Increased cardiac fibrosis is shown to be associated with cardiac conduction block and reentry in isolated-perfused animal and diseased human cardiac tissues [1-3] as well as in isolated Langendorff-perfused explanted human hearts with dilated cardiac myopathy [4, 5]. While alterations of cardiac conduction [6, 7] and the resulting reentrant wavefront of excitation[8] are uniformly accepted arrhythmic consequences of increased cardiac fibrosis, recent experimental findings in isolated whole heart studies, indicate that fibrosis may also importantly modulate the formation of cardiac afterpotentials notably early afterdepolarizations (EADs) that lead to triggered activity causing atrial fibrillation (AF) [9] and ventricular fibrillation (VF) [10-12]. These findings extend previous cardiac monolayer studies that showed myofibroblast coupling to cardiomyocytes through gap junction formation imparts enhanced automaticity to cardiomyocytes when coupled to a finite (critical) number of myofibroblasts [13]. Taken together these findings indicate that increased cardiac fibrosis promotes arrhythmias not only by the mechanism of reentry but also by the mechanism of triggered activity and enhanced automaticity potentially making cardiac fibrosis a highly effective antiarrhythmic target.

In this brief review we delineate step-by-step how the interaction of aged fibrotic ventricles with oxidative stress leads to the emergence of early afterdepolarizations (EADs), triggered ac-
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tivity and VF. We emphasize the importance of fibrosis as non-fibrotic hearts when stressed similarly or at even higher stress levels do no manifest any arrhythmic events. Specifically, we describe the dynamic scenario starting from cellular EADs that evolves to rapid focal ventricular tachycardia (VT) caused by triggered activity which then degenerates to VF. In the second part we point out briefly on recent experimental [14, 15] and the potential of drug-induced prevention and/or reduction of ventricular fibrosis as an antiarrhythmic strategy in humans [16-20].

The pathology of fibrosis

Fibrosis develops when the body's natural wound-healing process goes awry. Under normal (adaptive) conditions of wound healing, specialized cells known as fibroblasts become activated by transforming to myofibroblast. The myofibroblasts then undergo proliferation causing increased synthesis of collagen protein in the extracellular matrix composed predominantly of type I collagen and to a lesser extent type III collagen (normal wound healing process). What is initially an adaptive process, perhaps meant to enhance tensile strength, can progress to maladaptive (pathologic) conditions when the “healing” process persists with the development of diffuse and heterogeneous myocardial fibrosis characterized by a matrix made of increased collagen deposits and myofibroblast numbers [18, 21-23]. While resident cardiac fibroblasts may be activated and transformed to myofibroblasts there is also the potential of fibroblasts originating from endothelial cells, suggesting an endothelial-mesenchymal transition (EndMT). For example, it has been shown that the transforming growth factor-beta 1 (TGF-β1) induces endothelial cells to undergo EndMT, whereas bone morphogenic protein 7 (BMP-7) preserves the endothelial phenotype. The demonstration of the systemic administration of recombinant human BMP-7 (rhBMP-7) to significantly inhibit EndMT and the progression of cardiac fibrosis in mouse models of pressure overload may provide novel therapeutic approaches to control the progression of cardiac fibrosis [24].

Whole heart animal model of fibrosis and oxidative VF

In 1956, Harmann [25] made his groundbreaking observations on the role of reactive oxygen species (ROS) in the aging process. Thereafter, the concept of ROS became widely accepted in theories of aging [26]. While atrial and ventricular fibrosis may increase with aging, however, fibrosis per se does not promote cardiac arrhythmias [27-30]. Instead, fibrosis as we will see below, provides a substrate that when coupled to a mild form of stress that is of no arrhythmic consequence in non-fibrotic hearts, causes cardiac arrhythmias in the fibrotic heart. In this respect fibrosis becomes a significant risk factor for increased vulnerability to cardiac arrhythmias. We provide experimental evidence in isolated-perfused whole heart preparations that fibrosis acts as the “first hit,” a necessary and a required substrate, and that increased oxidative stress acts as a “second hit,” both hits being required to promote spontaneous VF. We call this the “double hit hypothesis” of spontaneous VF. We now describe the dynamic evolution as to how oxidative stress (hydrogen peroxide, H2O2) in aging-related fibrotic ventricles promotes spontaneous VF.

Oxidative stress

Hydrogen peroxide (H2O2) is shown to readily promote EADs and triggered activity in isolated rat and rabbit ventricular myocytes by increasing the late sodium current (INa,L) [31-33] However, this same stress fails to cause EADs in well-coupled, non-fibrotic cardiac tissue due to source-to-sink mismatches arising from cell-to-cell coupling. That is, a small current which is sufficient to reverse repolarization and cause an EAD in an isolated cardiac myocyte will be diluted into adjacent repolarizing myocytes (unless they are also simultaneously primed for an EAD), thereby suppressing the EAD. We investigated this issue by examining the effects of H2O2 on arrhythmias in Langendorff-perfused rat and rabbit hearts. Consistent with the predicted suppressive effects of well-coupled tissue on EADs, we found that oxidative stress with H2O2 failed to induce any ventricular arrhythmias in young adult rat or rabbit hearts with no fibrosis. In sharp contrast however, fibrotic aged rat and fibrotic middle-aged rabbit hearts, when exposed to similar levels of oxidative stress with 0.1 mM H2O2 readily induced VT/VF in these two species [10, 11]. Importantly it needs to be emphasized that in adult non-fibrotic hearts raising H2O2 concentration up to 2 mM still failed to induce arrhythmias in adult rat and adult (3-5 months old) rabbit hearts indicating that it is not the greater susceptibility of
aged hearts to oxidative stress relative to adult hearts that was responsible for the differential arrhythmic response in the two age groups of both species. These findings are consistent with the hypothesis that EADs are suppressed by cell-to-cell coupling in tissues, since \( \text{H}_2\text{O}_2 \) concentrations which consistently induce EADs and triggered activity in isolated rat and rabbit ventricular myocytes uniformly failed to induce EAD-related arrhythmias in non-fibrotic adult rat and rabbit intact hearts studied [10, 11].

**Fibrosis**

Histological analysis using Masson’s trichrome collagen showed marked increase in fibrosis in the aged (24–26 months) rat ventricles compared to adult ventricles averaging 45±26% of the ventricles. The distribution of fibrosis however, was highly heterogeneous, varying between 10% in the RV to 90% in the left ventricular (LV) endocardium and at intermediate level at the base of the LV epicardium. Middle-aged (3-5 years old) rabbit hearts however, had less extensive fibrosis averaging from 15% in the RV and mid LV wall to 25-35% in the anterior and posterior LV near the base of the heart. In contrast, fibrosis was minimal in adult rat hearts, averaging 4±2.5% of the ventricles, and 3±1% in adult rabbit ventricles (\( P<0.001 \) compared to aged rat and middle-aged rabbit ventricles)[10].

**From EADs to VF**

The mechanism of oxidative stress-mediated spontaneous VF (i.e., VF not induced by electrical stimulation) was studied systematically using optical activation map and single cell recordings with glass microelectrode in aged rats. For this purpose we used simultaneous voltage- and calcium (V-Ca) sensitive dyes to map activation pattern and define the underlying intracellular calcium dynamics (Ca\(^{2+}\)). In response to 0.1 mM \( \text{H}_2\text{O}_2 \), 66 of 70 of the aged rat hearts (94%) developed spontaneous VF after a mean perfusion time of 17.6±7.1 min (Figure 1). LV epicardial activation map showed that the VF was preceded by a transient period of focal VT (mean CL of 70±18 ms) which arose suddenly from regular sinus rhythm with a mean CL of 380±162 ms. Within 2 sec, the VT degenerated to sustained VF (CL of 55±16 ms) (Figure 1), requiring electrical shock for termination. However, spontaneous VT/VF reoccurred repeatedly after 2.3 min of cardioversion. The focal VT preferentially originated from the base of the LV anterior epicardium where the degree of fibrosis was intermediate (30-40%). This unique feature of spontaneous VF onset allowed us to capture the onset of the focal unstable VT during optical and glass microelectrode action potential recordings. Similar findings were observed in the middle-aged rabbit hearts when LV epicardial fibrosis averaged 35±16% of the LV. As in the case of aged rats, the VF in the middle aged rabbits was also preceded by a transient period of VT (mean CL of 130±20 ms) which arose suddenly from regular sinus rhythm with a mean CL of 460±80 ms.

**Mechanism of focal VT**

To gain insight into the cellular mechanisms of focal VT, after the cardioversion of the VF with an electrical shock, we continuously recorded with a roving glass microelectrode single cell action potentials from the epicardial surface of the LV base, where the focal VT frequently originated from to capture the onset of focal VT. With this approach, we show that the focal VT was initiated by an EAD-mediated triggered activity that arose from a mean take-off potential of -51±16 mV (Figure 2). The mean CL of the TA was 66±10 ms and was not significantly different from the mean CL of the VT (70±18 ms). The EAD preceded the QRS complex of a simultaneously recorded ECG by a mean of 8±4 ms and occurred during an isoelectric interval on the ECG indicating absence of electrical activity elsewhere in the heart during the EAD formation (Figure 2). The EADs arose when Ca\(^{2+}\) remained high relative to the diastolic resting level reflecting a slowed rate of decline of Ca\(^{2+}\). Maintained elevation of Ca\(^{2+}\) could activate the sodium-calcium exchanger (NCX) current providing a net inward depolarizing current [12] which along with the increased \( I_{\text{Na}-\text{L}} \) caused by \( \text{H}_2\text{O}_2 \) reduces the repolarization reserve and facilitates the emergence of reactivated L-type calcium (\( I_{\text{Ca-L}} \)) current causing EADs and triggered activity. These findings indicate that oxidative stress promotes EADs in areas of the LV where the level of fibrosis is intermediate by reducing ventricular repolarization reserve leading to EADs, triggered activity and focal VT.

**Why fibrosis is important? The source-to sink mismatch**

Our findings highlight the importance of cell-to-cell coupling on the ability of EADs to form and cause arrhythmias in tissue. While isolated ven-
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Figure 1: Spontaneous initiation of VT/VF in an aged rat heart exposed to 0.1 mM H$_2$O$_2$. Panel A is an ECG showing the last 5 sinus beats before the sudden onset of VT leading to VF. Panel B are voltage snap shots of the last beat of the VT (beats #1) and of the first two beats of the VF (beats #2 & #3). In each snap shot activation time in ms is shown at the bottom right with time zero (arbitrary) coinciding with the onset of beat #1. The red color in the snap shots represents depolarization and the blue repolarization as shown in panel E. The yellow arrows in the snap shots represent the direction of the wavefront propagation with double horizontal lines denoting the site of conduction block. The VT originates from a focal site at the LV base and propagates as single wavefront towards the apex and undergoes functional conduction block at site 3. The two lateral edges of the front however, continue to propagate laterally (snap shot 98 ms) forming figure-8 reentry (snap shot 108). During the second reentrant wavefront another wavefront emerges from the apical site of the LV (snap shot 122 ms) disrupting the activation pattern and signaling the onset of VF. Panel D, shows 3 optical action potentials (labeled 1, 2 and 3) recorded from sites identified on the heart silhouette (panel C). The two downward pointing blue arrows indicate the direction of propagation from site 1 to site 3 with the red downward pointing arrow showing block at site 3 followed by retrograde activation (upward pointing arrow). Notice the emergence of spatially discordant action potential duration (APD) alternans preceding conduction block at site 3 when the front with short APD (S) at site 1 encroaches a site (site 3) with long APD (L). S indicates short and L long APD. (From reference number 10, Morita et al)

Tricuspid myocytes readily develop EADs and triggered activity in response to oxidative stress, normal non-fibrotic tissue or hearts however can not generate EADs or triggered when exposed to similar or more intense stressful conditions. This discrepancy supports the supposition that cell-to-cell coupling is a potent mechanism suppressing EAD formation in tissue, by creating a source-to-sink mismatch that prevents local EAD currents generated by a small group of myocytes from reversing repolarization when they are electrotonically coupled to a large group of adjacent normally repolarizing myocytes.

To explore the effects of cell-to-cell coupling on EAD suppression, we performed computer simulations in 2D cardiac tissue in which normal
myocyte-to-myocyte coupling was disrupted by inserting fibroblasts into the tissue [10]. With too few fibroblasts, EADs and triggered activity were suppressed by cell-to-cell coupling, and with too many fibroblasts, while the EADs occurred but they were unable to propagate. With an intermediate myocyte-fibroblast ratio, however, EADs both formed and successfully propagated into the surrounding tissue. These findings agreed well with the experimental results in aged rat hearts, in which EADs and TA typically arose from regions with intermediate levels of fibrosis (30-40%), usually at the base of the LV epicardium. Two important factors tip the balance towards the emergence of EAD: 1) the process of synchronization of EADs.[34] and 2) increased cardiac fibrosis causing reduced gap junction coupling.[35] We recently described a synchronization mechanism for EADs based on the evidence that the irregularity of EADs is a form of dynamical chaos and that contiguous cells engaged in EAD formation actually synchronize over a finite length scale. However, when the tissue exceeds the critical size, electrotonic coupling can no longer globally synchronize EADs, resulting in regions of partial synchronization that shift in time and space. The regionally synchronized EADs then form premature ventricular complexes that propagate into recovered tissue without EADs causing premature ventricular depolarization and in the case of triggered activity focal VT. How EADs overcome electrotonic source-sink mismatches in tissue to trigger premature ventricular complexes remains incompletely understood. To study this question, we used a rabbit ventricular action potential model to simulate tissues in which a central area of contiguous myocytes susceptible to EADs was surrounded by unsusceptible tissue. In 1D tissue with normal longitudinal conduction velocity (0.55 m/s), the numbers of contiguous susceptible myocytes required for an EAD to trigger a propagating action potential was 70. In 2D tissue, the number of cells increases to 6940 and in 3D tissue to 696,910. The number of the cells decreases considerably when the gap junction conductance between the myocytes becomes reduced from 780 nS to 125 nS (6.25 fold) as might be expected to occur in fibrotic hearts. The decrease in the number of cells was proportionately more in 3D (93% reduction in the number of cells, i.e., 48,700 cells) than in 2D (84%, i.e., 1,100 cells) than in 1D (57%, i.e., 30 cells) [35]. These simulation studies have shown that the source-to-sink mismatch in well-coupled cardiac tissue provides a powerful mechanism to protect the heart from EAD-mediated arrhythmias. In contrast however when fibrosis develops and gap junctional couplings between myocytes decreases a considerable decrease in the number of myocytes needed to promote a PVD and VT occur presumably leading to increased vulnerability to VT/VF in intact fibrotic hearts.

Transition from focal VT to mixed reentrant and focal VF

The EAD-mediated focal VT degenerated within 3 sec of the onset to sustained VF (CL of 105±15 ms), requiring electrical shock for termination. The focal VT propagated as single wavefront from the base of the heart to the apex promoting spatially discordant action potential duration (APD) alternans (Figure 1) causing localized conduction block (wavebreak) midway between the base and the apex of the LV anterior epicardial surface. After the block, the wavefront continued to propagate laterally past the site of block forming a figure-8 type reentry, coinciding in time with the transition from the VT to VF on the ECG (Figure 1). This pattern of VT to VF transition occurred in about 70% of the VF episodes. In the remaining episodes the sites of the wavebreak could not be defined. Whether focal VTs seen in patients are also caused by EAD-mediated triggered activity remains to be elucidated [36,37-39].

Electrophysiological differences between young adult and aged rat hearts

Although disruption of normal cell-to-cell coupling by fibrosis provides a plausible explanation for the increased susceptibility of aged hearts to EADs and EAD-mediated arrhythmias, we cannot exclude the possibility that other aspects of aging-related remodeling (e.g. electrical, Ca²⁺ cycling, and gap junction remodeling) also make important contributions. Aging-related changes in electrophysiological and Ca cycling properties at the myocyte level may also be important in the genesis of EADs. In comparing young adult versus aged rat hearts, however, we found that H2O2 had no significant effect on action potential duration (APD) or the maximum APD restitution slope, both before and after exposure to H2O2. On the other hand, some differences in Ca²⁺ cycling between young adult and aged rat hearts were noted. The mean

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The rate constant of the Ca\(^{2+}\) transient decline (fit to a single exponential) was significantly longer in aged compared to young adult ventricles (P<0.05), both before and after H\(_2\)O\(_2\) exposure. Furthermore, there was greater regional heterogeneity in Ca\(^{2+}\) handling in the aged than young adult rat hearts, with the slowest decline rate constant clustering at the base of the LV compared to mid and apical sites. Overall, our findings in the aged fibrotic hearts with oxidative VF support a critical role of partial cellular uncoupling in EAD formation at tissue level caused by increased interstitial tissue fibrosis.

Has the time come for clinical trials to manage cardiac arrhythmias with anti-fibrotic therapy?

The link between ventricular fibrosis and ventricular arrhythmia risk suggests that targeting fibrosis may impart antiarrhythmic benefits. For example, a recent study found that in patients with hypertrophic cardiomyopathy, myocardial fibrosis as measured by the late gadolinium enhancement cardiovascular magnetic resonance (CMR) is an independent predictor of adverse outcome [40]. Interestingly these investigators found that the extent of myocardial fibrosis was an independent predictor for arrhyth-

Figure 2 Simultaneous microelectrode and ECG recordings at the onset of VT/VF in an aged rat heart exposed to 0.1 mM H\(_2\)O\(_2\). Panel A, onset of early afterdepolarization (EAD)-mediated triggered activity (TA) causing ventricular tachycardia (VT) 5 min after H\(_2\)O\(_2\) exposure. Note the smooth emergence of EAD (upward pointing arrow) during the isoelectric interval on the ECG followed by a run of 10 TA (downward pointing arrow) causing non-sustained VT on the ECG. The onset of the EAD precedes the QRS complex of the VT by 8 ms indicating absence of electrical activity elsewhere in the heart. Two additional short runs of VT with 4 beats each are also shown that follow a single subthreshold EAD (downward small arrow) with no TA. Panel B shows the degeneration of the TA to VF 15 min after H\(_2\)O\(_2\) exposure. (From reference number 10, Morita et al)
that the agent relaxin-1 reverses cardiac fibrosis and related cardiac dysfunction [50]. Relaxin is a potent antifibrotic peptide hormone that inhibits fibroblast activation (indicated by suppressed expression of α-smooth muscle actin) and collagen synthesis stimulated by angiotensin II or transforming growth factor-β [50].

It is hoped that these basic research findings will be translated to patients at risk of developing cardiac fibrosis-related related arrhythmias. To the extent that such a translation will be successful it is anticipated that a more rational and effective care of patients at risk of VT/VF may be developed.

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