Effects of anisotropy on the development of cardiac arrhythmias associated with focal activity

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Abstract The anisotropy that normally exists in the myocardium may be either enhanced in peri-infarction zones by loss of lateral cell connections or reduced by redistribution of gap junctions. To test how the degree of anisotropy affects the development of ectopic focal activity, we carried out computer simulations in which a model of an ectopic focus is incorporated as the central element of a two-dimensional sheet of ventricular cells. At low values of intercellular coupling conductance ($G_c$), the focus region is spontaneously active, but the limited intercellular current flow inhibits propagation. At high $G_c$, automaticity is suppressed by the loading effects of the surrounding cells. At intermediate $G_c$, the ectopic activity may propagate into the sheet. In the case of isotropic coupling, the minimum size of the focus region for propagation to occur (in terms of number of collaborating cells within the focus) is as small as approximately ten cells, and this number decreases with increasing anisotropy. Thus, the presence of anisotropy facilitates the development of ectopic focal activity. We conclude that the remodeling that occurs in peri-infarction zones may create a substrate that either facilitates (enhanced anisotropy) or inhibits (reduced anisotropy) the development of cardiac arrhythmias associated with ectopic focal activity.

Keywords Anisotropy · Computer simulations · Electrophysiology · Gap junctions · Heart · Ischemia · Ventricular cells

Introduction

Many experimental and theoretical studies have focused on action potential propagation in cell pairs, linear strands, or multi-dimensional structures of cardiac cells. Multi-dimensional cardiac tissue differs in conduction behavior from the simple linear models in several respects. Myocyte structure is asymmetrical, with average adult myocytes being 100–150 µm in length and 15–25 µm in width [13, 23], with the cellular arrangement of these cells giving the tissue a structural anisotropy. Anisotropy is further enhanced by the spatial distribution of gap junctions – ensembles of intercellular ion channels providing the electrical pathway between neighboring cells – such that the propagation velocity in the transverse direction is lower than that in the longitudinal direction, with ratios of 1 to 2.7 in the ventricle and up to 1 to 12 in the atrial crista terminalis [2, 11, 19, 36].

The interactions among heart cells with spatial variability in action potential properties and intercellular coupling are even more complex in the ischemic heart. Cellular electrophysiological dysfunction may lead to conduction slowing and/or block which may facilitate re-entrant arrhythmias, or, alternatively, to localized areas of ectopic activity which may generate non-reentrant arrhythmias. Numerous studies of healed infarctions in ventricular tissue have shown fractionated electrical activity indicative of the partial electrical uncoupling of groups of cells [10, 18, 37]. Direct evidence for reduced coupling between cells comes from immunohistochemical studies showing that there are smaller and fewer gap junctions in peri-infarction zones [24, 32]. Anisotropy may be enhanced by a particular loss of lateral connections compared to end-to-end connections among heart cells [24]. On the other hand, anisotropy may be reduced by a redistribution of gap junctions, tending to become...
randomly spread over the cell surfaces, and not in discrete intercalated disks at the end-to-end cell contacts as in normal myocardium [24, 32]. Many episodes of acute ischemia occur in such “abnormal” tissue, i.e., that has a spatially inhomogeneous distribution of intercellular coupling resulting from prior injury, which may serve as a substrate from which arrhythmias can arise [26].

The actual membrane properties of cells of a pathologically present ectopic focus are not clearly known. Recent microelectrode recordings from an ectopic focus area removed from a patient with atrial tachycardia [3] showed a group of depolarized, spontaneously active cells which, under the in vitro conditions of the recordings, showed considerable electrical uncoupling from the surrounding atrial cells. Several types of heart cells within the normal heart express spontaneous diastolic depolarization which can lead to focal activity either as a normal process (e.g., the sinoatrial node) or as an abnormal process (e.g., the atrio-ventricular node, the coronary sinus region, or the Purkinje system). In our previous work, in which we studied the electrical interactions between a spontaneously pacing cell model representing an ectopic focus and a real, isolated ventricular myocyte [20, 38], we used our previously published sinoatrial cell model [40] as a model of the ectopic focus. Even at this level of a pair of cells with different membrane properties, the anatomical properties that organize cardiac cells into functional groups could be seen to play a critical role in enabling propagation.

The multiple effects of anisotropy on an already propagating wave, produced by direct stimulation, have been widely studied, both in computer simulations [7, 15, 21, 34] and in cell culture systems with cultures grown on a directed collagen matrix to form an anisotropic network [5, 6]. We recently extended our “coupling clamp” technique in which we couple a real cell to a real-time simulation of a model cell to incorporate a real cardiac cell as the central element of a two-dimensional sheet of model cells [39]. In this sheet composed of 7x7 elements the coupling conductances may be different in the X and Y directions and a specific region of lack of coupling conductance may serve as a resistive barrier. We determined the critical size for the successful propagation from a directly stimulated real ventricular cell in the central location into the sheet, and found that this critical size was decreased when anisotropy was present compared to when it was not, and was further decreased when the central site of stimulation was close to a resistive barrier. Much less work has been done on the effects of anisotropy on the initiation of a propagating wave of excitation. The critical size of an automatic focus region was the subject of our previously published theoretical study [16], in which we simulated a two-dimensional sheet of atrial tissue with a centrally located sinoatrial node area of variable size. In this study, we showed that a partial electrical uncoupling between the automatic area and the quiescent area was required in order for the automatic area to maintain automaticity and still drive the quiescent area. Generation and propagation of the sinoatrial nodal impulse in a two-dimensional sheet of cells has also been studied in large-scale simulations on a massively parallel computer [1]. Winslow et al. [43] used similar simulation techniques to study the generation of ectopic activity in an isotropic two-dimensional sheet of atrial cells, into which a region of inhibited sodium–potassium pump activity was incorporated. They demonstrated that the local sodium overloading produced by the pump inhibition can induce propagating ectopic beats, with the critical size of the overload region for generating propagating ectopic beats depending on the cell-to-cell coupling magnitude.

The initiation of a propagating wave from an automatically active focus involves the input resistance of the syncytial tissue, the ability of the focus region to serve as a current source, and the sensitivity of the focus region to electrotonic interactions with the surrounding quiescent region which will tend to either abolish automaticity or prevent the initiation of propagation. In order to study the way in which anisotropic connectivity affects the initiation of a propagating wave, we carried out computer simulations using a two-dimensional sheet of ventricular cells into which a spontaneously active cell model is incorporated as a central focus element, and for which the intercellular conductances in the X and Y directions can be varied to introduce anisotropy.

**Materials and methods**

**Two-dimensional sheet**

The diagram of Fig. 1 illustrates the two-dimensional sheet we simulated composed of 13x13=169 elements. Each element (except those on the edges of the sheet) has resistive coupling to four adjacent elements and the (ohmic) coupling conductances are assumed to be a constant $G_y$ along the X-axis and a possibly different constant $G_x$ along the Y-axis. Elements for the central portion of the sheet are numbered along the X- and Y-axes as $-2, -1, 0, 1, 2$. The numbering of the entire sheet extends from $-6$ to $+6$ along each axis as indicated. The upper right quadrant (thick lines) consists of 49 elements for which the focus cell at $(0,0)$ (filled square in the center of the total sheet) is at the lower left position. We achieve horizontal and vertical symmetry by the condition that only the central element at $(0,0)$ or the 9 elements in the central 3x3 array (black and gray squares) are spontaneously active. From symmetry, we can then include all of the interactions among elements in the entire sheet by solving only for the potential and coupling currents of the 49 elements of the upper right quadrant, even if $G_x$ and $G_y$ are not equal to each other.

To represent all of these interactions, we let $I_x(i,j)$ be the (ohmic) current which flows to the right of element $(i,j)$, where $V_{ij}$ is the membrane potential of element $(i,j)$. For the elements on the left edge of the upper right quadrant [elements $(0,0), (0,1), \ldots, (0,6)$] we double the computed $I_x$ to include the current which, from symmetry, would have flowed in the opposite direction along the X-axis. Similarly, we let $I_y(i,j)$ be the current which flows up along the Y-axis and we double this current for the elements along the bottom edge of our upper right quadrant [elements $(0,0), (1,0), (2,0), \ldots, (6,0)$] to include the current which would have flowed in the opposite direction down the Y-axis for these elements. Thus, the central element (which has intrinsic spontaneous activity) gets both its $I_x$ and $I_y$ doubled, since it is on both edges. For the other elements in the upper right quadrant the definitions for $I_x$ and $I_y$ are as given except for the edge elements where the coupling current either to the right or up is zero [$I_x=0$ for elements $(6,0), (6,1), \ldots, (6,6)$ and $I_y=0$ for elements $(0,6), (1,6), \ldots, (6,6)$]. We thus reduce the problem of solving a sheet of 169 elements to a problem of solving for only 49 elements.
Membrane models and numerical integration

The membrane model we use for the spontaneously active element at the center of the sheet is our previously published sinoatrial node model [40] (SAN model), with its 32-pF membrane capacitance and currents scaled by a factor of 5 to account for differences in size between nodal and ventricular cells (focus model, central black square). The other elements are represented by the phase-2 ventricular cell model of Luo and Rudy [25] as updated by Zeng et al. [44] (LR model, open squares). Elements directly surrounding the central element (gray squares) are represented by either the focus model or by the LR model. The upper right quadrant of 49 elements (thick lines) is actually solved (see “Materials and methods”).

In this study we use a simple first-order numerical integration scheme with a fixed time step of 5 µs. At each time step, the (ohmic) coupling currents are calculated as described above and then are used together with the calculated membrane currents for each element to integrate the membrane potential for each element of the array. In this implementation of a two-dimensional array of cells, we have treated each element of the array (except the central element) as one isopotential cell. We allow the effective size of this central element to be varied, as if this element represents a small cluster of N cells infinitely well coupled to each other. The effective size of this central element can be varied by simply scaling the coupling currents associated with that element. To increase the effective size of this element by a factor of N, so that it represents N focus cells, its membrane potential is integrated with its computed coupling currents divided by N, as we have previously described in detail [41, 42].

We followed a protocol in which we first ran the simulation with the central element uncoupled from the sheet for two cycles of automaticity and then coupled the central element and the other elements of the sheet with desired values of \( G_x \) and \( G_y \), for at least 20 s to allow development of steady-state behavior (see Figs. 2, 3). The simulations were coded and compiled using Active Ada (version 5.3, Alsys, Boston, Mass., USA) and run on a personal computer equipped with a 450-MHz Pentium II processor. Typical computation time for a 20-s simulation was \( \approx 20 \) min.

Results

Activation of an isotropic sheet

Figure 2 illustrates the effects of coupling a central focus element into the 13×13 element sheet as diagrammed in Fig. 1. Each element, except the central one, is represented by the LR model and the central element is represented by the focus model with the size set to 8 (see Materials and methods). This size setting makes the central element equivalent to 8 focus cells with infinitely good coupling among these 8 cells such that they can act as a unit to provide current to the surrounding elements of the sheet. For this example we set \( G_x = G_y \), so that the sheet is isotropic. The results are plotted in three panels for \( G_x, G_y \) values of 15, 20, and 50 nS, respectively. Each panel shows results for elements 0, 1, …, 6 along the middle row of the sheet. The data displayed are the last two action potentials that occurred in the central element while it was uncoupled from the rest of the sheet, followed by the activity which develops over the \( \approx 19 \) s period after coupling was established.

In Fig. 2A (\( G_x, G_y = 15 \) nS), the focus element (0,0) shows two action potentials during the uncoupled period and then, after coupling has been turned on (vertical arrow), a train of action potentials which do not propagate into the sheet (thick dotted line). In the ventricular cells (1,0), (2,0), … (6,0) (thin solid lines), only subthreshold depolarizations are observed, which are largest in adjacent cell (1,0). At this value of coupling conductance, the focus element continues to pace upon coupling (at a
slower rate as compared to the uncoupled state due to the loading effects of the sheet), but it does not drive the sheet ("pace but not drive"). The low coupling conductance inhibits current flow into the other sheet elements and thus activation of the sheet does not occur.

If we now increase coupling conductance to 20 nS, and thus increase coupling current, we observe the results shown in Fig. 2B. The increased coupling current induces larger depolarizations in the adjacent cells, but these are still subthreshold. After a ≈6-s period of increasing interbeat intervals spontaneous activity of the focus element ceases due to increased loading effects as compared to the previous case of \( G_x = G_y = 15 \text{ nS} \). A further increase in coupling conductance to 50 nS (Fig. 2C) leads to almost instantaneous cessation of automaticity. After coupling is established there are only two action potentials in the focus element at (0,0) with a size factor of 10 into the two-dimensional sheet of Fig. 1 with \( G_x = G_y \) (isotropic sheet). Each panel shows the membrane potential (\( V_m \)) for elements 0, 1, …, 6 along the middle row of elements (\( Y = 0 \)), with the activity of the focus cell (element 0) shown as a dotted line. Coupling is turned on at the time indicated by the vertical arrows. Note axis breaks. A Failure of propagation ("pace but not drive") at \( G_x = G_y = 15 \text{ nS} \). B Successful propagation ("pace and drive") at \( G_x = G_y = 20 \text{ nS} \). C Failure of propagation ("not pace") at \( G_x = G_y = 50 \text{ nS} \).

Critical size factor

When we ran a large number of these simulations, always keeping \( G_x = G_y \), we were able to delineate the relationship between the value of \( G_x = G_y \) and a critical size factor for the central focus element (the size factor below which the sheet was not activated and above which the sheet was activated), as shown in Fig. 4 as the solid line and filled square symbols. This relationship thus defines a region of parameter values for which successful action potential propagation into the sheet is observed ("pace and drive", region labeled "PD"). The solid line with open squares marks the transition between the PND and NP regions. The horizontal dashed line with open squares marks the transition between the PND and NP regions. The critical size factor for successful propagation from the central focus element into the 13×13 element sheet as diagrammed in Fig. 1 with \( G_x = G_y \) (solid line with filled squares). Results for smaller and larger sheets are shown as a dashed line with filled circles (7×7 element sheet) and a dotted line with filled triangles (27×27 element sheet), respectively. Successful propagation occurs for parameter values in the region labeled "PD" ("pace and drive"). Failure of propagation occurs either as a failure to drive ("pace but not drive", region labeled "PND") or as a failure to pace ("not pace", region labeled "NP"). The solid line with open squares marks the transition between the PND and NP regions.
Effects of sheet size

To test whether the size of our two-dimensional sheet, i.e., the 13×13 element sheet as diagrammed in Fig. 1, is sufficient to simulate the electrotic loading effects of a sheet of ventricular cells on a central focus element, we repeated our simulations with larger and smaller arrays. The smaller sheet we used had 7×7 elements and thus each row (and column) had elements –3 through 3 instead of –6 through 6 (see Fig. 1). The larger sheet was composed of 27×27 elements with elements –13 through 13 along each row (and column). The results obtained with the larger sheet are plotted in Fig. 4 as a dotted line with closed triangles. At all values of coupling conductance tested the results were identical to those obtained with the “standard” 13×13 element sheet, indicating that the 13×13 sheet is sufficiently large. Actually, the results for the smaller sheet, plotted in Fig. 4 as a dashed line with closed circles, were very close to those obtained with the 13×13 and 27×27 arrays. Only at coupling conductances >40 nS, resulting in larger space constants, was the critical size factor slightly underestimated.

Two types of propagation failure

The other relationship shown in Fig. 4 with a solid line and open squares is the relationship obtained when we ran simulations to determine the type of action potential propagation failure in the “not-PD” region. This relationship marks the transition between two regions of parameter values with different types of propagation failure, labeled “PND” and “NP”. At low values of coupling conductance, only little coupling current flows into the sheet. As a consequence, the central focus element paces but it is not able to drive the sheet. Failure of action potential propagation thus occurs as “pace but not drive” (PND) as illustrated in Figs. 2A and 3A. At higher values of coupling conductance, the spontaneous activity of the central focus element is suppressed due to increased electrotic loading effects and propagation failure then occurs as “not pace” (NP) as illustrated in Figs. 2B,C and 3C.
(asynchronous) activation of the sheet that actually decreases the demand for current to activate the entire sheet. This is illustrated in Figs. 6 and 7.

In Fig. 6 we have plotted – for the action potentials of the central focus element indicated with asterisks in Fig. 5 – the total coupling current \( I_x + I_y \) for element (0,0) as well as its components flowing in the \( X \) direction \( I_x \) and \( Y \) direction \( I_y \). The total coupling current was plotted with a positive value leaving element (0,0), thus producing a repolarization. In Fig. 6D–F, we also included the associated voltage waveforms of elements (0,0), (1,0), and (0,1). The mechanisms for the activation of these adjacent elements are shown in the middle and right panels of Fig. 6.

In the middle panels of Fig. 6, we plotted the coupling current in the \( X \) direction for element (0,0) (Fig. 6B) and the membrane potential of element (1,0), which received this current (Fig. 6E). Changing the value of \( G_y \) from 20 nS to either 15 or 25 nS did not significantly change the magnitude of the coupling current in the \( X \) direction before activation of element (1,0) (Fig. 6B). However, the response of the membrane potential of element (1,0) to this current was quite different (Fig. 6E). When \( G_y \) was lowered to 15 nS, element (1,0) had a larger voltage response as compared to the symmetric case, which raised the membrane potential to threshold as shown by the thin solid line in Fig. 6E. The larger voltage response of element (1,0) when \( G_y \) was lowered was produced by the decreased load on this cell in the \( Y \) direction through its connections to elements (1,1) and (1,–1), and thus activation of element (1,0) occurred with the decreased \( G_y \) value (Fig. 6E, downward single arrow). This activation of element (1,0) then sent current back to element (0,0), preventing early repolarization of element (0,0) (Fig. 6D, downward single arrow). Thus, the activation of element (1,0) led to a greater membrane potential in element (0,0), which then activated, with delay, element (0,1), as

![Fig. 6A–F](image-url)
The right panels of Fig. 6 show the mechanism of successful activation of the array when $G_y$ is increased to 25 nS. Figure 6C shows the coupling current for element (0,0) flowing in the $Y$ direction to element (0,1). When $G_y$ is raised or lowered, there is a significant increase or decrease, respectively, of this coupling current. Increases in this coupling current produce a greater response of the membrane potential of element (0,1), as shown by the dotted line in Fig. 6F, and thus lead to activation of element (0,1) (upward single arrow). This activation of element (0,1) then sends current back to element (0,0) and prevents the early repolarization of this element, as shown in Fig. 6D (upward single arrow), which then allows more current to flow in the $X$ direction and produces a delayed activation of element (1,0), as shown by the dotted line in Fig. 6E (upward double arrow).

Figure 7 shows the activation pattern for the time period indicated by the horizontal bar in Fig. 5C on an expanded time scale. As in Fig. 5C, Fig. 7C shows the activation of elements 0, 1, ..., 6 in the central row (“row 0”) of the sheet. The spontaneous activity of the focus element (0,0) appears as a dotted line labeled “0”. The activity of the adjacent intrinsically quiescent element (1,0) is plotted as a thick solid line labeled “1”. Elements (2,0) through (6,0) activate left to right in time and are plotted as thin solid lines (labeled “2” through “6”). The other two panels of Fig. 7 show the activation of elements 0, 1, ..., 6 in rows 1 and 2 of the sheet (Fig. 7B and A, respectively). As indicated by the arrows in Fig. 7B and C, activation of the sheet is asymmetrical, with element (0,1) (thick solid line in Fig. 7B) being activated earlier than element (1,0) (thick solid line in Fig. 7C) as the lower coupling conductance along the $X$-axis inhibits current spread along the $X$-axis and thus delays activation along the $X$-axis. Propagation is more rapid along the $Y$-axis than along the $X$-axis, as illustrated by the earlier activation of element (0,2) (thin line labeled “0” in Fig. 7A) as compared to element (2,0) (thin line labeled “2” in Fig. 7C).

Once activation has succeeded in either direction from the central focus element, the current demand on this focus element is significantly reduced because the element in the direction in which propagation has occurred is no longer a current “sink” but now actually supplies current back to the central element (arrows in Fig. 7B, C). Thus, the presence of anisotropy, by “focusing” current predominantly into one direction of the sheet, shortens the time for which the central element must supply current to the sheet. The elements in the direction that activates first serve to increase the area of cells that have been excited, effectively increasing the size of the focus element as conduction can now proceed from a broader base. This effect is summarized in Figs. 8 and 9.

To delineate the relationship between the values of $G_x$ and $G_y$ and the success or failure of action potential propagation from the central focus element at (0,0) with a size factor of 8 into the two-dimensional sheet of Fig. 1 with $G_x=20$ nS and $G_y=25$ nS as shown in Fig. 5C on an expanded time scale. The three panels show activation of elements 0, 1, ..., 6 (left to right, labeled “0” through “6”) for A row 2 ($Y=2$), B row 1 ($Y=1$), and C row 0 (middle row, $Y=0$). Arrows in B and C point to the activation of element (0,1) (thick solid line in B) which prevents early repolarization of the focus element (0,0) (dotted line in C) and supports activation of element (1,0) (thick solid line in C).
propagation into the sheet, we ran a large number of the simulations illustrated in Fig. 5, maintaining a constant effective focus size. Figure 8 shows results for effective focus sizes of 8, 9, or 10 in panels A, B, and C, respectively. The three possible outcomes “not pace”, “pace but not drive”, and “pace and drive” are plotted as open circles, open squares, and filled diamonds, respectively. The open symbols along the diagonal in Fig. 8A again illustrate the propagation failing at effective focus size 9 (Fig. 8B) or 10 (Fig. 8C), successful propagation is only successful at a high degree of anisotropy (upper left and lower right diamonds). At relatively high values of $G_x$ (or $G_y$) >60 nS, propagation is only successful at a high degree of anisotropy (upper left and lower right diamonds). At intermediate values of $G_x$ (or $G_y$), only moderate degrees of anisotropy are required for successful propagation. At effective focus size 9 (Fig. 8B) or 10 (Fig. 8C), successful propagation can occur in the isotropic case (on-diagonal diamonds), but anisotropy still favors propagation as illustrated by the propagation failing at $G_x=G_y=40$ nS (total conductance of 80 nS), but being successful at $G_x=60$ nS and $G_y=20$ nS (total conductance also 80 nS).

Effects of anisotropy

To delineate the relationship between effective focus size and the degree of anisotropy we fixed $G_x$ at several different values from 15 to 40 nS as shown by the labels in Fig. 9 while we systematically varied $G_y$ and determined the critical size of the focus element that allows activation of the entire sheet. The thick dashed curve labeled “$G_x=G_y$” is the result previously shown as the solid line with filled squares in Fig. 4 for the isotropic state. For the lowest fixed value of $G_x$ (15 nS, solid line with filled circles) there is a very prominent peak in the relationship, with a maximum in the required critical size of the focus occurring at $G_y=15$ nS (the isotropic case). For fixed values of $G_x$ of 20 and 25 nS (dashed line with open squares and dotted line with filled triangles, respectively) there is also a maximum in this relationship at the values ($G_y=20$ and 25 nS, respectively) which make these arrays isotropic. For greater fixed values of $G_x$ (>25 nS, e.g., 40 nS, solid line with open diamonds) there is no such peak at the isotropy value as was seen for the lower values of $G_y$. Note that for each fixed value of $G_x$ ≤ 25 nS there is facilitation in the activation of the sheet (requiring a smaller effective focus size) as the value of $G_y$ is made either greater or smaller than the value for isotropy.

The absence of a peak at the isotropy value for fixed values of $G_x>25$ nS does not imply that the “focusing theory” does not hold for greater fixed values of $G_x$ and $G_y$. This is demonstrated in Fig. 10, where we have plotted results of simulations in which both $G_x$ and $G_y$ were varied to induce anisotropy, but in which total conductance in the X and Y directions ($G_x+G_y$) was constant.

For $G_x=60$ nS and $G_y=20$ nS we see a peak in the relationship at $G_y=40$ nS (solid line with open squares and filled circles, respectively). A value of 0.5 for $G_y/(G_x+G_y)$ (dashed vertical line) represents the isotropic case of $G_x=G_y$. A high degree of anisotropy corresponds with values of $G_y/(G_x+G_y)$ close to 0 ($G_y<<G_x$; focusing of coupling current in the Y direction) or close to 1 ($G_y>>G_x$; focusing of coupling current in the X direction). For total conductance values of either 50 or 80 nS the relationship shows a peak at $G_y/(G_x+G_y)$ value reflecting isotropy. Note that these peaks correspond with the filled triangle at $G_y=25$ nS and the open diamond at $G_y=40$ nS in Fig. 9, respectively. The critical focus size decreases with increasing degree of anisotropy, demonstrating that facilitation in the
were of equal effective size and coupled with the same instead of 4 (see Fig. 1), in which the 9 focus elements squares in Fig. 1), directly coupled to 12 ventricular cells squares in Fig. 1), with open triangles. The filled squares × constant, as it is in our simulations, but will increase magnitude of coupling within the focus region will affect the extent to which the surrounding ventricular cells impose an electrotonic load on the focus. Secondly, and more importantly, the number of direct cell contacts between the focus region and the surrounding cells is not constant, as it is in our simulations, but will increase with increasing focus size, thereby increasing the electrotonic load on the focus. To test to what extent the assumption of a single element focus had affected our results, we repeated our simulations for Fig. 4 with a 3×3 element focus (filled squares in Fig. 1), directly coupled to 12 ventricular cells instead of 4 (see Fig. 1), in which the 9 focus elements were of equal effective size and coupled with the same Gz and Gy values as the other elements in the sheet. These data, with focus size again expressed in terms of the effective number of ventricular cells in the focus region, are presented in Fig. 11 as solid lines with open and filled triangles. Figure 11 shows that results are quantitatively, but not qualitatively, different, with the main effect being an underestimation of the critical number of focus cells at higher values of coupling conductance. At lower values of coupling conductance, there is a shift of the relationship to the left. This may be produced by the smaller effective curvature of the wavefront (larger radius of the focus) as the initiation of propagation from the larger central focus encounters the surrounding ventricular cells. However, the conclusion that ectopic focal activity can originate from a region composed of only approximately ten cells still holds.

**Discussion**

Many studies have examined how structural changes to the myocardium caused by myocardial infarction form a substrate for arrhythmias by forming a structure for reentrant excitation. Such structures include enhanced anisotropy that contributes to the development of spiral waves and wavefront pivoting [8, 12], as well as linear regions of reduced coupling which produce a line of block around which the wavefront can propagate [4, 30]. The main point of the present study is to show how structural changes in the myocardium can affect the initiation of a propagating wave from an ectopic focus.

**Scaling of the model**

We have used a 49-element sheet to demonstrate the effects of anisotropy on the success or failure of propagation from a central focus element into the sheet. Each element of the sheet is represented by a single cell model, but the size of each element can be scaled such that it represents a number of N isopotential cells. With the assumption that each element within the sheet is isopotential, our model is restricted to low degrees of coupling. The values of coupling conductance we used are at least one order of magnitude smaller than the values of gap junctional conductance required for normal conduction in ventricular tissue [14, 31]. Under conditions of myocardial ischemia, discrete groups of surviving cells are separated from other groups of cells by connective tissue [10, 18, 37]. The slow conduction and fractionated waveforms that are characteristic of conduction in ischemic tissue are indications of the successive activation of groups of cells for which the intergroup conductance is limiting the conduction. Our use of a model of elements of variable sizes can be scaled to represent the conduction between groups of cells under the conditions that, within each group, the cells of the group are themselves well coupled to each other and essentially isopotential. That this condition is met under certain ischemic conditions is indicated by the brief duration of each compo-

![Fig. 11](image-url) Effective number of focus cells required for successful propagation into the 13×13 element sheet as diagrammed in Fig. 1 with Gz=Gy, either with a 1×1-element focus (dotted line with filled squares) or a 3×3-element focus (solid line with closed triangles). Successful propagation occurs for parameter values in the region labeled ‘PD’ (“pace and drive”). Failure of propagation either occurs as failure to drive (“pace but not drive”, region labeled “PND”) or as failure to pace (“not pace”, region labeled “NP”). The solid line with open triangles and the dotted line with open squares mark the transition between the PND and NP regions for the 3×3-element focus and the 1×1-element focus, respectively.
ponent of the fractionated signals and also from direct microelectrode recordings from cells within the groups [18]. Thus, the conduction between groups of 100 cells (“size 100 elements”) connected by a conductance of 5 µS would have the same characteristics of electrical loading and also the success or failure of conduction as one would observe between single cells (“size 1 elements”) connected by a conductance of 50 nS. In the case of the connections between large groups of cells, the interconnections might be a short bridge of cells, rather than a direct electrical connection between one cell of each group.

Our results show that there is a critical size for the central focus element to produce propagation out into the two-dimensional sheet. For a continuous one- or two-dimensional system, this would be analogous to the “liminal length” or “liminal area” that must be activated to produce propagation [9, 22, 27, 28]. To further the analogy for discontinuous tissue, the critical size of the central element might be considered the “liminal lump”.

In terms of the number of collaborating cells within the focus, the minimum size of the focus region for propagation to occur may be as small as approximately ten cells (Fig. 11). This number scales with the number of isopotential cells represented by each element of the sheet. If each element represents 100 cells, the “liminal focus size” would be ±1000 cells.

It should be emphasized that our model typically holds for conduction between poorly coupled (groups of) cells, as may be observed in peri-infarction zones and remodeled tissue. Our results should not be extrapolated to the case of the continuous conduction that occurs in normal, well-coupled ventricular tissue. In this case, cytoplasmic resistivity and cell geometry play a major role in determining the activation patterns [14, 35]. Simulation of these activation patterns implicates simulation of individual cells by various amounts of cytoplasmic resistors [31, 35].

Activation of an isotropic sheet

We have shown, using a computer model of a thin sheet of myocardium, that propagation from an ectopic focus into an isotropic sheet of ventricular elements requires that the focus be of sufficient size and that there is a biphasic relationship between size of the focus and the coupling conductance throughout the sheet (Fig. 4). When the coupling conductance is small, the ability of the focus to deliver current to the sheet is limited by the amount of coupling between the cells. Thus, the focus must be quite large to provide enough current to bring the neighboring cells to threshold and initiate a propagating wave. As the conductance increases, the critical size of the focus decreases to a minimum value at \(G_x = G_y = 25\) nS. At values of conductance larger than 25 nS, the size of the focus increases with increasing conductance. For these larger values of conductance there is an increased electrotonic load on the focus cell inhibiting its ability to spontaneously produce an action potential, thus requiring a larger size for the focus. The minimum critical size factor of the focus model required for successful propagation into an isotropic sheet of LR model cells is 8.6, and this minimum occurs at \(G_x = G_y = 25\) nS (Fig. 4). This result is consistent with the results of our recent experimental study [17], in which we demonstrate that a real atrioventricular nodal cell drives an isotropic model atrial sheet more easily than an isotropic model ventricular sheet. In this experimental study, we determined the critical size factor for successful propagation from a real atrioventricular nodal cell in the central location of an isotropic 7×7 sheet of LR model cells, and found that this critical size factor (also corrected for differences in cell size between nodal and ventricular cells; see “Materials and methods”) was 7.4±2.0 (mean ±SEM, \(n = 8\)) and occurred at \(G_x = G_y = 20\) nS.

The results summarized in Fig. 4 demonstrate that successful initiation of a propagating wave occurs only for intermediate values of coupling conductance. In their study of ectopic beating resulting from a centrally positioned region of sodium-overloaded cells in a two-dimensional sheet of atrial cells Winslow et al. [43] also observed a “window” of coupling conductance values allowing propagation of the ectopic beats (their Fig. 4). In a recent modeling study, Saiz et al. [29] obtained similar results. They modeled a strand of cardiac cells using the LR model where half the strand was set to be able to produce early afterdepolarizations (EADs) while the other half was set to control conditions. The coupling conductance between the two halves of the strand was varied and propagation of EADs as ectopic beats occurred only for intermediate values of conductance. At small values of conductance the EADs were unable to propagate and at large values of conductance the EADs were suppressed.

Activation of an anisotropic sheet

In a recent experimental study [39] we determined the critical size factor for successful propagation from a directly stimulated (not spontaneously active) real ventricular cell in the central location of a 7×7 sheet of LR model cells, and found that this critical size factor was \(\pm 7\) in the isotropic case of \(G_x = G_y = 30\) nS and decreased when anisotropy was present compared to the isotropic case. These results are consistent with the results of our present study, in which this size factor was \(\pm 9\) for a spontaneously active cell model as the central element of the sheet (Fig. 4). Our results are also consistent with our earlier study where we coupled our SAN model [40] to an isolated rabbit ventricular cell [38]. We showed that the SAN model had to be of sufficient size and that coupling conductance needed to be of an intermediate value for propagation of the focus to occur. The present study has theoretically extended that work, and shows that the two-dimensional geometry of a sheet of cells can alter the ability of an ectopic focus to propagate.
Altering the geometry of the sheet to include anisotropy enhances the ability of the ectopic focus to initiate a propagating wave by allowing a “focusing” of the coupling current which is then able to bring the neighboring cells to threshold. Introducing anisotropy into the sheet by either decreasing or increasing conductance in one direction causes an asynchrony in the propagation; thus, activation of a cell in the direction of the higher conductance decreases the load on the focus cell (because the cell in that direction has activated) and helps to provide current for activation of the rest of the sheet. This facilitation of propagation upon introduction of anisotropy by either decreasing or increasing \( G \), while maintaining a constant value of \( G \), is only observed at \( G \) values \(< 25 \text{ nS} \) (Fig. 9). At higher values of coupling conductances direct loading effects predominate and an increase in \( G \) leads to an increase in the critical focus size required for successful propagation. However, the “focusing theory” still holds, as shown in Fig. 10: under conditions of constant load (constant \( G + G \)), critical focus size is maximal in the isotropic case and decreases with increasing anisotropy.

Both resistive barriers and enhanced anisotropy are results of myocardial infarction [4, 30]. Aging can also increase the degree of anisotropy as was shown by Spach and Dolber [33]. They showed, in human atrial pectinate muscle, that increasing age increases the longitudinal versus transverse conduction velocity ratio and that this increase correlates with a greater amount of collagenous septa separating the muscle into small fiber bundles in the older muscle. Our simulation results suggest that this underlying structure caused by infarction or aging may facilitate the development of ectopic activity when the membrane properties of some portion of the myocardium are altered in such a way as to become spontaneously active in subsequent ischemic events. This arrhythmogenic effect may be counteracted by a reduction in anisotropy due to rearrangement of gap junctions in response to ischemia [10, 32].

Limitations

The present modeling study has several limitations. First, we chose the LR ventricular cell model as a specific ventricular cell model and the SAN model to create a specific ectopic focus model. The quantitative results of our simulations are of course dependent on the membrane models chosen. However, the conclusion that anisotropy may reduce the electrotonic load on the ectopic focus and thus facilitate propagation from an ectopic focus into a sheet of myocardium should not depend on the membrane models chosen, and should hold for any region of focal activity. Secondly, we chose to modulate effective focus size by scaling coupling currents, thus neglecting the effects of the increase in number of cell contacts with an increase in focus size. This approach allows direct comparison with our previous experimental data [17, 39], but it also affects the quantitative results of our simulations (see Fig. 11). Additionally, the requirement of symmetry to decrease the number of elements required for the simulation limits the distribution of conductances. Although it would be interesting to include heterogeneous distribution of coupling conductances into the sheet model, the current implementation is sufficient to show how the presence of anisotropy can enable an ectopic focus to initiate a propagating wave into a sheet.

Finally, our simulations are based on the assumption that a single cell is, for these low values of coupling conductance, very nearly isopotential, as demonstrated by the theoretical work of Shaw and Rudy [31]. Therefore each “element” is an intact cell and the actual spatial dimensions and orientation of the particular cells used do not affect the results. We have deliberately chosen lower values of coupling conductance than those occurring in “normal” ventricular tissue in order to emphasize the effects which might be observed in cells that are partly uncoupled, such as might have occurred by prior ischemia. Also, these low values of coupling conductance, and the resulting discontinuous conduction, allowed us to use relatively small arrays to demonstrate these effects.

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References


