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**Zusammenfassung des wissenschaftlichen Inhalts**

(Stephan E. Lehnart)

Calcium-induzierte Calcium-Freisetzung (CICR) beschreibt einen zentralen Mechanismus physiologischer Regulation von Herz-, Gehirn- und Pankreaszellen. Während CICR in ventrikulären Herzmyozyten (Kardiomyozyten) durch transversale Membraninvaginationen (T-tubuli) vermittelt wird, sind die Aktivierungsmechanismen in atrialen Kardiomyozyten weitgehend unklar. Brandenburg et al. identifizieren nun einen bislang unbekannten Mechanismus basierend auf ungewöhnlich grossen Membranstrukturen (axiale Tubuli) als Grundlage für ein neues Modell atrialer Erregungs-Kontraktions-Koppelung. Axiale Tubuli formen ausgedehnte Nanodomänen-Kontakte mit dem sarcoplasmatischen Reticulum (SR). Die axialen SR-Junktionen enthalten eine besonders grosse Zahl von Ryanodin-Rezeptor (RyR2) Calcium-Freisetzungskanälen, die eine besonders schnelle axiale Kontraktion in

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Axial tubule junctions control rapid calcium signaling in atria

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The canonical atrial myocyte (AM) is characterized by sparse transverse tubule (TT) invaginations and slow intracellular Ca2+ propagation but exhibits rapid contractile activation that is susceptible to loss of function during hypertrophic remodeling. Here, we have identified a membrane structure and Ca2+-signaling complex that may enhance the speed of atrial contraction independently of phospholamban regulation. This axial couplon was observed in human and mouse atria and is composed of voluminous axial tubules (ATs) with extensive junctions to the sarcoplasmic reticulum (SR) that include ryanodine receptor 2 (RyR2) clusters. In mouse AM, AT structures triggered Ca2+ release from the SR approximately 2 times faster at the AM center than at the surface. Rapid Ca2+ release correlated with colocalization of highly phosphorylated RyR2 clusters at AT-SR junctions and earlier, more rapid shortening of central sarcomeres. In contrast, mice expressing phosphorylation-incompetent RyR2 displayed depressed AM sarcomere shortening and reduced in vivo atrial contractile function. Moreover, left atrial hypertrophy led to AT proliferation, with a marked increase in the highly phosphorylated RyR2-pS2808 cluster fraction, thereby maintaining cytosolic Ca2+ signaling despite decreases in RyR2 cluster density and RyR2 protein expression. AT couplon “super-hubs” thus underlie faster excitation-contraction coupling in health as well as hypertrophic compensatory adaptation and represent a structural and metabolic mechanism that may contribute to contractile dysfunction and arrhythmias.

Introduction

Electrical and contractile dysfunction of the atria are frequent components of cardiac disease development, often culminating in atrial fibrillation (AF), a leading cause of ischemic stroke, which is predicted to triple in prevalence by 2050 (1). Among the risk factors associated with atrial dysfunction, hypertension is present in 60% to 80% of European patients (2). Using rapid pacing, key aspects of atrial remodeling and AF were reproduced in large animal models: whereas electrical and ionic changes occurred within minutes, intracellular Ca2+ overload was normal by 48 hours (3). Recently, high atrial pacing rates were shown to induce Ca2+-signaling silencing, preventing intracellular Ca2+ overload, albeit at the cost of depressed contractile function (4). Hypertrophy with depressed contractile function occurred as early as 2 days after pacing, in the absence of fibrosis or dilation in canine atria (5). These studies support the therapeutically relevant notion that electrical and metabolic disease mechanisms may cause contractile atrial dysfunction relatively early, preceding AF.

Atrial myocytes (AM) are distinguished from ventricular myocytes (VM) by their smaller diameter, which may explain why no or few transverse tubules (TT) could support adequate AM function (6–8). In contrast, VMs are characterized by high TT density and regularity in normal hearts, a feature lost in heart failure (HF) (9, 10). Given the regular presence of TTs at sarcomeric Z-lines, TT density is thought to determine uniform Ca2+ release in VMs (11). Paradoxically, despite sparse irregular TT components, contractile activation of atrial muscle from small and large animal species occurs faster than in ventricular tissue (12). Hence, there is a gap in understanding the mechanisms that underlie rapid activation of atrial contraction. In addition, even though atrial dysfunction and arrhythmias are recognized as a leading cause of cardiac disease burden, fundamental details about the subcellular mechanisms involved remain unclear (13).

In the absence of abundant TT membrane invaginations, AMs are thought to initiate excitation-contraction coupling (ECC) through Ca2+ release units (CRUs) at the surface sarcotubulemma, each including approximately 6 L-type Ca2+ channels (LCCs/Cav1.2) opposite a cluster of approximately 50 RyR2 Ca2+ release channels, separated by a subspace of approximately 15 nm width (14, 15). In this model, Ca2+ transients are activated through subsarcolemmal CRUs, leading initially to peripheral elevation of Ca2+, which travels toward the AM center through propagated Ca2+-induced Ca2+ propagation silencing, preventing intracellular Ca2+ overload, albeit at the cost of depressed contractile function (4). Hypertrophy with depressed contractile function occurred as early as 2 days after pacing, in the absence of fibrosis or dilation in canine atria (5). These studies support the therapeutically relevant notion that electrical and metabolic disease mechanisms may cause contractile atrial dysfunction relatively early, preceding AF.

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Results

AMs contain cell-wide TAT membrane networks. Although earlier electron microscopy–based (EM-based) representations of AMs (6), recent confocal data, and computational models (19) suggest that there are few, if any, TT invaginations, experimental studies have increasingly highlighted the presence of atrial TAT networks in several mammalian species, including humans (20–22). To clarify this apparent contradiction, we used membrane-preserved workflows developed for high-resolution live-cell imaging (9, 18) that revealed abundant TAT membrane structures in murine AMs (Figure 1A and Supplemental Figure 1A; supplemental material available online with this article; doi:10.1172/JCI88241DS1). We identified intact TAT networks in 100% of the AMs examined (n = 29), thus confirming the presence of preserved membrane structures. Using skeletonized images, we visualized individual TAT components and analyzed their orientations (Figure 1A, magenta). In confocal sections, the cell-wide TAT network density in AMs was lower (by ~24%) than in VMs (isolated from the same hearts), attributable chiefly to a significantly lower presence of TT components (Figure 1B), which dominate TAT networks in VMs (Supplemental Figure 1B). Despite smaller dimensions (Supplemental Figure 2) and a low TT abundance, mouse AMs contained significantly more ATs than VMs (Figure 1B). Thus, while cell type–specific differences in the abundance of TTs explain a moderately lower TAT network density, AMs showed a significantly higher abundance of AT components than did VMs.

Since confocal imaging suggested unusually large AT dimensions (Figure 1C, illustration), we used live-cell stimulated emission depletion (STED) super-resolution microscopy (nanoscopy) to assess the width of intact ATs and TTs (Supplemental Figure 3). At width (293.3 ± 7.7 nm) was significantly greater than TT width in AMs (Supplemental Table 1) and approximately 50% larger in AMs than in VMs (Figure 1C and Supplemental Table 1). This was further investigated using ET and 3D segmentation (23) of semithick sections (280 nm) of high-pressure-frozen AMs (Supplemental Figure 4). ET confirmed the presence of voluminous ATs containing multiple junctions with the sarcoplasmic reticulum (SR) (Figure 1D). The color rendering in Supplemental Figure 4 highlights extensive AT-SR junctions with a gap width of 15 nm or less that contained RyR2 densities (red) covering large regions of the AT surface area (Figure 1, D and E). While not perfectly circular in the cross section, we observed no difference in the mean eccentricity of ATs and TTs (0.62 ± 0.05, n = 11 and 0.64 ± 0.04, n = 9, respectively; P = 0.7). The measured volume/surface area ratio and width were significantly higher for ATs than for TTs in AMs (Figure 1F), consistent with the STED measurements. These independent techniques confirmed the presence of central-axial membrane structures of unusually large size and surface area that were distinct from those of VMs, with extensive SR junctions.

AT structures couple excitation to rapid intracellular Ca2+ release. TT invaginations, continuous with the surface membrane, contain LCCs juxtaposed with RyR2 clusters in the SR, and similar
structural features were evident for ATs in our EM AM data (Figure 1, D and E). To investigate the subcellular relations between RyR2 channels and AT structures, we immunolocalized RyR2 and caveolin 3 (Cav3). While colocalization of Cav3 and RyR2 is generally considered to indicate the presence of CRUs at SR contact sites (24), we observed no regular striated distribution due to sparse TT elements in AMs (Figure 1G). Instead, robust Cav3-positive AT structures were abundant in AMs from both mouse and human hearts (Figure 1G). Furthermore, Cav1.2 channel clusters were frequent and regularly associated with AT structures (Figures 1, H and I). Analysis of Cav1.2 clusters confirmed a substantial Tat-specific density, and the number of component-specific Cav1.2 clusters was significantly higher in AT components than in TT components (Figure 1J). Thus, central RyR2 clusters, separated by a subsurface gap from the peripheral RyR2 clusters described previously (19), colocalize in AMs with abundant AT structures that contain a high number of Cav1.2 clusters.

For AT structure-function analysis, we measured Ca²⁺ transients in AM using fluo-4 and the membrane-intercalculating dye di-8-ANEPPS using protocols established previously (9, 18). Transversal line scanning during electrical stimulation revealed robust Ca²⁺ rises, originating from AT sites (Figure 1K). Surprisingly, the spatiotemporal F25 profile (time point at which 25% of the maximal fluorescence is reached) indicated earlier Ca²⁺ transient onset at ATs than at surface (S) sarcolemmal sites (Figure 1K). To confirm this, we analyzed Ca²⁺ signal onset at AT, S, and off-membrane cytosolic (CY) locations by assessing the first derivative of fluorescence intensity (dF/dt) and determined the following latency profile: AT < S < CY (Figure 1L). In contrast, imaging planes that excluded visible AT structures showed principally different, U-shaped Ca²⁺ transient onset patterns (data not shown). This result is consistent with the steep S < CY latency gradients we observed (Figure 1L) and agrees with previous observations of U-shaped Ca²⁺ transients (4, 17, 25). To reproduce early Ca²⁺ signal onset at AT structures under direct voltage control, we patch clamped AMs (1 mM EGTA in the patch; Figure 1M). Analysis of Ca²⁺ signal onset by dF/dt during depolarization to 0 mV confirmed the significant AT < S latency (Figure 1N). In summary, electrically evoked Ca²⁺ transients originate deep inside AMs, from AT structures, and do so approximately 2 times earlier than Ca²⁺ transients on the surface membrane. The discovery of more rapid central Ca²⁺ release supports the presence of continuous electrical connectivity of ATs with the surface sarcolemma, presumably via sparse TTS as a conduit for ECC activation.

Human left atrial myocardium exhibits axial RyR2 cluster morphologies at ATs. To assess whether AT membrane structures are associated with junctional RyR2 clusters in human tissue as well, we developed dual-color immunolabeling protocols for STED nanoscopy of left atrial (LA) tissue sections. Cav3-labeled sections showed highly ordered rod-shaped AMs aligned with capillaries in atrial tissue (Figure 2A). Interestingly, AMs with robust intracellular Cav3 staining showed that multiple central ATs regularly intersected with transversal RyR2 cluster striations (Figure 2A), similar to our observations in isolated human and mouse AMs (Figure 1G). Importantly, higher magnifications inside tissue-embedded AMs showed distinct, axially extended RyR2 cluster morphologies, which tightly embraced the Cav3-labeled ATs (Figure 2B). Thus, central AT-associated (but not off-membrane CY) RyR2 clusters revealed a distinct morphology in human LA sections, in keeping with the possibility of unique axial couplon functions such as for the rapid Ca²⁺ release onset observed in murine AMs.

Differential regulation of atrial RyR2 clusters. Our finding that local Ca²⁺ release occurs earlier at AT sites suggests potentially unknown local regulatory mechanisms. In VMs, TT membranes were previously shown to contain β-adrenergic receptor (β-AR) and adenylyl cyclase (AC5/6) signaling structures, increasing cAMP concentrations locally (26, 27). Furthermore, cAMP sensing through PKA is directly tethered to each subunit of the homotetra-

Figure 2. ATs and junctions with RyR2 clusters are common in human tissue. Dual-color STED nanoscopy of 2 human LA tissue sections colimmunostained for Cav3 and RyR2. (A) Tissue section shows AMs aligned with capillaries (asterisks indicate the lumens). Magnified images (of red boxed area) show 1 Cav3-labeled AT membrane structure (arrowhead), which intersects with 6 transversal rows of RyR2 clusters delimited by the surface membrane on top. Scale bars: overview 10 μm; magnifications, 2 μm. (B) Intracellular high-power magnifications showing 2 parallel, axially aligned RyR2 cluster tracks, perpendicularly intersecting 6 transversal RyR2 rows. Arrowheads identify exemplary RyR2 clusters with typical axial couplon architectures. RyR2 clusters with elongated axial morphologies and Cav3-labeled structures are tightly spaced next to each other, indicating a functional role of AT-associated SR junctions. STED images are representative of 5 human LA samples. Scale bar: 2 μm.
Figure 3. Differential regulation of RyR2 cluster phosphorylation. (A) AMs coimmunostained for RyR2- and PKA-phosphorylated RyR2-pS2808. WT: highphos RyR2 clusters (yellow) aligned in central-axial string-of-pearls, intersecting transverse striations of lowphos RyR2 clusters (green); RyR2-S2808A−/−: zero PKA phosphorylation of RyR2 clusters, confirming phospho-epitope–specific labeling; WT + H89: attenuated highphos RyR2 cluster signals. Scale bars: 10 μm; magnified (×4) regions are indicated by yellow brackets. (B) Histogram showing bimodal frequency distribution of in situ pS2808/RyR2 normalized cluster signals in untreated WT AMs. The main peak indicates abundant lowphos versus fewer highphos RyR2 clusters represented by the shoulder. n = 22 AMs. (C) AMs colabeled for Cav3 and RyR2-pS2808. Highphos RyR2-pS2808 clusters alternated with Cav3 clusters in central AT structures. Scale bar: 10 μm; magnification ×4. (D) Image segmentation confirming highphos RyR2-pS2808 clusters aligned with Cav3-labeled TAT structures. (E) WT, RyR2-S2814A−/−, and WT AMs treated with the CaMK inhibitor AIP were coimmunostained for RyR2 and the CaMK-phosphorylated epitope RyR2-pS2814. Note the central highphos RyR2 clusters (yellow) versus lowphos RyR2 clusters (green). AMs from CaMK phosphorylation–incompetent RyR2-S2814A−/− mice showed no CaMK phosphorylation in situ. Magnification ×4. (F) Frequency histogram of the pS2814/RyR2 cluster distribution indicating abundant lowphos versus few highphos RyR2 clusters in control AMs. n = 19 AMs. (G) AMs coimmunostained for Cav3 and RyR2-pS2814. Highphos RyR2 clusters were associated with Cav3-labeled AT structures as confirmed by image segmentation (H). Scale bar: 10 μm, magnification ×4. (I) STED images comparing immunolabeled central RyR2 clusters in VMs versus AMs. Atrial RyR2 clusters identified by red circles show a higher density on Z-lines and a trend toward overall shorter NNDs. Images are representative of 18 VMs and 19 AMs. Scale bars: 500 nm. Box and whisker plot: boxes show lower and upper quartiles; whiskers represent the 5th and 95th percentiles. IF, immunofluorescence. (J) Confocal live AM imaging with di-8-ANEPPS and fluo-4 showing large AT-localized Ca2+ sparks occurring repeatedly. CaT, final 1-Hz pacing evoked Ca2+ transient; red triangles indicate AT-associated Ca2+ sparks; white triangles indicate surface membrane–associated Ca2+ sparks.
meric RyR2 channel complex (28). Since PKA phosphorylation of Ser 2808 and CaMK phosphorylation of Ser 2814 are powerful positive regulatory mechanisms of RyR2 sensitivity for Ca\textsuperscript{2+}-dependent activation (28–30), we investigated the nature of subcellular RyR2 phosphorylation using novel in situ immunofluorescence strategies.

Using Abs specific to either PKA-phosphorylated RyR2-pS2808 or CaMK-phosphorylated RyR2-pS2814, we identified distinct populations of highly phosphorylated (highphos) RyR2 clusters in AMs (Supplemental Figure 5A), but not in VMs from the same WT mouse hearts (Supplemental Figure 5B). In AMs, central highphos RyR2-pS2808 clusters were axially aligned in “string-of-pearls” arrangements, which regularly intersected with minimally phosphorylated (lowphos) clusters in transversal striations at Z-lines (Figure 3A). Molecular labeling specificity was confirmed by 2 independent approaches. First, RyR2-pS2808 epitope detection was obstructed by the blocking, but not by the control peptide, confirming specific phospho-Ab binding (Supplemental Figure 6). Second, AMs from mice expressing a PKA phosphorylation–incompetent mutation (Ryr2\textsuperscript{2808A/2808D}; hereafter referred to as RyR2-S2808A+/+–knockin mice) showed complete ablation of RyR2-pS2808 in situ cluster signals (Figure 3A).

Since local PKA activity, apparent through highphos RyR2 cluster phosphorylation, occurred in the absence of p-AR stimulation, we hypothesized that competitive pharmacological PKA inhibition would attenuate TAT-associated PKA activity. Indeed, live AMs treated with the cAMP-specific PKA inhibitor H89 (50 μM) showed blunted highphos RyR2 cluster signals (Figure 3A). This result indicates significant cAMP-dependent local PKA activity, sufficient to sustain PKA phosphorylation of highphos RyR2 clusters. Collectively, these data suggest a highly compartmentalized form of atrial RyR2 cluster regulation, which may functionally differentiate highphos from lowphos RyR2 channel populations.

To quantify the apparently heterogeneous in situ cluster phosphorylation, we analyzed the frequency distribution of the pS2808/RyR2 normalized cluster signals. Untreated WT AMs showed a major pS2808/RyR2 peak, indicating abundant lowphos clusters (Figure 3B). In contrast, highphos pS2808/RyR2 signals occurred at much lower frequencies, indicated by a right-skewed distribution (Figure 3B). VMs isolated from the same hearts showed a markedly different distribution (Supplemental Figure 5C). To explore whether axially aligned RyR2 highphos clusters are associated with AT structures in AMs, we coinmunostained AMs for Cav3 and RyR2-pS2808. This confirmed that Cav3-positive ATs were closely aligned with highphos RyR2 signals in string-of-pearls arrangements (Figure 3C). Furthermore, triple staining of Cav3, RyR2, and RyR2-pS2808 confirmed that RyR2 clusters that colocalized with Cav3-labeled AT structures were indeed highphos clusters (Supplemental Figure 7). In contrast, lowphos RyR2 clusters in transverse striations did not colocalize with AT or TT components.

Using a corresponding image segmentation strategy, we analyzed the spatial association of highphos versus lowphos RyR2-pS2808 clusters with Cav3-labeled TAT structures as the nearest neighbor distance (NND) (Supplemental Figure 8A). The NND was significantly smaller for highphos than for lowphos RyR2-pS2808 clusters (Supplemental Figure 8B). We visualized the segmentation analysis output and compared corresponding image regions and found that central highphos RyR2-pS2808 clusters were tightly associated with Cav3-positive AT structures (Figure 3D). While highphos RyR2-pS2808 clusters represented the smaller population in untreated AMs (Figure 3B), our analysis indicated their preferential association with AT structures, consistent with rapid junctional Ca\textsuperscript{2+} release at AT sites.

In addition, CaMK phosphorylation of the RyR2-S2814 site was identified as a mechanism of AF susceptibility in mice (31, 32), and both RyR2-S2808 and RyR2-S2814 phosphorylation changes were identified in atrial samples from patients with AF (33, 34). Consequently, we investigated RyR2-pS2814 phosphorylation in situ, finding axially aligned highphos clusters (Figure 3E). Confirming phospho-epitope–specific labeling, AMs from CaMK phosphorylation–incompetent RyR2\textsuperscript{S2814A/S2814A}–knockin mice (referred to herein as RyR2-S2814A+/+–knockin mice) showed complete ablation of RyR2-pS2814 cluster signals (Figure 3E). Furthermore, pretreatment with the CaMK-specific autoinhibitory peptide (AIP, 5 μM) blunted the RyR2-pS2814 highphos RyR2 cluster signal in situ (Figure 3E).

The pS2814-RyR2 frequency histogram showed a major peak and right-skewed distribution, indicating abundant lowphos and few highphos clusters (Figure 3F). Coinmunostaining of AMs for Cav3 and RyR2-pS2814 showed that highphos RyR2-pS2814 clusters aligned with AT structures (Figure 3G). While image segmentation of Cav3-labeled TAT structures and highphos RyR2-pS2814 clusters confirmed this association, some highphos clusters showed no TAT association (Figure 3H). While statistical analysis of RyR2-pS2814 to Cav3 spacing confirmed a specific correlation (Supplemental Figure 8C), NND analysis did not reach statistical significance between highphos and lowphos RyR2-pS2814 clusters using analogous thresholding procedures applied to RyR2-pS2808 clusters (Supplemental Figure 8D). Taken together, our data on untreated AMs suggest that locally increased CaMK activity can maintain RyR2-S2814 cluster phosphorylation at both TAT structures and other sites. Thus, both CaMK and PKA phosphorylate AT-associated RyR2 clusters in AMs, albeit through distinct molecular pathways converging locally in highphos clusters.

**Large elemental Ca\textsuperscript{2+} signals at AT sites.** Local control of elemental Ca\textsuperscript{2+} release (Ca\textsuperscript{2+} sparks) in CRUs may depend on the spatial interrelation between RyR2 clusters and AT membranes, as proposed previously (7). Using STED nanoscopy, we compared intracellular RyR2 cluster densities between AMs and VMs (Figure 3I). Interestingly, there was a nonsignificant trend (Mann-Whitney U test) toward a smaller yet highly variable NND between RyR2 clusters in AMs, suggesting a potentially higher central cluster density (Figure 3I). Since AT structures are significantly larger in AMs, their membrane surface may couple more and/or larger RyR2 clusters during local Ca\textsuperscript{2+} release than do TTs, as supported by ET and 3D segmentation (Figures 1, D and E).

To explore this hypothesis, we imaged fluo-4–loaded AMs costained with di-8-ANEPPS to determine whether Ca\textsuperscript{2+} sparks originate from distinct RyR2 cluster sites. Transversal line scanning during 1-Hz field pacing confirmed earlier central Ca\textsuperscript{2+} transient onset (analogous to Figure 1K), followed by unusually large post-pacing Ca\textsuperscript{2+} macro-sparks fired repeatedly from the same AT site (Figure 3J). Apparently, Ca\textsuperscript{2+} sparks originating at the surface membrane were smaller. Taken together, CRUs at AT structures
Figure 4. Ca\(^{2+}\) macro-sparks at AT sites are reproduced by RyR2 cluster modeling. (A) The final of 5 consecutive Ca\(^{2+}\) transients of a voltage-clamped AM (1-Hz pacing) during depolarization from –75 to 0 mV was magnified (corresponding dashed-line boxes) to visualize early Ca\(^{2+}\) signal onset locally. Block arrows indicate different Ca\(^{2+}\) spark locations at –50 mV, including off-membrane cytosolic sites (CY). Top 2D traces show Ca\(^{2+}\) spark events at AT, S, and CY locations, and examples highlighted by color are magnified by corresponding 3D surface plots (bottom panels): macro-spark (pink), ember (green), classic Ca\(^{2+}\) sparks (blue), and regenerative macro-sparks. Dot plot correlating Ca\(^{2+}\) spark full duration at half maximum (FDHM) and full width at half maximum (FWHM) to segregate classic (blue) from macro-sparks (green) and embers (pink). (B) Box plots comparing Ca\(^{2+}\) spark amplitude (Amp), FWHM, and FDHM in different locations. *P < 0.05, by ANOVA. (C) Atrial Ca\(^{2+}\) spark model composed of 1 transversal RyR2 cluster row. The indicated AT and S structures were associated with junctional highphos RyR2 clusters. Each cluster contained 35 RyR2 channels stochastically activated at –50 mV; both highphos and lowphos RyR2 clusters initiated SR Ca\(^{2+}\) release. Mathematical modeling generated the stochastic open time profiles of individual RyR2 channels for each cluster, which was used to calculate line-scan images of Ca\(^{2+}\) sparks with confocal resolution. Ca\(^{2+}\) sparks presented as 2D-flattened images (top) are highlighted by color, and the corresponding 3D surface plots are shown (bottom): classic Ca\(^{2+}\) sparks (blue), macro-spark (green), ember (pink), and regenerative macro-sparks (gray). \(N_{\text{RyR}}^{\text{c}}\) number of open RyR2 channels. (D) Box and whisker plots comparing Ca\(^{2+}\) spark amplitude, FWHM, and FDHM as the average modeling output. (E) Comparing the aggregate properties of calcium sparks show no significant differences between the AM imaging results and spark modeling output. *P < 0.05, by ANOVA (B and D). Box and whisker plots: boxes show lower and upper quartile; whiskers show the 5th and 95th percentiles.
Repeatedly from AT sites (Figure 4A). In contrast, Ca\(^{2+}\) sparks originating from non-AT–coupled sites appeared smaller (Figure 4A). 2D traces and 3D surface plots highlight the different types of Ca\(^{2+}\) spark events: macro-sparks (pink), long-lasting sparks (embers; green), classic sparks (blue), and regenerative macro-spark Ca\(^{2+}\) propagation (Figure 4A, compare color-rendered top and bottom panels). Central Ca\(^{2+}\) macro-sparks, apparently sequentially recruited, indicated that several RyR2 clusters may contribute to the same Ca\(^{2+}\) release event. The dot plot summarizes the differences can generate Ca\(^{2+}\) macro-sparks, suggesting a potential mechanism of local AT-dependent Ca\(^{2+}\) signaling in AMs.

To control LCC activity, voltage-clamped AMs underwent 5 depolarization cycles to equilibrate the SR Ca\(^{2+}\) load, followed by imaging of spontaneously occurring Ca\(^{2+}\) sparks. We used dF/dt onset analysis to identify early Ca\(^{2+}\) release sites (as shown in Figure 1M) and classified the principal Ca\(^{2+}\) spark locations accordingly. Following repolarization to –50 mV, Ca\(^{2+}\) macro-sparks, characterized by high amplitude, large width, and long duration, originated repeatedly from AT sites (Figure 4A). In contrast, Ca\(^{2+}\) sparks originating from non-AT–coupled sites appeared smaller (Figure 4A). 2D traces and 3D surface plots highlight the different types of Ca\(^{2+}\) spark events: macro-sparks (pink), long-lasting sparks (embers; green), classic sparks (blue), and regenerative macro-spark Ca\(^{2+}\) propagation (Figure 4A, compare color-rendered top and bottom panels). Central Ca\(^{2+}\) macro-sparks, apparently sequentially recruited, indicated that several RyR2 clusters may contribute to the same Ca\(^{2+}\) release event. The dot plot summarizes the differences.
ent types (grouped by colors, as above) of Ca\(^{2+}\) signals originating from AT sites: classic sparks, macro-sparks, and embers (Figure 4A). Comparison of the mean width and duration at different subcellular locations showed that Ca\(^{2+}\) macro-sparks originated more frequently from ATs (Figure 4B). This suggests that AT junctions can repeatedly generate Ca\(^{2+}\) macro-sparks and embers, generally coordinating larger local Ca\(^{2+}\) release events.

Quantitative RyR2 cluster modeling supports local and global AT coupling. To integrate the observed local and global Ca\(^{2+}\) signals mechanistically, we used an existing mathematical ECC model that incorporates the latest biophysical information regarding local RyR2 cluster dynamics (35). This model was modified to account for our data on AT-associated Ca\(^{2+}\)-signaling functions and discrete localizations of highphos RyR2 clusters at AT and S sites (Figure 4C). We built a 3D model of CICR within a single AM sarcomeric Z-region (M-line to M-line) containing CRUs with either lowphos or highphos RyR2 clusters (n = 35 RyR2 channels each), randomly arranged transversally to simulate AT-associated Ca\(^{2+}\) signals recorded in line-scan mode. RyR2 clusters, spontaneously activated by the local diadic subspace [Ca\(^{2+}\)]_J concentration or by LCC openings at the −50 mV holding potential, produced corresponding Ca\(^{2+}\) sparks and localized depletions in junctional SR Ca\(^{2+}\) concentration ([Ca\(^{2+}\)]_J) termed Ca\(^{2+}\) blinks (Supplemental Figure 9). Detailed visualization of the simulated Ca\(^{2+}\) spark behaviors and their spatiotemporal dimensions showed that Ca\(^{2+}\) macro-sparks and embers could be elicited in the model from AT highphos RyR2 clusters (Figure 4C). Systematic Ca\(^{2+}\) spark comparisons showed a significantly increased Ca\(^{2+}\) spark amplitude at AT sites and reproduced Ca\(^{2+}\) spark dimensions distinct from those at cytosolic and subsarcolemmal sites (Figure 4D). Average Ca\(^{2+}\) spark dimensions were quantitatively reproduced by computational modeling (Figure 4E). Thus, mathematical modeling supports the plausibility of a new Ca\(^{2+}\)-signaling mode, where-by AT-associated structures constitutively regulate atrial ECC through highphos RyR2 clusters, locally increasing the sensitivity for Ca\(^{2+}\)-dependent activation as a basis for Ca\(^{2+}\) macro-spark generation and local Ca\(^{2+}\) transient onset.

β-AR stimulation recruits nonjunctional RyR2 clusters. Lowphos RyR2 clusters might be selectively recruited during β-AR stimulation, as opposed to highphos clusters constantly saturated by tonic local PKA and CaMK phosphorylation. To characterize intracellular cAMP concentration changes in real time, we used transgenic mice that express a ubiquitous cytosolic fluorescence resonance energy transfer (FRET) sensor, Epac1-camps (36) (Figure 5A). Pairwise regional FRET sampling in AMs during isoproterenol (ISO) stimulation was used to detect central versus peripheral cAMP concentration changes (Figure 5A). The resulting traces, each from central and peripheral regions, showed comparable robust FRET signal responses as judged by the maximal amplitude of the intracellular cAMP rise after approximately 20-second ISO stimulation (Figure 5B). Also, the maximal rate of the cAMP rise (dF/dt) was not different between central or peripheral AM regions (Figure 5B). Given that cytosolic cAMP movement, at least in VMs, is much slower than free diffusion (37) and that principally similar cAMP concentration changes were present in the center and periphery of AMs, our data support a central, potentially AT-associated cAMP-generating mechanism.

To assess central cAMP-dependent PKA signaling in AMs, we studied the spatiotemporal dynamics of lowphos RyR2 cluster recruitment during acute ISO stimulation (1 μM). We analyzed RyR2-pS2808 signal intensities on RyR2 cluster striations, comparing central versus surface regions for temporal differences (Figure 5C). RyR2-pS2808 cluster phosphorylation on transversal cluster striations was significantly higher 20 seconds after ISO stimulation (Figure 5C), similar to the observed cAMP FRET signal dynamics. This supports the notion of faster and/or more efficient central cAMP–dependent signal transduction inside AMs (Figure 5C).

To maximally recruit RyR2 clusters, we combined β-AR stimulation (1 μM ISO) with selective phosphodiesterase PDE4D inhibition by 10 μM rolipram (RO). This combined ISO/RO treatment converted all lowphos into highphos RyR2-pS2808 clusters, in contrast to control AMs treated with carrier solution (Figure 5D). Consistent with maximal lowphos RyR2 cluster conversion, ISO/RO treatment significantly transformed the pS2808/RyR2 cluster population signal from a lowphos-dominated peak into a single highphos cluster peak (Figure 5D). In addition, ISO/RO treatment shifted the pS2814/RyR2 signal peak rightward (Figure 5E). Apparently, the peak RyR2/pS2808 highphos cluster conversion following ISO/RO stimulation is more pronounced. These data confirm that nonjunctional RyR2 cluster regulation in situ can be explained by bimodal cluster behavior through reversible catecholaminergic recruitment of the overall larger lowphos RyR2 cluster population, through both increased PKA and CaMK phosphorylation.

Since β-AR stimulation is a central inotropic, yet incompletely understood, mechanism in the atria, we studied intracellular Ca\(^{2+}\) transient behavior during 0.5-Hz pacing. AMs with abundant AT structures reproduced the early Ca\(^{2+}\) release profile led by AT site activation under control conditions (Figure 6A). ISO/RO-treated cells showed profoundly accelerated Ca\(^{2+}\) transients with flattened profiles, indicating more even activation of junctional and nonjunctional (CY) Ca\(^{2+}\) release sites (Figure 6A). While the earliest Ca\(^{2+}\) release sites were stably associated with either AT or S locations (shown by di-8-ANEPPS staining), the mean Ca\(^{2+}\) release onset was significantly accelerated at S, but not at AT, locations (Figure 6A). Thus, analysis of different Ca\(^{2+}\) release locations by derivative (dF/dt) showed that the main ISO/RO effect consists of accelerated cytosolic Ca\(^{2+}\) release (Figure 6A), compatible with nonjunctional lowphos RyR2 cluster recruitment leading to accelerated Ca\(^{2+}\) release spread during cAMP-dependent stimulation.

In VMs, cAMP is generated by transmembrane AC5/6, and β-AR stimulation represents the major pathway to increase AC5/6 activity (27). However, this has not been established in atrial muscle. We confirmed similar AC5/6 protein expression levels in murine atrial and ventricular tissue (Figure 6B). Interestingly, expression of the PKA catalytic subunit (PKA-C) was significantly higher in the atria (Figure 6B and Supplemental Figure 10), correlating with more robust PKA-C striations in AMs compared with those observed in VMs (Supplemental Figure 11). Since RyR2 channels and PKA-C were shown to colocalize in situ at Z-line striations (based on previous cAMP FRET measurements; ref. 28), and since the central RyR2 cluster density may be higher in AMs (Figure 3I), our data suggest an increased atrial capacity for cAMP-dependent regulation of RyR2 clusters. Indeed, immunoblots confirmed that the maximal change of substrate-normalized
RyR2-S2808 phosphorylation by PKA (Figure 6B) and RyR2-S2814 by CaMK (Figure 6C and Supplemental Figure 12) during ISO/RO treatment was significantly higher in AMs compared with that detected in VMs. In contrast, acute pharmacological ISO stimulation using a low nanomolar concentration (20 nm) resulted in a significant change in RyR2-S2808 phosphorylation, but no change in RyR2-S2814 phosphorylation (Supplemental Figure 12C). Taken together, our data indicate that synergistic regulation of lowphos RyR2 clusters by both PKA and CaMK occurs during maximal sustained cAMP generation through a highly efficient regulatory mechanism in AMs, with the potential for complete lowphos-to-highphos RyR2 cluster conversion.

Earlier data from rat hearts showed that phospholamban (PLN), an inhibitor of SR Ca\(^{2+}\) (SERCA2) pump function released by PKA phosphorylation of PLN-S16, is expressed at significantly lower levels in AMs than in VMs (16). In addition, SERCA2 expression is approximately 2-fold higher in AMs compared with expression in VMs, while the PLN/SERCA2 ratio is reduced by approximately 4-fold, suggesting decreased stoichiometric inhibition in atria (12, 16). We confirmed significantly higher SERCA2 levels, but since PLN levels were approximately 10-fold lower, the resulting PLN/SERCA2 ratio in mouse atria was reduced by approximately 50-fold compared with the ratio in ventricles (Figure 6B). Consequently, ISO/RO treatment induced only remote phosphorylation changes of PLN-S16. Overall, these data indicate a minor regulatory role of PLN for SERCA2 function in AMs (Figure 6B).

Since SERCA2 activity determines the SR transmembrane Ca\(^{2+}\) gradient underlying SR Ca\(^{2+}\) release, we measured steady-state Ca\(^{2+}\) transients during pacing. Consistent with previous studies (16), Ca\(^{2+}\) transients in mouse AMs showed significantly higher amplitudes and faster rise and decay rates compared with VMs (Supplemental Figure 13). We assessed the SR Ca\(^{2+}\) load by caffeine application (10 mM) and found an approximately 2-fold higher amplitude of the Ca\(^{2+}\) transient in AMs versus that in VMs (Figure 6D), consistent with previous observations in rat AMs (16).

Figure 6. Catecholaminergic stimulation accelerates atrial Ca\(^{2+}\) signaling. (A) Live imaging of AT and S membrane structures during Ca\(^{2+}\) transient onset in control- versus ISO/RO-treated AMs (0.5-Hz pacing). ISO/RO treatment accelerated transversal Ca\(^{2+}\) release propagation across the cytosol, compressing the F25 profile through a decreased cytosolic (CY) latency of early Ca\(^{2+}\) release (dF/dt). Data are representative of 9 control- and 17 ISO/RO-treated AMs. **P < 0.01 and ***P < 0.001, by Student’s t test. (B) Immunoblots comparing ISO/RO-treated (1 μM/10 μM) atrial (A) tissue from WT and phosphorylation-incompetent RyR2-S2808A+/+ mouse hearts; ventricular (V) tissue was compared for reference. In WT atria, significantly higher expression levels of both PKA-C and SERCA2 were detected, in contrast to significantly lower PLN levels. Atrial versus ventricular PKA phosphorylation changes were significantly increased for the RyR2-S2808 site, but not for the PLN-S16 site. Note the similar AC5/6 and RyR2 protein levels. Blots are representative of 3 individual experiments (see Supplemental Figure 10 for uncut blots). (C) Analogous RyR2-S2814–specific immunoblotting established in CaMK phosphorylation–incompetent RyR2-S2814A+/+ mouse hearts (see also Supplemental Figures 12A and 12B). Bar graph shows a significantly increased atrial RyR2-S2814 phosphorylation change in atrial tissue after ISO/RO (1 μM/10 μM) treatment. (D) SR Ca\(^{2+}\) load was significantly higher in AMs than in VMs measured with caffeine (10 mM). n = 8 AMs, 9 VMs. *P < 0.05 and ***P < 0.001, by Student’s t test.
always greater in AMs (Figure 7A). In contrast, PKA inhibition by H89 (1 μM) decreased the sarcomere-shortening amplitude and $V_{\text{max}}$ compared with that observed in untreated AMs (Figure 7A).

To investigate PKA-dependent SR Ca\(^{2+}\) uptake, we studied AMs from PLN-KO (Pln –/–) mice with constitutively upregulated SERCA2 function. While the shortening amplitude was not significantly different in Pln–/– AMs compared with that of ISO-stimulated AMs, the $V_{\text{max}}$ was significantly smaller and similar to that of control WT AMs (Figure 7A). In summary, PLN ablation had no significant effect on sarcomere shortening compared with unstimulated AMs, consistent with the overall low atrial PLN protein expression levels in AMs.

To probe the functional role of RyR2 channel phosphorylation, we analyzed AMs from PKA phosphorylation–incompetent RyR2-S2808A+/+ and CaMK phosphorylation–incompetent RyR2-S2814A+/+-knockin mice. Compared with WT cells, both RyR2-S2808A+/+ and RyR2-S2814A+/+ cells exhibited significantly decreased shortening amplitude and $V_{\text{max}}$ under basal (Figure 7A).
and ISO-stimulated (1 μM) conditions (data not shown). While both RyR2-knockin models showed decreased shortening and V_max under basal conditions, the absolute decrease was significantly greater in RyR2-S2808A/+/+ cells (Figure 7A). Given the more pronounced effect on sarcomere function in RyR2-S2808A/+/+ cells, we used LA echocardiography to explore the functional relevance of highphos RyR2 cluster ablation in vivo in RyR2-S2808A/+/+ mice. While age-matched RyR2-S2808A/+/+ mice with PKA phosphorylation–incompetent RyR2 channels exhibited normal LA diastolic diameters, fractional shortening was decreased (Figure 7B). Hence, genetic ablation of RyR2 PKA phosphorylation is sufficient to cause significant loss of atrial function in vivo, presumably through depressed sarcomere shortening in AMs. Since the M-band is generally considered a relatively elastic transversal sarcomere structure, we hypothesized that systolic M-band deformation could correlate with the Ca^{2+} transient pro-
files observed in AMs. To assess this, we imaged EGFP-labeled M-band striations using AM from knockin mice that express a C-terminally tagged titin-EGFP fusion protein (38). We found a significant transversal latency difference between peripheral and central M-band deformations during sarcomere shortening in live AMs under baseline conditions (Figure 7C; compare Supplemental Videos 1 and 2 or Supplemental Videos 3 and 4). Consistent with transversal lowphos-to-highphos RyR2 cluster conversion, acute ISO (1 μM) stimulation curtailed the M-band–specific latency difference (Figure 7C). The blunted M-band latency difference corresponded to the accelerated Ca²⁺ release spread observed in ISO/RO-stimulated AMs (Figure 6A).

We thus identified the following features of atrial sarcomere regulation: 1) axial sarcomere shortening is constitutively increased through tonic highphos RyR2 cluster regulation in AMs, contributing to rapid Ca²⁺ transient onset through AT-coupled highphos RyR2 clusters at baseline; 2) β-AR stimulation significantly increases sarcomere shortening and reduces transversal latency differences during sarcomere contraction.

**Atrial hypertrophy causes AT proliferation, leading to accelerated cytosolic Ca²⁺ release.** Atrial hypertrophy has been linked to depressed atrial contractility in patients and represents an important risk factor for AF, thromboembolic stroke, and HF. However, the mechanisms leading from atrial hypertrophy (frequently associated with systemically increased blood pressure) to contractile dysfunction, in particular prior to and/or independently of AF, remain unclear. We used a model of transaortic constriction (TAC) in mice to induce hypertrophy of the LA. Four weeks after TAC, echocardiography confirmed significantly increased LA inner diameters and decreased fractional LA shortening (Figure 8A and Supplemental Table 2). Furthermore, increased hypertrophic wall thickening (left ventricular anterior wall thickness in diastole [LV AWTh]) and increased heart weight (HW/BW) were observed (Supplemental Table 2). Histological analysis confirmed that TAC induced significant LA hypertrophy and dilation, but showed no right atrial (RA) pathology (Supplemental Figure 14, A–C).

To explore potential atrial mechanisms of contractile dysfunction, we analyzed AT-dependent Ca²⁺ transient behavior using transversal line scanning of AM from the left atrium of sham-operated and TAC hearts. Hypertrophied AMs from TAC hearts showed more numerous AT structures and considerably compressed Ca²⁺ transient latency profiles, indicating accelerated cytosolic Ca²⁺ release in post-TAC AMs (Figure 8B). Analysis of the Ca²⁺ transient latency profiles (ΔE/Δt) showed that off-membrane CY sites contributed significantly to the accelerated cytosolic Ca²⁺ release spread after TAC compared with that seen in sham AMs (Figure 8B). Hence, post-TAC cells are characterized by significantly accelerated cytosolic Ca²⁺ release through nonjunctional RyR2 cluster sites that are apparently activated through more numerous AT structures in substantially thicker AMs.

To elucidate hypertrophic changes in AT structures after TAC, we performed live-cell confocal membrane imaging. Hypertrophied AM from the left atrium showed an extensive proliferative adaptation of their TAT network compared with sham AMs (Figure 8C). For comparative TAT network analysis, the difference integral of all TAT component orientations was plotted for post-TAC minus sham, highlighting an increase in AT (0°) and a decrease in TT (90°) components in hypertrophied AMs after TAC (Figure 8C). Consequently, the post-TAC AT/TT ratio was significantly increased (Figure 8C). Notably, while atrial hypertrophy leads to select proliferation of AT components, this occurs on the background of significantly increased cell dimensions (Supplemental Figure 14D). Since AT structures couple highphos RyR2 clusters to rapid Ca²⁺ release in AMs, we next sought to analyze proteomic and in situ RyR2 cluster changes.

Immunoblotting showed significantly decreased post-TAC RyR2 protein levels (Figure 8D). Since RyR2 expression was not changed in LV tissue (data not shown), the observed post-TAC RyR2 channel downregulation appeared to be atria specific. Furthermore, PKA phosphorylation was significantly increased at the RyR2-S2808 phosphorylation site, but not at the CaMK phosphorylation site RyR2-S2814, in atrial tissue (Figure 8D). In contrast, 4 weeks after TAC in ventricular tissue, RyR2-S2814, but not RyR2-S2808, phosphorylation was substantially increased (data not shown), confirming an earlier study (39) and supporting the presence of atria-specific RyR2-S2808 phosphorylation changes. PKA hyperphosphorylation of RyR2 is known to increase the sensitivity for Ca²⁺ activation, which is generally consistent with altered cytosolic SR Ca²⁺ release in post-TAC AMs (Figure 8B). Additionally, decreased SERCA2 expression increased the PLN/SERCA2 ratio and may have depressed post-TAC SR Ca²⁺ uptake in AMs (Figure 8D). Of note, increased SR Ca²⁺ release in the context of increased RyR2 phosphorylation and decreased Ca²⁺ uptake may increase the propensity for cytosolic Ca²⁺ overload at higher heart rates.

In atrial hypertrophy with increased post-TAC AT proliferation, a parallel increase of highphos RyR2 clusters occurred, along with more numerous axial-central strings heterogeneously distributed across thicker AM cells from the left atrium (Figure 8E). Using isolated AMs from the LA of TAC hearts, we confirmed a significantly increased post-TAC hypertrophic cell size compared with sham-treated cells (Figure 8E and Supplemental Figure 14D). However, despite the observed increase in highphos RyR2 cluster strings, the cell-wide density of all RyR2 clusters (both lowphos and highphos) was significantly decreased after TAC (Figure 8E), in agreement with the observed decrease in RyR2 protein expression levels. Yet, the AM-specific fraction of highphos RyR2-pS2808 clusters was significantly increased (Figure 8E). This indicates a relatively strong increase in highphos, at the cost of a reduced lowphos RyR2 cluster population and can explain the observed Ca²⁺ transient changes through earlier nonjunctional cluster recruitment, contributing to overall rapid Ca²⁺ transient onset, as observed throughout hypertrophied AMs (Figure 8B). Taken together, the activation of SR Ca²⁺ release through more numerous AT structures and through an increased highphos RyR2 cluster fraction contributes to accelerated recruitment and earlier activation of adjacent nonfunctional lowphos clusters, indicating a sustained mechanism of subcellular Ca²⁺ transient adaptation in hypertrophied AMs, despite significantly decreased RyR2 channel expression.

**Discussion**

Here, we identify an atrial ECC “super-hub” mechanism that is based on a highly specialized membrane-protein network characterized by: 1) tubular membrane structures (ATs) with axial orientations that are 2) abundant in AMs and 3) connected to the
surface membrane through sparse TT components; 4) significantly larger-diameter AT structures compared with T'T' structures in AMs; 5) AT-coupled large junctional RyR2 clusters that are highly phosphorylated by PKA and CaMK and arranged in axial string-of-pearls; 6) triggering by ATs of rapid Ca\(^{2+}\) release from axial highphos R\(y\)R2 cluster couplons in AMs via Cav1.2 channels; and both 7) rapid central Ca\(^{2+}\) transient onset and 8) central Ca\(^{2+}\) macro-sparks that are activated via junctional highphos R\(y\)R2 clusters at AT sites. While these observations present a robust intracellular context for rapid Ca\(^{2+}\) signaling in AMs, they also lay the foundation for novel and, most important, atria-specific regulatory ECC mechanisms. Given the relevance and functionality of the membrane-protein network detailed above, we propose a signaling super-hub model that is structurally based on unusually large AT and axial couplon dimensions and that may change our perspective on cell-specific control mechanisms of atrial Ca\(^{2+}\) signaling and contractile regulation.

Our observations are supported by previously described dedicated AM isolation and high-resolution imaging workflows (18, 40), which led to our finding of preserved centrally located AT structures within TAT networks. Since AMs are generally thinner than VMs, it has been proposed that TTs are not needed for ECC in these cells (8, 19). However, when we normalized the total TAT network length to the cell area, we found AMs with intermediate (Figure 1B) rather than low or zero TAT densities, as previously reported in rodents (8). Furthermore, we found voluminous AT structures that were electrically connected to the surface sarcolemmal membrane, presumably via more sparsely present TTs. While overall similar, TAT structures were shown previously in a minority of cells isolated from atria (7, 41). Here, we identify ATs as regularly present and specific structural components that are key to ECC in AMs.

**Novel cell-specific properties: highly phosphorylated R\(y\)R2 clusters and SR load tune Ca\(^{2+}\) release at AT locations.** In living AMs, STED revealed that central R\(y\)R2 clusters may be spaced more tightly along Z-lines than are those in VMs. We show that the vast majority of transversely oriented R\(y\)R2 cluster striations have no nearby TT membrane and are therefore not triggered by the opening of local LCC. In theory, an increased density of transverse R\(y\)R2 cluster striations have no near-TT membrane, presumably via more sparsely present TTs. While overall similar, TAT structures were shown previously in a minority of cells isolated from atria (7, 41). Here, we identify ATs as regularly present and specific structural components that are key to ECC in AMs.

Our use of voltage-clamp and EGTA (1 mM) Ca\(^{2+}\) buffering allows us to conclude that electrical transmission via the TAT network underlies the more rapid ECC activation of AT-coupled junctional R\(y\)R2 clusters. Here, we identify highly and stably phosphorylated R\(y\)R2 clusters, which are to a large degree associated with AT membranes. At the R\(y\)R2 clusters, combined PKA and CaMK phosphorylation of R\(y\)R2-S2808 and R\(y\)R2-S2814, respectively, significantly increases the sensitivity of the Ca\(^{2+}\)-dependent activation of R\(y\)R2 (29, 30, 43). Consequently, junctional AT couplons are ideally positioned to rapidly initiate local SR Ca\(^{2+}\) release, as evidenced by Ca\(^{2+}\) macro-sparks. Furthermore, we show that nonjunctional SR locations contribute a functionally important and economic form of “inotropic reserve,” accessible through cAMP-dependent recruitment of lowphos R\(y\)R2 clusters and representing the first example, to our knowledge, of a differential regulatory mechanism of this protein in distinct SR domains.

We observed a high level of SR Ca\(^{2+}\) load in mouse AMs under baseline conditions, consistent with earlier observations (16). Within the context of highphos versus lowphos R\(y\)R2 clusters, a high SR Ca\(^{2+}\) content against the background of PLN-deficient AMs can explain the significantly larger central Ca\(^{2+}\) transient amplitude observed at baseline. Accordingly, we confirmed that SERCA2 protein levels were approximately 2 times higher in mouse atria than in ventricles, which was previously shown to stabilize Ca\(^{2+}\) cycling in ratAM (16). Importantly, minimal PLN protein levels in mouse atria resulted in a very low PLN/SERCA2 ratio. This is consistent with an earlier study showing that the transcription factor COUP-TFII suppresses PLN transcription in atria (44). Therefore, PKA-dependent recruitment of lowphos R\(y\)R2 clusters may represent a main inotropic SR regulatory mechanism in AMs. Indeed, PLN ablation did not significantly change the amplitude of sarcomere shortening in AMs from "Pkn\(^{−/−}\)K0 mice."

Using a simplified mathematical model of atrial Ca\(^{2+}\) sparks, we were able to reproduce the macro-spark behavior observed at the AM center, on the basis of stochastic R\(y\)R2 gating of lowphos and highphos clusters. As has been established before, the control of R\(y\)R2 sensitivity to cytosolic [Ca\(^{2+}\)], appears to depend on multiple factors and interactions. Recent investigations suggest that Ca\(^{2+}\) in the SR lumen ([Ca\(^{2+}\)\(_{\text{SR}}\)]) may exert a nonlinear action on the sensitivity of the R\(y\)R2 clusters to be triggered (45, 46). Thus, a higher [Ca\(^{2+}\)\(_{\text{SR}}\)] can underlie Ca\(^{2+}\) sparks, with a greater total amount of Ca\(^{2+}\) released per average event. Ca\(^{2+}\) macro-sparks may facilitate the local propagation of CICR, originating from axial couplons. The atrial super-hub provides a new concept, on the background of predominant expression of fast \(\alpha\)-myosin heavy
chain in atria, for both the rapid onset and extent of AM contraction, the potential for physiological regulation, and altered high-phosphorylated RyR2 cluster function in hypertrophy.

**Alterations of axial couplon function in atrial disease.** In rabbit AM after 5 days of rapid atrial pacing RyR2 protein expression was reduced by 50% (4), similar to our observations in mouse AM post-TAC (Figure 8D). This has given rise to the concept of a physiological antiarrhythmic adaptation (Ca²⁺-signaling silencing), attributed to a failure of central CICR due to reduced RyR2 channel numbers per cluster and despite RyR2-S2808 hyperphosphorylation (4). In contrast, after TAC, we found an additional decrease of approximately 30% in RyR2 cluster density, yet identified an increase of higmphos RyR2 clusters in more numerous strings, which contributed to an approximately 2-fold increase in the higmphos RyR2 cluster fraction. This suggests the presence of an adaptive mechanism that sustains cytosolic Ca²⁺ release through AT-coupled central RyR2 clusters after TAC, in stark contrast to the reported tachypacing-induced failure of central Ca²⁺ signal propagation (4). As hypertrophy and reduced contractility were identified in canine atria as early as 2 days after pacing (5), it will be important for future studies to explore the precise time course of transition from normal to hypertrophied atria. In contrast to our post-TAC model, a tachypacing HF model of persistent AF in sheep showed a complete loss of atrial TAT structures, consistent with advanced HF (20, 47). While we examined the LA response to TAC-induced atrial hypertrophy in the absence of any obvious arrhythmias (data not shown), the adaptive response was a profound increase in AT components, higmphos RyR2 clusters, and increased nonjunctional Ca²⁺ release.

**Clinical relevance of atrial super-hubs.** The use of STED nanoscopy and ET, combined with membrane-preserving workflows, has enabled visualization of AT structures throughout AMs. Instead of the TAT being absent in AMs as previously suggested (6, 20), we found that a robust and functionally relevant tubular membrane system was present in every AM studied. In mouse and human AMs, AT membrane structures are an essential component of axial couplings that contain extended RyR2 clusters. The functions of this complex assembly, described as super-hubs, are likely to play important roles in modulating atrial ECC, distinct from classic couplings in VMs. In contrast, RyR2 PKA phosphorylation is significantly increased in patients with chronic AF (33) and after short-term tachypacing (4). Both RyR2 single-channel function and PKA phosphorylation were partially normalized in ventricular samples from HF patients treated with β-AR blockers (48). Since β-AR blockers are used in approximately 90% of patients with AF (2), our finding of a super-hub mechanism of contractile regulation in AMs through higmphos RyR2 clusters may be instrumental in specifically targeting the (side) effects of GPCR-targeted drug compounds including β-AR blockers or currently available CaMK inhibitors on atrial contractility. Future work will need to examine the diverse functional roles of axial couplings and changes in RyR2 cluster regulation of hypertrophy in atrial cells, tissues, and the heart in vivo.

The causal relation between higmphos RyR2 clusters and rapid atrial contractile activation, identified through RyR2 phosphorylation-incompetent knockin animals, and the sustained structural and metabolic adaptation mechanisms in hypertrophy strengthen the rationale for the identification of early atrial disease mechanisms and, ultimately, for AM-directed treatment approaches.

**Methods**

Additional information is provided in the Supplemental Methods.

**Mouse cells.** AMs and VMs were isolated from adult mouse hearts using collagenase and protocols tuned to preserve subcellular structural integrity, as previously described (9, 18). RyR2-S2814A⁻/-knock-in mice were provided by Xander H.T. Wehrens (Baylor College of Medicine, Houston, Texas, USA); and PLN-KO mice were provided by Despina Sanoudou (University of Athens, Athens, Greece). Unless indicated otherwise, the genetic background was C57BL/6N.

**Human cells and tissues.** Atrial tissue samples were obtained from patients with sinus rhythm during bypass surgery, following written informed consent from the patients. Cell isolation (n = 5) occurred from atrial tissue by protocols described previously (49). Cells were plated on laminin-coated coverslips for immunofluorescence staining. For IHC, LA tissue was fixed in 4% paraformaldehyde (PFA), embedded in paraffin, cut into 4-μm-thick sections, deparaffinized, rehydrated, and antigens unmasked in 10 mM sodium-citrate buffer prior to Ab incubation.

**Live-cell confocal microscopy.** Confocal images were acquired with a Zeiss LSM 710 NLO microscope using a Plan-Apochromat 63x/1.40 oil objective and a pixel size of 100 nm. Myocytes attached to lamin-coated slides were stained with 50 μM di-8-ANEPPS (Molecular Probes) for 15 minutes, washed twice, excited at 488 nm, and emitted light detected above 510 nm as previously described (9). TAT networks in AMs and VMs were visualized using identical settings. Intracellular Ca²⁺ imaging with fluo-4 AM was combined with di-8-ANEPPS as described previously (9). For Ca²⁺ spark imaging and signal analysis, we used previously established protocols (50). Further details on the TAT component analysis, Ca²⁺ transient analysis, or patch-clamping are available in the Supplemental Methods.

**Live cell STED microscopy.** For super-resolution imaging of select TAT membrane tubules, we used a customized STED system and Matlab (The MathWorks) algorithms as described previously (9, 18). Further details are provided in the Supplemental Methods.

**ET.** AT and TT structures and isolated AM were preserved by high-pressure freezing with an HPM100 system (Leica) as described previously (51). ET at the Electron Microscopy core facility at the European Molecular Biology Laboratory (EMBL) Heidelberg (Heidelberg, Germany) and at the EM services of the University of Colorado Boulder (Boulder, Colorado, USA) was performed as described previously (23).

**Pharmacological assays.** AMs plated on laminin-coated coverslips were pretreated with 10 μM DMSO dissolved in DMSO (Sigma-Aldrich) for 3 minutes, followed by 1 μM ISO (Sigma-Aldrich) for 10 minutes (ISO/RO). For Western blot analysis, mouse hearts wereLangendörff perfused as described above, followed by immediate tissue freezing. For acute β-AR stimulation, 1 μM ISO was used if not stated otherwise. For in situ PKA inhibition, AMs were incubated for at least 15 minutes with H89 (Sigma-Aldrich). For in situ CaMK inhibition, cells were treated with 5 μM AIP (Sigma-Aldrich) for 30 minutes.

**FRET measurements.** In AMs from mice expressing the Epac1-camps FRET biosensor, intracellular cAMP was assessed during 0.1 μM ISO treatment as previously described (36). Additional details are provided in the Supplemental Methods.

**Sarcomere shortening.** Sarcomere shortening measurements used high optical contrast imaging according to the manufacturer’s instruc-


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