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InsP₃R-RyR channel crosstalk augments sarcoplasmic reticulum Ca²⁺ release and

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Abstract

 Ca^{2+} transients (CaT) underlying cardiomyocyte (CM) contraction require efficient Ca^{2+} coupling between sarcolemmal Ca^{2+} channels and sarcoplasmic reticulum (SR) ryanodine receptor Ca^{2+} channels (RyR) for their generation; reduced coupling in disease contributes to diminished CaT and arrhythmogenic Ca^{2+} events. SR Ca^{2+} release also occurs via inositol

1,4,5-trisphosphate receptors (InsP₃R) in CM. While this pathway contributes negligeably to Ca²⁺ handling in healthy CM, rodent studies support a role in altered Ca²⁺ dynamics and arrhythmogenic Ca²⁺ release involving InsP₃R crosstalk with RyRs in disease. Whether this mechanism persists in larger mammals with lower T-tubular density and coupling of RyRs is not fully resolved. We have recently shown an arrhythmogenic action of $InsP_3$ -induced Ca^{2+} release (IICR) in end stage human heart failure, often associated with underlying ischemic heart disease (IHD). How IICR contributes to early stages of disease is however not determined but highly relevant. To access this stage, we chose a porcine model of IHD, which shows substantial remodelling of the area adjacent to the infarct. In cells from this region, IICR preferentially augmented Ca²⁺ release from ion coupled RyR clusters that otherwise showed delayed activation during the CaT. IIC, in turn synchronised Ca²⁺ release during the CaT but also induced arrhythmogenic delaged afterdepolarizations and action potentials. Nanoscale imaging identified co-clustering of InsP₃Rs and RyRs, thereby allowing Ca²⁺-mediated channel crosstalk. Mathematical ... rdelling supported and further delineated this mechanism of enhanced InsP₃R-RyRs cc¹pling in MI. Our findings highlight the role of InsP₃R-RyR channel crosstalk in Ca²⁺ release and arrhythmia during post-MI remodelling.

Keywords:

Ischemic heart disease; myourdial infarction; Excitation contraction coupling; cardiac arrhythmia; Ca²⁺ homeoclasis, mositol 1,4,5-trisphosphate receptor.

Abbreviations:

Excitation contraction coupling (ECC), action potential (AP), delayed after depolarisation (DAD), inositol 1,4,5-trisphosphate (InsP₃), InsP₃ receptor (InsP₃R), InsP₃-induced Ca²⁺ release (IICR), cardiomyocyte (CM), myocardial infarction (MI), sarcoplasmic reticulum (SR), L-type Ca²⁺ channel (LTCC), ryanodine receptor (RyR), sarco/endoplasmic reticulum Ca²⁺ ATPase (SERCA), calsequestrin (CSQ), sodium/calcium (Na⁺/Ca²⁺) exchanger (NCX), Ca²⁺ calmodulin kinase II (CaMKII), Ca²⁺ transient (CaT), dull duration half maximum (FDHM), full width half maximum (FWHM), time to 50% peak amplitude (TF₅₀), Fourier Ring Correlation (FRC), T-tubule (TT), heart failure (HF), ischemic heart disease (IHD).

1. Introduction.

Cardiomyocyte contraction following action potential-(AP)-mediated depolarisation occurs via Excitation Contraction Coupling (ECC) [1, 2]. Activation of Ca²⁺ release from the (SR) via RyRs by Ca²⁺ entering the cell through L-type voltage gated Ca²⁺ channels (LTCC) on the sarcolemma is central to this process. Synchronised activation of RyRs distributed throughout the CM volume in turn generates an homogenous cell-wide increase in intracellular Ca²⁺ and the efficient contraction necessary for pump function [3-5]. This coordinated activation of RyRs throughout the CM is fac litated by T-tubular (TT) invaginations of the sarcolemma that place LTCC at locations opposing SR RyR clusters within 15 nm of each other, forming dyads or couplois [5, 6-8].

In rodents, TT are regularly spaced along the Z-lines of the sarcomeres, whereas, in disease, and in larger mammals, including pigs and in human, TT density is substantially lower [3, 5, 8-12]. With lower TT density, a greater proportion of RyR clusters are not associated with LTCCs (non-coupled, extra dypole) and therefore not directly activated by Ca²⁺ influx during the AP but by Ca²⁺ diffueing from coupled dyadic RyRs [13, 14]. Consequently, Ca²⁺ release synchrony across the cell during the CaT is reduced, resulting in lower ECC gain, global Ca²⁺ transient amplitude and contractile force [4]. Further, non-coupled RyRs display increased spontaneous Ca²⁺ release in HF, contributing to the initiation of Ca²⁺ waves, which via the electrogenic Na⁺/Ca²⁺ exchanger (NCX) and altered repolarising K⁺ currents, promote delayed after depolarisations (DADs) and APs that underlie potentially organ-wide arrhythmia [15-20].

Ventricular CMs also express inositol 1,4,5'-trisphosphate receptor (InsP₃Rs) Ca²⁺ release channels (the type 2 isoform, InsP₃R2) [21-25]; their expression and conductance are however substantially lower than RyRs [26-28]. While induction of inotropy and arrhythmogenic activity have been reported, effects of InsP₃-induced Ca²⁺ release (IICR) in healthy CM, are if detected, generally modest and vary between studies and species [23, 25, 29, 30]. In ventricular CMs from hypertrophic or failing rodent hearts, the influence of IICR on CM physiology is greater [23, 31, 32], with increases in Ca⁻ amplitude, diastolic Ca²⁺ and frequency of arrhythmogenic events following exposure to InsP₃ or neurohormones that promote InsP₃ generation such as Angiotensin II (AngII: a...⁴ indothelin (ET-1) reported [23, 29, 33, 34]. Despite increased expression in diseas 2. S Ca²⁺ release via InsP₃Rs is minor relative to via RyRs during the CaT. To explain now this small change in cytosolic [Ca²⁺] generated by IICR influences ECC, we and o here have proposed a mechanism that involves signal amplification by RyRs [23, 25, 35-37]. IICR is also reported to engage proximal NCX, thereby contributing to altered electrop physiology [31].

Despite extensive rode a studies, the role and mechanism of action of IICR in the pathophysiology of ventricular CM of larger mammals are not well defined. Given the substantial differences in the physiology and cellular architecture, including in the coupling of RyRs with the TT membrane, between rodent and large mammals [4, 38], analysis of the role of IICR in this context is vital. Previous studies suggested a contribution of IICR to the arrhythmogenic action of neurohormonal agonists in human HF [31]. However, these studies did not directly probe InsP₃R activity. Recently, by either direct introduction of InsP₃R activation to the arrhythmogenic action of GPCR agonists in end stage human HF [39].

Moreover, in HF, InsP₃ increased Ca²⁺ leak from the SR that on the background of lower SERCA activity present, led to SR Ca²⁺ content depletion and reduced CaT amplitude. Despite these advances, how IICR contributes to human CM physiology during the early stages of remodelling prior to HF observed during ischemic heart disease (IHD) is not determined. The mechanism by which IICR influences Ca²⁺ dynamics at the molecular level is also not fully understood. Given the high incidence of sudden cardiac death and arrhythmias in IHD patients and in the early period following MI, identifying mechanisms underlying pathology in this context is essential [20, 40]. Here we set out to test the hypothesis that IICR plays a dual role in pathological remodelling of CM, whereby is been rescued desynchronised Ca2+ release during ECC but also increased spontaneous Ca⁺⁺ release events that could in turn contribute to arrhythmia. To these ends, we chose an established preclinical pig model of ischemic heart disease where CM in the a ea adjacent to the infarct undergo substantial remodelling associated with increased arrhythmic activity [41]. Using Ca²⁺ imaging and patch clamp electrophysiology, we er to plished that Ca²⁺ signalling between InsP₃Rs and RyR is important in regulation of Ca^{2+} release and arrhythmogenic activity in post-MI remodelling. We further showed using DNA-PAINT [42] super resolution microscopy the best evidence of co-clustering of InsP₃Rs and RyRs, which we further showed by mathematical modelling could contribute to the functional interactions between these channels observed in MI. Notably, unlike in human, owing to retained SERCA function, IICR did not lead to increased Ca²⁺ leak and SR Ca²⁺ depletion and thus diminished CaT in pig MI CM. Together, these data provide key insights into the mechanism of action of InsP₃ signalling in the remodelled heart of large animals and how this may differ from CM in endstage heart failure.

2. Results

2.1 InsP₃ increases synchronicity of SR Ca²⁺ release during ECC without substantially affecting CaT properties.

We examined the influence of IICR on CaT in voltage clamped CM from Sham and MI pig as per the protocol in Figure 1A. Cardiac remodelling after MI determined by MRI is presented in Table S1 and is as previously described [41]. CaT amplitude and time to 50% decay (Figure 1Ci,Cii) were unaltered between Sham and MI CM and no effect of InsP₃ was detected. The synchronisation of Ca²⁺ release following cell depolarisation ouring the CaT was investigated by analysis of the average time to 50% amplitude of true CaT on a pixel by pixel basis (TF₅₀). TF₅₀ was significantly greater in MI than in Sham CM at baseline (Figure 1Ciii, left). Notably, InsP₃ application significantly reduced T^{E-1} in MI but not in Sham (Figure 1Ciii, right). Together, these data reveal the potential for IICR to modulate local Ca²⁺ release and its synchronicity during the CaT in post-M + CM remodelling.



Figure 1. InsP₃ accelerates **C**. ²⁺ re sase in **MI CM. A.** Cartoon of experimental protocol. **Bi.** Xt images of CaT recorded by confocal linescent again of Fluo-4 in Sham and MI CM in the presence and absence of InsP₃ at 1 Hz pacing. **Bii.** Normalised fluc escence traces (F/F0) of CaT described in i. **C.** Summary data of (i) CaT amplitude (ii) time to 50 % decay of the CaT (F50 decay) and (iii) pixel by pixel averaged time to 50% of CaT peak amplitude (TF₅₀). Left panel: example c averaged Xt image of 5 CaT and time to F₅₀ on a pixel by pixel plotted. Right panel: summary data of average **T** ₅₀ calculated as on the left panel. (Sham ±InsP₃ n_{cells}=15-18; N_{pigs}=6,6; MI ±InsP₃ n_{cells}=15,21; N_{pigs}=5,5). Statistical analysis was by a generalised mixed effect model with Tukey Post hoc analysis.

2.2 InsP₃ stimulates increased RyR-dependent sparks in post-MI ventricular CM.

Given that Ca^{2+} sparks are the fundamental units of Ca^{2+} release underlying the CaT, we examined whether they were influenced by IICR. Ca^{2+} sparks were recorded by linescan confocal imaging as in Figure 2A and B. At baseline, Sham and MI CM displayed a low

frequency of Ca²⁺ sparks (Figure 2C). InsP₃ significantly increased Ca²⁺ spark frequency in MI CM, whereas it was without effect in Sham (Figure 2B,C). Ca²⁺ spark amplitude, duration (FDHM) and width (FWHM) were unaffected by InsP₃ (Figure S2A). The contribution of RyRs to the effect of InsP₃ on Ca²⁺ sparks was next determined. RyR inhibition with tetracaine (Tet, 1 mM) significantly reduced Ca²⁺ spark frequency in both Sham and MI CM. <u>Under</u> <u>these conditions of RyR inhibition however, Ca²⁺ release events were observed in more cells</u> at a higher frequency in MI than in Sham (1 from 8 cells in Sham vs 6 from 12 cells in MI, p=0.0425 at frequency 0.0075±0.0075 vs 0.06±0.02 sparks/s/1 °0 µ n in sham vs MI; p=0.0441, n=8,12, N=3,3). The contribution of RyR to Ca²⁺ spark properties during InsP₃ stimulation was assessed by their analysis before and after Tet treat as in (Figure S2B). Amplitude of Ca²⁺ release events was significantly reduced in the presence of Tet whereas FWHM and FDHM were unaffected (Figure S2C). Top etrem is estimated that increased Ca²⁺ spark activity in MI CM requires both active RyRs and InsP₃Rs.

We next determined whether thr. incluased Ca²⁺ spark frequency in MI CM exposed to InsP₃ led to greater SR Ca²⁺ leak and JP Ca²⁺ store depletion. In the absence of InsP₃, no difference in spark-dependent Ca²⁺ 'eak from the SR, (spark mass (FWHM.FDHM.Amplitude) x frequency) [43] betweet Sham and MI CM was observed, whereas with InsP₃ it was significantly increased in MI (Figure 2D). Consistent with InsP₃Rs enhancing leak via RyRs, the effect of InsP₃ was prevented by Tet (Figure 2D). SR Ca²⁺ load was next measured by quantitating the integral of the NCX current during SR store exhaustion with caffeine [44]. In contrast to our findings in human HF, SR load was no different between Sham and MI CM, and no effect of InsP₃ on either Sham or MI CM was detected (Figure 2E). SERCA-mediated Ca²⁺ uptake rate was also assessed according to previous methods whereby the rate constant of decay of the electrically-evoked CaT was subtracted from the Caffeine-elicited

 Ca^{2+} elevation [45], and found to be no different between Sham and MI (Figure S2D). Together, these data support the notion that in MI CM, InsP₃ acts via RyR to increase Ca^{2+} sparks. Moreover, despite increased Ca^{2+} spark-dependent leak, and consistent with no effect of IICR on CaT amplitude, InsP₃ did not alter SR Ca^{2+} load.





Figure 2. InsP₃ increases spark 1. The angle of Values of Ca²⁺ stores in MI CM. A. Experimental protocol used for analysis of Ca²⁺ sparks. **B.** Example Xt images of Ca²⁺ sparks acquired by linescan imaging of voltage clamped MI CM in the preserve at a basence of InsP₃. **C.** Summary data of frequency Ca²⁺ spark/events under the conditions shown (for Sharine Liner's n_{cells}=19-24; N_{pigs}=6-9; MI ± InsP₃ n_{cells}=26-27; N_{pigs}=7-9; for Tet in the presence of InsP₃, Sham vs MI, n_{cells}=9, ?; N_{pigs}=3). **D.** Analysis of total spark mass (FWHM*FDHM*Amplitude*frequency) as a surrogate of spark-depende the example of experimental protocol showing a train of electrically evoked Ca²⁺ transients and the stimulation traces followed by caffeine-mediated Ca²⁺ increase and activation of NCX current (I_{NCX}). Right panel: Summary data of the integrated NCX current in Sham and MI CM in the presence and absence of InsP₃ (Sham ± InsP₃ n_{cells}=11,13; N_{pigs}=5,6; MI ± InsP₃ n_{cells}=10,13; N_{pigs}=4,6). Statistical analysis of C and D was by non-parametric ANOVA with Tukey post hoc analysis and E by a generalised mixed effect model with Tukey Post hoc analysis.

2.3 Alterations in InsP₃R expression and subcellular localisation are observed in MI.

The greater activity of InsP₃ in MI CM suggested an increase in expression and/or an alteration in subcellular localisation of InsP₃Rs. Increased InsP₃R activity could also be amplified through interaction with RyRs sensitised by CaMKII phosphorylation, which we have previously reported in this model [17]. Immunoblotting revealed a significant increase in InsP₃R2 in MI (Figure 3A) while RyR2 expression was unchanged. RyR2 phosphorylated at the serine 2814 CaMKII site (RyR2-pS2814) was however significantly increased (Figure S3A). InsP₃R2 mRNA levels (ITPR2) were unaltered as previously described [23], and RyR2 mRNA abundance was decreased (Figure S3B). Immunoblotting showed no significant changes in expression of SERCA2a or NCX (Figure S3C).

The subcellular localisation of InsP₃R2 relative to Ry, is and of both channels relative to sarcolemmal and TT membranes, labelled viction NCX antibody, was next determined (Figure 3B) [46]. Confocal images were deconvolved and InsP₃R and RyR clusters identified and analysed. RyRs were primarily distributed in a striated pattern along the Z line, where their distribution coincided with the sarcolemma/TT (Figure 3B,C). InsP₃Rs showed a similar distribution as RyRs, although a 'ower proportion of InsP₃Rs aligned with TT (Figure 3B,C). For both InsP₃R and R⁺R clusters, no significant difference in their location relative to membrane was detected n MI (Figure 3C). The similarity in the distributions of both channel types was supported by Mander's co-localisation analysis, which showed an overlap of ~ 5% of RyRs with InsP₃Rs and ~ 10% of InsP₃Rs with RyRs (Figure S3D). As cluster area correlates with cluster channel number [47], it was also analysed. RyR cluster area was almost double that of InsP₃R clusters (Figure 3D). No significant change in the area of InsP₃R or RyR clusters



was detected following MI, although for InsP₃Rs, this approached significance (Figure 3D).

Figure 3. InsP₃Rs are increased in expression in MI and show altered cellular distribution. A. Immunoblot analysis of InsP₃R and RyR in Sham and MI CM (N = 4 and 4 pigs). Left: InsP₃R2 (red), RyR2 (green) and CSQ (red) used as a loading control probed on the same blot are shown. Right: Densitometric analysis of InsP₃R2 and RyR2 normalised to the abundance of CSQ in the same blot. Analysis by Mann-Whitney test. **B.** Confocal images of an MI CM immunostained with antibodies against InsP₃R2 (magenta) and RyR2 (green). TT were stained with antibodies against NCX (white). Scale bar = 20 μ . Images are as follows: top left, deconvolved confocal image; top right, mask of deconvolved image showing distribution of clusters; bottom left, zoom of regions shown in confocal image; bottom right, zoom of masked image. A skeleton of the TT network is also shown in the bottom right image. **C.** Frequency distribution of InsP₃R2 and RyRs relative to a TT membrane in Sham vs MI (Sham n_{cells}=19; N_{pigs}=5; MI n_{cells}=20; N_{pigs}=5). **D.** InsP₃R2 and RyR2 cluster areas in Sham and MI plotted according to their location relative to the TT membrane. 2 way ANOVA with Bonferroni post-hoc analysis.

In pig MI CM, we have previously described increased propensity of Ca²⁺ release at sites distal to TT membranes (non-coupled) [48]. We therefore tested whether InsP₃Rs were differentially localised to coupled vs non-coupled RyR clusters. InsP₃Rs and RyRs were categorised according to their distance to the sarcolemma as coupled (<0.5 μ) or non-coupled (>2 μ) [46]. Although, no change in the number of either coupled or non-coupled InsP₃R clusters was observed, clusters area was significantly increased in MI at both regions (Figure 3E). RyR cluster area did not change at either region (Figure 3E). Together, this data suggested the potential for a greater influence of InsP₃Rs on RyR: in MI.

2.4 InsP₃R co-clustering with adjacent RyRs is increased in MI.

To examine whether InsP₃Rs are sufficient *y* close to RyR to influence their activity, their relative localisation was next detern. new by DNA-PAINT super resolution microscopy [42]. Fourier Ring Correlation (FRC) of the InsP₃R and RyR labelling data revealed a lateral resolution of ~46 nm [49] (Figure 34A). Consistent with our confocal imaging (Figure 3B), DNA-PAINT showed InsP₃Rs and RyRs to be localised in clusters that were arranged in a striated pattern (Figure 4A). Estimation of the number of receptors accommodated in the stained regions, as described [47, 50], revealed a broad distribution of cluster sizes (Supplementary Figure S4B,C), although many isolated receptors (cluster = 1 in histogram) were also detected. Since clustered channels are considered to underlie Ca²⁺ spark events [51], and that single point data could be contaminated by false-positive localisations, these were excluded from subsequent analysis. Applying these criteria revealed a significant increase in the number of InsP₃Rs per cluster in MI while RyR number per cluster was unchanged (Figure 4B). Using the the proportion of clusters with 2 or more





Figure 4. DNA-PAINT analysis of InsP_3R2 and RyR2 in Sham and MI. A. DNA-PAINT images of $InsP_3R2$ (magenta) and RyR2 (green) in sections cut from Sham (ai) and MI (bi) cardiac tissue. Scale bar = 1 μ . aii and bii, images show a zoom of the area highlighted in yellow in ai and aii. Aaiii and biii. Distribution of RyRs showing regions of clusters overlapping with $InsP_3R2$ clusters in white. Scale bar = 0.5 μ . **B.** Size of $InsP_3R2$ and RyR2 clusters comprising 2 or more channels represented as number of channels accommodated by labelled area. **C.** Proportion of $InsP_3R2$ or RyR2 clusters in clusters comprising 2 or more channels. **D.** Overlap of $InsP_3Rs$ with RyR clusters of 5 or more channels. **E.** Proportion of $InsP_3Rs$ within 30 nm of edge of RyR cluster. Sham $n_{sections}=29$; $N_{pigs}=3$; MI $n_{sections}=18$; $N_{pigs}=3$. Analysis by unpaired *t*-test with Welch correction. **F.** Cartoon summarising data from Figs 3 and 4 showing relative distributions of $InsP_3Rs$ and RyRs at coupled and non-coupled sites and their distribution in clusters.

channels as a measure of cluster size, InsP₃R cluster size was unchanged but RyR cluster size

was significantly increased in MI (Figure 4C). We next investigated whether InsP₃Rs and RyRs co-clustered. Analysis of the overlap of the distributions of the two channel types revealed that on average 2 or more InsP₃Rs overlapped with each RyR cluster, indicating the presence of mixed-channel clusters (Figure 4D). No difference in the number of overlapping InsP₃Rs per RyRs cluster was detected between Sham and MI. To account for InsP₃Rs that could influence RyR activity, we performed a second analysis whereby InsP₃Rs within 30 nm (scale of a single channel) of a RyR cluster were added to the InsP₁R population that overlapped with the RyR clusters. This InsP₃R population was significan the increase in MI (Figure 4E). Together, our analysis identifies for the first time natioscente co-clustering of InsP₃Rs and RyRs in CM, and moreover, an increase in InsP₃Rs previmal to RyRs in MI (Figure 4F).

2.5 Mathematical modelling predicts influence of non-coupled InsP₃Rs on Ca²⁺ spark generation.

Having shown increased overlap of InsP₃Rs with RyRs clusters, we explored using a mathematical model that incorporates the activity of both RyRs and InsP₃Rs in a Ca²⁺ release site (cluster) [52], whe her sensitisation of RyR by Ca²⁺ release via these InsP₃Rs could explain the increase in Ca²⁺ sparks seen in MI CM exposed to InsP₃ (Figure 5A and Figure S5A). We particularly examined whether InsP₃R-RyR-Ca²⁺ mediated crosstalk within a non-coupled site could facilitate the recruitment of this site by Ca²⁺ diffusing from nearby coupled sites, thereby providing a mechanism for the increased Ca²⁺ sparks in MI at these sites.

To this end, we built on our mathematical model that examined spontaneous spark activity at a dyadic Ca²⁺ release site with a fixed number of RyRs based on published data [51, 53] (orange blocks in Figure S5B) over a varying number of InsP₃Rs (0, 5, 10, or 20 per cluster; magenta blocks in Figure S5B) [52]. We extended this model to examine the effect of InsP₃Rs on Ca²⁺ release at a non-coupled (NC) site under the influence of coupled (C) sites placed equidistant (2 µm centre to centre) on either side (with 10 InsP₃Rs), which were either allowed to spontaneously activate (2 C 1 NC C 0 Triggers) or which were triggered, as in ECC, at 1 Hz (2 C 1 NC C 3 Triggers) (Figure 5B,C). Our choic e of using a reduced-order representation of the Ca²⁺ release site with 15 R/Rs, was justified by previous 3D simulations, which showed that the spatial distribut. In c f RyRs in a Ca²⁺ release site was not critical to the Ca²⁺ spark profile when >9 R⁻R, were in the cluster [54]. Ca²⁺ changes generated by the model are measured in terms of [Ca²⁺] since they are more informative than simulated fluorescence changes of the Ca²⁺ indicator Fluo-4, which is saturated by the [Ca²⁺] reaching 10s of µM present in microdomain around the Ca²⁺ release site [55, 56].

200 simulations were performed for each InsP₃R number condition with results obtained from a 2 s period after equinitial 1 s wait time for the system to achieve steady state (Figure 5B,C). Consistent with our recent findings [52] and with results presented here (Figure 2C), the chance of observing a spontaneous spark at the non-coupled site in isolation (1 NC 0 Trigger) increased with InsP₃R number (Figure 5C). Ca²⁺ spark amplitude decreased to a minor extent with the number of InsP₃R, which we attributed to a depletion of Ca²⁺ at the junctional SR and thus a reduction in the Ca²⁺ gradient between the cytosolic space adjacent to the release site and the SR lumen; whereas Ca²⁺ spark FDHM remained unchanged (Figure S6) [52]. To test whether InsP₃R-RyR co-clustering at non-coupled sites led to their



Figure 5. Mathematical modelling supports enhancement of Ca²⁺ sparks at non-coupled sites by InsP₃induced Ca²⁺ release. A. Cartoon showing localization of non-coupled (NC) and coupled (C) Ca²⁺ release sites and how this relates to the setup of the model used for analysis of impact of InsP₃Rs at non coupled sites on Ca²⁺ release at these sites. **B.** Examples of trials performed under the conditions indicated with [Ca²⁺]_i changes during the Ca²⁺ release events shown. The red arrowheads indicate timings of triggers used to stimulate Ca²⁺ release. The locations of coupled (C) and non-coupled (NC) sites on the simulations is indicated. **C.** Summary data showing percentage of simulations that display Ca²⁺ sparks under the indicated simulation conditions and differing number of InsP₃Rs. **D.** Analysis of the timing of Ca²⁺ sparks across the duration of the simulation in simulations presented in A with either 0, 5, 10 or 20 InsP₃Rs present in the non-coupled cluster. **Di.** Swarm plots of Ca²⁺ spark occurrences shown for all simulations. The time points of the triggers used in the 2 C 1 NC with 3 Triggers simulation condition are indicated by dashed lines and red arrow heads. **Dii.** Density function of the Ca²⁺ spark events relative to the Ca²⁺ trigger that is indicated by a dashed line and red arrowhead.

nearby coupled sites, the overall frequency as well as the timing of the Ca²⁺ spark events at

the non-coupled site was determined (including relative to the triggered spark in the 2 C 1

NC 3 Triggers condition) (Figure 5B-D). Consistent with this hypothesis, our model showed that spark frequency at the non-coupled site is increased in the presence of neighbouring coupled sites (Figure 5C, Di; 2 C 1 NC 0 Triggers). Importantly, and further showing the role of coupled sites in their activation, Ca²⁺ sparks were most frequent at or soon after triggers were imposed on the coupled sites (Figure 5Di; 2 C 1 NC 3 Triggers). An increased number of simulations with 2 sparks was also detected with increased InsP₃Rs and in the presence of triggers (Figure 5C). The positive effect of triggers on frequency of Ca²⁺ sparks was further validated by fitting a density function to the histogram of the timing of Ca²⁺ sparks that occurred within either 100 ms before or after the first and cecond trigger (Figure 5Dii). The increase in Ca²⁺ spark occurrence within 100 ms of the triggers was notably absent in the 2 NC 1C 0 Trigger case (Figure 5C), highlighting hit the increased occurrence of Ca²⁺ sparks after a trigger was not purely due to simulation stochasticity (Figure 5B-D). This model further predicted that Ca²⁺ release ct the non-coupled site is strongly determined by its distance from the coupled site (Figure S6B-E). Indeed, at intersite distances >4 µm, the activity of coupled sites no longer significantly influence that at non-coupled sites. Altogether, these data indicate that while IICR is sufficient to augment Ca²⁺ spark frequency at non-coupled sites in conation, it also facilitates Ca²⁺ spark generation at non-coupled sites when Ca²⁺ diffuses from activated coupled sites.

2.6 Diminished Ca²⁺ release at non-coupled sites during MI is enhanced by InsP₃.

Prompted by this modelling data and the harmonisation of Ca^{2+} release in MI by InsP₃ in Figure 1Ciii (decreased average TF₅₀ peak across linescan), we tested whether IICR differentially influenced Ca^{2+} release at coupled vs non-coupled sites in Sham and MI.

Release sites were classified as coupled or non-coupled based on their timing of activation during the CaT according to our previously described distance-map algorithm that correlates spatial and temporal Ca²⁺ release profiles [17, 46]. Coupled and non-coupled sites were activated with a TF_{50} <17.7 ms and TF_{50} >27.4 ms of the CaT respectively (Figure 6B). Events with a TF_{50} between these measures were omitted to allow better discrimination between the two regions [46]. The Xt image in Figure 6B shows the identification of Ca²⁺ sparks at coupled and non-coupled sites.

Ca²⁺ sparks were detected at both coupled and non-coupled sites in Sham CM, although their frequency was lower at non-coupled sites (Figure 6Ci,Cii). InsP₃ did not affect Ca²⁺ spark frequency at either location in these Sham CM. In MI CM, a low frequency of Ca²⁺ sparks was detected at both coupled and non-coupled sites in the absence of InsP₃ but in contrast to Sham, InsP₃ significantly increased event frequency, particularly at non-coupled sites (Figure 6Ci,Cii).



Figure 6. Ca²⁺ release at non-coupled sites is increased by InsP₃ in MI. A. Cartoon showing location of coupled and non-coupled sites. **B.** Xt image of CaT (middle) with time taken to reach 50% max along the linescan shown (left). An Xt image of the quiescent period subsequent to pacing of the same cell is shown. Ca²⁺ sparks at coupled and non-coupled sites are shown. Coupled sites are classified as those that reach TF₅₀ in less than 17.7 ms and non-coupled sites as those that take longer than 27.4 ms to reach TF₅₀. Right: Traces of CaTs in the coupled and non-coupled regions. **C.** Spark frequency at coupled and non-coupled sites in the presence/absence of InsP₃ in Sham and MI CM are shown. Ci. Spark frequency at Coupled release sites. Cii. Spark frequency at Non-Coupled release sites. **D.** Comparison of the rate of rise of the CaT at non-coupled sites that either exhibited Ca²⁺ sparks or not when InsP₃ was present (i.e. \pm activatable InsP₃Rs) in MI (Di) vs Sham (Dii) CM. **E.** % of linescan that has a TF₅₀ below 17.7 ms (early release sites) in Sham and MI \pm InsP₃. (Sham \pm InsP₃: n_{cells}=13,18; N_{pigs}=4,5; MI \pm InsP₃: n_{cells}=18,23; N_{pigs}=5,5). Statistical analysis was by a generalised mixed effect model with Tukey Post hoc analysis.

Whether IICR augmentation of non-coupled RyR activity acts to accelerate CaT rate of rise

uniquely at these sites was next investigated. The rate of rise of the CaT at non-coupled sites that exhibited Ca^{2+} sparks in response to $InsP_3$ (Sparks +; used as a surrogate of

functional InsP₃Rs) was determined and compared with non-coupled sites that did not show Ca^{2+} sparks (-) in response to InsP₃. Notably, non-coupled sites harbouring functional InsP₃R exhibited enhanced Ca^{2+} release kinetics during CaTs whereas no such increase in CaT kinetics occurred at sites without an increase in Ca^{2+} sparks after exposure to InsP₃ (Figure 6Di). InsP₃ was without effect on CaT rate of rise at non-coupled or coupled sites in Sham CM or at coupled sites in MI (Supplementary Fig S7), irrespective of the presence of Ca^{2+} sparks.

To determine whether IICR could recruit 'non-coupled' sites and decrease the latency of their activation during the CaT, we analysed whether $\ln SP_3$ increased the fraction of the linescan that reached TF₅₀ at <17.7 ms (coupled sites). It baseline, MI CM exhibited a lower proportion of early Ca²⁺ release sites than Superior (Figure 6E). In the presence of InsP₃ however, the % of early response sites was significantly increased, reaching a level similar to Sham. InsP₃ did not alter the responsiveness of Ca²⁺ release sites in Sham. These data together point to a special role for $\ln Sr_3$ Rs at non-coupled Ca²⁺ release sites in regulation of CM Ca²⁺ handling in MI. At whese locations, InsP₃Rs signal to RyRs enhancing their recruitment and throug's trais mechanism rescue the reduced synchronicity of Ca²⁺ release observed in MI to that in Jham.

2.7 Spontaneous activity is increased by InsP₃ in MI.

 Ca^{2+} waves arising from non-coupled RyR clusters via NCX lead to arrhythmogenic DADs and APs [17]. Whether IICR engagement of non-coupled RyRs could promote arrhythmogenic activity was therefore determined. To this end, using NCX currents (I_{NCX}) as a reporter of

Ca²⁺ waves, we first examined whether they were increased in MI CM by InsP₃. Conditions for increasing wave incidence were generated by pacing at 2 Hz for 1 min and incubation with Isoproterenol (Iso; 10 nM). I_{NCX} events were recorded during 40 s after pacing as per the protocol shown in Figure 7A [17]. In the absence of InsP₃, while both Sham and MI CM displayed I_{NCX} events, these events were significantly more frequent in MI (Figure 7B,C). This increased activity was not due to differences between Sham and MI in SR Ca²⁺ uptake rate (Figure S2D). In MI but not in Sham, InsP₃ induced an additional significant increase in the frequency of these events (Figure 7B,C). Given the positive relationship between I_{NCX} and APs, the effect of InsP₃ on I_{NCX} amplitude in MI CM was also measured. InsP₃ significantly increased I_{NCX} amplitude in MI CM exposed to Iso (Figure 7D).

Whether the increase in I_{NCX} events in MI CN is analated into increased frequency of DADs and APs was next determined. DADs and Ps were recorded as described above albeit, CM were paced and DADs and APs recorded under current clamp during the following 40 s period (Figure 7A)[17]. As no sportaneous activity was observed in the absence of Iso, experiments were only performed in its presence. At baseline, DAD frequency was significantly greater in Nits an in Sham CM but no effect of InsP₃ was detected for either condition (Figure 7E,F). A 's were also rarely detected in Sham or MI CM at baseline (Figure 7E,G). Notably however, AP frequency was uniquely significantly increased by InsP₃ in MI CM, with no effect in Sham detected (Figure 7G). This specific action of InsP₃ on APs in MI CM was also apparent when the incidence of DAD relative to AP was compared between Sham and MI (Figure 7H). Since greater DAD amplitude increases the propensity for AP generation, whether it was affected by InsP₃ was analysed (Figure 7I). While InsP₃ had no effect on DAD amplitude in Sham, it significantly increased DAD amplitude in MI, suggesting

that in MI, increasing DAD amplitude contributes to the mechanism by which $InsP_3$ increases APs occurrence.



Figure 7. InsP₃-induced Ca²⁺ release contributes to increased arrhythmogenic activity in MI **CM.** A. Cartoon of experimental protocol used for induction of arrhythmogenic Ca^{2+} waves and detection of DADs and APs. For induction of Ca^{2+} waves and the associated I_{NCX} events reported, cells were paced under voltage clamp and NCX activity monitored at resting membrane potential. For detection of DADs and APs, cells were paced in current clamp mode with current injections at 2 Hz for 90 s. CM were perfused with 10 nM lso. B. Summary data showing I_{NCX} events in Sham and MI CM under the conditions shown. (Sham \pm InsP₃: n_{cells}=7,6; N_{pigs}=3,3; MI \pm InsP₃: n_{cells}=13,8; N_{pigs}=4,3). **C.** Example Xt plots and associated voltage traces from MI CM in the presence of Iso ± InsP₃. 4 stimulation signals (CaT) are shown after which arrhythmogenic activity is observed in the form of Ca^{2+} waves and inward currents. **D.** Amplitude of <u>I_{NCX} events</u> in MI CM exposed to Iso ± InsP₃. t-test. E. Analysis of DADs and APs. Example current traces from MI CM exposed to Iso ± InsP₃. The blue arrow heads indicate DAD activity and the red arrowhead an AP. F. Incidence of DADs in Sham and MI CM in the presence and absence of InsP₃. G. Incidence of Ps in Sham and MI CM in the presence and absence of InsP₃. H. Incidence of APs and DADs as a percentage of total events. I. DAD amplitude in Sham and MI ± InsP₃. (Sham ± InsP₃: n_{cells}=1: ,10, N_{pigs}=4,4; ± InsP₃: n_{cells}=9,10; N_{pias}=4,4). Statistical analysis was performed in a hierarchical number with with Tukey post hoc analysis.

Whether alterations in membrane electrom siology contributed to the increased propensity for DAD and AP generation w. s also examined. No significant differences in resting membrane potential (RMP) or action potential duration (APD90) were detected between Sham and MI and no efficet c. InsP₃ was observed (Figure S7A,B). An increase in inward NCX current density have also been reported to underlie the effects of InsP₃ in mouse CM [31]. While a small effect of InsP₃ was detected at extreme negative membrane potentials (-100 and 122 mV) in MI CM, no significant effect of InsP₃ at physiological negative membrane potentials was observed in Sham or MI CM (Figure S7C,D). A significant Nickel-sensitive increase in outward current in MI CM following flash photolysis of InsP₃ was however detected by InsP₃. The increase in outward current elicited by InsP₃ in MI CM was lost in cells dialysed with 10 mM BAPTA to chelate intracellular Ca²⁺ demonstrating the Ca²⁺ dependence of the current activated by InsP₃ (Figure S7D).

3. Discussion

Unravelling mechanisms underlying altered Ca²⁺ handling during pathological cardiac remodelling is key to development of strategies to mitigate against the associated reduced cardiac function and arrhythmias that lead to sudden cardiac death. Here, in CM from a porcine preclinical model of ischemic heart disease, we show that enhanced Ca²⁺ release via InsP₃Rs contributes to enhanced arrhythmogenic activity, one altered Ca²⁺ handling. Specifically, we find that InsP₃R expression is increated in MI and through proximal localisation with and signalling to RyRs at extra dyauic locations, enhance their activity, thereby accelerating Ca²⁺ release during ECC and promoting spontaneous Ca²⁺ release events. These additional Ca²⁺ release ovents engage NCX, inducing membrane depolarisation and AP generation.

While a number of studies describe a rr le for InsP₃ in regulating ECC in rodents, particularly in disease, the relevance of IIC R to pathology in larger mammals, including in human is less investigated, and thus less clear. Although an arrhythmogenic action of InsP₃Rs was proposed in failing human CM over a decade ago, conclusions drawn are undermined by a number of weaknesses [31]. Particularly, conclusions were largely based on extrapolation of effects in mice, InsP₃R function was not directly analysed, and poorly selective InsP₃R antagonists were used to invoke InsP₃R activation as a mechanism underlying arrhythmias provoked by $G\alpha_q$ -coupled agonists. Since Xestospongin C, the InsP₃R inhibitor used, is also a potent inhibitor of SERCA [57], its application would increase SR Ca²⁺ leak and depletion, thereby masking the role of InsP₃Rs. More recently, we described an arrhythmogenic action of InsP₃ in CM and tissue from end-stage HF patients [39]. Unlike in the study of Signore et

al, we directly introduced InsP₃ into CM, avoiding potential off target effects of the inhibitors used to modify effects of GPCR agonists. Using 2-APB at a low concentration shown to have limited off target effects [33], we also demonstrated a potent role of InsP₃ in the arrhythmogenic action of Ang II in tissue wedges. Since our human study was performed on HF cells and tissue, an understanding of the influence of the GPCR/InsP₃R axis during earlier stages of cardiac remodelling during IHD in a large mammal model is still lacking. Insights into arrhythmia mechanisms at these earlier stages of disease, are however, vitally important and clinically relevant since sudden cardiac dea h ir IHD is highly prevalent in early stages after MI before frank HF [20, 40]. In this regard, the pig preclinical model used in this study is highly relevant, sharing features inclining increased arrhythmia propensity, observed during the substantial remodelling after MI in human but in the absence of associated co-morbidities [17].

GPCR agonists that we and others have previously used to stimulate InsP₃ production in CM also engage multiple kinase cascades, which in turn influence the activity of NCX, L-type current, myofilaments and NET with consequences for Ca²⁺ dynamics and contractility [25, 31, 34, 58, 59]. Since we binded in this study to specifically probe the influence of InsP₃R activation on CM physiclogy, we chose to directly activate InsP₃Rs with InsP₃. In line with previous observations in healthy rat and mouse, no effects of InsP₃ on CaTs were detectable [23, 33, 60, 61]. This contrasts with that in rabbit, where InsP₃ causes an increase in CaT amplitude and with our analysis in human HF where CaT amplitude was reduced [29]. The minimal contribution of InsP₃ to the CaT in these healthy pig CM is also consistent with the low abundance of InsP₃Rs in ventricular CM [17, 23, 27, 35, 60].

In large mammals, such as pig, and as in rodents, TT density and coupling of LTCC and RyRs is decreased in pathology leading to decreased synchronicity of Ca²⁺ release with

deleterious consequences for CaT generation and contraction [10, 17, 62]. In this model of early post-MI remodelling, while we cannot rule out LTCC redistribution, we previously did not detect alteration of I_{CaL} in MI and thus do not consider it a major contributor to reduced synchronicity of Ca²⁺ release in the MI CM described here [17]. Significantly, Ca²⁺ release heterogeneity was reduced by InsP₃, to a level approaching that in Sham. In this manner, G α_q -coupled agonists, acting via InsP₃ could elicit a beneficial effect to rescue diseaseassociated decline in cardiac function, analogous to that other provide for β -agonists, which recruits Ca²⁺ release at non-coupled sites [63]. In contrast to the mechanism for β stimulation however, which involves increasing store level, connancing I_{CaL} and sensitising RyR Ca²⁺ release channels via phosphorylation [1], store loading was unaffected by InsP₃, suggesting that RyRs are recruited through at outpernative mechanism involving increased [Ca²⁺] in their local vicinity arising from reigh bouring InsP₃Rs.

Analysis of Ca²⁺ sparks provided deeper insights into the remodelling and mechanism of action of IICR in MI CM, particularly how IICR could recruit non-coupled RyR clusters. Although no difference in Ca²⁺ spark activity was detected between Sham and MI at baseline, in response to ¹, 2^o₃, i/II CM exhibited significantly increased Ca²⁺ spark frequency, which was also apparent as a greater Ca²⁺ spark-dependent leak. The similar Ca²⁺ spark frequency frequencies in MI and Sham CM at baseline, which reflect our previous findings at low pacing frequency [17], would indicate that increased InsP₃R expression in MI CM does not in itself lead to greater basal Ca²⁺ release. Further, SR load was no different between Sham and MI and was not affected by InsP₃, indicating, that in MI CM, IICR did not act to deplete SR Ca²⁺ stores. These results contrast with our findings in human HF and those from a mouse CM transgenically overexpressing InsP₃R2, which show elevated baseline spark activity [39, 64]. In human HF, we found that Ca²⁺ spark frequency and and spark-dependent leak were

increased in the presence of InsP₃ leading to a depletion of the SR Ca²⁺ store and reduced CaT amplitude [39]. A key difference however between CM from MI pig and HF human is the substantially reduced activity and expression of SERCA-mediated Ca²⁺ clearance from the cytosol in human, which we show here is unchanged beteen Sham and MI pig CM. In the face of this reduced SERCA activity, the increased leak induced by InsP₃ is not resequestered ino the store but leads to SR Ca²⁺ depletion. In mouse, increased Ca²⁺ leak in the InsP₃R2 transgenic background was proposed to protect the SR from Ca²⁺ overload and Ca²⁺ wave initiation. This was not seen in human HF however, where Ca²⁺ waves were increased by InsP₃ [39]. Apart from substantial InsP₃F2 crearexpression (12 fold), InsP₃R2 transgenic mice only show mild hypertrophy and do not exhibit other aspects of a HF CM that would contribute to arrhythmogenic act vity such as altered repolarisation currents [64]. As we determined in human but not the rem pig, the InsP₃R transgenic mice also exhibit a substantial reduction in RyR expression, which would alter the relative contributions of these Ca²⁺ channels to Ca²⁺ handling.

In agreement with previous findings in rat and human proposing a role for RyRs in amplifying Ca²⁺ release vio mary₃Rs [23, 35, 39, 65], active RyRs were required for InsP₃ to increase Ca²⁺ spark frequency and spark dependent leak in MI CM. The premise of this hypothesis is that Ca²⁺ release from InsP₃Rs in the vicinity of co-localised RyRs leads to their sensitisation. Modelling data suggests that [Ca²⁺] in the 10s of μ M would be required at the location of RyRs for this mechanism to be effective [56]. Given that Ca²⁺ steeply declines with distance from its peak in the 100s of μ M at a Ca²⁺ channel mouth (e.g. RyR), for any neighbouring channel to influencing a RyR, it would need to be highly proximally localised [51, 56]. While this is established for RyRs in a cluster, single molecule data that places InsP₃Rs sufficiently close to RyRs is lacking. Here and elsewhere, we and others have shown

by confocal imaging localization of InsP₃Rs to the Z-line, where they colocalize to a certain extent with RyRs [23, 66, 67]. While in our recent analysis in human HF, we used STED microscopy to examine InsP₃R-RyR relative distributions [39], the resolution of this modality of microscopy may not be sufficient to resolve between clusters of different scales and to establish channel proximities. To overcome these issues, we employed DNA-PAINT super resolution imaging, which can achieve sub 10 nm resolution [42]. We definitively identified InsP₃Rs residing in amongst RyRs in a cluster, thereby en hling Ca²⁺-mediated channel crosstalk. The number of RyRs per cluster described here was similar to that previously described using super resolution approaches [14, 42, 47, 68]. However, although as previously reported in HF [14], the number of RyR channels per cluster were not increased in post-MI pig CM, we found an increased proportion of larger RyR clusters in MI. Cluster size has important consequences for spark generation in that larger clusters show greater fidelity in spark generation and thus make a greater contribution to spark frequency [53, 69]. In disease however, decreased organization and/or density of RyRs within the cluster leads to a reduced Ca²⁺ dependent coupling between RyRs and reduced kinetics of Ca²⁺ release [14, 70]. The increased proportion of RyR clusters comprising 2 or more channels together with the lack of change in RyR expression (established by immunoblot) in MI that we found are consistent with greater numbers of RyR in the clusters described. The identification of InsP₃Rs residing amongst RyRs in these clusters may represent a mechanism to rescue intra as well as inter cluster coupling, irrespective of whether the neighbouring clusters contain InsP₃Rs. Indeed, our modelling data shows that by providing extra Ca²⁺, by means of an increased number of InsP₃Rs, particularly in the immediate vicinity of RyRs, the coupling of stochastically opening RyRs in a cluster is increased, resulting in increased likelihood of a Ca²⁺ spark. Our modelling data also suggest that the presence of InsP₃Rs

within a RyR cluster serves to sensitise the cluster to Ca²⁺ diffusing from neighbouring coupled clusters during their activation by an electrical trigger. Consistent with these findings, our cellular analysis showed synchronization of Ca²⁺ release during the CaT by InsP₃ in MI. Further supporting this hypothesis, InsP₃ preferentially augmented Ca²⁺ sparks and Ca²⁺ release kinetics form non-coupled sites in MI. Moreover, InsP₃R were enriched at noncoupled sites in MI CM. In this way, enhanced IICR could act in positive as well as a deleterious role in disease. These conclusions regarding InsP₃I -RyR crosstalk agree with that previously hypothesised but were not substantiated due to he; bsence of super resolution imaging of InsP₃R distributions and the modelling apriloactes that we now report in this study [23, 35]. This preferential regulation of a^{2+} release at non-coupled sites is reminiscent of that which we previously des ribed in MI pig CM, where increased Ca²⁺ sparks at non-coupled sites were deprind int upon CaMKII activity and reactive oxygen species [17]. Indeed, consistent with the known phosphorylation of RyRs in cardiac pathology, we here observed increased levels of CaMKII phosphorylated RyRs in cardiac tissue from MI pigs. While our modelling data suggest that CaMKII is not required for Ca²⁺ release via InsP₃Rs to influence neighbouring RyR, CaMKII phosphorylated RyRs at noncoupled sites likely create a sensitised substrate upon which Ca²⁺ release via InsP₃Rs can preferentially act to augment RyR-mediated Ca²⁺ release at these sites. CaMKII can also phosphorylate InsP₃R2 [71], although since this phosphorylation event suppresses InsP₃R activity, it is unlikely to contribute to the increased InsP₃R activity observed in MI CM, which we describe in this manuscript. Given the importance of CaMKII phosphorylation of RyRs in cardiac pathology, contributing to enhanced Ca^{2+} leak and induction of arrhythmias [72, 73], future studies to assess the interaction between IICR, CaMKII, RyRs and Ca²⁺ release will be important to perform.

Arrhythmogenic electrophysiological activity was significantly increased by InsP₃ in MI CM. This effect was not due to a substantial increase in inward NCX current as previously reported or that is induced by GPCR agonists [31, 74]. Rather, and in line with findings in human HF [39], our data indicate that dysregulation of intracellular Ca²⁺ handling, including increased spontaneous Ca^{2+} release events via InsP₃R-RyR coupling, contribute through activation of NCX to the generation of arrhythmogenic DADs and APs [19]. Previously, we have shown that Ca²⁺ waves underlying arrhythmogenic octivity in MI arise from Ca²⁺ release at hyperactive non-coupled RyR clusters [17] The increased Ca2+ sparks at noncoupled sites together with the greater arrhythmogenic activity in MI CM in the presence of InsP₃ would suggest a similar model. Notably, and underlining the importance of IICR in disease, InsP₃ promoted the almost exc¹ usiv generation of APs in MI CM, whereas in Sham, DADs were more prevalent. Supporting increased activation of NCX as a mechanism by which InsP₃ crosstalk with RyRs envalue APs, DAD amplitude was greater in MI in the presence of InsP₃. The greater notential for InsP₃ to promote APs in MI also likely involves an alteration in repolarisation currents such as in I_{K1} , which we have previously reported in this model,[20]. Together, these data re-enforce the notion that mechanisms involving dysregulated Ca²⁺ release leading to membrane depolarisation via NCX are important contributors to the potentially fatal arrhythmic activity associated with remodelling after MI [19].

Conclusions. Here, we have demonstrated increased activity of the $InsP_3/InsP_3R/Ca^{2+}$ signalling pathway in ventricular CM from pig during remodelling after MI. This analysis in pig CM from early stages of IHD is an important step in understanding the role of

GPCR/InsP₃-signalling in disease evolution and occurrence of arrhythmias in the progression to HF in human. Our analysis also sheds light on the involvement of $G\alpha_q$ coupled GPCRs including AngII, ET-1 and catecholamines in cardiac pathophysiology. Not only are these mediators found to be elevated in cardiac disease, inhibitors of their activity, for example of the renin angiotensin system, decrease the incidence of arrhythmias in animal models [75] and in patients with IHD [76]. Our findings therefore identify an important and diseaserelated signalling pathway that may be targeted pharm cologically to prevent fatal arrhythmias.

Author Contributions

Conceptualisation Ideas; X.J, K.S, H.L.R. *Data* curation and Formal analysis; X.J., A.M., A.T., K.D., E.D., R.D.P., M.A., P.C., C.C., V.K. K.S., H.L.R. Investigation; X.J., A.M., A.T., K.D., E.D., R.D.P., M.A., P.C., C.C., V.R., K.S., H.L.R. Writing - Original Draft Preparation; X.J., H.L.R. Writing - Review & Editing; U authors. Resource provision; K.S., P.C., C.S., V.R., H.L.R. Supervision Oversight and Lectership; K.S., P.C., C.S., V.R., H.L.R.

Declaration of Competing interests

The authors declare no conflicts of interest.

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Methods

Animal model

A porcine model of myocardial remodelling after myocardial infarction (MI) and chronic ischemia as described previously was used [46]. Presence of infarct and expected alterations in hemodynamics properties determined by MRI are presented in Table S1. Animals were housed and treated according to the European Directive 2010/63/EU with protocols approved by the local ethical committee (Ethische Commissie Dierproeven, KU Leuven).

General Methods

Detailed reagents, methods for CM isolation, Ca²⁺ imagine with electrophysiology, immunoblotting, DNA-PAINT microscopy, RT-qPCR, mathematical modelling and statistical analysis are provided in the Extended Methods and in [35, 46, 48].

Solution

Graphical Abstract: Mechanism underlying pro-arrhythmic action of InsP₃**induced Ca**²⁺ **release.** Remodelling of cardiomyocyte TT and InsP₃R expression in healthy and post-MI heart is shown. **A.** in healthy heart, InsP₃R expression is low, and electrical stimulation (black arroes) induces Ca²⁺ entry via LTCC that activate Ca²⁺ release via RyRs on coupled dyads to generate a Ca²⁺ transient. **B.** In MI, TT are lost leading to increased non coupled RyRs and slower CaT are observed. InsP₃Rs expression is also increased. **C.** As in B but in the presence of increased InsP₃, Ca²⁺ release via RyRs, including non coupled RyRs, is increased leading to augmented Ca₂₊ transient kinetics and Ca₂₊ waves, which engage NCX, and if of sufficient magnitude, reach threshold for AP generation. Smaller Ca₂₊ waves produce DADs.



Highlights:

- InsP₃R expression is upregulated in a pig model of ischemic heart disease.
- InsP₃ increases Ca²⁺ sparks via RyR in post-MI pig cardiomyocytes.
- Ca²⁺ release via InsP₃Rs induces arrhythmogenic activity in MI pig cardiomyocytes.
- Nanoscale imaging reveals co-habitation of InsP₃Rs and RyRs within clusters.
- Mathematical modeling supports role for InsP₃Rs in RyR cluster activation in ECC.

Solution