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Electrophysiological Characterization of Subclinical and Overt Hypertrophic Cardiomyopathy by Magnetic Resonance Imaging-Guided Electrocardiography

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ABSTRACT

BACKGROUND Ventricular arrhythmia in hypertrophic cardiomyopathy (HCM) relates to adverse structural change and genetic status. Cardiovascular magnetic resonance (CMR)-guided electrocardiographic imaging (ECGI) noninvasively maps cardiac structural and electrophysiological (EP) properties.

OBJECTIVES The purpose of this study was to establish whether in subclinical HCM (genotype [G]+ left ventricular hypertrophy [LVH]–), ECGI detects early EP abnormality, and in overt HCM, whether the EP substrate relates to genetic status (G+/G-LVH+) and structural phenotype.

METHODS This was a prospective 211-participant CMR-ECGI multicenter study of 70 G+LVH-, 104 LVH+ (51 G+/53 G-), and 37 healthy volunteers (HVs). Local activation time (AT), corrected repolarization time, corrected activation-recovery interval, spatial gradients (G_{AT}/G_{RTc}), and signal fractionation were derived from 1,000 epicardial sites per participant. Maximal wall thickness and scar burden were derived from CMR. A support vector machine was built to discriminate G+LVH- from HV and low-risk HCM from those with intermediate/high-risk score or nonsustained ventricular tachycardia.

RESULTS Compared with HV, subclinical HCM showed mean AT prolongation (P = 0.008) even with normal 12-lead electrocardiograms (ECGs) (P = 0.009), and repolarization was more spatially heterogenous (G_{RTc} : P = 0.005) (23% had normal ECGs). Corrected activation-recovery interval was prolonged in overt vs subclinical HCM (P < 0.001). Mean AT was associated with maximal wall thickness; spatial conduction heterogeneity (G_{AT}) and fractionation were associated with scar (all P < 0.05), and $G_{+L}VH_{+}$ had more fractionation than $G_{-L}VH_{+}$ (P = 0.002). The support vector machine discriminated subclinical HCM from HV (10-fold cross-validation accuracy 80% [95% CI: 73%-85%]) and identified patients at higher risk of sudden cardiac death (accuracy 82% [95% CI: 78%-86%]).



Listen to this manuscript's audio summary by Editor-in-Chief Dr Valentin Fuster on www.jacc.org/journal/jacc. **CONCLUSIONS** In the absence of LVH or 12-lead ECG abnormalities, HCM sarcomere gene mutation carriers express an aberrant EP phenotype detected by ECGI. In overt HCM, abnormalities occur more severely with adverse structural change and positive genetic status. (J Am Coll Cardiol 2024;83:1042-1055) © 2024 The Authors. Published by Elsevier on behalf of the American College of Cardiology Foundation. This is an open access article under the CC BY-NC-ND license (http://creativecommons.org/licenses/by-nc-nd/4.0/).

From the ^aBarts Heart Centre, Barts Health NHS Trust, London, United Kingdom; ^bInstitute of Cardiovascular Science, University College London, London, United Kingdom; ^cMedical Research Council Unit for Lifelong Health and Ageing, University College London, London, United Kingdom; ^dCentre for Inherited Heart Muscle Conditions, Department of Cardiology, Royal Free London NHS Foundation Trust, London, United Kingdom; ^eElectrocardiology Section, School of Health and Wellbeing, University of Glasgow, Glasgow, United Kingdom; ^fCardiology Clinical and Academic Group, St George's University of London and St George's **P** roarrhythmic substrate in hypertrophic cardiomyopathy (HCM) is related to adverse cardiac structural change (hypertrophy, fibrosis, disarray, small vessel disease), and those with a sarcomere gene mutation (genotype [G]+ left ventricular hypertrophy [LVH]+ vs G-LVH+) have a higher incidence of sudden cardiac death (SCD).¹⁻³ Twelvelead electrogram (ECG) abnormalities occur in those with sarcomeric mutations without hypertrophy (G+LVH- also known as subclinical HCM) and predict future LVH.⁴ Both the origins and mechanisms underlying ventricular arrhythmia remain poorly understood.

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Abnormalities in ventricular activation and repolarization could be the link between structural changes and SCD.^{3,5} In this domain, electrocardiographic imaging (ECGI) provides detailed noninvasive electrophysiological (EP) assessment in intact hearts under physiological conditions⁶ and may be more sensitive to early changes (subclinical disease) than structural assessment by macroscopic imaging, ie, cardiovascular magnetic resonance (CMR) alone. ECGI is a high spatiotemporal resolution method of computing unipolar epicardial electrograms (UEGs) to panoramically map ventricular activation and repolarization.⁷ It has been validated ex vivo and in vivo, provided EP insights into multiple diseases, and shown potential to guide device and ablation therapies.7-13 Recent advances in this technique include our development of a fully washable and reuseable vest that has demonstrated high repeatability and allows noninvasive performance of ECGI at scale.¹⁴ Integration with advanced myocardial tissue characterization is therefore enabled by CMR. This approach, "CMR-ECGI," now permits fully coregistered simultaneous interrogation of the myocardial structure, scar, and EP properties.

We hypothesized that CMR-guided ECGI would detect the following: 1) subtle EP abnormalities in subclinical HCM; and 2) EP abnormalities in overt disease related to genetic status (G+ vs G-LVH+) and structural changes (maximal wall thickness [MWT], late gadolinium enhancement [LGE]).

METHODS

Ethical approval was obtained for each study site from the UK Research Ethics Committee (IRAS 227168), and all participants provided informed consent to participate in the study. An academic collaboration was formed for large-scale clinical deployment of ECGI technology (capturECGI Vest [Patent Application No. US 18/194 235] + body surface ECG + heart-torso geometry preprocessing¹⁴ led by University College London [UCL], United Kingdom; inverse solution led by Prof Rudy Lab, Washington University, St Louis, USA; digital 12-lead ECG analysis led by University of Glasgow, United Kingdom; and advanced artificial intelligence [AI]-powered CMR analysis + sequence optimization led by Barts Heart Centre, UCL, and National Institutes of Health, USA).

STUDY POPULATION. Adult participants (age 18-76 years) were recruited from cardiomyopathy clinics of 3 tertiary referral sites (Barts Heart Centre, St George's University of London, Royal Free London, United Kingdom). Genotype classification was performed according to American College of Medical Genetics criteria.¹⁵ Hypertrophic cardiomyopathy was defined as per European Society of Cardiology (ESC)/ American Heart Association guidelines (unexplained LVH: MWT >13 mm in any American Heart Association segment using either echocardiography or CMR in first-degree relatives of affected probands or else \geq 15 mm).^{1,16} HCM was diagnosed by an inherited cardiomyopathy expert in a recruiting site. Overt HCM patients with intermediate/high SCD risk were identified based on the ESC risk score.¹⁷ Subclinical HCM (G+LVH-) was defined in individuals having a pathogenic/likely pathogenic sarcomere gene variant identified by cascade screening but without LVH. Healthy volunteers recruited under the same ethical

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ABBREVIATIONS AND ACRONYMS

 ΔAT = dispersion of activation time

 $\Delta \mathbf{RT_c}$ = dispersion of repolarization time corrected

ARIC = activation recovery interval corrected

AT = activation time

G+/- = genotype positive/ negative

G_{AT} = gradient of activation time

G_{RTc} = gradient of repolarization time corrected

LGE = late gadolinium enhancement

LVH+/- = left ventricular hypertrophy positive/negative

RTc = repolarization time corrected

UEG = unipolar electrogram

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The authors attest they are in compliance with human studies committees and animal welfare regulations of the authors' institutions and Food and Drug Administration guidelines, including patient consent where appropriate. For more information, visit the Author Center.

protocol did not undergo genetic testing or Holter monitoring. Exclusion criteria were contraindication to CMR, needle-phobia, inability to consent, known or clinically suspected coronary artery disease, prior myectomy/alcohol septal ablation, implantable cardiac devices, persistent atrial fibrillation, high ventricular ectopic burden, poorly controlled hypertension (suboptimal control despite 2 antihypertensives) and significant valve disease. No patients were on cardiac myosin inhibitors. Recent (within 12 months) clinical Holter results and clinical echocardiograms identified participants with nonsustained ventricular tachycardia (≥3 beats of ventricular tachycardia at ≥120 beats/min) and LV outflow tract obstruction (gradient \geq 30 mm Hg under any condition).

ECGI ACQUISITION AND ANALYSIS. A resting 256lead ECG in the recumbent position was recorded for 5 minutes for signal averaging at a sampling frequency of 2,400 Hz using the fully reuseable, washable capturECGI vest and a high-performance amplifier system (g.HIamp 256 bundle GT-8016/ USBamp GT-0216; g.tec medical engineering GmbH).¹⁴ The 5-minute body surface recording was quality controlled using in-house Matlab software (Mathworks) (Figure 1). Signal averaging enhanced the signal quality, and low-quality signals were removed. The vest was replaced with an identically sized "mirror-vest" using 256 identically positioned magnetic resonance imaging lucent fiducial markers for scanning. Heart-torso geometry was obtained by segmenting an anatomical transaxial stack (see following text) using Amira-Avizo software (Thermo Fisher, version 2021.1) for electrode positions and epicardial meshes. Body surface recordings and heart-torso geometry underwent the inverse solution of electrocardiography (defined as the reconstruction of cardiac electrical activity from given body surface electrocardiographic measurements) using previously validated protocols.7 Computed electrodes arising from valve-plane (nonconducting tissue) were excluded. A total of 1,000 UEGs (Figure 1) were computed per heart to obtain activation time (AT) (the time from the earliest activation in the epicardium to steepest point of the QRS downslope), repolarization time corrected for heart rate (RTc) (the steepest point of the T-wave upslope¹⁸), and activation-recovery interval (ARIc) (the difference of AT and RTc). Heart rate correction used the Fridericia formula.¹⁹ Dispersions of AT and ARIC (Δ AT, Δ ARIC) were measured as the maximum-minimum of AT and ARIc, respectively, across the entire myocardium. Spatial gradients (gradient of activation time $[G_{AT}]$, gradient of repolarization time corrected [G_{RTc}]) were

computed for each epicardial site as the absolute value of the difference between neighboring sites (within 15 mm) divided by their distance, averaged across all neighbors. Signal amplitudes were defined as the peak to peak (maximum to minimum) of the QRS complexes. Parameters from all 1,000 electrodes were averaged per participant and maximal gradients were also recorded (max: G_{AT}/G_{RTc}). Subclinical HCM with max/mean $G_{RTc} > 2$ SDs above the mean were defined as having abnormal spatial repolarization. Signal fractionation was defined as number of UEGs with ≥ 2 negative deflections within the QRS complex.

CMR ACQUISITION AND ANALYSIS. All scans were performed at 3-T (Prisma, Siemens Germany) in UCL's Bloomsbury Center for Clinical Phenotyping immediately after body surface ECG recordings. Hearttorso geometry was obtained from a free-breathing 100 thin-slice (4 mm contiguous) Half-Fourier Acquisition Single-shot Turbo spin Echo imaging (HASTE) sequence.¹⁴ Cine imaging was conventional with retrospectively gated breath-held balanced single-shot free precession short- and long-axis views. These were segmented for cardiac volume and wall thickness using fully automated AI-based algorithms shown to exceed human precision.^{20,21} Free-breathing motion-corrected LGE data were acquired 5 minutes following intravenous contrast (Dotarem, Guerbet). LGE was quantified using the 5-SD technique and expressed as absolute mass in grams. Precontrast and postcontrast modified Look-Locker inversion recovery (MOLLI) T1 with sameday hematocrit for extracellular volume (ECV) were acquired and septal ECV was computed.²²

DIGITAL 12-LEAD ECG. A 12-lead ECG was acquired at rest using the Beneheart R3 (Mindray), and recordings were sent to University of Glasgow Core-Lab for automated analysis of electrical intervals (PR, QRS, QTc), QT dispersion, amplitudes, and the presence of ECG abnormalities.^{23,24} ECG abnormalities known to be relevant to disease progression in HCM^{4,25} were defined as follows: abnormal Q waves (≥2 contiguous leads and with minimum amplitude 0.3 mV or \geq 25% of the subsequent R-wave, or duration >40 ms); LVH criteria defined as Sokolow-Lyon or Cornell criteria and repolarization abnormalities (T-wave inversion of \geq 0.1 mV in \geq 2 contiguous leads and/or ST-segment depression $\ge 0.1 \text{ mV}$ in $\ge 2 \text{ contiguous leads}$; and ECG fractionation defined as notching within the QRS complex (including R', r', S notching, R notching, or fragmentation) in 2 contiguous leads.

STATISTICAL ANALYSIS. Statistical analysis was performed using Matlab (MathWorks, version May 11,



(Top) A 256-lead electrocardiogram (ECG) is recorded using the fully re-useable capturECGI vest. The mirror vest is positioned to match the electrode vest and the participant wears this during the dark blood (HASTE [Half-Fourier Acquisition Single-shot Turbo spin Echo imaging]) anatomical stack acquisition. Heart-torso geometry is obtained using segmentation of the epicardium and the markers on the mirror vest using Amira-Avizo software (Thermo Fisher). The inverse solution is performed according to previously described protocols thereby obtaining 1,000 computed unipolar electrograms. (Bottom Left) Activation time (AT) is defined by the steepest QRS downslope. Repolarization time (RT) is defined by the steepest part of QRS upslope. Both are referenced to earliest epicardial activation. Activation recovery interval (ARI) is the difference between RT and AT. Three points (circle, triangle, square) on AT and ARIc maps are shown and their corrected for heart rate. (Bottom right) AT/RT gradients: (ΔAT/ ΔRT) between adjacent orange and blue unipolar electrograms (UEGs) are shown. Gradients are measured as the difference in AT/RT between neighboring electrograms divided by their inter-electrode distance (d).

2022) and SPSS (IBM SPSS Statistics, version 28). The distribution of data was examined using histograms, Shapiro-Wilk tests and QQ plots. Normally distributed continuous variables were summarized as mean \pm SD and compared using independent samples Student's t-tests; non-normally distributed variables were presented as median (Q1-Q3) and compared using Mann-Whitney U tests. Multiple testing with Bonferroni was performed on a 3 comparisons per variable basis (P < 0.05/3). Categorical/binomial variables were expressed as absolute counts and percentages and compared using the chi-square/Fisher exact test as appropriate. Spearman's correlations were used to examine simple linear/monotonic relationships between structural variables, risk markers (nonsustained ventricular tachycardia [NSVT]/left ventricular outflow tract obstruction [LVOTO]) and ECG/ ECGI parameters. Multiple linear regression models were used to explore independent between-group differences by including a priori defined confounders as covariates, namely age, sex, LGE volume, and QRS duration for AT comparisons or QTc for ARIc comparisons. Additionally, differences in G+LVH- were adjusted for the presence vs absence of an abnormal resting 12-lead ECG. Variance inflation factor <3 excluded collinearity. Further details are provided in the Supplemental Methods. To discriminate subclinical HCM from health, a supervised machine learning approach using a support vector machine (SVM) classification was built. This incorporated all 12-ECGI biomarkers and was applied to the subclinical vs control data set followed by a 10-fold cross validation. Confusion matrices were generated to summarize model performance, and area under the receiver-operator curves (AUCs) were calculated using R package (ROCR). A similar SVM and receiver-operator curve approach explored whether in patients with overt HCM, those with high/intermediate risk (from the ESC Risk score) or those with prior NSVT could be discriminated from low-risk patients (Supplemental Methods).

RESULTS

A total of 211 participants were prospectively studied: 70 G+LVH- subjects, 104 LVH+ patients (51 G+LVH+, 53 G-LVH+), and 37 healthy volunteers (HVs), of whom 26 were younger and used for age-matched comparisons with G+LVH-, whereas 23 (with overlap) were older and used for age-matched comparisons with G+LVH+ (reported in Supplemental Table 1). A comparison of Drug-Free HCM and matched healthy volunteers are shown in Supplemental Table 2. Relationships between LV morphology and ECGI parameters are shown in Supplemental Table 3. Likely pathogenic/pathogenic variants for G+LVH– were present in the following genes: 41 (59%) *MYBPC3*, 16 (23%) *MYH7*, 5 (7%) *TNNI3*, 4 (6%) *TNNT2*, 1 (1%) *MYL2*, 1 (1%) *CSRP3*, 1 (1%) *ACTC1*, 1 (1%) *TPM1* and for G+LVH+: 28 (55%) *MYBPC3*, 13 (25%) *MYH7*, 4 (8%) *TNNI3*, 4 (8%) *TNNT2*, 1 (2%) *TNNC1*, and 1 (2%) *CSRP3*.

CLINICAL CHARACTERISTICS. Subclinical HCM (G+LVH-) vs HV. Compared with age-matched HVs, G+LVH- had similar age, sex, ethnicity, and body surface area (BSA). G+LVH- had similar LV cavity size and myocardium (MWT, LV mass, ECV, and LGE) but more hyperdynamic function. G+LVH- had a stronger family history of SCD.

G + LVH + vs G + LVH -. Compared with G+LVH-, G+LVH+ had similar BSA but were older, were more often male and with smaller cavity sizes, had more abnormal myocardium (higher MWT, LV mass, ECV, and LGE), and had higher ejection fraction.

G - LVH + vs G + LVH +. Compared with G+LVH+, G-LVH+ had similar sex and BSA; were older; had similar LV cavity sizes, function, and MWT; had greater LV mass; had lower ECV; and had similar LGE. There was no difference in LV morphology. G+LVH+ had a stronger family history of sudden cardiac death (Table 1).

12-LEAD ECG. Subclinical HCM (G+LVH-) vs HV. Compared with age-matched HVs, G+LVH- had shorter QRS intervals but otherwise similar electrical intervals, QRS amplitudes, and QT dispersion. G+LVH- had a higher prevalence of ECG abnormalities (31% [22] vs 8% [2]; P = 0.019, but not after correction for multiple comparisons).

G + LVH + vs G + LVH -. Compared with G+LVH-, G+LVH+ had similar heart rates, longer QRS and QTc intervals, greater QT dispersion, and greater Cornell amplitude (but similar Sokolow-Lyon). G+LVH+ also had a higher prevalence of ECG abnormalities including T-wave inversion (TWI), LVH by either ECG voltage criteria, ST depression, and fractionation.

G - LVH + vs G + LVH +. Compared with G+LVH+, G-LVH+ had longer QTc intervals, higher QRS amplitudes and a higher prevalence of LVH by either ECG voltage criteria. Other 12-lead ECG intervals and prevalence of ECG abnormalities were similar (QRS duration was longer in G-LVH+ but not after correction for multiple comparisons) (Table 1).

ECGI. G+**LVH**- **vs HV**. Compared with age-matched HVs, G+LVH- had slower ventricular conduction (more prolonged mean AT) (Table 2, Figure 2).

TABLE 1 Demographics, CMR, and 12-Lead ECG Variables in Younger HVs and HCM Cohorts											
					<i>P</i> Value						
	Younger HV ^a (n = 26)	G+LVH— (n = 70)	G+LVH+ (n = 51)	G–LVH+ (n = 53)	G+LVH– vs HV	G+LVH+ vs G+LVH–	G-LVH+ vs G+LVH+				
Demographics											
Age, y	35 (25-38)	36 (25-41)	52 (37-59)	59 (50-66)	0.85	<0.001	0.005				
Female	14 (54)	41 (59)	17 (33)	8 (15)	0.68	0.003	0.09				
White	21 (81)	57 (81)	43 (84)	36 (68)	0.94	0.68	0.051				
BSA, m ²	$\textbf{1.8}\pm\textbf{0.2}$	$\textbf{1.9}\pm\textbf{0.2}$	$\textbf{1.9}\pm\textbf{0.2}$	$\textbf{2.0} \pm \textbf{0.2}$	0.45	0.21	0.42				
BMI, kg/m ²	24 (22-27)	25 (22-28)	25 (24-28)	26 (24-28)	0.24	0.65	0.5				
Hypertension	0 (0)	4 (6)	9 (18)	19 (36)	0.29	0.029	0.035				
Diabetes	0 (0)	0 (0)	2 (4)	0 (0)	>0.9	0.17	0.15				
On lipid lowering	0 (0)	0 (0)	6 (12)	16 (30)	>0.9	0.005	0.021				
On antiarrhythmic	0 (0)	3 (4)	21 (41)	25 (47)	0.56	<0.001	0.54				
Family history of SCD	0 (0)	12 (17)	13 (25)	5 (9)	0.023	0.28	0.042				
Volumes and mass											
LVEDV index, mL/m ²	89.3 (74-109)	81.4 (70-94)	71.0 (63-80)	77.7 (68-85)	0.07	<0.001	0.087				
LVEF, %	65.9 (63-69)	70.9 (66-75)	77.7 (73-83)	78.6 (75-82)	0.001	<0.001	0.79				
MWT, mm	9.3 (8-11)	9.8 (9-11)	17.0 (15-21)	17.5 (16-22)	0.24	<0.001	0.24				
LV mass index, g/m ²	56.1 (47-61)	51.5 (43-62)	77.2 (67-92)	101.6 (82-127)	0.47	<0.001	<0.001				
LV morphology			()	()							
Isolated basal septal LVH	-	-	23 (45)	29 (55)	-	-	0.33				
Reverse septal curvature	-	-	22 (43)	15 (28)	-	-	0.11				
Concentric	-	-	0 (0)	1 (2)	-	-	0.32				
Mixed apical-ASH	-	-	6 (12)	8 (15)	-	-	0.62				
rissue characterization	26.0 (24.20)	271 (25.20)	20 4 (27 24)	26.0 (25.20)	0.70	-0.001	0.000				
ECV, %	26.9 (24-29)	27.1 (25-30)	29.4 (27-34)	26.8 (25-30)	0.79	<0.001	0.009				
LGE present			47 (92) 9 4 (2 15)	51 (96)	0.09	<0.001	0.43				
LGE mass, g	0 [0-0]	0 [0-4]	0.4 (3-13)	3.5 (3-14)	0.1	<0.001	0.32				
12-lead ECG: measures	0 [0 0]	0 [0 5]	4.5 (2 11)	5.5 (17)	0.12	.0.001	0.15				
HR beats/min	67 (60-75)	65 (57-76)	66 (56-72)	63 (57-73)	0.52	0.59	0.59				
ORS ms	90 (87-101)	86 (80-94)	96 (86-104)	98 (92-108)	0.005	<0.001	0.029				
PR ms	160 (138-175)	154 (140-170)	165 (147-181)	172 (152-196)	0.58	0.065	0.13				
OTc. ms	409 + 19	397 + 22	423 + 29	437 + 25	0.22	<0.001	0.011				
OT dispersion. ms	48 (24-62)	43 (20-62)	58 (38-77)	52 (36-68)	0.62	0.006	0.35				
Amp-Sokolow, mV	2.0 (1.5-2.2)	2.0 (1.6-2.7)	2.2 (1.8-3.1)	3.0 (2.3-4.5)	0.25	0.17	<0.001				
Amp-Cornell, mV	1.2 (0.9-1.5)	1.2 (1.0-1.6)	2.1 (1.3-2.7)	2.7 (1.9-3.1)	0.55	<0.001	0.001				
12-lead ECG: qualitative											
Abnormal ECG	2 (8)	22 (31)	41 (80)	48 (91)	0.019	<0.001	0.14				
Q waves	1 (4)	14 (20)	14 (27)	14 (26)	0.07	0.36	0.91				
TWI	0 (0)	5 (7)	34 (67)	40 (75)	0.33	<0.001	0.32				
LVH voltage criteria	0 (0)	6 (9)	19 (37)	34 (64)	0.33	<0.001	0.006				
ST-segment depression	0 (0)	0 (0)	20 (39)	30 (57)	>0.9	<0.001	0.076				
RBBB	0 (0)	0 (0)	1 (2)	3 (6)	>0.9	0.42	0.62				
LBBB	0 (0)	0 (0)	0 (0)	5 (9)	>0.9	>0.9	0.06				
Fractionation	1 (4)	3 (4)	10 (20)	18 (34)	>0.9	0.007	0.1				

Values are median (Q1-Q3), n (%), mean \pm SD. **Bold** *P* values are statistically significant. ^aThe remaining 11 healthy volunteers (HVs) (not shown here) were older to permit comparisons with the G+LVH+ group. ^bValues of LGE mass and LGE mass % for Younger HV and G+LVH- are median [range].

Amp = amplitude; ASH = asymmetric septal hypertrophy; BMI = body mass index; BSA = body surface area; ECG = electrocardiogram; ECV = extracellular volume; HR = heart rate; LBBB = left bundle branch block; LGE = late gadolinium enhancement; LVEDV = left ventricular end-diastolic volume; LVH = left ventricular hypertrophy; MWT = maximal wall thickness; QTc = corrected QT; RBBB = right bundle branch block; TWI = T-wave inversion.

Differences persisted after adjusting for age, sex, QRS duration, LGE volume, and the presence of abnormal ECG ($\beta = 0.42$ [95% CI: 0.2-0.6]; P < 0.001). Compared with HVs, G+LVH– with a normal 12-lead ECG (n = 48) also had slower ventricular conduction

(prolonged mean AT [39 ms (Q1-Q3: 35-44 ms) vs 35 ms (Q1-Q3: 31-41 ms)]; P = 0.009). There were no differences in repolarization duration (mean RTc/ARIc) or activation/repolarization dispersion (Δ AT/ Δ RTc). G+LVH– had more spatially heterogenous

TABLE 2 ECGI Parameters in HV, Subclinical HCM, G+LVH+, and G-LVH+											
					P Value						
	Younger HV (n = 26)	G+LVH- (n = 70)	G+LVH+ (n = 51)	G–LVH+ (n = 53)	G+LVH– vs HV	G+LVH+ vs G+LVH-	G-LVH+ vs G+LVH+				
Conduction											
Mean amplitude, mV	1.4 (1.1-1.6)	1.4 (1.2-1.7)	1.6 (1.2-1.9)	1.9 (1.4-2.6)	0.4	0.42	0.12				
Fractionation (n/1,000)	8 (0-22)	8 (1-20)	13 (2-27)	2 (0-10)	0.99	0.13	0.002				
Mean AT, ms	35 (31-41)	39 (35-45)	41 (37-45)	42 (38-47)	0.008	0.23	0.27				
ΔAT, ms	174 (153-196)	177 (154-196)	183 (162-208)	188 (159-205)	0.89	0.25	0.76				
Mean G _{AT} , ms/mm	0.42 (0.37-0.47)	0.40 (0.31-0.49)	0.40 (0.35-0.46)	0.44 (0.30-0.56)	0.38	0.82	0.33				
Max G _{AT} , ms/mm	4.7 (4.2-6.0)	4.9 (4.2-5.6)	5.1 (4.2-5.8)	5.1 (4.3-6.4)	0.74	0.54	0.27				
Repolarization											
Mean ARIc, ms	247 ± 18	245 ± 26	276 ± 29	281 ± 24	0.78	<0.001	0.29				
Δ ARIc, ms	180 (159-197)	181 (161-202)	183 (162-208)	193 (162-213)	0.62	0.62	0.63				
Mean RTc, ms	285 (274-297)	283 (273-292)	322 (296-340)	327 (305-345)	0.55	<0.001	0.21				
ΔRTc , ms	160 (145-179)	164 (147-183)	165 (150-178)	170 (149-193)	0.48	0.73	0.47				
Mean G _{RTc} , ms/mm	1.0 ± 0.2	1.1 ± 0.3	1.0 ± 0.3	1.0 ± 0.3	0.042	0.13	0.79				
Max G _{RTc} , ms/mm	9.5 (9.0-11)	11.2 (10-12.9)	11.3 (9.9-12.7)	11.1 (9.3-14.0)	0.005	0.79	0.92				

Values are median (Q1-Q3) or mean \pm SD. **Bold** *P* values are statistically significant.

 Δ = dispersion; Amp = amplitude; ARIc = activation recovery interval corrected; AT = activation time; G = gradient; max = maximum; RTc = repolarization time corrected for heart rate; other abbreviations as in Table 1.

repolarization (steeper G_{RTc} , occurring in 23% [11 of 48] of those with normal ECGs) (Figure 3, Table 2) but similar G_{AT} . There were no differences in signal amplitudes or fractionation.

G + LVH + vs G + LVH -. Compared with G + LVH -, G+LVH+ had prolonged ventricular repolarization (elevated mean ARIc and RTc), and differences persisted after adjusting for age, sex, LGE volume, and QTc (ARIc: $\beta = 0.31$ [95% CI: 0.2-0.5]; P < 0.001; RTc: $\beta = 0.28$ [95% CI: 0.2-0.5]; P < 0.001). All other ECGI parameters were similar (Table 2, Figure 2). Further analyses comparing HVs matched to G+LVH+ are included in the Supplemental Results. G - LVH + vs G + LVH +. Compared with G - LVH +, G+LVH+ had more signal fractionation (P = 0.002) (Table 2, Figure 4), and differences persisted after adjusting for age, sex, LGE volume, and ECG fractionation ($\beta = 0.39$ [95% CI: 0.2-0.6]; P < 0.001). All other global and local ECGI parameters were similar (Table 2, Figure 2).

ECG VS ECGI. Across all participants, AT was associated with QRS duration ($r_{\rm s} = 0.26$ [95% CI: 0.1-0.4]; P < 0.001). Mean ARIc and RTc were both associated with QTc (mean ARIc: $r_{\rm s} = 0.73$ [95% CI: 0.7-0.8]; P < 0.001; mean RTc: $r_{\rm s} = 0.73$ [95% CI: 0.7-0.8]; P < 0.001) and QT dispersion (ARIc: $r_{\rm s} = 0.16$ [95% CI: 0.01-0.30]; P = 0.03; RTc: $r_{\rm s} = 0.2$ [95% CI: 0.05-0.30]; P = 0.008). Δ RTc was associated with QT dispersion ($r_{\rm s} = 0.17$ [95% CI: 0.01-0.30]; P = 0.03).

ECG STRUCTURAL RELATIONSHIPS. 12-lead ECG. In G+LVH-, MWT associated with QRS duration

 $(r_{\rm s} = 0.42 [95\% \text{ CI: } 0.2-0.6]; P < 0.001)$, PR interval $(r_{\rm s} = 0.31 [95\% \text{ CI: } 0.06-0.50]; P = 0.012)$, and Cornell but not Sokolow-Lyon amplitude $(r_{\rm s} = 0.38 [95\% \text{ CI: } 0.1-0.6]; P = 0.002)$. In overt HCM (all LVH+), MWT associated with Cornell amplitude $(r_{\rm s} = 0.34 [95\% \text{ CI: } 0.1-0.5]; P < 0.001)$, presence of an abnormal ECG $(r_{\rm s} = 0.29 [95\% \text{ CI: } 0.1-0.5]; P = 0.019)$, and TWI $(r_{\rm s} = 0.26 [95\% \text{ CI: } 0.07-0.40]; P = 0.007)$. Also, LGE associated with PR interval $(r_{\rm s} = 0.26 [95\% \text{ CI: } 0.06-0.40]; P = 0.009)$, the presence of an abnormal ECG $(r_{\rm s} = 0.39 [95\% \text{ CI: } 0.2-0.5]; P < 0.001)$, and TWI $(r_{\rm s} = 0.40 [95\% \text{ CI: } 0.2-0.6]; P < 0.001)$.

ECGI RELATIONSHIPS TO RISK MARKERS. Maximal wall thickness. MWT associated with signal amplitude in HV ($r_s = 0.42$ [95% CI: 0.3-0.7]; P = 0.032), G+LVH- ($r_s = 0.26$ [95% CI: 0.02-0.50]; P = 0.028), and overt HCM ($r_s = 0.33$ [95% CI: 0.1-0.5]; P < 0.001). In G+LVH-, MWT associated with mean ARIC ($r_s = 0.24$ [95% CI: 0.003-0.50]; P = 0.046) and RTc ($r_s = 0.28$ [95% CI: 0.04-0.50]; P = 0.019). In overt HCM, MWT associated with mean AT ($r_s = 0.25$ [95% CI: 0.05-0.40]; P = 0.011), Δ AT ($r_s = 0.32$ [95% CI: 0.1-0.5]; P = 0.001), Δ ARIC ($r_s = 0.36$ [95% CI: 0.2-0.5]; P < 0.001), and mean RTc ($r_s = 0.24$ [95% CI: 0.04-0.40]; P = 0.015).

Scar burden. In overt HCM, LGE volume associated with fractionation ($r_s = 0.21$ [95% CI: 0.01-0.40]; P = 0.032) and local AT gradients (G_{ATmean} : $r_s = 0.27$ [95% CI: 0.08-0.40]; P = 0.005).

Left ventricular outflow tract obstruction. In total, 24 (23%) of overt HCM patients had LVOTO. No ECGI changes were associated with LVOTO.



SCD risk. To determine the relationship between ECGI abnormalities and surrogate markers of ventricular arrhythmia and risk in overt HCM, we describe a subgroup of 19 participants (18% of overt HCM) who had an intermediate/high SCD risk score or NSVT. Compared with overt HCM not meeting such criteria, these patients were similar in age (P = 0.29), sex (P = 0.74), and ethnicity (White: P = 0.80), but had a higher MWT (19.6 mm [Q1-Q3: 17-23 mm] vs

17.0 mm [Q1-Q3: 15-21 mm]; P = 0.005), more LGE (15.1 g [Q1-Q3: 8-28 g] vs 5.5 g [Q1-Q3: 2-13 g]; P < 0.001), and more spatially heterogenous conduction ($G_{ATmax} P = 0.007$). Other ECGI parameters were similar. After adjustment for MWT and LGE volume, those with NSVT/intermediate- or high-risk score continued to exhibit more spatially heterogenous conduction ($\mathcal{B} = 0.29$ [95% CI: 0.1-0.5]; P = 0.006). An exploratory analysis was performed



for all those with NSVT (n = 16) showing similar findings (Supplemental Results).

MACHINE LEARNING CLASSIFICATION OF ECGI HCM SUBTYPES. The SVM differentiated subclinical HCM from HV with an AUC of 0.96 (bootstrap 95% CI: 94%-98%; sensitivity 100% [95% CI: 93%-100%], specificity 91% [95% CI: 76%-98%], positive predictive value 96% [95% CI: 87%-99%], negative predictive value 100% [95% CI: 87%-100%], balanced accuracy 95.7%) (Supplemental Tables 5 to 7, Supplemental Figure 1) and an accuracy of 80% after 10-fold cross-validation (95% CI: 73%-85%). This ECGI biomarker panel was able to identify HCM patients at greater risk of SCD (because of prior NSVT or intermediate/high ESC SCD risk status) compared with low-risk patients, with an AUC of 0.97 (bootstrap 95% CI: 96%-98%; sensitivity 94% [95% CI: 71%-99%], specificity 100% [95% CI: 95%-100%], positive predictive value 100% [95% CI: 77%-100%], negative predictive value 95% [95% CI: 93%-99%], and balanced accuracy 97.2%) (Supplemental Tables 5 to 8) and an accuracy of 82% after 10-fold cross-validation (95% CI: 78%-86%).

DISCUSSION

This is a prospective multicenter study using CMRguided ECGI to simultaneously characterize structural and EP parameters across the spectrum of genotyped HCM. The main findings are as follows: 1) EP abnormalities exist in individuals with subclinical HCM, including in those with normal resting 12-lead ECGs; and 2) in overt HCM, EP abnormalities occur more severely with adverse cardiac structural features and positive genetic status (**Central Illustration**). There is well-described evidence that such EP abnormalities, consisting of spatially heterogenous repolarization, slowed discontinuous ventricular conduction, and signal



fractionation, are proarrhythmic,^{10,11,26,27} thus potentially affecting patient-specific risks for SCD in HCM. This is the largest clinical ECGI study ever performed and the only one to date to have studied subclinical HCM.

As novel HCM therapies show promise in disease modification with some suggestion that early expression of sarcomeric gene variants are potentially reversible,^{28,29} the discovery of sensitive biomarkers for early disease is an emerging priority. Structural changes adopted in modern-day HCM risk stratification (LA dilation, severe LVH, LVOTO, high LGE burden) tend to occur late. Furthermore, NSVT is the only EP characteristic considered in current risk stratification algorithms but has the highest predictive value for SCD; it is, however, uncommon in early disease and is a dichotomous risk marker.¹⁷ Here we show quantifiable EP abnormalities occurring in the absence of standard 12-lead ECG changes or measurable hypertrophy or fibrosis. Although cascade genetic screening allows the identification of a larger number of individuals at risk, HCM is characterized by variable penetrance and disease expression so detecting phenotype development is a priority. Due to the complexity of a 12-biomarker ECGI panel, a machine learning approach was used for discrimination between patient groups. This showed the potential of ECGI to discriminate subclinical HCM from healthy individuals and identify patients with HCM at higher risk of SCD. Future work will need to externally validate these discovered ECGI biomarker panels to understand their real-world diagnostic and prognostic potential.

Notably, EP abnormalities occurring in G+LVH– and G+LVH+ differed, likely reflecting disease heterogeneity and different phenotypic stages of disease. Longitudinal studies relating EP abnormalities to events are now possible using this more sustainable, time and cost-effective, and reuseable ECGI methodology. This may lay the foundation for ECGIbased risk stratification and provide support for decisions surrounding exercise prescription.

Ventricular conduction was slower in G+LVH– compared with HVs (increased AT) despite the absence of CMR markers of fibrosis and LVH (scar, ECV and MWT were similar to health). Importantly this occurred in a cohort predominantly without bundle branch block (2% in G+LVH+, none in G+LVH–). Furthermore, slowed ventricular conduction is missed by the 12-lead ECG occurring in both G+LVH– with normal and abnormal 12-lead ECG. In overt disease, ventricular conduction was not globally slower compared with matched HV (similar AT) but more regionally slowed (resulting in increased Δ AT) (Supplemental Results). This finding is consistent with invasive endocardial mapping findings in



nonsustained ventricular tachycardia. Severity of left ventricular hypertrophy (LVH) associated with regionally slowed conduction and prolonged repolarization.

overt HCM which showed lateral LV activation before the septum, longer action potential durations, prolonged stimulus-to-V times and reduced action potential upstroke velocity on patch-clamp tests of HCM myocardium.^{5,30} Mechanisms of ventricular conduction slowing in HCM are likely to include decreased electrical coupling, discontinuous propagation, and reduced conduction velocity. Our discovery that these occur in the absence of hypertrophy and fibrosis in subclinical HCM challenges the LVH-centric view of EP abnormalities and raises the possibility that electrical changes may be more closely related to myocyte disarray, ischemia, or electromechanical factors than actual LVH.^{28,31} However, a compounding effect of LVH is observed in overt disease, as demonstrated by the association of MWT and both activation time and dispersion. Slowed discontinuous ventricular conduction and local conduction disturbances are a well-known proarrhythmic prerequisite for ventricular arrhythmia initiation.^{10,11} Our findings provide a possible reason behind the increased incidence of ventricular arrhythmia observed with greater degrees of LVH.^{1,17,32} Slower ventricular conduction has been detected by ECGI in other disease states with primary structural substrate changes including arrhythmogenic cardiomyopathy (AC) and cardiac amyloidosis, and overlying the right ventricular outflow tract epicardium in Brugada syndrome (although this is primarily a channelopathy, epicardial focal fibrosis has also been observed on autopsy).⁸⁻¹⁰ Furthermore, in aborted SCD survivors and Brugada syndrome, ECGI has demonstrated greater conduction heterogeneity postexercise, further demonstrating its ability to detect abnormal pro-arrhythmic rate adaptation.³³

Repolarization in subclinical HCM was more spatially heterogenous (steeper repolarization gradients) when detected by ECGI, but globally, repolarization was largely normal on ECG (normal QTc, low prevalence of TWI/no ST-segment depression). Relationships between early LVH and prolonged repolarization can be observed in subclinical HCM where a positive relationship existed between MWT and ARIc. In overt disease where LVH is established, prolonged repolarization is detected in both ECG (elevated QTc) and ECGI (ARIc, RTc). In HCM mechanisms of repolarization abnormalities are likely related to ionic remodeling, which can be heterogenous because of LVH, calcium handling and sensitivity changes, which can result in action potential prolongation with early and late afterdepolarization leading to ventricular arrhythmia in structurally remodeled (hypertrophy, fibrosis, disarray) and therefore susceptible myocardium.^{30,31,34} The exaggerated spatially heterogenous repolarization noted in the absence of hypertrophy could be evidence of cellular and molecular changes occurring in response to sarcomeric mutation or prehypertrophic myocardial substrate changes.^{28,31} Repolarization gradients as seen in subclinical HCM have been observed in ventricular fibrillation survivors with structurally normal hearts.^{27,33} Spatially heterogenous repolarization is known to be proarrhythmic by supporting the conditions for asymmetric excitability and propagation of re-entry. This abnormality has also been demonstrated by ECGI in multiple arrhythmogenic diseases, including Brugada, AC, long QT syndrome, and early repolarization, and hypothesized in heart failure.9,10,26,35 Other ECG techniques quantifying repolarization abnormality in overt HCM include dynamic QTc changes on Holter monitoring, prolonged QTc as a predictor of ICD discharge, and QT dispersion; however, none have been especially useful in HCM, limited by lack of spatial information, technical limitations, and bias toward more advanced disease.36,37 With QTc being normal in subclinical disease, it is unlikely to afford benefits as an early disease biomarker.

Scar volume was related to spatial conduction heterogeneity (activation gradients and fractionation). This demonstrates the ability of ECGI to detect and quantify preclinical conduction discontinuities resulting from scar (not detected solely by LGE CMR). Conduction discontinuities most likely occur because of islands of viable tissue trapped within fibrotic tissue. Indeed, the fact that myocytes are electrotonically uncoupled will also serve to exaggerate cellular repolarization differences and increase repolarization gradients.38 Spatial conduction heterogeneity also associated with NSVT and SCD risk score independent of scar, showing the incremental value of ECGI for detecting proarrhythmic features. This EP abnormality has potential for arrhythmia formation^{3,9,11,38} through asymmetric loading on a propagating wave front favoring unidirectional block, creating dispersion of repolarization and facilitating re-entry. Both spatial conduction heterogeneity and fractionation are observed in AC scar, and fractionation is observed in both infarct and AC scar.^{9,11}

Despite similar MWT and LGE burden and a more benign 12-lead ECG phenotype, ECGI-detected fractionation was greater in genotype-positive HCM patients compared with G-LVH+. Greater fractionation in G+LVH+ could not be accounted for by bundle branch block, suggesting a genuinely greater area of tissue with fractionated electrograms. Exaggerated spatially heterogenous ventricular conduction is another plausible contributor to the greater ventricular arrhythmia seen in G+LVH+ HCM compared with G-LVH+.² Unexpectedly, we found more severe 12lead ECG abnormalities in G-LVH+ compared with G+LVH+ HCM (longer QRS duration, longer QTc, and greater QRS amplitudes). This mirrors data from a prior study showing G-LVH+ was characterized by greater lateral T-wave inversion.³⁹ Overall differences in ECG and ECGI between G- and G+ HCM could be in part related to different LV morphologies (greater LV mass despite similar MWT) and different tissue characterization (lower ECV) as described by large registry studies.^{2,40}

STUDY LIMITATIONS. ECGI provides epicardial EP mapping data only and does not therefore measure potentially important transmural electrophysiology; it also misses EP phenomena occurring within the septum. As expected, participants with subclinical HCM were 16 years younger than overt disease counterparts and medication use affects electrophysiology and may influence results. However, our sensitivity analyses (Supplemental Results) showed consistent EP changes even when comparing overt HCM and drug-free HCM to age-matched HVs.

Although ECGI-detected EP abnormalities found in our cross-sectional study have proven associations with ventricular arrhythmia in other diseases and show the potential to identify patients at greater risk of malignant arrhythmias, hold-out data set internal validation of the SVM was not possible because of the limited sample size. To mitigate this, we undertook 10-fold cross validation, which showed good accuracy of the combined panel. Longitudinal multicenter studies are now needed to externally validate these results and confirm their generalizability. No HV underwent genetic testing. G+LVH- subjects here were recruited in the context of familial evaluation, and these findings do not apply to those with pathogenic sarcomere variants as secondary genomic findings.

CONCLUSIONS

In the absence of LVH or abnormalities on the 12-lead ECG, HCM sarcomere gene mutation carriers express an aberrant electrophysiological phenotype detected by ECGI. In overt HCM, electrophysiological abnormalities occur more severely with adverse structural change and positive genetic status.

ACKNOWLEDGMENT Source code for ML is available via GitHub (see Supplemental Statistical Methods).

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PERSPECTIVES

COMPETENCY IN PATIENT CARE AND

PROCEDURAL SKILLS: In patients with HCM, CMRguided electrocardiographic mapping can detect pathogenic sarcomere variants even in the absence of myocardial hypertrophy or abnormalities on the standard 12-lead ECG.

TRANSLATIONAL OUTLOOK: Abnormalities detected by this method could identify patients with subclinical disease who might benefit from therapeutic intervention.

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KEY WORDS cardiac magnetic resonance imaging, ECG imaging, electrophysiology, hypertrophic cardiomyopathy

APPENDIX For expanded Methods and Results section, as well as supplemental tables and a figure, please see the online version of this paper.