

Differences between familial and sporadic dilated cardiomyopathy: ESC EORP Cardiomyopathy & Myocarditis registry

Folkert W. Asselbergs^{1,2*} , Arjan Sammani¹ , Perry Elliott^{3†} , Juan R. Gimeno^{4‡} , Luigi Tavazzi⁵, Michael Tendera⁶, Juan Pablo Kaski^{7‡} , Aldo P. Maggioni^{5,8} , Paweł P. Rubis⁹ , Ruxandra Jurcut^{10‡}, Tiina Heliö^{11‡} , Leonardo Calò¹², Gianfranco Sinagra¹³, Marija Zdravkovic¹⁴, Iacopo Olivotto¹⁵, Aušra Kavoliūnienė¹⁶, Cécile Laroche⁸, Alida L.P. Caforio^{17‡}, Philippe Charron¹⁸  Cardiomyopathy & Myocarditis Registry Investigators Group[†]

¹Department of Cardiology, University Medical Centre Utrecht, University of Utrecht, Heidelberglaan 100, Utrecht, 3584CX, The Netherlands; ²Institute of Cardiovascular Science and Institute of Health Informatics, Faculty of Population Health Sciences, University College London, London, UK; ³Barts Heart Centre, St Bartholomew's Hospital, University College London and Inherited Cardiac Diseases Unit, London, UK; ⁴Cardiac Department, Hospital Universitario Virgen de la Arrixaca, Murcia, Spain; ⁵GVM Care & Research, Maria Cecilia Hospital, Cotignola, Italy; ⁶Department of Cardiology and Structural Heart Diseases, School of Medicine in Katowice, Medical University of Silesia, Katowice, Poland; ⁷Centre for Inherited Cardiovascular Diseases, Great Ormond Street Hospital, UK and University College London Institute of Cardiovascular Science, London, UK; ⁸EUR Observational Research Programme, European Society of Cardiology, Sophia-Antipolis, France; ⁹Department of Cardiac and Vascular Diseases, Jagiellonian University Medical College, John Paul II Hospital, Krakow, Poland; ¹⁰Department of Cardiology, Emergency Institute of Cardiovascular Diseases C.C. Iliescu, Bucharest, Romania; ¹¹Department of Cardiology, Helsinki University Central Hospital Meilahti, Helsinki, Finland; ¹²U.O. Cardiologia, Policlinico Casilino, Rome, Italy; ¹³Cardiovascular Department, Azienda Sanitaria Universitaria Integrata Giuliano Isontina, Trieste, Italy; ¹⁴Clinical Hospital Center Bezanijska kosa, Faculty of Medicine, University of Belgrade, Beograd, Serbia; ¹⁵Cardiomyopathy Unit, Careggi University Hospital, Florence, Italy; ¹⁶Department of Cardiology, Lithuanian University of Health Sciences, Kaunas, Lithuania; ¹⁷Division of Cardiology, Department of Cardiological Thoracic and Vascular Sciences and Public Health, University of Padova, Padova, Italy; ¹⁸APHP, Centre de Référence des Maladies Cardiaques Héréditaires, ICAN, Hôpital Pitié-Salpêtrière, Sorbonne Université, Paris, France

Abstract

Aims Dilated cardiomyopathy (DCM) is a complex disease where genetics interplay with extrinsic factors. This study aims to compare the phenotype, management, and outcome of familial DCM (FDCM) and non-familial (sporadic) DCM (SDCM) across Europe.

Methods and results Patients with DCM that were enrolled in the prospective ESC EORP Cardiomyopathy & Myocarditis Registry were included. Baseline characteristics, genetic testing, genetic yield, and outcome were analysed comparing FDCM and SDCM; 1260 adult patients were studied (238 FDCM, 707 SDCM, and 315 not disclosed). Patients with FDCM were younger ($P < 0.01$), had less severe disease phenotype at presentation ($P < 0.02$), more favourable baseline cardiovascular risk profiles ($P \leq 0.007$), and less medication use ($P \leq 0.042$). Outcome at 1 year was similar and predicted by NYHA class (HR 0.45; 95% CI [0.25–0.81]) and LVEF per % decrease (HR 1.05; 95% CI [1.02–1.08]). Throughout Europe, patients with FDCM received more genetic testing (47% vs. 8%, $P < 0.01$) and had higher genetic yield (55% vs. 22%, $P < 0.01$).

Conclusions We observed that FDCM and SDCM have significant differences at baseline but similar short-term prognosis. Whether modification of associated cardiovascular risk factors provide opportunities for treatment remains to be investigated. Our results also show a prevalent role of genetics in FDCM and a non-marginal yield in SDCM although genetic testing is largely neglected in SDCM. Limited genetic testing and heterogeneity in panels provides a scaffold for improvement of guideline adherence.

Keywords Dilated cardiomyopathy; Sporadic; Familial; Genetic; Prognosis; Europe

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*Correspondence to: Folkert W. Asselbergs, Department of Cardiology, University Medical Centre Utrecht, University of Utrecht, Heidelberglaan 100, 3584CX Utrecht, The Netherlands. Tel: +31 88 75 570 65. Email: f.w.asselbergs@umcutrecht.nl

†Listed in the Appendix A.

‡European Reference Network for rare, low prevalence, and complex diseases of the heart (ERN GUARD-Heart).

Introduction

Dilated cardiomyopathy (DCM), defined as left ventricular dilation and systolic left ventricular (LV) impairment unexplained by coronary artery disease or abnormal loading conditions, is a leading cause of heart failure and heart transplantation with an estimated prevalence of ~36 cases per 100 000 in Europe.¹

Prior studies on clinical characteristics of patients with non-familial (sporadic) forms of DCM (SDCM) and familial DCM (FDCM) have reported that FDCM presents at earlier age but presented conflicting evidence regarding phenotype severity. Some studies found more favourable clinical profiles in FDCM compared with SDCM, whereas others report similar baseline phenotypes without any distinctive features.^{2–5} Due to genotype–phenotype associations, such as frequent ventricular arrhythmias in *LMNA* and *PLN* mutation carriers, heterogeneity in prognosis however may be expected between FDCM and SDCM.⁶ Others postulated that FDCM presents as SDCM and that proper active family screening is needed to distinguish true SDCM from undetected FDCM. This screening has been shown to effectively identify DCM and improve prognosis.³

Many studies have shown that DCM can be inherited as a genetic trait^{1,7–11} and that structural or functional LV abnormalities are present in 20% of asymptomatic relatives of patients with DCM.^{12,13} Genetic mutations are reported in more than a third of index patients with FDCM and in 8–25% of SDCM.^{8,14} The most common disease causing variants are found in genes coding for sarcomere proteins such as *TTN*, *MYH7*, and *FLNC*, and the nuclear envelope gene *LMNA*.^{15–17} Evidence also suggests that genetic predisposition in DCM may interact with extrinsic disease triggers such as toxin exposure (ethanol, chemotherapy, and cocaine), viral infection, and pregnancy.^{1,18,19} Nonetheless, toxin exposure may also be the sole putative trigger of DCM, for instance in alcoholic cardiomyopathy.¹⁸

The ESC EORP Cardiomyopathy & Myocarditis Registry is a prospective observational multinational survey of consecutive patients with cardiomyopathies.²⁰ To investigate the complexity in clinical characteristics and genetic yield, the current analysis of this registry aims to (i) study clinical cardiovascular differences in adult FDCM and SDCM, (ii) report the frequency of genetic testing across Europe, and (iii) report differences in genetic yield between familial and sporadic DCM.

Methods

The general policy as well as baseline results of the new EURObservational Research Programme (EORP) cardiomyopathy registry of the ESC have been previously published. In

short, the EORP cardiomyopathy registry is a multicentre registry where participating centres were asked to enter baseline, follow-up, and genetic data of about 40 consecutive patients with cardiomyopathy per centre.²⁰ Patients were included from 1 December 2012 until 30 December 2016.

Inclusion/exclusion criteria

Adult patients with DCM defined by ESC consensus criteria were studied. Specifically, (i) left ventricular ejection fraction <45% (>2 SD) and/or fractional shortening <25% (>2 SD), as ascertained by echocardiography, radionuclide scanning, or cardiac magnetic resonance imaging; and (ii) left ventricular end-diastolic diameter $>117\%$ of the predicted value corrected for age and body surface area (Henry's formula), which corresponds to 2 SD of the predicted normal limit +5%. Patients with heart failure attributable to coronary artery disease and clinically suspected or biopsy-proven myocarditis were excluded.^{20,21}

Definitions

Familial dilated cardiomyopathy (FDCM) was defined by the presence of two or more affected individuals in a single family or the presence of an index patient with DCM and a first degree relative with documented unexplained sudden cardiac death at <35 years of age. Patients that did not meet these criteria were deemed sporadic DCM (SDCM). Patients with missing data concerning familial status ($n = 315$) were compared with both FDCM and SDCM for clinical differences to account for bias. Primary outcome was defined as a composite of cardiovascular death, implantation of a ventricular assist device, or heart transplantation. Secondary endpoint was hospitalization for urgent cardiac reason. Genetic testing and variant classification were planned and performed according to clinician's judgement. Genetic variants and their classifications were reported by individual researchers representing their centres. Because lab techniques and genetic coordinates were not recorded, centralized variant classification was not possible. The definitions of included variables have been listed in prior EORP publications and are included in the Supporting Information, *Data S1*.^{20,22}

Statistical analysis

Univariable analysis was applied to both continuous and categorical variables. Continuous variables were reported as mean \pm standard deviation and/or as median and interquartile range (IQR) when appropriate. Among-group comparisons were made using a non-parametric test (Kruskal–Wallis). Categorical variables were reported as counts and percentages. Among-group comparisons were

made using a χ^2 test or a Fisher's exact test if any expected cell count was less than 5. Plots of Kaplan–Meier curves for primary outcome were performed. Cox proportional hazards model was used for survival estimates reporting hazard ratios (HRs) and 95% confidence intervals (95% CIs) in univariable and multivariable analysis. As the goal was to report covariates and their association to outcome rather than a clinical risk calculator, both multivariable results and variable selection ($P < 0.05$) were reported. To compare with external datasets, comparison of proportions was calculated using the "N-1" χ^2 test. A two-sided P -value of <0.05 was considered as statistically significant. For sensitivity analyses, probands of FDCM were compared with SDCM. Analyses were performed using SAS statistical software version 9.4 (SAS Institute, Inc., Cary, NC, USA).

Results

The cohort comprised 1260 patients, of whom 238 had FDCM, 707 SDCM, and 315 were unclassified (unknown). Patients with unknown status were compared with FDCM and SDCM. The analysis revealed the 'unknown' group to be similar to SDCM. These results and head to head group comparisons are available in the Supporting Information, *Data S2*.

The characteristics and treatment of patients with FDCM and SDCM are reported in *Table 1*. Compared with SDCM, patients with FDCM were younger (44 years [IQR 31–52] vs. 51 years [IQR 41–58], $P < 0.001$), had lower NYHA class ($P < 0.001$) and BNP and NT-proBNP levels ($P < 0.028$), less frequent left bundle branch block (9% vs. 22%, $P < 0.001$), smaller left ventricular end diastolic diameter ($P < 0.001$), and higher left ventricular ejection fraction (LVEF) (37% vs. 31%, $P < 0.001$). Sixty per cent of FDCM patients were index cases compared with 99% of SDCM patients ($P < 0.001$). Patients with SDCM had a higher burden of cardiovascular risk factors (hypertension, dyslipidaemia, diabetes, smoking, alcohol intake, and high BMI) than FDCM (all $P \leq 0.007$). A larger proportion of patients with SDCM received beta-blockers, diuretics, mineralocorticoid receptor antagonists, and other anti-arrhythmic agents ($P \leq 0.042$) compared with FDCM. A sensitivity analysis comparing solely FDCM index patients with SDCM showed that all characteristics except BMI remained significantly different between the groups (Supporting Information, *Data S2*, *Table 1.1A*).

Follow-up data were available in 1105 (88%) cases (median follow-up duration: 372 days; interquartile range [IQR] 363–428). There were no differences observed in primary and secondary outcomes or all-cause mortality when comparing FDCM ($n = 209$) to SDCM ($n = 611$) (Supporting Information, *Data S2*, *Table 3B*). Age at first primary event was also similar for FDCM and SDCM (*Figure 1*).

In multivariable analysis, BMI per unit decrease (HR 1.11; 95% CI [1.02–1.22]) and LVEF per % decrease (HR 1.08; 95% CI [1.03–1.11]) were associated with primary and secondary outcomes. After stepwise selection, NYHA class I/II versus III/IV (HR 0.45; 95% CI [0.25–0.81]) and LVEF per % decrease (HR 1.05; 95% CI [1.02–1.08]) were predictive (*Table 2*). There were no significant associations for all-cause mortality. Results of all analyses are available in the Supporting Information, *Data S2*, *Table 10*.

Genetic testing

In total, 214 out of the 1260 (17%) cases were genetically tested (58 out of 707 [8%] SDCM and 114 out of 238 [48%] of FDCM). In 110 cases, the information on genetic testing was missing. Genetic testing was less frequently performed in Eastern Europe and North Africa (respectively 4% and 0%), whereas North, South, and West Europe performed genetic testing in 21–27% of their population (*Table 3*).

In 63 out of the 114 (55%) tested cases of FDCM, at least one disease causing variant was reported compared with 13 out of 58 (22%) in SDCM. These variants were most prevalent in sarcomere ($n = 42$) and nuclear genes ($n = 27$), with most variants being discovered in *LMNA* (16% yield in FDCM and 3% yield in SDCM) and *MYH7* (14% yield in FDCM and 0% in SDCM) (Supporting Information, *Data S2*, *Table 7*).

Discussion

The three main findings of this study are (i) patients with FDCM present at younger age with a less severe phenotype and lower burden of cardiovascular risk factors but similar short-term prognosis to SDCM; (ii) patients with FDCM are genetically tested more frequently and with higher diagnostic yield than SDCM and; (iii) there are important differences in the use of genetic testing across European centres enrolled in the ESC cardiomyopathy registry. For a central illustration, these findings are summarized in *Figure 2*.

Disease burden

Overall, DCM patients included in this cohort had similar demographic and clinical characteristics compared with DCM populations described in the literature.^{2,4,5,12,23} However, in contrast to some studies, we found that patients with FDCM were diagnosed at a younger age and have a less severe phenotype than patients with SDCM but have similar cardiovascular prognosis.^{2,5} This was also seen in the prescribed medication, where we observed that patients with FDCM received less beta-blockers, diuretics and mineralocorticoid receptor antagonists. These findings

Table 1 Baseline table of patient characteristics, pharmacotherapy and outcome

	All (N = 1260)	FDCM (n = 238)	SDCM (n = 707)	P-value ^a
Age at diagnosis (years), median (IQR)	49 (40–58)	44 (31–52)	51 (41–58)	<0.01
Male				<0.05
NYHA class				<0.01
I	935 (74%)	165 (69%)	536 (76%)	
II	198 (19%)	65 (35%)	84 (14%)	
III	448 (43%)	79 (42%)	261 (44%)	
IV	316 (30%)	37 (20%)	187 (31%)	
Family history of SCD	87 (8%)	6 (3%)	65 (11%)	<0.01
Hypertension	479 (38%)	54 (23%)	288 (41%)	<0.01
Dyslipidaemia	472 (38%)	62 (26%)	274 (39%)	<0.01
Diabetes Mellitus	211 (17%)	25 (11%)	127 (18%)	<0.01
Alcohol use ≥ 1 units/day	174 (16%)	20 (10%)	124 (20%)	<0.01
Smoking (current and former)	507 (42%)	71 (31%)	323 (47%)	<0.01
Renal impairment	172 (14%)	24 (10%)	94 (13%)	0.20
History of Atrial Fibrillation	356 (28%)	63 (27%)	202 (29%)	0.53
History of Stroke	87 (7%)	12 (5%)	47 (7%)	0.38
History of resuscitation	61 (5%)	15 (6%)	32 (5%)	0.28
Atrioventricular block				0.02
1st	108 (9%)	20 (9%)	57 (8%)	
2nd	6 (1%)	3 (1%)	2 (0%)	
3rd	14 (1%)	6 (3%)	4 (1%)	
QRS Duration (ms), median, IQR	105 (92–130)	100 (90–112)	104 (90–130)	0.02
LBBB	219 (21%)	18 (9%)	133 (22%)	<0.01
RBBB	41 (4%)	6 (3%)	24 (4%)	0.53
LVEF (%), median (IQR)	31 (25–40)	37 (29–45)	31 (24–40)	<0.01
LVEDD (mm), median (IQR)	64 (58–70)	60 (54–67)	64 (59–70)	<0.01
Diastolic dysfunction (grades)				<0.01
Normal	63 (38%)	97 (20%)		
I (impaired relaxation)	57 (35%)	169 (35%)		
II (pseudonormal)	28 (17%)	108 (22%)		
III/IV (restrictive)	16 (9%)	109 (22%)		
Haemoglobin (g/dL), median (IQR)	14 (13–15)	14 (13–15)	14 (13–15)	0.23
BNP (pg/mL), median (IQR)	297 (72–717)	108 (36–555)	404 (106–718)	0.03
NT-proBNP (pg/mL), median (IQR)	1102 (312–3341)	589 (156–1663)	1276 (452–3527)	<0.01
Performed	259 (21%)	66 (28%)	148 (21%)	0.04
Abnormal	237 (19%)	57 (24%)	136 (19%)	0.04
Late gadolinium enhancement	153 (64%)	41 (65%)	92 (67%)	0.83
β-blockers	1130 (90%)	203 (85%)	644 (91%)	0.01
Diuretics	895 (72%)	123 (54%)	540 (77%)	<0.01
ACE-inhibitors or ATII-receptor blockers	1121 (89%)	208 (88%)	634 (89%)	0.329
Mineralocorticoid receptor antagonists	795 (63%)	119 (50%)	470 (67%)	<0.01
Other antiarrhythmics	361 (29%)	56 (24%)	215 (30%)	0.04

BMI, body mass index; LBBB, left bundle branch block; LVEDD, left ventricular end diastolic diameter; LVEF, left ventricular ejection fraction; NYHA, New York Heart Association; RBBB, right bundle branch block; SCD, sudden cardiac death.

Data are presented as number and percentages of valid. Patients with missing family status were not included in the subgroup SDCM in this table because their status was unknown. They have been included in the *Supporting Information, Data S2, Table 1* both as a separate group and combined with SDCM. Continuous data are presented as medians. Median presented with first and third interquartiles (IQR). Mean presented with ±standard deviation (SD).

^aFDCM versus SDCM.

may reflect of cascade family screening in patients with suspected FDCM. However, when comparing FDCM index patients with SDCM patients, the clinical differences remained significant. Because family screening has shown to effectively identify DCM patients at earlier stages of disease, potentially benefiting prognosis, genetic counselling remains an important pillar of care in patients with DCM and their relatives.³ Prognosis of DCM has been improving for the endpoint of cardiovascular death or heart transplantation/assist device implantation, arguably because of better cardiovascular care and the early identification and classification of disease.²⁴ Our observations however are more in line with earlier reports and less favourable

outcome, which may be caused because of a selection bias towards more expert/tertiary centres.²³

A relevant finding of this study was that patients with SDCM had a higher burden of cardiovascular risk factors, such as hypertension, hypercholesterolaemia, and smoking. Because it is well known that these risk factors are more prevalent with increasing age, and our patients with SDCM were significantly older than FDCM, this result might be expected.²⁵ However, the age difference between SDCM and FDCM does not fully account for the observed increase in risk factors. When comparing our patients with SDCM with external cohorts from the same age group, we still observe a significantly increased burden of cardiovascular factors

Table 2 Multivariable analysis for primary outcome

Variable	Pval hazard ratio	Hazard ratio (95% CI)
Age at diagnosis per year increase	0.040 (S)	0.973 (0.949; 0.999)
LBBB	0.635 (NS)	0.804 (0.327; 1.977)
NYHA class I or II versus III or IV	0.061 (NS)	0.483 (0.226; 1.035)
Alcohol intake 1 unit per day or more	0.618 (NS)	1.224 (0.553; 2.705)
Body mass index (kg/m^2) per unit increase	0.012 (S)	0.896 (0.823; 0.977)
Diabetes mellitus	0.029 (S)	2.815 (1.111; 7.131)
Hypertension	0.859 (NS)	1.070 (0.509; 2.251)
Smoking current or former	0.921 (NS)	0.964 (0.472; 1.970)
LVEF per % decrease	<0.001 (S)	1.073 (1.031; 1.117)

LBBB, left bundle branch block; LV, left ventricular ejection fraction; NS, not significant; NYHA, New York Heart Association; S, significant. Primary endpoint was defined as a composite of cardiovascular death, implantation of a ventricular assist device, or heart transplantation. Cox proportional hazards model is presented (hazard ratio's and 95% confidence intervals).

($P < 0.01$) in our patients with SDCM.^{25,26} In addition, the presence of common cardiovascular risk factors is also associated with worse outcome in SDCM. Smoking, for instance, independently increased sudden cardiac death in SDCM²⁷ and diabetes confers a two-fold to five-fold added risk for heart failure development, even after adjustment for other traditional risk factors such as coronary heart disease.²² Alcohol is also known to be the both sole cause of cardiomyopathy and may negatively affect cardiac function in mutation carriers. In a recent report including 716 DCM cases, excessive alcohol consumption resulted in an absolute LVEF reduction of $\pm 9\%$ in patients with a truncating *TTN* variant.¹⁸ These results support the hypothesis that cardiovascular risk factors and external toxic substances interplay with genetic frailty, (auto)immunity, or endocrinological disorders in the DCM phenotype.¹ Whether modification of cardiovascular risk factors provide opportunities for treatment remains to be investigated.

Differences in prognosis between FDCM and SDCM remain controversial with most follow-up studies of DCM not dividing populations in familial and non-familial categories.^{3,28,29} Conflicting results of prior studies may be due to the absence of reliable clinical or morphological parameters to differentiate between FDCM and SDCM.^{1,3,28,29} Genotype–phenotype associations affecting prognosis in DCM have been reported: mutations in *LMNA* and *RBM20* predispose for more/younger heart transplantsations.⁶ Mutations in *FLNC*, *LMNA*, and *PLN* have been associated with sustained ventricular arrhythmias in DCM.³⁰ Nonetheless, our analysis did not yield a significant difference in outcome between FDCM and SDCM.

Genetic mutations in dilated cardiomyopathy

Mutations in over 40 genes are causally related to DCM and explain up to 61% of cases in FDCM and up to 25% in SDCM.^{1,8,14,16,31} Our data confirm the higher yield in FDCM.⁸ At present, genetic screening is advised in patients with family history of DCM or a personal history of atypical features such as conduction/rhythm-disturbances,^{7,8,11,31} and yet less than 50% of all cases in the registry were tested.

Moreover, the fact that heterogeneous screening strategies were used may have impacted on the yield of testing.²⁰ For instance, mutations in Titin (*TTN*) have been implicated in up to 22% of SDCM, but this gene was only tested in 19% of our patients with SDCM.^{32,33} The selective nature of genetic testing may explain the high prevalence of mutations in genes coding for sarcomere genes and *LMNA*.

Genetic testing in Europe

We observed differences in genetic testing between European centres. Genetic testing was most prevalent in Northern, Southern, and Western Europe (around 20% of the cohort) and least common in Eastern Europe and North Africa. Importantly, Southern Europe included a majority (54%) of the patients with FDCM in this cohort, which could have led to sampling bias.^{34,35} These shortcomings should be accounted for in terms of external validity of this study. Furthermore, the regional differences in our study as well as differences described by prior EORP studies provide opportunities to improve guideline adherence in Europe.^{20,35,36} Even though guideline adherence is highest in secondary and tertiary centres and higher for cardiologists than other specialities, care may be fragmented in highly complex diseases such as inherited cardiomyopathies.³⁷ This fragmentation can cause unwanted protocol deviations that may deteriorate quality of care. Several strategies have been suggested to improve guideline adherence. These strategies include common care pathways, ongoing education, and focus groups.³⁷ In terms of genetic testing, actively looking for familial manifestations of disease in patients and relative and acquiring a detailed family history are quintessential and may improve genetic yield dramatically.

Study limitations

In contrast to controlled clinical studies, our data are likely to be heterogeneous given the nature of the study design. As the structure of the registry requested to introduce patients

Figure 1 Survival probability curves for primary outcome plotted over both time and age comparing SDCM to FDCM.

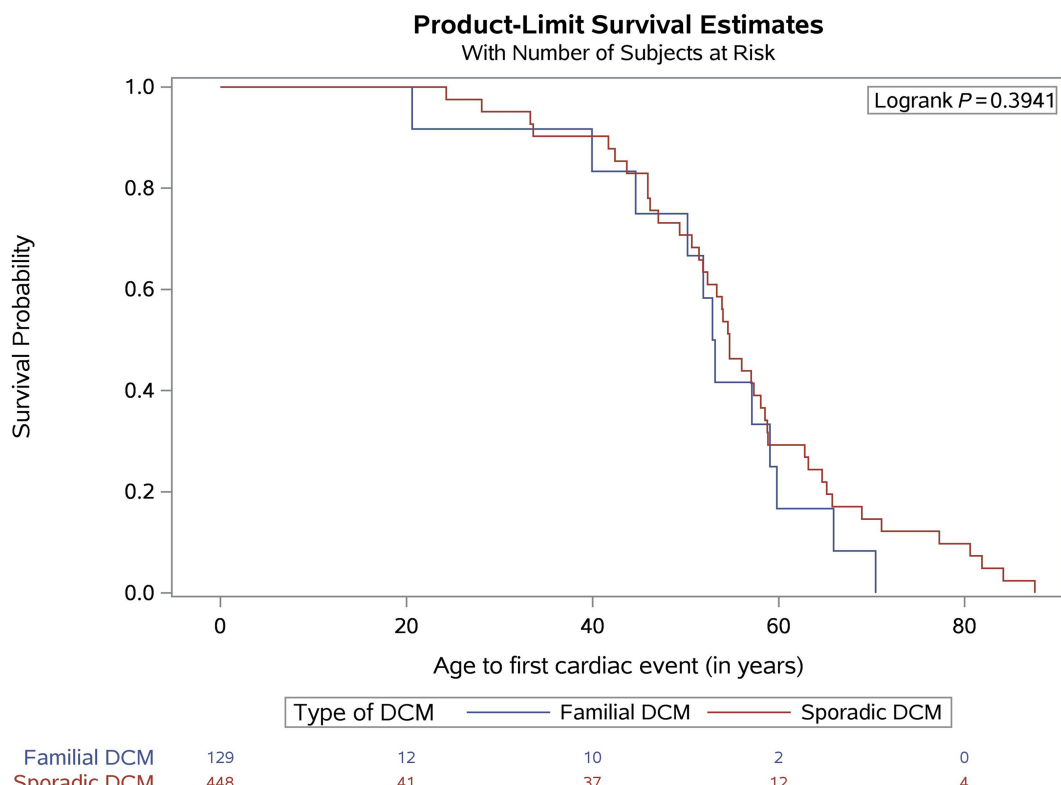
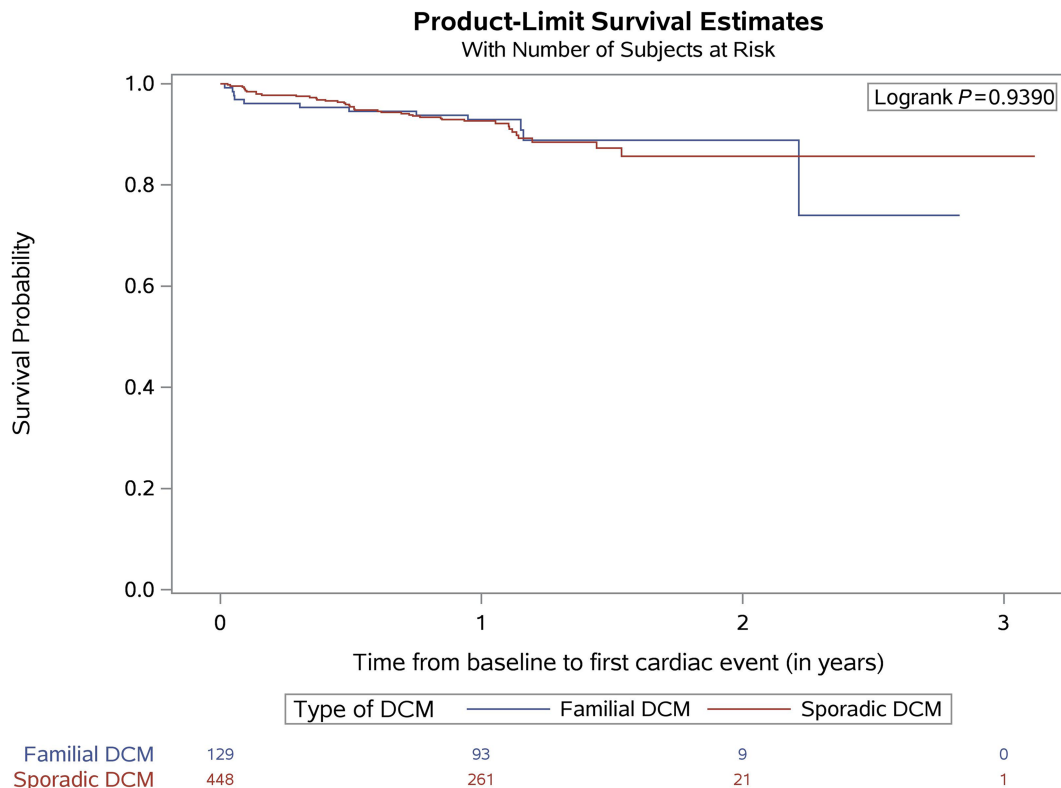
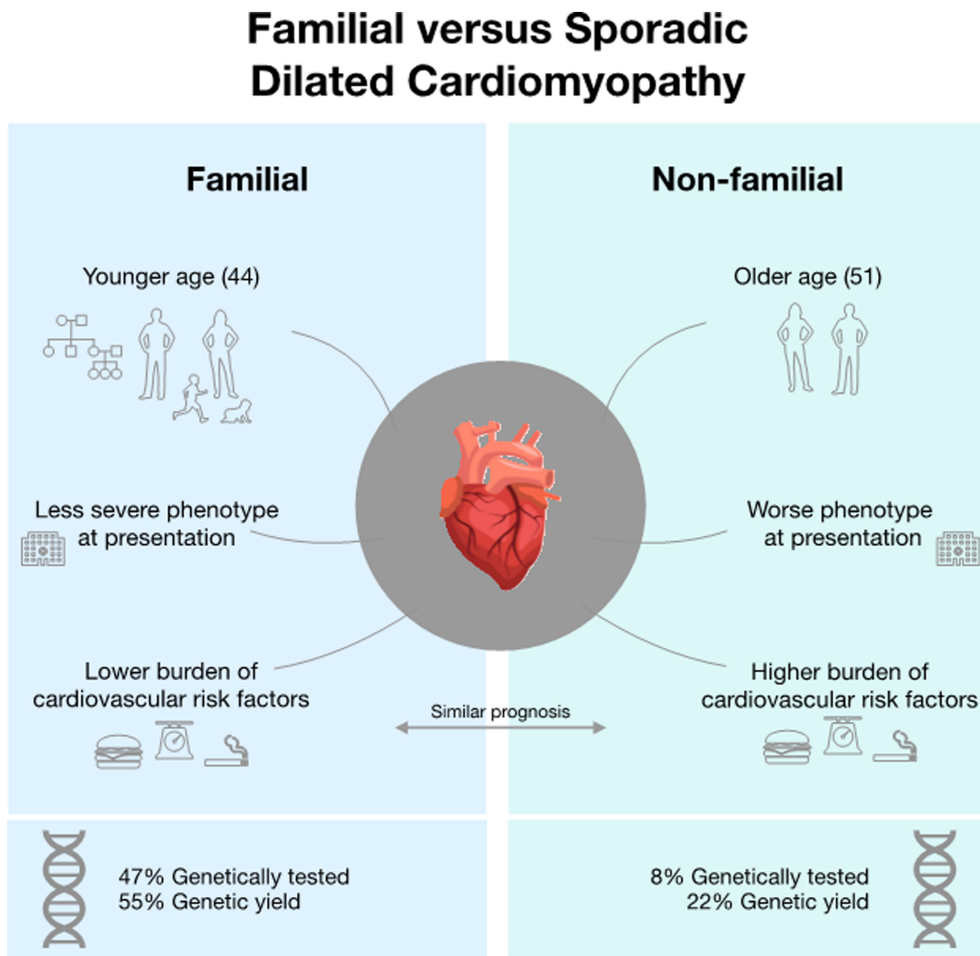


Table 3 Regional differences for genetic testing in DCM

	All (N = 1260)	FDCM (N = 238)	SDCM (N = 707)	P-value ^a
North Europe (n = 179)	37 (21%)	19 (56%)	11 (16%)	<0.001
South Europe (n = 425)	116 (27%)	74 (59%)	23 (14%)	<0.001
West Europe (n = 206)	48 (23%)	15 (56%)	24 (16%)	<0.001
East Europe (n = 302)	13 (4%)	6 (18%)	0 (0%)	<0.001
North Africa (n = 38)	0 (0%)	0 (0%)	0 (0%)	NC
Total (1150)	214 (17%)	114 (48%)	58 (8%)	<0.001

NC, not calculable.

In 110 cases, genetic testing was unknown.

^aFDCM versus SDCM.**Figure 2** Summary of the main findings of this study.

from expert/tertiary centres, this may have caused a selection bias. Second limitation is that not all genes were tested in all patients, and centralized variant classification was not repeated in this study. Given that genetic testing was unknown in 110 patients, this might constitute another limitation. Even though myocarditis is formally an exclusion criterion, inflammation-mediated reasons for DCM are

very difficult to diagnose. Therefore, we cannot exclude with certainty that a proportion of non-familial DCM might be suspected for inflammatory diseases. As the EORP study was not designed for inter-regional comparisons, and these analyses were not planned, such comparisons may be arbitrary and unrepresentative. Furthermore, sub-analyses using ethnicity/race to provide insight into heterogeneity of

populations are limited by poor accuracies of ethnicity/race in electronic health records. Importantly, even though we performed sensitivity analyses with and without patients with unknown classification, this still might have led to biased results. Lastly, the fact that no difference in outcome was detected in our DCM populations may be due to sample size and requires confirmation in larger studies.

Conclusions

Familial DCM and SDCM have significant differences at baseline. Patients with FDCM appear to present at earlier stage of disease. Patients with SDCM in this registry have less favourable clinical profile. Whether modification of associated cardiovascular risk factors provide opportunities for treatment remains to be investigated. Genetic testing confirms a prevalent role of genetics in FDCM but is largely neglected in SDCM. Although SDCM may probably be multifactorial, genetic influences need further understanding. Limited genetic testing provides a scaffold for improvement of guideline adherence.

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Conflict of interest

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Supporting information

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

Data S1. Centre Inclusion Criteria.

Data S2. Supporting information.

Appendix A.

EORP Oversight Committee

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