A comparison of cardiac magnetic resonance imaging peri-infarct border zone quantification strategies for the prediction of ventricular tachyarrhythmia inducibility

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Abstract

Background: Peri-infarct border zone (BZ) as quantified by late gadolinium enhancement (LGE) on cardiac magnetic resonance imaging (MRI) has been proposed as a risk stratification tool, and is associated with increased mortality. BZ has been measured by various methods in the literature. We assessed which BZ analysis best predicts inducible arrhythmia during electrophysiological study (EPS).

Methods: LGE was performed in 47 patients with coronary artery disease referred for EPS to assess for ventricular tachycardia (VT). LGE data was analyzed for BZ quantification by 3 previously published methods. Method I (BZ-I) used pixels 2–3 standard deviations over the mean of normal tissue, expressed as % of left ventricular mass, Method II (BZ-II, as described by Yan) and Method III (BZ-III, as described by Schmidt). EPS results were classified as negative (non-inducible) or positive (monomorphic VT — MVT).

Results: There were 47 subjects-age 61.7 years, 72% male. During EPS, 20 patients were non-inducible and 18 had induced MVT. Ejection fraction was not significantly different between non-inducible patients and those with MVT (34.1% vs. 28.5%, p = 0.13). BZ-I was significantly different (1.4% vs. 2.6%, p = 0.001), but not BZ-II (7.9% vs. 6.9%, p = 0.68) or BZ-III (2.7 g vs. 2.1 g, p = 0.88). Multivariate analysis demonstrated that only BZ-I was an independent predictor of EPS outcome after controlling for infarct size (OR 1.97 per % change, 95% CI 1.04–3.73, p = 0.04).

Conclusions: This study demonstrates significant variability between the published methods for measuring BZ. Also, BZ-I is a stronger predictor of inducible MVT during EPS than ejection fraction and infarct size. BZ may be another LGE marker of elevated risk of arrhythmia. (Cardiol J 2013; 20, 1: 68–77)

Key words: cardiac magnetic resonance imaging, ventricular tachycardia, sudden death

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Introduction

Sudden cardiac death (SCD) remains a pervasive modern health concern, with over 300,000 deaths annually in the United States alone [1]. Implantable defibrillators have proven to be an effective treatment modality in selected patients with coronary artery disease (CAD) [2–5]. The most widely used risk identifier for determining patients at risk for SCD is left ventricular ejection fraction (LVEF) [6]. While an effective parameter, LVEF has limited sensitivity and specificity; therefore identifying individuals at risk of SCD after myocardial infarction (MI) continues to be a challenging problem [7, 8]. SCD in patients with CAD is predominantly caused by ventricular tachycardia (VT) or ventricular fibrillation (VF) [4, 9]. The anatomic substrate defined by the infarction is a main component in the pathogenesis of these arrhythmias. Given cardiac magnetic resonance imaging’s (MRI) ability to delineate the infarction [10], infarct characterization by cardiac MRI (CMR) has become an evolving novel method for risk stratification [11–14].

Infarct size is a known determinant of the risk for occurrence of VT [15–17]. Recent human studies have demonstrated that infarct characterization by late gadolinium enhancement (LGE) on CMR is predictive of overall mortality [12] as well as inducibility of ventricular arrhythmia during electrophysiological study (EPS) [11, 18]. The peri-infarct border zone (BZ) has been defined histologically as the area of viable myocardium immediately adjacent to infarcted myocardium. This region has previously been shown in animal models to identify areas of abnormal enzyme activity, as well as altered electrical activation properties [19, 20]. Recently, “grey zone” imaging, has been introduced as a method for identifying the BZ by LGE and has been associated with increased mortality in a human population [14]. In delayed-enhanced CMR images, normal myocardium appears black and infarct white. The authors postulated that pixels with intermediate (grey) intensity identify a potentially arrhythmic heterogeneous zone of viable and nonviable peri-infarct myocardium, but this has not been confirmed pathologically. If this indeed represents the substrate for reentrant ventricular arrhythmias, there should be increased BZ among patients with inducible monomorphic VT (MVT) at EPS, a well accepted modality to demonstrate this substrate [21, 22]. Schmidt et al. [18] did identify BZ (in grams) to be a predictor of inducible VT, but infarct size was not. Interestingly, the techniques used for BZ quantitation in these latter 2 studies differed [14, 18]. Based on these recent data, we evaluated the predictive value of BZ relative to infarct size in our previously reported study of CMR determined infarct size and its relationship to inducible VT [11]. We hypothesized that BZ quantification by LGE would be more predictive of inducible arrhythmia during EPS than infarct size and ejection fraction.

Methods

This study was approved by the Northwestern University Institutional Review Board. This was a retrospective reanalysis of data from a prior study [11], in which 48 patients, all with known history of either chronic CAD or distant MI, were referred for EPS to assess for inducibility of VT for primary prevention risk stratification of sudden death. No patients had prior history of sustained ventricular arrhythmias.

In this study, these LGE data were re-analyzed for BZ quantification. One patient’s CMR could not be recovered and was excluded from this analysis. Patients underwent CMR within 32 ± 6 days of EPS, per the protocol of the prior study [11]. Patients were placed supine in a 1.5-T Magnetom Sonata scanner (Siemens, Medical Solutions, Malvern, Pennsylvania); fiberoptic electrocardiographic (ECG) leads were placed for scanner gating and a phased-array receiver coil was placed on the chest for imaging. All images were acquired using 10- to 15-s breath-holds. Short-axis cines were acquired from the base to apex, making sure to include the entire left ventricle (LV) using methods previously described [23, 24]. Typical CMR parameters were: matrix resolution 256, field-of-view 340–400 mm × 225–380 mm, slice thickness 6 mm, voxel size 1.33–1.56 mm × 1.56–2.43 mm × 6 mm. No parallel imaging was used. Repetition time (typically 59–60 ms) was selected to be 100 ms less than the R-R interval, to place the acquisition window within end-diastole, with 23–25 segments per cardiac cycle. Delayed-enhanced images were obtained more than 10 min following intravenous administration of a gadolinium-based contrast agent (0.2 mmol/kg, Magnevist, Berlex Pharmaceuticals, Wayne, New Jersey) using a T1-weighted, inversion-recovery, segmented fast gradient-echo pulse sequence [25]. The inversion time was adjusted throughout the scan to null normal myocardium. The methods used for the quantification of LVEF, infarct size, and infarct surface area have all been previously described [11]. Briefly, endocardial and epicardial borders of the myocardium were manually planimetered on the short-axis cine images for
each patient. Volumes were derived by summation of the pixel areas, followed by multiplication of in-plane resolution and the effective slice thickness. The LVEF was computed as (end-diastolic volume – end-systolic volume)/end-diastolic volume. LV mass was determined by subtracting endocardial volume from epicardial volume at end-diastole and multiplying by a density of 1.05 g/mL [18]. Infarct volume from epicardial volume at end-diastole and – end-systolic volume)/end-diastolic volume. LV mass was determined by subtracting endocardial mass from epicardial mass at end-diastole and multiplying by a density of 1.05 g/mL, the pixel values were converted to actual cardiac masses and surface areas.

Electrophysiologic study was performed using standard techniques. Programmed ventricular stimulation was performed using up to three extrastimuli at two right ventricular sites during 2 drive-cycle lengths. Study end points were either induction of sustained VT or completion of the study protocol. EPS results had been previously classified as negative (no inducible ventricular tachyarrhythmia) (n = 20), inducible MVT (n = 18), or inducible VF or polymorphic VT (PVT) (n = 9). Because inducible PVT/VF is considered a non-specific result during EPS, the primary analysis comparing LGE results, as well as comparing the different BZ calculation methods, evaluates the non-inducible group vs. those with inducible MVT, as in the original report. However, baseline characteristics and long-term outcomes are reported for all subjects.

Border zone has been previously quantified using two statistical techniques. Yan et al. [14] defined BZ by regions of signal intensity between 2 and 3 SD above normal myocardium. They then indexed this measure to the infarct size. Of note, using this index, there was an inverse relationship between infarct size and BZ (i.e. the larger the infarct, the larger the BZ).

All image analysis was performed by a single investigator, blinded to the results of prior analysis, clinical outcomes and to EPS results. The methods for image analysis have been described in detail [26]. Briefly, all short-axis delayed-enhancement images from apex to base were reviewed offline using ImageJ (National Institute of Mental Health). A custom macro for ImageJ was written for the semi-automated quantification of LGE BZ. The macro analyzes a single, post-contrast short axis image. The user first defines a region of LV remote from the infarct that appears normal. Next, epicardial and endocardial surfaces are defined. The macro generates a summation of all LV pixels that meet criteria for normal, BZ, or core infarct.

Due to significant variations in the published methods by which to calculate BZ, we next performed the BZ analysis by 3 different methods, which will be referred to as BZ-I, BZ-II, and BZ-III. Border zone-I was defined as all pixels in the endocardium that were between 2 and 3 SD above the mean of the normal region, and core infarct those above 3. Border zone and core infarct were reported as a percentage of the total LV mass. Border zone-II was defined as in method I, but the values of BZ were reported as a percent of infarct size as described by Yan et al. [14] In BZ-III, BZ and core infarct mass were calculated using the full-width half-maximum method as described by Schmidt et al. [18]. For all methods, pixels between 2 and 3 SD above the mean of the remote region that were not adjacent to areas of core infarction (≥ 3 SD above remote mean) were not included in the BZ calculation. Two example studies are shown in Figure 1, comparing all 3 methods of BZ calculation.

Follow-up data on implantable cardioverter-defibrillator (ICD) shocks and mortality were assessed for all 47 patients by referencing the social security death index, as well as hospital and ICD chart review. ICD programming was done at the discretion of the treating physician; the average VT or VF zone cut-off was 167 bpm, range 95–188. Appropriate therapy was considered any high-energy shock or anti-tachycardia pacing for sustained ventricular arrhythmia.

Statistical analysis

Results are expressed as mean ± SD, unless otherwise noted. Discrete variables were compared across patient groups using the χ², Fisher, or Spearman tests where appropriate. Continuous variables...
were compared with Student’s t-test, Kruskal-Wallis or Mann-Whitney tests, where appropriate. Analysis of non-parametric parameters (EF, infarct size, infarct surface area, LV mass, and BZ) utilize the Kruskal-Wallis test, and are reported as median value (1st quartile, 3rd quartile). Analysis of variance was performed to compare characteristics among all 3 EPS groups. Two separate multivariate logistical regressions were performed to identify independent determinants of EPS outcome for each method; one using BZ and infarct mass percentage alone, and one using BZ, infarct mass percentage and LVEF. A 2-tailed p-value < 0.05 was considered statistically significant. All analysis was performed using SPSS 16.0 (SPSS Inc, Chicago, IL). The authors had full access to the data and take responsibility for its integrity. All authors have read and agree to the manuscript as written.

Results

The study group consisted of 47 subjects with an average age of 61.7 ± 11.4 years and 72% were male. During EPS, 20 patients had no inducible ventricular arrhythmias, 9 had PVT or VF, and 18 had induced MVT. Baseline characteristics are shown in Table 1. There were no significant differences among the three groups in terms of history of MI, congestive heart failure, diabetes mellitus, or beta-blocker and ACE inhibitor use. Only age, gender, and a reported history of hypertension were significantly different between the 3 EPS outcome groups. QRS duration on baseline 12-lead ECG was 110 ± 6 ms in the non-inducible group, 121 ± 5 ms in the MVT group, and 118 ± ± 11 ms in the VF group, which was a non-significant difference.

CMR results

The differences in LGE results between the non-inducible and inducible EPS groups are shown in Table 2. As previously reported, LVEF was not significantly different between non-inducible patients and those with MVT (34.1% vs. 28.5%, p = 0.13), while infarct size (15.8% vs. 23.1%, p = 0.03) and surface area (104.4 cm² vs. 169.8 cm², p = 0.002) were. No areas of microvascular obstruction were noted. All patients except 3 had some evidence of LGE (2 in non-inducible group, 1 in VF group). Analysis was repeated excluding these 3 patients, but no results were significantly altered. All results shown include these 3 patients.
All 47 patients had BZ analysis performed by all 3 methods. Table 2 shows a comparison of BZ quantification as described in this study (BZ-I), in Yan et al. [14] (BZ-II), and in Schmidt et al. [18] (BZ-III). Border zone was significantly different between the non-inducible and MVT EPS groups using BZ-I (1.4% vs. 2.6%, p = 0.001), but not using BZ-II (7.9% vs. 6.9%, p = 0.68) or BZ-III (2.7 g vs. 2.1 g, p = 0.88). The relationship of BZ by each method to infarct size is demonstrated in Figure 3.

Several multivariate logistical regression models were evaluated, as shown in Table 3. The first model included both infarct size and BZ-I (model 1), which demonstrated that BZ-I was the only significant predictor (OR 1.84 per % change, 95% CI 1.03–3.28, p = 0.04). When including LVEF, infarct size and BZ-I (model 2), BZ-I was again the only independent predictor of EPS outcome (OR 2.07 per % change, 95% CI 1.08–3.98, p = 0.03). In a separate model that included BZ-I and infarct surface area (model 3), the confidence intervals for the OR for BZ-I (OR 1.71, 95% CI 0.98–3.01, p = 0.06) included 1, but is still consistent with the value of this parameter as a predictor of EPS outcome.

Another multivariate logistical regression model that included infarct size and BZ-II demonstrated that BZ-II was not a predictor of EPS outcome (OR 0.95 per % change, 95% CI 0.85–1.07, p = 0.395). When including both infarct size and LVEF in the model, BZ-II was still not a predictor (OR 0.95 per % change, 95% CI 0.85–1.06, p = 0.378). In model 4, which included BZ-I, BZ-II and BZ-III, only BZ-I was a significant, independent predictor of EPS outcome (OR 2.74 per % change, 95% CI 1.34–5.58, p = 0.006).

Another logistical regression model was performed that included LVEF, infarct size and BZ-III. This analysis demonstrated that only infarct size was an independent predictors of EPS outcome, and BZ-III was not (OR 0.93 per % change, 95% CI 0.78–1.11, p = 0.41). In a model that included only BZ-III and infarct size, BZ-III again was not a significant predictor (OR 0.92 per % change, 95% CI 0.77–1.10, p = 0.35).
In order to examine if BZ had a simply largely geometric relation to infarct size and surface area or was a unique measurement, its correlation with other infarct characteristics was evaluated. As shown in Figure 2, the original measurements of

**Figure 2.** Infarct size (% left ventricular mass) vs. infarct surface area [cm²].

**Table 3.** Univariate and selected multivariate analyses.

<table>
<thead>
<tr>
<th>LGE variable</th>
<th>OR (95% CI)</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Univariate</td>
<td></td>
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<tr>
<td>Age</td>
<td>1.04 (0.98–1.12)</td>
<td>0.216</td>
</tr>
<tr>
<td>History of CHF</td>
<td>0.43 (0.09–1.95)</td>
<td>0.274</td>
</tr>
<tr>
<td>Gender</td>
<td>0.11 (0.02–0.65)</td>
<td>0.015</td>
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<tr>
<td>Infarct size</td>
<td>1.07 (1.01–1.14)</td>
<td>0.026</td>
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<tr>
<td>Border zone-I</td>
<td>2.10 (1.21–3.65)</td>
<td>0.008</td>
</tr>
<tr>
<td>Border zone-II</td>
<td>0.95 (0.85–1.05)</td>
<td>0.310</td>
</tr>
<tr>
<td>Border zone-III</td>
<td>0.97 (0.83–1.14)</td>
<td>0.748</td>
</tr>
<tr>
<td>LVEF</td>
<td>0.95 (0.89–1.01)</td>
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<tr>
<td>Multivariate</td>
<td></td>
<td></td>
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<tr>
<td>Model 1</td>
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<td>1.84 (1.03–3.28)</td>
<td>0.040</td>
</tr>
<tr>
<td>Infarct size</td>
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<tr>
<td>Model 2</td>
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<td>0.029</td>
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<tr>
<td>Infarct size</td>
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<tr>
<td>LVEF</td>
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<tr>
<td>Model 3</td>
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<td></td>
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<tr>
<td>Border zone-I</td>
<td>1.71 (0.98–3.01)</td>
<td>0.061</td>
</tr>
<tr>
<td>Surface area</td>
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<tr>
<td>Model 4</td>
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<tr>
<td>Border zone-I</td>
<td>2.74 (1.34–5.58)</td>
<td>0.006</td>
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<td>Border zone-II</td>
<td>0.82 (0.65–1.04)</td>
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<tr>
<td>Border zone-III</td>
<td>0.88 (0.72–1.08)</td>
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<td>Model 5</td>
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<td>Gender</td>
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<tr>
<td>Border zone-I</td>
<td>1.63 (0.93–2.84)</td>
<td>0.087</td>
</tr>
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</table>

OR — odds ratio; CI — confidence interval; CHF — congestive heart failure; LVEF — left ventricular ejection fraction

**Figure 3.** Comparison of infarct size vs. the three different methods of border zone quantification. From the top to the bottom: Border zone as a percent of left ventricular mass (border zone-I), Border zone as a percent of infarct size (border zone-II), and border zone mass by full-width half-max method (border zone-III).

In order to examine if BZ had a simply largely geometric relation to infarct size and surface area or was a unique measurement, its correlation with other infarct characteristics was evaluated. As shown in Figure 2, the original measurements of
Long-term outcomes

While not powered for outcome data, long-term data were available for the original study participants. Average follow-up was 1383.8 ± 617.0 days. There were 14 deaths, 6 in the non-inducible group (30.0%), 2 in the PVT and VF group (22.2%), and 6 in the MVT group (33.3%). There was no significant difference in mortality among the groups. Cause of death was not known for all patients. There were 26 ICD implants (4 in the non-inducible group, 5 in the PVT/VF group and 17 in the MVT group). Five patients received appropriate ICD therapy for spontaneous ventricular arrhythmias (1 in the non-inducible group, 4 in the MVT group).

Border zone was not significantly different between patients alive vs. dead by any method; BZ-I (1.9% vs. 2.4, p = 0.6), BZ-II (6.9% vs. 9.4%, p = 0.3) or BZ-III (2.3 g vs. 1.4 g, p = 0.6). When age was controlled for, neither EPS outcome nor BZ was predictive of death or a composite endpoint of death or appropriate ICD therapy.

Discussion

The principle finding of this study is that BZ quantified by three different techniques on the same set of CMRs provides dramatically different results. The measurements of BZ-I, BZ-II, and BZ-III are not highly correlated with each other and the relationship with total infarct mass markedly differs so that BZ-II is inversely related to infarct mass while BZ-I and BZ-III are directly related. Furthermore, while BZ-I is an independent predictor of induction of MVT during EPS in this patient cohort, neither BZ-II nor BZ-III were. Further studies will be required to determine which of these measurements provides the most realistic physiologic measurement of the BZ, and which provides optimal prognostic information.

Ventricular tachycardia commonly arises from reentry around a fixed lesion, such as an infarct scar [21, 27]. The critical region for these VT has been shown histologically to be bundles of surviving myocytes in the immediate peri-infarct area — the border zone [21]. Intracardiac electrical mapping studies have confirmed that the peri-infarct area is typically involved in VT [28–31], and catheter-based mapping and ablation of these areas has been shown to be an effective therapy for VT [32]. Slow conduction, a required component of reentrant arrhythmias, is found in the border of healing infarcts [33, 34]. In addition, areas of patchy infarct that may provide the geometric substrate for reentry [15], as well as areas of mixed dead and viable myocardium [21] have all been identified in the peri-infarct zone.

There are various clinical tests to attempt to identify patients with this substrate for ventricular arrhythmias. Some tests identify anatomic features associated with the peri-infarct border zone — for example, infarct size might be expected to be related to the mass of BZ present. As infarct size, until recently, could not be accurately measured, LVEF has been used as a surrogate risk marker. Although many factors influence LVEF [35, 36], in general, larger infarcts are associated with lower LVEF. As this has been a readily clinically available tool, it has been extensively studied and shown to be a useful predictor of SCD [37, 38]. LGE measurement of infarct scar by delayed-enhancement with gadolinium is a more specific measure of infarct size, allowing for direct quantification of the infarct morphology, which may be linked to arrhythmogenesis [11]. EPS is a non-anatomic risk stratification tool that examines the electrical properties of the ventricle and identifies the presence of substrate for VT. BZ measurement, however, has the potential to be a direct, non-invasive measure of the anatomic substrate which has previously been only histologically defined.

While the data for the use of BZ as measured by LGE to predict cardiovascular risk appear promising, there are significant limitations in our understanding of this imaging finding. While histologic data have confirmed that the peri-infarct zone is an area of abnormal electrophysiologic properties [33], it is not proven that this is what is being represented by the intermediate-intensity pixels near or within a LGE identified infarct. Ideally, all of these pixels would be due to areas that are an admixture of viable and non-viable myocardium, which may be a substrate for arrhythmias. However, these pixels may also occur due to other imaging conditions and artifacts. Partial volume effects, where normal myocardium, blood pool, or epicardial fat may exist within the same voxel volume along with the myo-
cardiac infarct, can produce intermediate-intensity values. This may produce the appearance of BZ on LGE imaging, even if the actual tissue is not an admixture of viable and non-viable myocardium, and presumably not arrhythmogenic [39]. Partial volume can occur in the short axis plane along the border of the infarct, as well as in the Z direction from above or below the infarct. In addition, poor T1 nulling of normal myocardium, as well as low signal-to-noise ratios in general, can increase the amount of intermediate-intensity pixels, and thereby falsely elevate the BZ levels [40]. However, despite all the potential confounders in the measurement of BZ by LGE, these data as well as prior studies [14], suggest it has clear biologic implications. In our data, BZ had a positive, but poor correlation with both infarct size as well as infarct surface area, suggesting that there is a minimal amount of artifact from peri-infarct volume averaging. Techniques that may minimize these other causes of intermediate-intensity pixels should help further improve the specificity of BZ to identify pro-arrhythmic substrate.

Contemporary literature on the use of LGE imaging for the quantification of infarct BZ reveals a number of statistical techniques for the measurement and reporting of BZ burden. We analyzed our images using these different techniques to look for similarities and differences, as shown in Table 2. One technique was reported in a recent study by Yan et al. [14]; a cohort of 144 patients with CAD had CMR images taken and were followed for an average of 2.4 years. Yan et al. [14] defined BZ (method II) as regions of signal intensity between 2 and 3 SD above normal myocardium, indexed to infarct size (pixels with values 3 SD above). In this cohort, BZ-II was a univariate predictor of total mortality (p = 0.01) and cardiovascular mortality (p = 0.009), as was LVEF (p = 0.01) and LV end-systolic volume index (p = 0.002). A multivariable model showed that BZ-II and LV end-systolic volume index were the 2 strongest, independent predictors of all-cause and cardiovascular mortality. While these data demonstrate a link between BZ and mortality, little is known about the selection or actual cause of death for these patients, who exhibited a higher than expected mortality rate of 20% over 2.4 years. The mechanism for death was conjectured to be reentrant ventricular arrhythmias.

In the present study, we quantified BZ as reported by Yan et al. [14], but also as a percentage of the total LV mass, in a similar fashion as infarct size has been reported in the past. Reporting BZ as a percentage of the infarct size [14] may discard important information about the gross amount of arrhythmic BZ tissue. In both Yan's report and in this study, when BZ is indexed to infarct size (method II) as opposed to LV mass (method I), there is an inverse relationship between BZ % and infarct size, as patients with smaller infarcts have larger BZ to infarct size ratios. A patient with a relatively small amount of arrhythmic BZ tissue, with a very small infarct, would have a large reported BZ ratio by method II. This inverse relationship of BZ and infarct size in method II does not correlate to the observation that larger infarcts are more arrhythmogenic [11]. We felt that a more useful metric for BZ would take into account the total amount of arrhythmic substrate present, and should have a positive correlation with total infarct %. This lead to method I, reporting BZ as indexed to total LV mass.

Another technique for BZ calculation has been reported by Schmidt et al. [18]. This study examined a cohort of 47 patients with CAD prior to ICD implant for primary prevention of sudden death. These patients underwent CMR, as well as either EPS or non-invasive programmed stimulation to look for ventricular inducibility. They defined BZ (method III) as pixels greater than the peak value in the normal myocardium but less than 50% of the highest signal in scar tissue (visually defined). This was reported unadjusted, in grams of myocardium. In this cohort, BZ-III was a significant predictor of inducibility of MVT (p = 0.015), but not total infarct size.

The comparison of techniques in this study does not establish the superiority of one method over another; rather, it highlights the variety of reporting methods currently being employed for BZ quantification, and that they differ significantly. All 3 methods described are similar in that they are predictive of either mortality or EPS outcomes. This is likely due to a common shared underlying physiology that they indirectly or directly measure — be it actual histologic BZ, or some relation to infarct mass or surface area. Until histologic confirmation of BZ is available and is related to a particular CMR technique, this issue will likely remain ambiguous.

**Limitations of the study**

A significant limitation of this study is that it is non-randomized and utilizes a small sample size. With only 18 patients in the MVT group, the multivariate analysis can be of limited statistical accuracy, and should be considered only as thought provoking and requiring additional study. However, the univariate results should still be valid despite the small sample and of significant importance. In addition, EPS inducibility identifies the presence of
substrate for VT, but may not be an adequate surrogate for arrhythmic sudden death. Finally, this CMR dataset was acquired from 1998–2002, and the image quality is somewhat less than with modern techniques. However, this only underscores how the different BZ algorithms respond diversely to varying imaging artifacts in a clinical dataset.

Conclusions

This study provides additional information in the understanding of the mechanism by which BZ is associated with enhanced cardiovascular mortality. Previously, inducible MVT during EPS has been shown to be highly predictive of both sudden death as well as total mortality [22, 41]. Our data provide a confirmation of the link between BZ quantification and arrhythmias in a human cohort. We also demonstrated that BZ is not simply a surrogate measurement of infarct size or infarct surface area based on geometric relationships, but contains additional data. Border zone is a more specific predictor of inducible arrhythmia than LVEF, so has the potential for being an improved predictor of mortality due to SCD. Although no such relationship on outcome was identified in this study, this may be related to the limited sample size. Further studies are warranted, both to correlate imaging findings of BZ to histology, and to assess the prognostic significance of infarct characteristics such as infarct size and BZ.

Author’s contributions: JR designed study, created software for border zone analysis, performed analysis, and drafted manuscript. DL and EW conceived of study, assisted with the border zone analysis and manuscript creation. DB executed the original prospective study and data collection from which a large portion of this retrospective study was based on. AK, RP, and JG helped refine the study goals, and assisted with manuscript creation. All authors read and approve the final manuscript.

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