Mitochondrial fusion dynamics is robust in the heart and depends on calcium oscillations and contractile activity

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Mitochondrial fusion is thought to be important for supporting cardiac contractility, but is hardly detectable in cultured cardiomyocytes and is difficult to directly evaluate in the heart. We overcame this obstacle through in vivo adenosinergic transduction with matrix-targeted photoactivatable GFP and confocal microscopy. Imaging in whole rat hearts indicated mitochondrial network formation and fusion activity in ventricular cardiomyocytes. Promptly after isolation, cardiomyocytes showed extensive mitochondrial connectivity and fusion, which decayed in culture (at 24–48 h). Fusion manifested both as rapid content mixing events between adjacent organelles and slower events between both neighboring and distant mitochondria. Loss of fusion in culture likely results from the decline in calcium oscillations/contractile activity and mitofusin 1 (Mfn1), because (i) verapamil suppressed both contraction and mitochondrial fusion, (ii) after spontaneous contraction or short-term field stimulation fusion activity increased in cardiomyocytes, and (iii) ryanodine receptor-2-mediated calcium oscillations increased fusion activity in HEK293 cells and complementing changes occurred in Mfn1. Weakened cardiac contractility in vivo in alcoholic animals is also associated with depressed mitochondrial fusion. Thus, attenuated mitochondrial fusion might contribute to the pathogenesis of cardiomyopathy.

Cardiac contractions require a constant energy supply, which is provided by mitochondrial metabolism. ATP is needed for excitation-contraction coupling (ECC) for both contraction and relaxation in each cycle (1, 2). ECC-associated cytoplasmic 

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Significance

Mitochondrial function is supported by dynamic quality control processes, such as mitochondrial fusion. Cardiac contractility depends on mitochondrial metabolism, yet in cardiomyocytes, mitochondria are confined among myofilaments, raising questions about the possibility of mitochondrial physical communication. Here we demonstrate that mitochondrial continuity is robust and fusion is frequent in freshly isolated rat ventricular myocytes, manifesting both as rapid content mixing events between adjacent organelles and slower, often long-distance events. We show that mitochondrial fusion decreases dramatically in culture because of the decay in contractile activity and, more specifically, the underlying calcium oscillations, which involve mitofusin 1 (Mfn1) abundance. In addition, we show that attenuation of cardiac contractility in vivo in alcoholic animals is also associated with depressed mitochondrial fusion.


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confocal microscopy. We selected between three and five 25-μmDsRed and mtPA-GFP, and evaluated their distribution by two mitochondrial matrix-targeted soluble fluorescent proteins, tions among mitochondria in cardiac myocytes, we expressed isolated acute trains of stimulation (20, 31). Thus, it is plausible that cardiac contractile activity might exert some control over cardiac mitochondrial fusion. We speculate that the low frequency of mitochondrial fusion events previously documented in adult cardiomyocytes kept in culture for 48–72 h (16) might be a consequence of the decay in contractile and/or metabolic activity.

Contractile activity becomes depressed in dilated cardiomyopathy, a common cause of which is chronic alcohol consumption (32). Mitochondria are a primary target of alcoholic tissue injury (33), and altered mitochondrial ultrastructure has been reported in alcoholic cardiac cells (34). In addition, our previous studies documented attenuated mitochondrial fusion dynamics in both skeletal muscle fibers (17) and hepatocytes (35) isolated from ethanol (EtOH)-fed rats. To investigate whether suppressed mitochondrial quality control might be a factor in the progression of alcoholic cardiac muscle dysfunction, in the present work we evaluated cardiac mitochondrial dynamics in chronically EtOH-fed rats.

The technical challenges of measuring mitochondrial dynamics and the scanty information available on mitochondrial content exchange in contracting cardiomyocytes have precluded answering fundamental questions about mitochondrial quality control in the heart, including the specific mechanisms of content exchange, the dependence of content exchange events on ECC activity and the possible involvement of content exchange processes in cardiac disease. Here we applied photoactivatable fluorescent protein techniques to evaluate mitochondrial dynamics in cardiomyocytes and in intact heart.

Results
Matrix Continuity Among Mitochondria in Rat Cardiac Myocytes: Effect of Maturation State and Cell Culture. To study the interactions among mitochondria in cardiac myocytes, we expressed two mitochondrial matrix-targeted soluble fluorescent proteins, mtDsRed and mtPA-GFP, and evaluated their distribution by confocal microscopy. We selected between three and five 25-μm² square areas in each cell and illuminated them to achieve two-photon (2P) photoactivation (PA) of PA-GFP, and followed the spreading of mtPA-GFP out of the illuminated regions. The fluorescence decay within the PA areas represents the sum of diffusion within the mitochondrial network, mitochondrial fusion, and mitochondrial movement (17). Following the strategy described in a recent study (16), we first evaluated 24- to 48-h–cultured neonatal ventricular cardiomyocytes (NVCMs). NVCMs showed robust spreading of mtPA-GFP (Fig. 1 A and B, column 1 and Movie S1), reaching distant organelles within 5 min (quantified in Fig. 1 C).

In contrast to mitochondria in NVCMs, mitochondria in AVCMs rearranged into longitudinal columns of apparently unconnected organelles and subgroups of less well-organized perinuclear organelles (Fig. 1 A and B, columns 2–4 and Fig. S1). AVCMs were first transformed in vitro by adenoviral mtDsRed and mtPA-GFP. Cells were infected promptly after plating and imaged 24–36 h later, once the fluorescent protein expression reached detectable levels. We focused on the interactions among intermyofibrillar mitochondria, given their role in regulating contractile function and their strategic orientation toward the SR (4). Spreading of mtPA-GFP beyond the edges of the PA area (Fig. 1 A–C, column 2 and Movie S2) demonstrated some continuity...
among mitochondria in AVCMs; however, this diffusion was limited and only over short distances.

We reasoned that because culture conditions negatively affect contractile function (56), they also might alter mitochondrial dynamics. Thus, our next step was to perform in vivo adenoviral infection of the ventricular walls of adult rat hearts. Then, 7–10 d later, we isolated AVCMs and plated them for microscopic evaluation at 24–48 h postharvest or promptly after plating. Fig. 1 A–C, column 3 and Movie S3 show diffusion of mtPA-GFP comparable to that observed in in vitro-transformed AVCMs (column 2). Freshly isolated in vivo-transduced AVCMs exhibited significant mitochondrial continuity, characterized by both longitudinal and transversal diffusion of mtPA-GFP (Fig. 1A, column 4 and Movies S4 and S5), extending >5 μm away from the PA region (Fig. 1B). We also found a significant decrease in mtPA-GFP fluorescence within the PA regions, with 25.6% decay at 300 s, as opposed to 14.5% in cultured AVCMs (P < 0.01) (Fig. 1C).

We further evaluated the diffusion of mtPA-GFP above and below the PA focal plane in fresh AVCMs by means of Z-stack imaging at 8 min after PA. Mitochondria containing photostable mtPA-GFP were found even 12 μm below (~12 μm) the PA focal plane (Fig. 1D). In addition, at the +3 μm focal plane, a discrete row of mitochondria projected toward the periphery, extending 12 μm away from the PA region. Finally, high-magnification 3D-projection images (from different views) showed mitochondrial matrix continuity in both longitudinal and transversal orientations in a freshly isolated AVCM. We performed a comparable 3D analysis in in vitro- and in vivo-transduced, AVCMs cultured for 24–48 h and found a drastic decrease in mitochondrial continuity compared with freshly isolated myocytes (Fig. S1B). We conclude that mitochondria in AVCMs form complex networks with matrix content exchange, and that culture conditions cause suppression of mitochondrial communication. Moreover, in NVCMs, mitochondria are less ordered and communicate actively with their neighbors. **

Active Mitochondrial Fusion Dynamics in Freshly Isolated AVCMs.** We next evaluated the mitochondrial fusion activity in freshly isolated AVCMs transduced with the reporters in vivo. When mtPA-GFP is photoactivated, mtDsRed becomes photobleached in the same area, producing green-only mitochondria surrounded by red-only mitochondria. During the merging of green-only and red-only mitochondria, abrupt complementary changes in green and red fluorescence validate fusion pore opening and matrix content mixing driven by the concentration gradient. We observed fusion events among intermyofibrillar mitochondria, showing diverse orientations and kinetics (Fig. 2 A and B). Images of an AVCM before and 12 s after PA highlight the early spreading of PA-GFP (Fig. 2A), which indicates preexisting matrix continuity among individual mitochondria. Further examination of the distribution of mtPA-GFP in discrete mitochondria located in the proximity of the PA areas revealed abrupt increases in the area of the fluorescent protein, corresponding to mitochondrial fusion events; as an example, a series of organelles underwent sequential fusion events in longitudinal orientation (Fig. 2B). Another example shows a subregion of a PA area and its surroundings, presenting four sequential fusion events in both longitudinal and transversal orientations, characterized by a decrease in PA-GFP fluorescence in the donor mitochondrion and a parallel increase in a nearby acceptor mitochondrion (Fig. S2A and Movie S6). Noticeably, NVCMs also exhibited sequential fusion events among neighboring mitochondria (Fig. S2B). In quantitative analyses of the fusion events in AVCMs maintained under different culture conditions (Fig. S2C), freshly isolated AVCMs exhibited 1.4 ± 0.1 events/min, whereas cultured in vivo-transduced AVCMs showed four-fold fewer events, and cultured in vitro-transformed AVCMs showed almost no events (a rare event is shown in Fig. S2C). Thus, AVCM mitochondria regularly undergo fusion, and this activity decays in culture.

Transmission electron microscopy (TEM) in fixed heart muscles has demonstrated that intermyofibrillar mitochondria run parallel to myofibrils, mostly forming columns of individual mitochondria, accompanied by some transversally oriented organelles. Such a distribution in live AVCMs is also supported by our confocal microscopy data. Moreover, the fusion events occurred predominantly among longitudinally oriented organelles (Fig. 2D), although 32% of the interactions occurred transversally.

Slow Matrix Content Mixing Events in AVCMs.** Different from other cells that we have studied previously, including mouse embryonic fibroblasts (MEFs), H9c2 cells, hepatocytes, and skeletal muscle fiber cells (7, 17, 35), AVCMs exhibited diverse mixing kinetics of the matrix-targeted fluorescent proteins (Fig. 3). Two representative examples are shown in Fig. 3A and Movie S7. In the figure, the left panel shows an event with fast mixing kinetics between two longitudinally oriented mitochondria. This mixing event was completed within 12 s, as validated by the time course of both PA-GFP and DsRed in the donor and acceptor mitochondria. The complementary equilibration of both fluorescent proteins confirms that a fusion event occurred. The right panel shows a comparable fusion event example in terms of location relative to the PA region and size of both the PA-GFP donor and the acceptor. The transfer of the fluorescent proteins was slower, however, taking 70 s to reach equilibrium. The slow mixing kinetics applied to both fluorescent proteins, arguing against the possibility of an artifact associated with the conformation of either protein. Thus, based on the mixing kinetics, we discriminated between fast and slow fusion events, which are completed in <12 s and >12 s of mitochondrial matrix mixing, respectively. Overall, slow mixing kinetics fusion events are prevalent in AVCMs (Fig. 3D) and are likely to represent a tissue-specific feature of cardiac mitochondrial fusion.
We next searched for possible explanations for the slow matrix mixing kinetics. We found some evidence suggesting the IMM fusion pore might open in an intermittent fashion (Fig. 3B shows a two-step fusion event), similar to the fusion between exocytotic vesicles and the plasma membrane (37). Overall, 11% of the slow mixing kinetics events displayed detectable multistep mixing kinetics (Fig. S3A and Movie S8). Furthermore, the previously described narrow intermitochondrial nanotunnels (16) also might support slow content mixing. Indeed, we observed a nanotunnel-like structure growing out from a small PA-GFP–carrying mitochondrion and reaching a distant mitochondrion, which was associated with slow mixing kinetics (Fig. 3C). We also detected slow mixing kinetics between two non-neighbor mitochondria (Fig. S3A). In addition, we found narrow tubes apparently emerging from larger mitochondria (Fig. S3B and Movie S9). These examples demonstrate a fluorescence distribution consistent with the presence of nanotunnels, and demonstrate that these structures make mitochondrial soluble content exchange possible, accompanied by slow mixing kinetics.

Mitochondrial Fusion Dynamics in Whole Heart. To image mitochondria in intact tissue, in vivo-infected whole hearts were perfused and examined by confocal microscopy. The mitochondrial structure in the outer layer of myocytes in the ventricular wall was first visualized using tetramethylrhodamine, ethyl ester (TMRE), which accumulates in the mitochondria in a membrane potential-dependent manner (Fig. 4A). Cardiomyocytes of the ventricular wall of in vivo-transformed hearts exhibited mtDsRed fluorescence distribution comparable to TMRE localization (Fig. 4B). Approximately 40% of the myocytes expressed mtDsRed. Once we found a myocyte positioned parallel to the coverslip and showing well-defined organelles, we applied 2P illumination in discrete 25-μm² square regions and followed the time course after the PA of mtPA-GFP. Immediately after the end of PA, mtPA-GFP diffused into the surrounding mitochondria around the PA regions, mostly in longitudinal orientations (Fig. 4C). Interestings, at 6 min after PA, region 4 showed a new mitochondrion located 1 μm away from the corner of R4 and transversally oriented to the PA-GFP–containing mitochondria inside it, possibly indicating a fusion event. Owing to contraction, recording the kinetics of PA-GFP spreading was difficult; nonetheless, some decay in PA-GFP fluorescence was apparent even in the first 80 s after PA (Fig. 4D). Despite the complexity of the recording, we could identify fusion events. One example shows the mixing of mtPA-GFP and mtDsRed between two neighbor mitochondria (Fig. 4E). Therefore, these data confirm that the observed mitochondrial fusion activity found in isolated myocytes also occurs within cardiac muscle.
Motility of Mitochondria in Cardiac Myocytes. Given the relevance of mitochondrial movements as a facilitator of mitochondrial encounters and fusion events (7, 38), we studied motility in ventricular myocytes. NVCMs exhibited highly active mitochondrial movements throughout the cells (Fig. 5A). In contrast, cultured AVCMs showed minimal motility (data not shown). Moreover, mitochondria in the whole heart did not exhibit motility (Fig. 5C); however, some motility activity was present in freshly isolated AVCMs (Fig. 5B). In addition to Brownian-like movements (Fig. 5B), we found an isolated displacement motility event restricted to a small region (Fig. 5B, Lower). The limited motility observed in the AVCMs indicates that mitochondrial movements do not have a major role in supporting mitochondrial fusion in cardiac muscle.

Mitochondrial Fusion Depends on Excitation-Contraction Activity. The foregoing results indicate a negative effect of cell culture conditions on mitochondrial communication in AVCMs. The regular ECC activity decays in AVCMs after isolation and culture. This raises the possibility that ECC function is required to support mitochondrial fusion, which we tested more directly using the following approach. We first treated the cells with verapamil, an inhibitor of Ca\(^{2+}\) entry via L-type Ca\(^{2+}\) channels to suppress contractile activity in neonatal and freshly isolated ventricular myocytes. This treatment caused fewer cells to contract and contracting cells to show fewer contractions (Fig. 6A and B, Left), and partially decreased the mitochondrial fusion event rate in both NVCMs and freshly isolated AVCMs (Fig. 6A and B, Right). We also analyzed the rate of fusion events before and after the occurrence of a spontaneous contraction in AVCMs, and detected an increase in mitochondrial fusion frequency in the first 2 min after a spontaneous contraction (Fig. 6C). Finally, we evaluated the short-term effect of ECC activity...
in freshly isolated AVCMs by field-stimulating the cells for 5 min. (Fig. S4 shows a schematic presentation of the stimulation protocol.) After stimulation, we found a 60% increase in mitochondrial fusion compared with control cells (Fig. 6D). Thus, we conclude that regular ECC facilitates the mitochondrial fusion activity of AVCMs.

**RyR2-Mediated [Ca\textsuperscript{2+}] Oscillations Are Sufficient to Promote Mitochondrial Fusion.** Each ECC event in AVCMs is triggered by plasma membrane depolarization and an ensuing [Ca\textsuperscript{2+}], transient. To isolate the potential role of the depolarization and [Ca\textsuperscript{2+}], increase from contraction, we first used noncontracting H9c2 cardiac myoblasts. Depolarization evoked by increased KCl and [Ca\textsuperscript{2+}], elicited by a Ca\textsuperscript{2+} ionophore, ionomycin, failed to increase an mitochondrial connectivity (Fig. 7A and B). However, ionomycin causes a single and prolonged [Ca\textsuperscript{2+}], rise, whereas AVCMs display RyR2-mediated [Ca\textsuperscript{2+}], oscillations. A surrogate for these [Ca\textsuperscript{2+}], oscillations without contractile activity is provided by HEK293 cells expressing RyR2 (HEK-RyR2) in a tetracycline-inducible manner (39). Indeed, tetracycline-pretreated HEK-RyR2 cells displayed a moderately elevated average [Ca\textsuperscript{2+}], compared with nontreated cells (Fig. 7C), owing to an increase in the fraction of cells with [Ca\textsuperscript{2+}], spiking from 4.6% to 63.6%, manifesting in the form of baseline–spike [Ca\textsuperscript{2+}], oscillations (Fig. 7D). Simultaneous imaging of [Ca\textsuperscript{2+}], and mitochondrial dynamics (with mtDsRed and mtPA-GFP) revealed no difference in mitochondrial morphology between nonoscillating and oscillating HEK-RyR2 cells (Fig. 7E and F); however, the spread of the matrix-targeted fluorescent proteins, and specifically the number of mitochondrial fusion events, was enhanced in the oscillating cells (Fig. 7G). Interestingly, when caffeine (10 mM), a stimulator of RyR2, was added, a single and large [Ca\textsuperscript{2+}], transient occurred, and the oscillations stopped (Fig. 7H). This was followed by decreases in mitochondrial matrix continuity and fusion activity (Fig. 7I). Taken together, these results suggest that the effect of cardiac contractile activity on mitochondrial fusion could be mediated by the [Ca\textsuperscript{2+}], oscillations.

To identify the potential target of ECC activity/[Ca\textsuperscript{2+}], oscillations in the control of mitochondrial fusion, we quantified the abundance of OMM and IMM fusion proteins in freshly isolated and overnight-cultured AVCMs. Immunoblotting indicated significantly decreased Mfn1 in the overnight-cultured AVCMs, but no change in Mfn2 and Opal (Fig. S5A). To test for a cause-and-effect relationship between Mfn1 decrease and decay in mitochondrial fusion activity, we used an adenosine carrying a mouse Mfn1 expression construct (which shares 99% homology with rat Mfn1) in vitro-transfected nonoscillating AVCMs with mitochondrial fusion rescued mitochondrial continuity and fusion frequency compared with cells transformed with a control adenosine, AdLaCZ (Fig. S5C). The efficiency of AdMfn1 for actually rescuing the expression of Mfn1 was validated in MEF Mfn1\textsuperscript{-/-} cells in terms of both Mfn1 abundance and mitochondrial morphology (Fig. S5B). Complementing the results in AVCMs, HEK-RyR2 cells displayed higher Mfn1 levels compared with plain HEK cells (Fig. S5D). Thus, these results indicate that ECC activity/RyR2-mediated [Ca\textsuperscript{2+}], oscillations help maintain Mfn1 abundance as a mechanism to support mitochondrial fusion activity. A decrease in Mfn1 might account for the long-term effects of ECC activity, but is unlikely to explain the short-term changes in fusion activity observed on pacing in cultured AVCMs (Fig. 6D) and cessation of [Ca\textsuperscript{2+}], oscillations in HEK-RyR2 cells (Fig. 7I and Discussion).

**Suppression of Cardiac Mitochondrial Fusion by Chronic EtOH Exposure.** Environmental and dietary stressors target the heart’s contractile performance and represent a leading cause of death worldwide. Among these stressors, chronic EtOH consumption leads to dilated cardiomyopathy (32). Given that mitochondria are well-known targets of EtOH (33), and that our previous data have shown suppression of mitochondrial fusion in skeletal muscle by prolonged EtOH exposure (17), we performed experiments to evaluate the effect of chronic EtOH on mitochondrial fusion in cardiomyocytes. We first treated NCMVs in vitro with 50 mM EtOH for 48 h, which represents an in vitro model for chronic alcohol exposure. The EtOH-treated NCMVs exhibited a 40% decrease in mitochondrial continuity and a 75% decrease in mitochondrial fusion activity (Fig. 8A).

As an in vivo model, we used EtOH-fed (a 32% EtOH-containing diet for 6–9 mo) and pair-fed rats in which EtOH-induced contractile dysfunction was documented as decreased ejection fraction and fractional shortening (Materials and Methods). AVCMs isolated from the EtOH-fed rats showed unaltered resting mitochondrial membrane potential, as assessed by the potentiometric dye TMRE (EtOH: 23 ± 4% uptake, pair-fed: 21 ± 1% uptake; n = 5), though previous studies reported an increased sensitivity to Ca\textsuperscript{2+}-induced depolarization and permeability...
structure of normal and alcoholic AVCMs. Shown are representative images (arrowheads). Bar charts denote diffusion kinetics for control (cells) compared with EtOH-treated cells (Eisner et al. PNAS Early Edition).

Fig. 8. Effect of prolonged EtOH exposure on cardiac mitochondrial fusion dynamics. (A, Left) Diffusion of mtPA-GFP in control and EtOH-treated (50 mM for 48 h) NVCMs. Arrowheads indicate diffusion of mtPA-GFP due to pre-established connections (42 s) or newly formed connections due to fusion events (477 s). (A, Center) mtPA-GFP fluorescence decay in the PA area. (A, Right) Bar charts showing a greater percentage of decay at 300 s post-PA in control cells (n = 57 ROIs) compared with EtOH-treated cells (n = 59) (Upper Right) and a higher fusion event frequency in control cells (n = 63 cells) compared with EtOH-treated cells (n = 33) (Lower). (B) In vitro-transduced AVCMs from control and EtOH-fed rats (age 6–9 mo). Images of control and EtOH-treated myocytes show limited mtPA-GFP diffusion (arrowheads). Bar charts denote diffusion kinetics for control (n = 12) and EtOH (n = 10), expressed as the area covered by the PA-GFP PA areas. (C) Ultrastructure of normal and alcoholic AVCMs. Shown are representative images of longitudinal arrangements of sarcomeres (S) and mitochondria (M).

transition (33, 40). Here we infected in vitro the freshly harvested AVCMs with mtPA-GFP and mtDsRed, and found significantly reduced mitochondrial continuity in the cells derived from the EtOH-fed rats (Fig. S8). We next measured the levels of mitochondrial fusion proteins Mfn1, Mfn2, and Opal in mitochondrial fractions of control and EtOH-fed rats. No significant change was detected (Fig. S6), but in three of the five pairs Mfn1 was apparently somewhat decreased in the EtOH-fed rat, suggesting that EtOH can target mitochondrial fusion in the heart without altering the abundance of fusion proteins though a moderate decrease in Mfn1 might be involved.

To further evaluate the dysregulation induced by chronic EtOH exposure, we studied the ultrastructure of mitochondria in the ventricular muscles of hearts harvested from control and EtOH-fed rats (Fig. 8C). Mitochondrial cross-sections accounted for 34% and 37% in hearts from control and EtOH-fed animals, respectively (no significant difference); however, the individual mitochondrial cross-section areas in the hearts from EtOH-fed rats showed a tendency for enlargement (control: 0.74 ± 0.14 and EtOH: 0.81 ± 0.15 μm², n = 3, P = 0.056). In addition, mitochondria that showed signs of swelling, distortion, and/or resorption of cristae were more prevalent in the cardiac muscles of the EtOH-fed rats (Fig. S7). Using a scoring system for cristae abundance and form ranging from 0 (worst) to 4 (best), the mean score for control was significantly higher in control rats compared with the EtOH-fed rats (2.45 ± 0.23 vs. 1.92 ± 0.2, P = 0.05, paired t test) (Fig. S7B). Comparing the prevalence of individual score grades, grade 0 (no well-defined crista) and grade 1 (no crista in >50% of the mitochondrial area) were significantly more prevalent in the EtOH group (Fig. S7C). Thus, our results suggest that chronic EtOH consumption causes suppression of mitochondrial fusion activity in cardiac muscle and leads to distortion of the ultrastructure.

Discussion

We have demonstrated that in the adult heart, mitochondria exhibit active fusion dynamics to support matrix content exchange among individual organelles. Distinctively, content exchange among cardiac mitochondria often occurs with slower kinetics compared with that in neonatal heart or in any previously studied cell types. A systematic comparison of freshly isolated and cultured ventricular myocytes revealed that mitochondrial fusion activity decays rapidly when cardiomycocytes are transferred to cell culture. This effect might result from a decrease in contractile activity, as demonstrated by the ability to down-regulate and up-regulate mitochondrial fusion by suppressing and restoring cardiomycocyte contractions, respectively. More specifically, we have shown that oscillatory, but not sustained, Ca²⁺ transients significantly increase mitochondrial fusion frequency in RyR2-expressing HEK cells, suggesting that the Ca²⁺ component of the ECC alone can regulate cardiac mitochondrial communication, and that this mechanism seems to involve Mfn1. Finally, we also have shown that chronic alcohol-fed rats, which exhibit a cardiomyopathy phenotype, show impaired mitochondrial fusion and ultrastructural impairments.

Although mitochondrial fusion has been visualized in diverse differentiated cells, such as neurons (41), hepatocytes (35), and skeletal muscle fibers (17, 18), previous studies of cardiac mitochondrial fusion have been largely confined to evaluating cardiac function after genetic manipulation of the expression of fusion and fission proteins (13, 14, 21, 22). Mitochondrial “kissing” was recently described (16) in cultured AVCMs studied at 48–72 h postisolation. However, our study illuminates the active communication between intermyofibrillar mitochondria in noncultured adult cardiomycocytes, as well as in situ within cardiac muscle. This is a major step forward, because the fusion activity under these conditions is much higher than that in cultured primary cardiomycocytes, indicating that cardiac mitochondrial fusion activity has been underestimated until now.

The lesser mitochondrial continuity and motility observed in cultured AVCMs compared with NVCM is in accordance with our previous observations in differentiated and nondifferentiated cardiac-derived H9c2 cells and in skeletal myoblasts, and indicates decreased mitochondrial continuity and fusion in differentiated striated muscle (17, 42). Mitochondrial fusion is supported by mitochondrial motility, which brings mitochondria into contact to create the opportunity for fusion (7, 38). Multidirectional orientation patterns of microtubules and abundant mitochondrial movements occur in NVCMs (43), H9c2 cells (44), and skeletal myoblasts and myotubes (17). In contrast, in AVCMs and skeletal muscle fibers, mitochondria are spatially confined to the space among the myofilaments and have no opportunity to move toward distant mitochondria. This might account for the lower frequency of fusion events seen in cultured AVCMs and other differentiated muscle paradigms (17, 45). In addition to the structural confinement of mitochondria, in vivo and cell culture conditions seem to affect mitochondrial motility differently, with
mitochondria showing little movement in freshly isolated striated and vascular smooth muscle but vigorous displacement in proliferating cultured striated and smooth muscle cells (17, 46).

Despite the similar fiber organization in adult skeletal muscle fibers and AVCMs, the former exhibit 0.5 fusion event/min, compared with 1.4 fusion events/min in the latter. This difference may be linked to tissue-specific abundance of mitochondrial fusion and fission proteins; however, compared with skeletal muscle, cardiac muscle contains more abundant and larger mitochondria. Unexpectedly, we also found that the fusion-mediated exchange of soluble matrix content shows slower kinetics than that in skeletal muscle and other previously characterized paradigms. Based on our observations and the literature, several factors might explain the unusual complementation kinetics of cardiac mitochondria. First, our evidence for discrete and partial repetitive steps of mtPA-GFP transfer shows a similarity to neurotransmitter release by exocytosis, where fusion of neurotransmitter-filled vesicles with the plasma membrane occurs through partial opening of a fusion pore, leading to a so-called “kiss-and-run” between the membranes, which remain connected via a nanotube that opens up to a larger pore and then recloses to a nanotube (37). Thus, it might be speculated that the IMM undergoes fast intermittent fusion pore openings as well. Second, the observation of narrow connectors that resemble those previously described in heart mitochondria (16, 34), along with the results of Lavorato et al. (47) showing that in a model of catecholaminergic polymorphic ventricular tachycardia (RyR2-A4860G), nanotunnel structures coupled with slow matrix mixing events become more frequent, support the idea of nanotunnels mediating slow mitochondrial fusion in AVCMs. Third, ample ultrastructural evidence shows that the IMM is highly folded and the numerous cristae define narrow compartments of the matrix specifically in cardiac mitochondria (12, 48, 49), which might have more restricted communication compared with the matrix areas of mitochondria in other tissues. Thus, multiple nonexclusive mechanisms might contribute to the slow content mixing during fusion of heart mitochondria.

A striking observation of the present study is the time-dependent decay in mitochondrial fusion activity when AVCMs are placed into cell culture. The decreases in mitochondrial communality and fusion activity are accompanied by a previously reported loss of transversal tubules and, consequently, dysregulation of the ECC-associated Ca\(^{2+}\) transients (36) and suppressed contractile activity. Therefore, we reasoned that the decay in mitochondrial fusion activity might result from the loss of regular contractile activity or Ca\(^{2+}\) transients. Indeed, we found that attenuation of the Ca\(^{2+}\) transients and contractraction leads to depressed fusion activity, and that stimulation of contraction in cultured AVCMs positively affects mitochondrial fusion activity. Until now, it was thought that in contrast to vesicular fusion, mitochondrial fusion does not depend on Ca\(^{2+}\) (50); however, a number of posttranslational modifications regulate mitochondrial dynamics at different levels, including phosphoregulation of Drp1 (51), redox modification of Mfn2 (52), Opa1 proteolytic processing (24), and others. In the heart, mechanical forces and stretch activate various signaling pathways, including such as NADPH-derived ROS, leading to hypertrophy through the activation of diverse signaling programs (53). Whether or not these pathways extended to mitochondrial fusion regulatory proteins was unknown. Our present data show activation of mitochondrial fusion by contraction at both short-term and long-term levels.

An adaptation of mitochondrial communication to persistent contraction is in accordance with previous studies in skeletal muscle showing that exercise enhances the apparent communication among mitochondria (29) and induces expression of mitochondrial fusion proteins (30). Our data show that the amount of the mitochondrial fusion protein Mfn1 decreases in response to a lack of contractile activity in cultured AVCMs, and that the rescue of Mfn1 by exogenous expression in cultured AVCMs leads to a recovery of mitochondrial fusion, highlighting this protein as the key player in striated muscle mitochondrial fusion regulation, in agreement with our previous data in skeletal muscle (17). In contrast to Mfn2, Mfn1 has an exclusive mitochondrial fusion-related function in muscle, whereas Mfn2 is also involved in cardiac mitochondria–SR interactions (26).

We also show that RyR2-dependent oscillatory [Ca\(^{2+}\)]\(_{cyt}\) transients improve mitochondrial connectivity and fusion frequency and also support Mfn1 up-regulation, which may reflect a Ca\(^{2+}\)-induced transcriptional regulation. Extensive studies have shown regulation of gene expression by Ca\(^{2+}\) in striated muscle and neurons, also known as excitation-transcription coupling (54–56). RyR1- and IP3R-dependent calcium transients differentially control gene expression in skeletal muscle (54). In the heart, IP3R-induced Ca\(^{2+}\) transients lead to activation of nuclear factor of activated T cells and hypertrophy (55). Interestingly, repetitive oscillatory Ca\(^{2+}\) signals are more efficient than sustained transients for inducing Ca\(^{2+}\)-dependent transcriptional factors (57). Along the line of these findings, ECC-evoked Ca\(^{2+}\) transients might regulate mitochondrial fusion at a transcriptional level in the heart; however, we also observed a rapidly reversible effect of Ca\(^{2+}\) oscillations on the maintenance of mitochondrial fusion activity in HEK-RyR2-expressing cells. This observation, together with the rapid enhancement of mitochondrial fusion activity observed on pacing or spontaneous contractions in AVCMs, suggest a fairly direct and acute positive effect of Ca\(^{2+}\) oscillations on mitochondrial fusion.

Given that Mfn1 abundance is unlikely to increase and decrease within minutes, an alternative mechanism must be involved in the acute effect of Ca\(^{2+}\) oscillations, such as Ca\(^{2+}\)-dependent regulation of mitochondrial metabolism, which is central to fusion activity (20, 58, 59), or Ca\(^{2+}\) binding to some components of the fusion machinery or to one of its posttranslational modifiers. Importantly, stopping and restarting of ECC activity does not occur under physiological conditions; thus, we can only claim that the maintenance of mitochondrial fusion activity involves a Ca\(^{2+}\) oscillation/contraction-dependent factor in the heart. The potential pathophysiological significance of this finding is highlighted by the work of Lavorato et al. (47), who report that a mutation in RyR2 (A4860G), which alters the normal Ca\(^{2+}\) oscillation phenotype, causes a striking change in mitochondrial fusion dynamics.

Several lines of circumstantial evidence implicate mitochondria in either the initiation or progression of cardiomyopathy (60). Mitochondrial involvement in alcoholic cardiomyopathy is intriguing, because mitochondria represent a primary cellular target of alcohol (33). Alcoholic cardiomyopathy is characterized by reduced contractility, along with arrhythmias and fibrosis (61, 62). Alterations in hearts from alcoholic patients at the level of mitochondrial communication were suggested based on early ultrastructural studies (34), and recent studies have reported impairment of mitochondrial DNA integrity by EtOH (63). Given the relevance of mitochondrial fusion to mtDNA stability (5), our results provide a potential structural explanation of these latter findings. A loss of mtDNA integrity was also found in skeletal muscle-conditional knockout of Mfn1 and Mfn2 (64). Moreover, our recent findings in skeletal muscle from alcohol-fed rats show that chronic EtOH consumption leads to inhibition of mitochondrial fusion and mitochondrial calcium uptake efficiency, accompanied by dysregulation of ECC-derived calcium transients, leading to fatigue (17). Our present results indicate that in the heart, alcohol targets mitochondrial dynamic communication via a mechanism that might be linked to the alcohol-induced reduced contractility. Given the apparent bidirectional dependence between contractile and mitochondrial fusion
activities, the pathogenesis of cardiomyopathy might include contractile and mitochondrial fusion impairments that facilitate one another. Based on our findings, it might be speculated that targeting the fusion-fission process may help improve cardiac function in alcoholics.

Materials and Methods
cDNA Constructs and Adenovirus. mtDSRed and mtPA-GFP plasmid DNA have been described previously (7). Type V adenoviruses carrying mtDsRed, mtPA-GFP, LacZ, and mouse Mfn1 were produced by Vector Biolabs.

Cardiac Myocyte Isolation, Cell Lines, and Cultures. NVCVs and AVCMs (14, 65), H9c2 cells, WT cells, and RyR2-overexpressing HEK293T cells (39) were cultured as described in SI Materials and Methods.

Animal Treatment. The EtOH-consuming adult male Sprague–Dawley rat model is described elsewhere (17). All procedures were done in accordance with the National Institutes of Health’s Guidelines on the Use of Laboratory Animals and were approved by Thomas Jefferson University’s Committee on Animal Care. Cardiac function was monitored by echocardiography as described earlier (66). These studies validated significant EtOH-dependent decreases in fractional shortening (FS) and fractional shortening (EF) (EF: control, 78 ± 2%; ETOH, 72 ± 2%; P < 0.02; FF: control, 48 ± 2%; ETOH, 43 ± 2%; P < 0.02). Thus, in the chronic ETOH-fed rats, contractile dysfunction develops in the heart.

Myocardial Gene Transfer. In vivo adenovalyn transformation of adult rat hearts is described in SI Materials and Methods.

TEM. Rat hearts were fixed by perfusion of 2.5% (vol/vol) glutaraldehyde in 0.1 M Na-cacodylate buffer (pH 7.4) and stained as described in SI Materials and Methods, followed by TEM image acquisition and analysis (17).

Live Cultured Cell Fluorescent Ca2+ Imaging and Confoical Microscopy. Fura2-AM–loaded cells were evaluated by epifluorescence microscopy (17). Confocal microscopy imaging of live cells and whole-heart experiments are described in detail in SI Materials and Methods.

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Supporting Information

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SI Materials and Methods

Cardiac Myocyte Isolation and Culture. NVCMs were cultured in F-10 Ham medium supplemented with 10% HS, 5% FBS (In VitroGen), and penicillin/streptomycin as described previously (14). The NVCMs were plated on top of 2% (wt/vol) gelatin-coated glass coverslips at a density of 5 × 10^5 cells per 35-mm dish. AVCMs were isolated from 2- to 10-mo-old rats (65) and plated on laminin (Roche)-coated coverslips in M199 complete medium (0.2% BSA, 5 mM taurine, 2.5 mM creatine, 2 mM carnitine, and 1x insulin-transferrin-selenium; Invitrogen) containing 1 mM BDM. Both NVCMs and AVCMs were cultured at 37 °C with atmospheric air and 5% CO2. Fluorometric measurements of the mitochondrial membrane potential were performed in permeabilized AVCMs as described previously (67, 68).

Cell Lines and Cultures. H9c2 cells of passage 36–40 were grown in low-glucose DMEM (Thermo Fisher Scientific) containing sodium pyruvate and 10% FBS (Thermo Fisher Scientific), supplemented with penicillin/streptomycin (Lonza), L-glutamine (Thermo Fisher Scientific), and additional sodium pyruvate (Sigma-Aldrich). WT HEK293T cells and Flip-in HEK293T cells overexpressing RyR2 (39) were grown in DMEM and 10% FBS, supplemented with penicillin/streptomycin and L-glutamine. When induction of RyR2 was desired, 1 μg/mL tetracycline (Sigma-Aldrich) was applied.

Myocardial Gene Transfer. Male Sprague–Dawley rats (150–180 g) were anesthetized with 2–3% isoflurane. A small skin incision was made in the left front chest. After dissecting pectoral muscles, exposing the ribs, and externalizing the heart through the fourth intercostal space, multiple-point left ventricular free wall injections were performed with an insulin syringe (30g needle). The syringe carried 180 μL of mDsRed and mtPA-GFP mixed adenoviral particles (ratio, 1:2). Immediately after gene delivery, the heart was placed back into the thoracic cavity, followed by suturing of the ribs and skin. The procedure took less than 2 min. The animal were returned to their cages and were given appropriate analgesia. At 6–8 d after the surgery, the animals were killed, and hearts were harvested for whole-heart imaging or myocyte isolation. No damage to or change in viability of AVCMs was observed after adenovirus injection (69).

In Vitro Transfection and Transduction of Cardiomyocytes. At 18 h after plating, the NVCMs were incubated with Optitrim (In Vitrogen) for 6 h and then transfected with a 1:1 mix of mDsRed and mtPA-GFP (total 4 μg of DNA) and Lipofectamine 2000 at 1 mL per dish. After overnight incubation, the medium was replaced by 0.5% FBS–F-10 Ham medium supplemented with 20 μM 5-bromo-2′-deoxyuridine (Sigma-Aldrich). Freshly isolated AVCMs were in vitro-transformed with a 1:1 mixture of AdmDsRed and AdmPT-GFP (a total of 7 × 10^9 pfu per 35-mm dish) in 1 mL of 0.1% BSA-M199 medium, supplemented with antibiotics and 1 mM BDM. Freshly isolated in vivo-infected AVCMs were in vitro-transformed with AdmMfn1 (7 × 10^9 pfu per 35-mm dish) or AdmLacZ (5.7 × 10^9 or 2.4 × 10^9 pfu per 35-mm dish) in 1 mL of 0.1% BSA-M199 medium, supplemented with antibodies and 1 mM BDM. After overnight incubation, the medium was replaced by M199 complete medium.

Transfection of H9c2 and HEK293T Cells. Cells were plated on poly-d-lysine (EMD Millipore)-coated 25-mm glass coverslips (Thermo Fisher Scientific) in 1.8 mL of medium in 35-mm tissue culture dishes (Falcon; Corning). At 24 h after plating, cells were transfected with a 1:1 mix of mDsRed and mtPA-GFP in solution (total of 2 μg of DNA) diluted with 6 μL of XtremeGene9 (Fugene; Roche) in 200 μL of Opti-MEM (Thermo Fisher Scientific). Imaging was performed at 24–48 h after the transfection.

In Vitro Chronic Alcohol Exposure. At 24 h after plating, NVCMs were treated with 0.3% vol/vol 50 mM EtOH for 48 h. Culture dishes were placed in a 100-mm dish containing 1% ethanol/water solution to maintain an alcohol-rich atmosphere. EtOH was renewed every 12–24 h.

Short-Term Field Stimulation. Cardiomyocytes were field-stimulated with 2-Hz pulses of 2-ms each, for 5 min in a custom-made chamber (2 mL, with two parallel platinum electrodes positioned on each side of the light path, 1 cm apart) connected to a square pulse stimulator and an oscilloscope. Cells were perfused with a 0.25% BSA CaCl2-containing solution during electrostimulation. Cells were tested for electric pulse-induced contractile activity, stimulated for 5 min with perfusion, and imaged for another 5 min. Control cells were perfused without electrostimulation for 5 min and then imaged for another 5 min.

Western Blot Analysis. Here 30 μg of membrane-enriched fractions were loaded onto each lane of an 8% SDS/PAGE. Primary antibodies for Mn1 (kindly provided by Dr. Richard Youle, National Institutes of Health), Mn2 (Sigma-Aldrich), Opalin (BD Biosciences), COX IV (Cell Signaling Technology), and Prohibitin (Abcam) were used. Bound antibodies were detected by chemiluminescence or IRDye 800 secondary antibodies (LI-COR Biosciences).

Transmission Electron Microscopy. Rats were injected with 1,000 U of heparin i.p. 15 min before induction of anesthesia with nembutal, followed by opening of the chest and cannulation of the aorta with an 18-1/2g i.v. cannula for in situ washing of the coronary arteries with oxygenated Ringer’s solution supplemented with 0.1% procaine for 3–5 min at a perfusion rate of 40 mL/min. Subsequently, the perfusate was switched to 2.5% (vol/vol) glutaraldehyde in 0.1 M Na-cacodylate buffer (pH 7.4). All perfusates were maintained at 4 °C. The ventricle was subsequently cut into <2 × 2-mm pieces and kept in the fixative solution at 4 °C for 2 h. The samples were then washed with 0.1 M Na-cacodylate buffer and postfixed with partially reduced osmium tetroxide (2% OsO4 and 0.8% potassium ferrocyanide in 0.1 M Na-cacodylate) overnight. Multiple washing steps with water were followed by en bloc staining with 1% uranyl acetate for 1 h at 4 °C. Stained samples were dehydrated on an acetone dilution series and embedded in Spurr’s resin (Electron Microscopy Sciences) according to the manufacturer’s instructions. Sectioning, image acquisition, and analysis were performed as described previously (17).

Live Cultured Cell Fluorescent Ca2+ Imaging. For imaging of [Ca2+]i, the cells were first loaded with 2 μM Fura2-AM (Life Technologies) for 20 min at room temperature in 2.0% (wt/vol) BSA ECM, in the presence of 0.003% pluronic acid and 100 μM sulfipyrazone. After dye loading, the coverslip was mounted to a heated (35 °C) incubator chamber in ECM containing 0.25% BSA and sulfipyrazone. Detection of Fura2 fluorescence (excitation, 340 and 380 nm; emission, 540/50 nm) was achieved using an inverted microscope (40×, DMIRE 2; Leica), UApio objective (Olympus), and a CCD camera (ProEM 1024 EMCCD).
camera, MultiFluo system; Princeton Instruments). Image duplets (excitation, 340 and 380 nm for Fura2) were obtained every 3 s for 10 min. For H9c2 cells, treatment with 60 mM KCl or 5 μM ionomycin was applied at image 10.

**Image Analysis.** Image analysis was done using custom-designed Spectralyzer software or Zen 2010 (Carl Zeiss). Mitochondrial matrix continuity was evaluated via two approaches using Fiji. Spreading of mtPA-GFP from the area of photoactivation was evaluated by masking the 5 × 5-μm photoactivation areas and quantifying the time-dependent decay in fluorescence intensity. Alternatively, the entire area outside the photoactivated areas was masked. The images were thresholded using Fiji, and every pixel outside the PA area that exceeded the threshold by >10% was included in the mtPA-GFP area of spreading. Fusion events/min = #fusion events × 3 ROIs/#ROIs/min, unless stated differently.

**Confocal Microscopy.** Imaging of isolated cells was performed in a 0.25% BSA ECM consisting of 121 mM NaCl, 5 mM NaHCO₃, 4.7 mM KCl, 1.2 mM KH₂PO₄, 1.2 mM MgSO₄, 2 mM CaCl₂, 10 mM glucose, and 10 mM Na-Hepes, pH 7.4, at 35 °C. During the experiments, 10 mM BDM was added unless indicated otherwise. Image acquisition (LSM 780 NLO laser scanning microscope with GASP detectors; Carl Zeiss) was performed with excitation at 488 nm (mtPA-GFP) and 568 nm (mtDsRed) using a 63x/1.4 NA ApoPlan objective. PA-GFP was photoactivated in a two-photon mode using a Chameleon pulsed laser system (760 nm; Coherent). To create 3D-like reconstructions, a sequence of images was collected starting below the adherent surface and moving upward in 0.367-μm steps (18 steps). Imaging of H9c2 cells was performed in 2D in the same fashion. Cells were treated before imaging with 60 mM KCl or 5 μM ionomycin, or left untreated (control). A series of 125 images was obtained from each cell, with one image recorded every 4 s.

For whole-heart imaging, rats were injected i.p. with 1,000 U of heparin 15 min before being anesthetized with isofluorane 3–5%. The heart was isolated and cannulated through the aorta with a 16-g needle, followed by immediate reverse perfusion using a Langerdoff system with 37 °C carbogen pre-equilibrated CaCl₂-free Joklik’s medium (Lonza) for 6 min. When needed, TMRE (100 nM) was added during perfusion with the Joklik’s medium. The heart was then quickly transferred and connected to perfusion pump-fed tubing. Before imaging, the heart was perfused for 10 min with CaCl₂-free 0.25% BSA ECM supplemented with 10 mM BDM and 100 μM EGTA. During imaging, which took <7 min, the perfusion was stopped to avoid movement artifacts. Perfusion was continued between imaging periods. Imaging was performed as described for isolated cells using a 100× water objective.

For simultaneous study of the Ca²⁺ oscillations and GFP spreading, HEK293T cells transfected with mtDsRed and mtPA-GFP as described previously were loaded with 1.5 μM Fluo8-AM (TEFLabs) for 20 min at 37 °C in 2.0% (wt/vol) BSA ECM in the presence of 0.003% pluronic acid and 100 μM sulfinpyrazone. Imaging was performed in 0.25% BSA ECM and 100 μM sulfinpyrazone at 35 °C. Fluo8 was excited at 488 nm, and mtDsRed and mtPA-GFP were excited and bleached, respectively, as described previously. Fluo8 fluorescence was masked in the nuclear area.

**Statistics.** Data are reported as mean ± SE. The significance of differences was calculated using Student’s t test or the Mann–Whitney rank-sum test.
Fig. S1. Mitochondrial morphology in neonatal and adult cardiac myocytes and 3D reconstruction of mitochondria in cultured myocytes. (A) Cultured NVCMs and in vivo-transformed freshly isolated AVCMs expressing mitochondrial targeted DsRed. The two cells show significant differences in morphology as well as in mitochondrial distribution, size, and density. (B) A 3D reconstruction of mtPA-GFP PA regions (5 × 5 μm) and surrounding areas of 24- to 48-h-old myocytes transduced in vitro (Left) or in vivo (Right) at 8 min after PA.
Fig. S2. Examples of mitochondrial fusion events. (A) Diverse orientation fusion events in freshly isolated in vivo-transformed AVCMs. (Upper) Images of time series showing sequential fusion events in both transverse and longitudinal orientations. (Lower) Quantitative evaluation of mtPA-GFP fluorescence transfer between numbered organelles. (B) Sequential mitochondrial fusion events in NCVMs. (Left) A 5 × 5-μm region of mtPA-GFP PA. (Center) Zoom-in view of the rectangle denoted by the white-dashed line showing the time course of mitochondrial fusion events between mitochondria 1 and 2, followed by matrix mixing between mitochondria 2 and 3. (Right) Quantification of mtPA-GFP fluorescence transfer from donor 1 to acceptor 2, and concomitant mixing of mtDsRed from 2 to 1 (the latter was photobleached when mtPA-GFP was photoactivated by 2P illumination). At 100 s after the first event, a new two-way mitochondrial mixing is evidenced between mitochondria 2 and 3. (C) Mitochondrial fusion event in an in vitro-transformed, 48-h–cultured AVCM. (Upper) The area of PA. (Lower) Time series of a fusion event between two neighbor mitochondria. Quantification of mtPA-GFP mixing is shown in the bar chart, highlighting the changes in the PA-GFP levels between donor and acceptor mitochondria before and after the fusion event.
Fig. S3. Diverse fusion events in AVCMs. (A) Diversity of newly formed mitochondrial communications by means of fusion events. A fusion event starts when an mtPA-GFP donor mitochondrion reaches a non-neighbor acceptor (1) through an apparent narrow connector. The kinetics of this event are very slow. Next, mitochondrion 1 partially equilibrates its mtPA-GFP matrix content through an additional fusion event with mitochondrion 2. Note that mitochondria 1 and 2 are also apparently connected through a narrow structure. Moreover, transversally oriented connectors become evident between mitochondrion 1 and a transversally oriented mitochondrion. Finally, at 288 s post-PA, a new fusion event occurs between mitochondria 1 and 2, fully equilibrating their contents. Thus, this example indicates that heart mitochondria can use narrow connectors to exchange their contents, and that two organelles can undergo partial and then complete fusion events, suggesting the partial opening of fusion pores or incomplete fusion events, followed by fission and then fusion again. (B) An emerging nanotunnel in a freshly isolated AVCM. Dynamic tubular mitochondria appear to emerge from globular organelles. (Upper Left) Two PA regions. (Upper Right) A region in which apparent nanotunnels emerge from globular mitochondria. For mitochondria movement activity, red indicates negative changes, and blue indicates positive changes. (Lower) Time-lapse images of elongated mitochondria (2 and 3) emerging and retracting from one large globular mitochondrion, as shown in Movie S8. In addition, elongated mitochondrion 1 moves around another globular mitochondrion.

Fig. S4. Pacing scheme applied in Fig. 5D. Freshly isolated in vivo-transformed ventricular myocytes plated on top of laminin-coated coverslips were mounted in a custom-made chamber equipped with platinum electrodes. At the confocal microscope stage, the chamber was connected to a Grass stimulator, and 2-Hz square-shaped pulses were triggered (30 V, 2 ms) for 5 min. The stimulation buffer was 0.25% BSA CaCl₂ without BDM. After brief and careful washing, and loading of the same experimental chamber with 10 mM BDM-supplemented 0.25% BSA ECM, fusion events were evaluated for 5 min. Mock experiments involved the same treatment of the cells without field stimulation.
Mitochondrial fusion protein Mfn1 decays in cultured AVCMs and rescues fusion upon exogenous expression. (A) Freshly isolated AVCMs were plated on culture dishes for 2–8 h or kept in culture for at least 24–30 h. Samples were snap-frozen, and membrane extract was prepared for Western blot analysis and tested with the indicated antibodies. Specific bands were quantified with regard to the <8-h condition. n = 4 independent experiments. P < 0.05. (B) Mfn1 KO cells were transduced with AdMfn1, and at 24–28 h after infection, the cells were loaded with MitoTracker Red and evaluated by confocal microscopy (Left). Cell membrane extracts were evaluated by Western blot analysis to determine the abundance of Mfn1 in MEF WT, Mfn1 KO, and Mfn1 KO cells transduced with AdMfn1 (Right). Transformation of Mfn1 KO cells with AdMfn1 rescued the expression of Mfn1. (C) AdmptPA-GFP and AdmDsRed in vivo-infected AVCMs were infected in vitro with either control AdLacZ or AdMfn1 virus and then evaluated at 24–48 h after plating. (Left) Representative images of mtPA-GFP spreading at 300 s post-PA shows increased mitochondrial continuity for AdMfn1 transformed cells compared with control, as indicated by white arrowheads. (Right) Quantification of mitochondrial fusion for the LacZ- and Mfn1-infected AVCMs shows an increase in mitochondrial fusion for Mfn1 cells (n = 39) compared with LacZ cells (n = 40). **P < 0.01. (D) WT and RyR2-HEK293T cells noninduced or induced for 48 h with tetracycline (Tet) were used to prepare membrane-enriched extracts, which were evaluated on 8% PAGE and tested for Mfn1 protein levels. n = 2 independent experiments.
Fig. S6. Abundance of mitochondrial fusion proteins in control and EtOH-fed cardiac mitochondria. Shown is a representative Western blot of mitochondrial fusion proteins tested in rat heart mitochondrial extracts from control and EtOH-fed animals. Mfn1 and Mfn2 (n = 5 pair) and Opa1 (n = 2 pair) were tested.
Fig. S7. Higher prevalence of abnormal IMM folding with absent and/or irregular cristae in the cardiac mitochondria of the EtOH-fed rats. (A) Mitochondrial cristae morphology analysis in TEM images of longitudinally sectioned left ventricular wall samples. To quantitate potential alterations, a five-grade scoring system for cristae abundance and form in individual mitochondria was introduced as illustrated. (Inset) Example images for each grade and definition of the grades. (B and C) Bar graphs showing the cumulated overall mean scores (B) and the prevalence of the individual score grades across the analyzed mitochondrial population for each pair of animals (C). For one pair in which membrane contrast was exceptionally good, the analysis was performed by comparing a total of 179 mitochondria in five fields of 11 × 11 μm (4,400× direct magnification). For the other two pairs, higher magnification was needed. The number of mitochondria compared was 42 controls (32 2.3 × 2.3 μm and 3.2 × 3.2 μm fields at 21,000× and 15,000×) vs. 68 EtOH (23 3.2 × 3.2-μm fields) and 36 controls (24 2.3 × 2.3-μm fields) vs. 73 EtOH (32 fields ranging between 4.5 × 4.5 μm and 2.3 × 2.3 μm at 11,000–21,000× direct magnification).
Movie S1. A subregion of an NVCM expressing mDsRed and mtPA-GFP. On photoactivation of mtPA-GFP by means of 2P within a 5 × 5 μm region, mitochondrial continuity, fusion, and motility are evidenced by mtPA-GFP spreading, newly formed mitochondrial communications, and displacement, respectively. This 7-min video (frequency acquisition, 0.25 Hz, at 7 fps) shows highly active mitochondrial dynamics.

Movie S1

Movie S2. Mitochondrial dynamics in 24–48 h in vitro-transformed AVCMs expressing mtPA-GFP. mtPA-GFP was photoactivated in 2 5 × 5-μm regions, and the fluorescence spreading and newly formed communications among mitochondria were followed for 8.25 min. Frequency acquisition, 0.33 Hz, at 7 fps.

Movie S2
**Movie S3.** Mitochondrial dynamics in 24–48 h in vivo-transformed AVCMs expressing mtPA-GFP. mtPA-GFP was photoactivated in 3 $\times$ 5-$\mu$m regions, and the fluorescence spreading and newly formed communications among mitochondria were followed for 8.25 min. Frequency acquisition, 0.33 Hz, shown at 7 fps.

**Movie S4.** Mitochondrial dynamics in a freshly isolated in vivo-transformed AVCMs expressing mtDsRed and mtPA-GFP. mtPA-GFP was photoactivated in 5 $\times$ 5-$\mu$m regions, and the fluorescence spreading and newly formed communications among mitochondria were followed for 7 min. Frequency acquisition, 0.25 Hz, shown at 7 fps.
Movie S5. A subregion of a freshly isolated in vivo-transformed AVCM expressing mtDsRed and mtPA-GFP. Photoactivation of mtPA-GFP by means of 2P illumination within a 5 × 5-μm region. Mitochondrial continuity and fusion are evidenced by continuous and abrupt mtPA-GFP spreading, respectively. Video duration, 8.25 min; acquisition frequency, 0.33 Hz, shown at 7 fps.

Movie S6. Sequential mitochondrial fusion events in a freshly isolated in vivo-transformed AVCM. mtPA-GFP mixing in transverse and longitudinal orientations indicates mitochondrial fusion events in a region contiguous with a PA area. Video duration, 8.25 min; acquisition frequency, 0.33 Hz, shown at 7 fps.
Movie S7. Fast and slow kinetics of AVCM fusion events. Parallel videos showing representative examples described in Fig. 3, highlighting fast and slow fusion mixing kinetics. Each video lasts 300 s; acquisition frequency, 0.33 Hz, at 7 fps.

Movie S7

Movie S8. Diverse mitochondrial fusion events in a freshly isolated AVCM. Video corresponds to Fig. S3. Duration, 8.25 min; acquisition frequency, 0.33 Hz, at 7 fps.

Movie S8
Movie S9. Motility and emerging nanotunnel in a freshly isolated AVCM. Video corresponds to Fig. S4, showing the motility of elongated mitochondria followed by a longitudinal fusion event. Duration, 8.25 min; acquisition frequency, 0.33 Hz, at 7 fps.

Movie S9