Right Ventricular Strain Predicts Structural Disease Progression in Patients With Arrhythmogenic Right Ventricular Cardiomyopathy

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BACKGROUND: Arrhythmogenic right ventricular cardiomyopathy (ARVC) is an inherited condition associated with ventricular arrhythmias and myocardial dysfunction; however, limited data exist on identifying patients at highest risk. The purpose of the study was to determine whether measures of right ventricular (RV) dysfunction on echocardiogram including RV strain were predictive of structural disease progression in ARVC.

METHODS AND RESULTS: A retrospective analysis of serial echocardiograms from 40 patients fulfilling 2010 task force criteria for ARVC was performed to assess structural progression defined by an increase in proximal RV outflow tract dimensions (parasternal short or long axis) or decrease in RV fractional area change. Echocardiograms were analyzed for RV free-wall peak longitudinal systolic strain using 2-dimensional speckle tracking. Risk of structural progression and 5-year change in RV outflow tract measurements were compared with baseline RV strain. Of the 40 ARVC patients, 61% had structural progression with an increase in the mean parasternal short-axis RV outflow tract dimension from 36.2 to 38.5 mm ($P=0.022$) and 68% by increase in parasternal long-axis RV outflow tract dimension from 36.1 to 39.2 mm ($P=0.001$). RV fractional area change remained stable over time. Baseline RV strain was significantly associated with the risk of structural progression and 5-year rate of change. Patients with an RV strain more positive than −20% had a higher risk (odds ratio: 18.4; 95% CI, 2.7–125.8; $P=0.003$) of structural progression.

CONCLUSIONS: RV free wall strain is associated with the rate of structural progression in patients with ARVC. It may be a useful marker in determining which patients require closer follow-up and treatment.

Key Words: arrhythmogenic right ventricular cardiomyopathy ■ echocardiography ■ strain imaging
can be fatal. Noninvasive measures determined by the 2010 task force criteria (TFC) include structural and functional parameters for diagnosis of ARVC by echocardiography including RV outflow tract (RVOT) dimension and RV fractional area change (RV-FAC).4 Changes to these imaging parameters have been used to identify and follow disease progression.11 Echocardiography has emerged as an ideal modality because it is noninvasive, widely available,12 and safe for use in patients defibrillators, as is common in ARVC.

Echocardiographic myocardial strain has emerged as a principal method for the quantification of subclinical left ventricular (LV) dysfunction in a variety of clinical settings,13 including chemotoxicity14 and ischemic heart disease.15 The role of echocardiographic strain in evaluating RV dysfunction in patients with ARVC is not fully elucidated, although several studies have indicated a possible role for this technique. Prior studies report reduced RV free wall strain and strain rate in patients with ARVC compared with measurements in healthy controls.16–20 Limited data have evaluated the utility of various echocardiographic parameters in ARVC for predicting disease progression over time. In one study, variability in RV and LV contraction duration was associated with greater arrhythmia risk.21 A recent study in first-degree relatives of patients with ARVC found abnormal RV deformation in the subtricuspid region to be associated with disease progression.22 However, no studies have systematically characterized the role of traditional and novel measures of RV dysfunction as they relate to ARVC disease progression. Moreover, further characterization of the relationship between RV strain and structural progression over time in ARVC would be beneficial for risk stratification.

The purpose of this study was to evaluate echocardiographic free-wall longitudinal strain of the RV as a predictor of progression of structural disease in ARVC using data from patients who underwent serial echocardiograms enrolled in the Johns Hopkins ARVC registry. We hypothesized that worse peak RV strain and strain rate would be significantly associated with progression of disease, as defined by increasing RVOT dimension and decreased RV-FAC.4

**METHODS**

**Study Population**

The study population consisted of 40 patients with ARVC from the Johns Hopkins arrhythmogenic right ventricular dysplasia/cardiomyopathy registry (http://ARVD.com). All patients were diagnosed with ARVC and fulfilled the 2010 task force criteria.4 To be included in this study, patients must have undergone at least 2 separate echocardiograms >6 months apart that were digitally available and of sufficient quality for analysis. The first and last available examinations were used for analysis, and the first exam was considered the baseline echocardiogram. The study was approved by the Johns Hopkins Medicine institutional nonstandard abbreviations and acronyms
review boards, and patients in the Johns Hopkins University arrhythmogenic right ventricular dysplasia/cardio-myopathy registry provided written informed consent. Participants did not receive compensation for this study. The data that support the findings of this study are available from the corresponding author on reasonable request.

**Clinical Characterization**

Fulfillment of TFC was assessed for each patient, as similarly performed by Mast et al. All patients had complete transthoracic echocardiograms including dedicated RV apical views. LV ejection fraction was calculated on 2-dimensional parasternal long-axis (PLAX) images using the modified Quinones formula. Cardiac magnetic resonance was performed per standard ARVC protocols described elsewhere and was used only to assess fulfillment of TFC structural criteria.

All participants had undergone 12-lead ECG recordings as part of assessment of TFC. ECG recordings were analyzed for fulfillment of TFC, including depolarization criteria (ε waves and terminal activation duration ≥55 ms) and repolarization criteria (precordial T-wave inversion, V1–V6). In addition, signal-averaged ECG was evaluated for the presence of late potentials. History of arrhythmias (nonsustained or sustained ventricular tachycardia of criteria-defined axis and morphology) was also noted. Holter monitors were evaluated for premature ventricular complex count and were considered abnormal if >500 was observed in 24 hours, per the TFC.

All index patients also underwent genetic testing by molecular genetic screening of 5 ARVC-associated desmosomal genes (PKP2 [plakophilin-2], DSG2 [desmoglein-2], DSC2 [desmocollin-2], DSP [desmoplakin], and JUP [plakoglobin]), as well as nondesmosomal genes (TMEM43 [transmembrane protein 43] and PLN [phospholamban]).

**Assessment of Structural Progression**

Echocardiograms performed at baseline and follow-up were exported from Synapse Cardiovascular (FUJIFILM Medical Systems) in DICOM (Digital Imaging and Communications in Medicine) format and analyzed using Image-Arena v4 (TomTec Imaging Systems), a vendor-neutral imaging platform.

Measurements were obtained for proximal RVOT dimensions in both PLAX and parasternal short-axis (PSAX) views during end-diastole, as shown in Figure 1. RV-FAC was assessed as described in the TFC as the percentage of RV area decrease between diastole and systole obtained on apical imaging. All echocardiograms were analyzed by a single operator (S.W.), to exclude interobserver variability, and measurements were performed blinded to clinical data.

**Assessment of Strain and Strain Rate**

RV free-wall peak longitudinal systolic strain and strain rate were measured on DICOM images of an RV-focused apical 4-chamber view by endocardial speckle-tracking using the 2D Cardiac Performance Analysis module of Image-Arena. All available cardiac cycles were examined, and the best-quality clip with optimal tracking was chosen. The peak values of free-wall strain and strain rate were defined as the deepest points of the respective curves (Figure 2). The exact time of peak may vary but is generally around end systole for strain and early systole for strain rate. Furthermore, the peaks may not occur at the same time for the free wall and septum; however, only free-wall values were used in our analysis, and septal strain values were excluded. LV endocardial peak longitudinal systolic strain was measured using the AutoStrain package on similarly selected cycles with optimal tracking, where the endocardium was automatically recognized and fine-tuned manually at end diastole (first frame after mitral valve closure) and end systole (first frame after aortic valve closure). A sample image of RV strain on a patient with ARVC is shown in Figure 2. A similar technique is described by Hamada-Harimura et al.

**Statistical Analysis**

Continuous data are presented as mean±SD or median with interquartile range. Categorical variables are presented as counts and percentages. A 2-sided P<0.05
was considered significant. Differences between baseline and follow-up echocardiographic measurements were compared using paired $t$ tests for continuous variables and $\chi^2$ tests for categorical variables. Multivariable logistic regression (for risk of progression) and linear regression (for quartile of progression rate) adjusting for age and sex were used to assess the relationship between baseline RV strain and strain rate and structural progression. This was done separately for RVOT PSAX, RVOT PLAX, and RV-FAC. Statistical analysis was performed using Stata v14.2 (StataCorp).

**RESULTS**

Clinical characteristics for the patient population are shown in Table 1. All 40 participants fulfilled the TFC for definite ARVC. There were 29 probands and 11 affected family members; 21 patients (53%) were women. Mean age was 35.2±12.6 years at baseline, and median time to follow-up echocardiogram was 3.6 years (interquartile range: 1.3–6.8 years). Minimum age at baseline was 19 years, and maximum age was 73 years. A pathogenic or likely pathogenic variant was identified in 65% of the study population. Variants in **PKP2** were most common ($n=21$, 53%). From echocardiography, 23 patients (58%) met major criteria at baseline, whereas 29 patients (73%) met major criteria at follow-up. LV ejection fraction was 58±7.3% at baseline and 56±10.9% at follow-up ($P=0.035$). LV global longitudinal peak systolic strain was relatively stable during follow-up ($−20.1\%$ to $−19.3\%$, $P=0.253$).

Between baseline and follow-up studies, mean RVOT PSAX increased from 36.2 to 38.5 mm ($P=0.022$) with structural progression (increase in size) in 23 patients (60%); mean RVOT PLAX increased from 36.1 to 39.2 mm ($P=0.001$) with progression in 25 patients (67%); and mean RV-FAC was relatively unchanged, decreasing from 31.6% to 30.6% ($P=0.331$) with 21 patients (52%) classified as progressed. Baseline age, sex, LV ejection fraction, RV-FAC, RV basal diameter, and fulfillment of major structural TFC by echocardiography were compared according to progression status, and results are summarized in Table 2. Patients who exhibited progression by RVOT PSAX overall had lower RV-FAC (28.5±9.6% versus 38.1±6.6%, $P=0.002$), larger RV basal diameters (4.6±0.88 versus 3.9±0.42 cm, $P=0.005$), and a greater percentage of patients meeting major structural criteria by echocardiography (74% versus 27%, $P=0.004$) at baseline compared with patients who did not progress by RVOT PSAX. In addition, progressors by RVOT PSAX tended to be men, compared with nonprogressors (61% versus 27%, $P=0.039$). Patients who exhibited progression (decrease) in RV-FAC had significantly greater RV-FAC measurements at baseline (35.2±9.7 versus 27.7±8.6, $P=0.014$) compared with patients who did not progress by RV-FAC.

Table 3 shows the association of baseline RV strain and strain rate with the risk of structural progression and across quartiles of 5-year rate of change for each of the 3 parameters (adjusted for age and sex). Overall, both RV strain and strain rate were significantly associated with the likelihood of future structural progression and the rate of change of PSAX RVOT and PLAX RVOT dimensions. Patients with more negative strain (eg, $−22.7\%$ for PSAX quartile 1) or strain rate were...
less likely to progress, whereas less negative numbers (eg, −14.5% for PSAX quartile 4) were associated with a faster rate of structural progression. No significant differences in strain or strain rate were seen between structural progressors and nonprogressors or by quartile of rate of change according to RV-FAC. A comparison of distribution of baseline RV strain across each quartile is shown in Figure 3. We further defined an abnormal RV peak longitudinal systolic strain as being more positive (greater) than −20%. There were 24 patients (60%) with abnormal RV strain at baseline. These patients were at substantially greater risk of structural progression than those with a normal baseline RV strain: 18 times higher risk of RVOT PSAX progression (odds ratio: 18.4; 95% CI, 2.7–125.8; P=0.003) adjusting for age and sex and 3.7 times higher risk of RVOT PLAX progression (odds ratio: 3.65; 95% CI, 0.60–22.4; P=0.161).

### DISCUSSION

In this study, we evaluated 2-dimensional echocardiographic RV strain as a predictor of structural disease progression in ARVC. This study has several notable findings. First, our results indicate that many patients exhibited progression in RVOT dimensions (60% for PSAX and 67% for PLAX) and less so for RV-FAC during the follow-up period. This finding is expected given that ARVC is a progressive disease and because similar findings have been seen in prior studies that have evaluated for progression on serial echocardiograms. Mast et alreported progression of all 3 measures (PSAX RVOT, PLAX RVOT, and RV-FAC) in two-thirds of their study population over a mean follow-up time of 6.4 years, and this result was reproduced in our study. Our results also indicate that worse RV free-wall peak longitudinal systolic strain and strain rate are associated with increases in RVOT size, as obtained from PSAX and PLAX views. Specifically, when baseline strain and strain rate were compared for patients who demonstrated structural progression and those who did not, worse strain and strain rate were observed among the progressors. Higher quartiles of 5-year RV structural progression (by both PSAX RVOT and PLAX RVOT) were associated with diminished RV free-wall strain and strain rate at baseline (Table 3). Furthermore, patients with more abnormal RV strain (greater than −20%) had an 18-fold risk (P=0.002) of progression in PSAX RVOT. Thus, RV free-wall strain and strain rate appear to be predictive of structural progression of disease in patients with ARVC, and patients with relatively normal baseline values progressed slowly, whereas those with abnormal values progressed more rapidly. These measures may have important clinical utility because structural changes and progression are
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Regarding measurement of RVOT, prior studies have shown that RVOT measures by echo are reproducible and correlate with cardiac magnetic resonance imaging measures. One study reported high reproducibility, with interclass correlation values >0.91 for inter- and intraobserver correlation of 2-dimensional RVOT measurements on echo. In addition, RVOT dimension on transthoracic echocardiogram exhibited good correlation with the same measure on cardiac magnetic resonance (r = 0.87; 95% CI, 0.75–0.93; P < 0.0001) and good absolute agreement with a mean difference of 0.2 mm between the 2 imaging modalities. This metric was used in our study and was utilized by the Heart Rhythm Society 2010 TFC in the diagnosis of ARVC. Another study reported a 2-mm increase in RVOT dimension over time in a population of ARVC patients and was found to be significant, similar to our current findings. Based on the reproducibility of RVOT measurements and findings in prior studies, changes in RVOT size in this study of 2.3 and 3.2 mm on PSAX and PLAX views, respectively, are appropriately significant, and they correspond to relative changes of ~6.3% and 8.6% from baseline to follow-up.

It should be noted that patients who progressed by RVOT PSAX had lower baseline RV FAC, larger

Table 2. Comparison of Demographics and LVEF by Progression Status

|                         | RVOT PSAX Progression | RVOT PLAX Progression | RV-FAC Progression |
|-------------------------|------------------------|------------------------|---------------------|
|                         | Yes (n=23)             | No (n=15)              | P Value             |
| Age, y, mean±SD         | 34.5±12.5              | 33.6±10.7              | 0.825               |
| Male (%)                | 14 (61)                | 4 (27)                 | 0.039               |
| LVEF, %, mean±SD        | 57.6±8.2               | 60.8±5.0               | 0.187               |
| Baseline RV-FAC, %, mean±SD | 28.5±9.6            | 38.1±6.6               | 0.002               |
| Baseline RV basal diameter, cm, mean±SD | 4.6±0.88          | 3.9±0.42               | 0.005               |
| Major structural TFC satisfied by echocardiography at baseline, n (%) | 17 (74)          | 4 (27)                 | 0.004               |

Baseline age, sex, LVEF, RV-FAC, RV basal diameter, and fulfillment of major structural TFC by echocardiography were compared across progression status. P values were calculated using t tests (age, LVEF, RV-FAC, RV basal diameter) and χ² analysis (sex, major structural TFC by echocardiography). LVEF indicates left ventricular ejection fraction; PLAX, parasternal long axis; PSAX, parasternal short axis; RV, right ventricular; RV-FAC, right ventricular fractional area change; RVOT, right ventricular outflow tract; and TFC, Task Force Criteria.

Table 3. Association of Baseline RV Strain and Strain Rate With Risk and Rate of Structural Progression by PSAX-RVOT, PLAX-RVOT, and RV-FAC

|                         | Structural Progression | Quartile of 5-y Rate of Change (1=Best, 4=Worst) |
|-------------------------|------------------------|-------------------------------------------------|
|                         | Yes | No | P Value* | 1 | 2 | 3 | 4 | P Value† |
| PSAX RVOT               |     |    |         |    |   |   |    |         |
| RV strain, %, mean±SD   | −15.4±4.5 | −22.7±3.9 | 0.007       | −22.7±3.6 | −20.4±5.4 | −15.5±4.5 | −14.5±4.8 | 0.001     |
| RV strain rate, 1/s, mean±SD | −0.64±0.21 | −0.97±0.23 | 0.003       | −0.91±0.18 | −0.90±0.31 | −0.68±0.28 | −0.59±0.17 | 0.011     |
| PLAX RVOT               |     |    |         |    |   |   |    |         |
| RV strain, %, mean±SD   | −17.3±4.9 | −20.8±6.6 | 0.160       | −20.9±7.0 | −19.5±4.3 | −18.3±3.7 | −14.6±5.5 | 0.011     |
| RV strain rate, 1/s, mean±SD | −0.71±0.25 | −0.91±0.27 | 0.043       | −0.87±0.16 | −0.90±0.34 | −0.76±0.26 | −0.57±0.22 | 0.008     |
| RV-FAC                  |     |    |         |    |   |   |    |         |
| RV strain, %, mean±SD   | −18.6±5.7 | −16.8±6.4 | 0.505       | −15.9±7.0 | −17.0±6.1 | −18.9±6.7 | −19.2±4.2 | 0.219     |
| RV strain rate, 1/s, mean±SD | −0.79±0.32 | −0.71±0.24 | 0.497       | −0.83±0.28 | −0.74±0.22 | −0.84±0.40 | −0.78±0.20 | 0.254     |

RV strain and strain rate are compared between progressors and nonprogressors and across quartiles of 5-y rate of change. PLAX indicates parasternal long axis; PSAX, parasternal short axis; RV, right ventricular; RV-FAC, right ventricular fractional area change; and RVOT, right ventricular outflow tract.

*P values from multivariable logistic regression (adjusting for age and sex).
†P values from ordinal logistic regression (adjusting for age and sex).
RV basal diameters, and greater fulfillment of major structural criteria by echocardiography at baseline compared with patients who did not progress by RVOT PSAX (Table 2). These findings are consistent with the notion that ARVC is a progressive disease; patients who exhibited progression tended to have greater evidence of disease at baseline. In addition, these findings were not observed among progressors by RVOT PLAX. It is possible that variability in imaging technique is responsible for this discrepancy. However, it also suggests that RV strain and strain rate, in addition to other structural markers of disease, can be used to predict structural progression. Patients who progressed by RVOT PSAX tended to be male; however, the implications of this finding are unclear and perhaps incidental, as this association was not seen for RVOT PLAX and RV-FAC. Patients who progressed by RV-FAC tended to have larger RV-FAC at baseline, perhaps reflecting a larger “window” for decrease in RV function with higher RV-FAC at baseline.

In a prior study by Mast et al., subtricuspid RV deformation was evaluated in first-degree relatives of patients with ARVC, using criteria that incorporated strain. The authors concluded that abnormal RV deformation seemed to precede established signs of ARVC. However, the study performed by Mast et al. utilized prespecified deformation patterns to predict disease progression, whereas our study evaluated RV free-wall peak longitudinal systolic strain and strain rate. Our structural predictors for disease progression are related to the described deformation patterns but are arguably simpler to implement. Furthermore, the patient population in the study by Mast et al. consisted of first-degree relatives of patients with ARVC and excluded patients who fulfilled TFC for structural abnormalities. In contrast, our study evaluated patients who carried diagnoses of definite ARVC regardless of criteria fulfilled. Thus, our study expands the patient population in which strain imaging may be useful.

Speckle-tracking strain can also assess ventricular contraction homogeneity using mechanical dispersion, which is defined as the standard deviation of the time from the Q/R on ECG to peak negative longitudinal strain for all RV or LV segments. Prior studies have evaluated RV and LV mechanical dispersion, in addition to RV strain, to predict arrhythmias in ARVC and differentiate early ARVC from RVOT–ventricular tachycardia patients. Leren et al. found RV mechanical dispersion to be a marker of prior arrhythmic events in patients with early ARVC, but the authors also identified an association between diminished RV global longitudinal strain and arrhythmia events in a broader population that included patients with definite ARVC. Lie et al. also found an association between RV longitudinal strain and life-threatening ventricular arrhythmias but identified a stronger
relationship using LV mechanical dispersion. In contrast to these studies, our study validates the utility of strain techniques in the definite ARVC population and demonstrates the ability of strain to predict other markers of ARVC severity, namely, structural progression. In addition, compared with mechanical dispersion, RV free-wall strain is more widely available and less technically demanding to assess, making it more suitable for serial measurements over time to assess structural progression.

Results for RV-FAC disagree with our hypothesis; however, in our study population, the RV-FAC remained relatively stable during the follow-up period, with very few patients progressing. Saguner et al.\(^{26}\) found that reduced FAC was highly predictive of major adverse cardiovascular events in patients with ARVC. However, they also reported no association between an impaired tricuspid annular plane systolic excursion and subtricuspid regional wall motion abnormalities. Consequently, it is possible that strain and strain rate are more predictive of changes in the RVOT dimensions for patients with ARVC than for overall RV systolic function. Findings for RV-FAC may also have been limited by a relatively small patient population. In addition, the technical challenges of accurately determining RV-FAC on 2-dimensional echocardiographic imaging may have led to variability in assessment of RV function.

Limitations

This study has several limitations. First, it was a retrospective study from a single referral center with variable follow-up periods and relatively small sample size; however, because of the rare nature of ARVC, the study is still one of the largest of its kind. The measurement of strain and strain rate can be inconsistent depending on the user, image quality, software, or vendor of the echocardiography system. This is particularly true for the right ventricle, which remains challenging to assess quantitatively by echocardiography. However, we mitigated this by having only a single operator evaluate RV strain under the same software/hardware conditions. Furthermore, the operator was blinded to clinical data during analysis. It should also be noted that strain and wall dimension are related. However, apart from comparing against changes in RV-FAC, RV free-wall strain was used to identify changes in the RVOT, an anatomically separated location from where strain was measured. Last, echocardiography may be less sensitive than magnetic resonance imaging for determining structural progression of disease, given the ability of magnetic resonance imaging to identify segmental RV dilatation or hypokinesis with higher resolution.\(^{34}\) Nevertheless, echocardiography is more widely available clinically, particularly for ARVC patients, the majority of whom have defibrillators requiring special magnetic resonance imaging protocols.

CONCLUSIONS

Our study recognizes echocardiographic strain as a useful technique to predict structural progression of disease in ARVC. These findings suggest that echocardiographic strain and strain rate could be useful in identification of patients at risk of disease progression who may require closer follow-up and treatment.

ARTICLE INFORMATION

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