## HEMICHANNEL INVOLVEMENT IN CA2+ DYNAMICS AND CONTRACTILITY OF SMOOTH MUSCLE CELLS IN ACUTELY ISOLATED SMALL MESENTERIC ARTERIES

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Intracellular Ca<sup>2+</sup> mediates a variety of vascular endothelial and smooth muscle cell functions. Smooth muscle cells (SMC) respond to biological activators with oscillatory and propagating rises in  $[Ca^{2+}]_i$  that are highly organized in both time and space. Gap junctions (GJs) play a crucial role in the communication between vascular cells and in the synchronization of Ca<sup>2+</sup> signals thereby tightly controlling the level of vasoconstriction. Before being incorporated into GJs, connexin (Cx) hemichannels reside in the plasma membrane in a closed state. Recent evidence suggests that hemichannels can be opened by various messengers and conditions, thereby forming a pore that allows the passage of ATP and ions. Using confocal microscopy and the  $Ca^{2+}$  sensitive dye Fluo-3, we examined the role of hemichannels in dynamic Ca<sup>2+</sup> responses of SMC in intact acutely isolated small rat mesenteric arteries. Furthermore, we assessed the involvement of these signalling partners in contractile responses of small mesenteric arteries using a wire myograph for isometric tension measurements. Importantly, the experimental conditions were such that vasomotion, characterized by synchronized Ca<sup>2+</sup> signals, was avoided because in that case gap junctions between SMC and myo-endothelial gap junctions are expected to contribute. Norepinephrine (NOR, 3 µM) induced  $Ca^{2+}$  oscillations that were reduced in frequency by 98.4 % (p< 0.05) when exposed to carbenoxolone (CBX, 50 µM), a none specific Cx channel inhibitor. Gap27 (200 µM), a Cx mimetic peptide that blocks hemichannel responses (assayed by ATP release and dye uptake) after short incubation, reduced the spiking frequency by 96.4 % (p< 0.05). Suramin (200  $\mu$ M) and PPADS (75 µM), two P2Y receptor antagonists, decreased the spiking frequency by 90.5 % (p<0.05) and 96.4% (p<0.01) respectively. Apyrase (5 U/ml), an enzyme that rapidly degrades extracellular ATP, reduced the spiking frequency by 71.4 % (p<0.01). None of these agents affected the amplitude of the  $Ca^{2+}$  oscillations. Both gap27 (56.6 %, p<0.01) and CBX (53.4 %, p<0.05) reduced the NOR-induced contractions. Incubation with suramin decreased the NOR-induced contractions by 31.6 % (p<0.001). Our results suggest a role for Cx hemichannels and purinergic signaling in  $Ca^{2+}$  oscillations and contractility.