Abstract

Gap junctions play a pivotal role for the velocity and the safety of impulse propagation in cardiac tissue. Under physiologic conditions, the specific subcellular distribution of gap junctions together with the tight packaging of the rod-shaped cardiomyocytes underlies anisotropic conduction, which is continuous at the macroscopic scale. However, when breaking down the three-dimensional network of cells into linear single cell chains, gap junctions can be shown to limit axial current flow and to induce ‘saltatory’ conduction at unchanged overall conduction velocities. In two- and three-dimensional tissue, these discontinuities disappear due to lateral averaging of depolarizing current flow at the activation wavefront. During gap junctional uncoupling, discontinuities reappear and are accompanied by slowed and meandering conduction. Critical gap junctional uncoupling reduces conduction velocities to a much larger extent than does a reduction of excitability, which suggests that the safety for conduction is higher at any given conduction velocity for gap junctional uncoupling. In uniformly structured tissue, gap junctional uncoupling is accompanied by a parallel decrease in conduction velocity. However, this is not necessarily the case for non-uniform structures like tissue expansion where partial uncoupling paradoxically increases conduction velocity and has the capacity to remove unidirectional conduction blocks. Whereas the impact of gap junctions on impulse conduction is generally assessed from the point of view of cell coupling among cardiomyocytes, it is possible that other cell types within the myocardium might be coupled to cardiomyocytes as well. In this context, it has been shown that fibroblasts establish successful conduction between sheets of cardiomyocytes over distances as long as 300 μm. This might not only explain electrical synchronization of heart transplants but might be of importance for cardiac diseases involving fibrosis. Finally, the intriguing fact that sodium channels are clustered at the intercalated disc recently rekindled the provocative question of whether gap junctions alone are responsible for impulse propagation or whether electric field mechanisms might account for conduction as well. Whereas computer simulations show the feasibility of conduction in the absence of gap junctional coupling, a definite answer to this question must await further investigations into the biophysical properties of the intercalated disc.

Keywords: Heart; Impulse propagation; Gap junction; Tissue structure; Sodium channels

1. Introduction

Half a century ago, Weidmann [1] published his pioneering electrophysiological study which showed that cardiac Purkinje fibers behaved electrically similarly to axons. Two years after this work, however, electron microscopic observations of cardiac tissue led Sjöstrand and Andersson [2] to conclude that myocardial tissue is not, as was previously thought, an anatomical syncytium but that each cardiomyocyte is surrounded by a contiguous cell membrane. Based on both electrophysiological studies of the cable properties of Purkinje fibers and the observation that radioactive potassium readily spread along the tissue, this apparent contradiction between electrophysiological findings and cellular ultrastructure was resolved by postulating the presence of low-resistance pathways interconnecting the cardiomyocytes [3,4]. These findings and observations in other tissues led to the establishment of the concept of ‘gap junctions’ as the fundamental principle underlying impulse transmission and signaling molecule exchange among adjacent cells in a variety of tissues. It is the goal of this short review to summarize recent findings related to the role of gap junctions in the process of the spread of cardiac excitation on the cellular level based on both experimental studies and computer simulations and to briefly touch the controversial issue of whether gap junctions are the ‘truth and nothing but the truth’ for electrical impulse transmission between cardiomyocytes. For a review of the important effects of gap junctional remodeling and
genetic manipulations of connexin expression on impulse propagation in the heart, the reader is referred to the respective chapters of this spotlight issue.

2. General aspects of the role of gap junctions in propagation

The working myocardium can be regarded as a three-dimensional network of coupled excitable elements. In this network, the velocity and safety of the spread of excitation is dependent on both active and passive properties of the individual elements and on the connectivity of the network. Among passive properties, gap junctions play a pivotal role because they ultimately determine how much depolarizing current passes from excited to non-excited regions of the network. Thereby, they act as important determinants of the speed and safety of this process. A second important factor defining the passive properties of the network consists of the specific cellular architecture of cardiac tissue: whereas progressive uncoupling in a uniform network will result in a slowing of conduction velocity following the square root relationship, this does not necessarily have to be the case for non-uniform networks, i.e., for tissue structures where the size of a given excited region supplying depolarizing current (‘current source’) is ill matched to the amount of depolarizing current necessary to excite the regions ahead (‘current sink’). A long known example for such a source-to-sink mismatch is represented by the Purkinje–fiber–ventricular junction where a small source (Purkinje fiber) is coupled to a large sink (mass of ventricular tissue)[5–7]. Depending on the size of the mismatch, this results in either local conduction delays or unidirectional conduction blocks at the junction during anterograde conduction (Purkinje–fiber–ventricle). In contrast to the effects of gap junctional coupling on propagation in uniform tissue, partial uncoupling in such non-uniform tissue structures can actually be accompanied by a paradoxical increase in safety and velocity of conduction. Because of these marked differences of the effect of gap junctional coupling on conduction in uniform vs. non-uniform tissue, the two issues are discussed separately.

3. Gap junctions and impulse propagation in uniform tissue during normal coupling

Under physiological conditions, a given cardiomyocyte in the adult working myocardium is electrically coupled to an average of ~11 adjacent cells with gap junctions being predominantly localized at the intercalated discs at the ends of the rod shaped cells [8]. This particular subcellular distribution of gap junctions is a main determinant of anisotropic conduction in the heart even though, as shown recently, cell size on its own also significantly modifies the characteristics of transverse conduction [9] (for review, cf. Ref. [10]). It is generally accepted that macroscopic impulse conduction along the main fiber axis of cardiac tissue is continuous because of the low resistance coupling by gap junctions. However, when abstracting from the three-dimensional network to a chain of single cardiomyocytes, the results of the computer simulations shown in Fig. 1A suggest that propagation is actually saltatory and is composed of rapid excitation of individual cells followed by a transjunctional conduction delay [11,12]. When both gap junctional resistance and the resistivity of the myoplasm are set to levels resulting in macroscopic conduction velocities of ~50 cm/s, both conduction times along the entire cell soma and transjunctional propagation delays are ~100 μs. Whereas, at first glance, this difference seems to be moderate, one has to keep in mind that the gap junctional cleft is roughly 4 orders of magnitude shorter than the cell itself resulting in apparent local ‘conduction velocities’ of ~1 m/s along the cell and 0.1 mm/s across the junction. As shown in Fig. 1B, this saltatory type of conduction becomes highly accentuated during a tenfold reduction of gap junctional conductance (macroscopic conduction velocities of ~20 cm/s). In this case, cell somata are excited in a virtually simultaneous manner due to the cellular confinement of depolarizing

![Fig. 1. Computer simulation of microscopic impulse propagation. (A) Action potential upstrokes recorded during impulse propagation between two adjacent cells at the sites indicated in the insert. Under conditions of normal gap junctional coupling, the transjunctional conduction delay is roughly equal to the myoplasmic conduction times of the cells. (B) During a tenfold reduction of gap junctional coupling, myoplasmic conduction times are abbreviated whereas transjunctional conduction times are substantially increased. Redrawn with modifications from Ref. [12].](image-url)
current, whereas transjunctional conduction delays substantially increase to 500 μs. Experimentally, the question of microscopically discontinuous conduction in cardiac tissue has been addressed by combining patterned growth of cultured cardiomyocytes [13,14] and optical recording of transmembrane voltage with subcellular resolution [15]. As shown in Fig. 2, simultaneous optical recordings of action potential upstrokes in a single cell chain of cultured neonatal rat ventricular cardiomyocytes with subcellular resolution (15 μm) indicated the presence of an activation delay at the contact site between two end-to-end abutted cardiomyocytes, whereas conduction within individual cells was rapid and continuous [16]. Quantitative aspects of this phenomenon are shown in Fig. 3. In the case of single cell chains of cardiomyocytes (panel A), the difference between mean cytoplasmic (38 μs) and transjunctional (118 μs) conduction time as measured with detectors spaced 30 μm apart indicated the presence of a transjunctional activation delay of 80 μs [17]. Assuming an average cell length of 60 μm, transjunctional delays and cytoplasmic conduction times were of roughly equal magnitude, thus being in close agreement with the aforementioned computer simulation study. In contrast to the findings in single cell chains, the characteristic of conduction at the microscopic scale change drastically in strands several cells wide (Fig. 3B). There, mean cytoplasmic conduction times increased to 57 μs, whereas transjunc- tional conduction times simultaneously decreased to 89 μs, resulting in significantly reduced cell-to-cell propagation delays. This suggests that, in uniform multicellular tissue, conduction delays across longitudinally abutted cells largely disappear at the expense of increased cytoplasmic conduc-

tion times. This can be explained by the presence of lateral gap junctional coupling, which serves to average slight local advances/delays of the activation wavefront among laterally apposed cells. This lateral averaging, which can be expected to be even more pronounced in three-dimensional tissue, ultimately results in largely continuous conduction along the main fiber axis in intact uniform tissue.

4. Gap junctions and impulse propagation in uniform tissue during progressive uncoupling

Whereas, as noted above, propagation in uniform multicellular tissue under normal conditions is continuous, this changes during progressive uncoupling. As shown in Fig. 4, severe gap junctional uncoupling in a 250 μm wide cell strand not only drastically reduced conduction velocity from 36.7 to 0.3 cm/s, but further resulted in a meandering activation wavefront [18]. Meandering is induced by the presence of islands of completely uncoupled cells which causes the activation wavefront to follow a ‘zig-zag’ path of activation similar to that occurring in infarcted [19] and non-uniform anisotropic tissue [20]. Also at the cellular level, impulse propagation under these conditions is highly discontinuous as illustrated in Fig. 5. Whereas conduction is fast (43 cm/s) and continuous under control conditions (Fig. 5A), severe uncoupling (Fig. 5B) causes action potential upstrokes to rise in clusters with large inter-cluster activation delays which result in very slow overall conduction velocities of 1.1 cm/s (Fig. 5Bb). Moreover, when mapping the clustered action potential upstrokes to the cellular structure of the preparation (Fig. 5Bc), it becomes obvious that activation during severe uncoupling jumps from one region consisting of one or a few cardiomyocytes to the next and that completely uncoupled cells force the activation wavefront to meander. Thus, irrespective of scaling, meandering activation seems to be a basic principle governing impulse propagation during reduced gap junctional coupling in cardiac tissue.

When comparing minimal conduction velocities achieved by either a reduction of active membrane properties (~ 20 cm/s) [21–23] or by a reduction of gap junctional coupling (~ 1 cm/s; cf. data above), the fact that conduction velocities can be an order of magnitude slower during uncoupling suggests that this is a ‘safer way’ to reduce velocity than a decrease in excitability. The reason for this behavior has been analyzed in detail in computer simulation studies by Shaw and Rudy [12,24], which compared the characteristics of conduction slowing during a reduction of either excitability or gap junctional coupling. In order to analyze the robustness of conduction under these conditions, they defined the ‘safety factor’ for conduction as the ratio of the amount of charge produced by a given membrane patch during activation to the charge consumed during the activation process. By this definition, conduction fails when the safety factor drops below 1 and becomes increasingly stable as it rises.
Fig. 3. Differences in intra- and intercellular activation delays between single cell chains and strands several cells wide. (A) Single cell chains: (Aa) Schematic drawing of the cellular architecture of a chain of single cardiomyocytes. Three equally spaced photodetectors (black discs) were placed such that they recorded activation (corresponding to a fractional change in fluorescence of the voltage sensitive dye, $\Delta F/F$) either within cells (1→2; ‘cytoplasm’) or between cells (2→3; ‘junction’). (Ab) Optically recorded action potential upstrokes (above) and their derivatives (below) of a measurement during propagation from right to left. (Ac) Histograms of interdetector conduction times indicate coexistence of fast cytoplasmic and delayed transjunctional activation. (B) Multiple cell wide strands: (panels as in (A)). In the case of strands several cells wide, differences between cytoplasmic and transjunctional conduction times are substantially reduced indicating nearly continuous conduction along the preparation. Redrawn with modifications from Ref. [17].
above 1. As shown in Fig. 6A, a reduction of excitability, which was modeled by reducing sodium channel conductance, was accompanied by a decrease of conduction velocity from 54 to 17 cm/s before conduction failed. At the same time, the safety factor for conduction fell from $f_1.6$ to 1. As expected for a decrease in sodium channel conductance, the reduction of conduction velocity was paralleled by a tenfold decrease of maximal upstroke velocities (Fig. 6B). In contrast to these effects, a reduction of intercellular coupling showed marked differences (Fig. 7). First, whereas progressive uncoupling also induced a monotonic decrease of slowest conduction velocities reached before block (0.26 cm/s) were over an order of magnitude below those obtained during a reduction of excitability. Moreover, both maximal upstroke velocities and the safety factor for conduction showed a biphasic behavior with a substantial initial increase during progressive uncoupling. For both parameters, maxima close to twice control values were reached at conduction velocities in the few centimeters range corresponding to a $15$-fold reduced level of gap junctional coupling. This transient increase of maximal upstroke velocities and of the safety factor of conduc-
tion above control values is due to the fact that, with decreasing gap junctional conductance, the sodium inward current is increasingly confined to individual cells because less current is lost downstream. On the other hand, the decrease in the safety factor for conduction and maximal upstroke velocity at very low levels of gap junctional coupling is caused by the highly reduced axial current flow downstream, which causes long subthreshold charging of the
cells ahead and, concomitantly, a progressive inactivation of sodium channels.

In conclusion, whereas impulse propagation under physiologic conditions along single cell chains of cardiomyocytes is saltatory due to the recurrent increases in axial resistance at the sites of gap junctional coupling, this feature is lost in intact multicellular tissue due to lateral gap junctional coupling which serves to average local small differences in activation times of individual cardiomyocytes at the excitation wavefront. In multicellular tissue, saltatory conduction only reappears under conditions of critical gap junctional uncoupling. There it leads to a functional unmasking of the cellular structure and induces ultra-slow and meandering conduction, which is well known to be a key ingredient in arrhythmogenesis. In both experiments and computer simulations, partial gap junctional uncoupling was shown to result in conduction velocities which are over an order of magnitude slower than those obtained during a maximal reduction of excitability. The only feature of the characteristics of impulse propagation during severe uncoupling in computer simulation studies not reproduced routinely by experiments so far (for exception, cf. Ref. [25]) concerns the transient increase in maximal upstroke velocity. This is most probably due to the lack of specific uncoupling agents available, because increases in maximal upstroke velocities accompanying a reduction in conduction velocity have been found in mice with connexin43 null mutations [26].

Fig. 5. Effects of critical gap junctional uncoupling on impulse propagation characteristics at the cellular level. (A) Normal coupling: (Aa) Phase contrast picture of a 55-μm wide cell strand with white circles indicating the positions of individual photodetectors. (Ab) Evenly spaced isochrones of activation indicate fast and continuous activation at an overall velocity of 43 cm/s. (B) Critical gap junctional uncoupling: (Ba) Phase contrast picture of a 55-μm wide preparation with rings indicating the positions of photodetectors. (Bb) During propagation from left to right under conditions of severe gap junctional uncoupling, action potential upstrokes rise in clusters with large inter-cluster activation delays indicating a highly discontinuous type of conduction (average conduction velocity=1.1 cm/s). (Bc) Schematic representation of the path of activation and its dependence on the cellular tissue architecture. Regions showing quasi-simultaneous activation are numbered according to the cluster numbers indicated in (Bb). The hatched region indicates a completely uncoupled cell showing no electrical activation. Redrawn with modifications from Ref. [18].

Fig. 6. Computer simulation of the effects of a gradual reduction of membrane excitability on the characteristics of impulse propagation. (A) Dependence of conduction velocity and of the safety factor for conduction on membrane excitability. (B) Dependence of maximal upstroke velocity and maximal sodium inward current on membrane excitability. Redrawn with modifications from Ref. [12].

Fig. 7. Computer simulation of the effects of a gradual reduction of gap junctional coupling on the characteristics of impulse propagation. (A) Dependence of conduction velocity and of the safety factor for conduction on the degree of intercellular coupling. (B) Dependence of maximal upstroke velocity and maximal sodium inward current on the degree of intercellular coupling. Redrawn with modifications from Ref. [12].
5. Gap junctions and impulse propagation in non-uniform tissue

As pointed out above, the general principle of ‘less gap junctional coupling ≈ impaired conduction’ as found in uniform tissue architectures does not necessarily apply to impulse propagation along non-uniform structures as represented, e.g., by an abrupt tissue expansion. Such tissue structures induce propagation delays in anterograde direction due to the presence of a source-to-sink mismatch. If the mismatch is large enough, unidirectional conduction blocks ensue. Computer simulations of the characteristics of prop-

Fig. 8. Induction of successful bi-directional conduction across a tissue expansion by partial gap junctional uncoupling. (A) Schematic drawing of the experimental layout. A tissue expansion was locally superfused with the gap junctional uncoupler palmitoleic acid and impulse propagation characteristics across the expansion were assessed optically by detectors (black rings) along the central axis during anterograde (left) and retrograde (right) conduction. (B) Action potential upstrokes recorded under control conditions indicate decremental conduction during anterograde and normal conduction during retrograde stimulation, i.e., unidirectional conduction block. (C) Complete gap junctional uncoupling results in bi-directional conduction blocks at the boundaries of the superfusion. (D) During gradual re-coupling, successful bi-directional conduction is established. (E) At the end of the washout period, normal gap junctional conductance is reestablished as indicated by fast retrograde conduction and anterograde conduction block. Redrawn with modifications from Ref. [29].
The widths of the strands were chosen such that unidirectional conduction block occurred under control conditions (Fig. 8B). Initially, complete gap junctional uncoupling of the strand and the expansion was induced by local superfusion with palmitoleic acid (Fig. 8C). Thereafter, washout was started which resulted in a gradual increase of gap junctional coupling during which successful bi-directional conduction could be observed within a limited time window (Fig. 8D). After reaching normal levels of gap junctional coupling, unidirectional conduction blocks were reestablished (Fig. 8E). This illustrates that partial gap junctional uncoupling has the potential to remove unidirectional conduction blocks. This counter-intuitive behavior can be explained by differences in the dimensionality of the effect of gap junctional uncoupling on the source (strand) and the sink (expansion). Partial uncoupling of the essentially linear source will reduce its size in a linear proportional manner whereas equal uncoupling of the two-dimensional sink will reduce the size thereof following a square function. This overproportional reduction of the sink improves the source-to-sink mismatch up to the point of successful anterograde conduction. As can be expected for a gradual change in the balance between the source and the sink, overall conduction velocity in anterograde direction across a tissue expansion changes in a biphasic manner during progressive re-coupling. This is shown in Fig. 9 where the gradual increase in gap junctional conductance is initially accompanied by an increase and then by a paradoxical decrease in overall anterograde conduction velocity. How can this be explained? At the beginning of the establishment of successful anterograde conduction, conduction velocities are initially in the range of those observed in linear cell strands (cf. above). This suggests that conduction at these very low levels of gap junctional conductance is primarily determined by the degree of cell-to-cell coupling and not by the source-to-sink mismatch represented by the expansion. Accordingly, increases of gap junctional conductance led initially to an increase in conduction velocity. At the same time, however, the parallel increase in the size of the sink became more important and, ultimately, resulted in the paradoxical situation that overall conduction velocity decreased even though gap junctional conductance increased. Therefore, in contrast to the situation in uniform cell structures where a decrease in gap junctional coupling is invariably accompanied by a decrease of conduction velocity, overall conduction velocity across sites where planar wavefronts change to curved wavefronts (expansion, isthmus), initially show an increase which is followed by a decrease only during severe uncoupling.

Thus, whereas gap junctional uncoupling in general leads to an impairment of conduction and thereby contributes significantly to the generation of arrhythmias, there is another side to this coin for the case of discontinuous tissue architectures. In such structures, which are likely to be present in the border zone of healed infarcts [30] or in fibrotic tissue of the aged and/or hypertrophied myocardium [31], partial gap junctional uncoupling might actually remove unidirectional conduction blocks and therefore eliminate one of the key ingredients of reentrant arrhythmias. Accordingly, while improvement of gap junctional coupling in structurally diseased heart might be expected to act in an anti-arrhythmogenic manner by reducing the incidence of slowly conducting pathways, it might at the same time provoke arrhythmias by unmasking potential regions of unidirectional conduction blocks.

6. Gap junctional gating: is there an effect on impulse conduction?

One of the prominent biophysical features of gap junctions is their time- and voltage-dependent inactivation. Even though the time constants for inactivation (~150 ms at a transjunctional voltage gradient of 100 mV) [32] are extremely slow compared to transjunctional activation delays encountered during normal action potential propagation (~30 ms for multicellular strands, cf. above), recent computer simulations suggested that, under con-
ditions of severe uncoupling, junctional conduction times slightly increase when using a dynamic model of gap junctions and that conduction blocks occur at lower levels of resting transjunctional conductance than those found using a static model [33]. In another recent study using transfected neuroblastoma cells, inactivation kinetics of connexin43 were studied by imposing an action potential clamp instead of a rectangular voltage pulses on one of the cells [34]. These experiments showed that, following the peak of the action potential, junctional conductance decreased within 25 ms to 58% of control. Although these kinetics are faster than those reported earlier, the comparison of these inactivation times to transjunctional conduction times observed during steady state propagation under conditions of severe uncoupling (<5 ms) [18] suggests that gap junctional gating has only a small effect on overall conduction velocities. In addition to the effects of gap junctional gating on depolarizing current flow in the orthodromic direction, it would be interesting to know whether gating possibly affects the trailing part of activation, i.e., the repolarization. While it is tempting to speculate that partial inactivation might last into the repolarization phase, thereby channeling depolarizing current from the activation wavefront in the orthodromic direction, this mechanism is unlikely to exist because gap junctional conductance will rapidly recover during repolarization due to the reversal of polarity [35] and the decrease in size of the transjunctional voltage [34].

Thus, whereas gating possibly influences the degree of ultra-slow conduction to some extent during severe uncoupling, the role thereof during normal propagation is likely to be insignificant. However, because gating is dependent on a variety of factors such as the expression system used, transjunctional voltages present, and the type of gap junctions present (isoforms, connexon composition (homomeric, heteromeric) and connexon coupling (homotypic, heterotypic)) definitive answers to this question will require further studies, which take into account transjunctional voltage differences as they occur across a given junction during propagated activity.

7. Gap junctional coupling between cardiomyocytes and non-excitable cells

In the heart, approximately half of the cells consist of non-myocardial cells, among which fibroblasts constitute the largest fraction [36]. This number can be expected to increase as a result of cardiac diseases leading to fibrosis. Whereas the formation of excessive collagen sheets, which act as electrical insulators, has been recognized for a long time as being a cause for discontinuous conduction and the occurrence of arrhythmias [37], the question of whether the cellular constituents of fibrotic tissue, i.e., the fibroblasts, might affect impulse propagation directly by forming gap junctions with cardiomyocytes has been addressed only recently. It has been known for several decades that individual fibroblasts of cardiac origin can establish gap junctional communication with adjacent cardiomyocytes in culture [38,39]. In this context, it was shown that impulse propagation in monolayer cardiomyocyte cultures can be modified by grafting a layer of fibroblasts transfected with the voltage gated potassium channel Kv1.3 over them [40]. This co-culture resulted in conduction blocks in the cardiomyocyte monolayer, which were reversed upon application of specific blockers of the potassium channel. Moreover, it was shown in cell culture that fibroblasts adjacent to cardiomyocytes induce a decrease in maximal upstroke velocity [41] or a local slowing of conduction [42]. Whereas these findings illustrate that fibroblasts in intimate contact with cardiomyocytes can influence the electrophysiological behavior of the latter via gap junctions, the question arises whether such interactions might also occur over longer distances, i.e., whether fibroblasts are capable of relaying electrical activation between disparate sheets of cardiomyocytes. This issue was recently investigated with a cell culture model where patterned growth strands of cardiomyocytes were interrupted over defined distances by fibroblasts of cardiac origin [43]. The results of one of these experiments are shown in Fig. 10. In this particular preparation, the fibroblast insert had a length of 134 μm and propagating action potentials were elicited at 2 Hz on the left. As indicated by the optically recorded transmembrane voltage signals during propagated activity in Fig. 10C, action potential upstrokes were monotonically rising in the region of the cardiomyocytes, whereas they showed ‘double-humps’ in the region of the fibroblasts due to bidirectional electrotonic interaction with the proximal and the delayed activated distal cardiomyocyte strand. As shown by immunocytochemistry, this electrotonic interaction was based on the presence of both connexin43 and connexin45. As indicated by the plot of activation times along the preparation (Fig. 10D), the fibroblast insert induced a local propagation delay of 30 ms. This delay became as long as 68 ms for the longest inserts supporting impulse transmission (∼300 μm) resulting in apparent local ‘conduction velocities’ as low as 2.2 mm/s. Whereas it is tempting to speculate that such extremely slow conduction might be instrumental in the generation of arrhythmias in fibrotic hearts, studies with cell cultures have to be interpreted cautiously in regard to extrapolation of the results to intact tissue because there is as yet no firm proof of gap junctional coupling between cardiomyocytes and fibroblasts in-vivo. In contrast, a thorough investigation of this question found no evidence for robust gap junctional coupling in healthy intact tissue [44]. This raises the question whether fibroblasts in culture might undergo a phenotype switch to so-called myofibroblasts, which enables them to form gap junctions with cardiomyocytes. The conversion from fibroblasts into myofibroblasts, which are characterized by the expression of α-smooth muscle actin [45], has been described for
different types of fibroblasts in culture. In intact hearts, it was shown that fibroblasts convert into myofibroblasts after a local loss of cardiomyocytes [46]. Most interestingly, this conversion has been described to be accompanied by the expression of connexin43 in the case of breast cancer stroma cells [47] and myofibroblasts derived from corneal fibroblasts [48]. If such a conversion of fibroblasts into connexin expressing myofibroblasts should also occur in the heart under pathophysiologic conditions such as myocardial infarction [36], this would raise the interesting hypothesis that the ensuing coupling of non-excitable cells to cardiomyocytes might lead to very slow conduction and, thus, might constitute a possibly important new arrhythmogenic mechanism.

In conclusion, the observation that cardiomyocytes readily form functional gap junctions with heterogeneous cells and that this coupling supports the spread of excitation over extended distances may have implications beyond electrical interactions with fibroblasts as presented above. In particular, given the recent interest in using stem cells as a therapeutic approach for the diseased heart, the promiscuous gap junctional coupling is a prerequisite both for permitting orderly excitation sequences in the regions of grafted cells and for the intercellular exchange of signaling molecules.

8. Gap junctions, the “truth and nothing but the truth” for impulse propagation?

Whereas all of the above evidence stresses the importance of gap junctional coupling for impulse propagation under both physiologic and pathophysiologic conditions, the observation that the main ion channels underlying fast conduction, i.e., the sodium channels, are clustered at intercalated discs [49–53] is intriguing. If one were to design a cardiomyocyte, one would probably not plan to insert sodium channels at the intercalated disc because, among other reasons, they would face a highly restricted extracellular space which could be expected to be subject to large fluctuations in ion concentration and, consequently, in adverse changes in electrochemical driving forces. On the other hand and as formulated many years ago by Sperelakis and Mann [54], the fact that space is restricted at the intercalated disc could also act in favor of impulse propagation. These authors postulated an electric field mechanism of impulse transfer, which is based on the idea that activation of sodium channels at the intercalated disc could also act in favor of impulse propagation. These authors postulated an electric field mechanism of impulse transfer, which is based on the idea that activation of sodium channels at the intercalated disc results in a negative shift of the cleft potential between a given activated cardiomyocyte and a neighboring quiescent cardiomyocyte. This negative shift of the cleft potential reduces the transmembrane potential ‘seen’ by the sodium channels of the post-
junctonal membrane and, hence, brings this region to threshold with subsequent rapid activation of the entire cell [55]. Sperelakis and McConnell [51] further suggested that this mechanism might be supported by rapid potassium accumulation in the cleft during activation of the pre-junctional cell which would induce a depolarization of the post-junctional membrane cell to threshold. Obviously, this electric field mechanism is critically dependent on the radial shunt resistance of the intercellular cleft which has (i) to assume a value high enough to permit the build-up of a local extracellular negativity in the cleft region and (ii) which, at the same time, must permit establishment of a local circuit current large enough as to depolarize the quiescent cell. Under the assumptions of appropriate parameters, Sperelakis and McConnell showed in computer simulations that their model consisting of linear chains of single cardiomyocytes permits successful impulse propagation with conduction velocities up to ~ 40 cm/s (for review, cf. Ref. [51]).

Recently, the interplay between the effects of sodium channel clustering, cleft potentials and gap junctional coupling on impulse propagation have been further investigated by using a linear strand of cardiomyocytes represented by the Luo–Rudy model [52]. The effects of different combinations of cleft widths and degrees of sodium channel partitioning on conduction velocities achieved during progressive uncoupling are shown in Fig. 11. Under the assumption of a cleft width of 35 nm and sodium channels being present exclusively at the intercalated disc, these simulations showed that conduction velocities were substantially slower than those obtained with a non-cleft model during normal gap junctional coupling. This was explained by the occurrence of large negative cleft-potentials during activation, which induced a rapid and largely overshotting response of pre- and post-junctional membranes and, therefore, resulted in an attenuation of the sodium inward current and a concomitant slowing of conduction. With decreasing gap junctional coupling, the difference in conduction velocities between standard model and cleft-model became smaller until, at coupling levels <20%, the cleft-model performed increasingly better and supported conduction velocities of 15 cm/s even during a 10,000-fold reduction of gap junctional coupling. Whereas this finding of conduction in the virtual absence of gap junctional coupling points in the same direction as earlier findings by Sperelakis and colleagues, Fig. 11 also illustrates that only moderate changes in the parameters (cleft width 35→10 nm, sodium channel partitioning 100%→50%) resulted in a behavior which was only slightly different from the standard model. Whereas these studies show the possibility of conduction in the absence of gap junctional coupling, it is not clear whether these findings are relevant for intact tissue. Whereas clustering of sodium channels at the intercalated discs is undisputed for both intact cardiac tissue and cell cultures, there are no experimental data available regarding actual radial cleft resistances in cardiac tissue. Because a direct determination of the radial cleft resistance is out of reach of present experimental methods, an exact morphometric assessment of the three-dimensional architecture of the cleft between two adjoining cardiomyocytes might indirectly permit one to obtain an approximate estimate thereof. However, even if such studies should reinforce the computer simulation studies, there remains a number of open questions which are difficult to reconcile with the concept that electric field mechanisms alone are significantly involved in cardiac impulse conduction: (i) in the absence of substantial gap junctional coupling, space constants should be much shorter than what was generally reported in the past (for a critical discussion of this issue, cf. Ref. [51]); (ii) the electric field model cannot readily explain calcium inward current based conduction which occurs even though these ion channels are not clustered at the intercalated disc; (iii) it is difficult to understand how the experimentally firmly established effects of source-to-sink mismatches on impulse propagation could be explained by an electric field model in the absence of gap junctional coupling.

In conclusion, at least for now, it seems safe to conclude that gap junctions are “the truth and nothing but the truth” for impulse propagation in cardiac tissue. Nevertheless, the clustering of sodium channels at the intercalated disc together with the presence of Na–K-ATPases at the same location [56] remains a highly intriguing and interesting fact. It indicates that this part of the sarcolemma, even though facing a highly restricted extracellular space, might exhibit a specific function. However, answers as to the exact nature and relevance of this function will have to await further studies.

![Fig. 11. Dependence of conduction velocity on gap junctional coupling for 3 different models of cleft configuration and partitioning of sodium channels. Solid line: model with no cleft-effects but clustering of all sodium channels at intercalated disc. Dashed line: model with 10-nm wide cleft and even partitioning of the sodium channels between the intercalated disc and the surface sarcolemma. Punctate line: model with a 35-nm wide cleft and clustering of all sodium channels at intercalated disc. Redrawn with modifications from Ref. [52].](image)
9. Conclusions

Gap junctional coupling plays a crucial role for impulse propagation in cardiac tissue. It is well established that a reduction of gap junctional conductance as occurring, e.g., during ischemia is importantly contributing to arrhythmogenesis because of the resulting reduction of conduction velocities, which is more than an order of magnitude larger than that observed during a reduction of excitability. At the same time, however, partial gap junctional uncoupling has the capacity to remove unidirectional conduction blocks and therefore acts, at least over a certain range of partial uncoupling, in an anti-arrhythmic fashion. This suggests that electrical uncoupling is not inevitably followed by arrhythmias. Rather, it ultimately depends on the balance between pro- and antiarrhythmic effects of uncoupling whether a certain reduction of axial current flow in the setting of non-uniform tissue structures can give rise to reentrant excitation. A second possibly pro-arrhythmic mechanism caused by gap junctions in the diseased heart might be related to heterogeneous cell coupling between cardiomyocytes and fibroblasts. Whereas it has been shown in cell culture that fibroblasts are able to relay electrical activation with substantial delays over appreciable distances, it remains to be shown whether gap junctional coupling between heterogeneous cell populations is indeed present in the diseased heart. Finally, since the observation of a clustering of sodium channels at the intercalated disc, the hypothesis that there might exist gap-junction independent mechanisms of impulse propagation in cardiac tissue has received renewed attention. Whereas computer simulations show the feasibility of such concepts, their validation in intact cardiac tissue will depend on the development of adequate experimental approaches.

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Note added in proof

With the reference to in vitro findings of electrotonic coupling between heterogeneous cells in the heart (Section 7), it has been shown very recently that gap junctional coupling exists between fibroblasts and cardiomyocytes both in the sinoatrial node [57] and in the myocardium [58].

References


