Importance of genotype for risk stratification in arrhythmogenic right ventricular cardiomyopathy using the 2019 ARVC risk calculator

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Received 1 October 2021; revised 6 March 2022; accepted 25 April 2022

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Abstract

Aims
To study the impact of genotype on the performance of the 2019 risk model for arrhythmogenic right ventricular cardiomyopathy (ARVC).

Methods and results
The study cohort comprised 554 patients with a definite diagnosis of ARVC and no history of sustained ventricular arrhythmia (VA). During a median follow-up of 6.0 (3.1,12.5) years, 100 patients (18%) experienced the primary VA outcome (sustained ventricular tachycardia, appropriate implantable cardioverter deﬁbrillator intervention, aborted sudden cardiac arrest, or sudden cardiac death) corresponding to an annual event rate of 2.6% [95% conﬁdence interval (CI) 1.9–3.3]. Risk estimates for VA using the 2019 ARVC risk model showed reasonable discriminative ability but with overestimation of risk. The ARVC risk model was compared in four gene groups: PKP2 (n = 118, 21%); desmoplakin (DSP) (n = 79, 14%); other desmosomal (n = 59, 11%); and gene elusive (n = 160, 29%). Discrimination and calibration were highest for PKP2 and lowest for the gene-elusive group. Univariable analyses revealed the variable performance of individual clinical risk markers in the different gene groups, e.g. right ventricular dimensions and systolic function are signiﬁcant risk markers in PKP2 but not in DSP patients and the opposite is true for left ventricular systolic function.

Conclusion
The 2019 ARVC risk model performs reasonably well in gene-positive ARVC (particularly for PKP2) but is more limited in gene-elusive patients. Genotype should be included in future risk models for ARVC.

Structured Graphical Abstract

Key Question
What is the performance of the 2019 ARVC risk score in an external multicentre cohort?
What is the impact of the genotype on arrhythmic risk prediction in ARVC?

Key Finding
The 2019 ARVC risk score overestimates risk of sustained ventricular arrhythmias.
Better performance in gene positive (especially PKP2) versus gene elusive patients.
Variable performance of clinical risk markers between gene groups.

Take Home Message
The 2019 ARVC risk score performs better in gene positive patients.
Predictors of arrhythmic events perform differently between the various gene groups.

ARVC, arrhythmogenic right ventricular cardiomyopathy; AUC, area under the curve; ECHO, echocardiography; CMR, cardiac magnetic resonance; HR, hazard ratio; LGE, late gadolinium enhancement; LV, left ventricular; LVEDD, left ventricular enddiastolic diameter; LVEF, left ventricular ejection fraction; PLAX, parasternal long-axis; PVC, premature ventricular complex; RV, right ventricular; RVEDV, right ventricular end-
Arrhythmogenic right ventricular cardiomyopathy (ARVC) is a heritable heart muscle disorder characterized by ventricular arrhythmia (VA) that can lead to sudden cardiac death (SCD). The clinical diagnosis of ARVC is based on consensus criteria that have been developed to capture the right ventricular abnormalities typically manifested by patients. The genetic architecture of ARVC is diverse, with pathogenic variants first described in genes coding for desmosomal proteins and then, more recently, in genes encoding a number of non-desmosomal proteins. Although limited, current data suggest that clinical outcomes may differ between genotypes.

Implantable cardioverter defibrillator (ICD) therapy in patients with ARVC and documented sustained VA is associated with improved survival and is considered a Class 1 indication in current practice guidelines. However, the indications for primary prevention ICDs in individuals with ARVC that have no history of sustained VA are less certain and until recently were based on a subjective evaluation of clinical risk markers supported by evidence of variable quality. In a recent landmark study, a risk tool designed to provide individualized risk estimates was proposed but the model was agnostic to the underlying genetic cause of the disease.

In this study, we hypothesized that disease aetiology influences the risk of VA in ARVC and sought to determine the performance of the 2019 ARVC risk model in a large multicentre cohort of patients stratified by genotype. During the review of this paper, a correction to the 2019 ARVC risk model was published. Our analysis is based on the corrected risk score but data on the original model are presented in the Supplementary material online to illustrate the impact of the correction on its performance.

Methods

Study design and participating centres

This is an international, multicentre retrospective observational cohort study using data on consecutively evaluated patients with ARVC recruited from 17 centres in 7 countries (see Supplementary material online, Table S1). All participating centres specialize in the clinical management of cardiomyopathy patients. The study conforms to the Declaration of Helsinki and all centres have local ethical approval.

Study population

Patients were enrolled according to pre-specified inclusion criteria. Specifically: (i) a definite diagnosis of ARVC according to the 2010 task-force criteria (TFC); (ii) no history of sustained VA before first assessment at the participating centre; (iii) a follow-up period of at least 1 month; and (iv) age of diagnosis of 14 years or more.

Data collection and study variables

Study data were collected independently by each centre and managed using REDCap (Research Electronic Data Capture) electronic data capture tools hosted at University College London. Standard data collection procedures, in accordance with general data protection regulation, were followed (see Supplementary material online, Table S2). Data were collected at each centre following review of medical and death records using variables derived from those used by Cadrin-Tourigny et al. The time of diagnosis was set as the baseline timepoint. The baseline phenotypic data comprised the primary dataset used for most analyses. As not all individuals had cardiac magnetic resonance (CMR) imaging at their baseline timepoint, we collected a second dataset of all the phenotypic data at the time of a first CMR if performed during follow-up (CMR dataset). The latter was used only for data imputation and sensitivity analysis purposes (see Supplementary material online).

Genetic analysis

Clinical genetic testing is performed routinely at all participating centres using next-generation sequencing or direct sequencing of candidate genes associated with ARVC. In all probands, genes that have been shown to have a moderate or strong association with ARVC were analysed: plakophilin-2 (PKP2), desmoplakin (DSP), plakoglobin (JUP), desmoglein-2 (DSG2), desmocollin-2 (DSC2), transmembrane protein 43 (TMEM43), desmin (DES), and phospholamban (PLN). Genetic variants were classified according to the American College of Medical Genetics and Genomics guidelines following independent review (P.S. and A.P.). Where additional evidence of pathogenicity (e.g., segregation data) for novel variants was available from the contributing centre, they were re-classified accordingly. A list of all the identified variants and their respective classification are reported in Supplementary material online, Table S3. Gene-elusive and gene-positive genetic status was defined according to the absence or presence of a pathogenic or likely pathogenic variant in any of the genes tested in each patient.

Outcomes

The primary outcome of this study was the first VA during follow-up and was a composite of: (i) spontaneous sustained ventricular tachycardia (VT), defined as VT lasting ≥30 s or with haemodynamic compromise at ≥100 b.p.m. or terminated by electrical cardioversion; (ii) ICD intervention, defined as ICD shock or anti-tachycardia override pacing delivered in response to a ventricular tachyarrhythmia confirmed by intracardiac ECG data; (iii) SCD, defined as death of cardiac origin that occurred unexpectedly within 1 h of the onset of new symptoms or a death that was unwitnessed and unexpected; (iv) aborted sudden cardiac arrest, defined as SCD, that is reversed by cardiopulmonary resuscitation and/or defibrillation or cardioversion. Death from any other cause and heart transplantation was also recorded.

Predictors

Potential predictors similar to those used by Cadrin-Tourigny et al. were studied. All predictor variables were determined at the baseline timepoint or within 1 year of baseline but always before the arrhythmic outcome. Recent syncope was defined as cardiac syncope during the last 6 months before baseline.

General statistical methods

All data manipulation and analyses were performed using the Python programming language (Version 3.8, Python Software Foundation, https://
www.python.org/). Continuous variables were tested for normality of distribution by visual inspection of histograms and statistical normality tests (Shapiro–Wilk). Normally distributed variables are expressed as mean ± SD and non-normally distributed variables as median (25th, 75th percentiles). Categorical variables are reported as counts and percentages, as appropriate. The TableOne and Scipy libraries were used for the construction of summary statistics tables and for all comparisons.14 The Seaborn and Matplotlib libraries were utilized for data visualization.15 The Zephyr library was used to calculate incidence rates.16

Follow-up time was calculated as the difference in age between the baseline (specific to each dataset) and the age when the study endpoint or censoring was reached. The annual event rate was calculated by dividing the number of patients reaching the endpoint by the total follow-up period for that endpoint. The cumulative probability for the occurrence of an outcome was estimated using the Aalen-Johansen estimate in order to take into account competing risks.17,18 Competing events were defined as the occurrence of heart transplantation or non-arrhythmic death. The Lifelines library was used for all time-to-event analyses.19 Fine-Gray regression was used to model the impact of clinical predictors on the arrhythmic outcome, in the context of competing risks.20 Fine-Gray regression was performed through the cmprsk library from the R-project through an R to Python interface.21 Hazard ratios and 95% confidence intervals (CIs) were reported. Bonferroni correction was used to correct P-values when multiple comparisons were made. P-values < 0.05 were considered significant.

**Model validation**

The corrected 5-year ARVC risk score was calculated according to the proposed formula10–11:

\[ P(VA) = 1 - S_0(t)^{exp(LP)} \]

where \( S_0(t) \) is the baseline survival probability at the time \( t \), which is 0.8396 at the 5-year mark (\( t = 5 \)). The linear predictor was calculated as 0.488 × sex – 0.022 × age + 0.657 × history of recent cardiac syncope + 0.811 × history of NSVT + 0.170 × ln(24 h PVC count) + 0.113 × sum of anterior and inferior leads with TWI – 0.025 × RVEF where NSVT is non-sustained ventricular tachycardia; PVC, premature ventricular complex; TWI, T-wave inversion; RVEF, right ventricular ejection fraction.

Binary parameters (male sex, history of recent cardiac syncope, and history of NSVT) were considered as 1 = positive and 0 = negative. The original 5-year ARVC risk score differs in regard to the \( S_0(t) \) parameter (see Supplementary material online).

Model validation has been developed according to standard practices, assessing both discrimination and calibration.22 The discriminatory performance of the model was assessed using the Uno concordance index as obtained by the sklearn.metrics.concordance_index_ipcw function.23 Due to the time dependency of the outcomes, we also opted to use a time-dependent receiver operating characteristic curve analyses for the 5-year ARVC risk score.24 The sklearn.metrics.roc_auc_score function was used to calculate the time-dependent area under the curve at 5 years. In order to assess the model’s calibration, calibration plots were constructed using the sklearn.calibration.calibration_curve and seaborn.regplot functions.25 Bins of equal number of patients were created. All the model-validation analyses were repeated for each gene group. 95% CIs were obtained using a bootstrap procedure with 10,000 iterations of random sampling with replacement.

**Missing data**

Missing data were addressed for the model-validation analyses. The Missingno library was utilized to visualize missing data (Supplementary material online, Methods and Figure S1).26 Missing data were assumed to be missing at random and were imputed using the multiple imputation with chained equations method.27 The Sklearn library (impute.IterativeImputer) was utilized to perform data imputation.28 A total of 10 imputation rounds were performed before returning the imputations computed during the final round. A round is a single imputation of each feature with missing values. Sensitivity analyses were performed (Supplementary material online, Methods).

**Results**

A total of 554 patients were enrolled from 17 centres. Demographic, genetic, clinical, outcome characteristics of patients and missing data in either dataset are reported in Table 1.

**Patient outcomes**

During a median follow-up time of 6.0 (3.1, 12.5) years, 100 patients (18%) experienced the primary arrhythmic outcome (Figure 1) with a corresponding annual event rate of 2.6% (95% CI 1.9–3.3). The primary arrhythmic outcome consisted of spontaneous sustained VT in 37 (6.7%), appropriate ICD intervention in 52 (9.4%), aborted SCD in 2 (0.4%), and SCD in 9 (1.6%). At last follow-up, 17 (3.1%) patients had undergone heart transplantation and 43 (7.8%) had died. Causes of death were SCD in 16 (37%), heart failure in 11 (26%), non-cardiac in 11 (26%), and unknown in 5 (11%).

**Arrhythmogenic right ventricular cardiomyopathy risk score validation**

The corrected ARVC risk score was calculated in all 554 individuals. The median calculated corrected 5-year risk score was 17.2% (9.5–34.3%) (see Supplementary material online, Figure S3). Overlapping cumulative incidence of VA in the estimated-risk strata is shown in Figure 2. When fitting a multivariable Fine-Gray regression model with the same predictors as the 2019 ARVC risk model, sex \( (P = 0.021) \), recent syncope \( (P = 0.001) \), number of TWIs \( (P = 0.001) \), and log value for PVC count \( (P = 0.004) \) were found to be significant predictors, whereas age at baseline \( (P = 0.15) \), NSVT \( (P = 0.16) \), and RVEF from CMR \( (P = 0.16) \) were not significant predictors of VA as shown in Figure 2. Non-sustained ventricular tachycardia was identified more commonly in patients with ICD than those without (143/234, 61% vs. 120/320, 38%; \( P < 0.001 \)). Uno’s concordance index was 0.75 (95% CI 0.70–0.81). Calibration curve revealed a slope of 0.52 (95% CI 0.37–0.71) and an intercept of –0.01 (95% CI –0.05 to 0.02) suggestive of risk overestimation (Figure 2). As expected, the corrected version of the 2019 ARVC risk model resulted in lower risk estimates and less risk overestimation as compared with the original 2019 ARVC risk model (see Supplementary material online, Figure S6).

**Impact of genotype and sex on risk stratification**

Genetic analysis was available in 447 (80.7%) of the 554 patients enrolled in the study; a pathogenic or likely pathogenic variant was identified in 290 patients. For the purposes of this analysis, four major gene groups were studied: PKP2 (\( n = 118, 21.3% \), DSP (\( n = 79, 14\% \), other desmosomal gene (\( n = 59, 11\% \), and gene-elusive patients (\( n = 160, 28.8\% \)). We did not create a subgroup for non-desmosomal gene carriers due to the limited numbers of patients.
| Baseline          | NA (n = 554) | Overall (n = 554) | No VA (n = 454) | VA (n = 100) | P-value |
|-------------------|--------------|-------------------|-----------------|--------------|---------|
| Age (years)       | 0            | 41.0 (27.2, 53.1) | 42.8 (29.0, 54.0) | 37.0 (21.0, 49.0) | 0.002   |
| Male sex          | 0            | 302 (54.5)        | 231 (50.9)      | 71 (71.0)    | <0.001  |
| Genotype          | 0            |                   |                 |              | 0.136   |
| Gene elusive      |              | 157 (28.3)        | 138 (30.4)      | 19 (19.0)    |         |
| DES               | 5 (0.9)      | 4 (0.9)           | 1 (1.0)         |              |         |
| DSC2              | 11 (2.0)     | 9 (2.0)           | 2 (2.0)         |              |         |
| DSG2              | 27 (4.9)     | 24 (5.3)          | 3 (3.0)         |              |         |
| DSP               | 79 (14.3)    | 68 (15.0)         | 11 (11.0)       |              |         |
| FLNC              | 10 (1.8)     | 10 (2.2)          |                |              |         |
| JUP               | 21 (3.8)     | 16 (3.5)          | 5 (5.0)         |              |         |
| Multiple          | 13 (2.3)     | 8 (1.8)           | 5 (5.0)         |              |         |
| Not performed     | 104 (18.8)   | 79 (17.4)         | 25 (25.0)       |              |         |
| PKP2              | 118 (21.3)   | 91 (20.0)         | 27 (27.0)       |              |         |
| PLN               | 3 (0.5)      | 2 (0.4)           | 1 (1.0)         |              |         |
| RBM20             | 1 (0.2)      | 1 (0.2)           |                |              |         |
| TMEM43            | 2 (0.4)      | 1 (0.2)           | 1 (1.0)         |              |         |
| Ethnicity         | 1            |                   |                 |              | 0.476   |
| Caucasian         | 518 (93.7)   | 427 (94.3)        | 91 (91.0)       |              |         |
| African           | 15 (2.7)     | 11 (2.4)          | 4 (4.0)         |              |         |
| Asian             | 20 (3.6)     | 15 (3.3)          | 5 (5.0)         |              |         |
| Proband status    | 0            | 304 (54.9)        | 234 (51.5)      | 70 (70.0)    | 0.001   |
| Any symptoms      | 0            | 265 (47.8)        | 202 (44.5)      | 63 (63.0)    | 0.001   |
| Recent syncope    | 0            | 32 (5.8)          | 17 (3.7)        | 15 (15.0)    | <0.001  |
| Structural TFC—minor | 0             | 72 (13.0)        | 62 (13.7)      | 10 (10.0)    | 0.002   |
| Structural TFC—major | 290 (52.3)   | 222 (48.9)        | 68 (68.0)       |              |         |
| Tissue TFC—minor  | 0            | 11 (2.0)          | 10 (2.2)        | 1 (1.0)      | 0.693   |
| Tissue TFC—major  | 29 (5.2)     | 23 (5.1)          | 6 (6.0)         |              |         |
| Repolarization TFC—minor | 0         | 91 (16.4)        | 78 (17.2)      | 13 (13.0)    | 0.017   |
| Repolarization TFC—major | 261 (47.1)   | 201 (44.3)        | 60 (60.0)       |              |         |
| Depolarization TFC—minor | 0         | 198 (35.7)       | 169 (37.2)     | 29 (29.0)    | 0.132   |
| Depolarization TFC—major | 58 (10.5)   | 43 (9.5)          | 15 (15.0)       |              |         |
| Arrhythmia TFC—minor | 0           | 322 (58.1)       | 266 (58.6)     | 56 (56.0)    | <0.001  |
| Arrhythmia TFC—major | 96 (17.3) | 60 (13.2)         | 36 (36.0)       |              |         |
| Family history TFC—minor | 0         | 41 (7.4)          | 36 (7.9)        | 5 (5.0)      | 0.024   |
| Family history TFC—major | 367 (66.2) | 309 (68.1)        | 58 (58.0)       |              |         |
| RBBB              | 35           |                   |                 |              | 0.206   |
| None              | 433 (83.4)   | 355 (83.3)        | 78 (83.9)       |              |         |

Continued
| NA Overall (n = 554) | No VA (n = 454) | VA (n = 100) | P-value |
|---------------------|----------------|-------------|---------|
| iRBBB               | 41 (7.9)       | 37 (8.7)    | 4 (4.3) |
| cRBBB               | 45 (8.7)       | 34 (8.0)    | 11 (11.8)|
| LBBB                | 42             |             |         |
| None                | 477 (93.2)     | 393 (93.3)  | 84 (92.3)|
| LAFB                | 13 (2.5)       | 13 (3.1)    |         |
| LPFB                | 2 (0.4)        | 1 (0.2)     | 1 (1.1) |
| LBBB                | 12 (2.3)       | 7 (1.7)     | 5 (5.5) |
| IVCD                | 5 (1.0)        | 5 (1.2)     |         |
| QRS duration (ms)   | 148            | 100.0 (90.0, 112.0) | 100.0 (90.0, 110.0) | 100.0 (90.0, 120.0) | 0.306 |
| TAD (ms)            | 217            | 52.0 (40.0, 60.0) | 52.0 (40.0, 60.0) | 55.0 (40.0, 60.0) | 0.787 |
| Epsilon waves       | 0              | 51 (11.3)   | 34 (9.3) | 17 (19.5) |
| Number of TWIs      | 0              | 3.0 (1.0, 4.0) | 3.0 (1.0, 4.0) | 4.0 (2.0, 5.0) | <0.001 |
| Limb-lead voltages (mV) | 213     | 0.9 (0.6, 12) | 0.9 (0.6, 12) | 0.8 (0.5, 12) | 0.486 |
| Preocordial-lead voltages (mV) | 219 | 1.5 (1.0, 2.0) | 1.5 (1.0, 2.0) | 1.4 (1.0, 2.1) | 0.616 |
| PVCs (per 24 h)     | 77             | 1232.0 (382,3044) | 1126.5 (273,3022) | 1695.0 (983,3315) | 0.004 |
| Non-sustained VT    | 0              | 234 (42.2)  | 176 (38.8) | 58 (58.0) | 0.001 |
| RV dilatation (ECHO) | 41             |             |         | <0.001 |
| None                | 212 (41.3)     | 192 (45.9)  | 20 (21.1) |
| Mild                | 134 (26.1)     | 108 (25.8)  | 26 (27.4) |
| Moderate            | 103 (20.1)     | 79 (18.9)   | 24 (25.3) |
| Severe              | 64 (12.5)      | 39 (9.3)    | 25 (26.3) |
| RVOT-PLAX diameter (ECHO) | 249     | 33.0 (29.0, 37.0) | 32.0 (28.0, 36.8) | 36.0 (32.0, 39.0) | 0.001 |
| LVEDD (ECHO)        | 176            | 50.0 (47.0, 54.8) | 50.0 (47.0, 54.2) | 50.0 (46.0, 54.8) | 0.664 |
| LVEF (ECHO)         | 23             | 57.5 (50.0, 65.0) | 57.0 (50.0, 65.0) | 58.5 (51.0, 65.0) | 0.159 |
| RVEDV (CMR)         | 281            | 181 (140,219) | 178 (137,212) | 205 (168,239) | 0.008 |
| RVEF (CMR)          | 263            | 46.0 (39.5,55.0) | 48.0 (41.0,56.0) | 39.0 (35.0,45.0) | <0.001 |
| RV LGE              | 247            | 113 (36.8)  | 87 (33.5)  | 26 (55.3) | 0.007 |
| LVEDV (CMR)         | 265            | 154 (126,186) | 156 (128,186) | 145 (107,170) | 0.1 |
| LVEF (CMR)          | 253            | 57.0 (48.0, 63.0) | 56.5 (47.0,63.0) | 58.0 (52.0,62.5) | 0.308 |
| LV LGE              | 247            | 176 (57.3)  | 150 (57.9) | 26 (54.2) | 0.746 |
| Anti-arrhythmic at baseline | 1             |             |         | <0.001 |
| None                | 425 (76.9)     | 365 (80.4)  | 60 (60.6) |
| Amiodarone          | 35 (6.3)       | 25 (5.5)    | 10 (10.1) |
| Sotalol             | 64 (11.6)      | 46 (10.1)   | 18 (18.2) |
| Class IC            | 10 (1.8)       | 5 (1.1)     | 5 (5.1)   |
| Mexiletine          | 2 (0.4)        | 2 (2.0)     |         |
| Beta-blocker (excluding sotalol) at baseline, n (%) | 1             | 213 (38.5)  | 175 (38.5) | 38 (38.4) | 0.933 |

Continued
The clinical phenotypes of each genotype group are summarized in Figure 3 and Table 2. Desmoplakin patients had a higher median RVEF on CMR [51 (45,57) vs. 46 (38,55), \( P = 0.003 \)] and lower number of T-wave inversions in the inferolateral leads [1.0 (0.0–3.0) vs. 3.0 (1.5,5.0), \( P < 0.001 \)] than the rest of the cohort. Patients in the ‘other desmosomal gene’ group had the lowest frequency of non-sustained VT (17/59 vs. 217/495, \( P = 0.027 \)). Cumulative incidence for VA per gene group is shown in Figure 4. Overall, there was no significant difference (Gray’s test, \( P = 0.37 \)), but in pairwise comparisons, the gene-elusive group had significantly
Figure 2 Arrhythmogenic right ventricular cardiomyopathy risk model validation. Cumulative incidence curve for ventricular arrhythmia per strata of risk according to the corrected 2019 ARVC risk model 10,11 (top left). Forest plot of the impact of each parameter included in the model in a multivariable Fine-Gray regression analysis (top right). Time-dependent area under the curve at 5 years for prediction of the primary arrhythmic outcome (bottom left). Calibration plot showing the agreement between predicted and observed probabilities for the primary outcome (bottom right). Solid line represents results and dotted line represents reference line (bottom left) and perfect calibration (bottom right).

Figure 3 Violin plots (top) and barplots (bottom) of the six key variables for the arrhythmogenic right ventricular cardiomyopathy risk model grouped by genotype group and sex. 95% confidence intervals are shown in barplots.
Table 2  Demographic and clinical characteristics of patients in the main dataset according to the genotype subgroup

|                      | NA          | Overall 416 | PKP2 118 | DSP 79 | Gene elusive 160 | Other desmosomal gene 59 | P-value |
|----------------------|-------------|-------------|----------|--------|------------------|--------------------------|---------|
| Age (years)          | 0           | 42.0 (28.0,53.9) | 44.6 (29.8,51.9) | 41.0 (28.0,50.6) | 44.7 (30.0,55.0) | 40.0 (18.6,54.4) | 0.364   |
| Male sex             | 0           | 223 (53.6)  | 57 (48.3) | 31 (39.2) | 99 (61.9) | 36 (61.0) | 0.004   |
| Caucasian ethnicity  | 1           | 391 (94.2)  | 110 (93.2) | 74 (94.9) | 149 (93.1) | 58 (98.3) | 0.048   |
| Proband status       | 0           | 193 (46.4)  | 69 (58.5) | 48 (60.8) | 47 (29.4) | 29 (49.2) | <0.001  |
| Symptoms             | 0           | 202 (48.6)  | 46 (39.0) | 40 (50.6) | 84 (52.5) | 32 (54.2) | 0.101   |
| Recent syncope (<6 months) | 0     | 27 (6.5)    | 11 (9.3)  | 3 (3.8)  | 7 (4.4)  | 6 (10.2) | 0.172   |
| Structural TFC, major | 0           | 201 (48.3)  | 52 (44.1) | 16 (20.3) | 99 (61.9) | 34 (57.6) | <0.001  |
| Structural TFC, minor | 55 (13.2)  | 20 (16.9)   | 8 (10.1)  | 18 (11.2) | 9 (15.3)  |             |         |
| Tissue TFC, major    | 0           | 20 (4.8)    | 4 (3.4)   | 1 (1.3)  | 10 (6.2)  | 5 (8.5)   | 0.507   |
| Tissue TFC, minor    | 6 (1.4)     | 2 (1.7)     | 1 (1.3)   | 2 (1.2)  | 1 (1.7)   |             |         |
| Repolarization TFC, major | 0     | 199 (47.8)  | 78 (66.1) | 13 (16.5) | 76 (47.5) | 32 (54.2) | <0.001  |
| Repolarization TFC, minor | 71 (17.1) | 12 (10.2)   | 23 (29.1) | 25 (15.6) | 11 (18.6) |             |         |
| Depolarization TFC, major | 0      | 41 (9.9)    | 12 (10.2) | 1 (1.3)  | 15 (9.4)  | 13 (22.0) | 0.002   |
| Depolarization TFC, minor | 163 (39.2)| 42 (35.6)   | 39 (49.4) | 58 (36.2) | 24 (40.7) |             |         |
| Arrhythmia TFC, major | 0           | 49 (11.8)   | 14 (11.9) | 8 (10.1)  | 21 (13.1) | 6 (10.2)  | 0.062   |
| Arrhythmia TFC, minor | 265 (63.7)  | 68 (57.6)   | 62 (78.5) | 96 (60.0) | 39 (66.1) |             |         |
| Family history TFC, major | 0     | 297 (71.4)  | 106 (89.8) | 71 (89.9) | 72 (45.0) | 48 (81.4) | <0.001  |
| Family history TFC, minor | 24 (5.8)  | 1 (0.8)     | 2 (2.5)   | 16 (10.0) | 5 (8.5)   |             |         |
| cRBBB                | 20          | 33 (8.3)    | 5 (4.5)   | 1 (1.3)  | 18 (11.8) | 9 (16.4)  | 0.013   |
| LBBB                 | 20          | 10 (2.6)    | 3 (2.7)   | 6 (4.1)  | 1 (1.9)   |             | 0.024   |
| QRS duration         | 89          | 100.0 (90.0,110.0) | 97.5 (90.0,105.2) | 100.0 (90.0,107.2) | 104.0 (90.0,116.2) | 108.0 (97.5,121.0) | 0.004   |
| Terminal activation duration | 140     | 55.0 (40.0,60.0) | 52.0 (40.0,58.0) | 54.0 (40.0,60.0) | 55.0 (40.0,60.0) | 56.0 (40.0,64.0) | 0.822   |
| Epsilon waves        | 68          | 33 (9.5)    | 8 (8.0)   | 3 (4.5)  | 11 (8.7)  | 11 (20.0) | 0.025   |
| Number of TWIs       | 0           | 3.0 (1.0,4.0) | 4.0 (2.0,5.0) | 1.0 (0.0,3.0) | 3.0 (1.0,5.0) | 4.0 (2.0,5.0) | <0.001  |
| Limb-lead voltages (mV) | 140         | 0.9 (0.6,1.0) | 1.0 (0.7,1.1) | 0.7 (0.4,0.9) | 0.9 (0.7,1.2) | 0.8 (0.6,1.2) | <0.001  |
| Precordial-lead voltages (mV) | 146     | 1.5 (1.0,2.0) | 1.5 (1.0,2.0) | 1.5 (1.1,1.6) | 1.4 (1.1,1.9) | 1.6 (1.1,2.2) | 0.491   |
| PVCs (per 24h)       | 60          | 1243.5 (420.2,3121.8) | 1174.0 (182.8,2425.8) | 1604.0 (754.0,3121.0) | 1188.0 (196.0,3748.0) | 1100.0 (566.0,2500.0) | 0.111   |

Continued
Table 2  Continued

| NA Overall | PKP2 | DSP | Gene elusive | Other desmosomal gene | P-value |
|------------|------|-----|--------------|-----------------------|---------|
| Non-sustained VT | 0 169 (40.6) | 45 (38.1) | 39 (49.4) | 68 (42.5) | 17 (28.8) | 0.092 |
| RV dilatation (ECHO)—none | 29 174 (45.0) | 55 (51.4) | 51 (68.0) | 55 (36.9) | 13 (23.2) | <0.001 |
| RV dilatation (ECHO)—mild | 92 (23.8) | 15 (14.0) | 17 (22.7) | 43 (28.9) | 17 (30.4) | |
| RV dilatation (ECHO)—moderate | 75 (19.4) | 26 (24.3) | 6 (8.0) | 30 (20.1) | 13 (23.2) | |
| RV dilatation (ECHO)—severe | 46 (11.9) | 11 (10.3) | 1 (1.3) | 21 (14.1) | 13 (23.2) | |
| RVOT-PLAX diameter (ECHO) | 156 32.0 (28.8,37.0) | 32.0 (28.0,37.0) | 30.0 (27.0,32.8) | 33.0 (29.1,37.8) | 34.5 (31.2,37.8) | 0.001 |
| LVEDD (ECHO) | 114 50.0 (47.0,55.0) | 48.0 (45.0,52.5) | 53.0 (48.0,57.0) | 51.0 (47.0,55.0) | 48.0 (47.0,51.0) | <0.001 |
| LVEF (ECHO) | 19 58.0 (50.0,65.0) | 62.0 (55.8,66.2) | 52.0 (42.8,58.6) | 56.5 (50.6,64.2) | 60.0 (52.6,65.0) | <0.001 |
| ICD implantation | 0 209 (50.2) | 59 (50.0) | 54 (68.4) | 68 (42.5) | 28 (47.5) | 0.002 |
| Follow-up duration (years) | 0 6.0 (3.0,11.0) | 5.9 (2.9,10.9) | 5.0 (2.8,9.9) | 6.0 (3.4,12.9) | 7.0 (3.0,12.5) | 0.485 |
| VA | 67 (16.1) | 27 (22.9) | 11 (13.9) | 19 (11.9) | 10 (16.9) | |
| Heart transplantation | 0 7 (1.7) | 2 (2.5) | 5 (3.1) | | | 0.146 |
| Mortality | 0 21 (5.0) | 5 (4.2) | 4 (5.1) | 2 (1.2) | 10 (16.9) | <0.001 |
| VT ablation | 0 17 (4.1) | 3 (2.5) | 2 (2.5) | 7 (4.4) | 5 (8.5) | 0.247 |
| n (CMR dataset) | 288 | 116 | 80 | 62 | 30 | |
| RV EDV (CMR) | 40 182.0 (144.8,219.2) | 182.0 (147.8,213.2) | 170.0 (124.2,184.0) | 185.5 (145.8,229.8) | 197.0 (165.0,231.8) | 0.004 |
| RV EF (CMR) | 26 48.0 (40.0,56.0) | 49.5 (39.5,58.0) | 51.0 (45.0,57.0) | 46.0 (39.0,54.0) | 47.0 (38.2,56.0) | 0.066 |
| LV EDV (CMR) | 8 98 (35.0) | 26 (33.8) | 18 (29.0) | 39 (33.9) | 15 (57.7) | 0.073 |
| LV EF (CMR) | 23 156.0 (129.0,184.0) | 145.0 (128.5,168.0) | 180.0 (139.5,202.5) | 156.0 (132.0,190.0) | 149.0 (124.5,169.5) | 0.008 |
| LV EDV (CMR) | 12 58.0 (49.8,63.2) | 63.0 (58.0,67.0) | 53.0 (43.0,60.0) | 55.0 (49.0,61.8) | 60.0 (53.0,66.0) | <0.001 |
| LV EF (CMR) | 8 153 (54.6) | 23 (29.9) | 60 (96.8) | 55 (47.8) | 15 (57.7) | <0.001 |

Categorical variables are reported as n (%). Continuous variables are reported as mean ± SD or median (25th, 75th percentiles) according to the distribution normality. CMR, cardiac magnetic resonance; cRBBB, complete right bundle branch block; ECHO, echocardiography; ICD, intracardiac cardioverter defibrillator; LBBB, left bundle branch block; LGE, late gadolinium enhancement; LV, left ventricle; LVEDD, left ventricular end-diastolic diameter; LVEDV, left ventricular end-diastolic volume; LVEF, left ventricular ejection fraction; NA, not available; PLAX, parasternal long-axis; PVC, premature ventricular complex; RBBB, right bundle branch block; RV, right ventricle; RVEDD, right ventricular end-diastolic diameter; RVEDV, right ventricular end-diastolic volume; RVET, right ventricular ejection fraction; TAD, terminal activation duration; TFC, task-force criteria; TWI, T-wave inversion; VA, ventricular arrhythmia; VT, ventricular tachycardia.
higher cumulative VA incidence compared with the PKP2 group (Gray’s test, \( P = 0.02 \)). Annual VA rate was 3.6\% (95\% CI 2.2–5.0) in PKP2 patients, 2.3\% (95\% CI 0.9–3.6) in DSP patients, 2.1\% (95\% CI 0.8–3.5) in patients with variants in either DSG2, DSC2, or JUP and 1.5\% (95\% CI 0.8–2.1) in gene-elusive patients. Cumulative incidence of VA analysis of the cohort stratified by individual genes did reveal an overall difference (see Supplementary material online, Figure S4, Gray’s test, \( P = 0.03 \)). Male sex was associated with a higher cumulative incidence of VA (Figure 4, Gray’s test, \( P = 0.001 \)). Validation of the corrected 2019 ARVC risk score in the various gene groups revealed significant differences in the performance of the model. As shown in Figure 5, the best fit was demonstrated within the gene-positive group (Uno’s concordance index 0.82, 95\% CI 0.76–0.88; calibration curve slope 0.78, 95\% CI 0.53–1.06; calibration curve intercept \(-0.05, 95\% CI \(-0.10 \) to \(-0.01 \)) and the worst fit was found in the gene-elusive group (Uno’s concordance index 0.65, 95\% CI 0.57–0.74; calibration curve slope 0.27, 95\% CI 0.06–0.55; calibration curve intercept 0.04, 95\% CI –0.02 to –0.09). The best performance was seen for PKP2 (Uno’s concordance index 0.83, 95\% CI 0.75–0.91; calibration curve slope 0.73, 95\% CI 0.44–1.14; calibration curve intercept \(-0.03, 95\% CI \(-0.10 \) to 0.02\)) followed by DSP (Uno’s concordance index 0.80, 95\% CI 0.53–0.96; calibration curve slope 0.69, 95\% CI 0.00–1.37; calibration curve intercept \(-0.06, 95\% CI \(-0.14 \) to 0.05\)) and other desmosomal variants (Uno’s concordance index 0.73, 95\% CI 0.48–1.00; calibration curve slope 0.28, 95\% CI \(-0.06 \) to 0.77; calibration curve intercept 0.00, 95\% CI –0.09 to 0.08). Sensitivity analyses showed that these findings were recapitulated in the complete-case and CMR datasets but did not reach statistical significance due to smaller sample size (see Supplementary material online, Table S4).

The variable performance of the corrected 2019 ARVC risk model in different genotypes led us to hypothesize that there may be differences in the performance of individual parameters as predictors of risk among the different gene groups. Due to the limited number of events in each gene group we opted not to perform multivariable analyses, but instead conducted multiple univariate analyses of all the available clinical characteristics in each gene group. Significant differences were present among the gene groups (Figure 6). The PKP2 group had the most significant predictors and the DSP group the least among the clinical variables that were studied (Figure 6). Sex was mainly a PKP2 group-related risk predictor (Figure 6, see Supplementary material online). Ventricular parameters reached statistical significance almost only within the DSP group. Similarly, the utility of right ventricular parameters was limited in the non-PKP2 groups (Figure 6).

Discussion

In this multicentre cohort of patients with ARVC, we show that the corrected 2019 ARVC risk score\(^{10,11}\) has a reasonable discriminative ability for VA, but it suffers from risk overestimation for VA. The analysis by genotype shows that the corrected 2019 model performs best in patients with pathogenic or likely pathogenic gene variants, especially PKP2, but is more limited in patients with an elusive genetic status. Analyses of individual potential risk predictors among the gene groups revealed significant variability reflecting differences in clinical phenotypes (Structured Graphical Abstract).

Risk stratification in arrhythmogenic cardiomyopathy

Although the notion that all patients presenting with tolerated sustained VA have a high risk of SCD has been challenged, it remains standard practice to offer them ICDs.\(^{29}\) There has been less certainty about patient selection for primary prevention ICDs. A large number of clinical predictors have been suggested as risk markers in ARVC, but supporting evidence is inadequately validated and often based on small heterogeneous cohorts.\(^{30,31}\) The original 2019 ARVC risk model was the first systematic attempt to develop a validated clinical tool that provides individual risk estimates for sustained VA in patients with definite ARVC and no prior sustained VA.\(^{10}\) A correction of the baseline survival probability has recently been published.\(^{11}\) External validation of the corrected 2019 ARVC
Figure 5 Corrected arrhythmogenic right ventricular cardiomyopathy risk model validation per major genotype group. Calibration plot showing the agreement between predicted and observed probabilities for the primary outcome (left). Time-dependent area under the curve for prediction of the primary arrhythmic outcome (right). Solid line represents results and dotted line represents perfect calibration (left) and reference line (right).
Figure 6 Heatmap of univariate predictors in the full cohort and major gene groups displaying the hazard ratios and relevant statistical significance derived from univariate analyses.
risk score in our cohort revealed a generally good accuracy, but we observed a much lower annual event rate and a significant overestimation of risk compared with the original paper. Risk overestimation was even more pronounced in the original 2019 ARVC risk model. Other, smaller studies have reported a good performance of the model. Similarly, when all the risk predictors used in the 2019 ARVC risk score were fitted to a Fine-Gray regression model, only sex, syncope, the number of TWIs, and log PVC count remained independent risk factors, whereas NSVT, age at diagnosis, and RVEF on CMR were non-significant.

To provide close comparison, we sought to replicate the characteristics of the 2019 development cohort. Nevertheless, there may be some important biases in patient selection. For example, the participating centres in our study were predominantly cardiomyopathy units compared with the electrophysiology focused units that participated in the 2019 study, which could predispose to a more arrhythmia-prone population. This can potentially explain the lower annual incidence rate of VA and thus the risk overestimation that we have observed compared with the 2019 model development study. In addition, the genotype composition between the two studies differs; for example, the prevalence of PKP2 variants in our cohort is 21.7% compared with 48.9% in the 2019 study. The fact that PKP2 variant carriers exhibited a higher cumulative incidence of VA than gene-elusive patients might also affect the corrected 2019 ARVC risk model performance in our cohort.

Utilizing genotype information for precision arrhythmic risk stratification

While ARVC has been uniformly defined using the 2010 TFC diagnostic criteria, it is recognized that specific genotypes associate with different phenotypic features. For example, prognostic markers such as TWI and early age of disease onset are more commonly seen in patients with ARVC caused by desmosomal gene variants. Although some rare genotypes associated with particularly arrhythmic profiles exist, there has been limited evidence to link specific genes to arrhythmic outcome prediction. In a recent study by the Nordic ARVC Registry, PKP2 mutation carriers showed decreased arrhythmia-free survival. This is consistent with our findings suggesting a higher cumulative incidence for VA in PKP2 vs. gene-elusive patients (Figure 4).

In our cohort, we observed that gene-elusive patients have the lowest incidence of the primary outcome, whereas the incidence was similar between the major gene groups. Application of the corrected 2019 ARVC risk model in the specific gene groups revealed good performance within the gene-positive patients and particularly the PKP2 group with least overestimation of risk compared with the total cohort (mostly identified in the lower risk strata) in comparison with the gene-elusive group where the model had modest performance with significant risk overestimation across all risk strata. Model performance was intermediate for the DSP and other desmosomal groups but did not differ significantly with the other subgroups, possibly due to smaller patient numbers. The corrected 2019 ARVC risk model should be used with caution in non-PKP2 patients.

Due to restricted subgroup sizes, we were unable to perform multivariable analyses to study individual risk markers in different genotype clusters, but in a univariable analysis we did observe gene-specific differences in the association of some of the variables used by the 2019 ARVC risk model with VA. For example, significant risk was conferred by right ventricular parameters mainly in the PKP2 group. Interestingly, sex did not reach statistical significance in any of the non-PKP2 groups (Figure 6, see Supplementary material online, Figure S5). To our knowledge, this is the first demonstration of this phenomenon and as such generates hypotheses for future studies.

The main implication of our findings is that the incorporation of genotype is vital in future iterations of risk models in ARVC. Variables such as variant type and location and the presence of multiple variants have all been shown to affect phenotype and would likely demonstrate unique risk profiles. Ethnic differences might also add to this complexity. Similarly, forms of ARVC that affect predominantly the left ventricle may require bespoke risk prediction models.

Limitations

The limited size of the cohort did not allow for gene-specific analyses for all genes harbouring causative variants. Even in the groups with significant differences in model performance, a limited number of events was observed and the results should be interpreted cautiously. The lack of CMR at the baseline timepoint in a significant part of the cohort challenged the application of the corrected 2019 ARVC risk score, which was addressed using data imputation. Our study population is overwhelmingly Caucasian, and therefore extrapolation to other ethnic backgrounds should be done with caution. Information on ICD programming was not analysed but all centres are expert in ICD implantation and management and offered contemporary programming strategies to all patients. That said, considering the multicentre and retrospective nature of this study, patients are likely to have been exposed to varying ICD programming practices that may have influenced the frequency of the primary arrhythmic outcome. Non-sustained ventricular tachycardia has been defined as an ‘any exam diagnosis’ and therefore patients with an ICD have a higher likelihood of NSVT detection than those without.

Conclusion

The corrected 2019 ARVC risk score has a reasonable discriminative ability but suffers from risk overestimation. It performs best among gene-positive patients and especially in the PKP2 subgroup, but its utility is limited in gene-elusive patients. The predictive power of individual risk markers also varies by genotype. Future iterations of risk models in ARVC should incorporate genotype data.

Supplementary material

Supplementary material will be available at European Heart Journal online.

Acknowledgements

The authors are grateful to the patients and their families who have contributed to making this work possible. This study was supported by the Arrhythmogenic Cardiomyopathy Trust charity (London, UK) and the National Institute for Health Research University College London Hospitals Biomedical Research Centre. This study is
supported by Research Network on Cardiovascular Diseases (CIBERCV), ISCIII, Madrid, Spain.

Funding
This work has been supported by: a British Heart Foundation, clinical research training fellowship to A.P. (FS/18/82/34024); Regional Registry for Cardio-cerebro-vascular Pathology, Veneto Region, Venice, Italy; PRIN Ministry of Education, University and Research 2017ZWACS, Rome, Italy; and University Research Grants CPDA144300 and BIRD192170, Padua, Italy. Medical Research Council (MRC) UK Clinical Academic Research Partnership (CARP) to L.R.L. Instituto de Salud Carlos III Fondo Europeo de Desarrollo Regional «Una manera de hacer Europa» with research grants to E.Z. (PI18/01582, PI21/01282, La Fe Biobank PT17/0015/0043) and to J.R.G. (PI18/01231).

Conflict of interest: P.M.E. declares consultancies for Pfizer, Sarepta. G.S. declares consultancies at Novartis, Bayer, Astrazeneca, Boston Scientific, Vifor Pharma, Menarini, Akcea Therapeutics. The other authors declare that there is no conflict of interest.

References
1. Basso C, Corrado D, Marcus FI, Nava A, Thieme G. Arrhythmogenic right ventricular cardiomyopathy. Lancet 2009;373:1289–1300.
2. Marcus FI, McKenna WJ, Sherrill D, et al. Diagnosis of arrhythmogenic right ventricular cardiomyopathy/dysplasia: proposed modification of the Task Force Criteria. Eur Heart J 2010;31:806–817.
3. James CA, Jongbloed JDH, Hershberger RE, et al. International evidence based reappraisal of genes associated with arrhythmogenic right ventricular cardiomyopathy using the clinical genome resource framework. Circ Genom Precis Med 2021;14:e003273.
4. Bhonsale A, Groeneweg JA, James CA, et al. Impact of genotype on clinical course in arrhythmogenic right ventricular dysplasia/cardio-myopathy-associated mutation carriers. Eur Heart J 2015;36:847–855.
5. Rigato I, Bauce B, Rampazzo A, et al. Compound and digenic heterozygosity predicts life-time arrhythmic outcome and sudden cardiac death in desmosomal gene-related arrhythmogenic right ventricular cardiomyopathy. Circ Cardiovasc Genet 2013;6:533–542.
6. Protonotarios A, Anastasakis A, Panagiotakos DB, et al. Arrhythmic risk assessment in genotyped families with arrhythmogenic right ventricular cardiomyopathy. Europace 2016;18:610–616.
7. Corrado D, Wichter T, Link MS, et al. Treatment of arrhythmogenic right ventricular cardiomyopathy/dysplasia: an international task force consensus statement. Eur Heart J 2015;36:3227–3237.
8. Corrado D, Leonori L, Link MS, et al. Implantable cardioverter-defibrillator therapy for prevention of sudden death in patients with arrhythmogenic right ventricular cardio-myopathy/dysplasia. Circulation 2003;108:3084–3091.
9. Bosman LP, Samman A, James CA, et al. Predicting arrhythmic risk in arrhythmogenic right ventricular cardiomyopathy: a systematic review and meta-analysis. Heart Rhythm 2018;15:1097–1107.
10. Cadrin-Tourigny J, Bosman LP, Nezza A, et al. A new prediction model for ventricular arrhythmias in arrhythmogenic right ventricular cardiomyopathy. Eur Heart J 2019;40:1211–1213.
11. Corrigendum to: A new prediction model for ventricular arrhythmias in arrhythmogenic right ventricular cardiomyopathy. Eur Heart J 2022;e22181.
12. Harris PA, Taylor RP, Thielke R, et al. Research electronic data capture (REDCap)—a metadata-driven methodology and workflow process for providing translational research informatics support. J Biomed Inform 2009;42:377–381.
13. Richards S, Aziz N, Bale S, et al. Standards and guidelines for the interpretation of sequence variants: a joint consensus recommendation of the American College of medical genetics and genomics and the association for molecular pathology. Genet Med 2015;17:405–424.
14. Pollard TJ, Johnson AEW, Raffa JD, Mark RG. Tableone: an open source Python package for producing summary statistics for research papers. JAMA Open 2018;1:26–31.
15. Waskom M, Seaborn: statistical data visualization. J Open Source Softw 2021;6:3021.
16. Zivich P, zeplid, https://github.com/pzivich/zeplid
17. Aalen OO, Johansen S. An empirical transition matrix for non-homogeneous markov chains based on censored observations. Scand J Stat 1978;5:141–150.
18. Edwards JK, Hester LL, Gokhale M, Lesko CR. Methodologic issues when estimating risks in pharmacoepidemiology. Curr Epidemiol Rep 2016;3:285–296.
19. Davidson-Plon C. Lifelines: survival analysis in Python. J Open Source Softw 2019;4:1317.
20. Fine JP, Gray RJ. A proportional hazards model for the subdistribution of a competing risk. J Am Stat Assoc 1999;94:496–509.
21. Team RC. R: a language and environment for statistical computing. Vienna, Austria: R Foundation for Statistical Computing; 2021.
22. Ramspeck CL, Karger JK, Dekker FW, Zoccali C, van Diepen M. External validation of prognostic models: what, why, how, when and where? Clin Kidney J 2021;14:49–58.
23. Uno H, Cai T, Pencina MJ, D’Agostino RB, Wei LJ. On the C-statistic for evaluating overall adequacy of risk prediction procedures with censored survival data. Stat Med 2011;30:1105–1117.
24. Kamarudin AN, Cox T, Kolamunnage-Dona R. Time-dependent ROC curve analysis in medical research: current methods and applications. BMC Med Res Methodol 2017;17:53.
25. Waskom ML. Seaborn: statistical data visualization. J Open Source Softw 2021;6:3021.
26. blogs. A Missingno: a missing data visualization suite. J Open Source Softw 2016;3:547.
27. White IR, Royston P, Wood AM. Multiple imputation using chained equations: issues and guidance for practice. Stat Med 2011;30:377–399.
28. Buinick L, Louppe G, Blondel M, et al. API design for machine learning software: experiences from the scikit-learn project. arXiv [cs.LG] 2013. https://doi.org/10.48550/arXiv.1309.0238.
29. Cadrin-Tourigny J, Bosman LP, Wang W, et al. Sudden cardiac death prediction in arrhythmogenic right ventricular cardiomyopathy (ARVC): a multinational collaboration. Circ Arrhythm Electrophysiol 2021;14:e008509.
30. Orgeron GM, te Riele A, Tichnell C, et al. Performance of the 2015 international task force consensus statement risk stratification algorithm for implantable cardioverter-defibrillator placement in arrhythmogenic right ventricular dysplasia/cardio-myopathy. Circ Arrhythm Electrophysiol 2018;11:e005593.
31. Aquare GDO, De Luca A, Cappelletto C, et al. Comparison of different prediction models for the indication of implanted cardioverter-defibrillator in patients with arrhythmogenic right ventricular cardiomyopathy. ESC Heart Fail 2020;7:4080–4088.
32. Gasperetti A, Dello Russo A, Busana M, et al. Novel risk calculator performance in athletes with arrhythmogenic right ventricular cardiomyopathy. Heart Rhythm 2020;17:1251–1259.
33. Baudnoud P, Laredo M, Badenco N, et al. External validation of a risk prediction model for ventricular arrhythmias in arrhythmogenic right ventricular cardiomyopath. Can J Cardiol 2021;37:1263–1266.
34. Sen-Chowdhry S, Sirrs P, McKenna WJ. Desmoplakin disease in arrhythmogenic right ventricular cardiomyopathy: early genotype-phenotype studies. Eur Heart J 2005;26:1582–1584.
35. Xu Z, Zhu W, Wang C, et al. Genotype–phenotype relationship in patients with arrhythmogenic right ventricular cardiomyopathy caused by desmosomal gene mutations: a systematic review and meta-analysis. Sci Rep 2017;7:41387.
36. Verstraeten TE, van Lint FHM, Bosman LP, et al. Prediction of ventricular arrhythmia in phospholamban Arg14del mutation carriers-reaching the frontiers of individual risk prediction. Eur Heart J 2021;42:2842–2850.
37. Hodgkinson KA, Howes AJ, Boland P, et al. Long-term clinical outcome of arrhythmogenic right ventricular cardiomyopathy in individuals with a P5355L mutation in TMEM43 following implantable cardioverter-defibrillator therapy. Circ Arrhythm Electrophysiol 2016;9:e003589.
38. Antoniades L, Tsatsopoulou A, Anastasakis A, et al. Arrhythmogenic right ventricular cardiomyopathy caused by deletions in plakophilin-2 and plakoglobin (Naxos disease) in families from Greece and Cyprus: genotype–phenotype relationships, diagnostic features and prognosis. Eur Heart J 2006;27:2208–2216.
39. Christensen AH, Platonov PG, Jensen HK, et al. Genotype–phenotype correlation in arrhythmogenic right ventricular cardiomyopathy-risk of arrhythmias and heart failure. J Med Genet 2021. https://doi.org/10.1136/jmedgenet-2021-107911.
40. Corrado D, Perazzolo Marra M, Zorzi A, et al. Diagnosis of arrhythmogenic cardiomyopathy: the Padua criteria. Int J Cardiol 2020;319:106–114.