

Atrial fibrillation risk prediction from the 12-lead electrocardiogram using digital biomarkers and deep representation learning

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Aims

This study aims to assess whether information derived from the raw 12-lead electrocardiogram (ECG) combined with clinical information is predictive of atrial fibrillation (AF) development.

Methods and results

We use a subset of the Telehealth Network of Minas Gerais (TNMG) database consisting of patients that had repeated 12-lead ECG measurements between 2010 and 2017 that is 1 130 404 recordings from 415 389 unique patients. Median and interquartile of age for the recordings were 58 (46–69) and 38% of the patients were males. Recordings were assigned to train-validation and test sets in an 80:20% split which was stratified by class, age and gender. A random forest classifier was trained to predict, for a given recording, the risk of AF development within 5 years. We use features obtained from different modalities, namely demographics, clinical information, engineered features, and features from deep representation learning. The best model performance on the test set was obtained for the model combining features from all modalities with an area under the receiver operating characteristic curve (AUROC) = 0.909 against the best single modality model which had an AUROC = 0.839.

Conclusion

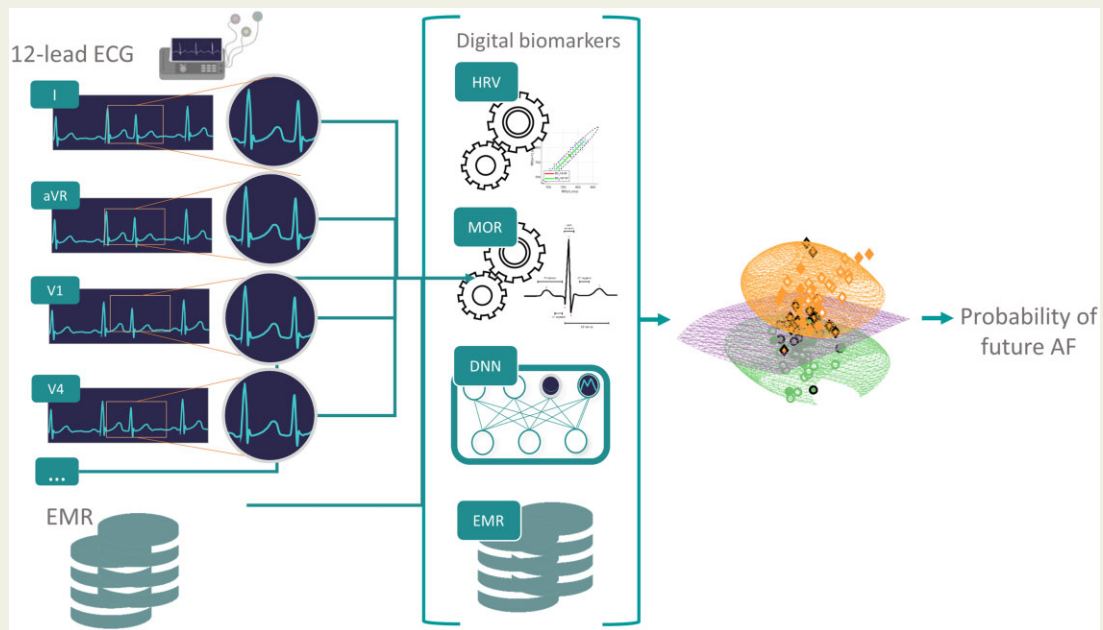
Our study has important clinical implications for AF management. It is the first study integrating feature engineering, deep learning, and Electronic medical record system (EMR) metadata to create a risk prediction tool for the management of patients at risk of AF. The best model that includes features from all modalities demonstrates that human knowledge in electrophysiology combined with deep learning outperforms any single modality approach. The high performance obtained suggest that structural changes in the 12-lead ECG are associated with existing or impending AF.

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Graphical Abstract



Keywords

Atrial fibrillation • Deep learning • Risk prediction

Introduction

Major cardiovascular and cerebrovascular events occur in individuals without known pre-existing cardiovascular conditions. Preventing such events remains a serious public health challenge. For that purpose, clinical risk scores can be used to identify individuals with high cardiovascular risks.^{1,2} However, available scoring scales have shown moderate performance.³ Despite being part of the routine evaluation of many patients in both primary and specialized care, the role of electrocardiogram (ECG) analysis in cardiovascular disease prediction and, hence, prevention is not clear.

Atrial fibrillation (AF) is the most common arrhythmia, with an estimated prevalence of 3% in adults.⁴ It is associated with quivering or irregular heartbeat that can lead to blood clots, stroke, heart failure, and other heart-related complications.⁵ Much interest has been given to developing novel algorithms for AF detection in the past decade.⁶ Because of its prevalence and clinical importance, AF is a good model for the development of new risk assessment algorithms. Wang and Wang⁷ highlight AF risk-prediction using big data and Machine learning (ML) as a new opportunity to improve AF management. Two recent work from Christopoulos *et al.*⁸ and Raghunath *et al.*⁹ made use of a deep learning approach using the raw 12-lead ECG signal as input for AF prediction. However, the deep learning approach limits the interpretation of the features found to be predictive of future AF. Furthermore, combining features engineered from the ECG morphology and heart rate variability (HRV) with deep representation learning features may improve the predictive performance. Overall, little research to date has investigated the feasibility of

cardiac abnormality risk prediction, i.e. predicting the future occurrence of a cardiac abnormality, using a data-driven approach based on the raw 12-lead ECG time series. Novel robust predictive models are likely to enhance clinical management of at-risk individuals and may also provide new insights into the aetiology of cardiac abnormalities. We introduce in this work a hybrid data-driven model combining demographics, feature engineering (or 'digital biomarkers') and deep representation learning obtained from the raw 12-lead ECG to predict the future occurrence of AF.

Methods

Database

The 12-lead ECG were obtained from the Telehealth Network of Minas Gerais (TNMG) between 2010 and 2017, a public telehealth system assisting 811 out of the 853 municipalities in the state of Minas Gerais, Brazil. The 12-lead ECG examination were performed mostly in primary care facilities using a tele-electrocardiograph manufactured by Tecnologia Eletrônica Brasileira (São Paulo, Brazil)—model TEB ECGPC—or Micromed Biotecnologia (Brasília, Brazil)—model ErgoPC 13. Recordings were also performed in emergency departments, ambulances and hospitals. Refer to Ribeiro *et al.*^{10,11} for more details. The original database had 1 773 689 patients. Patients less than 16 years old and with invalid ECG recording were excluded, resulting in 1 556 958 patients. The database used in this research is a subset from the TNMG that includes a total of $n = 1\,130\,404$ recordings from $m = 415\,389$ unique patients who had repeated 12-lead ECG examinations within the study period (2010–2017) and while also excluding patients with multiple recordings but all

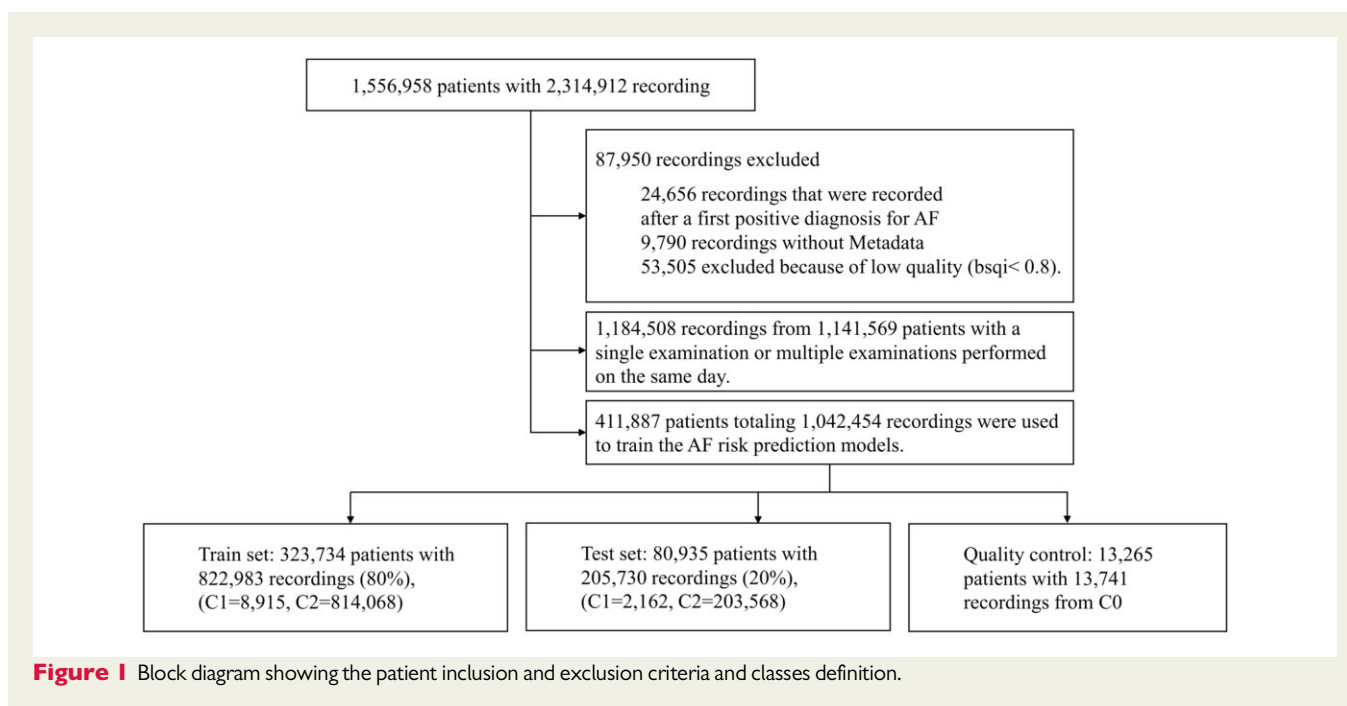


Figure 1 Block diagram showing the patient inclusion and exclusion criteria and classes definition.

being performed during the same day following the baseline examination. Within the selected subset, the median and interquartile number of recordings per patient was 2 (2–3), see [Supplementary material online, Figure S1](#). The median and interquartile age for the recordings were 58 (46–69), see [Supplementary material online, Figure S2](#), and 38% of the patients were males. The recording length is between 7 and 10 s and were originally sampled in frequencies ranging from 3 kHz and 1 kHz. We use the same pre-processing as Ribeiro et al.¹¹: we resample the recording at 400 Hz and zero-pad the signal so it becomes 4096 samples long (4096/400 Hz=10.24 s). All ECGs were interpreted by a team of trained cardiologists using standardized criteria,¹² in order to generate an ECG free text report. The reference diagnosis for the 12-lead ECG were originally obtained by testing the agreement between recognition of specific ECG diagnoses from the cardiologist reports (using natural language processing) and the automated diagnosis provided by the Glasgow ECG analysis software. In the case of disagreement, the decision was taken heuristically or through manual review.¹⁰

Considering the presence or not of AF, we defined C_0 as the class containing the recordings with a positive diagnosis for AF. A recording was included in class C_1 if it had a negative diagnosis for AF and there was, for the same patient, a positive AF diagnosis documented within 5 years, time interval distribution for those patients is presented in [Supplementary material online, Figure S3](#). A recording was included in class C_2 if it had a negative diagnosis for AF and there was no documented future positive diagnosis of AF for the corresponding patient. The mean follow-up time was 1.25 years. Recordings that were recorded after a first positive diagnosis for AF were discarded. This resulted in a total of 14 947 recordings in C_0 , 12 142 in C_1 and 1 078 545 in C_2 —see the block diagram in [Figure 1](#).

Data preprocessing

To automatically assess the quality of the raw ECG examples and discard flat or noisy examples, we included a signal quality preprocessing step (see [Supplementary material online, Note SN3](#)). Before computing the

ECG morphological biomarkers prefiltering of the raw ECG time series was performed. Specifically, we used a zero phase second-order infinite impulse response bandpass filter with the passband 0.67–100 Hz¹² to remove baseline wander and high frequency noise. We used a Notch filter at 60 Hz to remove the power-line interference (see [Supplementary material online, Figure S4](#)).

Digital biomarkers and representation learning features

Digital ECG biomarkers engineered from the heart rate variability (denoted 'HRV') and morphology of the ECG (denoted 'MOR') time series were engineered for each lead. In addition, the pretrained DNN¹¹ was used as a feature generator to include representation learning features (denoted 'DNN'). Briefly, the DNN structure of Ribeiro et al.¹¹ Consists of convolutional layer followed by four residual blocks with two convolutional layers per block. The output of the last block is fed into a fully connected layer with a Softmax function. The output of each convolutional layer is rescaled using batch normalization and fed into a rectified linear activation unit (ReLU). Finally, metadata (denoted 'META') were included as additional features. A summary diagram illustrating the overall 'hybrid' strategy taken is shown in [Figure 2](#). A total of 16 HRV-based features¹³ were engineered for each recording ([Supplementary material online, Table S1](#)). RR intervals were obtained from lead V1 using the epltd R-peak detector. For MOR features, a total of 36 different features were engineered; 12 features extracted from intervals duration ([Supplementary material online, Table S2](#)) and 24 from waves characteristics ([Supplementary material online, Table S3](#)) to describe the ECG morphology. These features were extracted for individual leads resulting in a total of 432 features. Some of these features were described in the work of Assaraf et al.¹⁴ and are based on the detection of fiducial points on the ECG waveform found using the popular open source wavelet algorithm.¹⁵ In addition, within the MOR feature set ([Table 1](#)), we included a measure of signal quality, bsqi, for each lead. Overall the number of MOR features per recording was 445. The DNN model presented in the

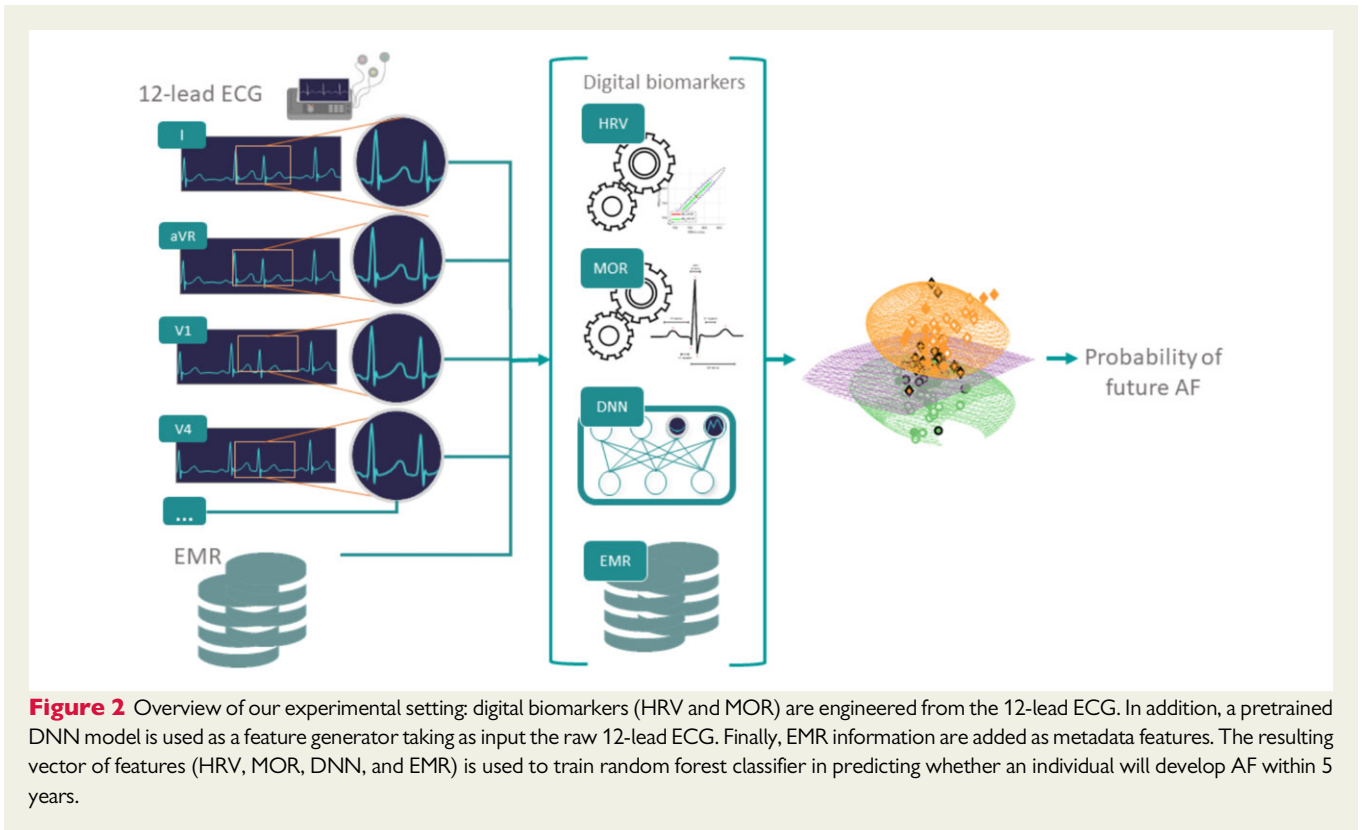


Table 1 The feature groups that compose each one of the presented models

	DNN	MOR	HRV	META	Number of features	Number of selected features
Model 1				X	15	—
Model 2			X		16	—
Model 3		X			445	80
Model 4	X				5120	340
Model 5	X	X	X		5581	400
Model 6	X	X	X	X	5596	180

Feature selection was performed using mRMR.

original work by Ribeiro *et al.*¹¹ was used as feature generators by using the features generated at the level of the first fully connected layer. For that purpose, the DNN was retrained using the 1 141 569 examples that had only 12-lead ECG recording(s) performed during the day of the baseline examination and thus were not used for our AF risk prediction experiment. In other words, the trained risk prediction models did not use any of the recordings used for training the DNN feature generator. Thus, there was no information leakage. For a 12-lead ECG recording, a total of 5120 DNN features were generated (Table 1). Finally, 15 META features (Supplementary material online, Table S4) were included (Table 1). This was to evaluate how patient information typically available from the hospital EMR may improve the model prediction. Although it is important to note that the META features used were self-reported by the patient at the time of the examination and thus are not directly obtained from an hospital EMR.

Performance statistics

The nonparametric Mann–Whitney rank test was used to determine whether individual features were significantly different between recordings in class C_1 and C_2 . The lower the P -value the stronger the evidence against the null hypothesis. For all features, a P -value cut-off at $P < 0.05$ was used for significance testing. In order to assess the performance of the models in correctly classifying individual examples the following statistics will be computed: sensitivity (Se), specificity (Sp), positive predictive value (PPV), the harmonic mean between the Se and PPV termed F_1 measure and the area under the receiver operating characteristic curve (AUROC). We estimated the confidence interval for AUROC using bootstrapping. That is, the AUROC was repeatedly computed on randomly sampled 80% of the test set (with replacement). The procedure was repeated 1000 times and used to obtain the intervals.

ML for risk prediction

Examples from C_1 and C_2 were assigned to train-validation and test sets in an 80:20% split and with stratification by class, age and sex, see [Supplementary material online, Table S5](#). ML models were trained to assess AF risk prediction defined as the ability of the classifier to predict the future development of AF within 5 years. A random forest (RF) classifier was trained with the biomarkers as input. To account for the class imbalance, the minority class was proportionally over-weighted. Feature selection was performed using the minimum redundancy maximum relevance (mRMR) algorithm.¹⁶ In brief, mRMR selects the set of relevant features while controlling for redundancy within the set of selected features. Hyperparameters were selected using BayesSearchCV algorithm implemented in the scikit-optimize library¹⁷ and five-fold cross-validation on the training set using 100 iterations. Hyperparameter and associated ranges for the search are listed in [Supplementary material online, Table S6](#). The behaviour of the model was also assessed on examples belonging to C_0 as it would be expected that the models predict a probability close to one for example in this class. A total of six models, listed in [Table 1](#) were evaluated. The threshold on the RF probabilistic output was defined on the ROC curves obtained in cross-validation and so that $Sp = 0.95$. Indeed, having some false positive (FP) predictions is acceptable because the consequences of interventions such as change in lifestyle, increases of some drugs to better treat risk factors such as diabetes will be minimal. Yet, the number of FP needs to be reasonable otherwise it would overburden the healthcare system with unnecessary follow-up examinations.

Survival analysis

We also performed Kaplan–Meier incidence-free survival analysis with the patients included in the test set and considering a new AF as the clinical endpoint. In order to build a per patient model (versus per recording), we only considered the model prediction for the baseline, AF free, 12-lead ECG ($n = 26\,982$). Patients who did not develop new AF were censored at the most recent encounter. A Cox proportional hazard ratio (HR) model was trained using as input variable the AF risk prediction model-predicted probability output. We computed the area under the

curve (AUC) for 5 years of new AF risk prediction, adjusted for age and gender. In a subsequent model, we adjusted for comorbidities and cardiovascular risk factors variables. In a first analysis, the patients were divided in five groups, according with quintiles of probability output of the ML model: (i) those with probability less than 0.2; (ii) those with probability between 0.2 and 0.4; (iii) those with probability between 0.4 and 0.6; (iv) those with probability between 0.6 and 0.8; and (v) those with probability greater than 0.8. In a second analysis, we considered two groups, based on the decision threshold found for the model.

Results

Overall, 53 505 out of 1 095 942 recordings (i.e. 4.9%) were excluded from the database after the signal quality step ([Figure 1](#)). [Table 2](#) summarizes the META data statistics for the patient's baseline recording. The patients in C_1 (72.0 ± 13.17) were significantly older than those in C_2 (55.0 ± 16.27) and there were significantly more males in C_1 (51.55%) vs. C_2 (37.61%).

Statistical analysis of the features

The Mann–Whitney rank test performed for individual features for C_1 and C_2 rejected the null hypothesis for 360 out of 445 MOR features, for 14 out of 16 HRV features and 13 out of 15 META features. Among the HRV features the minRR and PNN50 yielded the lowest P -values. Among the MOR features the Q, ST, J, and PR yielded the lowest P -values. Violin plots in [Supplementary material online, Figures S5 and S6](#) show the distributions of the HRV and MOR features with the lowest P -values. The median and standard deviation for HRV features and the top-20 MOR features are presented in [Supplementary material online, Tables S7 and S8](#), respectively. META features are summarized in [Supplementary material online, Table S9](#).

Table 2 The table summarizes the META data statistics

	C_0	C_1	C_2	P-value
Age (years), mean (SD)	72.00 (12.98)	72.00 (13.17)	55.00 (16.27)	<0.001*
Amiodarone, n (%)	197 (2.73)	165 (2.53)	2013 (0.51)	<0.001*
Diuretics, n (%)	2334 (32.3)	2072 (31.78)	85 025 (21.36)	<0.001*
Sex, male, n (%)	4011 (55.52)	3361 (51.55)	149 755 (37.61)	<0.001*
PAH, n (%)	3504 (48.5)	3198 (49.05)	143 902 (36.14)	<0.001*
Chagas, n (%)	432 (5.98)	384 (5.86)	9634 (2.42)	<0.001*
Beta blockers, n (%)	878 (12.15)	740 (11.35)	31 419 (7.89)	<0.001*
Family CHD, n (%)	1005 (13.91)	923 (14.16)	50 140 (12.59)	<0.001*
MI, n (%)	121 (1.67)	95 (1.46)	3207 (0.81)	<0.001*
COPD, n (%)	86 (1.19)	57 (0.87)	2423 (0.61)	<0.001*
DM, n (%)	561 (7.76)	542 (8.31)	28 633 (7.19)	<0.001*
Obesity, n (%)	335 (4.64)	391 (6.0)	22 868 (5.74)	0.001*
Dyslipidaemia, n (%)	287 (3.97)	268 (4.11)	15 363 (3.86)	0.031*
Calcium blockers, n (%)	0 (0.0)	0 (0.00)	95 (0.01)	0.155
Smoking, n (%)	367 (5.08)	387 (5.94)	24 627 (6.19)	0.486

It includes a summary for the baseline recordings in C_0 , C_1 , C_2 . With examples in C_0 being AF, examples in C_1 developing AF within 5 years and examples in C_2 not developing AF within 5 years. Mann–Whitney rank test was performed between C_1 and C_2 . Features ordered from most to least significant according to their P -value. The sign * indicates statistical significance (Mann–Whitney rank test, $P < 0.05$).

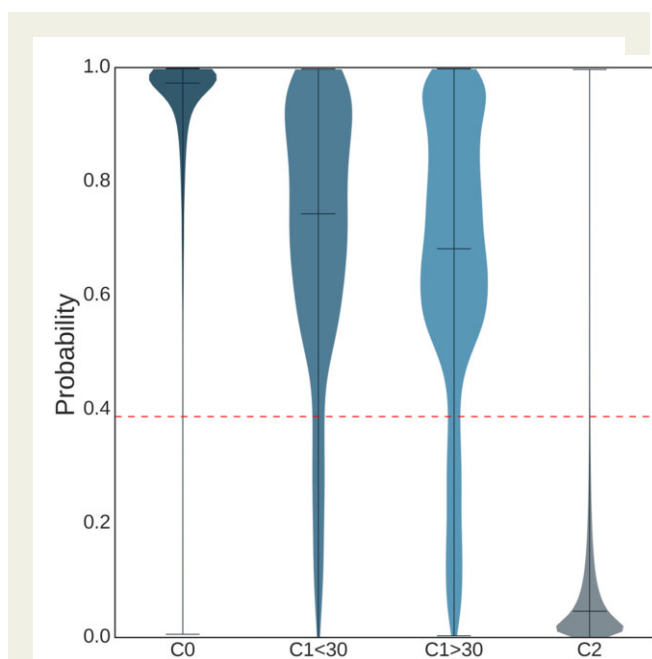


Figure 4 Violin plot showing the probability distribution for the output of the RF classifiers according to the reference class label, for 13 739 recordings in C_0 , 214 recordings in $C_1 < 30$, 1948 recordings in $C_1 > 30$ and 203 568 recordings in C_2 . The dotted red horizontal line depicts the probability decision threshold to classify an example as future AF or not.

Error analysis

Considering model 6, there was a total of 9517 FP and 873 false negative (FN). Violin plots representing the probability outputs for all FP per arrhythmia class are presented in [Supplementary material online, Figure S10](#). In particular, we observed that a large relative proportion of examples with 1dAVb (18.0%) and left bundle branch block (LBBB) (14.8%) rhythms were FP. The mean \pm standard deviation (SD) of time between the examination and a follow-up positive AF examination for true positive (TP) is 1.48 ± 1.23 years. Mean \pm SD for FN is 1.57 ± 1.20 years. A cardiologist (G.M.) with 5 years of clinical practice carefully reviewed 100 TP recordings within the test set blinded to any clinical or diagnosis information. Among the 100 TP, the cardiologist identified 50 recordings with a form of hypertrophy [Left ventricular hypertrophy (LVH)/Right ventricular hypertrophy (RVH)/Left atrial hypertrophy (LAH)/LVH], 24 recordings with AF/AFL, 16 recordings with premature atrial contraction (PAC)/premature ventricular contraction (PVC), 10 recordings with a form of bundle branch block (BBB) (LBBB/right bundle branch block), 25 recordings with other cardiac abnormalities, 4 recordings that were too noisy and could not be interpreted, and 2 recordings were normal sinus rhythm (NSR). It is important to note that a recording may present multiple cardiac abnormalities.

Survival analysis

The risk of development of AF for the two evaluated settings are shown in [Table 4](#) and adjusted survival curves in [Figure 5](#). In (A) the model indicates that patients with probability > 0.8 had higher risk of

Table 4 Risk of new AF

	HR	CI 95%	P – value
Adjusted by age and gender			
prob (0.2, 0.4]	5.222	4.343 – 6.279	<0.001
prob (0.4, 0.6]	12.063	9.933 – 14.651	<0.001
prob (0.6, 0.8]	20.967	17.053 – 25.779	<0.001
prob (0.8, 1]	43.764	36.186 – 52.928	<0.001
prob (0.387, 1]	10.719	9.389 – 12.237	<0.001

The table shows the hazard ratios (HRs) with for the different probability groups. The models were adjusted by different selection of variables. For the quintiles, the reference class is prob (0, 0.2]; for the dichotomic analysis, the reference class is (0, 0.387].

development AF (HR 43.77, 95% CI: 36.19–52.93; $P < 0.001$), considering as the reference those with probability less than 0.2. For b) the survival curve for probabilities > 0.387 had a higher risk of development AF (HR 10.72, 95% CI: 9.39–12.24; $P < 0.001$), compared to those with probability less or equal to the decision threshold. The Cox models adjusted by age and gender present a good performance in the prediction of new AF risk prediction, with an AUC of 0.87 (95% CI: 0.86–0.90) and (B) 0.84 (95% CI: 0.82–0.86). The adjusted survival curves for the model by different selections of comorbidities and cardiovascular risk factors are given in [Supplementary material online, Table S10](#). This analysis had similar results compared to the models adjusted by age and gender.

4. Discussion

Models performance

The results indicate that the ML models are indeed learning residual features that are predictive of the future development of AF. The best model was combining features from all modalities (META, HRV, MOR, and DNN) with AUROC = 0.909 (95% CI: 0.903 – 0.914). This highlights the value in combining engineered features that encapsulate Human knowledge in electrophysiology acquired over the past two centuries, DNN features which are less interpretable but enable to let the model to include features that are not intelligible to a human observer but are yet important for the task at hand, and finally metadata which encode demographic and medical information about the patient. The test set Se for individuals in C_0 was 0.970 and for individuals in C_1 it was 0.596 when using the decision threshold of 0.387. Although we found a high gap between training and test set performances, this is not uncommon when using RF and the performance between validation sets and the test set were close as seen on [Table 3](#). Furthermore, the Cox analysis showed that for different groups of probability results from the ML model further indicated that the highest probability resulted in the highest risk of AF future development (see [Table 4](#) and [Figure 5](#)).

Error analysis

The cardiologist review of 100 TP examples highlighted that a large number (24%) of the recordings correctly predicted as future AF were actually AF at the time of the examination and thus were likely

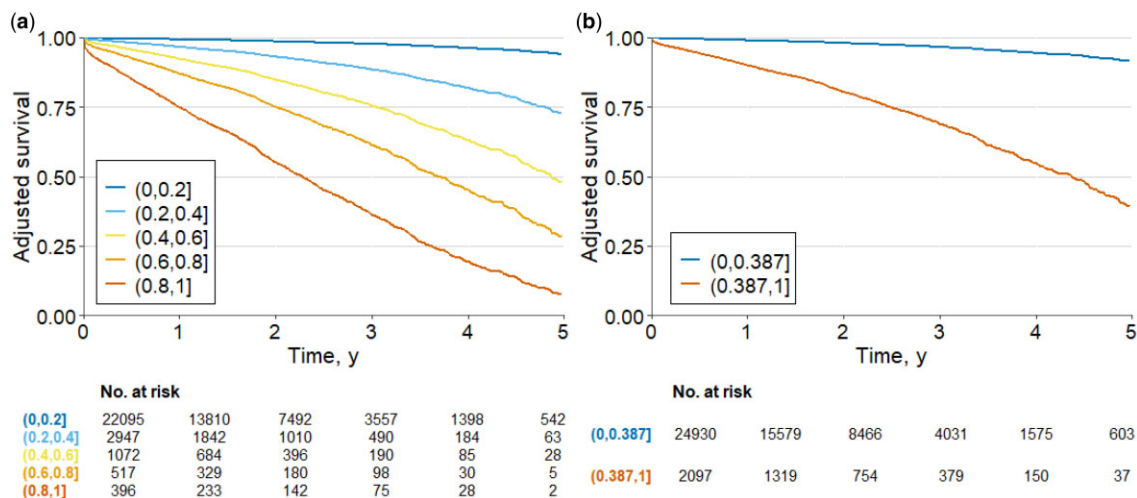


Figure 5 Adjusted survival curves for the risk of new AF. The plots display the survival curves for the different cohorts. The curves were computed from probability results of ML model. The models were adjusted by age and gender. In (A) the model with quintiles of probability and (B) the model with probability less or equal than 0.387 and those with probability greater than 0.387.

misdiagnosed. This number is coherent with the sensitivity of 0.769 reported in Ribeiro *et al.*¹⁰ for AF diagnosis by a cardiologist. Within this context, we foresee that our model may support the diagnosis of AF and avoid a significant number of misdiagnosis. Our second observation is that a number of TP had a type of hypertrophy, BBB or premature beats. This observation is coherent with previous reports that hypertrophy,¹⁸ ectopic beats,¹⁹ and BBB²⁰ are predictive of AF. Examples of three TP are displayed in Figure 6.

Clinical usability of the new ML model

The best model obtained AUROC = 0.909 (95% CI : 0.903 – 0.914), Se = 0.596, and Sp = 0.953. Practically speaking, with such a high Sp, and considering that a significant number of FP have another cardiac abnormality, this means that the number of FP is reasonable since intervention in FP will have limited consequences and the added need from the healthcare system is acceptable. The Se is more modest, yet it means that >50% of patients that will develop (or have a missed diagnosis of) AF can be identified which is a significant added value for the better management of these patients. Since examples in C₀ have AF then it is expected that the model will predict a high probability of AF risk and this was confirmed by our observations (Figure 4) where Se for individuals in C₀ was 0.970. Furthermore, when separating C₁ into two subclasses C₁(<30days) and C₁(≥30 days) we observed that the model indeed had a significantly higher probability distribution for C₁(<30 days) than C₁(>30 days)—see Figure 4 for the violin plot. In the work by Attia *et al.*²¹ individuals that developed AF within 31 days were considered as actually having AF at the baseline recording time but likely paroxysmal so that AF was not exerted at this time.

Overall, our findings are very promising. Predicting AF in a 12-lead ECG, a low cost and widely available exam, can help physicians in decision making. The model prediction could be used in the following fashion: patients with a high probability of future AF should first

undergo a careful review for the presence of AF. Eventually a second 12-lead ECG may be acquired at the same examination date in the case of doubt. This should result in lowering the number of misdiagnosis. Patients with no visible AF but with a high probability of future AF are at risk of imminent appearance of AF in a likely already diseased patient with paroxysmal AF and thus should be referred for a Holter ECG. If no Holter is available then a follow-up 12-lead ECG within the next few days should be scheduled. Individuals with no visible AF but predicted as future AF have a high risk to develop AF within the next 5 years and should regularly be examined. Although the model prediction has a good performance, the results are preliminary and it is still premature to make any changes in the current AF recommendations. Further research and external validation are necessary, but our data are a step forward in the field of AF screening and diagnosis.

Features importance

Through the analysis of feature importance in model 3, that includes features encoding prior known physiological information, we confirm the importance of the PR interval length^{22,23} and importance of QRS width and the R-wave amplitude as predictors of AF. AF is associated with functional or structural changes in the atrial myocardium.²⁴ In the ECG, these changes are often recognized in the P-wave abnormalities. Therefore, abnormal P-wave duration, P-axis and beat-to-beat P-wave variability have been described as AF predictors.²⁵ The association of the PR interval with AF is controversial in the literature. PR interval components (P-wave onset to P-wave peak duration, P-wave peak to P-wave end duration and PR segment) have different impacts on AF prediction. A prolongation of the PR-interval will be predictive of AF if the prolongation is mainly due to the P-wave duration. This also explains the observation that a short PR-segment is more predictive of AF, as in this case, the P-wave duration would contribute more to the overall PR interval.²⁶ Although AF is part of a

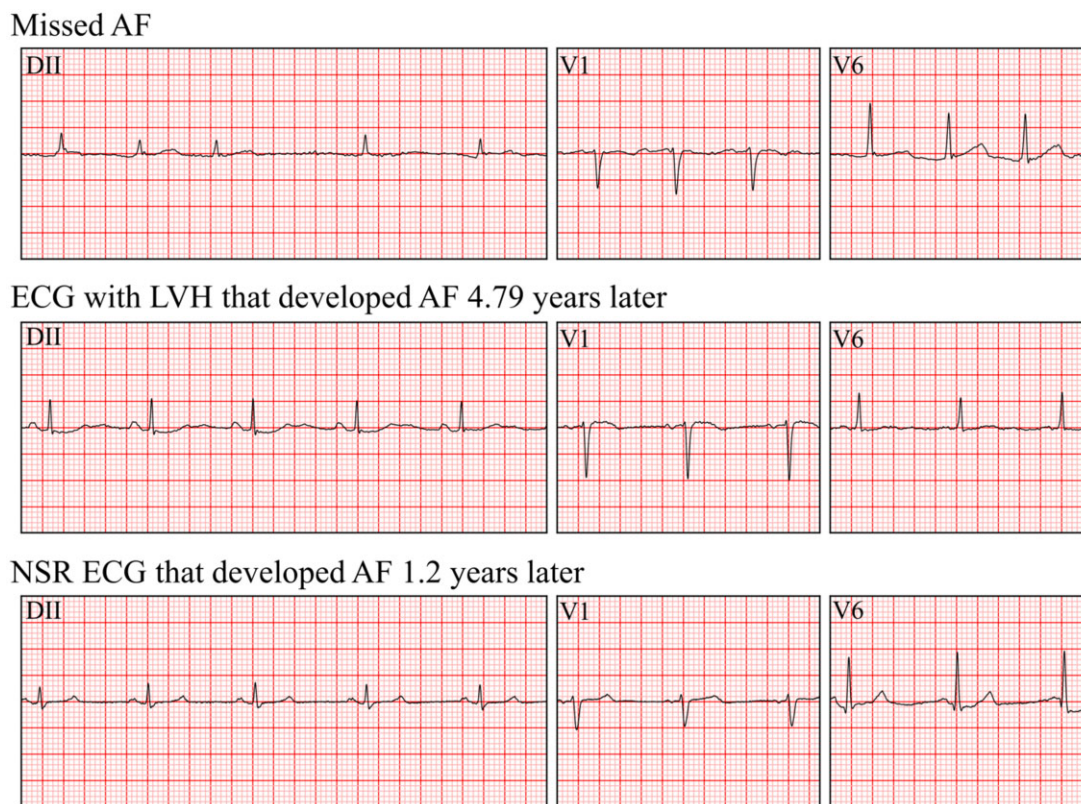


Figure 6 Example for three recordings from lead DII, V1, V6. All were re-annotated by cardiologist with 5 years of experience.

syndrome called atrial cardiomyopathy, some ventricular ECG abnormalities have been reported to be indirect predictors of atrial disease, QRS duration is a marker of structural modifications and ventricular remodelling, therefore, can be associated with left atrial size, supporting the concept of simultaneous processes in the atria and ventricles. Prolonged QRS duration reflected as fragmented QRS²⁷ and left ventricular hypertrophy¹⁸ have been associated with AF. Sokolow-Lyon voltage product ($[SV1 + RV5 / RV6] * QRS \text{ duration} \geq 371\,000 \mu\text{Vms}$) was related to incident AF, showing that both R-wave and QRS duration have an impact on AF prediction. When combining all features with model 6, age was the most important feature which is consistent with the knowledge that the likelihood of having AF significantly increases as a function of age.²⁸

Limitations

The main limitation of our study relates to the lack of evaluation of our model on a longitudinal cohort that was consistently and regularly followed-up for years, i.e. with a baseline examination and then followed-up regular visits for all the cohort. Also, it will be important to evaluate the models performance on external datasets in order to assess their generalization performances. Although it may not be possible to obtain the exact same metadata for external datasets we may be able to benchmark the performances of models 1–4 which only use the raw ECG signal as input. The second main limitation of our study is the lack of a better mapping with the patient's EMR. Indeed, only a limited number of EMR information were available as metadata

and these were self-reported by patients. A better access to EMR variables would also enable benchmarking our model against state-of-the-art AF risk scoring systems such as CHARGE-AF.²⁹ Our experimental setting including patients with repeated 12-lead ECG recordings and excluding those who did not, has an intrinsic bias towards individuals already having a cardiac abnormality or at risk for a cardiovascular disease. The clinical endpoints used in this study was documented AF development within 5 years. With documented corresponding to a future 12-lead ECG examination with a positive AF diagnosis. This label is not necessarily a gold standard as there exists a number of possible bias in that registration such as individuals that would develop silent AF after their latest documented visit but would not perform an additional 12-lead ECG examination. Finally, the model needs to be further validated on individuals presenting no cardiac abnormality at the time of the baseline examination but who later on developed AF.

Conclusion

It is the first study integrating feature engineering, deep learning and EMR metadata to create a risk prediction tool for the management of patients at risk of AF. This new hybrid model may also be extended for the risk prediction of other cardiac pathologies. The high performance obtained suggest that structural changes in the 12-lead ECG are associated with existing or impending AF. Our study has

important clinical implications for AF management. The system has the potential to be rapidly incorporated in the clinical practice, helping to detect those prone to develop AF and better manage these at-risk patients to prevent complications. For that purpose further validation of the model on the external test set longitudinal cohort is needed.

Supplementary material

Supplementary material is available at *European Heart Journal – Digital Health* online.

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Conflict of interest: none declared.

Data availability

Researchers affiliated to educational or research institutions might make requests to access the CODE dataset. Requests should be made to the corresponding author of this paper. They will be forwarded and considered on an individual basis by the Telehealth Network of Minas Gerais. If approved, any data use will be restricted to non-commercial research purposes. The data will only be made available on the execution of appropriate data use agreements.

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