
<td>Methodology not reflective of clinical practice, reducing general applicability of the results</td>
<td>Kiani et al. 2020 (58)</td>
</tr>
<tr>
<td>Pathology</td>
<td>N/A</td>
<td>Differentiation between healthy tissue and HCC</td>
<td>1,773 image features</td>
<td>Random forest</td>
<td>External validation</td>
<td>0.886</td>
<td>Different in ethological factors between two datasets used</td>
<td>Liao et al. 2020 (59)</td>
</tr>
<tr>
<td>Pathology</td>
<td>N/A</td>
<td>Grading of HCC</td>
<td>108 patients</td>
<td>ROI analysis, fractal dimensions</td>
<td>Internal validation (cross-validation)</td>
<td>Accuracy 95.97% (Atupelage et al. 2014); accuracy 95.51% (Atupelage et al. 2013)</td>
<td>Small sample size and lack of external validation. Inclusion of non-informative texture features into the classifiers</td>
<td>Atupelage et al. 2014 (60); Atupelage et al. 2013 (61)</td>
</tr>
<tr>
<td>Pathology</td>
<td>N/A</td>
<td>HCC diagnosis using hyperspectral imaging analysis</td>
<td>14 hyperspectral images</td>
<td>Deep learning</td>
<td>Internal validation (cross-validation)</td>
<td>0.950</td>
<td>Small sample size and single-centre character of the study</td>
<td>Wang et al. 2020 (62)</td>
</tr>
<tr>
<td>Pathology</td>
<td>N/A</td>
<td>HCC grading using multiphoton microscopy</td>
<td>217 images</td>
<td>CNN</td>
<td>Internal validation (cross-validation)</td>
<td>0.941 (0.913–0.968)</td>
<td>Small sample size, insufficient for deep learning purpose</td>
<td>Lin et al. 2019 (63)</td>
</tr>
</tbody>
</table>

**Notes:**
- CT: computed tomography; MRI, magnetic resonance imaging; US, ultrasound scan.
- HCC, hepatocellular carcinoma; AI, artificial intelligence; AUC, area under the curve; 95% CI, 95% confidence intervals; HepB, viral hepatitis type B; HepC, viral hepatitis type C; ANN, artificial neural network; ROI, region of interest; CNN, convolutional neural network; N/A, non-applicable; CT, computed tomography; MRI, magnetic resonance imaging; US, ultrasound scan.
techniques.

Whole-slide images using H&E stain were analysed by Kiani et al. to differentiate between cholangiocarcinoma and HCC, achieving 84.2% differentiation accuracy on an independent validation set (58). When this model was offered as assistance to experienced histopathologists, it increased accuracy when correct, but decreased accuracy when incorrect, bringing attention to possible negative consequences of introducing AI into clinical practice (58).

Such models, on top of having high diagnostic accuracy (88.6% in differentiation accuracy between healthy liver tissue and HCC), can also have prognostic value, predicting response to treatment, a crucial characteristic for an AI-driven model to have in the future, to facilitate its clinical applicability (59).

Pathological slides can also be used for HCC grading (low-grade HCC vs. high-grade HCC) based on the number of nuclei (60) or whole-slide texture analysis (61).

Finally, one of the future applications of AI-driven solutions is lab-free, real-time pathological diagnosis, suggested by Wang et al. (62). The deep learning model yielded 88.1% accuracy, using hyperspectral imaging instead of the gold standard frozen sections; a method which is both time and labour-intensive. Lin et al. investigated another alternative to lab-based tissue processing, multiphoton microscopy (MPM), which also shows potential, as the use of CNN resulted in 90% differentiation accuracy into low-, moderate- and high-grade HCC (63).

Prediction

The summary of literature on HCC prediction, based on treatment outcomes and overall survival is presented in Table 4.

Prediction of treatment outcomes

AI algorithms were successfully developed to predict response to and survival after transarterial chemoembolization (TACE). To forecast survival after TACE, an ANN model was produced based on all parameters used by ART, ABCR and SNACOR risk scoring systems, as well as, age, gender, type of TACE and type of imaging before the second TACE. When compared with abovementioned scores, risk prediction made by the ANN (AUC 0.83±0.06) was found to be significantly superior (P<0.001) to that of ART (AUC 0.54±0.08) (68). ANN model has also outperformed SNACOR and ABCR, however, the difference was not significant (P=0.201 and P=0.143 respectively). All of the other studies predicting response to TACE utilised analysis of imaging (69-71,87,88). When a combination of extracted MRI features, clinical information and therapeutic features were used to train logistic regression and random forest models, to classify patients as responders or non-responders, overall accuracy of 78% was reached (69). Moreover, two research groups developed computer tomography based convolutional neural network models (70,88). They predicted time to progression (TTP) based on follow-up CT radiological criteria (mRECIST), to divide patients as TACE-susceptible (TTP >14 weeks) or TACE-refractory (TTP <14 weeks), and classified patients into one of four groups: complete TACE response, partial response, stable disease or progressive disease respectively. The models achieved 74.2%, followed by 84.3% accuracy with AUC scores of 0.95–0.97 for individual prediction categories.

Finally, Liu and colleagues validated three (one deep and two ML), predictive models, based on radiomic features of CEUS scans (71). Deep learning model outperformed the other two in assigning patients in the validation cohort to either objective-response to TACE or non-response, reaching 0.93 AUC score.

AI solutions were also used to predict stereotactic body radiotherapy (SBRT) and radiofrequency ablation (RFA) outcomes. To challenge currently used dose-volume histogram (DVH) based metric and forecast hepatobiliary toxicity of SBRT more accurately, Ibragimov et al. trained a CNN model on healthy organ CT images, liver SBRT cases and nanodosimetric pre-treatment patient features (72). The approach used was general with only 36 HCC patients out of 125 cases in total. CNN model was moderately successful with AUC of 0.76 and overall accuracy similar to that of current DVH metric. Enriching the liver SBRT database and checking the performance of the toxicity prediction framework may improve the future performance of the model.

To predict 1-year and 2-year DFS of patients who underwent CT-guided percutaneous RFA used in early HCC stages, Wu and colleagues developed an ANN-based on 15 clinical features (73). The model was more accurate when anticipating 1-year DFS than 2-year DFS, with 85.0% and 67.9% accuracy, respectively.

Several AI predictive models were also validated to explore post-resection survival and HCC recurrence. Currently, the risk is calculated based on the histological analysis of resected specimens (74). Deep CNN models were shown to optimise this process by implementing the
<table>
<thead>
<tr>
<th>Domain</th>
<th>Sub-category</th>
<th>Notes</th>
<th>N</th>
<th>AI algorithm</th>
<th>Type of validation</th>
<th>AUC (95% CI)</th>
<th>Limitations</th>
<th>Reference</th>
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<tr>
<td>Treatment outcomes</td>
<td>TACE</td>
<td>Response to treatment based on clinical data and risk scoring systems</td>
<td>282</td>
<td>ANN</td>
<td>Internal validation (random sample split)</td>
<td>0.83±0.06</td>
<td>No independent external validation cohort</td>
<td>Mähringer-Kunz 2020 (68)</td>
</tr>
<tr>
<td></td>
<td>Response to treatment based on MRI and clinical data</td>
<td>36 HCC patients</td>
<td></td>
<td>Random forest</td>
<td>Internal validation (cross-validation)</td>
<td>Accuracy 78%</td>
<td>Small patient cohort</td>
<td>Abajian et al. 2018 (69)</td>
</tr>
<tr>
<td></td>
<td>Response to treatment based on CT images</td>
<td>105 patients</td>
<td></td>
<td>CNN</td>
<td>Internal validation (cross-validation)</td>
<td>Accuracy 74.2% (64–82%)</td>
<td>General model applied to multiple TACE chemotherapy regimens</td>
<td>Morshid et al. 2019 (70)</td>
</tr>
<tr>
<td></td>
<td>Response to treatment based on the contrast-enhanced US</td>
<td>130 patients</td>
<td></td>
<td>Deep learning</td>
<td>Internal validation (random sample split)</td>
<td>0.93 (0.80–0.98)</td>
<td>Limited sample size; single-centre retrospective data</td>
<td>Liu et al. 2020 (71)</td>
</tr>
<tr>
<td>SBRT</td>
<td>Hepatobiliary toxicity prediction based on CT images</td>
<td>125 patients and 2,644 images of human organs</td>
<td></td>
<td>CNN</td>
<td>Internal validation (cross-validation)</td>
<td>0.85</td>
<td>Limited liver SBRT database</td>
<td>Ibragimov et al. 2018 (72)</td>
</tr>
<tr>
<td>RFA</td>
<td>Disease-free survival prediction</td>
<td>252 1-year and 179 2-year DFS</td>
<td></td>
<td>ANN</td>
<td>Internal and external validation</td>
<td>0.84 for 1-year, 0.75 for 2-year DFS prediction</td>
<td>Uneven 1-year and 2-year DFS group sizes</td>
<td>Wu et al. 2017 (73)</td>
</tr>
<tr>
<td>Resection</td>
<td>Recurrence risk based on whole-slide histological analysis</td>
<td>522 patients</td>
<td></td>
<td>CNN</td>
<td>Internal and external validation</td>
<td>c-index 0.70</td>
<td>Overfitting (inferior performance on external validation set)</td>
<td>Sallard et al. 2020 (74)</td>
</tr>
<tr>
<td></td>
<td>Recurrence and progression-free survival based on immunological biomarkers</td>
<td>221 patients</td>
<td></td>
<td>Random forest</td>
<td>Internal validation (cross-validation)</td>
<td>0.80</td>
<td>No external dataset validation</td>
<td>Zhou et al. 2019 (75)</td>
</tr>
<tr>
<td></td>
<td>Survival based on CT images</td>
<td>470 patients</td>
<td></td>
<td>ANN</td>
<td>Internal and external validation</td>
<td>0.803</td>
<td>Most patients had hepatitis B-related HCC</td>
<td>Ji et al. 2019 (76)</td>
</tr>
<tr>
<td></td>
<td>Survival based on CT images</td>
<td>167 patients</td>
<td></td>
<td>Internal validation (cross-validation)</td>
<td>Accuracy 82%</td>
<td>No external dataset validation</td>
<td>Wang et al. 2017 (77)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Survival based on CT images</td>
<td>995 patients</td>
<td></td>
<td>Bayesian network</td>
<td>Internal validation (cross-validation)</td>
<td>Accuracy 57%</td>
<td>Lack of temporal information in the patient data</td>
<td>Xu et al. 2019 (78)</td>
</tr>
<tr>
<td></td>
<td>Microinvasion based on biomarkers and MRI</td>
<td>160 patients</td>
<td></td>
<td>Logistic regression</td>
<td>Internal validation (random sample split)</td>
<td>0.83 (0.71–0.95)</td>
<td>Normalisation of the signal intensities on MR images not performed</td>
<td>Feng et al. 2019 (79)</td>
</tr>
<tr>
<td></td>
<td>Survival based on BCLC criteria</td>
<td>976 patients</td>
<td></td>
<td>Classification and Regression Tree</td>
<td>Internal validation (random sample split)</td>
<td>c-index 0.604</td>
<td>Majority of patients had favourable liver function</td>
<td>Tsimigras et al. 2020 (80)</td>
</tr>
<tr>
<td>Transplantation</td>
<td>Recurrence based on clinical data and CT images</td>
<td>133 patients</td>
<td></td>
<td>Classification and Regression Tree</td>
<td>Internal validation (random sample split)</td>
<td>c-index 0.789 (0.620–0.957)</td>
<td>Retrospective design</td>
<td>Guo et al. 2019 (81)</td>
</tr>
<tr>
<td>Overall survival</td>
<td>N/A</td>
<td>Survival based on biomarkers in HepB-HCC</td>
<td>67 samples (40 patients)</td>
<td>Supervised ML</td>
<td>Internal validation (cross-validation)</td>
<td>Accuracy 78%</td>
<td>Small dataset</td>
<td>Ye et al. 2003 (82)</td>
</tr>
<tr>
<td></td>
<td>Survival based on DNA methylation patens</td>
<td>488 samples</td>
<td></td>
<td>Multiple techniques combined</td>
<td>External validation</td>
<td>Accuracy 63%</td>
<td>Reason for 40% of all patients being hard to predict remained unclear</td>
<td>Itzel et al. 2019 (83)</td>
</tr>
<tr>
<td></td>
<td>Survival based on DNA methylation patens</td>
<td>377 HCC, 50 control samples</td>
<td></td>
<td>Internal validation (cross-validation)</td>
<td>0.95 (mean 10-fold cross-validation score)</td>
<td></td>
<td>Limited validation outcomes reported</td>
<td>Dong et al. 2019 (84)</td>
</tr>
<tr>
<td></td>
<td>Survival based on gene-expression pathways</td>
<td>355 patients</td>
<td></td>
<td>Support vector machine</td>
<td>Internal and external validation</td>
<td>c-index 0.83</td>
<td>Class label of the TCGA HCC samples obtain using whole TCGA dataset</td>
<td>Fa et al. 2019 (85)</td>
</tr>
<tr>
<td></td>
<td>Survival based on clinical features</td>
<td>165 patients</td>
<td></td>
<td>ANN</td>
<td>Internal validation (cross-validation)</td>
<td>0.700</td>
<td>No external dataset validation</td>
<td>Santos et al. 2015 (86)</td>
</tr>
</tbody>
</table>

HCC, hepatocellular carcinoma; AI, artificial intelligence; AUC, area under the curve; 95% CI, 95% confidence intervals; HepB, viral hepatitis type B; ANN, artificial neural network; CNN, convolutional neural network; TACE, transarterial chemoembolization; SBRT, stereotactic body radiation therapy; RFA, radiofrequency ablation; BCLC criteria, Barcelona Clinic Liver Cancer criteria; DFS, disease-free survival; TCGA, The Cancer Genome Atlas; CT, computed tomography; MRI, magnetic resonance imaging.
analysis of whole-slide digitised histological slides (74). Two CNN algorithms reached similar efficiency, however, the combination of CNN with human input (tumour areas annotated by the pathologist) slightly outperformed the one without (AUC 0.78 vs. 0.75).

Immunological tumour biomarkers were also used as a tool for predicting survival, using three indices: Overall Survival (≤24 or >24 months), PFS (≤6 or >6 months) and recurrence/death producing AUC between 0.76 and 0.8 and accuracy over 85% (75).

Moreover, groups led by Ji and Wang validated CT-based ANN and deep CNN to predict survival (76,77). The first group developed a novel three-feature radiomic signature of contrast-enhanced CT image, where performance was improved by combining it with clinical features (c-index 0.63–0.69 vs. 0.73–0.801). Wang and colleagues employed multi-phase CT radiomics features together with clinical models to yield a combined model with AUC of 0.82. A Bayesian network-based approach was also used to predict the probability of post-resection HCC recurrence which considered respective recurrence evolution paths for clinical feature datasets (78).

At the same time, AI techniques were explored to provide information about the predictive power of particular biomarkers, which could guide decisions on liver resection. Feng et al. used MRI radiomics to predict microvascular invasion status of the tumours, an important factor for hepatectomy, reaching AUC of 0.83 for validation dataset (79). Furthermore, an AI model was applied to determine the prognostic weight of factors comprising the Barcelona Clinic Liver Cancer (BCLC) guidelines, which selected alpha-fetoprotein and Charlson comorbidity score as the most important preoperative factors of overall survival among BCLC-0/A patients, and radiologic tumour burden score for BCLC-B patients (80). These results have the potential to shape the next iteration of BCLC guidelines.

AI was also used to predict DFS following liver transplantation. CT radiomics and clinical risk factors were combined to train the model, which yielded c-index of 0.79 when tested on the validation cohort (81).

**Prediction of overall survival of HCC patients**

ML algorithms have also been used for the general prediction of survival in HCC patient population. The earliest studies used supervised ML to differentiate between metastatic and non-metastatic HCC in HepB positive patients based on gene expression, hence obtaining information about probable survival chance (82). It also identified osteopontin as a biomarker of metastatic HCC.

Recently, groups led by Itzel and Dong explored possibilities of using random gene sets and DNA methylation levels for survival prognostics (83,84), while Fa et al. followed disease-specific patterns in dysregulated gene-expression pathways instead of single genes (85). More studies followed with training predictive algorithms of clinical features of HCC patients to forecast survival. Santos et al. combined a cluster-based oversampling method with the neural network model to account for small and incomplete datasets, improving the AUC score from 0.69 to 0.75 (86).

**Discussion**

AI solutions have been applied in all aspects of medicine in recent years and HCC is no exception. AI has led to advances in detection of HCC (based on pre-malignant changes, imaging and biomarkers) due to its ability to analyse large datasets and integrate information efficiently. Biomarkers identified by the integration of multiple ‘omics’ datasets are especially promising, potentially leading to the identification of a biochemical tumour signature, revolutionising HCC detection in the future.

As AI algorithms have become more sophisticated, research emphasis shifted towards lesion characterisation, differentiation between various types of hepatic malignancies and stratification of patients into groups, based on the tumour stage or grade. Various datasets, such as radiological images or clinical and pathological data, can be used separately or in combination to provide accuracy superior to that of traditional statistical tools. What is more, AI-driven solutions can help in reducing interobserver variability when analysing imaging studies, leading to standardisation. Since most of the hepatocellular cancers develop on the background of chronic liver disease, future efforts in HCC characterisation should focus on accurate differentiation between pre-malignant changes and early malignancy, to provide the most clinical benefit.

AI methods for prediction of overall survival and treatment outcomes in HCC have emerged in the past two years and remain a dynamic area of study. Predictive potential of current models is higher for short-term outcomes rather than long-term survival, however, this approach offers an array of novel predictive tools to shape HCC guidelines and support clinical decision making. In
the future, models combining radiomic data with clinical features to provide characterisation and prognosis for HCC patients are likely to be implemented in the clinical practice to support decisions on treatment.

Limitations

AI has inherent limitations, and this holds true for its medical applications for HCC detection, characterisation and prediction. Most of the studies discussed in this review suffer from a small sample size, which is a major issue for deep learning algorithms, as they require large training dataset to perform well. What is more, a lot of the studies use single-institution data, often from tertiary care centres. As a result, despite achieving high AUC and accuracy, such AI algorithms often cannot be used outside a narrow context, ultimately hampering widespread clinical adoption of AI within HCC. Specific limitations also exist in each of the three domains discussed. Studies on detection, focus on specific subtypes of HCC and narrow populations (e.g., HBV positive patients), rendering the proposed AI algorithms are unable to perform screening for HCC on the level of the general population. Studies on characterisation are limited by lack of standardisation of biomarker assays, imaging techniques and histological specimen preparation, all of which contribute to difficulties in applying the results of the research in settings different than the original. Even though significant advancements in the implementation of HCC outcomes prediction have been made in the recent years, there are many outstanding questions. Interpretation of algorithm outcomes remains a major challenge as it is difficult to explain why the model fails to make accurate predictions for a proportion of cases.

This review is also not devoid of limitations. Firstly, the methodology of the study, being a literature review limits the applicability of its conclusions. Moreover, the heterogeneity of quantitative data and AI methodologies has not allowed for pooled analysis of outcomes, but only a qualitative synthesis of evidence. The three-domain classification that was adopted for the purposes for that review is also imperfect, as more recent studies often discuss potential uses of HCC across more than one domain, by utilising multiple types of data to inform clinical decisions. Moreover, selection bias might exist, and studies as studies with negative findings might not be published. Finally, this review only aimed at assessing studies in English, however, high-quality studies in other languages might exist.

Future research directions

Consistency in reporting and transparency in publishing AI algorithms can significantly improve the clinical value of studies exploring AI models applied to HCC. Discrepancies in data standards and diagnostic devices used across different treatment centres contribute to overfitting of AI models, which needs to be overcome to facilitate the development of AI solutions with general applicability. External validation of the AI algorithms should also be favoured over retrospective internal validation, further increasing the applicability of the AI-driven solutions. International and interdisciplinary collaboration is instrumental in approaching this issue, as shown in a recent study that investigated an AI model in breast cancer diagnosis using data from both the UK and US. What is more, widespread availability of source code for algorithms can speed the process of AI development and validation, also contributing to larger applicability of these solutions. Finally, effective communication between computer scientists, engineers and clinicians is crucial for generating research which can redefine the current practice to address unmet clinical needs.

Conclusions

AI will revolutionise the way we detect and characterise HCC, as well as predict the course of its development, however, it is still experimental. In recent years, the rise of big data has caused AI-driven solutions utilising clinical data, radiological images, biomarkers and pathology results to emerge and gradually improve in accuracy, however, their widespread introduction into the clinical practice has not occurred yet. Robust validation, large scale studies, multicentre cooperation, advocacy for AI and education on AI amongst clinicians are all necessary for AI models to take the next step, so that in the future, such models using multiple data modalities, have the chance of influencing HCC guidelines and shaping clinical practice.

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Footnote

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