

channel structure: connexin pores have constricted segments with which ABG-sugars interact, other than the size-selective filter. From the cytoplasmic side, ABG-G3 blocks Cx26 but not Cx32 or Cx26/Cx32 channels, and the wider ABG-G4 blocks with decreasing effectiveness  $Cx32 > Cx26 > Cx26/Cx32$ . From the extracellular side, ABG-G3 blocks Cx26/Cx32 better than Cx26, and does not block Cx32 channels, while ABG-G4 has no effect on any channels tested. If block were exclusively at the size-selective filter, the pattern of block from both sides of the pore should be identical and consistent with the PA-sugar pore sizing study. Instead, the data show that pore width varies as the selectivity filter is approached from one side or the other. Specifically, the pore lumen of homomeric Cx26 and Cx32 channels narrows on the cytoplasmic side of the selectivity filter. Intriguingly, heteromeric Cx26/Cx32 channels show unique and substantial narrowing extracellular to the selectivity filter. This is significant as most connexin channels *in vivo* are heteromeric, and heteromeric channels are selective amongst second messengers while the corresponding homomeric channels are not. Our data suggest a consequence of 'heteromericity' is that segment/s of a pore-lining domain are asymmetrically displaced toward the center of the lumen. Supported by GM36044, NS56509.

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##### Chemical Gating Mechanism Of Connexin26-containing Channels By Aminosulfonate

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Protonated taurine directly and reversibly inhibits homomeric and heteromeric Cx26-containing hemichannels but not homomeric Cx32 hemichannels. It is unknown if taurine interacts with Cx26 and/or Cx32 in heteromeric channels, which domains are involved, or if junctional channels are taurine-sensitive. These issues were addressed with channels composed of Cx26 and/or Cx32 with/without cleavable 3KDa carboxyl-terminal (CT) tags (T). Hemichannel activity was assessed in liposomes, and by extracellular dye-uptake in cells. In contrast to untagged hemichannels, Cx26T/Cx32 and Cx26T hemichannels were not taurine-sensitive, but Cx26/Cx32T hemichannels were. Tag cleavage (Tc, leaving 4aa at the carboxyl-terminus) restored taurine-sensitivity of Cx26Tc/Cx32 hemichannels, but taurine surprisingly narrowed rather than closed Cx26Tc hemichannels. Thus, the 3KDa CT tag blocks taurine-sensitivity, unless hemichannels also contain Cx32, and the short 4aa CT extension affects Cx26Tc channel open state. Taurine effects on junctional channels were assessed by intercellular dye-coupling. Taurine substantially reduced dye-coupling by Cx26 and Cx26/Cx32T channels, but not by Cx26T, Cx26T/Cx32 or Cx32T channels. Junctional channels therefore have identical taurine-sensitivity as their component hemichannels. An intracellular site for taurine action was shown by a membrane-impermeable blocker of taurine uptake. Thus, all data indicate taurine-induced pore closure utilizes the Cx26 CT. Taurine binding to Cx26-CT was assessed by natural-abundance <sup>13</sup>C-HMQC-NMR. Overlapping resonances of Cx26-CT peptide in the presence and absence of taurine indicate no direct taurine binding to Cx26-CT. Peptide 'elisa' showed a pH dependent interaction occurs between Cx26-CT and the carboxyl-terminal 20aa of the Cx26 cytoplasmic loop (Cx26-CL). Acidification increases the binding affinity of Cx26-CL and Cx26-CT peptides, and only the protonated form of taurine negatively affects this interaction, suggesting that its disruption leads to channel closure. Structural analysis of Cx26-CT and Cx26-CL peptides in the presence and absence of taurine are ongoing. Supported by GM36044, DC7470.

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##### Quantitative Experimental Measurements And Mathematical Modeling Of Multi-cellular Dynamics In The Islet Of Langerhans

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Multi-cellular interactions and dynamics are key to mammalian physiology. Many organs such as the heart or brain have complex functionality that is brought about by the interactions between constituent cells. To discover general rules and principles that govern multi-cellular behavior, we are studying a relatively simple mammalian multi-cellular system- the islet of Langerhans. The islet is located in the pancreas and provides the sole source of the hormone insulin for regulating blood glucose levels. Thus, understanding the function of the islet is of critical importance to effectively treat diabetes. In

this work, we have focused on understanding the electrical dynamics in the islet that underlie the coordinated pulsatile secretion of insulin. We have used quantitative microscopy to measure intra-cellular free calcium activity ( $[Ca^{2+}]_i$ ) simultaneously over a large cellular population of the islet. To provide an experimental axis, we introduced graded changes in the electrical coupling between beta cells by applying both chemical inhibitors of gap junction activity as well as a genetic knockout of the gap junction protein. Upon this reduction in electrical coupling, synchronization of pulsatile electrical activity decreased throughout the islet. Furthermore, the propagating  $[Ca^{2+}]_i$  waves, which serve to synchronize electrical oscillations, slowed and were disrupted as electrical coupling was reduced. Using a mathematical model of islet cell electrical activity and multi-cellular coupling, we can quantitatively reproduce this experimental data. This allows us to make quantitative predictions of the multi-cellular electrical behavior for any given level of electrical coupling, as well as expected behaviors under perturbations of other parameters, such as  $K^+$ -channel mutations. We can also hypothesize that this behavior is general for electrical coupling in other multi-cellular systems, potentially allowing us to predict the electrical dynamics in other neuro-endocrine cell systems.

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##### Triarylmethanes - a New Class of Connexin Blockers

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Connexins (Cx) are a family of proteins with 4 transmembrane regions, which are encoded by 21 genes in humans and which form hexameric connexons (= hemichannels). These connexons can either function as transmembrane ion channels or assemble into gap junctions that directly mediate signaling between adjacent cells by allowing the passage of ions, metabolites and signaling molecules up to 1 kd in mass. Both gap junctions and hemichannels play important roles in many tissues and have therefore been proposed as potential new targets for the treatment of epilepsy, cardiac arrhythmia and cancer. However, there are no specific and potent pharmacological tools to study the physiological as well as the pathophysiological role of connexins. The existing connexin modulators are either of low potency or cross-react with other ion channels. In order to identify potent and selective connexin blockers we screened a small library of compounds containing pharmacophores known to modulate other ion channels. From this library, we identified four new small molecule chemotypes including triarylmethanes (TRAMs) like clotrimazole and benzimidazoles like astemizole that inhibit Cx50 channels in a sub-type specific manner with IC50 values in the range of 1-10  $\mu$ M while having little or no effect on those formed by Cx46, Cx36 and Cx32. We are currently exploring the structure activity relationship (SAR) of TRAMs for Cx50 inhibition and have recently identified T66 (N-[(2-chlorophenyl)(diphenyl)methyl]-N-(1,3-thiazol-2yl)amine), which exhibits an IC50 of 3  $\mu$ M. In general, the SAR of the Cx50 inhibiting TRAMs significantly differs from the SAR of KCa3.1 blocking TRAMs. We propose T66 and its derivatives as novel pharmacological tool compounds that may be used to study the physiological role of connexins.

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##### Pro-arrhythmic Effects Of Fibroblast-myocyte Coupling In Simulated Cardiac Tissue

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Fibroblasts comprise the majority of non-cardiac cells in normal heart and mediate the structural remodeling underlying progressive fibrosis in cardiac diseases. Recent experimental studies have shown that fibroblasts can electronically couple to myocytes via gap junctions and alter myocyte electrophysiology. However, the implications for cardiac arrhythmias are incompletely understood. In this study, we used mathematical modeling and computer simulation to investigate how fibroblast-myocyte coupling affects the dynamics of action potential duration (APD), excitation-contraction coupling, and alternans. Our major findings are: 1) Fibroblast-myocyte coupling shortens APD when fibroblast membrane conductance is high and resting membrane potential is low, but prolongs APD for other choices of conductance and resting potential. 2) Depending on the membrane conductance and resting potential of fibroblasts, fibroblast-myocyte coupling can either promote or suppress APD alternans by steepening or flattening APD restitution. 3) When alternans is calcium-driven, fibroblast-myocyte coupling always promotes alternans, and can result in electromechanically discordant alternans. 4) In cardiac tissue, fibroblast-myocyte coupling slows conduction velocity and broadens its restitution, promoting