Sudden Death and Left Ventricular Involvement in Arrhythmogenic Cardiomyopathy

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Abstract

**Background:** Arrhythmogenic cardiomyopathy (ACM) is an inherited heart muscle disorder characterized by myocardial fibro-fatty replacement and an increased risk of sudden cardiac death (SCD). Originally described as a right ventricular (RV) disease, ACM is increasingly recognized as a biventricular entity. We evaluated pathological, genetic, and clinical associations in a large SCD cohort.

**Methods:** We investigated 5205 consecutive cases of SCD referred to a national cardiac pathology center between 1994 and 2018. Hearts and tissue blocks were examined by expert cardiac pathologists. Following comprehensive histological evaluation, 202 cases (4%) were diagnosed with ACM. Of these, 15 (7%) were diagnosed ante-mortem with dilated cardiomyopathy (DCM) (n=8) or ACM (n=7). Prior symptoms, medical history, circumstances of death, and participation in competitive sport were recorded. Post-mortem genetic testing was undertaken in 24/202 (12%). Rare genetic variants were classified according to American College of Medical Genetics and Genomics (ACMG) criteria.

**Results:** Of 202 ACM decedents (35.4±13.2 years; 82% male), 157 (78%) reported no prior cardiac symptoms. Forty-one decedents (41/202; 20%) were participants in competitive sport. The adjusted odds of dying during physical exertion were higher in males than females (OR 4.58; 95% CI 1.54-13.68; p=0.006) and in competitive athletes compared with non-athletes (OR 16.62; 95% CI 5.39-51.24; p<0.001). None of the decedents with an ante-mortem diagnosis of DCM fulfilled definite 2010 Task Force criteria. Macroscopic appearance of the heart was normal in 40/202 (20%) cases. There was left ventricular (LV) histopathological involvement in 176/202 (87%). Isolated RV disease was seen in 13%, isolated LV disease in 17%, and biventricular involvement in 70%. Among whole hearts, the most common areas of fibro-fatty infiltration were the LV posterobasal (68%) and anterolateral walls (58%). Post-mortem genetic testing yielded pathogenic variants in ACM-related genes in 6/24 (25%) decedents.

**Conclusions:** SCD due to ACM affects males predominantly, most commonly occurring during exertion in athletic individuals in the absence of prior reported cardiac symptoms. LV involvement is observed in the vast majority of SCD cases diagnosed with ACM at autopsy. Current Task Force criteria may fail to diagnose biventricular ACM prior to death.

**Key Words:** Arrhythmogenic cardiomyopathy; sudden cardiac death; left ventricular arrhythmogenic cardiomyopathy
Clinical Perspective

What is new?

- In this large comprehensive autopsy study, we demonstrate that left ventricular involvement is observed in most decedents with arrhythmogenic cardiomyopathy and the left ventricle is exclusively involved in nearly a fifth of cases.
- Age at death, sex, normal macroscopic appearance of the heart, and participation in competitive sport were not associated with the presence of left ventricular involvement.
- We describe diagnostic histopathological criteria for arrhythmogenic cardiomyopathy involving either or both ventricles.

What are the clinical implications?

- This study identified that the heart was macroscopically normal in 20% of decedents with ACM; expert pathological assessment, including histology, is therefore crucial to inform diagnosis in cases of initially unexplained sudden cardiac death.
- Left ventricular variants of arrhythmogenic cardiomyopathy may evade clinical detection using current diagnostic tools; this should be addressed in future revisions of Task Force criteria.
Introduction

Arrhythmogenic cardiomyopathy (ACM) is a genetic heart muscle disorder characterized by myocardial atrophy and fibrofatty replacement of the ventricular myocardium. Originally described as arrhythmogenic right ventricular cardiomyopathy/dysplasia (ARVC) (1,2), increased recognition of left ventricular (LV) involvement has recently led to adoption of the term ‘ACM’ (3,4). Arrhythmogenic cardiomyopathy has an estimated prevalence of 1:2000-1:5000 and is an important cause of sudden cardiac death (SCD) among young individuals including athletes (3,5,6). Typically, ACM follows a Mendelian autosomal dominant inheritance pattern, where disease-causing mutations in over thirteen genes have been identified (7).

Approximately half of the index cases with a clinical diagnosis harbor putative mutations in genes encoding desmosomal proteins: plakophilin-2 (PKP2), desmoplakin (DSP), desmocollin-2 (DSC2), junction plakoglobin (JUP), and desmoglein-2 (DSG2).

Clinical presentation of ACM is heterogeneous, and diagnosis can be challenging (3). This is reflected by the Task Force criteria, which integrate a number of structural, histopathological, electrocardiographic, familial, arrhythmic, and genetic parameters (6). Moreover, these criteria are derived from cohorts with predominantly right ventricular involvement (ARVC), thus can potentially fall short for a significant proportion of those with LV dominant or biventricular disease. Task Force criteria require the presence of fibrofatty replacement of the right ventricular (RV) free wall myocardium on endomyocardial biopsy or at autopsy (6), however, pathological and imaging studies have reported LV involvement ranging between 16% and 76% of cases (8-12). Cardiovascular Magnetic Resonance (CMR) may show late gadolinium enhancement, indicative of myocardial fibrosis, in a sub-epicardial or mid-myocardial distribution, usually within the LV inferior or inferolateral walls (13).
The purpose of this study was to report the pathological features of a large cohort of decedents experiencing SCD from ACM, with a particular focus on the presence of LV involvement. In addition, we sought to report on the clinical characteristics and genetic associations of the cohort.

Methods

The data, analytical methods, and study materials will not be made available to other researchers for purposes of reproducing the results or replicating the procedure. All unexpected sudden deaths in the UK are reported to the coroner, who determines the need for autopsy examination. The Cardiac Risk in the Young Center for Cardiac Pathology (CRY CCP) at St George’s, University of London provides a nationally recognized expert cardiac pathology service. Referral is initiated voluntarily by the coroner’s pathologist following an unexplained SCD, or if there is suspicion of an inherited heart condition. The center receives approximately 500 SCD cases per annum, just over half under the age of 35 years. A report to the coroner is issued within a two-week period.

All referrals to the CRY CCP undergo a full coroner’s pathologist autopsy. Since 2013, spleen samples have been collected for post-mortem genetic testing. Pathological findings and corresponding clinical information were entered into a database incorporating imaging from histological sections and autopsy/toxicology reports from UK coroners. Ethical approval for this study was obtained from the UK National Health Service Research Ethics Committee. Informed consent was provided by the next of kin at the time of referral.
Study group

We evaluated 5205 consecutive SCD cases referred to the CRY CCP between 1994 and 2018 (Figure 1). Inclusion criteria for this study were: 1) SCD defined by one of the following: an unheralded witnessed instantaneous death; a death preceded by a prodrome of acute cardiac symptoms up to one hour before death; or an unwitnessed case without a prior deterioration in the preceding 24 hours. 2) Pathological diagnosis of ACM at autopsy. 3) Absence of an extra-cardiac cause of death. 4) Negative toxicology screen. 5) Absence of obstructive coronary artery disease (atherosclerosis with residual luminal diameter <1mm²). Decedents with an ante-mortem diagnosis of cardiomyopathy were included.

Demographics and Clinical Characteristics

A questionnaire was sent to all referring pathologists and coroners’ officers relating to demographics, medical history, symptoms, family history, sports participation, and circumstances of death. Additional information was obtained by autopsy reports from the referring pathologist, primary care/hospital correspondence, and family member interviews. This was acquired within three months of initial referral to our center. Media and sports club correspondence was also reviewed for competitive sports participation. Decedents were considered competitive athletes if they engaged in an organized team or individual sport requiring regular training and participation in competitive events (14). For those with an ante-mortem diagnosis of cardiomyopathy, the clinical notes and investigations (including electrocardiogram (ECG), echocardiography, ambulatory cardiac monitoring, signal-averaged ECG, and CMR) were subsequently requested and analyzed according to the revised Task Force criteria (6).
Autopsy evaluation

Whole hearts (3018/5205; 58%) and cases with tissue blocks (2187/5205; 42%) underwent detailed histopathological analysis, including microscopic examination of tissue from both ventricles. Sections were fixed in formalin, embedded in paraffin, and stained with Hematoxylin and Eosin (H&E). The following areas were routinely examined: the right ventricular outflow tract; the anterior, lateral, and posterior right ventricle (mid-ventricular level); interventricular septum; anterior, lateral and posterior left ventricle; the three major coronary arteries; and the ascending aorta. The apices of both ventricles were not examined in the absence of an overt structural abnormality. Information from the referring pathologist’s autopsy was also recorded, incorporating macroscopic appearances of the heart, body weight, and heart weight. At least two tissue blocks from both ventricles were required for study inclusion.

ACM was defined on microscopic analysis as myocyte degeneration associated with intermixed fat and fibrosis (within the same microscopic field) from the subepicardial region inward or transmural in either or both ventricles (>20% in at least two tissue blocks of 4cm²; H&E stain). Disease involvement within the interventricular septum was defined by myocyte degeneration associated with intermixed fat and fibrosis from either side or within the mid-wall. This definition has been updated from a previous report (15). Figure 2 compares normal histological appearances in the right ventricle with a case meeting diagnostic criteria for ACM.

All cases included in the study were examined by the senior author (MNS), in addition to review by at least one other pathologist.

Post-mortem genetic testing

Following extraction of DNA from retained splenic tissue (n=24), cardiac gene panel testing was performed using the Illumina TruSight Cardio (174 genes) panel (or a custom Agilent SureSelect
with equivalent content) (*Supplemental Table 1*) and sequenced on the Illumina platform (NextSeq or HiSeq), as previously described and validated (16,17). Variant annotation was then undertaken in-house using SnpEff v4.3T (build 2017-11-24) (18), GRCh37.75 and ANNOVAR (version 2017-07-17) (19). Rare variants were defined as those with a minor allele frequency (MAF) cut off <0.01% in the Exome Aggregation Consortium (ExAC) general population database (exac.broadinstitute.org). Rare variants were then assessed for pathogenicity according to the American College of Medical Genetics and Genomics (ACMG) criteria (20). These criteria integrate a number of factors including population data, functional data, computational data, and segregation analysis to inform assessment of pathogenicity.

**Statistical analysis**

All variables were graphically inspected and summarized according to their nature and by outcomes of interest (i.e. mean, standard deviation, median) for continuous variables and proportions for binary or categorical variables. Data are presented as mean ± 1 standard deviation or percentages and appropriate tests (t tests, (Fisher) chi-squared) were employed for a preliminary flavor of variable associations.

There were two outcomes of primary interest, notably death occurring during physical exertion and LV histopathological involvement. Univariable and multivariable logistic regression analyses were used to understand the strength of the crude and adjusted associations between these two outcomes and available demographics and clinical features in the ACM SCD group. The Hosmer-Lemeshow test was used to assess models’ goodness of fit. A *P*-value of <0.05 was deemed statistically significant. Statistical analyses were performed using Stata IC/15 (StataCorp, College Station, TX, US). The author ICS holds responsibility for the statistical methodology applied to the data in this paper.
Results

Baseline demographics

Of 5205 cases of SCD referred to our unit, 202 (4%) were diagnosed with ACM. The majority of decedents from ACM were male (166/202; 82%) and white (182/202; 90%). The mean age of death was 35.4±13.2 years (median 34.5 years).

Clinical characteristics

The great majority of cases (187/202; 93%) did not have an ante-mortem diagnosis of cardiac disease. Only 15/202 (7%) decedents carried a diagnosis of cardiomyopathy, of which 7/15 (47%) were labelled as ACM and 8/15 (53%) as dilated cardiomyopathy (DCM). Six of the overall SCD cohort with ACM (6/202; 3%) had an implantable cardioverter defibrillator (ICD) in situ. Fifteen (7%) reported a family history of SCD under the age of 35 years, none of whom were diagnosed with cardiomyopathy in life. Two decedents with a positive family history had been referred for clinical investigation. Table 1 outlines clinical and pathological characteristics stratified by circumstances of death.

In most decedents with ACM (157/202; 78%), there were no reported symptoms prior to death; twenty (10%) recorded syncope prior to death, which was associated with an ante-mortem diagnosis of cardiomyopathy in six cases. The remaining decedents (25/202; 12%) documented a prior history of palpitations (8%), chest pain (2%), and pre-syncope (2%). Eighty-three (41%) of the overall cohort died during physical exertion, and 105 (52%) died at rest or during sleep. Decedents with an existing diagnosis of cardiomyopathy were more likely to have died at rest or during sleep compared with during physical exertion (87% vs 7%, respectively; p=0.004).

Circumstances of death were unspecified in 14/202 (7%); there was no significant pattern in missing data with respect to other variables. Among young (<35 years old) decedents with ACM,
there were more deaths during physical exertion compared with older decedents (50% vs 33%, respectively; p=0.01).

Pathological characteristics

In the majority of cases (120/202; 59%), the whole heart was received for analysis. Of the remaining cases with tissue blocks (41%), the mean number of blocks received across both ventricles was 8.3±3.2 (range 4-15). In 40/202 (20%), macroscopic examination of the heart did not yield an overt structural abnormality (Supplemental Figure 1). Heart weight and body weight were recorded in 193/202 (96%) and 131/202 (65%) cases respectively, and demonstrated a positive correlation (r=0.55 p<0.001, n=129). Overall, 27% of men recorded heart weights greater than 500g, and 23% of females greater than 400g. The association between body weight and heart weight was independent of LV histopathological involvement (n=129, p=0.74).

Following histological evaluation, none of the cases meeting diagnostic criteria for ACM showed myocardial inflammatory infiltrates.

Ventricular involvement

An overview of ventricular histopathological involvement is presented in Figure 3. Left ventricular fibrofatty infiltration was present in 176/202 (87%) decedents. Disease exclusive to the LV was observed in 35/202 (17%), and RV in 26/202 (13%), with 141/202 (70%) having evidence of biventricular disease. Among whole hearts referred to our center (120/202; 59%), the most common sites of LV disease involvement were the posterobasal wall (81/120; 68%) and anterolateral wall (69/120; 58%). In the RV, the anterolateral wall (77/120; 64%), and right ventricular outflow tract (57/120; 48%) were most frequently implicated. A logistic regression model found no significant association between LV disease and age at death (p=0.31), male sex
(p=0.58), macroscopically normal appearance of the heart (p=0.49), or competitive sport participation (p=0.27) (Supplemental Table 2).

**Competitive athletes**

Of 202 decedents with ACM, 41 (20%) participated in competitive sport and the majority of the decedents (35/41; 85%) were aged <35 years. Thirty-one athletes (31/41; 76%) were engaged in sports with a high dynamic component (Supplemental Figure 2). The vast majority (37/41; 90%) of athletes died during physical exertion, including 30 athletes (30/41; 73%) who died during participation in sport. In contrast, 46/147 (31%) of the non-athlete decedents died during physical exertion (90% vs 31%, respectively; p<0.001). There was no significant difference in disease distribution within the heart between athletes and non-athletes (p=0.10).

Multivariable analysis showed that male sex (OR 4.58; 95% CI 1.54-13.68; p=0.006) and participation in competitive sport (OR 16.62; 95% CI 5.39-51.24; p<0.001) were independent predictors of death during physical exertion (Table 2).

**Genetic testing**

Genetic testing was performed in all cases with post-mortem DNA available (n= 244 [12%]). A total of 27 rare variants were identified in 14/24 (58%) cases. Six decedents (6/24; 25%) hosted single pathogenic variants in known ACM genes: *DSP* (n=2; truncating mutations), *TMEM43* (n=2; missense mutations), and *PKP2* (n=2; frameshift mutations) (Table 3). The same pathogenic *TMEM43* variant (p.Ser358Leu) was identified in two unrelated individuals and has been described in a number of different cohorts in the ClinVar database (www.ncbi.nlm.nih.gov/clinvar). None of the decedents hosted more than one pathogenic variant. Four decedents harbored rare variants of uncertain significance (VUSs) according to ACMG criteria in desmosomal-related genes: one male with a pathogenic DSP variant hosted
both missense and splice-site variants in DSC2; a previously reported rare missense variant was identified in DSC2 was identified in one decedent; and novel missense variants in DSP and PKP (VUS) were identified in two individuals also harboring pathogenic variants in these genes. All VUSs are outlined in Supplemental Table 3.

**Clinico-pathological correlations**

Clinical investigations were available in 10 of the 15 (67%) decedents with an ante-mortem diagnosis of cardiomyopathy. Seven subjects were diagnosed with DCM of which four had ICDs in situ, and three decedents were diagnosed with ACM of which one had an ICD. Cardiac MRI had been performed in half and showed evidence of LV disease in 4/5 (80%). Modified Task Force criteria were applied retrospectively to all 10 cases. None of the decedents with a clinical diagnosis of DCM achieved a definite diagnosis of ACM using Task Force criteria (Table 4).

*Figure 4* shows ECG, CMR, and echocardiogram appearances in an individual with LV-exclusive ACM diagnosed with DCM prior to death. Of seven presumptive DCM diagnoses, none fulfilled major or minor TFC for imaging (*Supplemental Table 4*) whilst six (86%) exhibited T-wave inversion in at least one of the lateral leads (I, aVL, V5, and V6) and none demonstrated T-wave inversion in V1-V3.

Following autopsy examination, macroscopic appearances of the heart were abnormal in all 10 subjects. Post-mortem genetic testing was undertaken in only a few and identified a DSP pathogenic variant in one DCM case, and a PKP2 pathogenic variant in an individual with ACM.

**Discussion**

To our knowledge, we report the largest pathological series of ACM in SCD employing contemporary definitions. We demonstrate that comprehensive histopathological evaluation
identifies LV involvement in the overwhelming majority (87%) of individuals experiencing SCD from ACM. This appears to be independent of age at death, sex, normal macroscopic appearance of the heart, and participation in competitive sport.

**Clinical associations and implications**

While ACM affects both sexes, males are more often phenotypically affected with a higher mortality rate and an increased incidence of cardiac arrest as the initial manifestation (21,22). These observations are consistent with our cohort, which was mainly male (82%). Furthermore, whilst death during physical exertion is well described (15), 52% of deaths in our study occurred at rest or during sleep. Syncope is also a common presenting symptom and marker of SCD risk. A quarter of a US cohort of ACM patients (N=100) had a history of syncope (23), and an Italian registry of 301 ACM patients reported that syncope was associated with a three-fold increased risk of life-threatening arrhythmic events during follow-up (24). Nonetheless, most subjects in our study (78%) failed to report any cardiac symptoms prior to presenting with SCD, and only 10% had experienced some form of syncope.

Over half of our decedents diagnosed with cardiomyopathy in life were labelled as DCM, but showed pathological features of ACM at autopsy. The overlap between DCM and ACM may present several diagnostic challenges as reflected in the finding that none of our decedents with an ante-mortem diagnosis of DCM fulfilled current Task Force criteria for a definite diagnosis of ACM. Among the ante-mortem DCM cohort, six (86%) exhibited T-wave inversion in at least one of the lateral leads: I, aVL, V5, and V6; none demonstrated T-wave inversion in V1-V3. This highlights the phenotypic spectrum of ACM and the potential for biventricular disease to evade clinical diagnosis and subsequent risk management. We also identified a family history of young SCD (<35 years old) in 7% of the cohort, none of whom were diagnosed with
cardiomyopathy during life. Guidelines recommend comprehensive family evaluation in this setting and may have led to recognition of risk prior to death (25).

**Myocardial involvement at autopsy**

An early pathological study of 30 ACM hearts revealed LV involvement in 47% of their cohort (8). Patchy inflammatory infiltrates were also detected in 20 (67%) cases, a finding that was not replicated during histological examination of our study group. A later study of 42 cases showed macroscopic or microscopic features of LV disease in 76%, with a predilection for the septum and free wall (posteroseptal and posterolateral regions) (9). In keeping with this and other prior studies, the posterobasal wall was the commonest location of LV fibrofatty infiltration (68%) in our group. More recently, LV histological abnormalities were found in a lower proportion (32%) of 200 SCD cases attributed to ACM (10). However, fatty infiltration of the RV was considered pathological in this study, even in the absence of fibrosis. Our observations suggest that fatty infiltration of the RV should not be considered diagnostic for ACM. Our study also reports LV histopathological involvement in a much higher proportion of decedents with ACM (87%). The role of inflammation in ACM has yet to be fully resolved. The lack of myocardial inflammatory infiltrates detected among our cohort is contrary to earlier assertions linking inflammation to severity of outcome in ACM (26).

**Correlation with imaging studies**

The natural history of LV involvement in ACM is poorly understood. Our finding that LV involvement was not associated with age at death or macroscopic appearance of the heart at autopsy corroborates imaging studies that suggest LV involvement may occur at an early stage (27). Ghannudi et al (28) evaluated 21 patients with ACM and found 52% had LV involvement on CMR. The degree of RV impairment was similar between those with isolated RV
involvement and disease affecting the LV. Rastegar and colleagues also studied 78 ACM mutation carriers of whom 38 had structural abnormalities on CMR, including the LV in 55% (29). The higher proportion of cases with LV involvement in our cohort may reflect a greater sensitivity of histopathology for fibrosis.

**Competitive sport**

The link between athletic activity and SCD risk in patients with ACM is widely acknowledged (30). A previous study found that participation in competitive sport was associated with a two-fold increased risk of ventricular arrhythmia and SCD in ACM, but recreational sport followed a more benign course (31). Data from a prospective multinational registry of athletes with ICDs also identified ACM as the only diagnosis associated with appropriate shocks during competitive sport (32). Current guidelines recommend against participation in competitive and/or endurance sport among individuals meeting Task Force criteria (33). Our data show that most competitive athletes who died with ACM were engaged in sports with a maximum dynamic component, i.e. requiring the highest cardiac output, and the great majority (90%) of these deaths occurred during physical exertion. We found no difference in either RV-exclusive disease or LV involvement between athletes and non-athletes, suggesting that exercise-induced remodeling alone is insufficient to lead to the ACM phenotype.

**Genetic evaluation and implications**

To our knowledge, there have been no published studies evaluating the utility of post-mortem genetic testing in ACM. Diagnostic yield may be influenced by several factors, such as disease presentation, family history, and geographic region (34). Our use of a stringent MAF threshold and application of ACMG criteria may have resulted in a more conservative yield when compared with prior genetic studies (34,35). Among decedents unheralded with cardiac disease,
none had a reported family history of ACM. The only pathogenic variants identified in this group were in \textit{TMEM43} and \textit{PKP2}, the same missense variant being identified in both (unrelated) \textit{TMEM43} subjects. This variant had previously been reported in multiple ACM families internationally, is highly penetrant and has strong evidence for pathogenicity (36). Both probands exhibited histopathological involvement of the LV, consistent with prior data on the mutation, which is associated with a particularly malignant form of ACM (Arrhythmogenic Right Ventricular Cardiomyopathy Type 5). \textit{DSP} is also highly penetrant and associated with LV involvement and higher risk, especially truncating variants (37,38). Two subjects hosted truncating pathogenic \textit{DSP} variants, and both had prior diagnoses of DCM.

**Implications for the pathologist and clinician**

We demonstrated that 20\% of ACM cases were macroscopically normal on initial inspection, regardless of LV involvement. Our findings are in line with earlier studies that recognize a concealed phase in ACM, where SCD may occur in the absence of overt structural abnormalities (39). This reinforces the importance of histological analysis of hearts in all cases of SCD. Moreover, previous reports have described sudden death occurring in PKP2 mutation carriers exhibiting structurally normal hearts at autopsy (40,41); this may suggest a propensity for malignant arrhythmia in those with a pure electrical phenotype.

In 17\% of our cases, fibrofatty infiltration was identified exclusively within the LV, consistent with the previously described entity of ‘LV-Dominant’ ACM (13). LV-dominant ACM may manifest clinically with inferolateral T wave inversion on the 12-lead ECG, arrhythmias of LV origin, and suggestive appearances on cardiac imaging. Thus, the absence of RV structural abnormality may lead to poor recognition of ‘non-classical’ disease utilizing current diagnostic criteria. Moreover, the significant minority of ACM SCD cases with LV-
exclusive disease support inclusion of LV pathological and clinical markers in future revision of Task Force criteria.

Study limitations

This study relied on referral of SCD cases to an expert cardiac pathology center, therefore, there may be an element of referral bias as pathologists may choose to refer more challenging cases and potentially a higher proportion of cases with LV involvement. In 41% of cases, the whole heart was not sent to the center for analysis and macroscopic appearances of the heart were noted by the referring pathologist. Consequently, sampling technique may differ according to local protocols. However, tissue blocks from both ventricles were received in each case and histological diagnostic criteria consistently applied. Retrospective review of clinical information, competitive sports participation, and family member interviews could be subject to recall bias and may have underestimated the true extent of symptoms experienced in life that may otherwise have not been documented. Finally, whilst genetic testing was performed on all possible individuals with retained DNA for molecular autopsy, the group with genetic results is small so inference of genotype mediated risk is inappropriate. A larger genetic testing uptake in the cohort would facilitate important genotype-phenotype correlations and assessment of genetic yield, which would be an important area worthy of future research.

Conclusions

Most ACM-related SCDs occurred without preceding ante-mortem symptoms or relevant family history. Exercise-related SCD was observed in the majority of competitive athletes, with competitive sport participation conveying significantly increased odds of death during physical exertion. A fifth of cases demonstrated a macroscopically normal heart, emphasizing the importance of histopathological evaluation in cardiac autopsy. LV involvement was present in
the great majority of cases, with 17% exhibiting disease exclusive to the LV. Our study supports inclusion of left ventricular structural involvement in future revision of Task Force diagnostic criteria.

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Disclosures

None.
References


Table 1. Clinical and pathological characteristics stratified by circumstances of death. Values presented are mean±standard deviation, median, or n (%) as appropriate.

<table>
<thead>
<tr>
<th></th>
<th>Overall</th>
<th>Death during exertion</th>
<th>Death at rest / during sleep</th>
<th>P value</th>
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<tr>
<td>N (%</td>
<td>202</td>
<td>83 (41)</td>
<td>105 (52)</td>
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<tr>
<td>Age at death (years)</td>
<td>35.4±13.2, 34.5</td>
<td>32.7±12.5, 29</td>
<td>37.7±13.1, 37</td>
<td>0.01</td>
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<td>Male, n (%)</td>
<td>166 (82)</td>
<td>78 (94)</td>
<td>76 (72)</td>
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<td>White, n (%)</td>
<td>182 (90)</td>
<td>74 (89)</td>
<td>96 (91)</td>
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<td>Body weight, median</td>
<td>81.5±17.4, 81</td>
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<td>81.0±17.8, 80.55</td>
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<td>Diagnosis of cardiomyopathy ante-mortem, n (%)</td>
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<td>1 (1)</td>
<td>13 (12)</td>
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<td>Prior syncope, n (%)</td>
<td>20 (10)</td>
<td>8 (10)</td>
<td>12 (11)</td>
<td>0.69</td>
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<td>Asymptomatic status, n (%)</td>
<td>157 (78)</td>
<td>65 (78)</td>
<td>80 (76)</td>
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<td>FH SCD (&lt;35 years old), n (%)</td>
<td>15 (7)</td>
<td>5 (6)</td>
<td>9 (9)</td>
<td>0.58</td>
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<td>Macroscopically normal appearance of the heart at autopsy, n (%)</td>
<td>40 (20)</td>
<td>18 (22)</td>
<td>20 (19)</td>
<td>0.70</td>
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<td>Disease confined to right ventricle, n(%)</td>
<td>26 (13)</td>
<td>11 (13)</td>
<td>14 (13)</td>
<td>0.99</td>
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<tr>
<td>Disease confined to left ventricle, n(%)</td>
<td>35 (17)</td>
<td>15 (18)</td>
<td>19 (18)</td>
<td>0.99</td>
</tr>
<tr>
<td>Biventricular disease, n(%)</td>
<td>141 (70)</td>
<td>57 (69)</td>
<td>72 (69)</td>
<td>0.99</td>
</tr>
</tbody>
</table>

FH indicates family history; SCD, sudden cardiac death.
P values represent the strength of association between circumstances of death and the available clinical or demographic variables. Comparisons are made using appropriate tests upon each variable’s nature. Circumstances of death were unspecified for 14/202 individuals.
**Table 2.** Analyses of death during physical exertion among ACM decedents.

<table>
<thead>
<tr>
<th></th>
<th>Univariable Analysis</th>
<th>Adjusted Analysis</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Odds Ratio (95% CI)</td>
<td>P Value</td>
</tr>
<tr>
<td>Male</td>
<td>5.95 (2.19-16.19)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>White</td>
<td>0.77 (0.29-2.04)</td>
<td>0.60</td>
</tr>
<tr>
<td>Age at death</td>
<td>0.97 (0.95-0.99)</td>
<td>0.01</td>
</tr>
<tr>
<td>Body weight*</td>
<td>1.00 (0.90-1.11)</td>
<td>0.99</td>
</tr>
<tr>
<td>Heart weight†</td>
<td>1.02 (0.99-1.05)</td>
<td>0.16</td>
</tr>
<tr>
<td>Competitive Athlete</td>
<td>20.30 (6.84-60.34)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Left ventricular involvement</td>
<td>1.01 (0.43-2.35)</td>
<td>0.99</td>
</tr>
</tbody>
</table>

CI indicates confidence interval.
* per 5kg increase in Weight
† per 10g increase in Heart Weight
Table 3. Overview of pathogenic variants identified from post-mortem genetic testing

<table>
<thead>
<tr>
<th>Case ID</th>
<th>Age at death</th>
<th>Circumstances of death</th>
<th>Ante-mortem diagnosis</th>
<th>Pathology</th>
<th>Gene</th>
<th>cDNA change</th>
<th>ExAC MAF</th>
<th>ACMG</th>
</tr>
</thead>
<tbody>
<tr>
<td>Case 1 White Male</td>
<td>37</td>
<td>Died in sleep</td>
<td>ACM</td>
<td>Right ventricular ACM</td>
<td>PKP2</td>
<td>c.2198_2202delAC ACC</td>
<td>0</td>
<td>Pathogenic PVS1; PM2; PP3</td>
</tr>
<tr>
<td>Case 2 White Male</td>
<td>20</td>
<td>Died during exertion</td>
<td>None</td>
<td>Right ventricular ACM</td>
<td>PKP2</td>
<td>c.253_256delGAGT</td>
<td>0</td>
<td>Pathogenic PVS1; PM2; PP3</td>
</tr>
<tr>
<td>Case 3 White Female</td>
<td>26</td>
<td>Died at rest</td>
<td>DCM</td>
<td>Biventricular ACM</td>
<td>DSP</td>
<td>c.4395T&gt;G</td>
<td>0</td>
<td>Pathogenic PVS1; PM2; PP3</td>
</tr>
<tr>
<td>Case 4 White Male</td>
<td>28</td>
<td>Died at rest</td>
<td>DCM</td>
<td>Biventricular ACM</td>
<td>DSP</td>
<td>c.5269C&gt;T</td>
<td>0</td>
<td>Pathogenic PVS1; PM2; PP3</td>
</tr>
<tr>
<td>Case 5 White Male Athlete</td>
<td>20</td>
<td>Died during exertion</td>
<td>None</td>
<td>Biventricular ACM</td>
<td>TMEM43</td>
<td>c.1073C&gt;T</td>
<td>0</td>
<td>Pathogenic PS1; PM2; PM7; PP3</td>
</tr>
<tr>
<td>Case 6 White Male</td>
<td>45</td>
<td>Died at rest</td>
<td>None</td>
<td>Biventricular ACM</td>
<td>TMEM43</td>
<td>c.1073C&gt;T</td>
<td>0</td>
<td>Pathogenic PS1; PM2; PM7; PP3</td>
</tr>
</tbody>
</table>

ACM: Arrhythmogenic cardiomyopathy; DCM: Dilated cardiomyopathy; cDNA: Complementary DNA; ExAC: Exome Aggregation Consortium; MAF: Minor allele frequency; ACMG: American College of Medical Genetics and Genomics; PKP2: Plakophilin-2; DSP: Desmoplakin; TMEM43: Transmembrane protein 43; PVS: Pathogenic very strong; PS: Pathogenic strong; PM: Pathogenic moderate; PP: Pathogenic supporting.
Table 4. Clinico-pathological correlations among ACM decedents diagnosed with cardiomyopathy ante-mortem

<table>
<thead>
<tr>
<th></th>
<th></th>
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<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>A</td>
<td>ACM</td>
<td>37</td>
<td>Died at rest</td>
<td>Minor criterion: SAECG – 3 abnormal parameters</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Major criterion: Negative T waves in V1, V2, V3, and V4</td>
<td></td>
<td></td>
<td></td>
<td>BiV ACM</td>
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<tr>
<td>B</td>
<td>ACM (ICD in situ)</td>
<td>44</td>
<td>Died at rest</td>
<td>None</td>
<td></td>
<td></td>
<td></td>
<td>BiV ACM</td>
</tr>
<tr>
<td>C</td>
<td>ACM</td>
<td>37</td>
<td>Died in sleep</td>
<td>None</td>
<td></td>
<td></td>
<td></td>
<td>RV ACM PKP2†</td>
</tr>
<tr>
<td>D</td>
<td>DCM</td>
<td>49</td>
<td>Died during exertion</td>
<td>Minor criterion: Terminal activation duration of QRS ≥ 55ms in V1</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>Major criterion: Negative T waves in V4, V5, and V6</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>E</td>
<td>DCM</td>
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<td>Died at rest</td>
<td>None</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>F</td>
<td>DCM (ICD in situ)</td>
<td>48</td>
<td>Died at rest</td>
<td>None</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>G</td>
<td>DCM (ICD in situ)</td>
<td>35</td>
<td>Died at rest</td>
<td>None</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>H</td>
<td>DCM (ICD in situ)</td>
<td>26</td>
<td>Died at rest</td>
<td>None</td>
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<td></td>
<td></td>
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<tr>
<td>I</td>
<td>DCM (ICD in situ)</td>
<td>41</td>
<td>Died at rest</td>
<td>None</td>
<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td>J</td>
<td>DCM</td>
<td>56</td>
<td>Died in sleep</td>
<td>None</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

DCM: Dilated cardiomyopathy; ACM: Arrhythmogenic cardiomyopathy; TFC: Task Force Criteria (2010) (6); CMR: Cardiovascular Magnetic Resonance; Echo: Echocardiography; NSVT: Non-sustained ventricular tachycardia; PVCs: Premature ventricular contractions; LBBB: Left bundle branch block; BiV: Biventricular; ICD: Implantable cardioverter defibrillator; SAECG: Signal-averaged ECG; RV: Right ventricular; PKP2: Plakophilin-2; LV: Left ventricular; DSP: Desmoplakin; A-V: Atrial and ventricular; V: Ventricular.

* Task Force Criteria attained from different categories
† Pathogenic variant detected in gene following post-mortem genetic testing
Figure Legends

Figure 1. Selection of ACM cases from all SCD referred to pathology center.

Figure 2. Histological examination of ACM: Normal findings within the RV versus pathological criteria for ACM.

Figure 3. Ventricular disease involvement among all ACM decedents (n=202)(left). Distribution of fibrofatty infiltration among whole hearts referred to pathology center (n=120)(right). RV: Right ventricle; LV: Left ventricle; RVAL: RV anterolateral wall; RVOT: RV outflow tract; RVS: RV septum; RVPW: RV posterior wall; LVS: LV septum; LVAL: LV anterolateral wall; LVPW: LV posterior wall.

Figure 4. A case of LV-dominant ACM. A: Histological slide (Picrosirius red stain) demonstrating myocyte degeneration and fibrofatty infiltration within the posterolateral wall of the LV (extending transmurally). B: 12-lead ECG showing first degree AV-block, inferolateral T-wave inversion (arrows) and low voltage limb lead QRS complexes, prolonged terminal activation duration in V1, and ventricular bigeminy with fragmented, broad, ectopics of RBBB morphology and superior axis (arrow). C+D: Delayed-enhancement CMR images illustrating extensive LV delayed enhancement, including near transmural enhancement of the lateral wall and mid-wall enhancement of the anterior wall. E: Parasternal long axis view (echocardiography) showing severe LV dilatation (left ventricular end-diastolic dimension 71mm).
5205 SCD cases (1994-2018)

Pathological diagnosis of ACM n=202

Positive toxicology n= 243

Exclude Post-mortem genetic testing N=24 (12%)

Whole hearts n=120 (59%)

Tissue blocks n=82 (41%)

Competitive athlete 41/202 (20%)

Aged <35 at death n=101 (50%)

Aged ≥ 35 at death n=101 (50%)
Normal RV fatty infiltration

Isolated areas of fibrosis between trabeculae without intermixed adipocytes

Non-epicardial adipocytes and perivascular collagen within right ventricle

Normal / Non-Diagnostic

PATHOLOGICAL

Myocyte degeneration
Fat & fibrosis (co-located)
Subepicardial → Transmural
≥ 2 tissue blocks

≥20% Tissue Area
Biventricular Involvement
141/202 (70%)

RV: 26/202 (13%)
LV: 35/202 (17%)

RVAL 58%
RVS 32%
RVPW 43%
RVOT 48%

LVAL 58%
LVS 32%
LVPW 68%