Machine learning vs. conventional statistical models for predicting heart failure readmission and mortality

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Abstract

Aims This study aimed to review the performance of machine learning (ML) methods compared with conventional statistical models (CSMs) for predicting readmission and mortality in patients with heart failure (HF) and to present an approach to formally evaluate the quality of studies using ML algorithms for prediction modelling.

Methods and results Following Preferred Reporting Items for Systematic Reviews and Meta-Analyses guidelines, we performed a systematic literature search using MEDLINE, E-PUB, Cochrane CENTRAL, EMBASE, INSPEC, ACM Library, and Web of Science. Eligible studies included primary research articles published between January 2000 and July 2020 comparing ML and CSMs in mortality and readmission prognosis of initially hospitalized HF patients. Data were extracted and analysed by two independent reviewers. A modified CHARMS checklist was developed in consultation with ML and biostatistics experts for quality assessment and was utilized to evaluate studies for risk of bias. Of 4322 articles identified and screened by two independent reviewers, 172 were deemed eligible for a full-text review. The final set comprised 20 articles and 686 842 patients. ML methods included random forests (n = 11), decision trees (n = 5), regression trees (n = 3), support vector machines (n = 9), neural networks (n = 12), and Bayesian techniques (n = 3). CSMs included logistic regression (n = 16), Cox regression (n = 3), or Poisson regression (n = 3). In 15 studies, readmission was examined at multiple time points ranging from 30 to 180 day readmission, with the majority of studies (n = 12) presenting prediction models for 30 day readmission outcomes. Of a total of 21 time-point comparisons, ML-derived c-indices were higher than CSM-derived c-indices in 16 of the 21 comparisons. In seven studies, mortality was examined at 9 time points ranging from in-hospital mortality to 1 year survival; of these nine, seven reported higher c-indices using ML. Two of these seven studies reported survival analyses utilizing random survival forests in their ML prediction models. Both reported higher c-indices when using ML compared with CSMs. A limitation of studies using ML techniques was that the majority were not externally validated, and calibration was rarely assessed. In the only study that was externally validated in a separate dataset, ML was superior to CSMs (c-indices 0.913 vs. 0.835).

Conclusions ML algorithms had better discrimination than CSMs in most studies aiming to predict risk of readmission and mortality in HF patients. Based on our review, there is a need for external validation of ML-based studies of prediction modelling. We suggest that ML-based studies should also be evaluated using clinical quality standards for prognosis research. Registration: PROSPERO CRD42020134867

Keywords Machine learning; Heart failure; Readmission; Mortality; Hospitalization; Death; Statistical models; Prognosis

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Introduction

Despite major technological advances in the diagnosis, assessment, and management of cardiovascular disease, heart failure (HF) remains a major global public health concern, with an estimated prevalence of 64 million individuals around the world. HF hospitalizations have more than tripled in the last 30 years and are associated with high mortality. HF results in a substantial financial strain on the public health-care system and critically impairs the quality of life of those afflicted with it.

Patients with HF present with diverse clinical profiles such that physicians must assess a wide range of data to make fulsome assessments of their patients and to appropriately manage and predict prognosis. Artificial intelligence is a rapidly growing field in cardiovascular medicine that may aid in organizing past, current, and incoming data. Machine learning (ML) is a sub-field of artificial intelligence that comprises several algorithms such as artificial neural networks, random forests, decision trees, and other supervised or unsupervised models. These algorithms utilize existing and incoming data to identify patterns and predict future clinical events. Many studies have investigated the role of ML in the diagnosis of HF from electronic health records. However, we are still nascent in our understanding of the potential applications of ML in other aspects of patient management.

While there is interest in whether ML could improve our ability to predict outcomes, there have been few studies that have systematically reviewed the current literature comparing it with conventional statistical models (CSMs). To date, there is no systematic review that quantitatively compares ML with CSMs in HF prognosis. This systematic review presents an approach to summarize results graphically and a novel approach to formally evaluating the quality of these studies. The objective of this study was to review the performance of ML methods compared with conventional statistical models in the prognosis of hospitalized HF patients.

Methods

Literature search

This systematic review complies with Preferred Reporting Items for Systematic Reviews and Meta-Analyses (PRISMA) guidelines (Table S1). A comprehensive systematic literature search was performed in MEDLINE, EPUB, Cochrane CENTRAL, EMBASE, INSPEC, ACM, and Web of Science electronic databases for articles published between 1 January 2000 and 26 July 2020. The search terms included synonyms of HF such as cardiac failure, myocardial edema, and cardiac insufficiency, combined with synonyms or subcategories of ML terms such as neural networks, expert systems, and support vector machines. Search terms and keywords used in our search strategy are provided in Table S2. All titles and abstracts were manually filtered for outcomes of hospitalizations, readmissions, mortality, and related terms for inclusion in the study. If required, for clarity, we conducted a full-text review of the document to determine if relevant outcomes were examined. No restrictions were applied on language or sex.

Study selection

Two reviewers (S. S. and M. M.) independently screened all titles and abstracts. Only primary articles that compared ML with CSMs in the prognosis of initially hospitalized HF patients were considered for inclusion. This criterion was instituted to allow for the evaluation of hospital readmission outcomes and for a clear inception point for the assessment of mortality. Among studies deemed potentially relevant for a full-text review, articles were excluded if (i) the full-text manuscript could not be accessed or they were conference or symposium abstracts, (ii) the paper did not assess hospitalized HF patients, (iii) there was no comparison between ML and CSMs, or (iv) the outcomes examined did not include mortality or readmission. Discrepancies were resolved by the consensus of a group of reviewers (C. F., D. C., G. T., H. A. Q., and D. S. L.).

Data extraction

Data extraction included (i) author name and year of publication, (ii) country of data origin, (iii) specific patient population, (iv) distribution of age and sex in cohort, (v) sample size of developmental/derivation/training cohorts and validation/testing cohorts, (vi) internal and external validation methods, (vii) outcome of interest, (viii) ML algorithm, and (ix) classification and performance statistics (e.g. hazard ratio value, odds ratio, p-value, c-statistics, and calibration). Where the outcome was a composite of death or readmission, the study was included in analyses for both mortality and readmission. As model performance is often over-optimistic in the dataset in which it was derived, we used the c-indices from the validation set for our analyses. We also extracted the type of CSM methods employed, defining these approaches as logistic regression, Poisson regression, or Cox proportional hazards regression models.

Quality assessment

Two reviewers independently assessed the quality of each included study using a modified version of the CHARMS checklist, which is a validated review tool for quality evaluation. The CHARMS checklist was selected because it is applicable to both clinical studies and those published in the ML
Modifications to the CHARMS tool were made in consultation with experts in ML, epidemiology, and biostatistics, using frameworks developed in previously published studies. Studies received an overall score of low, moderate, or high risk of bias based on seven domains: (i) source of data, (ii) outcomes, (iii) candidate predictors, (iv) sample size/missing data, (v) attrition, (vi) model development, and (vii) model performance/evaluation. For the criterion of model evaluation, external validation was defined in the narrow sense, as previously described by Reilly and Evans, and defined by the Evidence-Based Medicine Working Group. A detailed description of the modified CHARMS checklist is available in Tables S3 and S4.

Statistical analysis

Continuous variables were reported as mean (standard deviation) or median (inter-quartile range) as published in the original reports. Categorical variables were reported as proportions. Extracted c-indices were not combined or pooled across studies because standard errors of the individual c-indices were not reported consistently in the original publications. Therefore, we constructed scatterplots with the c-indices for the CSMs on the x-axis and ML on the y-axis to show graphically the distributions between studies. We also determined the difference, \( \Delta \text{c-index} = \text{c-index}_{\text{ML}} - \text{c-index}_{\text{CSM}} \), and classified studies into three groups on the basis of \( \Delta \text{c-index} \leq 0 \) or \( > 0 \) but \( \leq 0.05 \), or \( > 0.05 \).

Results

Study characteristics

The initial literature search yielded 4322 articles, of which 3309 remained after exclusion of duplicates, and these articles underwent title and abstract screening. Full-text screening was performed for 172 prognostic studies with 20 articles included in the final set. The PRISMA flow chart is shown in Figure 1. The characteristics of each included study are shown in Table S5. In the final inclusion set, most studies were published from 2015 onward \( [n = 18 \ (90\%)] \), and more than half of the studies were from the USA \( [n = 11 \ (55\%)] \). Two studies utilized data from registry datasets, while the remainder used multicentre clinical datasets. The total sample size of the 20 studies was 686 842 patients. The weighted average age of reported means or medians of patients across all studies was 74 years, and the weighted proportion of women was 49%. Fifteen articles reported readmission outcomes, seven articles reported mortality outcomes, and two articles reported composite outcomes.

Machine learning methods

Seventeen of 20 studies incorporated a tree-type ML algorithm (Table S6). Two studies assessed survival time using random survival forests, which is an extension of the random forest ML approach. Support vector machines were the second most utilized ML algorithm \( (n = 9) \), followed by neural networks \( (n = 7) \). The remaining studies presented combinations of these with other ML algorithms such as deep learning, Bayesian techniques (including naïve Bayes classifiers and Bayesian networks), and \( K \)-nearest neighbour. Several studies employed multiple ML algorithms and compared them with one or more CSMs. Many studies took advantage of ensemble learning algorithms, which are ML techniques that aggregate the outcomes of multiple-based trained models, producing a unified general result for each data sample (e.g. random forests, gradient boosting machine, and boosted classification tree; Table S6). Ensemble learning techniques can be very powerful but lack interpretability, which is crucial in biomedical studies.

Conventional statistical model approaches

Logistic regression models were employed in 16 studies, three studies employed Cox regression models, and three studies employed a Poisson regression model. Some studies compared ML with previously derived, clinically validated models, including the Poisson-based Meta-Analysis Global Group in Chronic (MAGGIC) HF model, the Cox regression-based Get With The Guidelines HF (GWTG-HF) model, Seattle Heart Failure Model, MUerte Subita en Insuficiencia Cardiaca (MUSIC) risk score, SENIORS model, and the logistic regression-based LACE index. Nearly all studies that compared ML with one of the previously developed validated models also conducted comparisons with re-fit statistical models using one of the three CSM approaches described earlier. Only one study compared ML exclusively with one of the aforementioned clinical prediction models without developing de novo CSMs or re-fitting the model covariates in the new dataset.

Readmission

For the outcome of readmission, most studies reported superior performance using ML compared with CSMs. Of 15 studies examining readmission outcomes, 11 reported higher c-indices using ML, and one study reported higher c-indices at some (but not all time points). When each time point was counted separately, there were 21 comparisons, of which higher c-indices were reported using ML in 16. These outcome studies are depicted in Figure 2, where the dashed
line (blue diagonal) indicates equivalence between ML and CSMs. Figure 3 shows the magnitude of the differences (delta) between ML and CSMs. Four studies comprising comparisons at eight different time points reported a Δc-index > 0.05.

**Quality assessment**

Upon assessment using the modified CHARMS checklist (Tables S3 and S4), all studies demonstrated moderate to high risk of bias in at least three major categories of quality assessment. Low risk of bias was demonstrated in two major categories for 90% of the studies: ‘source of data’ and ‘outcomes’ (Table 1). However, 95% of studies demonstrated high risk of bias in the ‘sample size and missing data’ and ‘attrition’ (i.e. completeness of follow-up) domains. All studies demonstrated moderate risk of bias in the ‘model performance and evaluation’ domain. Critically, only one study performed a comparison of ML and CSMs in an entirely different external validation dataset. This study reported c-indices of 0.913 for ML, 0.835 for logistic regression, and 0.806 for MAGGIC. All other studies performed internal validation in the same dataset where the ML models were derived by...
cross-validation ($n = 11$), random split sample ($n = 8$), chronological split sample ($n = 2$), or bootstrap resampling ($n = 3$) techniques.

**Calibration**

Notably, although calibration is a component of model evaluation that is mentioned in the CHARMS checklist, only two studies reported ML and CSM calibration results.$^{18,30}$ Austin et al. found that neither CSMs nor ML had uniformly superior calibration; however, both logistic regression and random forests resulted in good calibration among subjects with a lower predicted probability of death.$^{30}$ Frizzell et al. found that while logistic regression was well calibrated, some ML methods (e.g. tree-augmented Bayesian network and gradient boosting model) were poorly calibrated when predicted readmissions were higher, and the miscalibration was observable in the validation set.$^{18}$

**Discussion**

In this systematic review, we found that ML methods had better performance than CSMs for prediction of readmission and mortality among patients with HF. All studies applied supervised learning algorithms predicting readmission and/or mortality. The most used method of supervised ML was tree-type ML algorithms, and logistic regression modelling was the most frequent conventional statistical approach. Unsupervised ML algorithms, which provide inputs with no pre-specified outcomes, were not utilized in any of the identified studies reviewed. Of the comparative studies, 90% demonstrated high risk of bias in at least two major domains of the modified CHARMS checklist, with $>60\%$ demonstrating high risk in at least three major domains. Importantly, most studies showing higher $c$-indices performed internal validation but lacked external validation (in even a narrow sense) in an independent dataset. Additionally, only a small minority of studies reported on calibration, which is an important component of predictive model development.

In the past decade, the incorporation of ML algorithms into prognostic models has increased. For example, multiple studies discuss the utility of ML in prognostic models for mortality following myocardial infarction.$^{30–32}$ ML has also been applied in cardiac diagnostics, to predict the occurrence of atrial fibrillation.$^{33}$ Recent reviews emphasize the tremendous interest in combining these techniques for clinical guidance and the need for additional prognostic studies.$^{34}$ The emergence of promising studies that leverage natural language processing or ML is illustrative of this burgeoning interest, but these early studies including patients with HF did not directly compare artificial intelligence algorithms with CSMs.$^{35,36}$

**Figure 2** Scatterplot of the highest reported $c$-index for machine learning and conventional statistical approaches for readmission studies. Circles of the same colour indicate different time points in the same study publication. CSM, conventional statistical model; ML, machine learning.
Figure 3 Cluster-bar plot of the difference in the highest reported c-index for machine learning and conventional statistical approaches for readmission studies. Bars of the same colour indicate different time points in the same study publication. Bars on the right side of the zero line indicate that c-indices were higher with ML; bars on the left of the zero line indicate c-indices were higher with CSMs. Outcomes: Ben-Assuli 2019 (1) = 90 day readmission; Ben-Assuli 2019 (2) = 60 day readmission; Ben-Assuli 2019 (3) = 30 day readmission; Mortazavi 2016 (1) = 30 day all-cause readmission, Mortazavi 2016 (2) = 30 day HF readmission, Mortazavi 2016 (3) = 180 day all-cause readmission, Mortazavi 2016 (4) = 180 day HF readmission, Sohrabi 2019 (1) = 1 month HF readmission, Sohrabi 2019 (2) = 3 month readmission. CSM, conventional statistical model; ML, machine learning.

| Study ID         | Source of data | Outcomes | Candidate predictors | Sample size/missing data | Attrition | Model development | Model performance/evaluation |
|------------------|----------------|----------|----------------------|--------------------------|-----------|-------------------|-----------------------------|
| Allam 2019       | L              | L        | M                    | H                        | H         | H                 | M                           |
| Austin 2010      | L              | L        | M                    | H                        | H         | L                 | L                           |
| Austin 2012      | L              | L        | L                    | H                        | H         | H                 | L                           |
| Avan 2019        | L              | L        | M                    | H                        | H         | M                 | M                           |
| Ben-Assuli 2019  | L              | M        | M                    | H                        | H         | M                 | M                           |
| Chen 2020        | L              | L        | H                    | H                        | H         | M                 | M                           |
| Frizzell 2017    | L              | L        | L                    | H                        | H         | L                 | M                           |
| Golas 2018       | L              | L        | M                    | H                        | H         | L                 | M                           |
| Kwon 2019        | L              | L        | M                    | H                        | H         | L                 | M                           |
| Liu 2020         | L              | L        | H                    | H                        | H         | L                 | M                           |
| Mahajan 2016     | L              | L        | H                    | H                        | H         | M                 | M                           |
| Mahajan 2020     | L              | L        | H                    | H                        | H         | M                 | M                           |
| McKinley 2019    | L              | M        | M                    | H                        | H         | M                 | M                           |
| Miao 2018        | L              | L        | M                    | H                        | H         | L                 | M                           |
| Mortazavi 2016   | L              | L        | M                    | H                        | H         | L                 | M                           |
| Padhukasahasram 2015 | L | M | H | H | H | L | M |
| Sohrabi 2019     | L              | M        | H                    | H                        | H         | H                 | H                           |
| Turgeman 2016    | L              | L        | H                    | H                        | H         | M                 | M                           |
| Wang 2020        | L              | L        | M                    | H                        | M         | M                 | H                           |
| Yu 2015          | L              | L        | H                    | H                        | H         | L                 | M                           |

H, high risk of bias; L, low risk of bias; M, moderate risk of bias.
While the interest in using ML in health care is growing exponentially, few studies have evaluated if it can potentially surpass CSMs in predictive performance. Christodoulou et al. compared ML algorithms with logistic regression in primarily non-cardiac disease conditions, and they found that ML did not perform better than logistic regression. However, this study explored health-care outcomes other than mortality or readmission, included diagnostic studies, did not include studies that utilized Cox regression or Poisson regression, and did not capture studies published in the computer science literature. In the specific clinical context of predicting mortality from gastrointestinal bleeding, a systematic review demonstrated higher c-indices and predictive capacity with ML compared with clinical risk scores. Another specific study aiming to predict bleeding risk following percutaneous coronary intervention reported that ML better characterized bleeding risk than a standard registry model. An important distinction of our review was that we included disease-specific studies that were published in the computer science literature, which were underrepresented in the aforementioned earlier comparisons of ML and CSMs. Finally, a recent comparison of ML vs. CSMs using the TOPCAT trial dataset found that ML methods had higher c-indices than CSMs for both readmission (0.76 vs. 0.73) and mortality (0.72 vs. 0.66). However, the study was restricted to heart failure with preserved ejection fraction in the setting of an ambulatory clinical trial and did not examine hospitalized HF cohorts and readmission outcomes, which were the focus of our report. Additionally, although a separate external validation was not performed, overall, the higher c-indices with ML were consistent with our reported findings.

CSMs have been used successfully in the clinical setting, and ML offers promise for clinical use. ML allows rapid examination of constantly expanding datasets and allows identification of patterns and trends not readily visible to clinicians. The advantages provided by ML are that it is flexible, is nonparametric, does not require a data model for the probability distribution of the outcome variable, does not require pre-specification of covariates, and can handle a large number of input variables simultaneously. Indeed, in our review, the maximum number of variables (or features) that could be input simultaneously was >3500. In the context of cardiology, this may allow clinicians to utilize these algorithms for high-performing prognostic models to enhance care of HF patients. There has been an extensive amount of research suggesting that improved ability to predict risk and implement transitional care interventions may improve HF patient outcomes and reduce readmissions. With improved accuracy of predictive modelling, clinicians and other health-care providers may be better equipped to offer the best care for each patient using individualized predictive data. Clinical decisions and discharge care for HF patients may be guided more effectively to reduce adverse events and improve quality of life.

Our study highlights the heterogeneity that currently exists in the literature for prognostic studies using ML, where many studies often did not report confidence intervals or standard deviations for their performance measures. The heterogeneity in the rigour of evaluation also extended to the lack of a priori cut points when validating prediction algorithms using ML. Importantly, we found that the term ‘external validation’ in the ML literature often referred to methods that would be considered ‘internal validation’ methods (e.g. split sample, cross-validation, or bootstrap resampling) using CSMs and accepted epidemiological standards. Thus, there is a strong need for future studies of prognostication using ML to use standardized reporting protocols, where the definitions of risk strata and a clear distinction between the derivation/training and validation/testing sets are explicitly stated.

Recommendations

To improve the quality of reporting of future comparisons of ML and CSMs, we recommend that standard errors of the differences in the c-statistics between the two analytical approaches should be reported, to allow pooling of multiple studies meta-analytically. Second, the calibration of ML algorithms was not provided in the majority of studies, but it should be routinely reported. As future studies expand to include high-dimensional and -omics data sources, the potential applications for ML, predictive analytics, and advanced statistical learning techniques are likely to grow. However, our study suggests that the same rigorous principles of model development, internal validation, considering potential sources of bias, and external validation should apply. Furthermore, collaborations between computer scientists, biostatisticians, and clinicians are necessary to ensure that the high quality and standards applicable to clinical prediction rules are also applied to ML methods.

Limitations must be acknowledged. Owing to the heterogeneity of reported performance and descriptive statistics, only a narrative synthesis was possible for this study. In addition, we used a modified version of the CHARMS checklist to conduct quality assessment, a tool that was not originally constructed to assess the quality of ML studies or for comparison between ML and CSMs. We modified the CHARMS in consultation with ML and biostatistical experts in order to best suit the objectives of the study and incorporated the framework from a previously published modification of the CHARMS as well. Other available approaches to assessing quality of prognostic studies, such as the TRIPOD, were considered, but its aim is to increase transparency of reporting, and from an operational standpoint, it was not ideally suited to evaluate prognostic studies using ML. The modified CHARMS checklist does set a high bar for predictive models because it was originally developed for assessing applications for use in clinical practice. However, there is often little
distinction between a pre-clinical and clinical predictive model from a mathematical standpoint. Rather, the major distinction is whether the model has undergone validation—initially in a narrow sense, followed by broad validation, and then in an impact analysis. Some of the CSMs were previously developed models, whose performance may be inferior to purpose-build CSMs tested in the derivation sample. However, in these studies, we also examined the performance of purpose-build CSMs derived in the study dataset, which was available in all but one study reviewed. Finally, we cannot exclude the possibility of publication bias, whereby studies showing an advantage of ML could be more likely to be published. However, given the novelty of ML in the health science literature, we would anticipate that studies showing either better performance with CSMs or ML would merit publication irrespective of the directionality of effects.

In conclusion, our study has shown that ML methods demonstrated an overall stronger predictive performance over CSMs for HF prognosis. The heterogeneity in reported outcomes and descriptive statistics warrants the need for established standards of reporting for ML studies. In particular, it is important to externally validate ML models and demonstrate that performance is preserved in new cohorts if the intention is to utilize them clinically.

Conflict of interest

The authors have no relevant disclosures.

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Supporting information

Additional supporting information may be found online in the Supporting Information section at the end of the article.

Figure S1. Scatterplot of the highest reported c-index for machine learning and conventional statistical approaches for mortality studies.

Figure S2. Cluster-bar plot of the difference in highest reported c-index for machine learning and conventional statistical approaches for mortality studies. Bars on the right side of the zero-line indicate that c-indices were higher with ML; bars on the left of the zero-line indicate c-indices were higher with CSM.

Table S1. PRISMA guidelines.
Table S2. Literature search strategy.
Table S3. Modified CHARMS checklist.
Table S4. Additional CHARMS criteria.
Table S5. Characteristics of included studies.
Table S6. Machine learning algorithms.

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