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**Challenges and opportunities in the application of artificial intelligence in gastroenterology and hepatology**

Christou CD *et al*. AI in gastroenterology and hepatology

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**Abstract**

Artificial intelligence (AI) is an umbrella term used to describe a cluster of interrelated fields. Machine learning (ML) refers to a model that learns from past data to predict future data. Medicine and particularly gastroenterology and hepatology, are data-rich fields with extensive data repositories, and therefore fruitful ground for AI/ML-based software applications. In this study, we comprehensively review the current applications of AI/ML-based models in these fields and the opportunities that arise from their application. Specifically, we refer to the applications of AI/ML-based models in prevention, diagnosis, management, and prognosis of gastrointestinal bleeding, inflammatory bowel diseases, gastrointestinal premalignant and malignant lesions, other nonmalignant gastrointestinal lesions and diseases, hepatitis B and C infection, chronic liver diseases, hepatocellular carcinoma, cholangiocarcinoma, and primary sclerosing cholangitis. At the same time, we identify the major challenges that restrain the widespread use of these models in healthcare in an effort to explore ways to overcome them. Notably, we elaborate on the concerns regarding intrinsic biases, data protection, cybersecurity, intellectual property, liability, ethical challenges, and transparency. Even at a slower pace than anticipated, AI is infiltrating the healthcare industry. AI in healthcare will become a reality, and every physician will have to engage with it by necessity.

**Key Words:** Artificial intelligence; Machine learning; Gastroenterology; Hepatology; Artificial neural networks; Support vector machine

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**Core Tip:** The opportunities that arise from the application of artificial intelligence/machine learning-based models in gastroenterology and hepatology include the establishment of targeted screening programs through the identification of patients prone to develop cancer, the development of non-invasive diagnostic tools, the improvement of the diagnostic accuracy, the development of treatment allocation frameworks based on predictions of outcomes for different treatment modalities, the development of models to ensure cost-effective use of resources, the development of triage tools for higher levels of care and decision-making tools for further treatment, based on individualized patient outcome predictions, and finally the development of predictive models of prognosis for patient and family counseling.

**INTRODUCTION**

Artificial intelligence (AI) is an umbrella term to describe any application where computer systems are used to perform tasks normally associated with human intelligence[1,2]. AI is a cluster of interrelated fields, including machine learning (ML) and probabilistic reasoning, planning and decision making, fuzzy systems, computer vision, natural language processing, knowledge representation, and neural networks (NN)[1]. Despite their differences, these fields are all driven by advancements in computing power and Big Data.

ML could be described as a model that learns from past data in order to predict future data[3]. The application of ML in healthcare was catalyzed by several healthcare trends, such as electronic healthcare records and patient summaries, genomic analyses and biomedical research, routine imaging, and telemedicine, that have transformed the healthcare industry into a data-rich science with data as an omnipresent concept[4–6]. Specifically, while in 2013, the healthcare industry produced 153 exabytes (1018 gigabytes) of data, it has been projected to reach 2314 exabytes in 2020[7]. Therefore, the healthcare industry generates an enormous amount of data that conform with the features that define Big Data: Volume, high-velocity, high-variety, and veracity, which cannot be analyzed or managed by traditional software[5,8,9]. AI promises to process and analyze these extensive repositories of data and turn them into meaningful insights.

Several studies have described AI as the potential solution to long-standing healthcare challenges such as increasing diagnostic accuracy, enhancing telemedicine, providing substantial cost reduction, promoting evidence-based medicine, facilitating targeted literature search, and delivering individualized care[10–14]. More importantly, AI could substantially alleviate the burden of diseases by reducing the associated mortality and morbidity through optimizing patient outcomes[15,16].

In gastroenterology and hepatology, physicians handle large amounts of clinical data and an extensive repository of imaging data generated from endoscopy, ultrasound, and computed tomography (CT). Therefore gastroenterology and hepatology are data-rich fields, and thus fruitful ground for AI applications with extensive research conducted in these fields, particularly regarding the prevention, diagnosis, management, and prognosis of diseases[17,18]. We aim to comprehensively review the applications of AI in gastroenterology and hepatology and identify the current challenges of utilizing AI in healthcare in an effort to explore ways to overcome them.

**SEARCH STRATEGY**

We conducted a comprehensive literature review of the Medline, Cochrane, and Scopus databases using the following algorithm: [(artificial intelligence OR machine learning OR deep learning OR neural networks OR support vector machine OR computer-assisted OR computer-aided) AND (gastroenterology OR hepatology OR esophageal OR small bowel OR large bowel OR gastric cancer OR capsule endoscopy OR polyps OR colonoscopy OR colorectal cancer OR gastrointestinal bleeding OR inflammatory bowel disease OR Crohn's disease OR celiac disease OR ulcers OR hepatocellular carcinoma OR cholangiocarcinoma OR liver fibrosis OR fatty liver OR chronic liver disease OR cirrhosis)]. Articles were reviewed for eligibility by the two authors independently (CC, GT), and disagreements were solved by a discussion between the two authors. Finally, the reference lists of eligible articles were reviewed to identify further related literature, including articles, books, and other forms of publication. We excluded studies written in a language other than English and publications of abstracts. The review of the literature was completed on January 27, 2021.

**AI CLASSIFIERS**

Before discussing AI's current applications and challenges in gastroenterology and hepatology, we briefly describe the AI classifiers in ML models that we identified as the most commonly used among the eligible articles. Specifically, we describe the basic features of the support vector machine (SVM), the artificial NN (ANN), and the convolutional NN (CNN). ML is divided into supervised and unsupervised based on whether the training data is labeled or not[19]. In other words, through supervised ML, a new data set is classified for an outcome based on previous data sets that trained the ML model, while in unsupervised ML, there is no outcome but rather an attempt to detect unknown patterns and correlations within the data[20].

SVMs are supervised learning models with associated learning algorithms trained to assign classes to new cases. SVMs require a training set of data where each case is pre-labeled regarding the outcome classification. The data points of the features of each case are treated as points in a high-dimensional space[21]. Based on these mapped points, separating hyperplanes are drawn, aiming to distinguish these points upon their labeled class[22]. The hyperplane with the maximum distance from the mapped data points, known as the maximum functional margin, is selected since it provides the SMV's full potential to classify new examples correctly[21,22]. SVMs typically perform linear classification analysis. For non-linear analysis, kernel function could introduce additional dimensions to the raw data; and thus turning a non-linear problem into a linear problem in a higher-dimensional space[23].

An ANN is an ML model inspired by the human brain's neuronal connections that consist of an input layer, an output layer, and a hidden layer between them[18]. ANNs are applied both in supervised and unsupervised ML[24]. When multiple hidden layers are inserted between the input and output layers, and the network's architecture becomes more complex with multiple interconnections, the concept of deep NN (DNN) emerges[24,25]. During DNN training, the model adjusts the weighted correlations between the nodes of the input layer and the nodes of the multiple hidden layers[18,20]. Within DNN, CNNs are biologically variants inspired by multi-layer perceptrons[26]. CNNs have been extensively applied for medical image analysis[17,18]. During the CNN model development, the images are preprocessed using multiple filters, and multiple feature maps are created in a process called convolution[26].

**APPLICATIONS OF AI IN GASTROENTEROLOGY**

AI/ML-based models have been extensively applied in the prevention, diagnostics, management, and prognosis of gastrointestinal (GI) diseases, including GI bleeding (GIB), inflammatory bowel diseases (IBDs), malignant and premalignant lesions, and other nonmalignant lesions/diseases such as gastroesophageal reflux, *Helicobacter pylori* (*H. pylori*) infection, ulcers, celiac disease, and intestinal hookworms.

***Prevention***

Table 1 summarizes the findings of the identified studies applying AI/ML models in the prevention of gastroenterological diseases. AI/ML-based software could be used in screening programs to identify high-risk patients not currently identified by standard screening guidelines. In a recent study, researchers developed seven different AI/ML models to identify patients at high risk of developing gastric cancer following the eradication of *H. pylori* infection[27]. The extreme gradient boosting (GB) model demonstrated the highest performance[27]. A CNN was developed in a different study to classify patients at low, moderate, and high risk of developing gastric cancer using endoscopic images[28]. A similar study used multiple classification and regression trees (CART) to develop a model that predicts future gastric cancer development[29]. Another notable example is the ColonFlag test, an ML algorithm that uses age, sex, and complete blood count data to identify patients at high risk of developing colorectal cancer[30,31]. In a recent study, among 254 individuals who underwent colonoscopy, 19 cancers (7.5%) and 22 (8.7%) advanced adenomas were identified in a population of patients who would otherwise have avoided screening[31]. These studies demonstrate how AI/ML-based models could be used in a real clinical setting to establish targeted screening programs.

***Diagnostics***

Table 2 summarizes the findings of the identified studies applying AI/ML models to diagnose gastroenterological diseases. AI/ML-based software is applied in Computed Aided Detection or Diagnosis (CAD) systems used in radiology to augment medical images' interpretation accuracy. A CAD typically includes the stages of preprocessing, extraction of features, and feature selection and classification[32]. CAD systems could also aid endoscopists in navigating through the different anatomical locations of the GI tract. Specifically, a model based on a CNN employed esophagogastroduodenoscopy images to recognize the anatomical locations with outstanding performance[33].

Regarding the diagnosis of gastroesophageal reflux, an ANN model was developed as a non-invasive diagnostic tool by employing only clinical data[34]. For patients with Barrett’s esophagus, a deep learning-based (DL) CAD system was developed to differentiate patients with malignant from patients with nondysplastic Barret’s esophagus[35]. The CAD system also identified the optimal site to perform a biopsy with an accuracy between 92% and 97%[35]. An SVM model was developed in another study employing white-light endoscopic imaging to identify early neoplastic lesions[36]. In a recent study, images from volumetric laser endomicroscopy were used to develop a CAD system to detect neoplastic lesions for patients with Barret’s esophagus[37]. A similar study using volumetric laser endomicroscopy developed an ML model for detecting neoplastic lesions[38]. Notably, the model outperformed the experts in volumetric laser endomicroscopy[38]. Finally, to diagnose esophageal squamous cell carcinoma, a CNN model was developed by employing endocytoscopic images as an alternative to biopsy[39].

For diagnosing patients with *H. pylori* infection, a CNN was developed in a study employing gastroscopic images[40]. A different CNN for *H. pylori* diagnosis was developed in a prospective pilot study analyzing images taken from the stomach's lesser curvature using either white light imaging, blue light imaging, or linked color imaging[41]. Notably, employing blue light and linked color imaging yielded significantly higher performance than white light imaging[41]. In a similar study, a CNN model employed images from upper GI endoscopy to detect *H. pylori* infection[42]. Finally, Shichijo *et al*[43] developed two different CNN models to diagnose *H. pylori* infection. The first, a 22-layered deep CNN, performed comparably with endoscopists, while the second, which classified images based on their location in the stomach, performed comparably with endoscopists in terms of sensitivity and specificity but demonstrated a significantly higher accuracy[43]. The time needed to analyze all the images was 3 min and 18 s and 3 min and 14 s for the two CNNs, while for endoscopists, the average time needed was 230.1 min[43]. A recent meta-analysis evaluating the performance of CNNs to diagnose infection with *H. pylori* concluded that it is currently equivalent to physicians[44].

Regarding the diagnosis of gastric cancer, an SVM-based CAD model was developed to identify early gastric cancer features in narrow-band imaging (NBI) gastroscopy[45]. In a different retrospective study, a CNN was developed to detect gastric cancer from endoscopic images. Even though the model managed to correctly classify 71 of the 77 gastric cancer lesions (92.2% sensitivity), it also falsely identified 161 non-cancerous lesions as gastric cancer, yielding a low positive predictive value of 30.6%[46]. Approximately half of these lesions were gastritis with an irregular mucosal surface or changes in color tone[46]. Finally, in a recent study, a non-invasive GB/ decision tree (DT) model employing only nonendoscopic parameters was developed to diagnose gastric cancer[47].

Regarding the diagnosis of celiac disease, five different AI/ML-based models were developed in a study using clinical data as a base for non-invasively diagnosing celiac disease[48]. Interestingly, the models were tested using 13 different algorithms and different input variables variations, resulting in 270 different tested models[48]. Among them, a model based on the Bayesian classifier demonstrated the highest performance[48]. A different study developed an SVM that employed images from the endomysial autoantibody test for IgA-class antibodies to classify patients with celiac disease[49]. Finally, in a different study, the authors aimed to develop several AI/ML-based models, which employ clinical data, to predict celiac disease in a group of patients who remained undiagnosed[50]. The models were unsuccessful, with only two models slightly outperforming random chance in predicting celiac disease[50].

In the diagnosis of IBDs, a recent study developed five different AI/ML-based models using data from the gut microbiome to classify patients with IBD[51]. The Random Forest (RF) model demonstrated the highest performance with AUROCs of 0.80 when bacterial taxa were used and 0.82 when operational taxonomic features were used[51]. Finally, the models were tested in distinguishing between Chron’s disease (CD) and ulcerative colitis (UC), with the RF model demonstrating an AUROC > 0.9[51]. A multicentered, genome-wide association study used data from single-nucleotide polymorphisms to develop several AI/ML-based models to classify patients with CD and UC[52]. For pediatric IBDs, an SVM-based model employed data from histologic and endoscopic images to classify pediatric patients with CD, UC, or unclassified IBD[53]. Other studies have employed images from wireless capsule endoscopy to build SVM models that identify patients with CD with reported accuracies between 80.2% and 100%[54–57].

Focusing on capsule endoscopy, in a recent study, magnetically controlled capsule endoscopy imaging was employed to develop a CNN that provides an automatic detection and classification system for gastric lesions[58]. These lesions included erosions, polyps, ulcers, submucosal tumors, normal mucosa, and xanthomas[58]. A different study used images generated by a wireless capsule to develop a CNN able to classify the small bowel motility among six distinct intestinal motility events[59]. In a recent study, an AI-assisted capsule endoscopy reading model was developed to assist lesion identification[60]. Notably, the model significantly shortened the reading time of images by trainees[60].

To distinguish between hemorrhagic and ulcerative lesions, a CNN was developed using images from small bowel capsule endoscopy[61]. In a similar, recent study, a DNN model used capsule endoscopy images to classify different small bowel lesions (erosions, ulcers, tumors, and vascular lesions)[62]. SVM-based models were developed in two other studies employing images from capsule endoscopy to identify peptic ulcers[63,64]. In three different studies, CNNs have been developed, employing images from capsule endoscopy to diagnose intestinal hookworms, GI angiectasia, and celiac disease[65–67]. Finally, a CNN was developed in a recent study to detect colorectal neoplasias in images from colon capsule endoscopy[68].

Regarding the identification of colorectal polyps in coloscopy, a CNN model was designed to analyze colonoscopy images and videos and identify colorectal polyps[69]. Interestingly, the model's sensitivity to identify flat isochromatic, less than 0.5 cm polyps, which are typically associated with a higher missing rate, was estimated at 91.65% per image[69]. A different study, employing short videos from colonoscopies, developed a CNN model that identifies colorectal polyps[70]. In another study, a CNN was developed employing videos and images from coloscopy to detect colorectal polyps in real-time[71]. Interestingly, when experts reviewed the colonoscopy videos, they managed to identify additional non-removed polyps with the assistance of the CNN model[71]. In a recent study, an automated polyp detection model based on CNN was developed to identify and then classify colorectal polyps into adenomas, hyperplastic, sessile serrated adenomas, and cancer[72]. The model managed to process the images at a speed of 20 m per frame[72].

Specifically for colorectal polyps classification, in a prospective study, an SVM-based model was developed using NBI and methylene blue staining images to classify diminutive rectosigmoid adenomas in real-time[73]. A prospective pilot study developed an SVM model that employed NBI images to detect and classify colorectal polyps based on vascularization features as neoplastic and non-neoplastic[74]. In a comparative study, an SVM model demonstrated comparable performance at classifying the neoplastic nature of diminutive polyps (< 10 mm) to that of experts in NBI colonoscopy but surpassed the performance of non-experts[75]. Two other studies developed SVM models that employed NBI images to classify colorectal lesions (neoplastic or non-neoplastic)[76,77]. Finally, an SVM model was developed in a retrospective study to diagnose invasive colorectal cancer based on NBI endocytoscopy images[78].

Except for SVMs, several studies employed CNNs for colorectal polyps classification. One comparative study developed a CNN employing NBI colonoscopy images to classify neoplastic polyps[79]. Notably, the model's performance was found comparable to that of experts but superior to the performance of non-experts[79]. In another study focusing on NBI colonoscopy, a CNN model was developed to classify adenomatous from non-adenomatous polyps[80]. A different study developed a CNN model to classify adenomas from hyperplastic polyps during NBI colonoscopy[81].

The performances reported by the majority of the AI/ML-based models surpass both the NPV threshold recommended by the American Society of Gastrointestinal Endoscopy (90%) for adenoma detection and the estimated pooled NPV reported in a meta-analysis conducted by the society (91%)[82,83]. Finally, we should mention that currently, the majority of the CAD systems that we reported have the shortcoming of manual segmentations of lesions. The endoscopists should identify the areas of interest before the model could analyze and attempt to classify. This weakness has been acknowledged by the European Society of Gastrointestinal Endoscopy[84]. Other obstacles to developing CAD systems constitute the lack of large datasets and the lack of variability in images. A recent study aimed to resolve this by developing a CNN that “adds” polyps to the images to increase the repository of images for training and advance the development of automated polyp detection models[85].

***Management***

Table 3 summarizes the findings of the identified studies applying AI/ML models for the management of gastroenterological diseases. In a recent study, data from baseline impedance, nocturnal baseline impedance, and acid exposure time were used as a base for a DT model to predict the treatment response with proton pump inhibitors for patients with gastroesophageal reflux disease[86]. The aim was to establish a decision-making framework for treatment allocation[86].

AI/ML-based models have been used in the management of malignant GI lesions. Regardinggastric cancer, a CNN has been developed to predict whether the early gastric cancer has invaded the mucosa and submucosa layers of the stomach and act as a decision tool for endoscopic resection[87]. Interestingly, the CNN model outperformed endoscopists[87]. In the same concept, a DNN was developed to classify gastric cancer based on invasion depth as a basis for treatment allocation[88]. Specifically, the model demonstrated accuracy for predicting T1, T1a, T1b T2, T3, T4, and an overall accuracy of 77.2%, 68.9%, 63.6%, 49.1%, 51.0%, 55.3%, and 64.7%, respectively[88]. In colorectal cancer, the identification of microsatellite instability significantly impacts the treatment allocation process. A DL model was developed to identify microsatellite instability directly from hematoxylin and eosin-stained whole slide image[89]. Notably, the model outperformed a group of pathologists[89]. Finally, an SVM model that predicted the lymph node metastasis status of patients with colorectal cancer was designed as a tool to identify patients who would benefit from additional treatment following the endoscopic resection of T1 tumors[90]. Notably, the model outperformed the staging systems endorsed by current guidelines[90].

Regarding the management of GIB, in a recent study, an ML model was developed using data from patients admitted to the intensive care unit (ICU) following a GIB to predict the need for transfusion[91]. The authors suggested that it could potentially be used as a decision-making tool to triage patients to the ICU or the ward[91]. A different study developed several AI/ML models for patients with acute GIB to predict four different outputs: The source of bleeding, the need for urgent blood transfusion, the need for urgent endoscopy, and the disposition[92]. The study aimed to establish a decision-making framework for the efficient management of GIB. The RF model outperformed all seven other models for all four outputs[92].

Focusing on the upper GIB, an ANN model was developed as a non-invasive triage tool for patients with upper GIB[93]. In the external cohort, the ANN performed similarly to the complete Rockall Score (includes endoscopic variables) in predicting stigmata of recent hemorrhage[93]. A different study developed a CART model regarding acute variceal hemorrhage to predict rebleeding and mortality and achieved the discrimination of three distinct prognostic groups of low, intermediate, and high risk[94]. Its performance was significantly superior to Child-Pugh and Model for End-Stage Disease (MELD) but comparable to a conventional logistic regression (LR) model[94].

Focusing on the lower GIB, two different ANN models were developed in a study aiming to predict severe acute lower GIB and the need for surgical intervention[95]. The first ANN significantly outperformed the Strate prediction rule for predicting severe bleeding (AUROCs: 0.98 *vs* 0.66)[95]. A different study employed nonendoscopic variables to develop a model based on a GB classifier to predict severe lower GIB, recurrent bleeding, and the need for clinical intervention[96]. On external validation, the model was found equally accurate to a conventional LR model for recurrent bleeding and the need for clinical intervention but superior in predicting severe lower GIB[96].

***Prognosis***

Table 4 summarizes the findings of the identified studies applying AI/ML models regarding the prognosis of gastroenterological diseases. Several SVM-based nomograms were developed in a study to predict distant metastasis for operated patients with oesophageal squamous cell carcinoma[97]. A different study employed clinicopathological data to develop an ANN model to predict the survival of patients with esophageal cancer operated with curative intends[98]. The model surpassed the Tumor–Node–Metastasis (TNM) model[98]. Regarding gastric cancer, a recent study developed five different AI/ML-based models to predict recurrence in operated patients[99]. Among the models, the RF demonstrated the best performance[99].

An ANN model was developed for patients with IBD, which used meteorological data to predict seasonal variations of onset and relapse in patients with CD and UC[100]. In the validation cohort, the model predicted the onset frequency and the frequency of relapse of the IBD with a mean absolute percentage error of 37.58% and 17.1%, respectively[100]. A study focusing on CD developed an ANN model to predict mucosal remission for patients treated with azathioprine 16 wk following treatment[101]. In a study focusing on UC, an ANN was developed employing clinical data to predict the patients with UC treated with cytoapheresis, who will eventually require operation[102].

Regarding the prognosis of GIB, a CART model was developed to predict in-hospital mortality of cirrhotic patients presenting with upper GIB[103]. In a multicentered study, an ANN model was developed employing pre-endoscopic variables to predict 30-d mortality in patients with non-variceal upper GIB[104]. Similarly, a prospective, multicentered study employed pre-endoscopic variables to develop an ANN model that predicts 30-d mortality in patients with non-variceal upper GIB[105]. The ANN significantly outperformed the Rockall scoring system (AUROCs: 0.95 *vs* 0.67)[105].

Regarding colorectal cancer, a recent study developed several ML models to predict the RAS and BRAF mutation status for patients with advanced colorectal cancer[106]. Notably, the ANN demonstrated the best performance[106]. In a different study employing clinicopathologic variables and data generated from immunochemistry, a least absolute shrinkage and selection operator regression model was developed to predict the lymph node metastasis status in a cohort of operated patients for T1 colorectal cancer[107].

***Opportunities of AI application in gastroenterology***

Therefore, the opportunities that arise from applying AI/ML-based software in gastroenterology include:

AI/ML models could be developed and integrated into the clinical setting to employ routinely collected data directly from the patient’s electronic health records and flag patients at high risk of developing certain GI diseases in real-time. Current efforts include the prevention of gastric[27–29] and colorectal cancer[30,31]. Such ML models could become the basis for tailoring targeted screening programs.

Endoscopy is the gold standard for the diagnosis of a plethora of GI diseases. AI/ML models employing non-invasive parameters that provide reliably accurate diagnosis could substitute endoscopy or significantly minimize its use, thus ameliorating the impact of endoscopy-related complications, significantly decreasing the cost for diagnosis, and providing an alternative to an unpleasant intervention for the patient. Current efforts include the diagnosis of gastroesophageal reflux disease[34], gastric cancer[47], celiac disease[48,50], and IBDs[51].

Expect from replacing endoscopy; AI/ML models could also improve its efficacy. CAD systems could facilitate navigating the GI tract and serve as the second “observer” for the endoscopist, an “observer” non-susceptible to distraction, which identifies lesions missed by the endoscopist. Particularly for capsule endoscopy, a CAD system that automatically detects and classifies lesions could significantly decrease the time required to evaluate the images by endoscopists while increasing diagnostic accuracy. Current efforts include the diagnosis of IBDs[53–57], gastric lesions[58], small bowel mobility disorders[59], small bowel lesions such as hemorrhagic and ulcerative lesions[60–64], intestinal hookworms[65], GI angiectasia[66], celiac disease[67], and colorectal cancer[68].

Physicians could use CADs to augment the accuracy of classifying polyps based on their neoplastic nature. As a result, the morbidity and mortality associated with failing to remove a neoplastic polyp could be lessened. At the same time, the complications related to removing a non-neoplastic polyp could be avoided. Therefore, CADs could significantly increase the cost-effectiveness of polyp management. Current efforts include the development of SVMs[73–76] and CNNs[79–81] for the classification of the neoplastic nature of colorectal polyps during NBI colonoscopy.

AI/ML models that accurately predict the response to different treatments could be used as a basis for individualized treatment allocation. Current efforts include the response of treatment with proton pump inhibitors for patients with gastroesophageal reflux disease[86], the mucosal remission for CD patients treated with azathioprine[101], and the need for operation for UC patients treated with cytoapheresis[102].

Particularly for GIB management, AI/ML models could be used to identify the source of bleeding[92] and as frameworks for decision-making, including the need to transfuse patients[92], perform emergent endoscopy[92,93] or emergent surgery[95,96], and finally, triage patients with severe GIB to the ICU[95,96].

Finally, AI/ML models could be used as predictive tools that stratify the risk of complications and predict overall survival and recurrence following treatment. Such models could be used to tailor individualized follow-up schedules and for patient and family counseling. Current efforts include the prediction of overall survival for patients with esophageal cancer[98], the risk of recurrence of operated patients with gastric cancer[99], and 30-d mortality of patients with non-variceal GIB[104,105].

**APPLICATIONS OF AI IN HEPATOLOGY**

***Prevention***

Table 5 summarizes the findings of the identified studies applying AI/ML models for the prevention of disease in the field of hepatology. In a recent study, a prospective cohort of apparently healthy volunteers was enrolled in a study, which developed a DT-based model to classify patients based on their risk of developing non-alcoholic fatty liver disease (NAFLD) and liver fibrosis[108]. A similar study used routinely collected laboratory and clinical parameters to develop several AI/ML-based models to identify patients with NAFLD in the general population[109]. In a different study, several AI/ML models were developed as potential tools for targeted screening for NAFLD[110]. The Bayesian network model demonstrated the highest accuracy, followed by an SVM model[110]. Notably, an LR model outperformed all the models. These studies are examples of how AI/ML-based models could be used in primary care as tools for detecting chronic liver diseases. In a different study, multiple AI/ML models were developed as a basis for a non-invasive tool for assessing the level of fibrosis progression in NAFLD patients[111]. Notably, the RF model demonstrated the highest performance[111]. A different study managed to develop an ML model for patients at high risk of developing NAFLD to classify patients with non-alcoholic fatty liver (NAFL) and non-alcoholic steatohepatitis (NASH)[112]. Several AI/ML models were developed in a different study to classify healthy individuals from patients with NAFL and NASH[113]. Among them, a GB tree model demonstrated the highest performance, followed by an RF model[113]. In another study, four different AI/ML-based models were developed utilizing data from biochemical and enzyme-linked immunosorbent assays to classify patients with NAFLD and alcoholic liver disease with or without cirrhosis[114].

AI/ML models that utilize data directly from the electronic health records from patients infected with hepatitis B (HBV) and hepatitis C (HCV) could be used as the basis for targeted screening tools for liver carcinomas. A GB model was developed in a study using only serum markers aiming to predict fibrosis and classify the stage of fibrosis in two cohorts of patients with HBV and HCV[115]. An ANN model was developed in another study that employed only routine clinical data to predict significant fibrosis for patients with HBV[116]. In a similar study, an ANN model was developed by employing only routine laboratory data from HBV patients to identify patients with cirrhotic liver[117]. A different study developed an ANN model employing only non-invasive data to predict advanced liver fibrosis in HCV patients[118]. A similar study developed several AI/ML-based models to develop a non-invasive tool for identifying HCV patients with advanced fibrosis[119]. Among the developed models, a model based on alternating DTs demonstrated the highest performance[119]. These studies demonstrate how AI/ML models using routine clinical data could be used to identify patients who will benefit from a sustained virological response (SVR) to delay chronic liver disease progression.

Regarding the development of HCC, a recent study investigated if a DNN could outperform conventional LR models in predicting HCC development in patients with HCV[120]. A different study using variables including demographic, laboratory results, and clinical findings, developed RF, DNN, and LR models to reliably predict the stage of liver cirrhosis in patients infected with HCV[121]. Accurately predicting patients with HCV prone to develop HCC could help identify patients who would be benefited from a targeted screening and are in a greater need of antiviral treatment to achieve an SVR.

***Diagnostics***

Table 6 summarizes the findings of the identified studies applying AI/ML models for diagnosis in the field of hepatology. Regarding chronic liver disease, a study used CT imaging data of patients with a confirmed liver fibrosis diagnosis to develop a CNN model for the staging of liver fibrosis[122]. Notably, the model outperformed the radiologists’ interpretation[122]. A different study employed ultrasound imaging of patients with fatty liver disease to develop an SVM model and an extreme learning machine model (a type of ANN) for diagnosis and risk stratification[123]. A different study that employed ultrasound shear wave elastography features developed an SVM model to identify patients with chronic liver disease[124]. Another study used images from real-time tissue elastography to develop four different AI/ML-based models to classify liver fibrosis[125]. Notably, the RF model outperformed the rest, followed by the KNN and the SVM models[125].

In a study, CT imaging was employed to develop an ANN model that differentiates between HCC, intrahepatic peripheral cholangiocarcinoma (CCA), hemangioma, and metastasis[126]. Interestingly, when radiologists evaluated the images, their performance significantly increased when they considered the ANN’s output, from an AUROC of 0.888 to one of 0.934[126]. Focusing on HCC diagnosis, a recent retrospective study developed a CNN that employed MRI images of patients with HCC. The model was trained with a combination of images that met the Liver Imaging Reporting and Data System (typical) and with images that did not (atypical)[127]. In a multicenter, retrospective study, a CNN was developed that employed MRI scans to identify HCC lesions[128]. Notably, the model surpassed the performance of less experienced radiologists in the diagnosis of small HCC lesions[128]. The model was able to analyze 100 images in just 3.4 s[128]. Finally, a different study aimed to develop a non-invasive ANN model that predicts the presence of microscopic vascular invasion and the tumor grade of HCC[129].

Regarding diagnosis of CCA and pancreatic adenocarcinoma, a study developed an ANN model using data generated by metabolomic and proteomic analyses of bile from patients undergoing endoscopic retrograde cholangiopancreatography aiming to classify patients with and without cancer[130]. A different study used data from the plasma levels of bile acids to develop six different AI/ML-based models to classify patients as having CCA or a benign biliary disease[131]. Among the six developed models, a model based on the Naive Bayes classifier demonstrated the highest performance[131]. Finally, in another study, an ANN model was designed to analyze images from magnetic resonance cholangiopancreatography to diagnose CCA with a reported accuracy of 92.8%[132].

***Management***

Table 7 summarizes the findings of the identified studies applying AI/ML models to manage hepatic diseases. With the guidance of AI/ML-based software trained by data generated from gene mutation biomarkers, serum markers, imaging, and the clinical setting, high-quality, evidence-based, and individualized treatments could be employed regarding chemotherapy, radiotherapy, and immunotherapy. In HBV management, a recent study developed several AI/ML-based models that use soluble immune markers to predict early virological relapse after discontinuation of nucleoside analogs treatment[133]. The model could be used to exclude patients at high risk of virological relapse from treatment cessation. In HCV management, a study utilized data from full-length HCV genome sequencing of variants of HCV to develop multiple AI/ML-based models that classify HCV anti-viral resistance variants[134]. Notably, the SVM model demonstrated the highest performance[134].

In HCC management, a recent study used data from DNA methylation profiling to develop an RF-based model that predicts 6-mo progression-free survival. Such models could be used to personalize patient surveillance[135]. In an international, multi-institutional study, a CART model was developed that aimed to create a framework beyond the Barcelona-Clinic-Liver-Cancer (BCLC) staging system, which is currently endorsed by guidelines for treatment allocation[136,137]. The model defined six distinct prognostic groups of patients based on predictive factors of overall survival that could be used as a framework for treatment allocation[137]. Notably, the radiologic tumor burden score, which is not integrated into the BCLC staging system, was found as the best predictor of long-term outcome for BCLC stage B patients[137]. These findings could help us reevaluate HCC management into a multidisciplinary, individualized approach that goes beyond the BCLC criteria[138].

In the management of patients with CCA, in an international study, a CART model was developed with solely preoperative variables to identify patients who would be more likely to benefit from surgery[139]. The model managed to isolate four distinct prognostic groups of patients with similar patient outcomes[139]. The authors concluded that this model could be used to inform presurgical decisions-for example, the use of neoadjuvant therapy for patients with poor prognoses[139]. In a different study, the researchers developed a DNN model to establish an AI framework through which specific prognostic groups could be used to identify which patients were more likely to benefit from different treatment modalities such as neoadjuvant chemotherapy or transarterial chemoembolization[140]. The framework was found to be significantly more accurate than the current guidelines of the American Joint Committee of Cancer[140]. Finally, a study developed an ANN to predict which patients with inoperable hilar CCA will develop early occlusion following a bilateral plastic stent placement[141].

***Prognosis***

Table 8 summarizes the findings of the identified studies applying AI/ML models regarding prognosis in the field of hepatology. An ANN model was developed in a study to identify patients with HBV cirrhosis at a high risk of developing esophageal varices[142]. In a different study, two different RF models were created using clinical data, the first to identify esophageal varices and the second to classify patients with esophageal varices that require treatment[143].

Regarding HCC, several studies have focused on developing AI/ML-based models that would reliably predict patient outcomes (survival and recurrence). A retrospective study compared the performance among an ANN, an LR model, and a DT to predict the 1-, 3-, and 5-year disease-free survival in patients with HCC following resection[144]. A similar study used a nationwide database to compare an ANN and an LR model in predicting the 5-year survival of patients with HCC following hepatic resection and concluded that the ANN model surpassed the performance of the LR model[145]. In a different study by the same department, an ANN model and an LR model were compared, but instead, the outcome was in-hospital mortality, and the ANN was found superior to the LR model[146]. Interestingly, the study reported that the surgeon volume was the best single predictor of in-hospital mortality[146]. Regarding predictors, a different study comparing ANN and LR models reported that the ANN model identified a greater number of significant predictors than the LR model, except for outperforming it in survival predictions[147]. In a prospective study that aimed to compare an ANN model's performance to the performance of traditional staging systems in predicting survival for patients with early HCC, the ANN model outperformed the Hepato-Pancreato-Biliary Association’s, the TNM 6th, and the BCLC staging systems with higher reported AUROCs in all training, internal validation, and external validation cohorts[148]. In a recent study, a GB survival classifier-based model was developed to stratify the risk of an HCC-related death into three distinct categories[149]. Finally, in another recent study, an ANN identified albumin-bilirubin grade as the most important prognostic factor for the survival of patients with HCC treated with the combination of transarterial chemoembolization and sorafenib as initial treatment[150]. A systematic review aimed to compare the performance of AI/ML-based software and that of traditional linear prediction models in predicting survival for patients with HCC concluded that AI/ML models provided enhanced accuracy[151].

Except for survival, other studies have developed AI/ML models to predict the recurrence of HCC following therapeutical treatment. An interesting study employed several AI/ML methods, including Artificial Plant Optimization, SVM, and RF to predict HCC recurrence following radiofrequency ablation[152]. A recent study developed a CNN employing histopathologic images to predict recurrence in operated HCC patients[153]. A different study developed an SVM model to predict the recurrence in a group of patients who underwent radiofrequency ablation[154].

Regarding primary sclerosing cholangitis, a team of researchers derived and validated a risk estimate tool based on GB algorithms to predict the outcomes of the disease[155]. In a different study, an ML model was developed to predict survival curves for patients with primary sclerosing cholangitis following liver transplantation[156]. The *P*-value of the χ2 test of the distributional calibration was 1, indicating excellent calibration of the model[156].

Focusing on liver transplantation, a team developed an ANN model combined with genotyping for microsatellite mutations/deletion to predict HCC recurrence in a cohort of patients receiving a liver transplant[157]. Several other studies have focused on the survival of individual liver grafts following transplantation[158–160]. In a multi-centered study, the authors developed an ANN model to predict the 3-mo graft survival and loss[160]. Interestingly, their model outperformed all extensively validated scores, including the MELD, the donor risk index (DRI), the survival outcome following liver transplantation, and the balance of risk, with their performance significantly lower (AUROCs range: 0.42-0.67)[160]. In a different study, an RF model was developed to predict the 30-d failure graft. Notably, this model too outperformed MELD and DRI[158]. These findings could help reevaluate our thinking regarding the current models of recipient-donor matching.

***Opportunities of AI application in hepatology***

Therefore, the opportunities that arise from applying AI/ML-based software in hepatology include:

Regarding primary care, ML models could be integrated into the clinical setting and flag individuals in the general population at high risk of developing chronic liver disease in real-time by employing routinely collected data from the electronic health records. Current efforts include models that identify patients at high risk to develop NAFLD, NASH, fibrosis, and cirrhosis[108–114]. These models could be used to design targeted screening programs.

Particularly for patients with chronic HBV and HCV infection, AI/ML models could be used to stratify each individual's risk to progress through the several stages of cirrhosis and develop HCC. Multiple studies have been conducted in this regard for both HBV-related[115–117] and HCV-related[115,118–121] cirrhosis. Such models could be used to identify patients in greater need of SVR and tailor individualized follow-up schedules.

In diagnosis, AI/ML models provide the opportunity for increased diagnostic accuracy of various hepatic diseases and in multiple diagnostic modalities (such as US/CT/MRI imaging and histologic images). Current efforts include CAD models for the diagnosis of chronic liver disease, HCC[127–129], and CCA[130–132].

AI/ML models that accurately and reliably predict the response to specific treatments could be used to tailor evidence-based frameworks for individualized treatment allocation. Current efforts include the development of frameworks for the management of HCC[136,137] and CCA[139,140].

Regarding prognosis, AI/ML models could be used to reliably predict complications, in-hospital mortality, overall survival, and recurrence following treatment. Current efforts include the prediction of in-hospital and HCC-related death[146,149], of overall survival of HCC patients following surgery[144,145,147,148], and HCC recurrence following surgery or RFA[152–154]. Such models could be used to tailor individualized follow-up schedules and for patient and family counseling.

Particularly for liver transplantation, AI/ML models could be used to predict graft failure or cancer recurrence for patients transplanted for cancer. Such models could be used to reevaluate and optimize our current practices regarding recipient-donor matching for graft allocation[157–160].

**CURRENT CHALLENGES OF AI APPLICATIONS**

***Intrinsic bias and accuracy***

The level of accuracy of AI systems is mostly dependent on the quality of the training dataset. Existing biases and prejudices in the training data set will inadvertently be built into the algorithms limiting the AI/ML-based software's accuracy[161]. Discrepancies in the data collection process, imperfections of standardization, and incorrectly labeled cases become part of the algorithms' training and are thus integrated into the end product. Two of the most common biases found in these models are: (1) Spectrum bias; and (2) Overfitting. Spectrum bias occurs when the patients (whose generated data were used during the training and internal validation of these models) do not constitute a representative sample of the target population[162]. On the other hand, overfitting refers to the tendency of models to be customized for the training data[162]. The performance of the model is thus exaggerated for the training dataset but is significantly inferior in new datasets[162]. CNNs, which are extensively used in gastroenterology and hepatology, are particularly vulnerable to overfitting[26]. Another substantial bias source is that physicians often misrecord data in the electronic health records, sometimes even the chief complaint[163]. These biases significantly impact the performance of AI/ML-based models and undermine their applications. A measure to alleviate the impact of biases is the standardization of data collection methods while establishing evaluation systems that scrutinize underlying biases and check data accuracy[15,164]. Another crucial step to tackle these biases is the external validation of models, in a clinical setting, from data generated from patients prospectively enrolled in the study[165]. Initial implementation of AI/ML-based models in the clinical setting should occur on a small scale, similar to phase I and phase II of the clinical trials[166]. Existing tools, such as the PROBAST tool, could be used during the development of AI/ML-based models to comprehensively assess the risk of bias[164,167]. The American Medical Association has recently acknowledged the need to identify and address bias in data when testing or deploying AI/ML-based software to avoid introducing or exacerbating health care disparities[168]. Therefore, the real challenge is to develop an AI/ML-based software that taps into the true potential hidden by the data without picking up the biases.

Several other limitations affect the performance of these models. An example is the limited resolution of capsule endoscopy images compared to other types of endoscopy[18]. A different limitation is the in silico nature of most of the currently developed models, which significantly impacts the expectations of similar performances in a real clinical setting. Another limitation of these algorithms is that they utilize a series of variables that in the real clinical setting are derived in a series of careful decisions made by physicians and are not readily available as a complete set of data. Therefore, these variables' presence or absence could become a significant shortcoming of these models in a real clinical setting, making them impractical to use. Finally, there is a lack of consistency in the metrics used to assess performance (sensitivity, specificity, AUROC, accuracy, *etc.*), limiting the ability to draw meaningful comparisons between the models[17].

***Data protection and cybersecurity***

Similarly, with all patient data, those utilized in AI/ML-based software should conform with the seven principles described in Article 5 of the General Data Protection Regulation. Specifically, the personal data are required to: (1) Be processed fairly, lawfully, and transparently; (2) Be relevant, adequate, and limited to the intended purpose; (3) Be collected for explicit, specific, and lawful purposes; (4) Be accurate and up to date; (5) Permit identification for only as long as necessary; (6) Ensure appropriate security; and (7) Demonstrate compliance and accountability[169]. A particular challenge for AI is the resistance to the concept of utterly electronic tracking of healthcare records due to the belief that it exposes the vast amount of stored sensitive health record data to massive disclosures[170]. These concerns are not entirely unjustifiable if one considers examples such as the transfer of data from 1.6 million patient records from the Royal Free Nathional Health Service Foundation Trust to Google DeepMind, which was later ruled illegal[171]. Nevertheless, any effort or policy towards paper-based data due to data protection concerns could substantially undermine the application of AI modes in healthcare.

The healthcare industry is a particularly attractive target for cyberattacks as it contains sensitive personal data and financial information. Several physical and technical safeguards have been implemented under the Health Insurance Portability and Accountability Act to protect against the breach of sensitive patient data[172]. However, from the application of AI technology in the healthcare industry, new vulnerabilities and dangers emerge except traditional cybersecurity concerns. If ML models can learn from data, they could also be fooled by data. Data could be introduced malevolently in the algorithms to manipulate the developed AI/ML models into making wrong decisions with currently unknown ramifications to patient outcomes[173]. In a recent study, the authors demonstrated how attackers could use DL to add or remove lung cancer tumors in CT scans[174]. This study demonstrated how both a group of radiologists and a state-of-the-art deep AI model were particularly susceptible to the attack[174]. How could we, therefore, be confident that the AI/ML model has not been compromised? Evidently, healthcare facilities will have to develop additional information technology infrastructure to shield the healthcare system from these new threats.

***Intellectual property***

The Food and Drug Administration (FDA) recently acknowledged that it receives a high volume of submissions regarding AI/ML-based software marketing, with the list of already approved algorithms increasing rapidly[175,176]. A challenging point will be determining the law framework and regulatory standards that AI/ML-based software should follow. The key is classifying AI/ML software as "medical device", "service", or as "product", and achieving this requires a careful evaluation of the intentive use of AI/ML-based software. AI/ML-based software that aims to assist physicians in diagnosing, interpreting, and treatment decisions could be classified as medical devices and fall under the respective regulations[12]. In 2019, the FDA announced its aim to review AI/ML-based software regulation and has recently published the AI/ML– Based Software as a Medical Device Action Plan[175,177]. The action plan published by the FDA focuses on five pillars aimed to facilitate innovation and advance AI/ML-based software that are classified as medical devices. These include: (1) A tailored regulatory framework; (2) Good ML practice that could be achieved by consensus standards efforts; (3) A patient-centered approach incorporating transparency to users that takes into account usability, trust, equity, and accountability; (4) Regulatory science methods related to algorithm bias and robustness, and finally; and (5) Real-world performance[175]. When it comes to law and regulation, the real challenge is finding the golden snitch between too much regulation that strangles innovation and creation and too little regulation, which could have unexpected and devastating consequences for healthcare, and by extension, the well-being of patients.

Another interesting point regarding intellectual property and safety is the substantial divergence of the AI/ML-based software from the original product years after its approval and distribution[178]. What are the rights of developers on the product following its purchase? Since the original product constantly changes by learning from the clinical setting's data, the deviated model years after the purchase and the original software could be seen as two entirely different products. The first is protected under copyright law. However, the latter has now produced intellectual property on its own since it encompasses data generated by the healthcare facility[173]. Who could therefore claim legal rights over the final product? Also, who ensures the credibility and safety of the model as it deviates from the original product? Clearly, there is a need for lifecycle regulation of AI/ML-based software that ensures postapproval guardrails such as built-in audits[165]. Alternatively, time-limited authorizations could be employed to allow the FDA to perform periodic audits to review the accumulated modifications to the initial product[165].

***Liability***

Except for intellectual property, another legal challenge that arises from applying AI/ML-based software in the clinical setting is liability. A quite intriguing concept is that AI's findings and decisions could become legally binding in the future. As AI/ML-based models evolve and become more sophisticated, it is fair to assume that they will eventually surpass physicians, at least in specific tasks. How could then, the physicians justify ignoring the decision presented by AI? Especially when their decisions are made solemnly based on data and lack any sense of subjectivity. And who is liable when the followed decision made by the model causes injury? Currently, there is no legal precedent that assigns liability in a case where the injury was inflicted on a patient due to an erroneous output generated by AI/ML-based software[179].

To avoid malpractice liability, a physician is required to provide medical care at the same level as a competent physician of the same specialty while considering the available resources[180]. However, when an AI/ML-based software recommendation is involved, the concept of liability becomes more complicated. In an insightful recent legal analysis, the authors provide eight scenarios based on the combinations of whether the AI recommendation follows standard care and/or is accurate, the physician follows or rejects that recommendation, and whether a medical injury occurs[179]. The authors conclude that since current law shields physicians from liability when the standard care is followed, it also incentivizes physicians to minimize AI's actual usefulness, transforming AI into a confirmatory tool rather than a tool to augment the level of care[179]. Until a comprehensive legal framework regarding liability is developed, healthcare facilities would justifiably hesitate to adopt AI technologies due to the fear and unawareness of how they will expose the facility and its staff to liability[181].

***Ethical challenges and transparency***

AI could become the third participant in the physician-patient relationship and potentially undermine the trust between them. First, the idea that data are shared with third parties for AI model development could lead patients to withhold information from physicians and become less transparent[182]. Second, AI/ML-based models cannot act like compassionate human beings, which is an integral part of a physician's clinical life. Therefore, AI-driven decision-making neither encompasses an understanding of the patient’s needs nor respects the patient’s wishes nor demonstrates empathy, nor realizes when a patient feels discomfort or requires some rest or a hand to hold on to. Therefore, retaining the trust in the physician-patient relationship could be proven challenging in the AI era.

Another challenge is patient consent in AI applications. Clearly, it is practically impossible to acquire informed consent from each and every patient whose data are used during the development and validation of AI/ML-based software. In any case, we cannot predict how the algorithms would use specific data points during the training of the model and whether data of a specific patient have any significant impact on the model as a whole[183]. However, when AI/ML-based models are used in the clinical setting, especially for patient recording (in computer vision), patients should be adequately informed, and explicit consent should be acquired[184].

Achieving equilibrium between ensuring a high level of care and avoiding privacy encroachment could be challenging. For example, with computer vision advancement, monitoring could be used to detect any deviations from the optimal bedside practices such as patient mobilization and hand hygiene practices, which leaves patients vulnerable to identification[185,186]. Practices such as data minimization (collecting the least data needed) could address these challenges[185].

Changing the physicians' and patients’ stance towards AI technologies could be proven a herculean task. Currently, around 2/3 of the population feels uncomfortable using their data to improve care quality and are unfavorable to AI/ML-based software performing tasks typically performed by physicians[183]. There are justifiable concerns that certain biases included in the training data due to racism, sexism, and socioeconomic inequality would be integrated into the AI/ML-based software. A notorious example is the COMPAS algorithm, which was found to falsely flag black people as likely to re-offend[187]. This was aggravated when developers made the argument that the algorithm was protected under intellectual property law and thus is not open to scrutiny[187]. These examples could erroneously undermine the trust towards AI altogether. A trustworthy AI/ML-based software should be built around the principles of transparency, credibility, auditability, reliability, and recoverability[188].

The last but not least challenge in applying AI/ML-based software in healthcare is the lack of transparency, which is currently a major concern. The lack of transparency is demonstrated by the non-interpretability associated with these algorithms, described as a “black-box”, where the inner logic is hidden[18]. This violates a fundamental tenet of medical ethics that physicians should comprehend at least the basic features of the devices they use and could undermine the trustworthiness of these technologies[189]. Thus, those who develop AI/ML-based models and physicians should collaborate to reach at least a degree of explainability on how AI/ML-based algorithms reach decisions. Attention maps and saliency region are examples of methods that could lessen the lack of interpretability of these models[190]. A slightly different and intriguing point of view is that withholding the widespread application of AI/ML-based software, due to their opacity, when their application could significantly benefit patients is, in fact, unethical[183]. With most of the population being prejudiced against AI, developing trust for these technologies would require several steps, including addressing bias, increasing transparency, communicating with the patients their role in provided care, protecting data privacy, and developing a robust regulatory framework.

**CONCLUSION**

Even though multiple efforts for AI's integration into healthcare have been made, they all originate from high-income countries[191]. Understandably, since all AI/ML-based software is data-driven, the opportunities to participate in the AI ecosystem for low- and middle-income countries, where the healthcare system does not generate extensive electronic, standardized data, are significantly limited. Not being included in the development of these models would introduce a significant spectrum bias, which could undermine these models' application in low- and middle-income countries. Thus, despite being described as the key to healthcare equity, AI could become another brick in the wall of inequality. A challenge for the future would be to include low- and middle-income countries in AI model development initiatives.

Even though these AI/ML-based software's central idea is that they could surpass physicians in performing certain tasks, the existing literature comparing AI/ML-based models and physicians' performance is limited[192]. A recent meta-analysis, which included multidisciplinary comparative studies, identified only 25 studies meeting their inclusion criteria from all medical specialties[193]. The authors concluded that currently, DL models' diagnostic performance in detecting diseases from medical imaging is equivalent to that of healthcare professionals[193]. Clearly, there is a contrast between the abundance of studies developing and validating ML algorithms and the lack of studies comparing these models' performance to physicians' performance at the investigated task. Thus, there is a crystalline need for further comparative studies in the clinical setting.

Another challenge for the future would be to combine AI with other emerging fields, such as 3D printing and bioprinting, augmented reality, novel biomarkers such as microRNAs, and robotics[194–198]. Envision a world where all these technologies and their applications are unified and integrated into everyday clinical practice. AI could be the means through which this new sophisticated, complex clinical setting is handled. The future AI clinician rejects oversimplifying an inherently complex field but instead embraces the complexity.

AI can outperform physicians in the ability to precisely quantify correlations even in domains where physicians possess in-depth knowledge[173]. However, AI models are simply tools like any other and should be treated as such. Tools with their accuracy, sensitivity, and specificity in performing certain tasks, tools that carry biases, and whose findings should be evaluated in conjunction with other clinical and paraclinical findings. Reliance on AI should not exclude non-quantifiable information from decision-making[183]. Clinical reasoning and critical thinking should not be subsumed by AI at the altar of technological advancement. AI's integration into healthcare should not replace the physicians' intelligence but rather augment it. Thus, we should aim for AI-assisted and not AI-driven clinical practice[14]. Finally, AI systems should be applied in healthcare along with equally advanced evaluation systems, which properly assess their ramifications within the clinical practice environment, investigate their unintended consequences, and, most notably, evaluate their impact on patient outcomes[163]. An example of an evaluative initiative is the Digital Health Innovation Action Plan launched by the FDA that aims to assess medical software under development based on five excellence criteria: product quality, patient safety, clinical responsibility, cybersecurity responsibility, and proactive culture[199].

Despite being on the front line for several decades, AI still comes short of delivering the presumable solutions to long-standing healthcare problems. AI winters, where AI funding is significantly reduced, demonstrate the frustration that AI is not evolving at a pace investors would feel comfortable sustaining the funding[1]. AI may seem like a field that constantly overpromises but usually underdelivers, given its current impact on healthcare. In this review, we have highlighted the current challenges that we believe restrain the extensive application of AI in healthcare in an attempt to explore ways to overcome them.

Nevertheless, even at a slower pace, AI will eventually "infiltrate" not only the hospital setting but rather the healthcare industry as a whole, from central healthcare facilities to private practice and telemedicine, from medical schools and teaching hospitals to pharmaceutical companies, and even healthcare policy-makers in government. AI will become a reality; everyone will have to conform with, to avoid becoming obsolete. Therefore, sooner or later, physicians will have to engage with the field of AI by necessity. The role of physicians in this upcoming "revolution", however, remains to be seen. Will we be shaping this "revolution", or will we be mere observers?

***Limitations***

Finally, our review has few limitations. First, our study is a narrative review, and despite our best effort to follow a carefully designed search strategy to provide a comprehensive review of the current literature, our study is prone to selection bias due to its nature. Second, each study described in our study carries its own risk of bias and faces several limitations described in each study. We did not use any bias risk assessment tool to systematically evaluate if a study was worth inclusion rather than the followed search strategy and our judgment. Again this makes our review prone to selection bias. In addition, the majority of our included studies were performed in silico, and the performances reported could deviate substantially when the models are applied in a real clinical setting. Finally, with few exceptions, the majority of results report a superior or at least equivalent performance of AI/ML-based algorithms compared to the performance of conventional statistic models, widely used staging systems, extensively investigated and validated scores, and physician knowledge and reasoning, which have dominated decision-making in healthcare for decades. Thus, at least in the concept of publication bias, our findings should be interpreted with caution.

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**Table 1 Artificial intelligence applications in gastroenterology: Prevention**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **Parameters employed** | **AI classifier** | **Sizes of the training/validation sets** | **Outcomes** | **Performance** |
| Leung *et al*[27] | Laboratory results, clinicopathological parameters | Several | 64238/25330 patients | Risk of gastric cancer development following *H.pylori* eradication | 0.53-0.972,6, 59.3-98.13,6, 51.5-93.64,6 |
| Nakahira *et al*[28] | Laboratory results, clinicopathological parameters, endoscopic images | CNN | 7826/454patients | Stratify risk of gastric cancer development | --- |
| Taninaga *et al*[29] | Laboratory results, clinicopathological parameters, endoscopic images | CART | 1144/287 | Prediction of future gastric cancer | 63.4-94.81,6, 0.736-0.8742,6 |
| Goshen *et al*[31] | Laboratory results, clinicopathological parameters | DT, RF, GB | 688 flagged patients | High risk of CRC development | ---- |

1Accuracy (%).

2Area under the receiver operating curve or c-index.

3Sensitivity (%).

4Specificity (%).

5Training.

6Internal validation.

7External validation/testing. CART: Classification and regression tree; CNN: Convolutional neural network; CRC: Colorectal cancer; DT: Decision tree; GB: Gradient boosting; RF: Random forest.

**Table 2 Artificial intelligence applications in gastroenterology: Diagnosis**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **Diagnostic Modality** | **AI classifier** | **Sizes of the training/validation sets** | **Outcomes** | **Performance** |
| Takiyama *et al*[33] | Esophago-gastro-duodenoscopy imaging | CNN | 1750/4357 | Anatomical classification among larynx, esophagus, stomach, and duodenum | 0.99-1.002,7 |
| Pace *et al*[34] | Laboratory results, clinicopathological parameters | ANN | 159 patients | Diagnosis of gastroesophageal reflux disease | 67.86-1001,6 |
| de Groof *et al*[35] | Esophageal endoscopic images | DNN | 1247/2976/807/807 patients | Classification of malignant from nondysplastic Barret’s esophagus | 88.21,6, 87.5-88.81,7, 87.63,6, 90.0-92.53,7, 88.64,6, 82.5-87.54,7 |
| van der Sommen *et al*[36] | White-light endoscopic imaging | SVM | 44 patients with Barret’s esophagus | Diagnosis of early neoplastic lesions | Per image: 62-903,6, 65-904,6,Per patient: 52-1003,6, 74-964,6 |
| Struyvenberg *et al*[37] | Volumetric laser endomicroscopy imaging | Several | 29 patients with Barret’s esophagus | Diagnosis of neoplastic lesions | 0.83-0.942,6 |
| Swager *et al*[38] | Volumetric laser endomicroscopy imaging | Several | 60 images | Diagnosis of neoplastic lesions | 0.89-0.952,6 |
| Kumagai *et al*[39] | Endocytoscopic imaging | CNN | 4715/15207 | Diagnosis of esophageal squamous cell carcinoma | 90.91,7, 0.72-0.902,7, 39.4-46.43,7, 98.2-98.44,7 |
| Zheng *et al*[40] | Endoscopic images | CNN | 1507/452 patients | Diagnosis of *H.pylori* infection | 84.5-93.81,6, 0.93-0.972,6, 81.4-91.63,6, 90.1-98.64,6 |
| Nakashima *et al*[41] | Endoscopic images | CNN | 162/60 patients | Diagnosis of *H.pylori* infection | 0.66-0.962,6 |
| Itoh *et al*[42] | Endoscopic images | CNN | 149/30 images | Diagnosis of *H.pylori* infection | 0.9562,6, 86.73,6, 86.74,6 |
| Shichijo *et al*[43] | Endoscopic images | CNN | 32308/114817 | Diagnosis of *H.pylori* infection | 83.1-87.71,7, 81.9-88.93,7, 83.4-87.44,7 |
| Kanesaka *et al*[45] | NBI | SVM | 126/81 NBI images | Diagnosis of gastric cancer | 96.31,6, 96.73,6, 95.04,6 |
| Hirasawa *et al*[46] | Endoscopic images | CNN | 13584/22967 | Diagnosis of gastric cancer | 92.23,7 |
| Zhu *et al*[47] | Laboratory results, clinicopathological parameters, cancer biomarkers | GB/DT | 496/213 patients | Diagnosis of gastric cancer | 85.91,5, 831,6, 0.912,6, 883,5, 873,6, 83.44,5, 84.14,6 |
| Tenório *et al*[48] | Laboratory results, clinicopathological parameters | Several | 178/38 | Diagnosis of celiac disease | 71.5-801,6, 0.71-0.842,6, 69-823,6, 67-804,6 |
| Caetano Dos Santos *et al*[49] | Endomysial autoantibody test for IgA-class antibodies images | SVM | 2597 images (training:validation = 7:3) | Diagnosis of celiac disease | 96.8-98.851,6, 82.84-98.913,6, 98.81-99.404,6 |
| Hujoel *et al*[50] | Laboratory results, clinicopathological parameters | Several | 408 undiagnosed patients | Diagnosis of celiac disease | 0.49-0.532,6 |
| Manandhar *et al*[51] | Gut microbiome data | RF | 1429 fecal 16S metagenomic data subjects | Diagnosis of IBD | 0.80-0.822,6 |
| Wei *et al*[52] | Single nucleotide polymorphisms data | Several | 60828 samples | Classifification of CD and UC | 0.782-0.8662,6 |
| Mossotto *et al*[53] | Capsule endoscopy, histologic imaging | SVM | 239/487 pediatric patients | Classifification of CD, UC, and unclassified IBD | 71-82.71,5, 0.78-0.872,5, 83.31,7, 83-853,7 |
| Xia *et al*[58] | Capsule endoscopy imaging | CNN | 697/1007 patients, 822590/2013657, images | Classification among different types of gastric lesions | 77.1-861,7, 0.80-0.902,7, 96.2-1003,7, 56.5-76.24,7 |
| Seguí *et al*[59] | Capsule endoscopy imaging | CNN | 50 videos | Classification of small bowel mobility events | 961,6 |
| Park *et al*[60] | Capsule endoscopy imaging | CNN | 139 videos, 200000 images (training:validation: test = 6:2:2) | Small bowel lesion identification | 80.29-98.341,6, 0.9992,5, 0.9982,6,7 |
| Hwang *et al*[61] | Capsule endoscopy imaging | CNN | 7556/57607 images | Classification of hemorrhagic and ulcerative lesions | 96.62-96.831,7, 95.07-97.613,7, 96.04-98.184,7 |
| Otani *et al*[62] | Capsule endoscopy imaging | DNN | 167/407 patients | Classification among different types of small bowel lesions | 0.950-0.9962,6, 0.884-0.9282,7 |
| Yuan *et al*[63] | Capsule endoscopy imaging | SVM | 20 patients, 340 images (training:validation = 8:2) | Diagnosis of peptic ulcers | 92.651,6, 94.123,6, 91.184,6 |
| Karargyris *et al*[64] | Capsule endoscopy imaging | SVM | 80 frames | Diagnosis of peptic ulcers | 753,6, 73.34,6 |
| He *et al*[65] | Capsule endoscopy imaging | CNN | 11 patients, 440000 images | Diagnosis of intestinal hookworms | 88.51,6, 0.8952,6, 84.63,6, 88.64,6 |
| Leenhardt *et al*[66] | Capsule endoscopy imaging | CNN | 600/600 images | Diagnosis of gastrointestinal angiectasia | 1003,6, 964,6 |
| Zhou *et al*[67] | Capsule endoscopy imaging | CNN | 21 videos | Diagnosis of celiac disease | 1003,6, 1004,6 |
| Yamada *et al*[68] | Colon capsule endoscopy imaging | CNN | 15933/47847 | Diagnosis of colorectal neoplasias | 83.97, 0.9022,7, 793,7, 874,7 |
| Wang *et al*[69] | Colonoscopy imaging | CNN | 5545 images/271137 images/6127 images/1387 videos/547 videos | Identification of colorectal polyps | 0.9842,7, 88.24-1003,7, 95.40-95.922,7 |
| Misawa *et al*[70] | Colonoscopy imaging | CNN | 411/35 short videos | Identification of colorectal polyps | 76.51,6, 0.872,6, 903,6, 63.34,6 |
| Urban G *et al*[71] | Colonoscopy imaging | CNN | 8641 images/207 videos | Identification of colorectal polyps | 96.41,7, 0.9912,7 |
| Ozawa *et al*[72] | Colonoscopy imaging | CNN | 20431/70777 images | Identification of colorectal polyps, Classification of colorectal polyps | 90-973,7, 47-981,7 |
| Mori *et al*[73] | NBI and methylene blue staining images | SVM | 466 diminutive polyps | Classification of diminutive rectosigmoid adenomas | NPV(%): 93.7-96.5 |
| Tischendorf *et al*[74] | NBI | SVM | 209 colorectal polyps | Classification of colorectal polyps | 903,6, 70.24,6 |
| Gross *et al*[75] | NBI | SVM | 434 colorectal polyps | Classification of small colorectal polyps | 93.11,6, 95.03,6, 90.34,6 |
| Kominami *et al*[76] | NBI | SVM | 118 colorectal polyps | Classification of colorectal polyps | 93.21,6, 93.03,6, 93.34,6 |
| Misawa *et al*[77] | NBI endocytoscopy | SVM | 979/100 endocytoscopy, images | Classification of colorectal polyps | 901,6, 84.53,6, 97.64,6 |
| Takeda *et al*[78] | NBI endocytoscopy | SVM | 5543/200 endocytoscopy, images | Diagnosis of invasive CRC | 94.11,6, 89.43,6, 98.94,6 |
| Chen *et al*[79] | NBI | CNN | 2157/2847 | Classification neoplastic from hyperplastic polyps | 96.33,7, 78.14,7, NPV(%): 91.57 |
| Komeda *et al*[80] | NBI | CNN | 1200/600 images | Classification of adenomatous from non-adenomatous polyps | 75.11,6 |
| Byrne *et al*[81] | NBI | CNN | 223/407 videos | Classification of adenomas from hyperplastic polyps | 941,7, 0.952,7, 983,7, 834,7, NPV(%): 977 |

1Accuracy (%).

2Area under the receiver operating curve or c-index.

3Sensitivity (%).

4Specificity (%).

5Training.

6Internal validation.

7External validation/testing. ANN: Artificial neural network; CD: Chron’s disease; CNN: Convolutional neural network; CRC: Colorectal cancer; DNN: Deep neural network; DT: Decision tree; GB: Gradient boosting; IBD: Inflammatory bowel disease; NBI: Narrow-band imaging; NPV: Negative predictive value; RF: Random forest; SVM: Support vector machine; UC: Ulcerative colitis.

**Table 3 Artificial intelligence applications in gastroenterology: Treatment**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **Parameters employed** | **AI classifier** | **Sizes of the training/validation sets** | **Outcomes** | **Performance** |
| Rogers*et al*[86] | Data from baseline impedance, nocturnal baseline impedance, and acid exposure time | DT | 335 patients | Prediction of treatment response with proton pump inhibitors for patients with gastroesophageal reflux disease | 0.31-0.9382,6 |
| Zhu *et al*[87] | Endoscopic images | CNN | 790/2037 images | Invasion of gastric cancer at the mucosa and submucosa layers of the stomach | 89.161,7, 0.942,7, 76.473,7, 95.564,7 |
| Kubota *et al*[88] | Endoscopic images | DNN | 800/90 images | Invasion depth of gastric cancer | 64.71,6 |
| Yamashita *et al*[89] | Hematoxylin and eosin-stained WSI | DNN | 100/156/4847 | Identificication of CRC microsatellite instability | 0.9312,6, 0.7792,7, 763,7, 66.64,7 |
| Ichimasa *et al*[90] | Laboratory results, clinicopathological parameters | SVM | 590/1007 | Prediction of lymph node metastasis status | 691,7, 0.8212,7, 1003,7, 664,7 |
| Levi *et al*[91] | Laboratory results, clinicopathological parameters | RFE | 14620 patients | Prediction of the need for transfusion following GIB | 50.21-74.881,6, 0.7858-0.81412,6, 69.17-92.773,6, 35.02-79.824,6 |
| Chu *et al*[92] | Laboratory results, clinicopathological parameters | Several | 122/67 patients | Prediction of the source of GIB | 69.7-94.31,6, 0.658-0.9992,6, 90.1-98.03,6, 89-1004,6 |
| Prediction of the need for blood resuscitatio | 64.7-94.11,6, 0.381-0.9932,6, 90.3-93.93,6, 18.4-95.54,6 |
| Prediction of the need for emergent endoscopy | 62.7-83.31,6, 0.404-0.9132,6, 80.1-89.13,6, 13.8-85.74,6 |
| Prediction of disposition | 58.4-89.71,6, 0.324-0.9722,6, 81.9-92.93,6, 18.4-90.94,6 |
| Das *et al*[93] | Laboratory results, clinicopathological parameters | ANN | 194/1936/2007 patients | Prediction of major stigmata of recent hemorrhage | 891,3,4,6, 771,7, 963,7, 634,7 |
| Prediction of the need for emergent endoscopy | 811,3,6, 611,7, 943,6, 824,6, 484,7 |
| Augustin *et al*[94] | Laboratory results, clinicopathological parameters | CART | 164/1037 patients | Stratification of risk of rebleeding and mortality following acute variceal hemorrhage | 0.81-0.832,7 |
| Loftus *et al*[95] | Laboratory results, clinicopathological parameters | ANN | 103/44 patients | Prediction of severe lower GIB | 0.9792 |
| Prediction of the need for surgical intervention | 0.9542,6 |
| Ayaru *et al*[96] | Laboratory results, clinicopathological parameters | GB | 170/1307 | Prediction of severe lower GIB | 781,6, 831,7 |
| Prediction of recurrent bleeding | 881,6, 881,7 |
| Prediction of the need for intervention | 881,6, 911,7 |

1Accuracy (%).

2Area under the receiver operating curve or c-index.

3Sensitivity (%).

4Specificity (%).

5Training.

6Internal validation.

7External validation/testing.ANN: Artificial neural network; CART: Classification and regression tree; CNN: Convolutional neural network; CRC: Colorectal cancer; DT: Decision tree; DNN: Deep neural network; GB: Gradient boosting; GIB: Gastrointestinal bleeding; RFE: Recursive feature elimination; WSI: Whole-slide image.

**Table 4 Artificial intelligence applications in gastroenterology: Prognosis**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **Parameters employed** | **AI classifier** | **Sizes of the training/validation sets** | **Outcomes** | **Performance** |
| Yang *et al*[97] | Laboratory results, immunomarkers, clinicopathological parameters | SVM | 319/164 patients | Distant metastasis of oesophageal squamous cell carcinoma following surgery | 69.5-80.11,6, 44.7-67.23,6, 81.6-97.74,6 |
| Sato *et al*[98] | Laboratory results, clinicopathological parameters, tumor characteristics | ANN | 395 patients (training:validation: test = 53:27:20) | 1-year and 5-year survival of patients with esophageal cancer following surgery | 0.883-0.8842,7, 78.1-80.73,7, 84.7-86.54,7 |
| Zhou *et al*[99] | Laboratory results, clinicopathological parameters, tumor characteristics | Several | 2012 patients (training:validation = 8:2) | Recurrence of gastric cancer following surgery | 0.790-0.9622,5, 0.771-0.7952,6 |
| Peng *et al*[100] | Meteorological data | ANN | 901 patients | Variations of onset and relapse of IBDs | ---- |
| Hardalaç *et al*[101] | Clinicopathological parameters, treatment data | ANN | 129 patients (training:validation:test = 80:10:10) | Prediction of mucosal remission for CD patients treated with azathioprine | 58.1-79.11,6, 0.527-0.8832,6 |
| Takayama *et al*[102] | Clinicopathological parameters, treatment data | ANN | 54/36 patients | Prediction of the need for operation for UC patients treated with cytoapheresis | 963,6, 974,6 |
| Lyles *et al*[103] | Laboratory results, clinicopathological parameters | CART | 884 patients | Prediction of in-hospital mortality of upper GIB in cirrhotic patients | ---- |
| Grossi *et al*[104] | Laboratory results, clinicopathological parameters | ANN | 807 patients | 30-d mortality of patients with non-variceal upper GIB | 81.2-89.01,6, 0.872,6, 81.5-93.33,6, 80.9-84.74,6 |
| Rotondano *et al*[105] | Laboratory results, clinicopathological parameters | ANN | 2380 patients | 30-d mortality of patients with non-variceal upper GIB | 96.81,6, 0.952,6, 83.83,6, 97.54,6 |
| Shi *et al*[106] | CT radiomics | Several | 124/35 patients | Prediction of the presence of RAS and BRAF mutations in CRC | ANN: 871,5, 711,6, 0.90-0.952,5, 0.792,6 |
| Kang *et al*[107] | Laboratory results, immunomarkers, clinicopathological parameters, tumor characteristics | LASSO | 221/95 patients | Prediction of lymph node metastasis status in operated patients for T1 CRC | 0.7952,5, 0.7652,6 |

1Accuracy (%).

2Area under the receiver operating curve or c-index.

3Sensitivity (%).

4Specificity (%).

5Training.

6Internal validation.

7External validation/testing. ANN: Artificial neural network; CART: Classification and regression tree; CD: Chron’s disease; CRC: Colorectal cancer; CT: Computed tomography; GIB: Gastrointestinal bleeding; IBD: Inflammatory bowel disease; LASSO: Least absolute shrinkage and selection operator; SVM: Support vector machine; UC: Ulcerative colitis.

**Table 5 Artificial intelligence applications in hepatology: Prevention**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **Parameters employed** | **AI classifier** | **Sizes of the training/validation sets** | **Outcomes** | **Performance** |
| Goldman *et al*[108] | National database of routine annual health check-ups | DT-based | 12019 patients | Risk of NAFLD and cirrhosis | 84.50-85.731,6, 0.7740-0.84862,6 |
| Yip *et al*[109] | Laboratory results, clinicopathological parameters | Several | 500/422 | Identify patients with NAFLD | 0.87-0.902,5, 0.78-0.882,6, 55.48-94.523,5, 51.69-92.373,6, 58.47-91.534,5, 50.99-90.464,6 |
| Ma *et al*[110] | Laboratory results, clinicopathological parameters | Several | 10508 patients (training:validation = 9:1) | Identify patients with NAFLD | 49.47-82.921,6, 20.2-68.03,6, 54.4-94.64,6 |
| Sowa *et al*[111] | Laboratory results, clinicopathological parameters | Several | 126 morbidly obese patients (training:validation = 9:1) | Fibrosis in NAFLD patients | 791,6, 30.8-60.03,6, 77.0-92.24,6 |
| Canbay *et al*[112] | Laboratory results, clinicopathological parameters | EFS | 164/122obese patients | Classification of NAFLD and NASH | 0.73392,5, 0.70282,6 |
| Fialoke *et al*[113] | National database of routine annual health check-ups | Several | 108139 patients (training:validation = 4:1) | Classification among healthy, NAFLD, and NASH | 77.2-79.71,6, 0.842-0.8762,6, 74.5-77.43,6 |
| Sowa *et al*[114] | Data from biochemical and enzyme-linked immunosorbent assays | Several | 133 patients (training:validation = 9:1) | Classification of NAFLD and ALD | DT: 89.02-95.11,6, 74.19-94.123,6, 96.08-98.044,6,RF: 0.8932-0.98462,6, SVM: 0.9058-0.91182,6 |
| Wei *et al*[115] | Laboratory results, clinicopathological parameters | GB | 576 HBV patients, (training:validation = 7:3), 3687 HCV patients | Classification of fibrosis/cirrhosis in HBV patients | 0.904-0.9742,5, 0.871-0.9182,6, 79-883,5, 78-843,6, 86-924,5, 854,6 |
| Classification of fibrosis/cirrhosis in HCV patients | 0.797-0.8492,7 |
| Wang *et al*[116] | Laboratory results, clinicopathological parameters | ANN | 226/1136/1167 HBV patients | Classification of significant fibrosis | 0.8832,5, 0.8842,6, 0.9202,7 |
| Raoufy *et al*[117] | Laboratory results, clinicopathological parameters | ANN | 86/58 HBV patients | Classification of liver cirrhosis | 91.381,6, 0.8982,6, 87.53,6, 924,6 |
| Piscaglia *et al*[118] | Laboratory results, clinicopathological parameters | ANN | 414/96 HCV patients | Classification of significant fibrosis | 45.8-86.51,6, 0.872,5, 0.932,6, 30.4-1003,6, 30.1-98.64,6 |
| Hashem *et al*[119] | Laboratory results, clinicopathological parameters | Several | 22690/16877 HCV patients | Classification of significant fibrosis | 66.3-84.41,6, 0.73-0.762,6 |
| Ioannou *et al*[120] | Clinical/laboratory data extracted directly from electronic health records | DNN | 48151 patients with HCV-related cirrhosis (training:validation = 9:1) | HCC development in HCV cirrhosis | 0.759-0.8062,6 |
| Emu *et al*[121] | Laboratory results, clinicopathological parameters | Several | 1385 patients HCV (training:validation = 4:1) | Stage of liver cirrhosis | 97.228-97.8311,6 |

1Accuracy (%).

2Area under the receiver operating curve or c-index.

3Sensitivity (%).

4Specificity (%).

5Training.

6Internal validation.

7External validation/testing. ALD: Alcoholic liver disease; ANN: Artificial neural network; DNN: Deep neural network; DT: Decision tree; EFS: Ensemble feature selection; GB: Gradient boosting; HBV: Hepatitis B; HCC: Hepatocellular carcinoma; HCV: Hepatitis C; NAFLD: Non-alcoholic fatty liver disease; NASH: Non-alcoholic steatohepatitis RF: Random forest; SVM: Support vector machine.

**Table 6 Artificial intelligence applications in hepatology: Diagnosis**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **Diagnostic Modality** | **AI classifier** | **Sizes of the training/validation sets** | **Outcomes** | **Performance** |
| Choi *et al*[122] | CT imaging | CNN | 7461/4216/2987/1727 patients | Liver fibrosis staging (F0-F4) | 83.11,5, 80.81,6, 74.4-80.21,7 |
| Classification among significant fibrosis, advanced fibrosis, and cirrhosis | 92.1-95.01,6,7, 0.95-0.972,6,7, 84.6-95.53,6,7, 89.9-96.64,6,7 |
| Kuppili *et al*[123] | US imaging | ELM, SVM | 63 patients | Diagnosis of FLD | ELM: 81.7-92.41,6, 0.81-0.922,6, 85.10-91.303,6, 78.52-92.104,6, SVM: 76.14-86.421,6, 0.74-0.862,6, 76.80-88.203,6, 74.52-86.304,6 |
| Gatos *et al*[124] | US shear wave elastography imaging | SVM | 126 patients | Classification of chronic liver disease from healthy patients | 87.31,6, 0.872,6, 93.53,6, 81.24,6 |
| Chen *et al*[125] | Real-time tissue elastography imaging, age, sex | Several | 513 patient (training:validation = 3:1) | Classification of liver fibrosis | 80.44-82.871,6, 79.67-92.973,6, 46.25-82.504,6 |
| Matake *et al*[126] | Clinicopathological parameters, CT imaging | ANN | 120 patients | Classification among four types of focal liver lesions | 0.9612,6 |
| Oestmann *et al*[127] | Multiphasic MRI scans | CNN | 150/10patients | Classification of HCC and non-HCC lesions | 94.11,5, 87.31,6, 0.9122,6 |
| For HCC: 92.73,6, 82.04,6 |
| For non-HCC: 82.03,6, 92.74,6 |
| Kim *et al*[128] | MRI scans | CNN | 4555,6/547 patients | HCC detection | 0.972,6, 943,6, 994,6, 0.902,7, 873,7, 934,7 |
| Cucchetti *et al*[129] | Laboratory results, clinicopathological parameters, radiological data, histological data | ANN | 175/75patients | MVI | 0.922,5, 91.01,6 |
| Histopathological Grade | 0.942,5, 93.31,6 |
| Urman *et al*[130] | Metabolomic and proteomic analyses of bile | Several | 139 patients | Classification of CCA and pancreatic adenocarcinoma | 0.98-1.002,6, 88-94.13,6, 92.3-1004,6 |
| Negrini *et al*[131] | Plasma bile acids profiles | Several | 112 patients (training:validation = 4:1) | Classification of CCA and benign biliary disease | 68.2-86.41,6, 0.77-0.952,6, 64-793,6, 63-1004,6 |
| Logeswaran[132] | MRCP | MLP | 55/5937 images | CCA diagnosis | 92.8-96.31,6, 83.64-90.141,7 |

1Accuracy (%).

2Area under the receiver operating curve or c-index.

3Sensitivity (%).

4Specificity (%).

5Training.

6Internal validation.

7External validation/testing. ANN: Artificial neural network; CCA: Cholangiocarcinoma; CNN: Convolutional neural network; CT: Computed tomography; ELM: Extreme learning machine; FLD: Fatty liver disease; HCC: Hepatocellular carcinoma; MLP: Multi-layer perceptron; MRCP: Magnetic resonance cholangiopancreatography MRI: Magnetic resonance imaging; MVI: Microvascular invasion; SVM: Support vector machine; US: Ultrasound.

**Table 7 Artificial intelligence applications in hepatology: Treatment**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **Parameters employed** | **AI classifier** | **Sizes of the training/validation sets** | **Outcomes** | **Performance** |
| Wübbolding *et al*[133] | Analyze soluble immune markers | Several | 28/497 HBV patients | Prediction of early virological relapse | 0.73-0.892,6, 0.59-0.672,7 |
| Haga *et al*[134] | WGS of HCV | Several | 86/87 HCV patients | Classification of HCV variants resistant to antiviral drugs | 0.5-0.9372,5, 0.597-0.9542,6 |
| Bedon *et al*[135] | DNA methylation profiling | RF-based | 300/74 HCC specimens | 6-mo progression-free survival | 67.1-80.61,5, 64.8-80.21,7 |
| Tsilimigras *et al*[137] | Laboratory results, clinicopathological parameters, tumor characteristics | CART | 976 HCC patients | Determining factors of prognostic weigh preoperatively within the BCLC staging system | --- |
| Tsilimigras *et al*[139] | Laboratory results, clinicopathological parameters, tumor characteristics | CART | 1146 CCA patients | Determining factors of prognostic weigh preoperatively | --- |
| Jeong *et al*[140] | Laboratory results, clinicopathological parameters | DNN | 1421/2347 | Intrahepatic CCA susceptible to adjuvant therapy following resection | 0.842,5, 0.782,7 |
| Shao *et al*[141] | Clinicopathological parameters | ANN | 288 CCA patients (training:validation = 8:2) | Predict early occlusion following bilateral plastic stent placement | 0.96482,5, 0.95442,6 |

1Accuracy (%).

2Area under the receiver operating curve or c-index.

3Sensitivity (%).

4Specificity (%).

5Training.

6Internal validation.

7External validation/testing. ANN: Artificial neural network; BCLC: Barcelona clinic liver cancer; CART: Classification and regression tree; CCA: Cholangiocarcinoma; DNN: Deep neural network; HCC: Hepatocellular carcinoma; HCV: Hepatitis C; RF: Random forest; WGS: Whole-genome sequencing.

**Table 8 Artificial intelligence applications in hepatology: Prognosis**

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| **Ref.** | **Parameters employed** | **AI classifier** | **Sizes of the training/validation sets** | **Outcomes** | **Performance** |
| Hong *et al*[142] | Laboratory results, clinicopathological parameters | ANN | 197 HBV patients (training:validation = 4:1) | Development of esophageal varices in HBV cirrhosis | 87.821,6, 93.753,6, 71.704,6 |
| Dong *et al*[143] | Laboratory results, clinicopathological parameters | RF | 238/1097 | Identification of esophageal varices | 0.842,5, 0.822,7 |
| Classification of esophageal varices requiring treatment | 0.742,5, 0.752,7 |
| Ho *et al*[144] | Laboratory results, clinicopathological parameters, surgery parameters | ANN, DT | 427, 354, and 297 patients for 1-, 3-, and 5-year survival (training:validation = 8:2) | 1-, 3-, and 5-year disease-free survival | ANN: 0.963-0.9892,5, 93.5-96.33,5, 91.6-97.94,5, 0.774-0.8642,6, 70.0-78.73,6, 54.2-92.74,6 |
| Following surgical resection |
| DT: 0.675-0.8252,5, 19.6-94.83,5, 45.8-97.94,5, 0.561-0.7182,6, 0-88.53,6, 37.5-96.44,6 |
| Shi *et al*[145] | Laboratory results, clinicopathological parameters, tumor characteristics | ANN | 22926 patients | 5-year survival following surgical resection | 96.571,6, 0.8852,6, 97.431,7, 0.8712,7, 74.233,7, |
| Shi *et al*[146] | Laboratory results, clinicopathological parameters, surgery parameters | ANN | 22926 hepatectomies | In-hospital mortality following surgical resection | 97.281,6, 0.842,6, 95.931,7, 0.822,7, 78.403,7, 94.574,7 |
| Chiu *et al*[147] | Laboratory results, clinicopathological parameters, tumor characteristics | ANN | 434, 341, and 264 patients for 1-, 3-, and 5-year survival, (training:validation = 8:2) | 1-, 3-, and 5-year overall survival, following surgical resection | 98.5-99.51,5, 0.980-0.9932,5, 99.7-1003,5, 96.2-99.24,5, 72.1-85.11,6, 0.798-0.8752,6, 71.4-88.63,6, 50.0-82.14,6 |
| Qiao *et al*[148] | Laboratory results, clinicopathological parameters, tumor characteristics | ANN | 362/1816/1047 patients | Survival following surgical resection | 0.8552,5, 80.003,5, 73.404,5, 0.8322,6, 78.673,6, 75.704,6, 0.8292,7, 77.423,7, 78.084,7 |
| Liu *et al*[149] | Laboratory results, data from immunochemistry of peripheral blood mononuclear cells, tumor characteristics | GB survival classifier | 136/566/1057 | Risk of HCC-related death | 0.8442,5, 0.8272,6, 0.8062,7 |
| Zhong *et al*[150] | ALBI/CTP stage | ANN | 319 / 617 / 1247 | Survival of patients treated with chemoembolization and sorafenib | ALBI-based: 0.7162,7, 0.8232,7 |
| CTP-based: 0.7792,7, 0.6932,7 |
| Divya and Radha[152] | Laboratory results, clinicopathological parameters, tumor characteristics | APO, SVM, RF | 152 patients | Recurrence following RFA | 95.51,6, 95.13,6, 95.84,6 |
| Yamashita *et al*[153] | Hematoxylin and eosin-stained WSI | CNN | 299 / 536/1987 WSIs | Recurrence following Surgical Resection | 0.7242,6, 0.6832,7 |
| Liang *et al*[154] | Laboratory results, clinicopathological parameters | SVM | 83 patients | Recurrence following RFA | 73-821,6, 0.60-0.692,6, 77-863,6, 73-824,6 |
| Eaton *et al*[155] | Laboratory results, clinicopathological parameters | GB-based | 509/278 patients with primary sclerosing cholangitis | Classify risk of primary sclerosing cholangitis-related complications | 0.962,6, 0.902,7 |
| Andres *et al*[156] | Laboratory results, clinicopathological parameters, donor characteristics | PSSP system | 2769 patients | Survival following transplantation for primary sclerosing cholangitis | ---- |
| Rodriguez-Luna *et al*[157] | Genotyping data from microsatellite mutations/deletions | ANN | 19 transplated patients | Post-transplant HCC recurrence | 89.51,6 |
| Lau *et al*[158] | Laboratory results, clinicopathological parameters, donor characteristics | ANN, RF | 90/90 transplants | Graft failure/primary nonfunction | ANN: 0.734-0.8352,6 |
| RF: 0.787-0.8182,6 |
| 3-mo graft failure | ANN: 0.5592,6, R6: 0.7152,6 |
| Briceño *et al*[160] | Laboratory results, clinicopathological parameters, surgical parameters, donor characteristics | ANN | 1003 liver transplants | 3-mo graft failure | 0.806-0.8212,6 |

1 Accuracy (%).

2Area under the receiver operating curve or c-index.

3Sensitivity (%).

4Specificity (%).

5Training.

6Internal validation.

7External validation/testing. ALBI: Albumin-bilirubin; ANN: Artificial neural network; APO: Artificial plant optimization; CNN: Convolutional neural network; CTP: Child–Turcotte–Pugh; DT: Decision tree; GB: Gradient boosting; HBV: Hepatitis B; HCC: Hepatocellular carcinoma; PSSP: Patient-specific survival prediction; RF: Random forest; RFA: Radiofrequency ablation; SVM: Support vector machine; WSI: Whole-slide image.



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