Increased Wave Break During Ventricular Fibrillation in the Epicardial Border Zone of Hearts With Healed Myocardial Infarction

Toshihiko Ohara, MD; Keiko Ohara, MD; Ji-Min Cao, MD; Moon-Hyoung Lee, MD; Michael C. Fishbein, MD; William J. Mandel, MD; Peng-Sheng Chen, MD; Hrayr S. Karagueuzian, PhD

Background—The action potential duration (APD) restitution hypothesis of wave break during ventricular fibrillation (VF) in the epicardial border zone (EBZ) of hearts with chronic myocardial infarction is unknown.

Methods and Results—VF was induced by rapid pacing, and the EBZ with the two adjoining sites (right ventricle and lateral left ventricle) were sequentially mapped in random order in 7 open-chest anesthetized dogs 6 to 8 weeks after left anterior descending artery occlusion and in 4 control dogs. At each site, 3 seconds of VF was mapped with 477 bipolar electrodes 1.6 mm apart. The number of wave fronts and approximate entropy were significantly (P<0.01) higher in the EBZ than all other sites in both groups independent of the rate of invasion of new wave fronts and epicardial breakthroughs. The higher wavelet density in the EBZ was caused by increased (P<0.01) incidence of spontaneous wave breaks. There was no difference between the two groups in either reentry period (80 episodes) or VF cycle length. Reentry in the EBZ had a smaller core perimeter, slower rotational speed, and a small or no excitable gap (P<0.01), often causing termination after one rotation. The dynamic monophasic action potential duration restitution curve in the EBZ had longer (P<0.01) diastolic intervals, over which the slope was >1. Connexin43-positive staining was significantly (P<0.01) and selectively reduced in the EBZ.

Conclusions—A selective increase in wave break and alteration of reentry occur in the EBZ during VF in hearts with healed myocardial infarction. Increased wave break in the EBZ is compatible with the action potential duration restitution hypothesis. (Circulation. 2001;103:1465-1472.)

Key Words: myocardial infarction ■ fibrillation ■ action potentials ■ waves
by permanent left anterior descending coronary artery (LAD) occlusion distal to the first diagonal branch.15 Dogs were then restudied 6 to 8 weeks later in the open-chest state.1,16,17 Four dogs with no LAD occlusion served as control.

**Monophasic Action Potential Restitution**

Dynamic monophasic APD restitution curves were constructed in all mapped regions of MI and the control groups.12,18

**Computerized Mapping During VF**

VF was induced by rapid pacing as described previously.19 Three seconds after the onset of each VF episode, 3 seconds of data were acquired with a 3.2×3.8-cm plaque containing 477 bipolar electrodes 1.6 mm apart.20 The VF was then converted by electrical shocks of 15 J with internal paddles. After 5 to 10 minutes of recovery, VF was reinduced and activation mapped from another site. The order of mapping sites was randomized. Activation pattern and wave front number were determined by dynamic visualization.3,20 The VF cycle length (CL) was measured on each channel over the entire 3 seconds of VF in each dog and was averaged. Typically, in each dog, >400 channels were of acceptable quality for VF CL calculations, for a total of >9000 beats in each dog (20 to 22 cycles over 400 channels).1,16,17 Data are presented as mean VF CL in each group.

**Measurement of Reentry Core Perimeter**

Once a reentrant wave front was detected, the inner edge of the core was traced for one complete rotation, and core perimeter and average conduction velocity (CV) around the core were measured.1,12,16,21 The CV of nonreentrant wave fronts, along and across the fiber, were also measured during VF and during regular pacing by dividing conduction time over distance in a region of consecutively activated adjacent electrodes.22 The effective refractory period (ERP) during VF was estimated by perpendicular wave-wave interaction as described before1,12 and during regular pacing at 400-ms CL by the extrastimulus method.2,11,12

**Approximate Entropy**

Three seconds of a bipolar electrogram sampled at 1 kHz (3000 points) was used for approximate entropy (ApEn) calculations (temporal complexity), as described previously.23

**Histological Studies**

Ventricles were sectioned from apex to base in 1- to 1.5-cm-thick transverse slices, and Cx43 was stained with antibodies by the ABC method (Santa Cruz Biotechnology Inc).24 The percentage of positive Cx43 staining was calculated in ≥20 different fields in each section by Image-Pro software (Media-Cybernetics). Infarct size was determined from hematoxylin and eosin transverse sections by planimetry.15,25

**Statistical Analysis**

All statistical analyses were performed with GB-STAT,26 Student’s t tests for single comparison, and ANOVA for multiple comparisons with Newman-Keuls post hoc analyses. The null hypothesis was rejected at a value of P≤0.05. Results are expressed as mean±SD.

**Results**

**Infarct Size and Location**

In all dogs LAD occlusion caused MI in the anteropapical site sparing 30 to 100 epicardial cell layers over the infarct (EBZ) (Figure 1). The mean infarct size was 19±3% of the total (225±15 g) left ventricular (LV) mass.

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**Figure 1.** Histopathological and immunostaining with Cx43. A, Gross anatomy of transmural section of infarcted LV (8 weeks after LAD occlusion). Whitish areas are scar tissue. B, Hematoxylin and eosin microscopic section showing EBZ (Epi) over deeper scar tissue (infarct). C and E, Longitudinally and transversely oriented normal myocardium showing numerous Cx43 stains. D and F show similar sections in EBZ, with reduced Cx43 staining. Bar is 50 μm.
During regular pacing, no conduction block occurred over the entire EBZ in each dog, indicating that the observed blocks during the VF were functional in nature. The ERP during the VF, estimated by the method of perpendicular wave-wave interaction, was significantly longer (P < 0.01) in the EBZ than in the anterior LV of control dogs (115 ± 6 ms versus 87 ± 11 ms). Significantly fewer (P < 0.001) epicardial breakthroughs occurred in the EBZ than at all other sites in both groups (Figure 2 and 3). Epicardial breakthroughs occurred 6.75 ± 0.9/s in the normal zones of both groups (11% of all wave front activity), whereas in the EBZ, the breakthroughs occurred only 1.72 ± 0.75/s, accounting to 0.73 ± 0.14% of overall wave front activity. The rate of invasion of new wave fronts from adjoining lateral regions was not significantly different in the EBZ compared with normal non-MI anterior LV (48.3 ± 6.6/s versus 51 ± 5.3/s, P = NS). The average num-

### Table 1. Wave Front and Electrophysiological Properties in Dogs With Chronic Myocardial Infarction

<table>
<thead>
<tr>
<th></th>
<th>RV</th>
<th>EBZ</th>
<th>Lateral LV</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Threshold, mA</td>
<td>0.57 ± 0.26</td>
<td>0.81 ± 0.30</td>
<td>0.58 ± 0.22</td>
<td>§</td>
</tr>
<tr>
<td>ERP, ms</td>
<td>168.2 ± 11.3</td>
<td>188.0 ± 18.0</td>
<td>168.0 ± 16.3</td>
<td>§</td>
</tr>
<tr>
<td>CL during VF, ms</td>
<td>134.1 ± 18.0</td>
<td>135.2 ± 17.3</td>
<td>136.0 ± 18.9</td>
<td>NS</td>
</tr>
<tr>
<td>Wavelet n/10 ms</td>
<td>3.08 ± 0.27</td>
<td>4.29 ± 0.24</td>
<td>2.98 ± 0.29</td>
<td>††</td>
</tr>
<tr>
<td>ApEn</td>
<td>0.212 ± 0.010</td>
<td>0.251 ± 0.024</td>
<td>0.184 ± 0.013</td>
<td>*‡¶</td>
</tr>
<tr>
<td>CV along, cm/s</td>
<td>55.5 ± 14.4</td>
<td>35.3 ± 6.4</td>
<td>57.4 ± 14.7</td>
<td>††</td>
</tr>
<tr>
<td>CV across, cm/s</td>
<td>35.7 ± 6.2</td>
<td>28.7 ± 5.6</td>
<td>36.3 ± 5.5</td>
<td>†§</td>
</tr>
<tr>
<td>Anisotropic ratio</td>
<td>1.66 ± 0.56</td>
<td>1.19 ± 0.24</td>
<td>1.68 ± 0.55</td>
<td>††</td>
</tr>
<tr>
<td>Reentry episodes</td>
<td>20</td>
<td>17</td>
<td>15</td>
<td></td>
</tr>
<tr>
<td>Core perimeter, cm</td>
<td>2.65 ± 0.36</td>
<td>1.95 ± 0.26</td>
<td>3.06 ± 0.26</td>
<td>††**</td>
</tr>
<tr>
<td>Reentry period, ms</td>
<td>100.5 ± 9.8</td>
<td>100.3 ± 10.4</td>
<td>102.8 ± 12.6</td>
<td>NS</td>
</tr>
<tr>
<td>Reentry CV, cm/s</td>
<td>25.9 ± 3.0</td>
<td>19.2 ± 2.7</td>
<td>29.5 ± 2.9</td>
<td>††**</td>
</tr>
<tr>
<td>Max slope</td>
<td>2.27 ± 0.5</td>
<td>3.61 ± 0.34</td>
<td>1.36 ± 0.14</td>
<td>†§¶</td>
</tr>
<tr>
<td>DI &gt; 1, ms</td>
<td>16.0 ± 0.9</td>
<td>23.2 ± 1.3</td>
<td>12.8 ± 1.9</td>
<td>*¶</td>
</tr>
</tbody>
</table>

ERP was determined during regular pacing at 400-msec CL by the extrastimulus method; wavelet n/10 ms is the number of independent wave fronts in the mapped region counted in 10-ms increments over a 3-second VF interval (300 frames).

EBZ vs lateral LV, *P < 0.01, †P < 0.001.
EBZ vs RV, ‡P < 0.05, §P < 0.01, ¶P < 0.001.
RV vs lateral LV, ¶P < 0.05, **P < 0.001.

### Table 2. Wave Front and Electrophysiological Properties in Dogs With No Myocardial Infarction

<table>
<thead>
<tr>
<th></th>
<th>RV</th>
<th>Anterior LV</th>
<th>Lateral LV</th>
<th>P</th>
</tr>
</thead>
<tbody>
<tr>
<td>Threshold, mA</td>
<td>0.48 ± 0.12</td>
<td>0.52 ± 0.17</td>
<td>0.50 ± 0.14</td>
<td>NS</td>
</tr>
<tr>
<td>ERP, ms</td>
<td>169.5 ± 6.5</td>
<td>171.0 ± 4.1</td>
<td>168.0 ± 16.3</td>
<td>NS</td>
</tr>
<tr>
<td>CL during VF, ms</td>
<td>131.4 ± 3.3</td>
<td>134.1 ± 4.1</td>
<td>131.3 ± 4.3</td>
<td>NS</td>
</tr>
<tr>
<td>Wavelet n/10 ms</td>
<td>2.96 ± 0.05</td>
<td>3.03 ± 0.06</td>
<td>2.98 ± 0.05</td>
<td>NS</td>
</tr>
<tr>
<td>ApEn</td>
<td>0.189 ± 0.018</td>
<td>0.184 ± 0.012</td>
<td>0.186 ± 0.026</td>
<td>NS</td>
</tr>
<tr>
<td>CV along, cm/s</td>
<td>53.5 ± 14.0</td>
<td>52.7 ± 12.0</td>
<td>53.0 ± 13.0</td>
<td>NS</td>
</tr>
<tr>
<td>CV across, cm/s</td>
<td>36.5 ± 5.5</td>
<td>38.6 ± 5.9</td>
<td>36.0 ± 6.0</td>
<td>NS</td>
</tr>
<tr>
<td>Anisotropic ratio</td>
<td>1.50 ± 0.44</td>
<td>1.50 ± 0.37</td>
<td>1.52 ± 0.44</td>
<td>NS</td>
</tr>
<tr>
<td>Reentry episodes</td>
<td>10</td>
<td>10</td>
<td>8</td>
<td></td>
</tr>
<tr>
<td>Core perimeter, cm</td>
<td>3.10 ± 0.44</td>
<td>3.12 ± 0.40</td>
<td>3.00 ± 0.40</td>
<td>NS</td>
</tr>
<tr>
<td>Reentry period, ms</td>
<td>100.0 ± 12.2</td>
<td>104.6 ± 7.9</td>
<td>101.3 ± 7.4</td>
<td>NS</td>
</tr>
<tr>
<td>Reentry CV, cm/s</td>
<td>30.1 ± 2.4</td>
<td>30.3 ± 3.8</td>
<td>29.1 ± 2.5</td>
<td>NS</td>
</tr>
<tr>
<td>Max slope</td>
<td>1.53 ± 0.13</td>
<td>1.58 ± 0.17</td>
<td>1.45 ± 0.13</td>
<td>NS</td>
</tr>
<tr>
<td>DI &gt; 1, ms</td>
<td>13.7 ± 1.5</td>
<td>14.3 ± 1.5</td>
<td>13.7 ± 2.1</td>
<td>NS</td>
</tr>
</tbody>
</table>

ERP was determined during regular pacing at 400-msec CL by the extrastimulus method; wavelet n/10 ms is the number of independent wave fronts in the mapped region counted in 10-ms increments over a 3-second VF interval (300 frames).

RV vs anterior LV; RV vs lateral LV; anterior LV vs lateral LV.
The number of wavelets in the mapped region was significantly ($P<0.001$) higher in the EBZ than all other sites in both groups. (Tables 1 and 2). The higher number of wave fronts in the EBZ compared with other sites resulted from the more frequent spontaneous breakups. Breakup occurred when a wave front encountered refractory tissue from a previous activation (Figure 2B and Figure 3C). The mean number of wave breaks (counted over 3 seconds at 10-ms intervals) was significantly ($P<0.001$) higher in the EBZ than in the left anterior wall in the non-MI group ($19.3\pm2.5/s$ versus $8\pm1.4/s$). To ensure reproducibility of the observed results, wave front number and ApEn (see below) determinations were repeated by the same investigator and by a second investigator blinded to the origin of the data (a total of 33 episodes of VF in 7 dogs). A linear regression analysis that was run between the intraobserver and interobserver measurement outcomes yielded a correlation coefficient of 0.97 and 0.96, respectively, for both measurements.

**Figure 2.** Selected snapshots obtained sequentially from 3 different sites in a dog 8 weeks after LAD occlusion. A, Plaque in RV; at time 0, two wave fronts enter mapped area (arrow), with third breakthrough wave front in middle right of plaque. These fronts propagate to edges of plaque without breakups. B, In EBZ, total of 4 wave fronts. Two wave fronts entering plaque (arrows) undergo breakups with conduction blocks occurring in middle of both fronts. Third (bottom) and fourth (middle right) wave fronts undergo block (bottom) and collision, with subsequent annihilation in both cases. C, In normal noninfarcted LV, 3 wave fronts (time 0) propagate downward (arrows) and coalesce together before exiting plaque. Most recent activated site is shown in red, then pink, then yellow, then green, and finally, blue. Persistence of each color is 10 ms. Selected bipolar electrograms (D, E, and F) are shown next to each panel with similar CLs.

### Wave Front Collisions

In addition to head-and-tail interactions, two wave fronts could collide head on or roughly perpendicular to each other. The consequences of head-head wave front collisions were different in the EBZ compared with all other sites in both groups. Collision in the EBZ ($5\pm1/s$, $P=NS$, compared with EBZ) caused annihilation in 54% of the episodes and survival in 46% of episodes with propagation in a direction perpendicular to the direction of collision. In the normal anterior LV however, collision ($7\pm1/s$, $P=NS$, compared with EBZ) caused annihilation in 71% of the episodes and survival in 29%, with subsequent propagation of the wave front in a direction perpendicular to the direction of collision. The differences in the outcome of the head-on collisions (annihilation and survival) were significantly ($P<0.01$) different in the EBZ compared with similar LV anterior site in the non-MI control group.
The VF CL was not different at all three sites in both groups; however, the ApEn was significantly \((P<0.05)\) higher in the EBZ than all other sites in both groups (Tables 1 and 2).

Reentrant Wave Front Characteristics During VF
A total of 80 episodes of complete reentrant activations were captured during VF in both groups (Figure 4). The core perimeter of the reentry was significantly \((P<0.01)\) shorter and had slower speed in the EBZ compared with all other sites in both groups \((P<0.05; \text{Tables 1 and 2 and Figure 4})\). The core of the reentry in the EBZ became roughly circular instead of elliptic, which was seen at all other sites in both groups (Figure 4). The similar reentry period at all sites narrowed or closed the excitable gap at the EBZ because of the longer ERP in the EBZ during VF. The smaller excitable gap during reentry in the EBZ caused the reentry to terminate after only 1 rotation (4 episodes) or after an incomplete “reentry” (10 episodes). Two to 3 consecutive rotations during reentry were seen in a total of 11 episodes at normal sites in both groups.

CV of Nonreentrant Wave Fronts During VF
A total of 179 CV measurements of nonreentrant wave fronts were made during VF in both groups. At all sites, the CV was significantly slower in the EBZ both along and across the fiber orientation (Tables 1 and 2). Although the directional differences in CV were decreased in the EBZ compared with normal, CV still remained significantly \((P<0.05)\) faster along than across the fiber orientation in the EBZ.

Dynamic APD Restitution Relation
The maximum slope and the range of DIs within which the slope of the APD restitution curve remained \(>1\) were significantly higher in the EBZ than all other sites in both groups (Table 1 and Figure 5).

Cx43 Distribution
Immunostaining showed a selective and significant \((P<0.001)\) reduction of Cx43 distribution in the EBZ. Cx43 was reduced both in the end-to-end and side-to-side locations (Figure 1). Cx43 was \(0.77\pm0.17\%\) of the total tissue area in the normal zones in both groups versus \(0.17\pm0.06\%\) in the EBZ \((P<0.001)\) (Figure 1). At distances 3 to 5 mm away from the EBZ, however, staining of Cx43 increased and became normal 1 cm away from the EBZ. These findings are consistent with the earlier electron micrographs of gap junctions and connexin staining in hearts with healed MI.

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**Figure 3.** Selected snapshots obtained sequentially from 3 different sites in a dog with no LAD occlusion. All panels were obtained from similar sites as in Figure 2. At all sites, note presence of relatively large wave fronts (arrows) that either collide with each other (A), join with each other (B), or propagate in different directions (C) to exit from plaque site. Selected electrograms from each site are shown next to panels (D, E, and F). Color code as in Figure 2.
Discussion

Major Findings
The increased incidence of spontaneous wave break in the EBZ during VF compared with adjoining normal sites and to comparable sites in control non-MI dogs constitutes a major finding in the present study. A second major finding of this study is the loss of the normal relation in the EBZ between ERP and reentry period. Although in the normal ventricle lengthening of the ERP lengthens the period of reentry, the longer ERP in the EBZ, however, failed to lengthen the reentry period. Finally, the slower CV in the Cx43-deficient EBZ appeared to prevent faster activation during VF (CL shortening) that might have been caused by the denser EBZ wavelets.

Why Regional Differences in Wave Front Dynamics?
The difference of the APD restitution properties in the EBZ compared with adjoining sites might provide a basis for the observed regional differences in wave front dynamics. Simulation studies showed that an increase in the range of DIs of the APD restitution curve over which the slope is $>1$ promotes wave front breakups. The dynamic APD restitution relation, which closely resembles the restitution during VF, showed that the EBZ cells manifest higher maximum slopes and a significantly wider range of DIs over which the slope remained $>1$. Although the restitution hypothesis is compatible with the higher incidence of wave break in the EBZ, additional mechanisms of wave break may also be operative in the EBZ. For example, increased ERP and partial cellular uncoupling (decreased Cx43) might increase vulnerability to wave break. As the number of wavelets in the EBZ increased, so did the complexity of VF, as indicated by the increase in the ApEn. This finding is consistent with previous reports on isolated porcine right ventricle (RV). Whereas complexity and wave front density increased, the VF CL however did not change, an effect probably caused by the severe CV slowing in the EBZ that offsets the potential of more frequent activations by the denser wave fronts.
Dynamics of Reentry in EBZ

Severe conduction slowing along and across the fiber in the Cx43-deficient EBZ changed the elliptical shape of the reentry core to a more or less circular one. Such a change in the shape of the reentry core may be caused by a significant ($P<0.01$) decrease in the anisotropic ratio from $1.6 \pm 0.5$ in normal sites in both groups to $1.19 \pm 0.24$ in the EBZ (Table 1). The slowing of the CV in the EBZ during reentry, however, did not prolong the period of the reentry because of a concomitant reduction in the core size of the reentry (scaling). Although the mechanism of core shrinking in the EBZ remains to be defined, we speculate that reduced gap-junctional Cx43 might promote such an effect because the longer ERP in the EBZ would have increased and not decreased the core size. More work is needed to determine the mechanism(s) by which the reentry core size versus reentry period relation becomes altered in the EBZ.

Clinical Relevance

Although speculative, it is possible that increased susceptibility to wave break observed during VF in the EBZ might be related to increased vulnerability to VF in patients with chronic MI, because increased propensity to wave break might lead to VF. This speculation, however, needs mapping data on human ventricles with chronic MI to ascertain the wave break hypothesis of VF in humans.

Limitations of the Study

Because recordings from different sites during a VF episode were not simultaneous, it could be argued that the observed regional differences in wave front dynamics might be caused by differences in different VF episodes. However, the presence of similar wave front dynamics, reentry morphology, and CV at a given site on repeat VF episodes (data not shown) and randomized acquisition of data from different sites refutes this possibility. It may be argued that our maps are 2D and not 3D, raising doubt on the interpretation of the EBZ maps. However, the EBZ in our model is a relatively thin rim of tissue made of about 30 to 60 cell layers (sometimes much less) (Figure 1), with tissue beneath the EBZ made essentially of nonviable scar tissue. As the result, the EBZ remains relatively immune to interference by transmurally conducted wave fronts, validating the interpretation of the EBZ maps.

Conclusions

We conclude that there is increased incidence of wave break during VF in the EBZ of hearts with healed MI. The mechanism of increased vulnerability to wave break is compatible with the APD restitution hypothesis.

Acknowledgments

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References


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