Artificial intelligence in clinical care amidst COVID-19 pandemic: A systematic review

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Abstract

The worldwide health crisis caused by the SARS-Cov-2 virus has resulted in >3 million deaths so far. Improving early screening, diagnosis and prognosis of the disease are critical steps in assisting healthcare professionals to save lives during this pandemic. Since WHO declared the COVID-19 outbreak as a pandemic, several studies have been conducted using Artificial Intelligence techniques to optimize these steps on clinical settings in terms of quality, accuracy and most importantly time. The objective of this study is to conduct a systematic literature review on published and preprint reports of Artificial Intelligence models developed and validated for screening, diagnosis and prognosis of the coronavirus disease 2019. We included 101 studies, published from January 1st, 2020 to December 30th, 2020, that developed AI prediction models which can be applied in the clinical setting. We identified in total 14 models for screening, 38 diagnostic models for detecting COVID-19 and 50 prognostic models for predicting ICU need, ventilator need, mortality risk, severity assessment or hospital length stay. Moreover, 43 studies were based on medical imaging and 58 studies on the use of clinical parameters, laboratory results or demographic features. Several heterogeneous predictors derived from multimodal data were identified. Analysis of these multimodal data, captured from various sources, in terms of prominence for each category of the included studies, was performed. Finally, Risk of Bias (RoB) analysis was also conducted to examine the applicability of the included studies in the clinical setting and assist healthcare providers, guideline developers, and policymakers.

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1. Introduction

The World Health Organization (WHO) declared on March 11th, 2020 the COVID-19 outbreak, emerged in December 2019 in Wuhan, China [1] resulting at the time of writing in more than 3 million deaths and 150 million cases worldwide. The most critical steps in assisting healthcare professionals to save lives during this pandemic are early screening, diagnosis and prognosis of the disease. Several studies have been conducted using Artificial Intelligence (AI) techniques to optimize these steps on clinical settings in terms of quality, accuracy and time. AI techniques, employing Deep Learning (DL) methods, have demonstrated great success in the medical imaging domain due to DL’s advanced capability for feature extraction [2]. Apart from the medical imaging domain, AI techniques are widely used to screen, diagnose and predict prognosis of COVID-19 based on clinical, laboratory and demographic data.

Early clinical course of SARS-CoV2 infection can be difficult to distinguish from other undifferentiated medical presentations to hospital and SARS-CoV-2 PCR testing can take up to 48 h for operational reasons. Limitations of the gold-standard PCR test for COVID-19 have challenged healthcare systems across the world due to shortages of specialist equipment and operators, relatively low test sensitivity and prolonged turnaround times [3]. Hence, rapid identification of COVID-19 is important for delivering care, aiding proper triage among patients admitting to hospitals, accelerating proper treatment and minimizing the risk of infection during presentation and waiting hospital admission time. Several studies have been conducted to face the need of early screening by using AI methods [4,5].

Challenges on COVID-19 diagnosis are also present due to the difficulties of differentiating Chest X-Ray radiographs (CXRs) with COVID-19 pneumonia symptoms from those with common pneumonia and insufficient empirical understanding of the radiological morphology in CT scans of this new type of pneumonia among other. Moreover, CXR or CT-based diagnosis may need laboratory confirmation. Therefore, there is an imperative demand for accurate methods to assist clinical diagnosis of COVID-19. Multiple studies using AI techniques have been conducted towards this direction, to extract valuable features from CXRs or CTs [6,7], to use clinical data and laboratory exams [8,9] or to even combine both imaging quantitative features and clinical data to result in accurate diagnosis [10,11]. Finally, prognosis is an essential step towards assisting healthcare professionals to predict ICU need,
mechanical ventilator need, hospitalization time, mortality risk or severity assessment of the disease.

The objective of our study was to conduct a systematic literature review on published and preprint reports of Artificial Intelligence techniques developed and validated for screening, diagnosis, and prognosis of the coronavirus disease 2019. Studies that developed AI prediction models for screening, diagnosis or prognosis that can be applied in the clinical setting were included (see Fig. 1). Screening studies describe prediction models developed for early identification of COVID-19 infection, whereas diagnostic studies propose prediction models developed to establish a diagnosis of the disease. In these studies, several predictors were recognized, including clinical parameters (e.g., comorbidities, symptoms) laboratory results (e.g., hematological, biochemical tests), demographic features (e.g., age, sex, province, country, travel history) or imaging features extracted from CT scans or CXRs. Identification of the most prominent predictors was also part of our analysis. Furthermore, novel technologies incorporated in AI techniques were investigated to determine the current state of research in developing AI prediction models. Additionally, the advantages of using imaging, clinical and laboratory data or the combination of those were analyzed. To achieve this objective, each study was analyzed in terms of COVID-19 positive patients included in the primary datasets, AI methods employed, predictors identified, validation methods applied, and performance metrics used. Finally, a Risk of Bias (RoB) analysis was conducted to examine the applicability of the included studies in the clinical setting and support decisions made by healthcare providers, guideline developers, and policymakers.

The paper is organized as follows. In Section 2, we describe the used methods in the approach and protocols including the description of the AI algorithms performance metrics, and the inclusion–exclusion criteria of the reviewed studies. In Section 3, results are presented on the primary datasets, AI algorithms, validation methods, as well as prediction models developed for screening, diagnostic and prognostic purposes. In this section, we also provide results on the most prominent predictors for each category of the included prediction models. Moreover, the results of the RoB assessment, are provided in Section 3. In Section 4, we discuss the results and the limitations related to the applicability of the developed prediction models and we identify possible future directions aiming at enhancing the adoption of AI-based prediction models in clinical practice.

2. Methods

A. Review approach and protocols

In this systematic literature review, we followed the guidelines of the Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) protocol to ensure transparent and complete reporting (see Fig. 2) [12]. This study focused on peer-reviewed publications, as well as preprints published in English, that applied AI techniques to develop prediction models for diagnosis or prognosis of COVID-19. A systematic literature search was conducted for collecting research articles available from January 2020 through December 2020, using the online databases PubMed, Nature, Science Direct, IEEE Xplore, Arxiv and medRxiv. By combining appropriate keywords with Boolean operators, the following expression was formed:

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[('artificial intelligence' OR 'AI') OR ('machine learning' OR 'ML') OR ('deep learning' OR 'DL')] AND ('hospital' OR 'clinical' OR 'healthcare system') AND ('triage' OR 'early screening' OR 'diagnosis' OR 'mortality prediction' OR 'severity assessment') AND ('covid-19' OR 'sars-cov-2' OR 'Coronavirus' OR 'pandemic') AND ('prediction models')
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Title and abstract screening, full text review, data extraction and Risk of Bias Analysis were conducted by two independent reviewers using Covidence [13], a software for systematic review management.

![Fig. 2. PRISMA (preferred reporting items for systematic reviews and meta-analyses) flowchart.](image-url)
Data were extracted with Covidence software using a customized extraction form. The extraction form included the following fields for each included study: Covidence Study ID, Lead Author, Title, Database source, Country (the country of dataset origin), Hospital Name, No of hospitals, Start date, End date, Outcome, No of days for mortality prediction, Type of AI model, AI Methods used, Type of input data, Source of input data, Sample Size of input data, Predictors, Study design, Number of participants for model development (with outcomes), Total Number of COVID-19 positive patients, Population description, Validation method, Number of participants for model validation (with outcomes), Performance (Area under the curve (AUC%), Accuracy (Acc%), Sensitivity (SEN %), Specificity (SPE%), Positive Predictive Values/ Negative Predictive Values (PPV/NPV %), (95% CI)), Code availability, Limitations, Ethical Considerations, Risk of Bias for participants/ predictors/ outcome/ analysis/, overall risk of bias.

The performance of each AI model was reported in terms of metrics defined using the number of True Positives (TP), True Negatives (TN), False Positive (FP) and False Negatives (FN) [14], as follows:

1) AUC is the area under the Receiver Operating Characteristic (ROC) curve, which plots the true positive rate against the false positive rate. This metric is a standard method for evaluating medical tests and risk models [15].

2) Accuracy is the percentage of cases correctly identified calculated by:

\[ Acc = \frac{TP + TN}{TP + FN + FP + TN} \]

3) Sensitivity is the rate of true positives. It measures the proportion of true positives that the model predicts accurately as positive [16], expressed by:

\[ SEN = \frac{TP}{TP + FN} \]

4) Specificity is the rate of true negatives. It measures the proportion of true negatives that the model accurately predicts as negative [16], calculated by:

\[ SPE = \frac{TN}{FP + TN} \]

5) The positive predictive value (PPV), can be expressed as the ratio of the true positives to the sum of the true positives and false positives and NPV is defined as the ratio of the true negatives to the predicted negatives [17].

In this systematic review, we used PROBAST (Prediction model Risk Of Bias ASsessment Tool) [18], to assess the risk of bias and applicability of the included studies with a focused and transparent approach. PROBAST protocol is organized into the following four domains: participants, predictors, outcome, and analysis. These domains contain a total of 20 signaling questions to facilitate structured judgment of ROB, which was defined to occur when shortcomings in study design, conduct, or analysis led to systematically distorted estimates of model predictive performance.

B. Inclusion - Exclusion criteria

Studies that reported the use of AI techniques, including but not limited to techniques from the AI subfield of Machine Learning (ML) and techniques from the ML subfield of DL for developing predictive models for Triage, Diagnosis and Prognosis (such as disease progression, mortality prediction, severity assessment) were included in this systematic review. Cohorts, retrospective cohorts, randomized controlled trials, diagnostic test accuracy, single-centered or multicentered retrospective studies were selected for further analysis.

Restrictions were applied concerning the setting of the studies. If the outcome of the studies could not be applied on a clinical setting, these studies were excluded. Concerning the type of participants, studies that did not include COVID-19 patient data, were excluded. Additionally, studies affecting mental health were excluded. Finally, studies that did not use exclusively AI, ML, or DL to develop these type of prediction models were also excluded.

3. Results

In this review, 879 titles were screened and 101 studies presenting 101 AI-based models for screening, diagnosis and prognosis of COVID-19, were included for full-text review. A significant increase in the number of studies published in the 3rd and 4th quarter of the year was observed (see Fig. 3). We identified in total 14 models for screening (5 based on medical imaging), 38 diagnostic models for detecting COVID-19 (31 based on medical imaging) and 50 prognostic models (7 based on medical imaging) for predicting ICU need, ventilator need, mortality risk, severity assessment or hospital length stay (see Fig. 4). The results are presented in Tables 1-8, including Lead Author of each included study, Country (in case datasets from specific hospitals were used), outcome of the study, number of COVID-19 Positive Patients (CPP) included in the development of the model, AI methods, validation methods and performance of developed prediction models. Missing values of CPP were mainly found in imaging studies that did not specify the number of CT scans or CXR images corresponding to each COVID-19 positive patient. There are in total 11 studies [9,15,19,20,21,22,23,24,25,26,27] with unclear reporting of the number of CPP included.

A. Primary multimodal datasets

Several datasets were identified consisting of multimodal data (e.g., demographic, clinical, imaging) among the included studies. In total 70 studies used COVID-19 positive patient data derived from hospitals from various countries (41 in China, 12 in United States, 6 in Italy, 4 in Brazil, 3 in South Korea, 3 in UK and one study for each of the following countries: France, Spain, Germany, Singapore, Greece, Denmark, Mexico and Israel).

Apart from patient datasets derived from hospitals in the above countries, various publicly available online databases of COVID-19 and pneumonia CXRs ad CT scans were used in the included studies such as:

1) Italian Society of Medical and Interventional Radiology (SIRM) COVID-19 database [28]:

SIRM COVID-19 database reports 384 COVID-19 positive radiographic images (CXR and CT) with varying resolution. Out of 384
radiographic images, 94 images are chest X-ray images and 290 images are lung CT images. This database is updated in a random manner and until 9th December 2020, 115 confirmed COVID-19 cases were reported in this database. This dataset was used by three included studies [29,30,31].

2) Novel Corona Virus 2019 Dataset [32]:

This is a public database in GitHub by collecting 319 radiographic images of COVID-19, Middle East respiratory syndrome (MERS), Severe acute respiratory syndrome (SARS) and ARDS from the published articles and online resources. In this database, they have collected 250 COVID-19 positive chest X-ray images and 25 COVID-19 positive lung CT images with varying image resolutions. This dataset was used by twelve studies [5,30,33,19,20,34,35,36,37,21,22,23].

3) J. Kaggle chest X-ray database [38]:

This is a very popular database, which has 5,863 images chest X-ray images of normal, viral and bacterial pneumonia with resolution varying from 400p to 2000p. Two included studies [30,22] used this dataset.

4) K. COVID-19 Radiography Database [39]:

This database was created for three different types of images classified as chest x-ray images belonging to patients infected with COVID-19, chest x-ray images of cases with viral pneumonia and Chest x-ray images of healthy persons. There are currently 1200 COVID-19 positive images, 1341 normal images, and 1345 viral pneumonia images. This dataset was used by [40] and [24].

5) L. COVID-19 cases open database [41]:

This database contains temporal acquisitions for 42 patients with up to 5 X-ray images per patient, with ground truth annotation as a therapeutic issue for each patient: death or survival. The ground truth annotation can aid in developing prognostic models like the one presented by Fakhfakh et al. [42] to classify multi-temporal chest X-ray images and predict the evolution of the observed lung pathology based on the combination of convolutional and recurrent neural networks.

Moreover, regarding the type of input data employed in the developed models, several studies used demographic, clinical and imaging data or even a combination of those multimodal data types. Fang et al. [43] study, reported to be among the first attempts to fuse clinical data and sequential CT scans to improve the performance of predicting COVID-19 malignant progression in an end to end manner. Liu et al. [44] also used a combination of quantitative CT features of pneumonia lesions with traditional clinical biomarkers to predict the progression to severe illness in the early stages of COVID-19. Additionally, Chassagnon et al. [45] reported an AI solution for performing automatic screening and prognosis based on imaging, clinical, comorbidities and biological data.

B. AI algorithms

The most frequently used AI algorithms for classification purposes in all studies were Random Forests (RF), Linear Regression (LR), Support Vector Machines (SVM), Convolutional Neural Networks (CNN), Decision Trees (DT) and XGBoost (XGB) which are in general six widely used classification methods. Random Forest is a tree-based learning algorithm that utilizes decision trees rising from the training subset which are selected randomly to solve a classification problem [16]. A logistic regression model predicts the probability of a categorical dependent variable occurring [16]. The SVM model seeks to find the hyperplane that has a maximal distance between two classes [16]. CNN is one of the most widely used deep neural networks with multiple layers, including convolutional layer, non-linearity layer, pooling layer, and fully connected layer. CNN has an excellent performance in machine learning problems, especially in imaging studies [46]. DTs are one of the most popular approaches for representing classifiers, expressed as a recursive partition of the instance space [47]. XGB generates a series of decision trees in sequential order; each decision tree is fitted to the residual between the prediction of the pre-

![Fig. 4. Included AI based prediction models.](image-url)
| Study, Country, Outcome                              | No. of CPP* | AI methods         | Predictors                                                                                           | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/Predictors/Outcome/Analysis/Overall |
|----------------------------------------------------|-------------|-------------------|------------------------------------------------------------------------------------------------------|--------------|-----------------------------------------------------------------------------------------------|-------------------------------------------------------------------|
| Yang et al. [4], USA, Early and rapid identification of high-risk SARS-CoV-2 infected patients | 1,898       | LR, DT, RF, GBDT  | age, gender, race and 27 routine laboratory tests                                                   | 5-FCV        | AUC 0.854 (95% CI: 0.829–0.878)                                                                  | L U H H H                                                        |
| Li et al. [63], China, Screening based on ocular surface features | 104         | DL                | Imaging features                                                                                   | 5-FCV        | AUC 0.999 (95%CI, 1670.997–1.000, SEN 98.2, SPE 97.8)                                           | U U U U H                                                        |
| AS Soltan et al. [3], UK, Early detection, Screening | 437         | multivariate LR, RF, XGBoost | Presentation laboratory tests and vital signs                                                         | TTS, 10-FCV  | ED model: AUC 0.939, SEN 77.4, SPE 95.7Admissions model: AUC 0.940, SEN 77.4, SPE 94.8Both models achieve high NPP (>99) | H H H H H                                                        |
| Nan et al. [57], China, Early screening             | 293         | DL, LR, SVM, DT, RF | 4 epidemiological features, 6 clinical manifestations (muscle soreness, dyspnea, fatigue, lymphocyte count, WBC, imaging features) | TTS          | AUC 0.971, Acc 90, SPE 0.95 (LR optimal screening model)                                        | H U H H H                                                        |
| Soares et al. [58], Brazil, Screening of suspect COVID-19 patients | 81          | ML, SVM, SMOTE Boost, ensembling, k-NN | Hemogram: (Red blood cells, MCV, MCHC, MCH, RDW, Leukocytes, Basophils, Monocytes, Lymphocytes, Platelets, Mean platelet volume, Creatinine, Potassium, Sodium, CRP, Age | unspecified  | AUC 86.78 (95%CI: 85.65–87.90), SEN 70.25 (95% CI: 66.57–73.12), SPE 85.98 (95%CI: 84.94–86.84), NPV 94.92 (95% CI: 94.37–95.37), PPV 44.96 (95% CI: 43.15–46.87) | L U H H H                                                        |
| Feng et al. [59], China, Early identification of suspected COVID-19 pneumonia on admission | 32          | ML, LR (LASSO), DT, Adaboost | lymphopenia, elevated CRP and elevated IL-6 on admission                                             | 10-FCV       | AUC 0.841, SPE 72.7                                                                            | H H H H H                                                        |
| Wu et al. [60], China, Early detection              | 27          | RF                | 11 key blood indices: TP, GLU, Ca, CK-MB, Mg, BA, TBIL, CREA, LDH, K, PDW platelets, leukocytes, eosinophils, basophils, lymphocytes, monocytes, |
|                                                    |             |                   |                                                                                                      | 10-FVC, Ext. Val. 10-FCV | Acc 95.95, SEN 95.12, SPE 96.97                                                                | L L L L H                                                        |
| Banerjee et al. [61], Brazil, Initial screening    | 81          | RF, ANN           |                                                                                                      | 10-FCV       | AUC 0.95                                                                                       | H H H H H                                                        |
| Peng et al. [62], China, Quick and accurate diagnosis | 32         | SRLSR, non-dominated radial slots-based algorithm, ARMED, GFS, RFE | 18 diagnostic factors: WBC, eosinophil count, eosinophil ratio, 2019 new Coronavirus RNA (2019-n-CoV), Amyloid-A, Neutrophil ratio, basophil ratio, platelet, thrombocytocrit, monocyte count, procalcitonin, neutrophil count, lymphocyte ratio, lymphocyte count, monocyte ratio, MCHC, Urine SG | not performed  | not performed                                                                                   | L L U H H                                                        |

*CPP = COVID-19 Positive Patients, Abbreviations of medical terms included in this Table are provided in the Appendix.

**L: Low, H: High, U: Unclear
vious decision tree and the target value, and this is repeated until a predetermined number of trees or a convergence criterion is reached [48]. AI techniques based on the above algorithms or other algorithmic approaches for classification purposes that were identified among the reviewed studies included GDCNN (Genetic Deep Learning Convolutional Neural Network) [74], CRT (Classification and Regression Decision Tree) [116], ET (Extra Trees) [15], GBDT (Gradient Boost Decision Tree) [4,71], GBM light (Gradient Boosting Machine light) [100,115], Adaboost (Adaptive Boosting) [27], Boost Ensembling [58,45,59], k-NN (K-Nearest Neighbor) [5,9,58,70,106,112], NB (Naïve Bayes) [9], BNB (Bernoulli Naïve Bayes) [5], GNB (Gaussian Naïve Bayes) [5,27], Inception Resnet (Inception Residual Neural Network) [117], LDA (linear discriminant analysis) [43], RBF (Radial Basis Function) [45], LSTM (Long-Short Term Memory) [43].

Moreover, the use of AI techniques for data preprocessing purposes was reported in the reviewed studies. In particular, AI techniques, including ARMED (Attribute Reduction with Multi-objective Decomposition Ensemble optimizer) [62], GFS (Gradient boosted feature selection) [62], MMRMR (Maximum Relevance Minimum Redundancy) [84], and RFE (Recursive Feature Elimination) [62] were used for feature selection. Regression models, including L1LR (L1 Regularized Logistic Regression) [17], LASSO (Least Absolute Shrinkage and Selection Operator) [50,59,84,89,112], CoxPH (Cox Proportional Hazards) [100], and SRLSR (Sparse Rescaled L1 L2 Shrinkage and Selection Operator) [50,59,84,89,112], CoxPH were also employed for feature selection.

The use of Nadam optimizer (NesterovAccelerated Adaptive Moment optimizer) for model optimization was reported in two studies [33,117], while application of SMOTE (Synthetic Minority Oversampling Technique) for data augmentation was reported in one study [58]. Finally, NLP (Natural Language Processing) was employed in three studies [7,48,86], for data mining purposes.

Regarding novel technologies used in the included studies, explainable AI (XAI) methods, and Federated Learning (FL) were investigated. XAI refers to the ability to explain to a domain expert the reasoning that enables the algorithm to produce its results and is deemed increasingly important in health AI applications [49]. FL, the reasoning that enables the algorithm to produce its results and is deemed increasingly important in health AI applications [49], is a nascent field for data-private multi-institutional collaborations, where model-learning leverages all available data without sharing data between hospitals, by distributing the model-training to the data-owners and aggregating their results [51].
| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/ Predictors/ Outcome/Analysis/ Overall |
|-------------------------|------------|------------|------------|--------------|-------------------------------------------------------------------------------------------------|-------------------------------------------------------------------|
| Chen et al. [92], China, Diagnosis | 51 | DL | Imaging features | TTS | Acc 95.24, SEN 100, SPE 93.55, PPV 84.62, NPV 100 | H U L H H |
| Rahimzadeh et al. [33], Diagnosis | 118 | DNN, Nadam optimizer | Imaging features | TTS | Acc 99.50 | L U L H H |
| Roy et al. [72], Italy, Diagnosis | 17 | DL | Imaging biomarkers | 5-FCV | F1-score 65.9 | H U H H H |
| Zhou et al. [73], China, Diagnosis | 35 | DL, CNN | Imaging features | Ext. val. | TTS | Acc 91.66, SEN 90.80 | H U H H H |
| Ter-Sarkisov et al. [93], China, Diagnosis | 150 | DL, CNN | Imaging features | Int.- ext. val. | TTS | Acc 92.5, SEN 92 | U U H H H |
| Babukarthik et al. [74], Early diagnosis | 102 | GDCNN | Imaging features | TTS | Acc 98.84, SEN 90, SPE 97.0 | H H H H H |
| Minaee et al. [20], Diagnosis | unclear | CNN | Raw images without feature extraction | TTS | SEN 99.5 (95% CI: 99.3–99.7), SPE 95.6 (95% CI: 94.9–96.2) | L H H H H |
| Yan et al. [94], China, Diagnosis | 206 | CNN | Imaging features | TTS | SEN 96.4 (95% CI: 88–100), SPE 88.4 (95% CI: 82–94) | U U H H H |
| Lorkwani et al. [95], India, Diagnosis | 55 | NN | Imaging features | TTS | Acc 95.8, SEN 96, SPE 100.00 | U U U H H |
| Jin et al. [96], China, Screening (early detection) | 751 | DL, CNN | Imaging features | TTS | Acc 98.7, SEN 99.58, SPE 100.00 | U U U H H |
| Ko et al. [29], South Korea, Diagnosis | 20 | 2D DL | Imaging features | TTS, ext.val. | TTS | Acc 98 | H H H L H |
| Ezzat et al. [34], Diagnostic imaging | 99 | Hybrid CNN | Not applicable | TTS | Acc 98 | H U H H L |
| Ouchicha et al. [40], Diagnosis | 43 | DCNN | Imaging features | 5-FCV | Acc 97.20, SEN 96 (95% CI: 90–98), SPE 97 (95% CI: 91–98) | U U L L H |
| Xiong et al. [75], China, Diagnosis | 521 | DL, CNN | Imaging features | 5-FCV | Acc 97.4 | U U H H H |
| Li et al. [2], China, Diagnosis | 468 | DL, CNN | Imaging features | TTS | Acc 97.6 | L U H H H |
| Mahmud et al. [25], China, Diagnosis | unclear | DCNN | Imaging features | 5-FCV | Acc 97.4 | L U H H H |
| Li et al. [97], China, Diagnosis | 305 | NN | Imaging features | 5-FCV | Precision 93% | U U L L H |
| Sun et al. [98], China, Diagnosis | 1495 | LR, SVM, RF, NN | Imaging features | 5-FCV | ACC 91.79, SEN 93.05, SPE 89.95 | L U L H H |

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/ Predictors/ Outcome/Analysis/ Overall |
|-------------------------|------------|------------|------------|--------------|-------------------------------------------------------------------------------------------------|-------------------------------------------------------------------|
| Ozturk et al. [35], Diagnosis (automatic COVID-19 detection) | 125 | DNN | Imaging features (1. Ground-glass opacities (GGO) (bilateral, multifocal, subpleural, peripheral, posterior, medial and basal) 2: A crazy paving appearance (GGOs and inter-/intra-lobular septal thickening 3. Air space consolidation. 4. Broncho vascular thickening (in the lesion). 5. Traction bronchiectasis Imaging features) | 5-FCV | Acc 98.08, SEN 95.13, SPE 95.03, F1-score 96.51 | H U H H H |
| Zhang et al. [7], United States, Diagnosis | 2060 | DNN (CV19-Net), NLP | Imaging features | TTS | AUC 0.92 (95% CI: 0.91–0.93), SEN 88% (95% CI: 87–89), SPE 79% (95% CI: 77–80) | H U H H H |
| Borkowski et al. [37], Diagnosis | 103 | Automated ML platform Microsoft CustomVision | Imaging features | Ext. val. with US Dep. of Vet. Affairs (VA) PACS | Acc 97, SEN 100, SPE 95, PPV 91, NPV 100 | H H H H H |
| Xu et al. [6], China, Diagnosis | 432 | DL, CNN, FL | Imaging features | 5-FCV | SEN 77.2, SPE 91.9 | L U U H H |
C. Validation methods

The most frequent validation methods used in the included studies were training test split (TTS) (34 studies), 5-fold cross-validation (FCV) (23 studies) and 10-fold cross-validation (19 studies). Other studies performed internal and external validation methods. Internal validation techniques, are advocated when no other data than the study sample are being used, to estimate the potential of overfitting in the performance of the developed model [52]. External validation is used to adjust or update the model in other data than the study sample [52]. TTS validation method splits the data into training and testing datasets based on a predefined threshold. FCV is the traditional method for estimating the future error rate of a prediction rule constructed from a training set of data [53]. To estimate the potential for overfitting and optimism in a prediction’s model performance, internal validation techniques are advocated, meaning that no other data than the study sample are being used [52]. External validation uses new participant level data, external to those used for model development, to examine whether the model’s predictions are reliable and adequately accurate, in individuals from potential population for clinical use [54]. All validation methods used in the included studies, are reported in Tables 1-7.

D. Models developed for screening purposes

We identified 14 models for screening COVID-19 (see Tables 1 and 2). Five screening models were based in medical imaging [5,30,31,55,56] using CXRs and CT scans from public databases or hospitals in China. Wang et al. [56] were the first to develop and validate a deep learning algorithm on the basis of chest CT scans of 1647 COVID-19 positive patients acquired from fever clinics of five hospitals in Wuhan, China, for rapid triaging, achieving AUC 0.953 (95% CI 0.949–0.959), SEN 0.923 (95% CI 0.914–0.932), SPE 0.851 (0.842–0.860), PPV 0.790 (0.777–0.803) and NPV 0.948 (0.941–0.954). The rest of the screening studies [4,3,22,24,60,61,62] used as input data -among other- demographic data, comorbidities, epidemiological history of exposure to COVID-19, vital signs, blood test values, clinical symptoms, infection-related biomarkers, and days from illness onset to first

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**Table 4 (continued)**

| Study, Country, Diagnosis, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias** | Participants/ Predictors/ Outcome/ Analysis/ Overall |
|----------------------------------|-------------|------------|------------|--------------|-----------------------------------------------------------------|---------------|-----------------------------------------------|
| Wang et al, [59], Diagnosis      | 266         | DCNN       | Imaging features | Int. val. | Acc 93.3, SEN 91.0, PPV 98.9                                      | L             | U U H H                                      |
| Elaziz et al, [21], Diagnosis    | unclear     | CNN        | Imaging features | Ext. val. | Acc 98.09                                                       | H             | U U L H                                      |
| Gozes et al, United States, China, Diagnostic imaging | unclear | DCNN | Imaging features | TTS       | AUC 0.996, (95%CI: 0.989–1.00), SEN 98.2, SPE 92.2              | U             | U H H H                                      |
| Salman et al, [22], Diagnosis    | unclear     | DL, CNN    | Imaging features | TTS       | Acc 100, SEN 100, SPE 100, PPV 100, NPV 100                      | U             | U U H H                                      |
| Liu et al, [24], Diagnosis       | unclear     | FL (Federated Learning) | Imaging features | Comparison with and without FL 10-   | Comparison with and without FL 10- | U             | U L L H                                      |
| Castiglioni et al, [76], Italy, Diagnosis | unclear | CNN | Imaging features | Int. – ext. | Acc 89.5, SEN 87, SPE 88                                        | H             | U U H H                                      |
| Padma et al, [23], Diagnosis     | unclear     | ML, CNN    | Imaging features | TTS       | Acc 99                                                          | U             | U U H H                                      |
| Zhang et al, [100], China, Diagnosis | 752    | Light GBM, CoxPH | Imaging features | 5-FCV     | AUC 0.9797 (95% CI: 0.966–0.9904), Acc 92.49, SEN 94.93, SPE 91.13 | U             | U U H H                                      |
| Wang et al, [77], China, Diagnosis | 79         | DL, CNN    | Imaging features | Int. – ext. |                                                     | U             | U H H                                         |

*CPP = COVID-19 Positive Patients, Abbreviations of medical terms included in this Table are provided in the Appendix.

**L: Low, H: High, U: Unclear
### Table 5

Results for prognostic models – part 1.

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/Predictors/Outcome/Analysis/Overall |
|-------------------------|-------------|------------|------------|-------------|-------------------------------------------------------------------------------------------------|------------------------------------------------------------------|
| Muhammad et al. [9], South Korea, Recovery prediction, disease progression | unclear | DT, SVM, NB, LR, RF, K-NN | unclear | 5-FCV | Acc 99.85 (Decision Tree) H U U U H | |
| Cheng et al. [78], United States, Severity Assessment, (risk prioritization tool that predicts ICU transfer within 24 h) | 1987 | RF | respiratory failure, shock, inflammation, renal failure | TTS, 10-FCV | AUC 79.9 (95% CI: 75.2–84.8), Acc 76.2 (95% CI: 74.6–77.7), SEN 72.8 (95% CI: 63.2–81.1), SPE 76.3% (95% CI: 74.7–77.9) | H H H H H |
| Kim et al. [79], South Korea, ICU need prediction | 4787 | 55 ML models developed, (XGBoost model revealed the highest discrimination perf.) | age, sex, smoking history, body temperature, underlying comorbidities, activities of daily living (ADL), symptoms | TTS | AUC 0.897, (95% CI 0.877–0.917) | |
| Yadaw et al. [101], United States, Mortality prediction | 4802 | ML, RF, LR, SVM, XGBoost | age, minimum oxygen saturation over the course of their medical encounter, type of patient encounter (inpatient vs outpatient and telehealth visits) | TTS | AUC 91 | L H H H H |
| Klann et al. [102], USA, France, Italy, Germany, Singapore, Severity assessment | 4227 | ML | PaCO2, PaO2, ARDS, sedatives, d-dimer, immature granulocytes, albumin, chlorhexidine, glycopyrrolate, palliative care encounter | 5-FCV, TTS | AUC 0.956 (95% CI: 0.952, 0.959) | U U H H H |
| Navlakha et al. [103], United States, Severity assessment in cancer patients (predicting severity occurring after 3 days) | 354 | ML, RF, DT | 40 out of 267 clinical variables (3 most important individual lab variables: platelets, ferritin, and AST (aspartate aminotransferase) | 10-FCV | AUC 70–85 | L H H H H |
| Shashikumar et al. [104], United States, Mechanical ventilation need prediction (24 h in advance) | 777 | DL | vital signs, laboratory values, sequential-organ failure assessment (SOFA) scores, Charlson comorbidity index scores (CCI) index, demographics, length of stay, outcomes | Ext. val., 10-FCV | AUC 0.918 | L H H L H |
| Bertsimas et al. [80], Greece, Italy, Spain, United States, Mortality risk | 3,927 | XGBoost | Increased age, decreased oxygen saturation (<93%), elevated levels of CRP (>130 mg/L), blood urea nitrogen, blood creatinine | Cross-validation | AUC 0.90 (95% CI: 0.87–0.94) | H H U L H |

### Results for prognostic models – part 2

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/Predictors/Outcome/Analysis/Overall |
|-------------------------|-------------|------------|------------|-------------|-------------------------------------------------------------------------------------------------|------------------------------------------------------------------|
| Youssef et al. [105], UK, Severity assessment (ventilation, ICU need prediction), early warning system | 472 | GBT, LR, RF | Vital signs: Heart Rate, Oxygen Saturation, Respiratory Rate, Systolic Blood Pressure Temperature, AVPU/Venous blood tests: Albumin, ALK, Phosphatase, ALT, APTT, Basophils, Bilirubin Creatinine, CRP, Eosinophil, Haemocrit, Haemoglobin, INR Lymphocytes, Mean Cell Vol., Monocytes, Neutrophils Platelets, Potassium, Prothrombin Time, Sodium, Urea, WCs, eGFR | 5-FCV | AUC 0.94 (best performance, GBT) | U H H H H |
| Vaid et al. [50], United States, Mortality Prediction | 4029 | FL, LR, federated LASSO, federated MLP | gender, age, ethnicity, race, past medical history (such as asthma), lab tests (white blood cell counts) | 10-FCV | AUC 0.693–0.805 | L L H H H |
| Karthikeyan et al. [106], China, Mortality prediction | 370 | ML, XGBoost, k-NN, LR, RF, DT | neutrophils, lymphocytes, LDH, hs-CRP, age | 5-FCV | Acc 96 (16 days in advance) | H U U U H |
Table 5 (continued)

Results for prognostic models – part 2

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/ Predictors/ Outcome/Analysis/ Overall |
|-------------------------|-------------|------------|------------|--------------|-------------------------------------------------------------------------------------------|---------------------------------------------------------------|
| Jimenez-Solem et al. [81], Denmark, UK, Severity assessment (ICU need, mechanical ventilation need), disease progression, mortality prediction | 3944 ML, RF | age, gender, BMI, hypertension, diabetes saturation values, lab-oratory values (lymphocyte counts, CRP, WBC counts, Haemoglobin), variables related to comorbidities (number of comorbidities, presence of cardiovascular pathologies and/or arterial hypertension), radiological values computed through CovidNet, and presence of symptoms (vomiting/nausea or dyspnea or respiratory failure). | Ext. val. (UK biobank), 5-FCV | AUC 0.678–0.818 (hospital admission), AUC 0.587–0.736 (ICU admission), AUC 0.625–0.674 (mortality prediction) | H H H U H |
| Casiraghi et al. [107], Italy, Risk prediction | 301 ML, RF | 3944 ML, RF | AUC 0.587–0.736 (ICU admission), AUC 0.625–0.674 (mortality prediction) | | |
| Burdick et al. [82], United States, Ventilation need prediction within 24 h and triage | 197 ML, XGBoost | diastolic blood pressure (DBP), systolic blood pressure (SBP), heart rate (HR), temperature, respiratory rate (RR), oxygen saturation (SpO2), white blood cell (WBC), platelet count, lactate, blood urea nitrogen (BUN), creatinine, and bilirubin | Ext. val. | AUC 0.866, SEN 90, SPE 58.3 | H U H H H |

Results for prognostic models – part 3

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/ Predictors/ Outcome/Analysis/ Overall |
|-------------------------|-------------|------------|------------|--------------|-------------------------------------------------------------------------------------------|---------------------------------------------------------------|
| Cai et al. [83], China, Severity assessment, ICU need and length of stay prediction, O2 inhalation duration prediction, sputum NAT-positive prediction and patient prognosis | 99 ML, RF | CT quantification | 7 continuous laboratory variables: blood routine test, serum biochemical (including glucose, renal and liver function, creatine kinase, lactate dehydrogenase, and electrolytes), coagulation profile, cytokine test, markers of myocardial injury, infection-related makers, other enzymes | 10-FCV | AUC 0.945 (ICU treatment), AUC 0.960 (prognosis/partial recovery vs pro-longed recovery) | H U U H H |
| Wu et al. [84], China, Mortality prediction | 58 ML, LR, mRMR, LASSO LR | COVID-19 patient’s geographical, travel, health, and demographic data | Cross validation | SEN 98 (95% CI: 93–100), SPE 91 (95% CI: 84–99) | H L L H H |
| Iwendi et al. [27], Severity and outcome prediction | unclear ML, DT, SVM, GNB, Boosted RF, AdaBoost DT, RF, ET (extra trees) | COVID-19 patient’s geographical, travel, health, and demographic data | unspecified Acc 94 | H U H H H |
| Gerevini et al. [15], Italy, Mortality prediction | unclear DT, RF, ET | Age, sex, C-Reactive Protein (PCR), Lactate dehydrogenase (LDH), Ferritin (Male/Ferritin (Female), Troponin-T, White blood cell (WBC), D-dimer, Fibrogenin, Lymphocyte (over 18 years old patients), Neutrophils/Lymphocytes, Chest XRay-Score (RX) | Cross validation | AUC 90.2 for the 10th day | H U U L H |
| Yan et al. [108], China, Severity assessment | 404 ML, XGBoost | 3 biomarkers that predict the survival of individual patients: LDH, lymphocyte, high-sensitivity C-reactive protein (hs-CRP), 8 features had a positive association with mortality (high risk: consciousness, male sex, sputum, blood urea nitrogen [BUN], respiratory rate [RR], D-dimer, number of comorbidities, and age) and 6 features were negatively correlated with mortality (low risk: platelet count[PLT], fever, albumin [ALB], SpO2, lymphocyte, and chronic kidney disease [CKD]), | cross-validation | Acc 90 | L L H H H |
| Gao et al. [71], China, Mortality prediction | 2160 LR, SVM, GBDT, NN | | Int. val. cohort | AUC 0.9621 (95% CI: 0.9464–0.9778) | H H L L H |

(continued on next page)
### Table 5 (continued)

Results for prognostic models – part 3

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/ Predictors/ Outcome/Analysis/ Overall |
|------------------------|-------------|------------|------------|--------------|-------------------------------------------------------------------------------------------------|---------------------------------------------------------------|
| Ma et al. [85], China, Mortality prediction | 305 | ML, RF, XG Boost | LDH, CRP, age | 4-FCV | AUC 0.951 | H L U H H |

Results for prognostic models – part 4

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/ Predictors/ Outcome/Analysis/ Overall |
|------------------------|-------------|------------|------------|--------------|-------------------------------------------------------------------------------------------------|---------------------------------------------------------------|
| Schwab et al. [109], Brazil, Prognosis for Hospitalization need, ICU need | 558 | LR, NN, RF, SVM, XGBoost | Predicting SARS-CoV-2 test results (first 3): presence of arterial lactic acid measurement, age, urinalysis count | TTS | AUC 0.92, (95% CI 0.81–0.98), sensitivity 75(95% CI 67–81), specificity 49 (95% CI 46–51) for COVID-19 positive patients that require hospitalization| L H H H H |
| Hao et al. [48], United States, Early prediction of level-of-care requirements (hospitalization/ ICU/ mechanical ventilation need prediction, | 2566 | NLP, RF, XGBoost, SVM, LR | Vital signs, age, BMI, dyspnea, comorbidities (Most important for hospitalization), Opacities on chest imaging, age, admission vital signs and symptoms, male gender, admission laboratory results, diabetes (most important risk factors for ICU admission and mechanical ventilation) | TTS | Acc 88 (hospitalization need), Acc 87 (ICU care), Acc 86 (mechanical ventilation need) | L L L L U |
| Wu et al. [87], China, Severity assessment, triage | 299 | LR | 1) Clinical features: age, hospital employment, body temperature and the time of onset to admission 2) Laboratory features: Lymphocyte (proportion), neutrophil, (proportion), CRP, lactate dehydrogenase (LDH), creatine kinase, urea and calcium2) CT semantic features: lesion-Most prominent predictors: age, lymphocyte (proportion), CRP, LDH, creatine kinase, urea and calcium symptoms and comorbidities | 5-FCV | AUC 0.84–0.93, Acc 74.4–87.5, SEN 75.0–96.9, SPE 55.0–88.0 | H H H H H |
| Li et al. [16], Fatality prediction | 30,406 | DL, LR, RF, SVM, XGBoost | age, gender | TTS | Acc> 90, SPE> 90, SEN< 40 | H H H H H |
| Nemati et al. [110], Predict patient length of stay in the hospital | 1182 | SVM, GB, DT | top predictors: D-dimer, G2 Index, neutrophil: lymphocyte ratio, C-reactive protein, lactate dehydrogenase | 5-FCV | AUC 0.968 (95% CI: 0.87–1.0) | L H U L H |
| Zhu et al. [111], China, Mortality prediction | 181 | DL, DNN | | 10-FCV | Ext. val. 1) Critical event prediction: AUC 0.80 (3 days), 0.79 (5 days), 0.80 (7 days), 0.81 (10 days) 2) Mortality prediction: AUC 0.88 (3 days), 0.86 (5 days), 0.84 (10 days) | L H H H H |

Results for prognostic models – part 5

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc%), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/ Predictors/ Outcome/Analysis/ Overall |
|------------------------|-------------|------------|------------|--------------|-------------------------------------------------------------------------------------------------|---------------------------------------------------------------|
| Wollenstein-Betech et al. [113], Mexico, predict :(1) hospitalization, (2) mortality (3) ICU need (4) ventilator need | 20,737 | SVM, LR, RF, XGBoost | age, gender, diabetes, COPD, asthma, immunosuppression, hypertension, obesity, pregnancy (hospitalization need prediction), chronic renal failure, tobacco use, other disease, SARS-CoV-2 test results | 10-FCV | AUC 0.63, Acc 79 (mortality prediction performance), AUC 0.74 (hospitalization prediction performance), AUC 0.55, Acc 89 (ICU need prediction performance), AUC 0.58, Acc 90 (ventilator need prediction performance) | L H H H H |
| Wang et al. [17], China, Mortality prediction of hospitalized COVID-19 patients | 375 | L1LR, L1SVM | LDH, lymphocytes percentage, hs-CRP, Albumin | TTS | Acc> 94, F1-score 97 | H L U H H |
### Table 5 (continued)

#### Results for prognostic models – part 5

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc %), Sensitivity (SEN %), Specificity (SPE %), PPV/NPV (%), (95% CI)) | Risk of Bias**: Participants/Predictors/Outcome/Analysis/Overall |
|-------------------------|-------------|-------------|-------------|--------------|--------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------|
| Jiang et al. [88], China, Severity assessment, Outcome prediction | 53 | ML, LR, k-NN, DT, RF, SVM | Most predictive features: alanine aminotransferase (ALT), myalgias, hemoglobin (red blood cells) Less predictive features: lymphocyte count, white blood count, temperature, cycle threshold, creatinine, gender, CRP, age, fever, CK, LDH, Gli, AST, K+, N+ | 10-FCV | Acc 70–80 | H U H H H |
| Sun et al. [114], China, Severity assessment | 336 | SVM | 36 clinical indicators (mainly thyroxine, immune related cells and products including CD3, CD4, CD19, CRP, high-sensitive CRP, leukomonocytes and neutrophils), age, GSH, total protein | TTS | AUC 0.999, SEN 77.5, SPE 78.4 | L H L H H |
| Razavian et al. [115], United States, Outcome prediction | 3,317 | LR, RF, Light GBM | TTS | AUC 90.8, (95% CI: 90.7–91.0) | L U H L H |
| Liang et al. [89], China, Early triage of critically ill (Critical illness was defined as a composite event of admission to an ICU or requiring invasive ventilation, or death.) | 1590 | DL, LASSO algorithm | Three int. ext. val. | AUC 0.911 (95% CI, 0.875–0.945) | H H U L H |
| Gong et al. [90], China, Early identification of cases at high risk of progression to severe COVID-19 | 72 | LR, DT, RF, SVM | Ext. val. | AUC 0.912, (95% CI: 0.846–0.978), SEN 85.7, SPE 87.6 | H H H H H |
| Pourhomayoun et al. [70], 76 countries, Mortality prediction | 117,000 | SVM, ANN, RF DT, LR, k-NN | demographic features (age, sex, province, country, age, travel history), comorbidities(diabetes, cardiovascular disease), patient symptoms (chest pain, chills, colds, conjunctivitis, cough, diarrhea, discomfort, dizziness, dry cough, dyspnea, emesis, expectoration, eye irritation, fatigue, gasp, headache, lesions on chest radiographs, little sputum, malaise, muscle pain, myalgia, obnubilation, pneumonia, myelofibrosis, respiratory symptoms, rhinorrhea, somnolence, sputum, transient fatigue, weakness) | 10-FCV | Acc 93.75, SEN 90, SPE 97 | U L U H H |
| Abdulaal et al. [91], UK, Mortality risk (point-of-admission scoring system) | 398 | ANN | demographics, comorbidities, smoking history, presenting symptoms | Cross validation | AUC 90.12, Acc 86.25, SEN 87.50, SPE 85.94 (95% CI: 61.65–98.45), 85.94 (95% CI: 74.98–93.36), PPV 60.87 (95% CI: 45.23–74.56), NPV 96.49 (95% CI: 88.23–99.02) | H L H H H |
| Assaf et al. [116], Israel, Severity assessment risk prediction model | 162 | ML, NN, RF, CRT | clinical, hematological and biochemical parameters at admission | 10-FCV | Acc 92.0, SEN 88.0, SPE 92.7 | L H H H H |
| Fang et al. [43], China, Early warning to predict malignant progression | 104 | linear discriminant analysis (LDA), SVM, multilayer perceptron (MLP), LSTM | Troponin, Brain natriuretic peptide, White cell count, Aspartate aminotransferase, Creatinine, and Hypersensitive C-reactive protein. | 5-FCV | AUC 0.920, (95% CI: 0.861, 0.979) | H H H H H |

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admission. One study only [63], used ocular surface photographs (eye-region images) as input data demonstrating that asymptomatic and mild COVID-19 patients have distinguished ocular features from others.

In screening studies, the most prominent predictors were age, platelets, leukocytes, monocytes, eosinophils, lymphocytes, CRP, WBC from routine blood tests and imaging features from CXR and CT images. Only one study [57] reported as predictors 4 epidemiological features (relationship with a cluster outbreak, travel or residence history over the past 14 days in Wuhan, exposure to patients with fever or respiratory symptoms over the past 14 days who had a travel or residence history in Wuhan, exposure to Table 5 (continued)

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc %), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias** |
|------------------------|-------------|------------|------------|--------------|--------------------------------------------------------------------------------------------------|----------------|
| Fakhfakh et al. [42], Prognosis | 42 | RNN, CNN SVM, LR | Unclear | unspecified | TTS Acc 95.3, AUC 96.0, SEN 94.0, SPE, 94.0, F1-score 93.3 | U U U H H |
| Zhu et al. [66], China, Disease progression prediction | 408 | SVM, LR | Imaging features | 5-FCV | Acc 92 | L U U H H |
| Qi et al. [67], China, Hospital stay prediction (Short-term (<10 days), long-term (>10 days)) | 31 | RF | Imaging features | 5-FCV | Acc 83.91 | H U U H H |
| Xiao et al. [68], China, Severity assessment, disease progression | 408 | DL, CNN, ResNet34 (RNN) | Imaging features | 5-FCV | Acc 90.97 (95% CI: 0.968–1.00), Acc 97.4 | L U U H H |
| Cohen et al. [36], Severity assessment for COVID19 Pneumonia | 80 | NN | CXR features | not performed | Acc 81, SEN 88, SPE 78 | H U U H H |
| Salvatore et al. [69], Italy, Prognosis prediction (discharging at home, hospitalization in stable conditions, hospitalization in critical conditions, death) | 98 | LR | Imaging features | not performed | Ext. val. | L U U U U |
| Liu et al. [44], China, Severity assessment | 134 | CNN | Imaging features (APACHE-II, NLR, d-dimer level) | Ext. val. | AUC 0.93, (95% CI: 0.87–0.99) | L U U U U |

*CPP = COVID-19 Positive Patients, Abbreviations of medical terms included in this Table are provided in the Appendix.
**L: Low, H: High, U: Unclear

### Table 6

Results for Diagnostic and Prognostic Imaging models.

| Study, Country, Outcome | No. of CPP* | AI methods | Predictors | Val. methods | Performance (AUC, Accuracy (Acc %), Sensitivity (SEN%), Specificity (SPE%), PPV/NPV (%), (95% CI)) | Risk of Bias** |
|------------------------|-------------|------------|------------|--------------|--------------------------------------------------------------------------------------------------|----------------|
| Chassagnon et al. [45], France, Quantification, Staging and Prognosis of COVID-19 Pneumonia | 693 | DL, 2D-3D CNN, RBF SVM, Linear SVM, AdaBoost, RF, DT, XGBoost | 15 radiomics features: imaging from the disease regions (5features), lung regions (5features) and heart features (5features), biological and clinical data (6features: age, sex, high blood pressure (HBP), diabetes, lymphocyte count, CRP level), image indexes (2features: disease extent and fat ratio). | TTS | Acc 70, SEN 64, SPE 77 (Holistic Multi-Omics Profiling & Staging), Acc 71, SEN 74, SPE 82 (AI prognosis model performance) | L U L U U |

*CPP = COVID-19 Positive Patients, Abbreviations of medical terms included in this Table are provided in the Appendix.
**L: Low, H: High, U: Unclear
patients with fever or respiratory symptoms over the past 14 days who had a travel or residence history in other areas with persistent local transmission, or community with definite cases) and 6 clinical manifestations (muscle soreness, dyspnea, fatigue, lymphocyte count, white blood cell count, imaging changes of Chest X-ray or CT).

E. Diagnostic prediction models

Thirty-eight diagnostic prediction models for detecting COVID-19 were identified out of which 32 were based on medical imaging using CXRs or CT scans as input data. Results are presented in Tables 3 and 4. The rest of the diagnostic studies [64,8,9,10,65,11,45] used as input, among other data, age, gender, demographics, symptoms, routine blood exam results, and clinical characteristics. The largest dataset of COVID-19 positive patients (CPP) used in the included diagnostic imaging studies was 2060 patients (5806 CXRs; mean age 62 ± 16, 1059 men) [7]. In this retrospective study, a deep neural network, CV19-Net, was trained, validated, and tested on CXRs, to differentiate COVID-19 related pneumonia from other types of pneumonia, achieving AUC 0.92 (95% confidence interval [CI]: 0.91, 0.93), SEN 88% (95% CI: 87%, 89%) and SPE 79% (95% CI: 77%, 80%). In the non-imaging diagnostic studies, the largest number of CPPs was 845 patients admitted at an Italian hospital from February to May 2020 [64]. Routine blood tests of 1,624 patients were exploited in this study to develop Machine Learning models to diagnose COVID-19 patients, achieving an AUC ranged from 0.83 to 0.90. The most prominent predictors were age, WBC, LDH, AST, CRP and calcium [64].

Most frequently reported predictors in the included studies for identification or diagnosis of COVID-19 cases were age, lymphocytes, WBC and quantitative and radiomic features derived from CXR and CT images.

F. Prognostic prediction models

We identified 50 prognostic models (7 based on medical imaging) [36,42,66,67,68,69,44] for predicting hospitalization need (8 studies), ICU need (10 studies), ventilator need (8 studies), mortality risk (17 studies), severity assessment (16 studies), recovery prediction or disease progression (9 studies) or hospital length stay (3 studies). The results are presented in Table 5 and Table 6. Table 7 presents the results for one diagnostic and prognostic model. The first study to jointly predict the disease progression and the conversion time, which could help clinicians to deal with the potential severe cases in time or even save the patients’ lives, used Support Vector Machines (SVM) and Linear Regression (LR) methods on 408 chest CT scans from COVID-19 positive patients [66]. The largest dataset of COVID-19 positive patients included in the prognostic studies was 117,000 patients worldwide. In this study, Support Vector Machine, Artificial Neural Network, Random Forest, Decision Tree, Logistic Regression, and K-Nearest Neighbor were used to predict the mortality rate based on the patients’ physiological conditions, symptoms, and demographic information [70].

Most of the prognostic prediction models did not report how many days in advance they can produce predictions. Only fourteen studies reported predictions with a time range varying from 12 h to 20 days before the outcome. Highest time frame prediction prior to the outcome was reported by Gao et al. [71] who presented a mortality risk prediction model for COVID-19 (MRPMC) that uses patients clinical data on admission to stratify them by mortality risk, which enables prediction of physiological deterioration and death up to 20 days in advance.

Most frequently reported predictors for prognosis of COVID-19 cases were age, CRP, lymphocyte, LDH and imaging features derived from CXR and CT images.

G. Risk of Bias assessment

The results of the RoB analysis (H: High, L: Low, U: Unclear) were provided for each included study in Tables 1-8. In total 98 prediction models were at high overall risk of bias and three studies [44,45,48] were at unclear risk of bias, indicating that further research needs to be conducted to fully apply them in the clinical practice. In the participants domain, 30 studies had high risk of bias and 21 unclear. Sources of bias in the participants domain varied from small or incomplete datasets to exclusion criteria indicating the need of further data collection to test the generalizability of the developed AI models to other patient populations [2,7,15,16,17,35,37,43,48,57,59,65,69,71–91]. Thirty studies had high RoB in the “predictors” domain related to different ways of definitions and assessment for all participants or predictors availability.

Finally, RoB analysis was high for 99 studies and unclear for two studies [44,48]. Most frequent reasons for assessing high analysis using the PROBAST protocol were number of participants, missing data on predictors and outcomes and exclusion criteria for participants reported as limitations in the included studies.

4. Discussion

In the present systematic review, we included 101 studies that developed or validated screening, diagnostic and prognostic prediction models that can be applied in clinical practice and were published from January 1st, 2020 to December 30th, 2020. Even though most of the studies reported high performance algorithms, results of the RoB analysis conducted in the present review indicate that application in clinical practice may be problematic. Limitations related to the applicability of the developed prediction models were reported by several studies. The most prominent limitation reported was the use of a single data source (one hospital from one geographical area) for the algorithm’s training [4,10,11,15,17,30,56,58,59,62,63,64,69,44,76,83,84,91,92,78,103,107,106,108,109,112,114,115,116,118]. Generalizability of the trained models can be enhanced by adding multiple data sources in future studies.

Concerning the results of the present review, a clear distinction between prediction models that relied on imaging features and models that relied on clinical or laboratory data, proved to be evident amongst the included studies. In particular, medical imaging studies were most prominent for diagnostic purposes (31 out of 37) and least prominent for prognostic purposes (7 out of 50). Models developed for screening purposes were rather few (14) compared to the other two categories and distinction between the use of medical images and clinical or laboratory data was not clear (5 and 9 respectively). Analysis regarding the handling of unbalanced data and application of appropriate performance metrics across the developed models was not feasible due to the fact that different validation methods were applied. Student’s t-test was deployed to investigate differences in dataset sizes among included studies, in terms of total number of participants and COVID-19 Positive Patients. Such comparison between datasets employed in screening, diagnostic and prognostic models did not display statistically significant differences (p > 0.05). Average number of total participants included in datasets of non-imaging medical studies proved to be significantly higher than in datasets of medical imaging studies (11,033.45 versus 1,528.27, p = 0.04). This was not identified for the number of COVID-19 Positive Patients between these two categories (p > 0.05). Regarding the application of performance criteria for evaluation purposes in the included studies, some dissimilarities were observed between models developed for different purposes. In particular, AUC score was
the most prevalent metric in models developed for prognostic purposes (69.38%) and screening (71.14%) purposes, but not for diagnostic purposes (35.12%). The most widely used metric in models for diagnostic purposes was SEN (70.27%), followed by SPE (59.45%) and AUC score (76.78%) being the most common. Comparison between different types of prediction models that employ heterogeneous predictors is exceedingly difficult, taking into consideration that performance criteria applied were not similar. Additionally, since most prediction models were trained on a specific localized dataset, the evaluation of AI techniques used and the importance of predictors cannot be discerned through meta-analysis of the results presented in each study. This heavily underlines the need for global collaboration and data sharing, to enable the development of validated benchmarks for the evaluation of newly introduced AI techniques. This would expedite the application of new prediction models in clinical practice, especially in times of extreme urgency such as the COVID-19 pandemic.

Concerning the type of multimodal input data used in the developed models, three studies [43,45,44] demonstrated the advantages of using the combination of clinical features and image features (CT scans, CXR images), indicating that both CT scans and clinical data are of paramount importance to the diagnosis and prognosis of COVID-19. Moreover, developing AI diagnostic and prognostic models in an end-to-end manner enables the use of raw data without the need for manual design of feature patterns or interference of clinicians. Therefore, future AI studies can explore more methods of fusing clinical and image features as well as developing end-to-end models for use in the clinical practice.

Regarding novel technologies, we investigated the use of explainable AI and Federated Learning in the developed prediction models. Evaluation of the AI methods presented in this review study in terms of explainability proved to be difficult, due to the lack of uniform adoption of interpretability assessment criteria across the research community [119]. A multidisciplinary approach of combining medical expertise and data science engineering in future studies might be necessary to overcome this difficulty [119]. The integration of explainability modalities in the developed models can enhance human understanding on the reasoning process, maximize transparency and embellish trust towards the models’ use in clinical practice [120]. Therefore, XAI techniques could prove to be critical in tackling volatile crises like the COVID-19 pandemic and as such, it is of paramount importance that XAI should be taken into consideration in future works. Moreover, clinical adoption of FL is expected to lead to models trained on datasets of unprecedented size, thus having a catalytic impact towards precision and personalized medicine [51]. FL shows promise in handling the new coronavirus electronic health record data to develop robust predictive models without compromising patient privacy [50] and can inspire more research on future applications.

One of the limitations of the present review is the rapid increase of new COVID-19 related AI models in the literature, challenging the completion of a full list of available studies. Another limitation is the fact that there were 31 included pre-prints which may differ from the final versions once accepted for official publication. A follow-up of the included studies at the time of writing indicated that 7 of the included pre-prints had already been peer-reviewed [34,59,45,70,76,81,102].

Based on the results of the Risk of Bias analysis, further research needs to be conducted to decrease the sources of bias in the included studies. Future studies can investigate the role of the most prominent predictors such as age, platelets, leukocytes, monocytes, eosinophils, lymphocytes, CRP, LDH, WBC from routine blood tests, imaging features from CXR and CT images. Most prominent AI methods (RF, LR, SVM, CNN, DT and XGBoost) along with the aforementioned predictors can be used as a leading approach in developing and validating future screening, diagnostic and prognostic prediction models that can be applied in clinical practice.

5. Conclusion
Artificial Intelligence methods are critical tools for utilizing the rapidly growing body of COVID-19 positive patient datasets, with a vast contribution in the fight against this pandemic. These multimodal datasets may include collected vitals, laboratory tests, comorbidities, CT scans or CXRs. In the present systematic review, we discussed the applicability and provided an overview of the AI-based prediction models developed by the rapidly growing literature, which can be used for screening, diagnosis, or prognosis of COVID-19 in the clinical setting. Limitations and considerations regarding the design and development of said prediction models were identified and future directions were proposed. Moreover, novel technologies such as explainable AI and Federated Learning could prove to be critical in tackling volatile crises like the COVID-19 pandemic. Increased collaboration in the development of the AI prediction models can enhance their applicability in the clinical practice and assist healthcare providers and developers in the fight against this pandemic and other public health crises.

CRediT authorship contribution statement
Eleni S. Adamidi: Conceptualization, Methodology, Formal analysis, Investigation, Data curation, Visualization, Writing - original draft, Writing - review & editing.
Konstantinos Mitsis: Conceptualization, Methodology, Formal analysis, Investigation, Data curation, Visualization, Writing - original draft.
Konstantina S. Nikita: Conceptualization, Methodology, Supervision, Writing - review & editing.

Declaration of Competing Interest
The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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