Artificial intelligence and machine learning (ML) models are rapidly being applied to the analysis of cardiac computed tomography (CT). We sought to provide an overview of the contemporary advances brought about by the combination of ML and cardiac CT. Six searches were performed in Medline, Embase, and the Cochrane Library up to November 2021 for (i) CT-fractional flow reserve (CT-FFR), (ii) atrial fibrillation (AF), (iii) aortic stenosis, (iv) plaque characterization, (v) fat quantification, and (vi) coronary artery calcium score. We included 57 studies pertaining to the aforementioned topics. Non-invasive CT-FFR can accurately be estimated using ML algorithms and has the potential to reduce the requirement for invasive angiography. Coronary artery calcification and non-calcified coronary lesions can now be automatically and accurately calculated. Epicardial adipose tissue can also be automatically, accurately, and rapidly quantified. Effective ML algorithms have been developed to streamline and optimize the safety of aortic annular measurements to facilitate pre-transcatheter aortic valve replacement valve selection. Within electrophysiology, the left atrium (LA) can be segmented and resultant LA volumes have contributed to accurate predictions of post-ablation recurrence of AF. In this review, we discuss the latest studies and evolving techniques of ML and cardiac CT.
**Introduction**

Recent advancements in computed tomography (CT) and data science have fostered the development of machine learning models across several domains within cardiology. Clinical implementation of dual-energy CT systems has improved diagnostic accuracy, reduced calcium blooming artefact, enabled identification of atherosclerotic plaque composition, and decreased the radiation and contrast required for scans, while also paving the way for the identification of novel imaging biomarkers and radiomic profiles. New 256- and 320-slice CT systems significantly reduce radiation doses by achieving a full volume acquisition in one to two cardiac cycles. This reduces cardiac motion artefact, improves image quality and diagnostic accuracy, and enables better quantitative analysis.

These newer systems are relatively expensive and further research is needed into their full potential. In this review, we provide an up-to-date summary of the evolving machine learning (ML) techniques used in conjunction with cardiac CTs, including: (i) coronary artery imaging [fractional flow reserve (FFR), coronary artery calcium (CAC), and plaque characterization], (ii) epicardial adiposity quantification, (iii) aortic stenosis (AS), and (iv) atrial fibrillation (AF).

**Terminology**

A paucity of universally accepted terms and the relationships between ML and other aspects of artificial intelligence (AI) can lead to misunderstanding. Artificial intelligence is an umbrella term given to any algorithm mimicking a human being’s method of problem-solving. Machine learning falls under this category by using probability and statistics to make predictions based on data. Table 1 shows examples of specific tools used. The process of ML starts with patient data and finishes with a final prediction as follows: (i) data collection, (ii) pre-processing, (iii) application of the ML algorithm, and (iv) optimization of the aforementioned steps. Machine learning algorithms can be further classified based on whether they require input ‘training data’ that comprises the original patient data and a corresponding data class ‘label’. The volume and quality of
‘training data’, in combination with the appropriateness of the statistical algorithm applied, correlates with the utility of an ML model. Algorithms that require ‘training data’ are termed ‘supervised learning’ algorithms and are discussed in this review; in contrast to ‘unsupervised learning’ that do not require ‘training data’.

**Methods**

We performed six searches of Medline, Embase, and the Cochrane Library up to November 2021 for original articles containing human subjects pertaining to the use of ML in (i) CT-fractional flow reserve (CT-FFR), (ii) AF, (iii) AS, (iv) plaque characterization, (v) fat quantification, and (vi) CAC score (figure 1). The following terms were used, including MeSH terms, synonyms, and abbreviations (CAC score/ fractional flow reserve/ atrial fibrillation/ aortic stenosis/ coronary plaque/ fat quantification) AND (machine learning OR neural network OR k-nearest neighbour OR random forest) AND (computer tomography). Studies utilizing deep learning algorithms other than convolutional neural network (CNN) were excluded. Duplicates were removed from each search, before titles and abstracts were screened by two authors for each search. Studies were selected if they were original articles describing the use of ML and cardiac CT in each topic. Articles identified are summarized in Tables 2–7.

**Applications of machine learning in cardiac computed tomography**

**CT-fractional flow reserve**

The degree of stenosis on coronary CT angiography (CCTA) does not always correlate with functional flow restriction. For stable coronary artery disease (CAD) with invasive physiological assessment using FFR or instantaneous wave-free ratio (iFR) remains the invasive gold standard in assessing FFR or instantaneous wave-free ratio (iFR) remains the invasive gold standard in assessing flow-limiting lesions, with an FFR ≤ 0.8 or iFR < 0.89 suggesting the need for follow on percutaneous coronary intervention. Advancements in computational fluid dynamics have allowed for the estimation of FFR from CCTA imaging data, resulting in the development of CT-FFR protocols. Using numerous iterations of CNN algorithms, CT-FFR has consistently been demonstrated to be superior to CCTA in assessing flow-limiting lesions with an average area under the curve (AUC) of 0.89 (Table 2). Early work demonstrated that this technique can reduce processing durations by 80-fold compared with physics-based computations, in addition to being less computationally demanding. Nevertheless, Itu et al. was trained on synthetic phantoms and thus lack certain physiological traits that may detrimentally affect clinical accuracy. Moreover, the study by Xu et al. demonstrated the effect of poor image quality and tachycardia on the performance of the algorithm. Indeed, performance was substantially decreased in low-quality images vs. high-quality images, subjectively determined by expert readers (AUC: 0.80 vs. 0.93, respectively). Moreover, in a multicentre study by Tesche et al., performance was also impacted by the CAC burden. Performance of CT-FFR, per vessel, was significantly affected at higher Agatston scores. This appeared to be due to a negative dose–response effect on specificity with higher CAC scores. In 2021, The National Institute for Health and Care Excellence updated its guidance recommending the use of CT-FFRML, provided by companies such as HeartFlow, as it is non-invasive, considered to deliver high diagnostic accuracy, whilst having the potential to be cost-effective. In conjunction, contemporary American and European guidelines also support the use of CT-FFRML.

**Calcium scoring**

Coronary artery calcium predicts cardiovascular events. Low dose electrocardiogram-gated non-contrast CT imaging (CCT) is an effective and non-invasive way for quantifying CAC, having a high sensitivity and negative predictive value for obstructive CAD. Coronary artery calcium is traditionally measured in Agatston scores, which grade calcium severity by multiplying the area of calcification by CT attenuation in Hounsfield units yielding an estimated total CAC burden. Agatston scores correspond to calcium burden, as so: 1–100 mild; 101–400 moderate; and >400 severe. Machine learning has been used for the automation of CAC identification and scoring with subsequent risk categorization of CAD or future cardiac events; easing the burden on reporting clinicians thereby saving both time and resources (Table 3). The use of gradient boosting algorithms has had success in predicting prognosis for patients with suspected cardiovascular disease. In a large retrospective cohort by Nakanishi et al., ML-derived predictions with combined

**Table 1** An overview of algorithms used in machine learning with summary definitions and benefits

| Algorithm | Overview |
|-----------|----------|
| Logistic regression | Determines the probability of a particular class for a discrete variable. A simple algorithm with extensive applications. |
| Support vector machines | Uses ‘kernel mapping’ to set boundaries of data classes. Can be used for hand-written characters and text categorization but is limited in larger datasets. |
| k-nearest neighbour | Classifies data based on the classes of the k closest data points (where k is a positive, whole number). Simple and easy to implement. |
| Random forest | A collection of decision trees that iteratively split data based on binary criteria. The output is a combination of the results of each single decision tree. A major advantage is its ability to prioritize more important characteristics of the dataset. A highly versatile classifier that works well with small datasets. |
| Convolutional neural networks (U-Net) | A convolutional neural network (CNN) is a deep learning algorithm that captures the essence of data using a filter based on convolution. This is used extensively in image processing applications. U-net is a specific form of CNN architecture that utilizes fewer training images to provide more accurate segmentation. |

**Table 2**

| Algorithm | Overview |
|-----------|----------|
| Logistic regression | Determines the probability of a particular class for a discrete variable. A simple algorithm with extensive applications. |
| Support vector machines | Uses ‘kernel mapping’ to set boundaries of data classes. Can be used for hand-written characters and text categorization but is limited in larger datasets. |
| k-nearest neighbour | Classifies data based on the classes of the k closest data points (where k is a positive, whole number). Simple and easy to implement. |
| Random forest | A collection of decision trees that iteratively split data based on binary criteria. The output is a combination of the results of each single decision tree. A major advantage is its ability to prioritize more important characteristics of the dataset. A highly versatile classifier that works well with small datasets. |
| Convolutional neural networks (U-Net) | A convolutional neural network (CNN) is a deep learning algorithm that captures the essence of data using a filter based on convolution. This is used extensively in image processing applications. U-net is a specific form of CNN architecture that utilizes fewer training images to provide more accurate segmentation. |

**Table 3**

| Algorithm | Overview |
|-----------|----------|
| Logistic regression | Determines the probability of a particular class for a discrete variable. A simple algorithm with extensive applications. |
| Support vector machines | Uses ‘kernel mapping’ to set boundaries of data classes. Can be used for hand-written characters and text categorization but is limited in larger datasets. |
| k-nearest neighbour | Classifies data based on the classes of the k closest data points (where k is a positive, whole number). Simple and easy to implement. |
| Random forest | A collection of decision trees that iteratively split data based on binary criteria. The output is a combination of the results of each single decision tree. A major advantage is its ability to prioritize more important characteristics of the dataset. A highly versatile classifier that works well with small datasets. |
| Convolutional neural networks (U-Net) | A convolutional neural network (CNN) is a deep learning algorithm that captures the essence of data using a filter based on convolution. This is used extensively in image processing applications. U-net is a specific form of CNN architecture that utilizes fewer training images to provide more accurate segmentation. |

**Table 4**

| Algorithm | Overview |
|-----------|----------|
| Logistic regression | Determines the probability of a particular class for a discrete variable. A simple algorithm with extensive applications. |
| Support vector machines | Uses ‘kernel mapping’ to set boundaries of data classes. Can be used for hand-written characters and text categorization but is limited in larger datasets. |
| k-nearest neighbour | Classifies data based on the classes of the k closest data points (where k is a positive, whole number). Simple and easy to implement. |
| Random forest | A collection of decision trees that iteratively split data based on binary criteria. The output is a combination of the results of each single decision tree. A major advantage is its ability to prioritize more important characteristics of the dataset. A highly versatile classifier that works well with small datasets. |
| Convolutional neural networks (U-Net) | A convolutional neural network (CNN) is a deep learning algorithm that captures the essence of data using a filter based on convolution. This is used extensively in image processing applications. U-net is a specific form of CNN architecture that utilizes fewer training images to provide more accurate segmentation. |
data were superior to (i) clinical data, (ii) CAC score, and (iii) CT variables alone. This was consistent with Commandeur et al., who performed prospective analysis of 1912 individuals and found ML-derived predictions to be superior to traditional atherosclerotic cardiovascular disease risk algorithm and CAC score. These predictive ML algorithms also predict obstructive CAD with a high degree of accuracy (AUC: 0.77; sensitivity: 100 ± 0.0% and specificity 69.8 ± 3.6%). Automated identification of CAC score has been performed using k-nearest neighbour, CNN and gradient boosting ML with reasonably good accuracies (sensitivity: up to 72% and false positive rate: as low as 0.48 errors per scan; sensitivity: up to 72% and false positive rate: as low as 0.48 errors per scan; and AUC: 0.67–0.85, respectively). It has also been proposed that CAC score can be predicted from clinical variables.

CCT-based, whole heart and vessel-specific CAC scoring algorithms have been developed to include Agatston, mass, and volume scores. They use a k-nearest number classifier with forward feature selection on vessels identified from an atlas-based approach with relatively high degrees of sensitivity and low false-positive rates. Similar vessel-specific volume-based CAC scores were achieved in another study using random forest algorithms with fuzzy spatial features to achieve total intraclass correlation coefficients of 0.99 and an accuracy of 1.0 K in risk class assignment, at a 10 s run time. Lossau et al. have developed CNN trained on simulated cardiac motion images, aimed to automate the estimation and correction of coronary motion in coronary computed tomographic angiography (CCTA) scans, with small degrees of error. This approach may be useful in the CCTA calculation of CAC; however, the results were based on a small dataset of 12 clinical cases.

**Plaque characterization**

Nine studies were identified pertaining to plaque characterization by cardiac CT and the use of ML (Table 4). Earlier studies demonstrated that non-calcified plaques could be identified using ML, with extreme gradient boosting algorithms proving superior to topological soft-gradient detection methods (AUC 0.92 vs. 0.87, respectively). Masuda et al. also showed that their algorithmic approach performed better than the median CT number. Validated methods of ascertaining morphological characteristics of plaques using ensemble methods and multi-task CNNs have been produced. Using similar boosted ensemble algorithms, studies have managed to identify culprit stenotic lesions, predict individuals at risk of rapid coronary plaque progression, and retrospectively predict individuals at risk of major adverse cardiovascular events (MACE), with high degrees of accuracy (AUC: 0.77; 0.83; and 0.96, respectively). The CAD reporting and data system is designed to classify severely obstructed coronary lesions on CCTA. Muscogiuri et al. have demonstrated that a deep learning CNN algorithm can classify over 5 times faster than expert readers, although with an accuracy of between 60% and 86%. It has also been demonstrated that analysis of plaque characteristics can predict MACE and other clinically relevant composite.
| Study       | Design and aim                                                                 | Algorithm used | Participants | Outcome                        |
|------------|-------------------------------------------------------------------------------|----------------|--------------|---------------------------------|
| Itu 2016   | In vitro-validated, in vivo-tested, diagnostic accuracy comparison of CT-FFR<sub>ML</sub> vs. invasive FFR and CT-FFR<sub>CFD</sub> | CNN            | 87           | AUC: 0.90 Accuracy: 83.2%       |
|            |                                                                                |                |              | Sensitivity: 81.6%              |
|            |                                                                                |                |              | Specificity: 83.9%              |
|            |                                                                                |                |              | PPV: 68.9%                      |
|            |                                                                                |                |              | NPV: 91.2%                      |
|            |                                                                                |                |              | Time: 2.4 s                      |
| Coenen 2018| Multicentre, retrospective, diagnostic accuracy comparison of CT-FFR<sub>ML</sub> vs. invasive CCTA and CT-FFR<sub>CFD</sub> | CNN            | 351          | AUC: 0.84 Accuracy: 85%         |
|            |                                                                                |                |              | Sensitivity: 77%                |
|            |                                                                                |                |              | Specificity: 89%                |
|            |                                                                                |                |              | PPV: 76%                        |
|            |                                                                                |                |              | NPV: 89%                        |
| Tesche 2018| Single-centre, retrospective, diagnostic accuracy comparison of CT-FFR<sub>ML</sub> vs. CT-FFR<sub>CFD</sub> and QCA | CNN            | 85           | AUC: 0.91 Accuracy: 90%         |
|            |                                                                                |                |              | Sensitivity: 90%                |
|            |                                                                                |                |              | Specificity: 95%                |
|            |                                                                                |                |              | PPV: 90%                        |
|            |                                                                                |                |              | NPV: 95%                        |
|            |                                                                                |                |              | Time: 40.5 min                  |
| Xu 2020    | Investigation of the impact of image quality, BMI, sex, HR, and calcium on CT-FFR<sub>ML</sub> diagnostic accuracy vs. CCTA and invasive FFR | CNN            | 437          | AUC: 0.80 LQ: 0.80 HQ: 0.93       |
|            |                                                                                |                |              | Accuracy: 83%                   |
|            |                                                                                |                |              | Sensitivity: 78%                |
|            |                                                                                |                |              | Specificity: 86%                |
|            |                                                                                |                |              | PPV: 82%                        |
|            |                                                                                |                |              | NPV: 83%                        |
|            |                                                                                |                |              | Time: 40.5 min                  |
| Zreik 2020 | Retrospective study investigating automatic calculation of CT-FFR<sub>ML</sub> (FFR cut off < 0.9) | CNN            | 187          | AUC: 0.87 Accuracy: 80%         |
| Baumann 2020| Single-centre, retrospective, diagnostic accuracy comparison of CT-FFR<sub>ML</sub> vs. iFR | CNN            | 40           | AUC: 0.96 Accuracy: 95%         |
|            |                                                                                |                |              | Sensitivity: 92%                |
|            |                                                                                |                |              | Specificity: 96%                |
|            |                                                                                |                |              | PPV: 92%                        |
|            |                                                                                |                |              | NPV: 96%                        |
|            |                                                                                |                |              | Time: 11 min                    |
| Lossnitzer 2020 | Single-centre, retrospective, diagnostic accuracy comparison of CT-FFR<sub>ML</sub> vs. invasive FFR and CCTA | CNN            | 88           | AUC: 0.96 Accuracy: 93%         |
|            |                                                                                |                |              | Sensitivity: 94%                |
|            |                                                                                |                |              | PPV: 93%                        |
|            |                                                                                |                |              | NPV: 94%                        |
|            |                                                                                |                |              | Time: 23.9 min                  |
| Li 2021    | Single-centre, retrospective, diagnostic accuracy comparison of CT-FFR<sub>ML</sub> vs. invasive FFR and CCTA | CNN            | 73           | AUC: 0.93 CT-FFR vs. CCTA vessel-level |
|            |                                                                                |                |              | AUC: 0.957 vs. 0.599, P < 0.0001 |
|            |                                                                                |                |              | Accuracy: 90.4%                 |
|            |                                                                                |                |              | Sensitivity: 93.6%              |
|            |                                                                                |                |              | Specificity: 88.1%              |
|            |                                                                                |                |              | PPV: 85.3%                      |
|            |                                                                                |                |              | NPV: 94.9%                      |
| Morais 2021| Single-centre, retrospective, diagnostic accuracy comparison of CT-FFR<sub>ML</sub> vs. invasive FFR | CNN            | 93           | AUC: 0.93 Sensitivity: 87%       |

Continued
Table 2 Continued

| Study       | Design and aim                                                                 | Algorithm used | Participants | Outcome                                                                 |
|-------------|--------------------------------------------------------------------------------|----------------|--------------|-------------------------------------------------------------------------|
| Renker 2021 | Multicentre, retrospective post hoc per-vessel diagnostic accuracy analysis of MACHINE registry comparing of CT-FFR<sub>ML</sub> vs. invasive FFR and CCTA | CNN            | 330          | Specificity: 86% 
PPV: 73% 
NPV: 94% 
Overall average (LAD, LCx and RCA) 
AUC: 0.784 
Sensitivity: 78.4% 
Specificity: 77.2% 
PPV: 64.7% 
NPV: 86.6% |

Time is reported as an approximation of total time required for analysis. Statistics are per patient (per vessel).

AUC, area under the curve; CFD, computational fluid dynamics; CNN, convolutional neural networks; HR, heart rate; HQ, high-quality images; LQ, low-quality images; low Agatston score, >0 to <100; high Agatston score, >400; QCA, quantitative coronary angiography; iFR, instantaneous wave-free ratio.

outcomes<sup>35</sup> with high degrees of accuracy (AUC: 0.96 and 0.797, respectively).

**Epicardial adipose tissue quantification**

The epicardial adipose tissue (EAT), being the fat contained between the pericardium and surface of the myocardium, is involved in a complex interplay with the coronary arteries. It is thought that dysfunctional pro-inflammatory adipokines mediate the development of an elevated risk of CAD and MACE.<sup>69</sup> Another effective use of ML in the analysis of cardiac CT output is in the fully-automated identification and quantification of EAT. Studies have done this using numerous algorithmic approaches (<Table 5>), achieving accuracies up to 98.5%,<sup>36</sup> with excellent correlation with expert readers (Pearson’s correlation, r > 0.924),<sup>37–40,42</sup> and almost identical intra-study dice similarity coefficients (DSCs).<sup>36,39,43,44</sup> A similar technique has been used in combination with a fat radiomic profile (FRP) derived from biopsy and CCTA data of perivascular adipose tissue in a retrospective study by Oikonomou et al.<sup>41</sup> to predict MACE at a 5-year follow-up superior to traditional risk stratification tools with an AUC of 0.880 with FRP and an AUC of 0.754 without FRP.

**Aortic stenosis**

Transcatheter aortic valve replacement (TAVR) is a successful percutaneous intervention for the treatment of severe AS, that is increasingly being used in lower surgical-risk patients.<sup>70</sup> For successful deployment of a TAVR device, pre-operative CT imaging is used to derive various anatomical features of the aortic valve to guide optimal device size selection in order to limit paravalvular regurgitation, coronary obstruction, and conduction disturbance.<sup>47,71</sup> Automated segmentation of the aortic annulus perimeter has been reported using several methods (<Table 6>). Elattar et al.<sup>47</sup> developed a method using thresholding, morphological operators, and fuzzy classification to achieve identical DSC coefficients (0.95 vs. 0.95) at over 13-times faster-processing speeds vs. expert reader. This method, however, did not perform segmentation of the valve leaflets themselves. Al et al.<sup>49</sup> developed a bespoke regression tree-based algorithm to localize all eight aortic valve landmarks required for pre-operative assessment of TAVR procedures, yielding a mean localization error of 2.04 mm and a run time of 12 ms compared with an inter-observer variability of 2.38 mm. To enable segmentation of aortic valve landmarks, Al Abdullah et al.<sup>49</sup> developed a regression tree-based algorithm, yielding high accuracies (mean localization error: 2.38 mm), fast run times (12 ms), and close comparability to expert readers (inter-observer variability: 2.38 mm). Moreover, this model was trained on a generalizable population of patients with variable valvular calcification.<sup>49</sup> To address the computational modelling of valve biomechanics, Liang et al.<sup>48</sup> developed a novel method utilizing CT imaging for the reconstruction of 3D valve geometries with built in mesh correspondence. This approach used linear coding and shape dictionary learning based on k-nearest number algorithms to achieve patient-specific reconstructions with mean discrepancies of 1.57 mm. A limitation of this study was the lack of patients with severe AS, and thus it lacks the impact of valvular calcification on valvular biomechanics.<sup>46</sup> More recently, in a small number of patients with hypertrophic obstructive cardiomyopathy undergoing surgery, CNN models have been used to automatically segment the cardiac structure.<sup>55</sup> This cut time required down from 3 h manually segmenting to 5 min, although one of the two cases did require some manual adjustment.<sup>55</sup>

As mortality following TAVR can vary widely, ML can also be used to predict post-procedural survival and thus identify individuals who are likely to benefit from the intervention. Using Gradient boosting ML and Cox proportional hazard regression models, it has been possible to predict survival to an AUC of 0.72–0.79.<sup>52,54</sup> is superior to manual scoring systems (TAVI2-Score: 0.56 and CoreValve Score: 0.53),<sup>72</sup> and the predictive capacity appears to persist up to 5 years.<sup>54</sup>

**Atrial fibrillation**

Computed tomography imaging is used in pre-operative mapping prior to ablation for AF to assess left atrium (LA) chamber size and pulmonary vein (PV) anatomy. However, the task of isolating the LA and deriving volumes manually is time-consuming. Studies
| Study               | Design and aim                                                                 | Algorithm used                                      | Participants                               | Outcome                                                                 |
|---------------------|--------------------------------------------------------------------------------|----------------------------------------------------|--------------------------------------------|------------------------------------------------------------------------|
| Ilgum 2007          | Accurate, automated identification of CAC scores                                | $k$-nearest neighbour and feature selection scheme | 76 female participants                     | Sensitivity 73.8%                                                      |
| Shahzad 2013        | Automatic detection of whole heart calcium lesions, at 1.5 and 3.0 mm slice spacing | $k$-nearest neighbour                               | 366 patients (training 57%, testing 43%)  | - False-positive rate: 0.1 errors per scan                               |
| Wolterink 2016      | Accurate, automated identification of CAC scores                                | Paired convolutional neural networks                | 250 patients (60% training, 40% testing)    | Detection by paired convolutional neural networks identified more lesions than individual observers: |
|                     |                                                                                |                                                    |                                            | - Sensitivity: 67–72%                                                  |
|                     |                                                                                |                                                    |                                            | - False-positive rate: 0.48–1.69 errors per scan                       |
| Al’Aref 2017        | Accurate, automated identification of CAC score                                 | Gradient boosting machine learning                 | 35 281 patients (CONFIRM registry) (70% training, 30% testing) | AUC                                                                    |
|                     |                                                                                |                                                    |                                            | - CAC score 0.84                                                       |
|                     |                                                                                |                                                    |                                            | - CAC score 1–100: 0.67                                                 |
|                     |                                                                                |                                                    |                                            | - CAC score 101–400: 0.74                                               |
|                     |                                                                                |                                                    |                                            | - CAC score >400: 0.85                                                  |
| Nakanishi 2017      | Retrospective analysis of the capability of ML-determined CAC, clinical data and CT variables vs. each individual factor in predicting coronary heart disease or cardiovascular death. | - 66 636 participants without cardiovascular disease from the Multi-Ethnic Study of Atherosclerosis (MESA) | AUC                                                                 |
|                     |                                                                                |                                                    |                                            | - ML (all variables): 0.85                                              |
|                     |                                                                                |                                                    |                                            | - Clinical data only: 0.83                                              |
|                     |                                                                                |                                                    |                                            | - CAC score only: 0.81                                                 |
|                     |                                                                                |                                                    |                                            | - CT variables only: 0.82                                               |
| Durlak 2017         | Automated CAC labelling system vs. expert reader                                | Atlas-based feature approach and random forest classifier | 40 patients                      | ICC: 0.99                                                             |
|                     |                                                                                |                                                    |                                            | Accuracy: 1.0 $\kappa$                                                 |
|                     |                                                                                |                                                    |                                            | Run time: 10 s                                                         |
| Lossau (née Elss)   | Use of ML to improve interpretability through reducing motion artefact by predicting motion direction. | CNN                                                 | 19 clinical datasets                             | Motion direction error: 34.9 ± 1.21                                     |
|                     |                                                                                |                                                    |                                            | Motion magnitude error: 1.86 ± 0.11 mm                                   |
| Commandeur 2020     | Prospective analysis of the capability of ML-determined CAC score and other variables in predicting MI or cardiac death. | Extreme gradient boosting                           | 1912 participants without cardiovascular disease | AUC                                                                 |
|                     |                                                                                |                                                    |                                            | - ML: 0.82                                                            |
|                     |                                                                                |                                                    |                                            | - ASCVD: 0.77                                                          |
|                     |                                                                                |                                                    |                                            | - CAD: 0.77                                                           |
| Al’Aref 2020        | ML model using CAC and clinical factors to improve prediction of obstructive CAD. | Boosted ensemble algorithm                         | 35 281 patients (CONFIRM registry) (80% training, 20% testing)          | AUC                                                                 |
|                     |                                                                                |                                                    |                                            | - ML: 0.77                                                           |
|                     |                                                                                |                                                    |                                            | - CAD consortium clinical score: 0.73                                  |
|                     |                                                                                |                                                    |                                            | - CAC score: 0.87                                                     |
|                     |                                                                                |                                                    |                                            | - UDF score: 0.68                                                     |
| Glowacki 2020       | ML model prediction of obstructive CAD following CAC score.                     | Gradient boosting machine learning                  | 435 patients                                | Sensitivity 100 ± 0.0%                                                 |
|                     |                                                                                | Binary logistic regression                          |                                            | Specificity 69.8 ± 3.6%                                                |
| Lee 2020            | Retrospective analysis to ascertain best ML algorithm to predict CAC score from clinical variables. | - XGBoost: 0.82                                     | 2133 participants without cardiovascular disease | AUC                                                                 |
|                     |                                                                                | - CatBoost: 0.75                                    |                                            | - XGBoost: 0.82                                                       |
|                     |                                                                                | - Binary logistic regression: 0.59                  |                                            | - CatBoost: 0.75                                                      |

Testing includes validation. Statistics are per patient.
ML, machine learning; CAC score, coronary artery calcium score; CNN, convolutional neural networks; AUC, area under the curve; ASCVD, atherosclerotic cardiovascular disease risk algorithm; CAD, coronary artery disease; UDF score, updated Diamond–Forrester score; ICC, intraclass correlation coefficient.
have demonstrated CNN algorithms that can automatically segment the LA with 99% accuracy vs. expert reader, and compartmentalize the LA into individual sub-sections using marginal space learning-based object segmentation with minimal error (Table 7).

Post-ablation recurrence of AF has a rate of ca. 45%; Firouzina et al. successfully used random forest classifiers to identify

Table 4 Summary of articles investigating the use of ML in cardiac CT determined plaque characterization

| Study       | Design and aim                                                         | Algorithm used                                      | Population | Outcome                      |
|-------------|------------------------------------------------------------------------|----------------------------------------------------|------------|------------------------------|
| Wei 2014²⁷  | Retrospective, automated detection of non-calciﬁed plaques, grouped by vessel diameter | Topological soft-gradient detection method          | 83 patients| AUC: 0.87 ± 0.01
|              |                                                                        |                                                    |            | Sensitivity: 70–90%
|              |                                                                        |                                                    |            | False-positive rate: 1.39–3.16 per scan
|              |                                                                        |                                                    |            | Information gain ratio
|              |                                                                        |                                                    |            | • Low-density non-calciﬁed plaques: 0.097
|              |                                                                        |                                                    |            | • Plaque length: 0.092
|              |                                                                        |                                                    |            | • Plaque volume: <0.001
| Dey 2018²⁸  | Prospective, multicentre trial performing semi-automated quantiﬁcation of calciﬁed and non-calciﬁed plaques, and plaque length and volume | Ensemble classiﬁcation approach with LogitBoost and single-node decision trees | 80 patients (90% training, 10% testing) |
| Masuda 2019²⁹ | Retrospective comparison of ML-determined plaque characterization vs. median CT number | Extreme gradient boosting                           | 78 patients| AUC
|              |                                                                        |                                                    |            | • ML: 0.92 (95% CI: 0.86–0.92)
|              |                                                                        |                                                    |            | • Median CT number: 0.83 (95% CI: 0.75–0.92)
| Zreik 2019³⁰ | Retrospective, detection, characterization and assessment of stenosis | Multi-task recurrent convolutional neural network | 163 patients (60% training, 40% testing) | Accuracy
|              |                                                                        |                                                    |            | • Detection and characterization: 0.77
|              |                                                                        |                                                    |            | • Stenosis: 0.80
| Al’Aref 2020³¹ | Case-control study identifying culprit lesions with multiple models | Boosted ensemble algorithm                          | 468 patients at high-risk of ACS (80% training, 20% testing) | AUC of best model: 0.77
| Han 2020³²   | Retrospective cohort study identiﬁcation of individuals at risk of rapid coronary plaque progression | Boosted ensemble classiﬁcation (LogitBoost)        | 1083 patients who underwent serial CTs in the PARADIGM registry (70% training, 30% testing) | (95% CI: 0.60–0.76)
| Muscogiuri 2020³³ | Automated categorization to Coronary Artery Disease Reporting and Data System (CAD-RADS) guidance using three models | CNN                                                 | 208 patients | Sensitivity: 47–82%
|              |                                                                        |                                                    |            | Speciﬁcity: 58–91%
|              |                                                                        |                                                    |            | Negative predictive value: 74–92%
|              |                                                                        |                                                    |            | Positive predictive value: 46–69%
|              |                                                                        |                                                    |            | Accuracy: 60–86%
| Tesche 2021³⁴ | Retrospective prognostication using clinical parameters and ML-derived plaque characteristics at 5-year follow-up | Boosted ensemble algorithm (RUSBoost)              | 361 patients with suspected CAD | AUC 0.96
| Yang 2021³⁵  | Retrospective prognostication using clinical parameters and ML-derived plaque characteristics at 5-year follow-up | Boruta algorithm and hierarchical clustering        | 1013 vessels | Sensitivity 0.97
|              |                                                                        |                                                    |            | Speciﬁcity 0.86
|              |                                                                        |                                                    |            | AUC for low FFR of best model: 0.797 (P < 0.001)

Testing includes validation. Statistics are per patient. 95% CI, 95% conﬁdence interval; CNN, convolutional neural network; MACE, major adverse cardiovascular events; ML, machine learning; AUC, area under the curve; CAD, coronary artery disease.
| Study                  | Design and aim                                                                 | Algorithm used                                      | Population | Outcome                                                                 |
|-----------------------|--------------------------------------------------------------------------------|-----------------------------------------------------|------------|-------------------------------------------------------------------------|
| Rodrigues 2016        | Prospective, automatic segmentation of mediastinal and epicardial adipose tissue using several algorithms compared with manual segmentation | CNN, probabilistic models, and decision tree algorithms | 20 patients | Random forest classification was superior Accuracy: 98.3% DSC for mediastinal and EAT: 0.98 |
| Norlén 2016           | Automatic pericardial segmentation and epicardial adipose tissue quantification vs. expert readers | Multi-atlas technique and random forest classification combined into a Markov random field | 30 examinations (SCAPIS study) (training 67%, testing 33%) | Pearson’s correlation vs. two experts: r > 0.998 Segmentation time: 52 s |
| Rodrigues 2017        | Prediction of mediastinal and epicardial adipose tissue volumes vs. expert readers | Rotation forest algorithm using multilayer perceptron Regressor | 50 examination images | Pearson’s correlation: 0.988 Relative absolute error: 14.4% Root relative squared error 15.7% |
| Commandeur 2018       | Fully automated assessment of mediastinal and epicardial adipose tissue vs. expert readers | CNN | 250 participants (80% training, 20% testing) | Pearson’s correlation • EAT: 0.924 • Mediastinal adipose tissue: 0.945 DSC • EAT: 0.823 • Mediastinal adipose tissue: 0.905 |
| Commandeur 2019       | Fully automated quantification and assessment of progression at follow-up of mediastinal and epicardial adipose tissue vs. expert readers | CNN with TensorFlow framework | 850 participants (80% training, 20% testing) | Pearson’s correlation vs. expert reader • Quantification: r > 0.973 • Progression at follow-up: r = 0.905 Quantification mean time: 1.57 s • Radiomic features linked to expression of inflammatory, fibrotic and vascularity genes • Fat radiomic profile provided superior MACE prediction at 5-year follow-up relative to traditional risk stratification • Fat radiomic profile elevated in patients with MI relative to matched controls |
| Oikonomou 2019        | Prediction of cardiac risk by analysis of radiomic profile of coronary perivascular adipose tissue (three studies) | Random forest | 312 patients | |
| Chernina 2020         | Retrospective, automatic vs. semi-automatic vs. expert radiologist for acquisition of EAT volume | 3D convolutional network | 452 (78% training, 22% testing) | Pearson’s correlation • ML vs. semi-automatic: r > 0.95 • ML vs. expert radiologists: r > 0.98 Median DSC pericardial fat: 0.88 Median DSC myocardium: 0.96 Consistency with contour, ICC: 0.97; P < 0.05 |
| He 2000b              | Retrospective, simultaneous myocardial and pericardial fat quantification | 3D deep attenuation U-Net (DAU-net) | 422 patients with suspected CVD (testing) | Sensitivity: 0.91 Specificity: 0.95 ML median DSC pericardial fat: 0.93 Manual control median DSC pericardial fat: 0.92 |
| He 2000a              | Retrospective, automatic vs. manual segmentation of epicardial adipose tissue | 3D deep attenuation U-Net (DAU-net) | 200 patients | Demonstrated automated adipose tissue analysis. Median DSC pericardium/muscle: 0.96 |
| Kroll 2021            | Retrospective comparison of CAC scores and pericardial fat in coronary calcium CT scans | Multi-resolution U-Net 3D network | 1066 patients at intermediate risk of CAD (9% training, 91% testing) | |

*Testing includes validation. Statistics are per patient. Accuracy was defined in Rodrigues* as (true positive + true negative/total population). CNN, convolutional neural networks; DSC, dice similarity coefficient; EAT, epicardial adipose tissue; MACE, major adverse cardiovascular events; MI, myocardial infarction; ML, machine learning.*
| Study            | Design and aim                                                                 | Algorithm used                                      | Participants | Outcome                                                                                                                                                                                                 |
|------------------|--------------------------------------------------------------------------------|-----------------------------------------------------|--------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Grbic 2013[46]   | Retrospective, automated prediction of aortic annulus perimeter and area       | —                                                   | 11           | Accuracy: 1.30 ± 0.03 mm Predicted implant size error: 1.75 ± 0.4 mm Aortic annulus error: 1.32 mm ‘errors in predicted implant deployment were of 1.74 ± 0.4 mm in average and 1.32 mm in aortic valve annulus region, which is almost three times lower than the average gap of 3 mm between consecutive implant sizes.’ |
| Elattar 2014[47] | Automated segmentation of the aortic root                                    | Connected component analysis and fuzzy classification | 20           | DSC • ML: 0.74 ± 0.39 mm • Expert reader: 0.68 ± 0.34 mm Time • ML: 12 min Expert reader: 20 min                                                                                                                                 |
| Liang 2017[48]   | Automated reconstruction of the aortic valve                                  | Neighbour-constrained segmentation                  | 10           | Mean discrepancy ML vs. expert reader: 1.57 mm                                                                                                                                                           |
| Al Abdullah 2018[49] | Automated identification of aortic valve landmarks                           | Randomized regression tree-based algorithm (colonial walk) | 71           | Mean localization error: 2.04 mm Inter-observer variability: 2.38 mm Time • ML: 12 mss • Expert reader: 4 min                                                                                                                                 |
| Astudillo 2019[50] | Retrospective, automated prediction of aortic annulus perimeter and area     | CNN                                                 | 473 patients (75% training, 25% testing) | Difference between predicted values and device size selected: Area • ML: 3.3 ± 16.8 mm² • Expert reader: 1.3 ± 21.1 mm² Perimeter • ML: 0.6 ± 1.7 mm • Expert reader: 0.2 ± 2.5 mm The difference between manually obtained aortic annulus measurements and those produced by the automated method were comparable to intra-operator variability |
| Theriault-Lauzier 2020[51] | Automated location and orientation of the aortic valve annular plane          | CNN                                                 | 94 patients with severe AS | Relative measurement error • Annular area: 4.73 ± 5.32% Annular perimeter: 2.46 ± 2.94%                                                                                                                     |
| Agasthi 2021[52] | Retrospective, predictive modelling of 1-year life expectancy of TAVR candidates | Gradient boosting ML (caret R package)              | 1055         | AUC 1 year: 0.72                                                                                                                                                                                          |
| Kang 2021[53]    | Predictive modelling to diagnose AS using CT features of aortic valve calcium | Least absolute shrinkage and selection operator (LASSO), random forests, and eXtreme Gradient boosting (XGBoost) | Retrospective study of 408 patients (240 with and 168 without severe AS) | 3/9 radiomics prediction models were successful in showing greater ability to distinguish AS. Differences for all models were not statistically significant (P > 0.05)                                                                 |
| Maeda 2021[54]   | Retrospective, predictive modelling of life-expectancy of TAVR candidates     | Cox proportional hazard regression                  | 388 (259)    | AUC 1 year: 0.79                                                                                                                                                                                          |
morphological traits on 3D fractal features to predict the risk of AF recurrence from pre-ablation contrast CTs (AUC: 0.87). This is likely because LA wall thickness and scarring depth that can be detected pre-procedure, relate to ablation success. Atta-Fosu et al. employed a similar technique using Gradient boosted classifiers (XGBoost) and found a lower AUC for shape alone (0.67) that was similar when combined with clinical features (0.78). In addition, it has been reported that post-ablation AF recurrence secondary to non-PV triggers can also be predicted with a similarly high degree of performance (AUC: 0.88). Given the utility of LA volumes measurements obtained by cardiac CT, it has been incorporated into a recently validated ATLAS score to predict AF recurrence after first PV isolation radiofrequency PV isolation ablation. Indeed, the application of CNN algorithms to the measurement of LV volume on

**Table 6** Continued

| Study            | Design and aim                                                                 | Algorithm used | Participants | Outcome                  |
|------------------|-------------------------------------------------------------------------------|----------------|--------------|--------------------------|
| Shirakawa 2021   | Proof-of-concept automated precise segmentation from CT of cardiac structure in the pre-operative assessment of patients with HOCM | CNN            | training, 129 testing) | 3 years: 0.76 5 years: 0.78 |
|                  |                                                                                |                | 2            | ML segmentation was ca. 36 faster |

| Testing includes validation. Statistics are per patient. ML, machine learning; DSC, dice similarity coefficient; AUC, area under the curve; CNN, convolutional neural network; HOCM, hypertrophic obstructive cardiomyopathy.

| Study          | Design and aim                                                                 | Algorithm used | Population | Outcome                  |
|----------------|-------------------------------------------------------------------------------|----------------|-------------|--------------------------|
| Zheng 2014     | Retrospective subsection segmentation of the left atrium                      | Marginal space learning-based object segmentation | 687 datasets | Mean mesh error |
| Bratt 2019     | Retrospective prediction of AF using left atrial volume vs. expert reader      | CNN (U-Net)    | 1000 patients undergoing routine CT thoraces (50% training, 50% testing) | AUC: 0.77 (95% CI: 0.71–0.82) |
| Chen 2020      | Retrospective detection and segmentation of the left atrium vs. expert reader | CNN (U-Net)    | 518 patients who underwent pulmonary vein ablation | Accuracy: 99.0% Sensitivity 99.3% Specificity: 98.7% |
| Liu 2020       | Retrospective prediction of post-ablation AF recurrence due to non-pulmonary vein triggers | CNN (U-Net) (ResNet34) | 521 patients (73% training, 27% testing) | AUC: 0.88 ± 0.07 Accuracy: 88.6% ± 2.3 Sensitivity 75.0% ± 5.8 Specificity 95.7% ± 1.8 |
| Firouznia 2021 | Retrospective prediction of post-ablation AF recurrence using morphological analysis of the left atrial myocardium and pulmonary veins | Random forest | 203 patients | AUC: 0.87 (95% CI: 0.82–0.93) |
| Deepa 2021     | Prospective ML detection of epicardial fat within the left atrium              | CNN            | 10 patients | Accuracy: 89.22% Sensitivity: 90.18% Specificity: 88.52% |
| Atta-Fosu 2021 | Retrospetive investigation of left atrial shape differences and prediction of post-ablation AF recurrence | Gradient boosted classifier (XGBoost) | 68 patients | AUC for shape features from the SOI: 0.67 AUC for clinical parameters: 0.71 |

| Testing includes validation. Statistics are per patient. AUC, area under the curve; AF, atrial fibrillation; CNN, convolutional neural network; DSC, dice similarity coefficient; ML, machine learning; SOI, shape of interest.
routine non-gated chest CT have been able to effectively predict AF. Given the morbidity and mortality associated with undiagnosed paroxysmal AF and the increasing use of thoracic CT imaging this may be a worthwhile add-on.

Discussion and limitations

Given the black-box nature of commercial ML tools, we may not be able to fully analyse the reasoning behind the outputs of these complex models, and as such may not easily identify implicit biases within a given dataset or methodology. Algorithms lack context and causality for their predictions. This may be less of an issue for algorithms which aim to automate calcium measurements but would be very significant for example in predictive algorithms for AF status or neural networks to simulate device biomechanics for TAVR.

Candidate selection and accurate labelling for the training of models are the most crucial steps in the development of ML protocols. Disparities in these factors between studies may explain variability in results demonstrated in Tables 2–7. Utilizing large multicentre studies, such as Nakanishi et al.,73 in predicting coronary heart disease events from CTS from the Multi-Ethnic Study of Atherosclerosis cohort, or Coenen et al.,6 for assessing the diagnostic accuracy of CT-FFRML within the MACHINE consortium, is a useful start in the optimization of models for a broader patient population and may account for labelling issues in training datasets. Reproducibility can also be hampered by a requirement for specific CT scanner capabilities, the use of distinct imaging protocols, and other methodological heterogeneity. Machine learning has already been applied to automate image quality assessment in CCTA studies in a reproducible manner, which may provide a tool to stratify clinical trials to the levels of image quality. Another challenge that is apparent from the findings of this review is the lack of standardization in metrics used to analyse outcomes (i.e. AUC, dice coefficients, or accuracy). Though the chosen metric is matched to the task it is undertaking, for example, AUC for classification or DC for segmentation, this hampers comparability. Many ML models exist to address the same task with varying metrics of performance and results, as evidenced by Tables 2–7. Approaches such as that undertaken by Lopes et al.,74 who compared several ML models on a single large standardized dataset, need to increasingly be undertaken to provide more insights into an optimal methodology for diagnostic and prognostic reliability. With the introduction of a new datasets, the models will need to be continually retrained and in so doing new features may need to be accounted for.

Conclusion

Application of ML protocols to cardiac CT output has many benefits in automating time-consuming calculations, risk stratification and prognostication, and in pre-operative procedure planning across several pathologies including CAD, epicardial adiposity quantification, AF, and AS. Machine learning provides exciting advances in CCT- and CCTA-based calcium scoring and in near real-time analysis of flow-limiting lesions on CT-FFR. ML-CT-derived measurements and predictive prognostics may assist patient selection for radiofrequency ablation in patients with refractory AF. ML-CT may guide device selection and improve pre-procedural processes for TAVR candidates. Though far from replacing the bedside physician, efforts to incorporate these novel models into clinical practice may reduce time and resources while at the same time improving patient outcomes.

Lead author biography

Dr Jonathan J. H. Bray is an Academic Junior Doctor and The Training Manager for the British Junior Cardiologists Association (BJCA) Starter committee. Jonathan intercalated in Physiological Sciences at the University of Bristol in 2016. He has published 17 peer-reviewed articles, given seven international or national presentations and been awarded almost £3000 as part of a number of awards and prizes. He is a solicited peer review of several high impact journals and contributes as an author to the Cochrane collaborative. He teaches three courses at Cardiff and Swansea University as Honorary Tutor and Research Fellow.

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