Cardiac repolarization alternans is a disturbance in the normal rhythm of the heart that has been identified as a precursor to potentially lethal arrhythmias including ventricular tachycardia and ventricular fibrillation. Alternans is seen clinically as a beat-to-beat alternation of action potential durations and intracellular calcium transient magnitudes. Recent experimental studies have reported “subcellular alternans,” in which distinct calcium transients, either in phase (electromechanically [EM]-concordant) or out of phase (EM-discordant), depending on the temperature and/or species. Although alternans has been mechanistically linked to the formation of potentially fatal reentrant arrhythmias, significant uncertainty remains regarding its mechanism and effective therapy aimed at its treatment or prevention has yet to be developed. Although much recent experimental evidence has identified instabilities in intracellular Ca cycling (ie, sarcoplasmic reticulum calcium release/reuptake) as the primary cause of alternans, uncertainty remains regarding the contribution of membrane voltage ($V_m$) instability (caused by insufficient time for recovery of sarcolemmal ion channels between excitations) to the total instability seen in alternating cells.

Alternans has been reported at multiple spatial scales, including spatially discordant alternans (SDA) in tissue and counterphase Ca alternans in adjacent cells. Most recently, several studies have reported “subcellular alternans,” in which Ca transients in distinct regions of an individual myocyte alternate out of phase. Importantly, the large gradi-
ents of calcium concentration formed between adjacent out-of-phase subcellular regions have been shown to predispose the cell to the formation of propagating waves of intracellular calcium release, which are thought to induce arrhythmogenic afterdepolarizations.9–13 Subcellular alternans has been reported several times in intact tissue,8,14 as well as in isolated cells,10,11,15 but the cause of this phenomenon remains unclear.

Two mechanisms have been the most frequently proposed to account for the formation of subcellular alternans. The first and most prevalent attributes these subcellular differences in Ca alternans either directly or indirectly to the presence of preexisting subcellular heterogeneity in calcium-cycling components along the length of the cell.9–11,14 In one recent example, Aistrup et al explained the subcellular alternans they observed in isolated rat hearts using a mathematical model containing significant heterogeneity in Ca-release properties in discrete regions along the length of the cell. In their model, a pacing protocol that reverses the phase of alternans (in both APD and Ca transients) can induce subcellular alternans by reversing the phase of Ca alternans in some regions earlier than others.14

A different hypothesis was proposed by Shiferaw and Karma, in which subcellular alternans results from a purely dynamical mechanism, in the absence of such preexisting subcellular heterogeneity.16 They proposed a well-known dynamical pattern-forming instability (a Turing instability) as a mechanism for subcellular alternans, wherein “a fluctuation that causes the amplitude of Ca alternans to increase in a small region of the cell generates APD alternans that in turn inhibits the growth of Ca alternans far from that localized fluctuation.”16 Their theoretical analysis, however, applied to only EM-discordant myocytes, and has yet to be validated experimentally.

Here, we extend this theory to predict a novel means of dynamically inducing subcellular alternans in EM-concordant myocytes and validate this hypothesis in a computational model, as well as in isolated guinea pig ventricular myocytes. Although physiological subcellular alternans may result from a combination of both dynamical and anatomic factors, here we present the first in vitro evidence that subcellular alternans can be dynamically induced without fixed heterogeneity. Furthermore, we hypothesize that this same dynamical mechanism may contribute to subcellular alternans seen during static pacing in intact tissue.

**Alternans Control**

A body of literature has developed around the idea of terminating alternans for antiarrhythmic purposes. The most studied approach uses nonstatic pacing in which the pacing cycle length (T) is adjusted following each AP by an amount proportional to the existing APD alternans magnitude (the difference in duration between the last 2 APs, \( A_n \) and \( A_{n-1} \)), according to the following equation:

\[
\Delta T = \frac{g}{2} (A_n - A_{n-1})
\]

where g is the control gain.17–19 Using this simple “alternans control” algorithm, the otherwise long diastolic interval following short APs is shortened and the normally short diastolic interval following long APs is lengthened. Because of a strong dependence of APD on the preceding diastolic interval (APD restitution), these continued cycle length perturbations force the APDs of an alternating cell toward an unstable period-1 rhythm. As described below, we hypothesize that alternans control dynamically induces subcellular alternans in EM-concordant cells, thereby providing a controlled means to investigate its formation in real cardiac myocytes.

**Methods**

An expanded Methods section is available in the Online Data Supplement at http://circres.ahajournals.org.

Briefly, computational modeling was conducted in model cells in which a single myocyte consists of a chain of 75 individual sarcomeres (each with its own calcium concentrations) coupled by calcium diffusion (as in Ref. 14, 16). The membrane voltage of all sarcomeres is explicitly synchronized, and simulations were conducted in a 400-cell cable of such model cells, paced from the left end. This model has the distinct advantage that it is easily tuned to exhibit either positive or negative Ca→V_m coupling as well as V_m-driven, Ca-driven, or Ca-driven (hereafter “mixed instability”) alternans (as described in Data Supplement). In all figures and movies, Ca alternans is quantified as \( \Delta c = (-1)^n (c_n - c_{m-1}) \), where \( c_n \) is the maximum [Ca^{2+}]_i of each sarcomere during beat n. APD alternans is quantified as \( \Delta \text{APD} = (-1)^n (A_n - A_{n-1}) \), where \( A_n \) is the APD of beat n.

Experimental studies were performed in isolated left ventricular myocytes from adult Hartley guinea pigs loaded with fluo-4/acetoxymethyl ester fluorescent calcium indicator. Cells were paced at cycle lengths sufficiently fast to induce steady-state APD alternans (range, 200 to 400 ms), following which alternans control pacing (varying the cycle length in real-time using Eq. 1) and calcium imaging were initiated. All experiments were performed at room temperature.

**Results**

**Dynamical Mechanism for Subcellular Alternans in EM-Concordant Myocytes**

We hypothesized that alternans control can induce subcellular alternans in EM-concordant myocytes by a mechanism anal-
Figure 1. In the presence of fluctuations in Ca alternans ($\Delta c$) between sarcomeres, alternans control pacing induces a marked subcellular alternans pattern in the EM-concordant model cell. The graphs are snapshots from Online Video 2 showing the formation of subcellular alternans in the model myocyte when alternans control is applied. Note that in this example, a large gradient was added to initial Ca concentrations (and therefore Ca alternans magnitude [$\Delta c$]) only for the sake of clarity (as seen in the first graph). The progression to subcellular alternans is described in the text. Note that the node (the point at which $\Delta c=0$, circle) is seen to migrate to the midpoint, such that the 2 regions of opposite phase Ca alternans offset one another and a period-1 whole-cell average Ca transient (and therefore a period-1 APD) results. This simulation was performed in the mixed-instability model, $T^*=300$ ms (where $T^*$ indicates target cycle length [during alternans control pacing]).

This is seen in Figure 1 (and Online Video 2), in which a model cell (described in Methods) is paced with alternans control. Note that in this simulation, for the sake of clarity, a large, smooth gradient was added to initial Ca concentrations (and thus Ca alternans magnitude); similar results are seen with the incorporation of only small random noise (as was done in all other simulations in this study). When such a cell is far away from its period-1 rhythm, alternans control will force all regions of the cell toward their unstable period-1 Ca transient rhythm (Figure 1, first and second graphs). As control continues, however, some regions will have reached their period-1 Ca rhythm (Ca alternans magnitude = 0) before the whole-cell average Ca transient is at a period-1 rhythm. Because the APD is still alternating (due to its dependence on the whole-cell average Ca transients), alternans control will continue to perturb the cycle length, reversing the phase of Ca alternans (changing the sign of Ca alternans magnitude) in those regions that had the lowest initial Ca alternans magnitude (Figure 1, third graph). For such regions of the cell, with Ca alternans now of opposite phase to that of the whole-cell average Ca alternans (and thus the APD alternans), the cycle length perturbations delivered by alternans control will now amplify these counterphase Ca alternans. The magnitude of Ca alternans in these regions will grow until the regions of opposite phase Ca alternans balance, such that the whole-cell average Ca alternans magnitude (and thus $\Delta APD$) will approach 0 (Figure 1, fourth graph). As this point is approached, alternans control will deliver progressively smaller perturbations to the cycle length (Eq. 1), approaching static pacing. Even with precisely static pacing, however, period-1 calcium dynamics are unstable (without such instability there would have been no alternans in the first place); Ca alternans will therefore grow in magnitude at all points along the cell, regardless of phase (Figure 1, graphs 4 and 5), forming a marked subcellular alternans.

This mechanism predicts subcellular alternans will occur in EM-concordant myocytes during alternans control but only in the presence of spatial variations in intracellular Ca concentrations (which can be satisfied by stochastic noise in initial conditions) and only if alternans is primarily driven by Ca-cycling instabilities.
Single-Cell Modeling Results

Static Pacing
When the model is tuned to exhibit EM concordance (positive Ca→Vm coupling), static pacing is unable to cause subcellular alternans at any pacing cycle length. The small degree of noise included in initial Ca concentrations quickly smoothes out through Ca diffusion between sarcomeres, which become approximately synchronized within 75 beats. To exhibit subcellular alternans during static pacing (by a Turing-type instability), the model must be tuned to EM discordance (negative Ca→Vm coupling) and Ca-driven alternans, as previously reported. Interestingly, with this parameter regime, the model never exhibits APD or whole-cell Ca alternans. The Ca transients of the 2 halves of the cell are always either at a period-1 rhythm or are exactly out of phase, making the whole-cell average Ca transient a period-1 rhythm over all pacing cycle lengths (data not shown), inconsistent with most single-cell electrophysiology studies, in which APD alternans is seen with rapid pacing.

Alternans Control Induces Subcellular Alternans in an EM-Concordant Model Cell
When the model is tuned to exhibit EM concordance, though subcellular alternans is never seen with static pacing, the model does undergo a more characteristic rate-dependent alternans bifurcation into a stable period-2 APD and whole-cell concordant Ca alternans rhythm. As seen in Figure 2 and Online Video 3, alternans control forces the period-2 APD and whole-cell Ca transients toward their unstable period-1 rhythm (Figure 2A, black traces). Interestingly, this transition to a period-1 rhythm occurs simultaneously with a symmetry-breaking transition from whole-cell concordant Ca alternans to subcellular alternans, in which the 2 halves of the cell are out of phase (causing a period-1 whole-cell average Ca transient rhythm; Figure 2A and 2B). Along the length of the model cell, Ca alternans is seen to be largest at 2 terminal antinodes surrounding a single, central node with no Ca alternans. Because Vm (and thus APD) is explicitly dictated by the ionic currents averaged over the length of the cell (in this model), the effects of calcium-sensitive membrane currents in each of the 2 halves average out to produce a period-1 APD. When control is released by terminating the infinitesimally small perturbations being applied to the static pacing interval, this transition is traversed in the opposite direction, causing a smooth return to the original whole-cell concordant Ca alternans rhythm (not shown).

Subcellular alternans occurs in the EM-concordant model at all pacing cycle lengths at which alternans is seen, whenever alternans control is successful. As predicted, this effect is robust over a large range of alternans control gains above a low threshold (g>0.7 for most pacing cycle lengths) and is not seen without alternans control or in the absence of noise in initial-condition Ca concentrations. Subcellular alternans was induced in this model whenever Ca-cycling dynamics were sufficiently unstable (both the mixed instability and Ca-driven models) but, as predicted, never occurred when alternans was attributable to Vm instability. Subcellular alternans became unstable and reverted to whole-cell Ca alternans with cessation of alternans control for all cycle lengths tested.

Isolated Cell Studies
To test our hypotheses in vitro, we applied alternans control to isolated guinea pig ventricular myocytes while imaging intracellular calcium. Subcellular alternans has previously been reported in statically paced intact guinea pig heart, even though guinea pig myocytes are well characterized as EM-concordant both as isolated cells and in tissue. Isolated left ventricular cells (from all myocardial regions) were paced at cycle lengths sufficiently fast to induce steady-state alternans (range 200 to 400 ms), after which alternans control was initiated. A total of 51 trials were conducted in 16 different cells (an average of 4 trials were performed in each cell). All recordings showed EM concordance throughout. Alternans control successfully suppressed alternans (in both APD and whole-cell average Ca transient magnitude) in 45 of the 49 trials in which it was applied, and 42 of those 45 successful trials showed subcellular Ca alternans (as in Figure 3A through 3C; Online Videos 5 and 6). Subcellular alternans was never seen without control, and none of the trials in which control failed showed subcellular alternans. This effect was seen at all cycle lengths tested (at which alternans occurred), independent of alternans control gain, and was robust to changes in patch-clamp configuration (perforated versus ruptured patch) and dye affinity (Fluo-4 versus Fluo-4 FF).

In 13 of 17 trials in which alternans control was released (static pacing resumed) during imaging, subcellular alternans...
transitioned back to whole-cell concordant alternans (as in Figure 4; Online Videos 5 and 6). In the 4 trials in which the subcellular alternans persisted after control was released, the APDs diverged to stable alternans after imaging had ended, implying that a longer imaging period would have captured a return to whole-cell concordant Ca alternans as well.

Six of the 16 cells tested showed more than 1 node (and more than 2 antinodes) of Ca alternans during alternans control in one or more trials, in patterns similar to those seen in the mathematical model (as in Figure 3A through 3C, Online Figure I, and Online Video 6). The majority of cells showed a similar pattern and number of nodes in each trial, but 5 of the 16 cells exhibited different numbers of nodes in different trials at the same cycle length (as seen in Figure 3B and 3D).

One-Dimensional Fiber Simulations
We further hypothesized that previous reports of subcellular alternans in statically paced, intact tissue may also be accounted for by our proposed dynamical mechanism. In particular, the cycle length perturbations induced by the alternans control protocol are directly analogous to the local activation times of cells away from the pacing site in whole heart given sufficiently steep conduction velocity (CV) restitution. We therefore predicted that given sufficiently steep CV restitution, subcellular alternans can be induced by the same dynamical mechanism in EM concordant tissue during static pacing.

To test this hypothesis, we implemented a cable of 400 model cells, each one itself a cable of 75 sarcomeres. The cells were tuned to be EM-concordant, with calcium-driven alternans (as described in the Online Methods) and contained no fixed heterogeneity in Ca-release properties. The left-most 5 cells of the cable were statically paced for 300 beats at a slow cycle length (T/H = 400 ms, at which there is no alternans at any spatial scale) and then for an additional 200 beats at a faster cycle length (either 330 or 320 ms). When the pacing cycle length was decreased to 330 ms, CV-restitution was not steep enough to induce tissue-level SDA. In this case, alternans is seen in the Ca transients of every sarcomere, as well as the APD of every cell, but subcellular alternans is not observed (Figure 5A). In contrast, when the pacing cycle length is decreased to 320 ms, tissue-level SDA arises in both the APD and the whole-cell average calcium transient (Figure 5B, top and middle). Additionally, over the course of the 200 beats at this faster cycle length, a prominent subcellular alternans transiently develops in the cells near the node of tissue-level SDA (Figure 5B, bottom). Figure 5C shows a pseudo–line scan image of a single cell near the node of SDA (the cell with the most significant subcellular alternans). The patterns of subcellular alternans formed were similar to those seen in previous experiments in intact rat heart (Ref. 8, 14), with nodes (dashed lines) migrating along the length of the cell as the simulation progressed until the entire cell has reversed phase of Ca alternans. Note that in this model the membrane voltage is explicitly synchronized for all sarco-
Subcellular Ca alternans has been reported multiple times in the intact heart as well as in isolated cardiac myocytes, but the mechanism for this striking phenomenon remains unclear. Although previous studies have invoked subcellular heterogeneities in Ca-cycling characteristics to explain the occurrence of subcellular alternans, we believe that the rich cardiac myocyte dynamics (which are known to contribute to alternans) at the cellular and tissue level cannot be discounted. An elegant theoretical study proposed a dynamical mechanism for this phenomenon in real cells, allowing in vitro confirmation for the first time.

**Alternans Control Induces Subcellular Alternans in EM-Concordant Myocytes**

When alternans control pacing was applied to an EM-concordant mathematical model, subcellular alternans was induced in a robust manner. Notably, subcellular alternans was induced in this manner without the need for fixed heterogeneity, thus predicting that a purely dynamical mechanism can induce subcellular alternans. Our in vitro studies similarly demonstrated that alternans control robustly induces subcellular alternans in (EM-concordant) isolated guinea pig ventricular myocytes. We believe that although there is undoubtedly fixed subcellular heterogeneity in real myocytes, our results support a primarily dynamical induction of subcellular alternans in our experiments and do not suggest the presence of heterogeneity of the type previously proposed to account for subcellular alternans formation.

Several experimental observations support these conclusions. (1) A change from static pacing to alternans control was sufficient to robustly induce a smooth transition to subcellular alternans. This agrees with our theoretical and mathematical modeling work, in which this pacing algorithm can induce subcellular alternans in cells by a purely dynamical mechanism. (2) Release of alternans control (resumption of static pacing) during subcellular alternans caused this rhythm to lose stability and return to a period-2 APD and whole-cell concordant Ca alternans rhythm. It is important to note that successful alternans control (as was achieved before release in Figure 4) forces a cell toward its period-1 rhythm with progressively smaller perturbations to the cycle length, culminating in very small changes to the pacing interval around the target cycle length (nearly static pacing). As such, the transition from alternans control to static pacing is a smooth transition. (3) The nodes of subcellular alternans were distributed approximately symmetrically about the midline, as has been reported in several other studies. In contrast, our mathematical modeling results indicate that in the presence of the large, fixed gradients of calcium-cycling components previously proposed to cause subcellular alternans (Online Figure I, B), the pattern of subcellular alternans would be strongly dependent on the distribution of this underlying heterogeneity. We are unaware of any previous work that suggests the presence of the symmetrical gradients of subcellular heterogeneity that would be required for the symmetrical pattern of subcellular alternans seen in our experimental studies (and those of others) to be caused by underlying heterogeneity alone. (4) The patterns of subcellular alternans were seen to vary in different trials of the same cell at the same cycle length (5/16 cells; as in Figure 3B and 3D). This is consistent with the dynamical model of Online Figure I, A, but inconsistent with the predictions of the model when large, fixed gradients of calcium-cycling components
were included (Online Figure I, B). Notably, with only a small degree of stochastic fixed heterogeneity in calcium-cycling components, the model was seen to exhibit different patterns in different trials with nodes symmetrically distributed about the midline, consistent with our experimental results. Future studies using confocal imaging will be needed to precisely determine 3D cellular morphology and the location of nodal lines during subcellular alternans.

Notably, the occurrence of subcellular alternans during alternans control is only predicted when Ca-cycling instability is greater than instability attributable to membrane voltage dynamics (if this were not the case, all Ca transients would alternate secondary to APD alternans and would thus be synchronized). The occurrence of subcellular alternans in our guinea pig myocytes therefore supports the findings of several previous studies indicating that alternans is primarily attributable to Ca-cycling dynamics.1 Testing for subcellular alternans during alternans control pacing may represent a useful tool for determining the primary source of instability in other cell types/species, as well as at different pacing cycle lengths.

**Subcellular Alternans May Be Dynamically Induced in Intact Tissue During Static Pacing**

The dynamical induction of subcellular alternans in EM-concordant myocytes may also occur physiologically, and may contribute to rate-dependent subcellular alternans seen in statically paced intact tissue.8,14 In tissue, at sufficiently fast pacing rates, the effect of steep CV restitution produces an effect similar to alternans control in the local activation times of cells distant from the pacing site. Specifically, if APD alternans occurs at the pacing site during static pacing, CV-restitution dictates that “long” APs will propagate away faster than “short” APs. Cells away from the pacing site will thus experience effectively nonstatic pacing, and the excitations following their “long” APs arriving late (because they are carried by a more slowly traveling, “short” AP) and those following their “short” APs arriving early (because they are carried by a faster, “long” AP). The further away from the pacing site, the greater the effect of propagation velocity on cycle length, and the stronger is this alternans control-like effect. Indeed, these cycle length alternations can be strong...
enough to entirely eliminate APD alternans (forming a period-1 node) and, even further away, reverse the phase of APD alternans relative to that at the pacing site (forming tissue-level SDA). Notably, such cycle length alternations were observed concomitant with the subcellular alternans recently observed during static pacing of rat heart.¹⁴

Indeed, we found that when tissue-level SDA is induced by static pacing in a simulated 1D fiber of EM-concordant myocytes, subcellular alternans is dynamically induced in those cells adjacent to the tissue-level node (where cycle length alternations are the most pronounced). Although there are clearly many dynamical and anatomic differences between isolated cells and the intact heart, we believe that the proposed dynamical mechanism may contribute to the formation of physiological subcellular alternans.

It is important to note that although subcellular alternans can be induced dynamically in isolated cells, the physiological formation of subcellular alternans is likely governed by a rich interplay of dynamical as well as anatomic factors. Still, we believe that our identification of a robust, dynamical mechanism by which subcellular alternans can be induced in isolated cardiac myocytes not only provides novel insights into the mechanism of this phenomenon in the real heart but also will facilitate future studies into the formation and consequences of subcellular alternans and biological pattern formation.

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MATERIALS AND METHODS

Cell model

A spatially extended version of the Shiferaw-Sato-Karma ventricular myocyte model was implemented in C++. This model, described in Ref. 1, consists of a chain of 75 sarcomeres, each of which has its own Ca-cycling dynamics (and concentrations) and is coupled to its neighbors via compartment-specific Ca-diffusion. $V_m$ in this model is synchronized over the length of the cell, influenced by the average states of all sarcomeres. Unless stated otherwise, all simulations were initialized to steady-state conditions with Gaussian noise added to all calcium concentrations ($\sigma$=1% of steady state values). To model fixed heterogeneity, changes were made to each sarcomere’s SR release load dependence ($u$), either by the addition of Gaussian noise ($\sigma$= 0.1sec$^{-1}$) or by varying this parameter sinusoidally along the length of the cell (as in Fig. 3B). Note that although physiological heterogeneity and Ca-transient fluctuations are likely present at scales smaller than the sarcomere, for reasons of computational tractability a more finely discretized model was not used. Based on the generality of the mechanism proposed, we do not predict this would lead to qualitatively different results.

$Ca \rightarrow V_m$ coupling (and thus EM concordance) in the model was controlled by adjusting the degree of Ca-induced inactivation of L-type calcium channels ($\gamma$), where $\gamma$=0.7 causes positive $Ca \rightarrow V_m$ coupling (and EM-concordant alternans), and $\gamma$=1.5 causes negative $Ca \rightarrow V_m$ coupling (and EM-discordant alternans), as in Ref. 1). The degree of voltage and calcium-driven instability were controlled by adjusting the time constant of recovery from inactivation of the L-type calcium channels ($\tau_i$) and the load dependence of sarcoplasmic reticulum calcium release ($u$), respectively. In the single cell
studies, the “$V_m$-driven alternans” model corresponded to $\tau_f=60\text{ms}$ and $u=5\text{sec}^{-1}$, the “Ca-driven alternans” to $\tau_f=20\text{ms}$ and $u=22\text{ sec}^{-1}$, and the “mixed-instability” model $\tau_f=40\text{ms}$ and $u=11\text{sec}^{-1}$ (as in Ref. 1, 2). Model code is available upon request.

A 1-dimensional cable of 400 of these model cells—each one itself a cable of 75 sarcomeres (as above)— was simulated using the standard cable equation

$$\frac{\partial V}{\partial t} = -\frac{I_{ion}}{C_m} + D \frac{\partial^2 V}{\partial x^2},$$

where $I_{ion}$ is the total membrane current, $C_m$ is the membrane capacitance, and $D = 1 \times 10^{-3} \text{ cm}^2 / \text{ms}$ is the effective diffusion constant for membrane voltage ($V$). This cable equation was integrated using an operator splitting method$^3$ with a space step of $\Delta x = 0.15\text{cm}$ and a time step of 0.05ms. Because the CV restitution curve of this model is very flat, we have increased the time constant of the slow inactivation variable of the sodium current ($\tau_j$) by a factor of five, as has been done previously (e.g. Refs. 4, 5). This modification steepens the CV restitution curve and promotes the occurrence of tissue-level spatially discordant alternans. Parameters were tuned such that model cells were EM-concordant ($\gamma=0.7$) and alternans was calcium-driven ($\tau_f=50\text{ms}$ and $u=13\text{ sec}^{-1}$, as in Ref. 1). Gaussian noise ($\sigma=0.5\%$ of steady-state calcium concentrations) was added to every sarcomere’s calcium concentrations before each stimulus. To ensure propagation, the leftmost five cells were stimulated.

**Calcium Imaging**

Cells were loaded with the cell permeant (acetoxymethyl) form of the calcium sensitive dye Fluo-4 or its low-affinity counterpart, Fluo-4 FF (to minimize calcium buffering by the dye itself). Epifluorescence was recorded at 1000Hz, digitized at 14 bits, and stored for offline analysis using one camera of an 80X80 pixel CardioCCD-sm Dual-Camera
Imaging System (RedShirtImaging, LLC, Decatur, GA) and RedShirtImaging Cardioplex software on a Dell workstation running Microsoft Windows. All calcium traces shown have been low-pass filtered at 20Hz. Data acquisition and lamp shutter opening were controlled electronically by RTXI on a Dell workstation running Linux. The same RTXI process that acquired/processed the patch-clamp data controlled imaging. Data analysis for voltage and calcium data was performed with custom C++ and Matlab programs.

**Cellular Electrophysiology**

Myocytes were isolated from the left ventricle of adult (400-600 gram) Hartley guinea pigs using the Langendorff retrograde perfusion method, as described previously. Whole-cell voltage recordings were obtained using the Amphotericin B perforated patch technique, as described previously, with an A-M Systems Model 2400 patch clamp amplifier (Carlsborg, WA). Each myocyte was paced in current-clamp mode by 2ms square pulses of twice diastolic threshold current at a cycle length of 1000ms until stable APDs (defined as time to 90% repolarization) were seen. The pacing cycle length was then decreased to a rate sufficiently fast to induce steady-state alternans (from 200 to 400ms in different cells), at which point alternans control pacing was initiated (by varying the pacing cycle length using Eq. 1) and calcium imaging begun. Note that we non-specifically selected left ventricular cells and therefore likely studied myocytes from different regions. As such, alternans thresholds varied from cell to cell, as is commonly reported (e.g. Ref. 8). Shutter openings were limited to less than 30 seconds to minimize the effects of phototoxicity/photobleaching. The gain ($g$) of alternans control was
adjusted (from 1.5 to 2.5) for each cell as needed to force the cell to a period-1 APD rhythm. An average of 4 trials were conducted in each myocyte.

Experiments were performed at room temperature, with constant superfusion of Tyrode’s solution at 1-2 ml per minute. Stimulus-timing control and data acquisition used custom plugins for the Real Time Experimental Interface system (RTXI; www.rtxi.org) running on a real-time Linux platform at 5kHz.

SUPPLEMENTAL RESULTS

Subcellular alternans is induced in the presence or absence of fixed subcellular heterogeneity, but with qualitatively different characteristics

Previous studies have proposed a mechanism for subcellular alternans in which fixed heterogeneities in calcium-cycling properties account for the differences seen along the length of the cell in subcellular alternans.9-11, 14 Although we found that alternans control causes subcellular alternans in the absence of fixed subcellular heterogeneity (Fig. 2), we predicted that even in the presence of such heterogeneity the proposed mechanism would lead to subcellular alternans, but with qualitatively different characteristics. To this end, we employed a Monte Carlo approach analyzing the characteristics of subcellular alternans in the presence and absence of fixed subcellular heterogeneities in calcium-cycling components (Fig. S1).

We ran 200 simulations of the cell model, in which Gaussian noise was added to the initial calcium concentrations of each sarcomere (as described in Materials and Methods). In 100 of the simulations, fixed heterogeneity in calcium-cycling components was incorporated by varying the load dependence of each sarcomere’s SR Ca-release (u)
sinusoidally along the length of the cell (with an amplitude and wavelength similar to the discontinuous heterogeneity used in Ref. 14).

Without fixed heterogeneity, different patterns of subcellular alternans (different numbers of nodes) were seen in different trials (differing only in their initial conditions): 80/100 trials resulted in 1 node, 20/100 resulted in 2 nodes (Fig. S1A; Supplemental Video 4). Even greater numbers of nodes were seen when larger noise was added to initial calcium concentrations (not shown). Importantly, although the initial point(s) of subcellular alternans node formation (phase reversal) were random, in these simulations the node(s) were always seen to migrate to a distribution in which the whole-cell average Ca-alternans magnitude is nearly 0, with symmetry about the midline of the cell (as seen in Fig. S1A), consistent with the proposed mechanism.

In the presence of gradients of fixed heterogeneity, alternans control still led to the formation of subcellular alternans, but with qualitatively different characteristics. Every trial of a given pattern of fixed heterogeneity always resulted in the same number of nodes, regardless of wavelength or position of these variations (Fig. S1B). For the patterns shown in Fig. S1B, there were always 2 nodes (located at the inflection points of the sinusoidal variation in underlying Ca-cycling dynamics), regardless of initial Ca-concentration conditions. Note that qualitatively similar results were obtained in the presence of large, discontinuous (rather than smooth) regions of fixed subcellular heterogeneity (as were used in Ref. 14; simulations not shown). Also note that when fixed heterogeneity was instead incorporated as small, stochastic differences in each sarcomere’s SR Ca-release load dependence ($\nu$), the results were similar to those obtained in the absence of fixed heterogeneity.
SUPPLEMENTAL FIGURE CAPTIONS

Supplemental Figure 1. Alternans control induces subcellular alternans in the EM-concordant cell model both in the absence (A) and presence (B) of fixed heterogeneity in calcium-cycling components (u). The fixed calcium-cycling components (top), initial intracellular calcium concentrations (middle), and steady-state Ca-alternans magnitude (bottom, Δc, defined as in Fig. 1) are shown for each sarcomere, as well as the whole-cell average steady-state Ca-alternans magnitude (c_{max}, bottom, right). Δc_{max} is calculated as the difference in peak concentration of the whole-cell average Ca-transient between two successive beats at steady-state. A, Without fixed heterogeneity, different patterns of subcellular alternans form in different trials (as in Supplemental Videos 3 and 4). The results of two trials of the same model cell are shown, in which small differences in initial Ca-concentrations led to different patterns of steady-state subcellular alternans. Note that the nodes migrate to a distribution such that alternans control is able to very successfully eliminate whole cell Ca-alternans. B, When sinusoidally varying heterogeneity in calcium-cycling components (u) is included, alternans control still induces subcellular alternans, but the number and position of nodes are dictated by the distribution of this underlying heterogeneity. Shown are the results from three different model cells in which the position of the underlying heterogeneity is shifted—regardless of variations in initial conditions, the number and position of nodes is constant for a given underlying structure and alternans control is unable to entirely eliminate whole-cell Ca-alternans. All simulations were conducted in the EM-concordant, mixed-instability model at T=300ms.
SUPPLEMENTAL VIDEO CAPTIONS

Supplemental Video 1. Starting from steady-state alternans, alternans control is applied to a perfectly homogeneous cell model (i.e. no fixed heterogeneity and no noise in Ca-concentrations) with 75 sarcomeres. Because there are no differences between adjacent sarcomeres, the cycle length perturbations of alternans control are appropriately timed to force every sarcomere’s Ca-transients to a period-1 rhythm ($\Delta c = 0$). At the same time, APD-alternans (dependent on the whole-cell average Ca-alternans) is forced towards a period-1 rhythm ($\Delta APD = 0$). This simulation was performed in the mixed-instability model, $T^*=300$ms.

Supplemental Video 2. In the presence of fluctuations in Ca-alternans between sarcomeres, alternans control causes a marked subcellular alternans pattern (as described in text). In this simulation, a large gradient in Ca-concentrations (and thus Ca-alternans magnitudes ($\Delta c$)) was added to the steady-state (alternans) conditions, after which alternans control was applied. Note that the node (the point at which $\Delta c = 0$, circle) migrates to the midpoint, such that the two regions of opposite phase Ca-alternans offset one another and a period-1 whole cell average Ca-transient (and therefore a period-1 APD) results. Note that in this example, such a large, smooth gradient in initial Ca-alternans magnitude was added for the sake of demonstration clarity only. This simulation was performed in the mixed-instability model, $T^*=300$ms.

Supplemental Video 3. Beginning from steady state alternans, the same structurally homogeneous model cell as in Videos 1 and 2 was paced with alternans control after Gaussian noise was added to the Ca-concentrations of each sarcomere. In the presence of
such initial fluctuations in Ca-concentrations (and therefore Ca-alternans magnitudes ($\Delta c$)), alternans control induces subcellular alternans. Although the point of initial phase reversal is random (dependent on the initial noise), the node (where $\Delta c = 0$, circle) is always seen to migrate such that the whole-cell average Ca-alternans magnitude=0. In this example, a two-region (one node) subcellular alternans is formed. When different noise is added, the point of initial phase reversal is different, and different patterns of subcellular alternans can result (as in Supplemental Video 4). This simulation was performed in the mixed-instability model, $T^*=300$ms.

**Supplemental Video 4.** In the same cell model as in Supplemental Video 3, when different noise is added to initial Ca-concentrations, a different pattern of subcellular alternans results when alternans control is applied. Note that the nodes of subcellular alternans (where $\Delta c = 0$, circle) migrate such that they are symmetric about the midline at steady-state and a period-1 whole-cell average Ca-transient and APD rhythm results. This simulation was performed in the mixed-instability model, $T^*=300$ms.

**Supplemental Video 5.** An isolated guinea pig ventricular myocyte was statically paced to induce steady-state APD (not shown) and Ca-alternans. With static pacing, the Ca-alternans is seen to be approximately synchronized throughout the cell. Upon initiation of alternans control (at approximately time=6s), however, the Ca-alternans began to transition to a 2-region subcellular alternans rhythm in which the two halves of the cell (red and blue traces) alternated out-of-phase, causing an approximately period-1 APD and whole-cell average Ca-transient rhythm (black trace). After this period-1 rhythm was achieved, static pacing was resumed (at approximately time=57s), and the subcellular alternans began to transition back to a whole-cell concordant Ca-alternans and period-2
APD rhythm. This experiment was conducted at a cycle length of $T=340\text{ms}$. The traces shown are the average fluorescence of the whole cell (black) and each of the two regions diagrammed (red and blue).

**Supplemental Video 6.** Alternans control was used to induce subcellular alternans in an isolated guinea pig ventricular myocyte (prior to the video), at which point static pacing was resumed (at the beginning of this movie). The period-1 whole-cell average Ca-transient (black trace) returns to a period-2 rhythm as the three-region subcellular alternans pattern (red, blue, and green traces) returns to a whole-cell concordant Ca-alternans. Note that there is an abrupt increase in the Ca-fluorescence in a portion of region 2 on beat 8 relative to beat 6. Without confocal imaging, we cannot exclude the possibility that a spontaneous SR Ca-wave reset the phase of Ca-alternans in this region. By the proposed mechanism, however, such Ca-waves are not predicted to interfere with the formation of subcellular alternans during alternans control pacing and, indeed, may hasten its onset. This experiment was conducted at a cycle length of $T=300\text{ms}$. The traces shown are the average fluorescence of the whole cell (black) and each of the two regions diagrammed (red and blue).

**SUPPLEMENT REFERENCES**


