The yield of post-mortem genetic testing in sudden death cases with structural findings at autopsy

Running title: Post-mortem genetic testing

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ABSTRACT

Sudden cardiac death (SCD) is often associated with structural abnormalities of the heart during autopsy. This study sought to compare the diagnostic yield of post-mortem genetic testing in (1) cases with structural findings of uncertain significance at autopsy to (2) cases with autopsy findings diagnostic of cardiomyopathy. We evaluated 57 SCD cases with structural findings at cardiac autopsy. Next-generation sequencing using a panel of 77 primary electrical disorder and cardiomyopathy genes was performed. Pathogenic and likely pathogenic variants were classified using American College of Medical Genetics (ACMG) consensus guidelines. In 29 cases (51%) autopsy findings of uncertain significance were identified whereas in 28 cases (49%) a diagnosis of cardiomyopathy was established. We identified a pathogenic or likely pathogenic variant in 10 cases (18%); in 1 (3%) case with non-specific autopsy findings compared to 9 (32%) cases with autopsy findings diagnostic of cardiomyopathy (p=0.0024). The yield of genetic testing in SCD cases with autopsy findings consistent with cardiomyopathy is comparable to the yield in cardiomyopathy patients that are alive. Genetic testing in cases with findings of uncertain significance offers lower clinical utility than in cardiomyopathy, with lower yields than detected previously. This highlights the need for stringent evaluation of variant pathogenicity.

1. INTRODUCTION

Sudden cardiac death (SCD) is an important cause of death in the young. Predominant causes vary by age and include coronary artery disease, cardiomyopathy and primary electrical disorders¹. SCD and cardiac arrest in the young due to cardiomyopathies are, however, uncommon both in the general population² and during participation in competitive sport ³. They remain important because of their inherited nature and hence potential value in family screening. In 30-40% no diagnosis can be established at autopsy despite toxicological and histopathological analysis^{4,5}. These cases are classified as sudden arrhythmic death syndrome (SADS) and we have recently reported a 13% yield of pathogenic and likely pathogenic variants from post-mortem genetic testing (i.e. molecular autopsy) in a large set of 302 SADS cases⁵. In the current study, we aimed to determine the diagnostic yield of genetic testing in cases with autopsy findings of uncertain significance and compare them to cases with autopsy findings diagnostic of cardiomyopathy.

2. MATERIALS AND METHODS

2.1. Study population

Suitable cases were included from two cohorts of SCD cases as previously described ⁵. Cohort 1 consisted of a population-based coronial series (Cardiac Inherited Disease Registry, Auckland, New Zealand 2000-2009, [n=11]^{6,7}; consecutive referrals for autopsy (Royal Brompton Hospital, UK 2007-2011 [n=6], Sheffield Children's Hospital, UK 1985-2001 [n=1]); and consecutive referrals at the time of familial cardiac evaluation following a SCD (St George's and Lewisham Hospitals, UK 2009-2011 [n=3]; Academic Medical Centre (AMC), Netherlands 1995-2011 [n=2]). Cohort 2 consisted of 34 consecutive referrals for autopsy to St George's Hospital, UK 2012-2015. The study was approved by Research Ethics Service Wandsworth (Ref: 10H/0803/72). SCD was defined as death within one hour of onset of symptoms, or an unwitnessed death with the individual being seen in good health within 24 hours of death. Clinical history, prior symptoms, circumstance of death and family history were collected by direct contact with next of kin and from coroners' and pathologists' reports.

2.2. Autopsy evaluation

Cardiac autopsy was performed by locally recognised expert pathologists and included histological analysis. Cases where than classified into those (a) with autopsy findings of uncertain significance and (b) cases with autopsy findings diagnostic of cardiomyopathy. Findings of uncertain significance were defined as cardiac structural changes identified at autopsy of uncertain relevance to the cause of death. Adjudication of autopsy findings was performed by two of the authors using both recent guidelines for autopsy investigation of sudden cardiac death⁸ and as previously described⁹ and then confirmed by an expert pathologist (MNS).

2.3. Next generation sequencing and quality control

Next-generation-sequencing was performed as previously described ⁵. In summary, target regions were captured by in-solution hybridization target capture, using the Agilent SureSelect system (cohort 1) or the Illumina TruSight Cardio system (cohort 2). For SureSelect capture, custom hybridization-capture probes were designed using the Agilent eArray platform to target 201 genes implicated in cardiovascular disease. RNA baits targeted all exons of all Ensembl (v54) transcripts, including untranslated regions, with a 100-bp extension into adjacent introns. The TruSight Cardio Kit (Illumina) consists of comprehensive coverage of 174 genes (coding sequence region only) with known associations to 17 inherited cardiac conditions¹⁰. Libraries were prepared according to manufacturer's instructions and sequenced on the Illumina HiSeq (cohort 1) or NextSeq (cohort 2). Variant were annotated using ANNOVAR¹¹. We generated alignment summary metrics, coverage and callability metrics. We considered a base "callable" if it was sequenced with minimum read depth = 10x, base quality ≥ 20 , and mapping quality ≥ 10 . Target base callability of $\geq 90\%$ was achieved in 90% and 91% of samples from cohort 1, 2, respectively. Gene level callability is shown in Supplementary Table 1. We analyzed 77 genes (Supplementary Table 2) present in both capture systems that have been previously associated with primary electrical diseases or cardiomyopathies.

2.4. Classification of genetic variants

Variants with a minor allele frequency (MAF) >1/10,000 in the Genome Aggregation Database (gnomAD), synonymous variants not located at splice sites, and non-truncating variants in *TTN* were excluded. We applied American College of Medical Genetics (ACMG) consensus guidelines¹² to classify variants as (a) pathogenic, (b) likely pathogenic or as (c) variant of unknown significance (VUS). This included evidence from population data; identification of the variant in clinically affected patients in public and in-house disease databases; functional evidence; and concordance between the gene, type of variant and disease mechanism. Variants were adjudicated by two investigators as previously described⁵. Pathogenic and likely pathogenic variants according to ACMG guidelines were considered likely to cause disease in the patient. Genetic variants were submitted to ClinVar (<u>www.ncbi.nlm.nih.gov/clinvar/</u>) and the following accession number was obtained UB5428733.

2.5. Statistical analyses

Comparison was performed using the Mann-Whitney U test for non-normally distributed continuous variables and the chi-square or Fisher's exact test, as suitable, for categorical variables. All statistical analyses were performed using R (version 3.2.1.), and a p < 0.05 was used to indicate statistical significance. Percentages reported for clinical parameters are based on nonmissing data (n=50).

3. RESULTS

In total, 28 SCD cases with a diagnosis of cardiomyopathy and 29 cases with findings of uncertain significance during autopsy were included (median age of 29 and 32 years respectively, p-value=0.7659, **Table 1**). The most prevalent circumstances of death in the overall cohort were during sleep (n=15, 30%) or rest (n=14, 28%) with death occurring during exercise in 10 cases (20%). No statistically significant difference was observed in demographic and clinical characteristic between cases with autopsy findings of uncertain significance and cases with autopsy findings diagnostic of cardiomyopathy. Out of 28 SCD cases diagnosed as cardiomyopathy 6 (21%) had a premorbid diagnosis of cardiomyopathy and 3 (11%) had a family history of cardiomyopathy.

The main diagnoses established at autopsy were hypertrophic cardiomyopathy (HCM), dilated cardiomyopathy (DCM) and arrhythmogenic cardiomyopathy (ACM) in 7, 9 and 12 cases respectively (**Figure 1A**). Autopsy findings of uncertain significance were identified in 29 cases (**Figure 1B**), namely: idiopathic fibrosis (n=8), idiopathic left ventricular hypertrophy (i.e. without disarray) (n=11), or a combination thereof (n=8) and isolated mitral valve prolapse without associated myocardial fibrosis and extensive myxomatous degeneration (n=2).

Genetic testing uncovered a total of 40 rare protein-altering variants (MAF <1 in 10,000) in 23 different genes after variant filtering (**Supplementary Table 3**). In 33 patients (58%) no rare variant was identified: 20 cases with autopsy findings of uncertain significance and 13 cases diagnosed with cardiomyopathy. Manual curation against ACMG guidelines yielded 2 pathogenic and 8 likely pathogenic variants in 10 cases in the overall cohort, with an overall yield of 18% (10 of 57 patients, **Table 2**). The majority of the yield was from cases diagnosed with cardiomyopathy (9 of 28 [32%], 95% confidence interval: 18-51% **Figure 1C**) compared with non-specific findings (1 of 29 [3%], 95% confidence interval: 0.006-17%,

p=0.0024, **Figure 1D**). The two pathogenic variants were in *PKP2* and *TMEM43*, whereas the likely pathogenic variants were identified in *MYBPC3* (n=4), *DSP* (n=2), *TNNI3* (n=1) and *TTN* (n=1). In one patient who died at the age of 26 with ACM one likely pathogenic variant and one rare VUS were identified in *DSP* (**Table 2, Supplementary Table 3** case P1). One case with non-specific findings (idiopathic fibrosis) hosted the likely pathogenic *TTN* variant suggesting that this may be an early presentation of DCM as suggested by recent data from Finland¹³. No pathogenic or likely pathogenic variants were identified in genes associated with primarily electrical disorders.

4. **DISCUSSION**

This study indicates that the yield of pathogenic and likely pathogenic variants following molecular autopsy in cases with cardiomyopathy (32%) is comparable to the yield in living cardiomyopathy patients ¹⁴ but substantially greater than in cases with non-specific findings (3%). However, the yield in cases with non-specific findings did not differ significantly from that in our SADS cases from the same original SCD cohorts (13%) (p=0.2312)⁵. Across all groups we used a stringent MAF threshold of 1 in 10,000 and then applied ACMG variant classification guidelines ¹². Hertz et al. have previously studied cases with non-specific structural changes at autopsy ¹⁵ and reported a much higher prevalence (29%) of rare variants with likely functional effects. This had relied upon a higher MAF threshold (1 in 100) and computational approaches to classify variant pathogenicity which may have overestimated the yield. Interestingly, Papadakis et al. evaluated 41 families of SCD cases with non-specific findings at autopsy and diagnosed 45% with primary electrical diseases on clinical grounds alone without the availability of post-mortem genetic testing ⁹. This was similar to the results of clinical evaluation in families of SADS cases. The main clinical diagnosis was, however, the Brugada syndrome, where the diagnostic return of genetic testing is only approximately 20%¹⁶. It is therefore possible that by chance pathogenic or likely pathogenic variants for Brugada syndrome were not identified in the current cohort, regardless of whether Brugada syndrome may have been an important cause of death.

The most common uncertain findings in our study were idiopathic left ventricular hypertrophy (n=11), idiopathic fibrosis of the myocardium (n=8) or a combination thereof (n=8). It remains unclear whether these uncertain findings during autopsy are innocent bystander or truly involved in sudden death. Both idiopathic left ventricular hypertrophy as well as myocardial fibrosis may present as a partial phenotypic expression of an underlying

cardiomyopathy in association with sudden death. This has been suggested for myocardial fibrosis by a recent Finnish study, where the authors identified a higher yield than our study of pathogenic or likely pathogenic cardiomyopathy associated variants in 10% of sudden death cases in whom myocardial fibrosis was noted during autopsy (in genes associated to ACM [n=2], HCM [n=4], and DCM [n=4]). ¹³ In line with our findings, no variants were identified in genes associated to primary electrical disorders.

As we only screened our cohort for 77 genes, we cannot exclude the possibility of pathogenic variants in other genes that are associated to these subtle findings during autopsy and predispose to sudden death. Furthermore, a polygenic architecture with a combination of genetic variation, underlying these findings and sudden death cannot be excluded. In addition, rather than a genetic cause, sudden death could be the results of environmental factors (e.g. infection) with or without interaction with predisposing genetic factors.

4.1. Limitations

This study is limited by the small number of cases included (n=57) and lack of comprehensive clinical evaluation of relatives. The combined yield of genetic and clinical screening of relatives is therefore unknown. Although autopsies were performed at different centres by locally recognised experts using international guidelines, this may have resulted in non-uniformity of autopsy results. Unfortunately, quantitative criteria for autopsy diagnoses are still limited. Although included cases were either consecutive referrals or from population studies, we acknowledge that there may be an element of referral bias

5. CONCLUSION

This study highlights the importance of applying stringent criteria to classify variants, resulting in a lower yield of actionable genetic results in cases with non-specific findings at

autopsy. Currently, whilst molecular autopsy in these cases offers less clinical utility than in cardiomyopathy, its yield is not significantly less than in SADS cases based on these data. Larger studies coupled with family findings are necessary to elucidate the importance of monogenic diseases and the role for genetic testing in this group.

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FIGURE LEGEND

Figure 1. Overview of structural finding during autopsy and yield of genetic testing.

Legend: Figure summarizing autopsy findings and yield of genetic testing in cases with autopsy findings diagnostic of cardiomyopathy (**A** and **C**) and cases with autopsy findings of uncertain significance (**B** and **D**). *Abbreviations*: ACM, arrhythmogenic cardiomyopathy; CAD, coronary artery disease; DCM, dilated cardiomyopathy; HCM, hypertrophic cardiomyopathy; IF, idiopathic fibrosis; ILVH, idiopathic left ventricular hypertrophy; LP, likely pathogenic; MVP, mitral valve prolapse; P, pathogenic

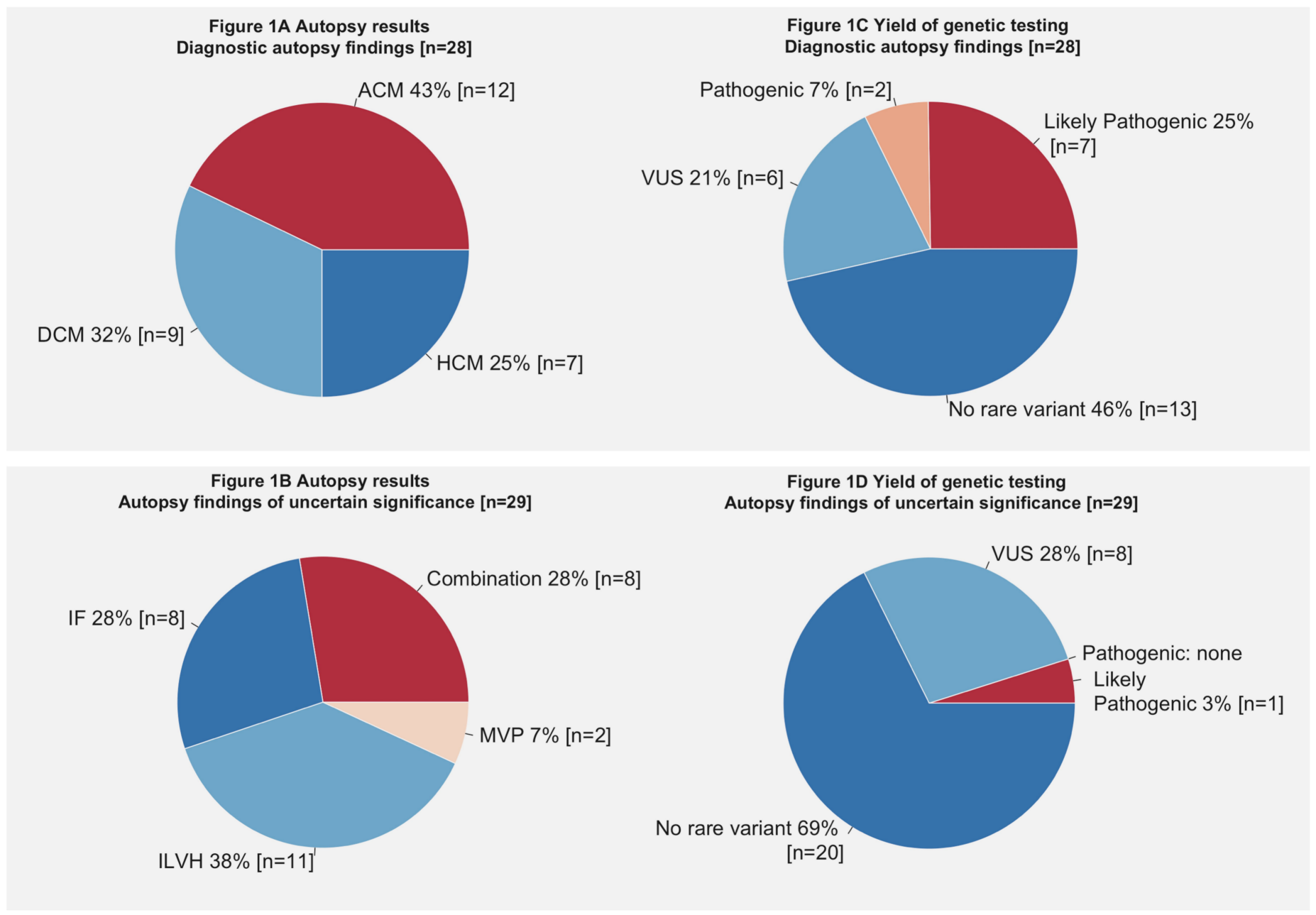


 Table 1. Demographic and clinical characteristics in the sudden death cohort

	Autopsy findings							
Variable	uncertain significance (n=29) †	diagnostic of cardiomyopathy (n=28) †						
Age at death, median age [IQR], years	32 [21-38]	29 [24-38]						
Male gender, no. (%)	86% [25/29]	68% [19/28]						
Circumstances of death, no./total no. (%) §								
Exercise and extreme emotion	25% [6/24]	19% [5/26]						
Sleep and rest	63% [15/24]	54% [14/26]						
Light Activity	12% [3/24]	27% [7/26]						
Symptoms prior to death, no./total no. (%) §								
Syncope and/or Seizures	19% [5/26]	7% [2/27]						
Palpitations	8% [2/26]	11% [3/27]						
None	73% [19/26]	81% [22/27]						
Family history of cardiomyopathy, no./total no.	7% [2/29]	11% [3/28]						

Legend: * Positive outcome was defined as a pathogenic or likely pathogenic variant according to the ACMG guidelines (11).

† No statistically significant difference was observed in demographic and clinical characteristic between cases with autopsy findings of uncertain significance and cases with autopsy findings diagnostic of cardiomyopathy.

 \S Please note missing data for these parameters and that percentages are based on non-missing data.

Table 2. Overview of pathogenic and likely pathogenic variants in the sudden death cohort

Case	Age at death (yrs)	Sex	Circumstances of Death	Symptoms Before Death	Autopsy finding	Gene	Chr	Start*	End	Ref/Alt	MAF‡ dbSNP	cDNA change*	Amino acid change*	ACMG Classification	ACMG criteria †
P1	26	F	Light Activity	Seizures	ACM	DSP	6	7580818	7580818	T/G	None	c.4395T>G	p.(Y1465*)	Likely Pathogenic	PVS1.PM2
P2	28	М	Light Activity	None	ACM	DSP	6	7581692	7581692	C/T	None	c.5269C>T	p.(Q1757*)	Likely Pathogenic	PVS1, PM2
Р3	20	М	Exercise	None	ACM	TMEM43	3	14183165	14183165	C/T	None rs63750743	c.1073C>T	p.(S358L)	Pathogenic	PS3,PS4,PM2,PP3
P4	37	М	Sleep	None	ACM	PKP2	12	32955433	32955438	GGGTGT/G	1.6x10 ⁻⁵	c.2066_2070delACACC	p.(H689Pfs*8)	Pathogenic	PVS1, PS4
Р5	29	М	Light Activity	Seizures	DCM	TNNI3	19	55665462	55665462	C/T	4x10 ⁻⁵ rs397516354	c.485G>A	p.(R162Q)	Likely Pathogenic	PS4,PP5,PP3
P6	29	М	Exercise	None	HCM	MYBPC3	11	47373030	47373030	A/AC	None	c.51dupG	p.(S18Tfs*31)	Likely Pathogenic	PVS1,PM2
P7	19	F	Sleep	Palpitations	НСМ	MYBPC3	11	47359112	47359112	T/C	None rs375675796	c.2432A>G	p.(K811R)	Likely Pathogenic	PM1,PM2, PP3,PP5
P8	19	М	NA	None	НСМ	MYBPC3	11	47372895	47372905	CGTGTGC CCTCT/C	None/rs397515925	c.177_187delAGAGGGCACAC	p.(E60Afs*49)	Likely Pathogenic	PVS1
P9	46	F	Sleep	None	HCM	MYBPC3	11	47372960	47372960	C/CG	None	c.121dupC	p.(R41Pfs*8)	Likely Pathogenic	PVS1,PM2
P10	40	М	NA	NA	IF	TTN	2	179440001	179440001	T/A	None	c.70858A>T	p.(R23620*)	Likely Pathogenic	PVS1,PM2

Table legend: * Genomic position refers to GRCh37. The following transcripts were used: DSP: NM 004415.4; MYBPC3:NM 000256.3; PKP2:NM 00105242.2; TNNI3:NM 000363.4; TMEM43:NM 024334.2; TTN:NM 001267550.2.

† See ACMG guidelines 12 for further information on classification criteria.

* MAF is based on total number of individuals in the Genome Aggregation Database (gnomAD).

Abbreviations: ACM, arrhythmogenic cardiomyopathy; Alt, alternate allele; cDNA, coding DNA; Chr, chromosome; DCM, dilated cardiomyopathy; F, female; HCM, hypertrophic cardiomyopathy; IF, idiopathic fibrosis; LP, likely pathogenic; M, male; MAF, minor allele frequency; NA, not available; P, pathogenic; Ref, reference allele.