Relationship Between Electrocardiographic Findings and Cardiac Magnetic Resonance Phenotypes in Arrhythmogenic Cardiomyopathy

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Background—The new designation of arrhythmogenic cardiomyopathy defines a broader spectrum of disease phenotypes, which include right dominant, biventricular, and left dominant variants. We evaluated the relationship between electrocardiographic findings and contrast-enhanced cardiac magnetic resonance phenotypes in arrhythmogenic cardiomyopathy.

Methods and Results—We studied a consecutive cohort of patients with a definite diagnosis of arrhythmogenic cardiomyopathy, according to 2010 International Task Force criteria, who underwent electrocardiography and contrast-enhanced cardiac magnetic resonance. Both depolarization and repolarization electrocardiographic abnormalities were correlated with the severity of dilatation/dysfunction, either global or regional, of both ventricles and the presence and regional distribution of late gadolinium enhancement. The study population included 79 patients (60% men). There was a statistically significant relationship between the presence and extent of T-wave inversion across a 12-lead ECG and increasing values of median right ventricular (RV) end-diastolic volume (P<0.001) and decreasing values of RV ejection fraction (P<0.001). The extent of T-wave inversion to lateral leads predicted a more severe RV dilatation rather than a left ventricular involvement because of the leftward displacement of the dilated RV, as evidenced by contrast-enhanced cardiac magnetic resonance. A terminal activation delay of >55 ms in the right precordial leads (V1-V3) was associated with higher RV volume (P=0.014) and lower RV ejection fraction (P=0.053). Low QRS voltages in limb leads predicted the presence (P=0.004) and amount (P<0.001) of left ventricular late gadolinium enhancement.

Conclusions—The study results indicated that electrocardiographic abnormalities predict the arrhythmogenic cardiomyopathy phenotype in terms of severity of RV disease and left ventricular involvement, which are among the most important determinants of the disease outcome. (J Am Heart Assoc. 2018;7:e009855. DOI: 10.1161/JAHA.118.009855.)

Key Words: cardiac magnetic resonance imaging • cardiomyopathy • electrocardiography • late gadolinium enhancement

Arrhythmogenic cardiomyopathy (ACM) is an inherited heart disease characterized by fibrofatty myocardial replacement, which may act as a substrate for life-threatening ventricular arrhythmias and cause progressive ventricular systolic dysfunction.1-3 Previously referred to as arrhythmogenic right ventricular (RV) cardiomyopathy because of the predominant involvement of the RV in the originally reported phenotype, the recognition of biventricular and left-dominant disease variants has led, in recent years, to use of the current designation of ACM, which better reflects the broader phenotypic spectrum of the disease.2-5

Conventionally, the diagnosis of ACM relies on clinical criteria that encompass electrocardiographic repolarization and depolarization abnormalities, arrhythmic events, mor-phofunctional ventricular alterations, histopathologic changes, and familial genetic background.3 A 12-lead ECG is an integral part of the evaluation of affected patients, and depolarization/repolarization abnormalities are commonly observed, with a reported prevalence of up to 85%.6 The most striking electrocardiographic abnormalities in patients with ACM include T-wave inversion in anterior or anterolateral leads; prolongation of right precordial QRS duration with a delayed S-wave upstroke (terminal activation delay, >55 ms); QRS fragmentation and postexcitation ε waves (ie, small-amplitude potentials occurring at the end of the QRS complex/beginning of the ST segment); and low QRS voltages in limb leads.3,6 The underlying structural and functional ventricular abnormalities and the clinical and

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Clinical Perspective

What Is New?

- We studied the relationship between electrocardiographic abnormalities and contrast-enhanced cardiac magnetic resonance findings in a cohort of patients with a definite diagnosis of arrhythmogenic cardiomyopathy, according to 2010 International Task Force criteria.
- We found that routine electrocardiography has the potential to predict the disease phenotype, with particular reference to the severity of right ventricular disease and the left ventricular involvement, which are among the most important clinical determinants of the disease outcome.
- Specifically, extent of T-wave inversion and prolonged terminal activation delay predicted higher right ventricular dilation and lower right ventricular ejection fraction, whereas low QRS voltages in limb leads indicate the presence of late gadolinium enhancement/myocardial fibrosis of the left ventricular wall.

What Are the Clinical Implications?

- The study indicates that routine electrocardiography provides both diagnostic and prognostic information in patients with arrhythmogenic cardiomyopathy.
- According to our study results, the electrocardiographic pattern of low QRS voltages in limb leads, which was shown to be a more accurate predictor of left ventricular involvement than T-wave inversion in the inferolateral leads, should be included among the International Task Force criteria for diagnosis of biventricular arrhythmogenic cardiomyopathy.

Methods

Study Population

The data, analytic methods, and study materials will be available to other researchers for purposes of reproducing the results or replicating the procedure on reasonable request. The study included 79 consecutive patients with a diagnosis of ACM, according to the 2010 International Task Force (ITF) criteria, who underwent CE-CMR from July 2006 to March 2016. In patients enrolled before 2010, the diagnosis was established according to the original 1994 ITF criteria and confirmed using the 2010 revised criteria. CE-CMR findings were not considered for diagnosis. The study was approved by an institutional review committee, and the subjects gave informed consent.

Electrocardiographic Evaluation

A standard 12-lead ECG was recorded at the time of CE-CMR study and evaluated using a digital procedure. The electrocardiographic tracing was interpreted by 2 observers (A.Z. and B.B.) who were unaware of clinical data and CE-CMR findings. Ambiguous cases were reviewed by a third expert (D.C.). Standard measurements included heart rate; rhythm; PR interval; QRS axis, voltage, and duration; and ST-segment and T-wave abnormalities. The following specific electrocardiographic parameters were evaluated: (1) postexcitation waves, defined as small-amplitude potentials occurring at the end of the QRS complex/beginning of the ST segment; (2) terminal activation duration (TAD), defined as prolongation of the terminal part of the QRS (from the nadir of the S wave to the end of QRS) >55 ms in leads V1 to V3, in the absence of complete right bundle branch block; (3) low (<0.5-mV) QRS voltages in limb leads, including both negative and positive components; (4) T-wave inversion (TWI) ≥0.1 mV in depth in ≥2 contiguous leads. According to the presence and extent of TWI across the 12 leads, 3 subgroups were defined: (1) absence of TWI in right precordial leads (V1-V3); (2) TWI limited to right precordial leads (V2-V3); and (3) TWI in right precordial leads, which extended to left precordial (V4-V6) or inferior (II, III, or aVF) leads. The presence of late gadolinium enhancement/myocardial fibrosis of the left ventricular wall.

Cardiac Magnetic Resonance

Acquisition protocols

CMR was performed on a 1.5-T scanner (Magnetom Avanto; Siemens Healthcare, Germany) using a comprehensive dedicated protocol. All patients underwent detailed CE-CMR; study protocol included postcontrast sequences. Biventricular morphofunctional assessment was performed by a set of steady-state free precession sequence cine loops in sequential short-axis views (slice thickness, 6 mm; gap, 0 mm;
repetition time, 2.5–3.8 ms; echo time, 1.1–1.6 ms; average in-plane resolution, 1.5 × 2.4 mm; flip angle, 45°–60°; temporal resolution, 40–45 ms) and long-axis views (2-, 3-, and 4-chamber views) of the LV from the semilunar valve plane through the apex. The RV was evaluated with balanced transaxial Steady-state free precession cine images from the outflow tract to the diaphragm, with a 2-chamber view. After IV administration of contrast agent (gadobenate dimeglumine, Multihance; Bracco; 0.2 mmol/kg of body weight), 2-dimensional segmented fast low-angle shot inversion recovery sequences, after 10 minutes, were acquired in the same views of the cine images, covering the entire ventricles (repetition time, 5.4–8.3 ms; echo time, 1.3–3.9 ms; average in-plane spatial resolution, 1.4–1.5 × 2.2–2.4 mm; 6-mm slice thickness; 2-mm gap; and flip angle, 20°–25°). Inversion times were adjusted to null normal myocardium using a Look-Locker sequence, and images were repeated in 2 separate phase-encoding directions to exclude artefacts.

Ventricular size and systolic function analysis

Global ventricular volumes and systolic function were calculated from the short-axis cine images, excluding papillary muscles from the myocardium, using a computer-aided analysis package (CMR42; Circle International, Calgary, Alberta, Canada). Ventricular volumes were indexed by body surface area. Analysis of RV global and regional wall motion abnormalities (akinesia or dyskinesia) was based on 5 RV segments: inferior wall, anterolateral wall, RV outflow tract, apex, and inflow tract. Analysis of LV global and regional wall motion abnormalities was based on the 17-segment model proposed by the American Heart Association.

Late gadolinium enhancement assessment

The presence and regional distribution of late gadolinium enhancement (LGE) were visually assessed independently by 2 experienced observers (M.D.L. and B.G.) who were blinded to patient clinical data. Ambiguous cases were reviewed by a third expert (M.P.M.). To exclude artefacts, LGE was deemed present if only visible in 2 orthogonal views (short- and long-axis views). Areas of LGE were allocated to the American Heart Association 17-segment model for the LV and to the 5 regions of the RV.

Displacement of RV toward lateral precordium

Assessment of the degree of lateral displacement of the RV apex toward the left axilla was performed according to a previously reported protocol. In brief, in the long-axis 4-chamber view, the cardiac apex angle was calculated as the angle of the interventricular septum with respect to the thoracic midline; the ratio of lateral displacement was calculated as the ratio between the distance from line A (passing through the midpoint of the sternum and thoracic vertebrae) to the cardiac apex along the internal thoracic cage and the distance from the cardiac apex to line B (orthogonal to line A and passing through the left axilla).

Statistical Analysis

Data are expressed as median with 25th to 75th percentiles because normality could not be assumed for any variable. Categorical differences among groups were evaluated by the χ² test or the Fisher exact test, as appropriate. Differences among continuous variables were compared using the Mann-Whitney U test or the Kruskal-Wallis test, as appropriate. A 2-tailed probability value of 0.05 was considered statistically significant. The interobserver reliability analysis for electrocardiographic findings and CMR qualitative parameters was measured by Cohen’s κ coefficient. All analyses were performed using SPSS 23 (SPSS Inc, Chicago, IL).

Results

Baseline Clinical and Electrocardiographic Characteristics

The study sample included 79 patients (60% men; median age, 33 years), whose baseline clinical and electrocardiographic characteristics are summarized in Table 1. A normal ECG was found in 7 patients (9%), whereas the others had ≥1 repolarization and depolarization abnormalities. In 18 subjects (23%), there were no TWIs in right precordial leads (V1-V3); in 11 subjects (14%), TWIs were confined to V1 to V3; in 50 subjects (63%), TWIs extended to other leads (to the left precordial leads V4-V6 in 16 [20%]; to the inferior leads II, III, and aVF in 8 [10%]; and to both the left precordial and inferior leads in 26 [33%]). Thirteen patients (16%) exhibited postexcitation “ε” waves. After excluding 12 patients with a right bundle branch block pattern, 27 of the remaining 67 patients (40%) had a prolonged TAD (>55 ms). Low QRS voltages in limb leads were observed in 16 patients (20%). The interobserver reliability for TWI assessment and low voltages in limb leads was κ = 1 (95% confidence interval [CI], 1.000–1.000; P = 0.906 [95% CI, 0.802–1.000; P < 0.001]); and for the ε wave, κ = 0.816 (95% CI, 0.641–0.990; P < 0.001).

CMR Findings

Detailed CE-CMR findings are reported in Table 1. At CE-CMR, 61 patients (77%) showed RV dilatation (median RV end-diastolic volume, 109 mL/m²) and 37 (47%) showed RV dysfunction (median RV ejection fraction, 47%), according to the cutoff proposed by the 2010 ITF criteria. A dilated LV was found in 33 subjects (42%), and a reduced LV ejection fraction.
was found in 22 subjects (29%). At postcontrast sequences, LGE was found in 71 patients (90%): the RV was involved in 14 patients (18%), the LV was involved in 16 patients (20%), and both ventricles were involved in 41 patients (52%). In all patients with RV LGE, there was an associated regional wall motion impairment with akinesia or dyskinesia. In all patients with LV LGE a nonischemic pattern of distribution was shown, involving the subepicardial/midmural layers and sparing the subendocardium. The interobserver reliability for the LV LGE was \( j = 0.907 \) (95% CI, 0.803–1.000; \( P < 0.001 \)); and for the RV LGE, \( j = 0.798 \) (95% CI, 0.655–0.941; \( P < 0.001 \)).

Comparison Between Electrocardiographic and CMR Findings

The relationship between electrocardiographic and CE-CMR findings is summarized in Tables 2 through 5. The presence of postexcitation \( \varepsilon \) waves did not correlate with RV and LV dilatation/dysfunction or with LGE (Table 2). Patients with a prolonged TAD (>55 ms in V1-V3) had a larger RV end-diastolic volume (116 versus 99 mL/m\(^2\); \( P=0.014 \)) and a lower RV EF (41% versus 47%; \( P=0.226 \)). LV EF also decreased in patients with prolonged TAD (38% versus 44%; \( P=0.228 \)).

Table 1. Clinical Characteristics of the Overall Sample (n=79)

| Characteristics | Value |
|-----------------|-------|
| Age, y          | 33 (20–48) |
| Male sex        | 47 (60) |
| Family history of sudden death | 13 (16) |
| Clinical symptoms |      |
| Chest pain      | 5 (6) |
| Palpitations    | 42 (53) |
| Syncope         | 5 (6) |
| Dyspnea         | 8 (10) |
| International Task Force criteria |      |
| 2 Major criteria | 42 (53) |
| 1 Major and 2 minor criteria | 33 (42) |
| 4 Minor criteria | 4 (5) |
| Electrocardiographic characteristics |      |
| Normal ECG      | 7 (9) |
| Depolarization abnormalities |      |
| Postexcitation \( \varepsilon \) wave | 13 (16) |
| Prolonged terminal activation duration (>55 ms) | 27/67 (40) |
| Low (<0.5-mV) QRS voltages in limb leads | 16 (20) |
| Repolarization abnormalities |      |
| No TWI in right precordial leads (V1-V3) | 18 (23) |
| TWI in right precordial leads (V1-V3) only | 11 (14) |
| TWI in right precordial (V1-V3) plus left precordial (V4-V6) or inferior (II, III, and aVF) leads | 50 (63) |
| Cardiac magnetic resonance features |      |
| Functional analysis |      |
| RV EDV, mL/m\(^2\) | 109 (94–120) |
| RV EF, %        | 47 (37–56) |
| No. of RV segments with WMA | 2 (1–4) |
| RV volume*      |      |
| No dilatation   | 18 (23) |
| Minor criteria dilatation | 19 (24) |
| Major criteria dilatation | 42 (53) |
| RV systolic function |      |
| RV EF >45%      | 42 (53) |
| RV EF >40%<45%  | 11 (14) |
| RV EF <40%      | 26 (33) |
| LV EDV, mL/m\(^2\) | 85 (73–95) |
| LV EF, %        | 58 (50–63) |
| LV dilatation (≥90 mL/m\(^3\)) | 33 (42) |
| LV dysfunction (≥50%) | 22 (29) |

Table 2. Relationship Between Postexcitation \( \varepsilon \) Wave and CMR Parameters

| Parameters                        | \( \varepsilon \) Wave |
|----------------------------------|-----------------------|
|                                  | No (n=66) | Yes (n=13) | \( P \) Value |
| RV EDV, mL/m\(^2\)              | 108 (94–118) | 109 (98–150) | 0.315 |
| RV EF, %                         | 49 (37–58) | 41 (35–49) | 0.226 |
| RV WMA, no. of segments          | 2 (1–4) | 2 (1–4) | 0.968 |
| RV LGE                           | 45 (68) | 10 (77) | 0.744 |
| LV EDV, mL/m\(^2\)              | 86 (73–95) | 85 (70–105) | 0.802 |
| LV EF, %                         | 59 (51–62) | 51 (44–64) | 0.228 |
| LV LGE                           | 46 (70) | 11 (85) | 0.334 |

Values are expressed as number (percentage) of patients or median (25th–75th percentile). CMR indicates cardiac magnetic resonance; EDV, end-diastolic volume; EF, ejection fraction; LGE, late gadolinium enhancement; LV, left ventricular; RV, right ventricular; WMA, wall motion abnormality.
lower RV ejection fraction (41% versus 50%; \( P=0.053 \)) than those without a prolonged TAD. There was no association between prolonged TAD and LV dimension, function, or LGE (Table 3). The relationship between the presence and extent of right precordial TWI and CMR parameters is shown in Table 4. There was a statistically significant trend toward increasing RV end-diastolic volume (92, 94, and 117 mL/m², respectively; \( P=0.001 \)) and decreasing RV ejection fraction (57%, 49%, and 41%, respectively; \( P=0.001 \)) among patients with no TWI, with TWI confined to the right precordial leads, and with TWI extending to the left precordial and/or inferior leads (Figure 1). The degree of lateral displacement of the RV apex toward the left axilla was also related to the presence and extent of right precordial TWI (Figure 2). The median number of RV segments showing regional wall motion abnormalities (1, 2, and 3, respectively; \( P<0.001 \)) and the presence of RV LGE (50%, 46%, and 82%, respectively; \( P=0.007 \)) correlated with the presence and extent of right precordial TWI. Instead, none of the LV parameters differed among the 3 groups. Two patients had TWI in both inferior leads (L1, L2, and aVF) and lateral precordial leads (V4-V6), but none in right precordial leads (V1-V3). Both patients had a slight dilation and systolic dysfunction of the LV with evidence of LGE; one also showed low QRS voltages in limb leads. The RV was normal in one patient and appeared slightly dilated in the other.

The presence of low QRS voltages in limb leads was associated with the presence of LV LGE (65% versus 100%; \( P=0.004 \)) but not with RV LGE (68% versus 75%), reaching a high specificity (100%) with a low sensitivity (28%) (Figure 3). Among the 57 of 79 patients with LV LGE, those with low QRS voltages showed a higher median number of affected LV segments (6 [5–9] versus 3 [2–7]; \( P<0.001 \)). No other functional and structural CMR parameters correlated with low QRS voltages (Table 5).

**Discussion**

A 12-lead ECG is a fundamental part of the evaluation of patients with ACM. Inverted T waves in right precordial leads, delayed S-wave upstroke, e waves, low-QRS voltages in limb leads, and intraventricular conduction defects are the most striking electrocardiographic abnormalities in patients with ACM with an overt disease phenotype.\(^1\)\(^2\)\(^15\) Accordingly, this study showed that a surface 12-lead ECG was abnormal in 91% of patients with ACM who had a diagnosis of definitive ACM, according to the 2010 ITF criteria.\(^3\) The present study was designed to investigate the relationship between electrocardiographic abnormalities and structural/functional remodeling of both ventricles on the basis of CE-CMR findings in patients with ACM. The main results were that electrocardiographic repolarization and depolarization abnormalities may predict the severity of RV dilatation/dysfunction and the coexistence of LV involvement, which are among the most important determinants of the disease phenotypic appearance and clinical outcome.\(^8\)\(^9\)

**Electrocardiographic Repolarization Abnormalities**

Previous studies demonstrated that T waves in ACM typically become negative with the disease worsening and that negative T waves in the right precordial leads (V1-V3/V4) reflect the fibrofatty myocardial replacement, resulting in RV dilatation and dysfunction.\(^15\) Our study results confirm and extend these previous observations by demonstrating a relationship between the presence and extent of negative T waves and the degree of RV dilatation and dysfunction, as evaluated by cine CMR sequences.

In previous studies, the extent of negative T waves across a 12-lead ECG was shown to predict the amount of RV electroanatomic scar, as evaluated by the voltage mapping at the CARTO (Biosense Webster, Inc) system.\(^16\)\(^17\) However, the contributing role of LV scar involvement to left precordial TWI was not assessed because the map was limited to the RV. The present study allowed us to analyze the relationship between electrocardiographic repolarization abnormalities and the relative involvement of the RV, LV, or both, in terms of either end-diastolic volumes and ejection fraction by cine CMR sequences or LGE/myocardial scar by postcontrast sequences for tissue characterization. A major study result was that the extent of TWI toward the left precordial leads (V4-V6) and/or inferior leads was associated with a more severe RV dilatation or dysfunction, whereas it did not predict

**Table 3. Relationship Between Prolonged (>55 ms) TAD and CMR Parameters**

| Parameters               | TAD >55 ms | \( P \) Value |
|--------------------------|------------|---------------|
|                         | No (n=40)  | Yes (n=27)    |
| RV EDV, mL/m²            | 99 (92–115)| 116 (95–133)  | 0.014 |
| RV EF, %                 | 50 (44–59)| 41 (33–56)    | 0.053 |
| RV WMA, no. of segments  | 2 (1–3)    | 3 (1–4)       | 0.119 |
| RV LGE                   | 25 (63)    | 19 (70)       | 0.506 |
| LV EDV, mL/m²            | 82 (72–93)| 85 (72–100)   | 0.288 |
| LV EF, %                 | 59 (53–64)| 56 (49–61)    | 0.088 |
| LV LGE                   | 29 (73)    | 19 (70)       | 0.850 |

Values are expressed as number (percentage) of patients or median (25th–75th percentile). Patients with complete right bundle branch block were excluded.\(^3\) CMR indicates cardiac magnetic resonance; EDV, end-diastolic volume; EF, ejection fraction; LGE, late gadolinium enhancement; LV, left ventricular; RV, right ventricular; TAD, terminal activation duration; WMA, wall motion abnormality.
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According to International Task Force criteria, cutoff values were ejection fraction; %LatD, ratio of lateral displacement; LGE, late gadolinium enhancement; LV, left ventricular; RV, right ventricular; TWI, T-wave inversion; WMA, wall motion abnormality.

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Values are expressed as number (percentage) of patients or median (25th–75th percentile). α indicates cardiac apex angle; CMR, cardiac magnetic resonance; EDV, end-diastolic volume; EF, ejection fraction; %LatD, ratio of lateral displacement; LGE, late gadolinium enhancement; LV, left ventricular; RV, right ventricular; TWI, T-wave inversion; WMA, wall motion abnormality.

*According to International Task Force criteria, cutoff values were >100 mL/m² (men) and >90 mL/m² (women).

The involvement of the LV, either hemodynamic or structural, contrary to what one would expect.

The CE-CMR sequences also allowed us to find the plausible mechanism responsible for this finding. A previous electrocardiographic and vectorcardiographic correlation study reported that RV enlargement induces a backward displacement of the LV, with posterior dislocation of the LV electrical forces. Using the technique previously reported by Brosnan et al., we provided imaging evidence that the RV was displaced toward the axilla and a greater proportion of the ventricle was positioned under the electrocardiographic leads placed more laterally, as a consequence of its dilatation. Hence, the traditional LV leads (V4–V6) explored the electrical activity of the RV rather than that of the LV. This accounts for the strong association between the extent of TWI toward the lateral precordial leads and the severity of RV dilatation (and dysfunction) leading to leftward displacement.

The electrogenesis of T-wave inversion in ACM and its relation to the severity of RV disease remain to be elucidated. On the basis of our findings, one can speculate that replacement of RV myocardium by fibrofatty tissue induces a RV remodeling, with both hemodynamic alterations leading to ventricular dilatation and dysfunction and electrical changes of transmural and/or regional activation and repolarization times. These electrophysiological changes may become critical enough to cause reversal of repolarization gradients and lead to T-wave inversion.

**Table 4. Relationship Between Presence and Extent of Right Precordial TWI and CMR Parameters**

| Parameters                  | No TWI in Right Precordial Leads (n=18) | TWI in Right Precordial Leads Only (n=11) | TWI in Right Precordial-Left Precordial and/or Inferior Leads (n=50) | P Value |
|-----------------------------|----------------------------------------|------------------------------------------|-----------------------------------------------------------------|--------|
| RV EDV, mL/m²               | 92 (75–103)                            | 94 (90–115)                              | 117 (104–130)                                                   | <0.001 |
| RV EF, %                    | 57 (54–63)                             | 49 (38–60)                               | 41 (34–49)                                                      | <0.001 |
| RV WMA, no. of segments     | 1 (0–2)                                | 2 (1–3)                                  | 3 (2–4)                                                        | <0.001 |
| RV dilatation*              | 6 (33)                                 | 9 (82)                                   | 46 (92)                                                         | <0.001 |
| RV dysfunction (RV EF <45%) | 1 (6)                                  | 4 (36)                                   | 32 (64)                                                        | <0.001 |
| Displacement angle (α), °    | 40 (39–42)                             | 36 (32–39)                               | 52 (44–60)                                                     | 0.02   |
| Displacement ratio %LatD, %  | 60 (56–63)                             | 52 (41–53)                               | 97 (57–100)                                                    | <0.001 |
| LV EDV, mL/m²               | 92 (77–96)                             | 89 (75–100)                              | 82 (71–91)                                                     | 0.192  |
| LV EF, %                    | 57 (49–64)                             | 59 (52–63)                               | 58 (48–62)                                                     | 0.787  |
| LV dilatation (≥90 mL/m²)   | 11 (61)                                | 5 (46)                                   | 17 (34)                                                        | 0.131  |
| LV dysfunction (≤50%)        | 7 (39)                                 | 1 (9)                                    | 14 (28)                                                        | 0.221  |
| RV LGE                      | 9 (50)                                 | 5 (46)                                   | 41 (82)                                                        | 0.007  |
| LV LGE                      | 13 (72)                                | 7 (64)                                   | 37 (74)                                                        | 0.786  |

Values are expressed as number (percentage) of patients or median (25th–75th percentile). α indicates cardiac apex angle; CMR, cardiac magnetic resonance; EDV, end-diastolic volume; EF, ejection fraction; %LatD, ratio of lateral displacement; LGE, late gadolinium enhancement; LV, left ventricular; RV, right ventricular; TWI, T-wave inversion; WMA, wall motion abnormality.

**Table 5. Relationship Between Low Voltages of QRS (<0.5 mV) in Limb Leads and CMR Parameters**

| Parameters                  | Low QRS Voltages | P Value |
|-----------------------------|------------------|---------|
|                             | No (n=63)        | Yes (n=16) |       |
| RV EDV, mL/m²               | 109 (94–118)     | 105 (89–137) | 0.864 |
| RV EF, %                    | 49 (38–56)       | 42 (30–60)  | 0.320 |
| RV WMA, no. of segments     | 2 (1–4)          | 3 (2–4)    | 0.349 |
| RV LGE                      | 43 (68)          | 12 (75)    | 0.764 |
| LV EDV, mL/m²               | 85 (74–96)       | 86 (71–92) | 0.403 |
| LV EF, %                    | 59 (51–64)       | 53 (46–59) | 0.087 |
| LV LGE                      | 41 (65)          | 16 (100)   | 0.004 |
| LV LGE, no. of segments     | 3 (2–7)          | 6 (5–9)    | <0.001 |

Values are expressed as number (percentage) of patients or median (25th–75th percentile). CMR indicates cardiac magnetic resonance; EDV, end-diastolic volume; EF, ejection fraction; LGE, late gadolinium enhancement; LV, left ventricular; RV, right ventricular; WMA, wall motion abnormality.

Electrocardiographic Depolarization Abnormalities

The electrocardiographic abnormalities of ACM resulting from delayed RV activation/conduction include incomplete (rarely complete) right bundle branch block, prolongation of right precordial QRS duration, delayed S-wave upstroke with a terminal activation delay of >55 ms, and postexcitation ε waves (ie, small-amplitude potentials occurring at the end of the QRS complex/beginning of the ST segment). These electrocardiographic changes reflect areas of slow...
intraventricular conduction, in which the underlying substrate consists of regions of surviving myocardium interspersed with fatty and fibrous tissue. These can cause fragmentation of the electric activation of the ventricular myocardium and predispose to reentrant ventricular arrhythmias. Experimental studies have showed that susceptible desmosomal-gene carriers may harbor arrhythmogenic mechanisms at a molecular and cellular level because of a cross talk between altered desmosomes and both voltage-gated sodium-channel and gap junction proteins. Loss of expression of desmosomal proteins may induce electrical ventricular instability by a concomitant sodium channel dysfunction with current reduction, as a consequence of the interaction of these molecules at the intercalated discs. These findings have raised concerns that desmosomal-gene mutation carriers may experience life-threatening ventricular arrhythmias before fibrofatty myocardial replacement, structural/functional ventricular abnormalities develop, and sudden cardiac death may occur during the so-called concealed phase of the disease. However, clinical outcome studies did not support this hypothesis, showing that phenotypic expression is a prerequisite for malignant arrhythmic events and sudden cardiac death in ACM.

Among depolarization/activation abnormalities, in our study, abnormal prolongation of TAD (>55 ms) was the most important electrocardiographic predictor of RV dilatation and dysfunction. An abnormal TAD was observed in 40% of our study patients and was significantly associated with a worse RV disease, characterized by greater cavity dimension and more reduced systolic function. The TAD interval, which, according to the original definition, extends from the nadir of the S wave to the end of all depolarization deflections, is an expression of the global RV activation time, including other depolarization abnormalities, such as prolonged QRS duration, delayed and fractionated S-wave upstroke, and ε waves.

The ε waves are an uncommon finding in patients with advanced disease and are usually associated with the positivity of other diagnostic criteria. In the present study,

Figure 1. Electrocardiographic and contrast-enhanced cardiac magnetic resonance (CMR) findings of a representative case of right-dominant (classic) arrhythmogenic cardiomyopathy variant. A, Basal ECG showing T-wave inversion in right precordial leads (V1-V4). B, End-diastolic frame of cine CMR sequence in long-axis 4-chamber view showing a dilated right ventricle (end diastolic volume, 127 mL/m²) with a severely reduced ejection fraction (25%). The postcontrast orthogonal images in long-axis (C) and short-axis (D) views show late gadolinium enhancement as midwall stria in the midseptum (white arrow). In C, late gadolinium enhancement is also visible in the anterolateral, mid, and apical regions of the right ventricular wall, with segmental transmural involvement (white arrowheads) associated with regional dyskinesia (not shown).
postexcitation e waves were found in only 16% of patients, and any statistical association with structural or functional CE-CMR abnormalities was not demonstrable.

The finding that depolarization abnormalities were unrelated to LV involvement was expected, given that all these electrocardiographic depolarization parameters are confined
to right precordial leads and their alterations indicate a right intraventricular conduction defect.

**Low QRS Voltages in Limb Leads**

The electrocardiographic pattern of low QRS voltages in the limb leads with normal precordial amplitudes is the result of conditions that either alter voltage transmission or impair voltage generation from the myocardium to the skin electrodes, such as pericardial or large pleural effusion, infiltrative cardiomyopathy, and obesity. Several studies reported anecdotal evidence that low QRS voltages in limb leads may reflect the presence of a segmental LV scar/LGE, which can be evidenced by CE-CMR. Actually, CE-CMR has emerged as the imaging test that allows the “in vivo” detection and the characterization of LV involvement in ACM, in terms of dilatation and systolic dysfunction assessed by cine CMR sequences and/or fibrofatty scar detected by postcontrast CMR sequences. On the basis of the presence and extent of the coexistent LV involvement, the spectrum of ACM phenotypes has broadened to include “biventricular” and “left-dominant” variants, in addition to the classic “right-dominant” form. The LV lesions consist of fibrofatty myocardial replacement, which may affect the free wall, the interventricular septum, or both, either diffusely or regionally, with a predilection for the posterolateral regions. Like RV lesions, the wave front of fibrofatty myocardial replacement in the LV progresses from the epicardium to the endocardium, with scar tissue typically confined to subepicardial/midmural layers. The presence of LV involvement was originally considered a sign of advanced disease, being an end-stage manifestation and leading ultimately to biventricular pump failure. More recent genotype-phenotype correlation studies have shown early and greater LV involvement related to some genetic defects, such as desmoplakin-gene mutations.

The present study extended previous observations by showing that the electrocardiographic pattern of low QRS voltages may predict LV involvement in the context of ACM with a specificity of 100%, although its sensitivity did not...
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Conclusion

The study results indicate that electrocardiography has the potential to predict the ACM phenotype, with particular reference to the severity of RV disease and LV involvement, which are among the most important clinical determinants of the disease outcome. Specifically, the extent of TWI toward left precordial and inferior leads and prolonged TAD predicted higher RV dilation and lower RV ejection fraction, whereas low QRS voltages in limb leads predicted the presence of LV LGE/sca. These findings support the concept that routine electrocardiography in ACM provides both diagnostic and prognostic information. According to the 2010 IFT diagnostic criteria for ACM, the electrocardiographic pattern of TWI in V4, V5, or V6 is the only proposed electrocardiographic criterion for LV involvement. Our study results suggest considering the electrocardiographic pattern of low QRS voltages in limb leads, which was shown to be a more accurate predictor of LV involvement.

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