ORIGINAL ARTICLE

Role of late gadolinium enhancement cardiovascular magnetic resonance in the risk stratification of hypertrophic cardiomyopathy

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ABSTRACT

Objective Myocardial fibrosis identified by late gadolinium enhancement (LGE) cardiovascular magnetic resonance (CMR) in patients with hypertrophic cardiomyopathy (HCM) is associated with adverse cardiovascular events, but its value as an independent risk factor for sudden cardiac death (SCD) is unknown. We investigated the role of LGE-CMR in the risk stratification of HCM.

Methods We conducted a prospective cohort study in a tertiary referral centre. Consecutive patients with HCM (n=711, median age 56.3 years, IQR 46.7–66.6; 70.0% male) underwent LGE-CMR and were followed for a median 3.5 years. The primary end point was SCD or aborted SCD.

Results Overall, 471 patients (66.2%) had myocardial fibrosis (median 5.9% of left ventricular mass, IQR: 2.2–13.3). Twenty-two (3.1%) reached the primary end point. The extent but not the presence of fibrosis was a significant univariable predictor of the primary end point (HR per 5% LGE: 1.24, 95% CI 1.06 to 1.45; p=0.007 and HR for LGE: 2.69, 95% CI 0.91 to 7.97; p=0.073, respectively). However, on multivariable analysis, only LV-EF remained statistically significant (HR: 0.92, 95% CI 0.89 to 0.95; p<0.001). For the secondary outcome of cardiovascular mortality/aborted SCD, the presence and the amount of fibrosis were significant predictors on univariable but not multivariable analysis after adjusting for LV-EF and non-sustained ventricular tachycardia.

Conclusions The amount of myocardial fibrosis was a strong univariable predictor of SCD risk. However, this effect was not maintained after adjusting for LV-EF. Further work is required to elucidate the interrelationship between fibrosis and traditional predictors of outcome in HCM.

INTRODUCTION

Hypertrophic cardiomyopathy (HCM) is a common myocardial disorder that in a minority of patients is associated with sudden cardiac death (SCD). Data from observational studies suggest that implantable cardioverter-defibrillator (ICD) therapy effectively prevents SCD, but also shows that the majority of ICD recipients do not receive appropriate therapies and have a high incidence of device-related complications. There is, therefore, a need to improve the identification of patients at risk. Current approaches to risk stratification rely on the assessment of a small number of clinical features that have been shown in observational cohort studies to associate with a greater risk of SCD. In aggregate, these clinical risk markers are associated with a substantially increased incidence of SCD in long-term studies, but are poorly predictive when they occur in isolation.

Myocardial fibrosis has been implicated in the pathogenesis of SCD in HCM. There has therefore been considerable interest in the use of late gadolinium enhancement (LGE) cardiovascular magnetic resonance (CMR) to detect and quantify myocardial fibrosis in vivo. Initial prospective cohort studies with short-term follow-up and therefore low event rates have suggested that the presence and amount of fibrosis as detected by LGE-CMR is of potential prognostic importance. These studies focused on surrogate end points, and despite the use of broad composite outcome measures, were inadequately powered to address SCD risk or to adjust for the impact of confounding variables. Thus, the role of LGE-CMR in the risk stratification of HCM for SCD and cardiovascular mortality remains unresolved. We therefore sought to determine the independent prognostic significance of myocardial fibrosis in a large consecutive HCM cohort with long-term follow-up.

METHODS

Study population

From September 2000 to June 2011, 815 consecutive patients with HCM referred for CMR at Royal Brompton Hospital were considered for enrolment. Reasons for referral included diagnostic evaluation, confirmation of diagnosis, family screening and assessment of disease severity. HCM was diagnosed in accordance with standard clinical guidelines. Patients were excluded if they had previous myectomy or alcohol septal ablation (n=21); previous myocardial infarction (n=14); or concomitant medical conditions to CMR and gadolinium-based contrast agents (n=12). Eight patients withdrew consent for study participation and 49 were lost to follow-up, resulting in a final study cohort of 711 patients. The study was performed in accordance with the Declaration of Helsinki and was approved by the National...
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Research Ethics Service, with written informed consent obtained from all patients.

Image acquisition
Images were acquired using 1.5 T scanners (Avanto/ Sonata, Siemens, Erlangen, Germany) with full myocardial coverage. Balanced steady-state free-precession sequences were used to obtain breath-hold cine images in three long-axis planes, followed by a contiguous stack of short-axis slices from the atrioventricular ring to the apex. Late enhancement images were acquired ~10 min after the administration of 0.1 mmol/kg intravenous gadolinium contrast agent (gadopentetate dimeglumine/gadobutrol, Bayer, Berlin, Germany) with an inversion recovery-prepared gradient-echo sequence. Inversion times were optimised to null normal myocardium with images acquired in two orthogonal phase-encoding directions to exclude artefact.

Image analysis
LV volumes, function, mass and EF were measured using a semiautomated threshold-based technique (CMRTools, Cardiovascular Imaging Solutions, London, UK). All volume and mass measurements were indexed to body surface area. Late enhancement was defined as an area of high signal intensity on a background of adequately nullified myocardium present in two orthogonal phase-encoding directions. The amount of LGE was quantified from a short-axis stack of images with dedicated software using the full-width at half-maximum method (CMR42, Circle Cardiovascular Imaging, Calgary, Canada). This was expressed as present or absent, and its amount was quantified as a percentage of LV mass as previously described.

Definition of outcome events and follow-up
All events were adjudicated by the consensus of an independent end point committee blinded to the CMR results. The primary end point was SCD or aborted SCD. The secondary end points were: a composite of cardiovascular mortality (defined as death due to heart failure, SCD, or non-haemorrhagic stroke), aborted SCD or cardiac transplantation; and all-cause mortality.

SCD was defined as unexpected death either within 1 h of cardiac symptoms in the absence of progressive cardiac deterioration; during sleep; or within 24 h of last being seen alive. Cardiac symptoms in the absence of progressive cardiac deterioration were defined as death associated with unstable progressive deterioration of pump function despite active therapy. Aborted SCD was diagnosed in patients who received an appropriate ICD discharge for ventricular arrhythmia or had a non-fatal episode of ventricular fibrillation (VF) or spontaneous sustained ventricular tachycardia (VT) (>30 s in duration) causing haemodynamic compromise and/or requiring cardioversion.

Survival status was determined through the UK National Strategic Tracing Service. The date and cause of death were determined using a combination of death certificates; post-mortem data when available; communication with patients’ primary care physicians and cardiologists; and hospital records. All patients were evaluated for non-fatal events using a standardised telephone and/or postal questionnaire. The patients’ primary care physicians and cardiologists were contacted to verify questionnaires and to obtain details of potential aborted SCD events, including the results of relevant ICD interrogations.

Statistical analysis
Continuous variables are expressed as mean±SD for normally distributed variables and as medians with IQR for non-parametric data. Differences between parametric continuous variables were assessed using Student t tests, and for non-parametric data, the Mann-Whitney U test. Categorical data are presented as frequencies and percentages, with differences assessed using the χ² and Fisher’s exact tests as appropriate. The start of follow-up was defined as the date of initial CMR evaluation. Patients were censored at the time of their last clinical follow-up. For composite end points, only the first event in each patient was considered for analysis. Univariable cumulative survival analysis was performed using the Kaplan-Meier method with a log-rank test to assess the significance of any difference between groups. To adjust for potential confounders, multivariable Cox proportional hazards modelling was carried out using covariates identified as significant on univariable analysis or based on previous clinical experience or models. Survival models were assessed for collinearity; interactions between covariates; and to ensure validity of the proportional hazards assumption. The number of covariates examined was restricted to maintain ≥10 events per degree of freedom. Two-tailed values of p<0.05 were considered statistically significant. Statistical analysis was performed using Stata SE V.12.0 (StataCorp, College Station, Texas, USA).

RESULTS
Study population
The baseline demographic and clinical characteristics of the study population are summarised in table 1. The median duration of follow-up was 3.5 years (IQR: 2.1 years to 5.6 years) amounting to 2852 patient-years of follow-up. The median age at study entry was 56.3 years. The majority of patients were male (70.0%) and minimally symptomatic (92.7% New York Heart Association (NYHA) class I/II). Patients with fibrosis were more likely to have extreme hypertrophy (LV wall thickness ≥30 mm) and non-sustained VT at baseline, giving rise to a greater proportion with two or more risk factors for SCD.

CMR findings
Patients with fibrosis had more severe hypertrophy compared with those without fibrosis exhibiting significantly higher maximum LV wall thickness and indexed LV mass (table 2). At baseline, LV-EF was significantly lower in those with fibrosis than those without.

Figure 1 illustrates some representative examples of patterns and severity of fibrosis seen in the study cohort. Two-thirds of the cohort exhibited LGE (66.2%) with a median amount of 5.9% of LV mass (IQR: 2.2 to 13.3). A maximum end-diastolic wall thickness of ≥19 mm was found on receiver operator characteristic analysis to optimally predict the presence of LGE with a sensitivity of 65.0%, specificity of 77.5% and Harrell’s c-statistic of 0.71 (95% CI: 0.68 to 0.75). The positive and negative predictive values were 85.0% and 53.0%, respectively, in our cohort. For patients with LV-EF ≤55%, the median amount of LGE was 8.0% of LV mass (IQR 2.9 to 31.4).

Primary outcome
During follow-up, 22 (3.1%) of the 711 patients reached the primary outcome measure of SCD or aborted SCD (table 3). Overall, 18 (3.8%) patients in the fibrosis group versus 4 (1.7%) in the no fibrosis group reached the primary outcome (HR LGE: 2.69, 95% CI 0.91 to 7.97, p=0.073) (figure 2). On univariable analysis, the amount of fibrosis was a significant predictor of outcome (HR per 5% LGE 1.24, 95% CI 1.06 to 1.45, p=0.007). A previous history of sustained VT/VF (HR: 7.76, 95% CI 1.79 to 33.57; p=0.006), and LV-EF (HR: 0.92,
Increased risk (lysis, only LV-EF emerged as an independent predictor (table 4, figure 2). Univariable analysis (table 4). However, on multivariable analysis (table 4), 95% CI 0.89 to 0.95; p<0.001) were also statistically significant on univariable analysis (figure 3). Increasing amounts of fibrosis were associated with increased risk (figure 4) but this relationship did not hold after adjusting for LV-EF (table 4).

### Secondary outcomes

In total, 38 (5.3%) of the 711 patients reached the principal secondary composite outcome measure of cardiovascular death (25 (3.5%) or aborted SCD (13 (1.8%)). Of the 25 cardiovascular deaths, 14 (56%) were due to heart failure, 9 (36%) to SCD and 2 (8%) to thromboembolic stroke (table 3). In the fibrosis group, 30 (6.4%) patients versus 8 (3.3%) in the no fibrosis group reached this outcome (HR for LGE: 2.24, 95% CI 1.03 to 4.89; p=0.043) (figure 4). Univariable analysis revealed that the amount of fibrosis was also a significant predictor (HR per 5% of fibrosis group, LV-ESV index—mL/m²: 1.26, 95% CI 1.12 to 1.41; p<0.001) together with LV-EF (table 3, figure 4), and non-sustained VT (see online supplementary table S2). However, after adjusting for the effects of LV-EF and non-sustained VT, neither the presence nor the amount of fibrosis remained statistically significant (see online supplementary table S1).

During follow-up, 55 (7.7%) of the 711 patients reached the secondary end point of all-cause mortality. Of the 55 deaths, 25 (45.5%) were due to cardiovascular causes. In contrast to the cardiovascular mortality composite, neither the presence nor the amount of fibrosis were significant univariable predictors (see online supplementary table S1, figure 2). For heart failure mortality, there were too few events to permit valid multivariable Cox regression analysis. However, on exploratory univariable analysis (table 3), similar to the primary and all-cause mortality end points, the amount but not the

### Table 1  Baseline clinical and demographic characteristics of the study cohort

<table>
<thead>
<tr>
<th>Characteristic—n (%)</th>
<th>No fibrosis</th>
<th>Fibrosis</th>
<th>All patients</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Characteristic—n (%)</td>
<td>240 (33.8)</td>
<td>471 (66.2)</td>
<td>711</td>
<td></td>
</tr>
<tr>
<td>Median follow-up—years (IQR)</td>
<td>4.0 (2.9, 5.8)</td>
<td>3.2 (1.9, 5.5)</td>
<td>3.5 (2.1, 5.6)</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Median age at study entry—years (IQR)</td>
<td>56.0 (46.1, 66.4)</td>
<td>56.9 (47.1, 66.8)</td>
<td>56.3 (46.7, 66.6)</td>
<td>0.686</td>
</tr>
<tr>
<td>Median age at diagnosis—years (IQR)</td>
<td>55.7 (45.5, 66.2)</td>
<td>54.7 (45.9, 65.5)</td>
<td>55.0 (45.8, 65.6)</td>
<td>0.742</td>
</tr>
<tr>
<td>Male sex—n (%)</td>
<td>164 (68.3)</td>
<td>334 (70.9)</td>
<td>498 (70.0)</td>
<td>0.478</td>
</tr>
<tr>
<td>Atrial fibrillation—n (%)</td>
<td>3 (1.3)</td>
<td>19 (4.2)</td>
<td>22 (3.2)</td>
<td>0.045</td>
</tr>
<tr>
<td>Apical phenotype—n (%)</td>
<td>60 (25.5)</td>
<td>66 (14.2)</td>
<td>126 (18.0)</td>
<td>&lt;0.001</td>
</tr>
</tbody>
</table>

Risk factors for SCD

- Sustained VT/VF—n (%) | 3 (1.3) | 11 (2.4) | 14 (2.0) | 0.326 |
- Family history of SCD—n (%) | 25 (10.5) | 51 (10.9) | 76 (10.8) | 0.859 |
- LV wall thickness ≥30 mm—n (%) | 0 (0) | 32 (6.8) | 32 (4.5) | <0.001 |
- Resting LVOT obstruction ≥30 mm Hg—n (%) | 65 (27.1) | 143 (30.4) | 208 (29.3) | 0.364 |
- Non-sustained VT—n (%) | 4 (1.7) | 34 (7.3) | 38 (5.4) | 0.002 |
- Unexplained syncope—n (%) | 19 (8.1) | 51 (11.0) | 70 (10.0) | 0.220 |
- Number of risk factors for SCD—n (%)* | 0 | 1 | 2± | 142 (59.2) | 232 (49.3) | 374 (52.6) | 0.005 |
- NYHA functional class—n (%) | 157 (66.5) | 282 (60.8) | 439 (62.7) | 0.331 |
- I | 65 (27.5) | 145 (31.3) | 210 (30.0) | 0.005 |
- II | 12 (5.1) | 35 (7.5) | 47 (6.7) | 0.005 |
- III | 2 (0.9) | 2 (0.4) | 4 (0.6) | 0.005 |
- IV | 30 (14.1) | 203 (46.1) | 272 (39.0) | <0.001 |

Medications at baseline—n (%) | 69 (30.9) | 203 (46.1) | 272 (39.0) | <0.001 |

| β-blocker | | | | |
| Ca²⁺-channel blocker | 30 (14.1) | 79 (19.6) | 109 (17.7) | 0.088 |

*Risk factors for SCD are as listed above.

LVOT, left ventricular outflow tract; NYHA, New York Heart Association; SCD, sudden cardiac death; VT, ventricular fibrillation; VT, ventricular tachycardia.

### Table 2  Baseline cardiovascular magnetic resonance findings for the study cohort

<table>
<thead>
<tr>
<th>CMR parameters—n (%)</th>
<th>No fibrosis</th>
<th>Fibrosis</th>
<th>All patients</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>CMR parameters—n (%)</td>
<td>240 (33.8)</td>
<td>471 (66.2)</td>
<td>711</td>
<td></td>
</tr>
<tr>
<td>Maximum end-diastolic LV wall thickness—mm</td>
<td>16.5±3.7</td>
<td>20.8±5.2</td>
<td>19.4±5.1</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>LV-EDV index—mL/m²</td>
<td>67.8±14.5</td>
<td>69.8±16.2</td>
<td>69.1±15.7</td>
<td>0.123</td>
</tr>
<tr>
<td>LV-ESV index—mL/m²</td>
<td>15.9±7.5</td>
<td>19.3±10.3</td>
<td>18.2±9.6</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>LV EF—%</td>
<td>77.1±7.3</td>
<td>73.3±9.5</td>
<td>74.6±9.0</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>50–59%</td>
<td>5 (2.1)</td>
<td>25 (5.3)</td>
<td>30 (4.2)</td>
<td>0.021</td>
</tr>
<tr>
<td>≥60%</td>
<td>232 (97.5)</td>
<td>434 (92.3)</td>
<td>666 (94.1)</td>
<td>0.123</td>
</tr>
<tr>
<td>LV mass index—g/m²</td>
<td>87.8±25.1</td>
<td>108.4±40.1</td>
<td>101.9±37.0</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Extent of LGE—% of LV mass</td>
<td>9.5 (9.9)</td>
<td>6.3 (9.2)</td>
<td>9.2 (9.2)</td>
<td>0.021</td>
</tr>
<tr>
<td>Median—% of LV Mass (IQR)</td>
<td>5.9 (2.2, 13.3)</td>
<td>2.2 (0, 9.1)</td>
<td>2.2 (0, 9.1)</td>
<td>0.021</td>
</tr>
</tbody>
</table>

All values are mean±SD unless otherwise stated.

CMR, cardiovascular magnetic resonance; EDV, end-diastolic volume; ESV, end-systolic volume; LGE, late gadolinium enhancement.
The presence of fibrosis was a significant univariable predictor of outcome (HR per 5% LGE: 1.23, 95% CI 1.02 to 1.48; p<0.028). No statistically significant interaction was detected between LV-EF and the amount of LGE for either primary end point (HR: 1.01, 95% CI: 0.99 to 1.03, p=0.303) or either of the secondary end points (cardiovascular mortality HR: 1.00, 95% CI 0.99 to 1.02, p=0.680; and all-cause mortality HR: 1.00, 95% CI 0.99 to 1.01, p=0.959).

DISCUSSION

The pressing need to improve risk stratification in HCM has driven the search for better biomarkers that interrogate the underlying substrate responsible for heart failure and SCD. Myocardial fibrosis can be assessed non-invasively using LGE-CMR and is thought to play a key role in arrhythmogenesis and the development of contractile dysfunction, providing a strong pathophysiological rationale for evaluating it as a biomarker of risk in HCM. In the present study, CMR evidence of fibrosis was identified in two-thirds of patients and the amount was an important predictor of SCD or aborted SCD risk but did not provide independent prognostic information incremental to LV-EF. Similar results were obtained with respect to the principal secondary composite end point of cardiovascular mortality or aborted SCD.

The present findings add substantially to our earlier work, and two other previous smaller studies that have examined the prognostic utility of LGE-CMR in HCM. We previously found that the presence and the amount of fibrosis predicted adverse cardiovascular outcomes in 217 patients with HCM. In this earlier work, our primary composite end point

<table>
<thead>
<tr>
<th>Table 3 Summary of outcomes for the whole cohort and stratified according to the presence/absence and the amount of fibrosis</th>
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<tbody>
<tr>
<td><strong>Outcome measure, n (%)</strong></td>
</tr>
<tr>
<td>----------------------------</td>
</tr>
<tr>
<td>SCD or aborted SCD</td>
</tr>
<tr>
<td>Cardiovascular death or aborted SCD</td>
</tr>
<tr>
<td>Heart failure death</td>
</tr>
<tr>
<td>SCD</td>
</tr>
<tr>
<td>Aborted SCD</td>
</tr>
<tr>
<td>Thromboembolic stroke</td>
</tr>
<tr>
<td>All-cause mortality</td>
</tr>
</tbody>
</table>

SCD, sudden cardiac death.

*HRs per 5% of LV mass with late gadolinium enhancement.

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Figure 1 Representative examples of the patterns and spectrum of late gadolinium enhancement/fibrosis seen in hypertrophic cardiomyopathy. Diffuse fibrosis of the basal anteroseptum and inferoseptum (A); mixture of patchy and diffuse fibrosis of the interoseptum and inferolateral walls (B); focal insertion point and mid-wall fibrosis (C); and extensive dense patchy fibrosis of the basal anteroseptum and inferoseptum (D).
groups. In the present much larger study with longer follow-up, there were 22 SCDs or aborted SCDs and 38 cardiovascular deaths or aborted SCDs in total, allowing us to make more robust inferences on the prognostic value of fibrosis.

Bruder et al13 conducted the only other published prospective LGE-CMR study to date using cardiovascular mortality as an end point. They were able to demonstrate that fibrosis was a statistically significant univariable predictor of outcome, but with only a total of 16 events, similar to our own previous work, their study was not sufficiently powered to demonstrate the independent prognostic significance of fibrosis with respect to SCD or cardiovascular mortality over and above potential confounders.9 In particular, as in the present study, fibrosis was highly associated with significant hypertrophy, defined in terms of maximal LV wall thickness and indexed LV mass. There was also a significant excess of patients with non-sustained VT in their fibrosis group relative to those without fibrosis. The latter association has also been noted in numerous earlier studies.9 11 12 15 22 23

In the only other published prospective LGE-CMR study, Maron et al14 evaluated the clinical significance of fibrosis in 202 patients with HCM using a composite primary end point of cardiovascular death or progressive heart failure, which was defined as a change in NYHA class. Despite the use of this broad end point, and the inclusion of antiarrhythmic pacing as a SCD-equivalent, only 11 adverse cardiovascular events occurred over a mean follow-up of 1.9 years. Their event rate was therefore insufficient to identify either the presence or the amount of fibrosis as even an univariable predictor of outcome.9 However, in keeping with the present study, a statistically significant inverse association was found between LV systolic function as measured by LV-EF and the presence and amount of fibrosis.14 The latter finding is also in accordance with the results of Rubinstein et al.15 Their retrospective cohort study also identified the presence of fibrosis as a significant univariable predictor of SCD or appropriate ICD discharge. However, as only eight events occurred among their 424 patients after a mean of 3.6 years follow-up, in common with previous studies,13 14 they were underpowered to adjust for statistically significant differences in baseline characteristics between those with and without fibrosis.13

In the present study, LV-EF emerged as the most powerful independent predictor of SCD risk as well as cardiovascular and all-cause mortality. Although widely used to guide the management of patients with ischaemic heart disease and non-ischaemic dilated cardiomyopathy,23 26 the assessment of LV systolic function is not currently emphasised in guidelines for the risk stratification of HCM.17 In the largest series to date, Harris et al17 examined the prevalence and significance of adverse LV remodelling in HCM.27 Of their retrospectively identified cohort of 1259 consecutive patients with HCM, 44 (3.5%) had evidence of LV systolic dysfunction defined by echocardiography as LV-EF ≤ 50% at rest. In our cohort, the prevalence of LV dysfunction defined using an equivalent CMR threshold of LV-EF ≤ 50% was similar in 23 (3.2%) of 711 patients. However, of these, 5 (21.7%) reached the primary end point of SCD or aborted SCD, and 9 (39.1%) reached the principal secondary cardiovascular composite outcome in comparison with two-thirds of patients in the Harris cohort, although their higher incidence of adverse cardiovascular events may reflect their longer follow-up duration. Of note, 9 (81%) of a subgroup of 11 of their end-stage patients had evidence of extensive fibrosis on CMR or at autopsy. In the present study, 21 of 23 patients (91%) with LV dysfunction had CMR evidence of fibrosis. The superior predictive value of LV-EF over and above fibrosis may reflect the fact that it provides a better synthesis of the various ventricular effects of HCM,

Figure 2 Kaplan–Meier estimates of event-free survival for the primary outcome measure of sudden cardiac death or aborted sudden cardiac death (A); the principal secondary end point of cardiovascular mortality or aborted sudden cardiac death (B); and all-cause mortality (C) stratified according to the presence and absence of fibrosis. LGE, late gadolinium enhancement.
which include fibrosis and genetically mediated contractile dysfunction and energetic abnormalities.\(^2\)\(^,\)\(^3\)\(^,\)\(^8\)

Although the poor prognosis of end-stage or incipient end-stage disease is recognised, LV dysfunction is not sufficiently emphasised in current clinical guidelines for risk stratification.\(^1\)\(^7\)\(^,\)\(^2\)\(^9\)

As a consequence of LV hypertrophy, LV end-diastolic volumes tend to be small in HCM, resulting in large fractional volume changes in systole. Thus, even an apparently borderline EF may tend to be small in HCM, resulting in large fractional volume changes. Thus, even an apparently borderline EF may be a harbinger of incipient adverse outcomes analogous to the situation of LV volume overload due to chronic mitral regurgitation.

Limitations

Our study was conducted at a single tertiary centre. While this facilitated the use of a standardised scanning and follow-up protocol, it also incurred the possibility of selection bias. However, referrals to our centre are drawn from an extensive network of district general hospitals spanning the entire south-east and the wider UK, mitigating against this possibility.

Although the event rates observed in the present study were considerably higher than those found in community-based cohorts,\(^1\)\(^\,\)\(^4\)\(^\,\)\(^6\)\(^\,\)\(^9\)\(^,\)\(^1\)\(^3\)\(^\,\)\(^1\)\(^4\)\(^,\)\(^1\)\(^5\)\(^,\)\(^1\)\(^6\) they are comparable with those seen in other studies of patients who have come to clinical attention.\(^1\)\(^7\)\(^,\)\(^1\)\(^8\)\(^,\)\(^1\)\(^9\)\(^,\)\(^2\)\(^0\)

Our data are therefore representative of patients encountered by cardiologists in clinical practice. Indeed, high-risk patients may have been inadvertently selected out from our cohort. For instance, those with an ICD at baseline were necessarily excluded as this represents a contraindication to CMR. Similarly, patients with previous gradient reduction therapy were also excluded as this introduces confounding iatrogenic scar, potentially lowering the event rate observed.

The amount of LGE was a statistically significant univariate predictor of the primary and principal secondary end points. This implies by definition that our study was adequately powered to detect a statistically significant difference in outcome based on the amount of LGE. However, on limited further modelling, the amount of LGE did not emerge as an independent risk factor incremental to other factors. While we cannot exclude the possibility that a significantly larger sample size and/or longer duration of follow-up would be able to tease out a separate independent effect, we believe that this implies that the effect size that we would be trying to detect is very small and therefore the positive predictive value of the amount of LGE over and above existing markers is likely to be very limited.

Although 49 patients were lost to follow-up, we can confirm from death certificate records that none of them died within the UK. Nevertheless, we cannot exclude the possibility that these patients may have emigrated overseas or while within the UK, received an ICD which was then subsequently liable to discharge. However, generously assuming a device implantation rate of 10%, only ~five patients would have received an ICD. With an annual discharge rate for primary prevention of 3.6% per annum with a cumulative 5-year probability of 17%,\(^5\) only ~one patient is likely to have received an appropriate shock over 5 years (our median follow-up was 3.5 years). Therefore, it is unlikely that statistically significant events were not captured that would have altered our data interpretation. Furthermore, among the 49 patients lost to follow-up, 24 (49.0%) had no fibrosis versus 240 (33.8%) of the 711 patients with full follow-up (p=0.0426). Thus, significantly more patients without LGE were lost to follow-up, making it even more unlikely that undetected events would have altered our assessment.

Finally, the LGE-CMR technique exploits relative differences in signal intensity between fibrotic and ostensibly normal myocardium. The full-width at half-maximum technique has been shown to be the most reproducible quantification method.\(^2\)\(^0\)

### Table 4

<table>
<thead>
<tr>
<th>Predictor</th>
<th>Hazard ratio</th>
<th>95% CI</th>
<th>p Value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Male</td>
<td>1.42</td>
<td>0.52 - 3.84</td>
<td>0.493</td>
</tr>
<tr>
<td>New York Heart Association class</td>
<td>1.02</td>
<td>0.54 - 1.93</td>
<td>0.951</td>
</tr>
<tr>
<td>LV wall thickness ≥30 mm</td>
<td>7.76</td>
<td>1.79 - 33.57</td>
<td>0.006</td>
</tr>
<tr>
<td>LV outflow tract obstruction ≥30 mm Hg</td>
<td>0.50</td>
<td>0.17 - 1.47</td>
<td>0.207</td>
</tr>
<tr>
<td>Non-sustained ventricular tachycardia</td>
<td>1.71</td>
<td>0.40 - 7.38</td>
<td>0.472</td>
</tr>
<tr>
<td>Unexplained syncope</td>
<td>0.84</td>
<td>0.19 - 3.59</td>
<td>0.809</td>
</tr>
<tr>
<td>Male</td>
<td>1.00</td>
<td>0.55 - 1.80</td>
<td>0.995</td>
</tr>
<tr>
<td>Maximum LV end-diastolic wall thickness (mm)</td>
<td>1.01</td>
<td>0.93 - 1.09</td>
<td>0.845</td>
</tr>
<tr>
<td>Indexed LV end-diastolic volume (per 10 ml/m²)</td>
<td>1.17</td>
<td>0.89 - 1.54</td>
<td>0.257</td>
</tr>
<tr>
<td>Indexed LV end-systolic volume (per 10 ml/m²)</td>
<td>1.88</td>
<td>1.45 - 2.45</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>LV EF (%)</td>
<td>0.92</td>
<td>0.89 - 0.95</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>LV mass index (per 10 g/m²)</td>
<td>1.07</td>
<td>0.99 - 1.17</td>
<td>0.104</td>
</tr>
<tr>
<td>Presence of fibrosis</td>
<td>2.69</td>
<td>0.91 - 7.97</td>
<td>0.073</td>
</tr>
<tr>
<td>Amount of fibrosis (per 5% of LV mass)</td>
<td>1.24</td>
<td>1.06 - 1.45</td>
<td>0.007</td>
</tr>
<tr>
<td>Presence of fibrosis as a candidate</td>
<td>0.92</td>
<td>0.89 - 0.95</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>LV EF (%)</td>
<td>2.98</td>
<td>0.66 - 13.44</td>
<td>0.155</td>
</tr>
<tr>
<td>Presence of fibrosis</td>
<td>0.92</td>
<td>0.89 - 0.95</td>
<td>&lt;0.001</td>
</tr>
<tr>
<td>Amount of fibrosis as a candidate</td>
<td>1.10</td>
<td>0.92 - 1.31</td>
<td>0.299</td>
</tr>
</tbody>
</table>
Nevertheless, there is no widespread consensus on the best quantification strategy to use and a paucity of histological validation for the LGE technique in the setting of cardiomyopathy.11 12 13

![Graphs and Figures](http://heart.bmj.com)

**Figure 3** Kaplan–Meier estimates of event-free survival for the primary outcome measure of sudden cardiac death or aborted sudden cardiac death (A); the principal secondary end point of cardiovascular mortality or aborted sudden cardiac death (B); and all-cause mortality (C) stratified according to LVEF.

**Figure 4** Predicted 5-year risk of reaching the study outcome measures according to amount of fibrosis (A) and LVEF (B). ASCD, aborted sudden cardiac death; CV, cardiovascular; AC, all-cause; SCD, sudden cardiac death.

**Key messages**

**What is known on this subject?**

Multiple small retrospective and prospective cohort studies using composite outcome measures have suggested that the presence and amount of myocardial fibrosis, as detected by late gadolinium enhancement (LGE)-cardiovascular magnetic resonance (CMR), are associated with adverse outcomes in hypertrophic cardiomyopathy (HCM). However, none of these studies were of sufficient size and/or duration to be able to address the prognostic significance of myocardial fibrosis with respect to sudden cardiac death (SCD) risk, leaving the role of LGE-CMR in HCM risk stratification unresolved.

**What might this study add?**

This is the first study to definitively confirm an association between the amount of fibrosis, but not its mere presence, and SCD risk. However, after adjusting for other factors such as LV-EF, the amount of fibrosis did not retain independent significance.

**How might this impact on clinical practice?**

The amount of myocardial fibrosis is an important predictor of SCD risk in HCM, but may not provide information incremental to other factors. In contrast, LV dysfunction emerged as an independent risk factor and should be given greater emphasis in clinical guidelines as a harbinger of increased SCD risk.


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CONCLUSIONS
The amount of myocardial fibrosis is an important predictor of outcome in HCM but may not provide information incremental to LV-EF. The latter is an important prognosticator for outcome in HCM, which is underemphasised in current guidelines. Further work is required to elucidate the interrelationship between myocardial fibrosis and other markers of risk in HCM.

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Contributors
SKP and DIP conceived the study and were responsible for study design. WJM, ICM, MRC, MR, ROH and PE advised on study design. TFI, AJ, AG, AM, SR, TEC, BD, JK, FFA, RW, AV, CS, SAC and ROH were involved in data gathering, analysis, interpretation as well as drafting of the manuscript. MR was responsible for all data and statistical analysis. TFI drafted the initial manuscript. All authors were involved in the critical revision of the manuscript for important intellectual content.

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Competing interests
SKP has received honoraria from Bayer-Schering. DIP is a consultant to Siemens and a director of Cardiovascular Imaging Solutions. Royal Brompton Hospital has a research collaboration agreement with Siemens. MRC has received speaker fees from Roche Diagnostics and Pfizer; grants from Medtronic and ResMed; and is a consultant to Novartis, Medtronic, ST Jude Medical, Boston Scientific, GE Medical and Servier. None of the other authors report any disclosures relevant to this work.

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Royal Brompton and Harefield NHS Trust Ethics Committee.

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