Localized translation and sarcomere maintenance requires ribosomal protein SA in mice

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Cardiomyocyte sarcomeres contain localized ribosomes, but the factors responsible for their localization and the significance of localized translation are unknown. Using proximity labeling, we identified ribosomal protein SA (RPSA) as a Z-line protein. In cultured cardiomyocytes, the loss of RPSA led to impaired local protein translation and reduced sarcomere integrity. By employing CAS9-expressing mice, along with adeno-associated viruses expressing CRE recombinase and single-guide RNAs targeting *Rpsa*, we knocked out *Rpsa* in vivo and observed mislocalization of ribosomes and diminished local translation. These genetic mosaic mice with *Rpsa* knockout in a subset of cardiomyocytes developed dilated cardiomyopathy, featuring atrophy of RPSA-deficient cardiomyocytes, compensatory hypertrophy of unaffected cardiomyocytes, left ventricular dilation, and impaired contractile function. We demonstrated that RPSA C-terminal domain is sufficient for localization to the Z-lines and that if the microtubule network is disrupted RPSA loses its sarcomeric localization. These findings highlight RPSA as a ribosomal factor essential for ribosome localization to the Z-line, facilitating local translation and sarcomere maintenance.

Introduction

Local translation has several benefits over diffuse or unlocalized protein production. Importantly, it allows protein synthesis closer to the site of action, is more economic, may prevent protein aggregation, and can facilitate processes such as cotranslational assembly (1). Local translation requires the presence of the mRNA and the ribosome as well as additional translation factors and cofactors. While the localization of mRNAs and the RNA-binding protein involved, particularly in neurons, have been studied (2), the factors involved in the localization of cytoplasmic ribosomes are less clear.

The sarcomere is the basic contractile element of cardiomyocytes. Because its protein constituents have a limited half-life, this complex must be continuously maintained (3). Failure in sarcomeric proteostasis may lead to heart failure because failing human hearts have reduced sarcomeres and sarcomeric protein content (4–6). We previously showed that ribosomes are localized on both sides of the Z-line of the sarcomere and that proteins are translated there, leading us to propose that sarcomeres are maintained by the localized

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translation of proteins within them (7). We have also demonstrated that the microtubule cytoskeleton and kinesin are essential for localized translation in the heart (7, 8). Similar observations were made for skeletal muscles (9). Yet, the importance of localized translation for sarcomere maintenance has not been demonstrated, and the ribosomal factors required for localization are unknown.

Here, we used proximity labeling to elucidate the proteomic composition of sarcomeric Z-lines in an unbiased manner and identified ribosomal protein SA (RPSA), a protein with reported ribosome and laminin receptor functions (10). We show that loss of RPSA in cultured cardiomyocytes resulted in the loss of local protein translation and in reduced sarcomere integrity. Using conditional and unconditional CRISPR-associated protein 9expressing (CAS9-expressing) mice and adeno-associated viruses (AAVs) expressing CRE recombinase in cardiomyocytes with single-guide RNAs (sgRNAs) targeting Rpsa we knocked out Rpsa in cardiomyocytes in vivo and show that the loss of RPSA results in mislocalization of ribosomes and reduced local protein translation. These genetic mosaic mice, where only a portion of cardiomyocytes are transduced and have Rpsa knockout, developed a dilated cardiomyopathy with atrophy of Rpsa-knockout cardiomyocytes, compensatory hypertrophy of untransduced cardiomyocytes, left ventricular dilation, and reduced contractile function. RPSA is known to directly bind the microtubular system, and we showed that its C-terminal domain is sufficient for localization to Z-lines and that if the microtubule network is disrupted RPSA loses its sarcomeric localization. These data show that RPSA, the first ribosomal factor responsible for localizing cytoplasmic ribosomes identified to our knowledge, is required for localization of ribosomes to the Z-line, for the local translation there, and for the maintenance of the sarcomere.

Results

Proximity-dependent labeling identifies components of translational machinery at sarcomeric Z-lines. To investigate the proteomic composition of sarcomeric Z-lines and identify the factors involved in localized translation, we employed the BioID2 proximity labeling assay using Cypher (LDB3), a known Z-line protein, as bait. For this purpose, we generated a fusion construct between Cypher and the promiscuous biotin ligase BioID2. We then transduced neonatal rat ventricular cardiomyocytes (NRVMs) with an adenoviral vector encoding for Cypher-BioID2 or control virus expressing BioID2 alone. To confirm the correct subcellular localization of our fusion construct, we first performed immunofluorescence staining that demonstrated that Cypher-BioID2 fusion construct closely matched the known localization pattern of endogenous Cypher at the sarcomeric Z-lines, whereas BioID2 alone displayed a more diffuse cytoplasmic staining (Figure 1A). We then used 3 independent samples for each group, each containing 12 million cardiomyocytes, and purified biotinylated proteins using streptavidin-coated magnetic beads after 18 hours of incubation with biotin. Western blot analysis of the extracted cardiac proteins, using a streptavidin probe confirmed the successful biotinylating of proteins and showed distinct sets of labeled proteins in the Cypher-BioID2 sample compared with the BioID2-alone control samples (Figure 1B).

Pulled down proteins were trypsin digested on beads and subjected to mass spectrometry analysis. The proteomic analysis identified 135 proteins significantly enriched in the Cypher-BioID2 sample compared with the BioID2-alone control using a cutoff of more than 2-fold enrichment (\log_2 fold change ≥ 1) and ttest *P* < 0.05 (Figure 2A and Supplemental Table 1; supplemental material available online with this article; https://doi.org/10.1172/ JCI174527DS1). As expected, LDB3 (Cypher), which served as the bait, was the most abundant and most significantly enriched protein in the bait group. Proteins known to be localized to the Z-line, such as α -actinin (ACTN2) and titin (TTN), were also significantly enriched in the bait group. On the other hand, highly expressed proteins known to be localized to the M-line, such as MYOM1 (Myomesin1) or OBSCN (Obscurin), were significantly more abundant in the control group, demonstrating that we have specifically enriched for a Z-line nanodomain. Additional proteins significantly enriched in the bait group included RPSA and ribosomal protein L38 (RPL38).

To gain insight into the protein groups associated with the Cypher-BioID2 significantly enriched proteins (fold change \geq 2 and P < 0.05), we performed an enrichment analysis using gene ontology (GO) terms. The analysis indicated that the Cypher-BioID2-associated proteins were significantly enriched in several terms, including ribosome and sarcomere Z-line, further confirming our previous observation that the Z-line region contains a protein translation nanodomain (Figure 2B). There is an inherent overlap among GO terms, as a given protein can be associated with multiple terms. We therefore also employed an unsupervised clustering approach with the aim of identifying distinct clusters of enriched GO terms. This analysis revealed two major clusters of protein

functions. The first cluster consisted of pathways related to translational machinery, such as ribosomal proteins and ribonucleoprotein complexes. The second cluster included pathways involved in cytoskeletal elements, particularly those associated with sarcomeric structures (Figure 2C). All Cypher-BioID2-enriched proteins are also shown in a STRING analysis (Supplemental Figure 1).

Of the identified proteins, we decided to concentrate on RPSA. RPSA is considered a ribosomal protein that has acquired the function of laminin receptor during evolution. Although its function in the heart is not known, it was reported to bind both ribosomes and the microtubular system (10). We also studied RPL38, a known component of the 60S subunit of the ribosome. We performed immunofluorescence staining of RPSA in adult rat cardiomyocytes as well as 3-dimensional reconstruction of Z-stacked images showing that RPSA was aligned on both sides of the α -actinincontaining Z-lines (Figure 2, D and E, and Supplemental Video 1). This result further validates our proximity labeling approach and confirms that RPSA is localized to sarcomeric Z-lines.

Loss of RPSA disrupts localized translation and sarcomeric integrity in cultured cardiomyocytes. The localization of RPSA on both sides of the Z-line suggested that in cardiomyocytes this protein has a ribosomal rather than a laminin receptor function. We therefore asked if RPSA is required for localized protein translation in the sarcomere and for sarcomere maintenance. We used siRNA to knockdown *Rpsa* in NRVMs or knocked down *Rpl38*, a known ribosomal protein, as a control. The efficacy of the *Rpsa* knockdown was validated by quantitative PCR (RT-qPCR), which demonstrated an approximately 85% reduction in *Rpsa* and approximately 82% reduction in *Rpl38* mRNA levels compared with the nontargeting negative control siRNA group (Figure 3A). A reduction in RPSA protein levels was also confirmed using Western blotting (Figure 3B).

To assess the effect of *Rpsa* or *Rpl38* knockdown on localized protein translation, the control and knockdown cells were incubated with O-propargyl-puromycin (OPP), a puromycin analog that labels newly translated proteins, for 60 minutes. Cardiomyocytes were fixed, and OPP-incorporated nascent peptides were labeled using Alexa Fluor 488 by a Click reaction and immunostained with antibodies against sarcomeric ACTN2 to examine the sarcomeric structure. This analysis revealed a major disruption in localized protein synthesis with knockdown of *Rpsa* but not with knockdown of *Rpl38* or control siRNA knockdown (Figure 3C). The sarcomere structure on the other hand appeared largely intact.

To quantify these changes, we used an automated image processing approach inspired by the Sarcomere Texture Analysis method (11). These custom tools segment cells from multichannel immunofluorescence images and employ algorithms to quantify two aspects of the sarcomeric structure, namely homogeneity and colocalization of different markers. Homogeneity score is a measure of the uniformity of signal distribution in the image, with a higher score indicating a more diffuse signal distribution, and a lower score indicating a more localized distribution (Figure 3D). This unbiased automated analysis showed that the knockdown of *Rpsa* resulted in a statistically significant increase in homogeneity in the OPP signals (Figure 3E), indicating a loss of localized protein translation with a shift toward more diffuse translation. While our visual analysis did not reveal a major disruption in the sarcomere organization, our automated analysis of ACTN2

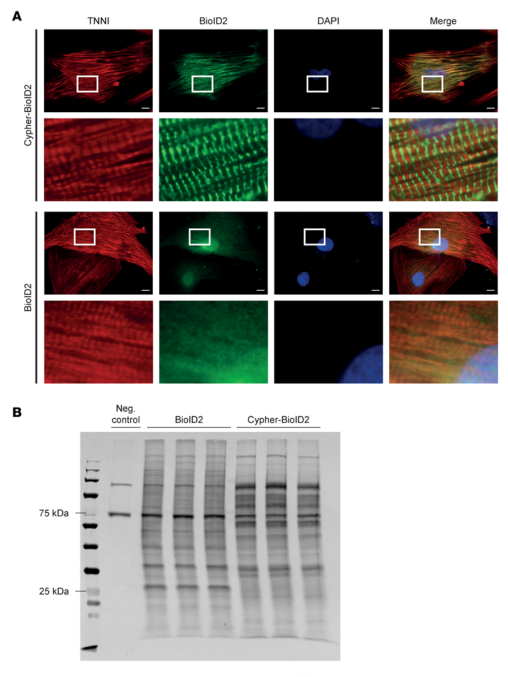


Figure 1. Cypher-BioID2 is localized to the Z-lines and biotinylates specific proteins. (A) Representative immunofluorescence images of NRVMs transduced with adenoviruses encoding for either Cypher-BioID2 (bait construct) or BioID2 (control construct), with the boxed area magnified to show Z-line localization of Cypher-BioID2 and a diffuse cytoplasmic localization for BioID2 alone. Samples were immunostained for cardiac troponin (red), BioID (green), and DAPI (blue). Scale bar: 10 μm. The higher-magnification images were magnified 5.5 times the original. (B) Western blot analysis of biotinylated proteins using IRDye800-Streptavidin showing different biotinylated patterns for cardiomyocytes transduced with BioID2 or Cypher-BioID2 and minimal biotinylation of the untransduced negative control sample. Each lane was loaded with 100 μg protein lysate.

staining demonstrated an impairment (Figure 3F). The automated analysis also revealed a significant decrease in the colocalization between the ACTN2 and OPP signals after *Rpsa* knockdown (Figure 3G), indicating a much greater loss of sarcomeric localized translation relative to the changes in sarcomeric structure. In contrast with these findings, the loss of RPL38 did not affect the localization of translation or the integrity of the sarcomere. Altogether, these data indicate that the loss of RPSA results in the

loss of sarcomeric Z-line protein translation and a likely secondary loss of sarcomere structure in cardiomyocytes.

Loss of RPSA results in cardiac dysfunction in vivo. We next wanted to study the consequences of RPSA loss in cardiomyocytes in vivo. We first used mice generated by crossing the Rosa26-Lox-stop-Lox-Cas9-knockin mice, which have a floxed-STOP cassette preventing the expression of the downstream CAS9 endonuclease from a CAG promoter knocked in the Rosa26

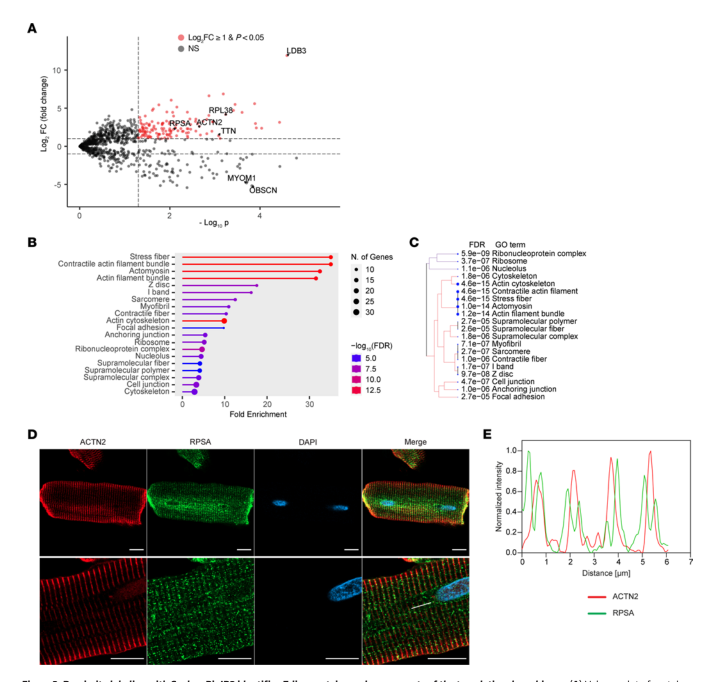


Figure 2. Proximity labeling with Cypher-BiolD2 identifies Z-line proteins and components of the translational machinery. (A) Volcano plot of proteins identified by mass spectrometry showing the \log_2 fold change in protein abundance with Cypher-BiolD2 versus BiolD2 alone and the t test $-\log_{10}$ (P value) of the difference significance. Dashed horizontal lines show the ≥2 and ≤2 cutoffs for fold change, and the vertical dashed bar shows the P < 0.05 cutoff for significance. Proteins highlighted in red are those with a fold change ≥2 and P < 0.05 that were used for further pathway enrichment analysis. Other proteins were denoted as nonsignificant and are shown in gray. Several proteins identified as significant are highlighted. (B) Pathway enrichment analysis of BiolD2 significantly enriched proteins (fold change ≥2 and P < 0.05) showing fold enrichment of GO terms associated with cytoskeletal elements of the sarcomere and with ribosomes and protein translation. The number of proteins identified for each term is shown as a circle, and the $-\log_{10}$ of the false discovery rate (FDR) is color coded. (C) Unsupervised clustering of enriched GO terms showing identified proteins were predominantly associated with two groups − cytoskeletal sarcomere proteins or ribosomal components. The enrichment FDR of specific GO terms is shown. (D and E) Representative immunofluorescence images (D) and line scan (E, white line) analysis of isolated adult rat ventricular cardiomyocytes stained for α-actinin (red), RPSA (green), and DAPI (blue), showing localization of endogenous RPSA to both sides of the Z-line identified with α-actinin. Scale bar: 10 μm.

locus, and the β -actin-driven CRE recombinase strain, resulting in mice with ubiquitous and strong expression of CAS9 under the control of the CAG promoter in multiple tissues, including the heart. Studies at The Jackson Laboratory showed that constitutive CAS9 expression does not result in any detectable toxicity or

morphological abnormalities (12). To achieve *Rpsa* knockout we injected neonatal (3–5 days after birth) mice with AAVs encoding for sgRNAs targeting exons 3, 4, and 5 of the *Rpsa* gene, in equal proportions (Supplemental Figure 2A). The resulting mice are genetic mosaics in which the fraction of cardiomyocytes with

Rpsa knockout in the heart depends on the transduction efficiency, and we refer to these mice as unconditional Rpsa-knockout mice. For controls, we injected mice with an AAV encoding for a nontargeting sgRNA sequence. We confirmed the CRISPR/CAS9-mediated gene editing in the Rpsa gene in the mouse heart tissue using a T7 endonuclease I (T7E1) assay at the age of 3 months (Supplemental Figure 2B). We analyzed these unconditional Rpsa-knockout mice at the age of 3 months. Echocardiography analysis was consistent with dilated cardiomyopathy with increased diastolic and systolic left ventricular dimensions, decreased wall thickness, and reduced systolic function (Figure 4, A-F). Gravimetric analysis supported these findings, showing an elevated heart weight-to-body weight ratio (Figure 4G).

To ensure that the phenotype resulted from cardiomyocyte cell autonomous loss of Rpsa we next used the parental Rosa26-Lox-STOP-lox-Cas9-knockin mouse line. These mice were similarly transduced with AAVs that encode for the sgRNAs targeting Rpsa, or control sgRNA, and also expressed HA-tagged nuclear localized CRE under the control of the chicken cardiac troponin T promoter to delete the STOP cassette and express CAS9, specifically in cardiomyocytes. We also confirmed the CRISPR/ CAS9-mediated gene editing in the Rpsa gene in the mouse heart tissue using a T7E1 assay in these conditional mice at the age of 3 months (Supplemental Figure 2C). We analyzed mice with cardiomyocyte-specific CRE expression at the age of 1, 3, and 6 months by echocardiography (Figure 5, A-F). Interestingly, the fractional shortening was mildly increased in the Rpsa-knockout mice at the age of 1 month. The cause of this increase is not clear and may represent a compensatory neurohumoral response to pathological changes in these hearts. By the age of 3 months there was a trend toward dilation and reduced systolic function, and the same phenotype as in the unconditional Rpsa-targeted mice was fully manifested at the age of 6 months, with significant chamber dilation and reduced function. The delay in the development of overt cardiomyopathy in the conditional Rpsa-knockout mice may be due to decreased efficiency of the troponin T-driven CRE-expressing AAVs compared with the germline β-actin-driven CRE or due to the added effect of Rpsa deletion in noncardiomyocytes in the unconditional Rpsa-knockout mice. Nevertheless, these data confirm that Rpsa knockout induces cardiomyocyte cellautonomous dilated cardiomyopathy phenotype.

We next assessed changes in gene expression in ventricle tissue using RT-qPCR at the age of 1 and 3 months in these mice (Figure 5, G-P). Although echocardiography did not show significant changes at the age of 1 or 3 months, the fetal genes *Nppa* and *Nppb*, encoding ANP and BNP, respectively, and *Myh7* were upregulated, suggesting cardiac stress. Similarly, markers of the fibrotic reaction *Fn1* and *Col1a1* were upregulated. *Acta1* trended for upregulation and *Myh6* trended for downregulation, but these changes were not statistically significant. At the age of 3 months, *Nppa* and *Myh7* remained elevated, as was the Myh7/Myh6 ratio, but *Fn1* and *Col1a1* were no longer significantly elevated. There were no significant changes in *Tnni1* or *Tnni3* levels. These findings are consistent with dilated cardiomyopathy that started to develop at an early age.

For microscopic analysis, we first stained cardiac sections from both *Rpsa*-knockout and control mice at the age of 6 months with antibodies for HA-tag and wheat germ agglutinin (WGA) (Figure

5, Q-T). As the mice are genetic mosaics, only some cardiomyocytes were transduced and showed Rpsa knockouts, and we used the HA staining to identify the transduced cardiomyocytes through their HA-CRE nuclear signal. In control mice, HA-positive and HAnegative cells displayed no significant difference in area, width, or length. However, in Rpsa-knockout mice, HA-positive cells were significantly smaller compared with HA-positive cells in control mice and had significantly smaller area, width, and length. On the other hand, HA-negative cells in Rpsa-knockout mice were significantly larger in area, width, and length compared with HA-negative cells in control mice. The percentage of HA-positive cardiomyocytes in Rpsa-knockout hearts was approximately 15% and did not differ between the age of 1 and 6 months (Supplemental Figure 3, A and B), and similarly, we could not find apoptotic cardiomyocytes on TUNEL staining (Supplemental Figure 3C). Together, these findings show cell atrophy in Rpsa-knockout cardiomyocytes and a compensatory hypertrophy of the untransduced cardiomyocytes in the mosaic Rpsa-knockout hearts, without any evidence for significant cell death of Rpsa-knockout cardiomyocytes.

Loss of RPSA results in loss of localized translation and loss of ribosome localization in vivo. Next, we wanted to see if loss of RPSA resulted in the loss of localized protein translation in cardiomyocytes in vivo. We intraperitoneally injected 3-month-old Rpsa-knockout and control mice with OPP (0.049 mg/g), and after 1 hour, we sacrificed the mice, harvested the hearts, and performed cryosections. The OPP labeling was performed by a Click reaction on the sections, and the cells were costained for the HA-tag to identify the nuclear HA-CRE in the transduced cells. As we have previously shown (7), the OPP staining displays a characteristic striated pattern, reflecting localized sarcomeric protein translation in wild-type cardiomyocytes. When compared with HA-positive cardiomyocytes in control mice, Rpsa-knockout cardiomyocytes lacked this localized pattern indicating that the loss of RPSA markedly reduced sarcomeric localized translation (Figure 6, A and B). The cross-striated pattern of the OPP signal was quantified by measuring the mean amplitude of a line-scan across cardiomyocytes' long axis, and Rpsa-knockout cardiomyocytes showed significantly lower amplitude indicating a more diffuse signal (Figure 6C). Quantification of the density of the OPP signal showed that while translation activity was reduced in Rpsa-knockout cardiomyocytes, as compared with cardiomyocytes transduced with control sgRNA, the OPP signal was still significantly higher in Rpsa-knockout cardiomyocytes than in mice with Click reactions but without OPP (NC.1) or mice with OPP but no Click reaction (NC.2), (Figure 6D). These analyses indicate that loss of RPSA results in disruption of localized translation at the sarcomeres but not a complete abrogation of protein translation.

The loss of sarcomeric localized translation is expected to result in the loss of sarcomeric proteins. To assess this loss, we immunostained sections with antibodies against ACTN2 or myosin heavy chain (MYH), together with HA-tag to identify transduced cardiomyocytes. This analysis showed that the cytoplasm of *Rpsa*-knockout cardiomyocytes largely lacked these sarcomeric proteins, with some remaining signal in the cell periphery (Figure 6, E-J). To further confirm and quantify the loss of sarcomeric proteins, we knocked down *Rpsa* expression in cultured NRVMs with *Rpsa* or control siRNAs. While the knockdown of *Rpsa* resulted in

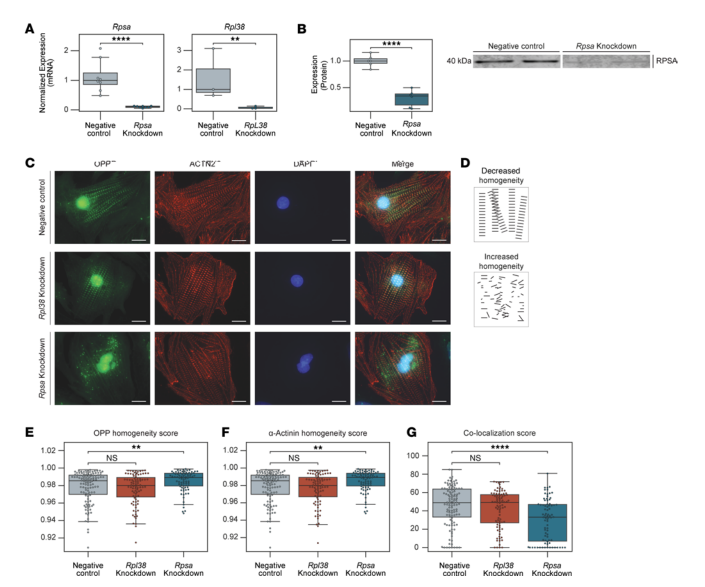


Figure 3. *Rpsa* knockdown disrupts localized translation in cultured cardiomyocytes. (A) Gene expression analysis with RT-qPCR of *Rpsa* and *Rpl38* following siRNA-mediated knockdown compared with control siRNA. Expression was normalized to *Gapdh*. n = 3, and 8 biological samples per experimental group pooled from 3 independent experiments. (B) Western blot analysis with densitometric analysis of RPSA protein expression showing an approximately 70% reduction in RPSA protein level compared with control following siRNA-mediated knockdown. n = 6 biological samples per experimental group from 3 independent experiments. (C) NRVMs were incubated with OPP to label nascent proteins 48 hours after treatment with negative control (top), *Rpl38* (middle), or *Rpsa* (bottom) siRNAs. Representative images of staining for OPP-labeled nascent proteins (green), α-actinin (red), and DAPI (blue) show disruption of localized protein synthesis following *Rpsa* knockdown but not following *Rpl38* knockdown or under control conditions. Scale bar: 10 μm. (D) Cartoon illustrating a structure with decreased homogeneity (top, score = 0.959) and a structure with increased homogeneity (bottom, score = 0.977). (E and F) Homogeneity scores for translation from the OPP signal (E) and for the sarcomeric structure from the α-actinin signal (F), showing increased homogeneity implying decreased localization following *Rpsa* knockdown but not following *Rpl38* knockdown or control siRNAs. (G) Colocalization analysis between the OPP and the α-actinin signal showing significantly decreased colocalization between translation and the Z-line after knockdown of *Rpsa*. n = 79, 87, and 116 cells per group pooled from 4 independent experiments. ** $P \le 0.01$, **** $P \le 0.0001$, by unpaired Student's t test (A and B) and 1-way ANOVA test (E-G). Data are presented as individual values, with box plots displaying the median and 25th and 75th percentiles.

a significant loss of the sarcomeric proteins MYBPC3, ACTN2, and TNNI3 or the nonsarcomeric protein tubulin, the levels of the mitochondrial proteins VDAC1 or SDHA remained mostly unchanged (Figure 6, K and L).

To verify that RPSA was indeed associated with the ribosomal complex we performed polysome profiling from normal mice heart extracts. Western blot analysis from these fractions showed that RPSA was associated with both monosomes (80S) and polysomes (Supplemental Figure 4A). The association of RPSA with polysomes

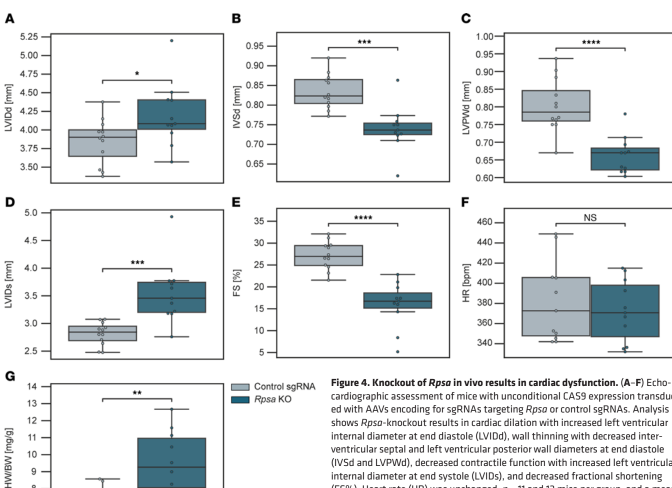
indicates it is part of the ribosome complex actively engaged in mRNA translation. Costaining of sections from control mice administered OPP for RPSA showed that RPSA was localized on both sides of the OPP-incorporated nascent proteins at the sarcomere (Supplemental Figure 4, B and C), further supporting the association of RPSA with active translation adjacent to the sarcomere Z-line.

We next asked if the loss of localized translation resulted from impaired ribosome localization in *Rpsa*-knockout cardiomyocytes. We used dual single-molecule fluorescent in situ

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cardiographic assessment of mice with unconditional CAS9 expression transduc-(IVSd and LVPWd), decreased contractile function with increased left ventricular internal diameter at end systole (LVIDs), and decreased fractional shortening (FS%). Heart rate (HR) was unchanged. n = 11 and 12 mice per group, and a mean of 3 measurements for each mouse. (G) Gravimetric analysis showing increased heart weight normalized to body weight (HW/BW) after Rpsa knockout. n = 11and 12 mice per group. * $P \le 0.05$, ** $P \le 0.01$, *** $P \le 0.001$, **** $P \le 0.0001$, by unpaired Student's t test. Data are presented as individual values with box plot displaying the median with 25th and 75th percentiles.

hybridization (smFISH) with probes directed at the ribosomal 18S RNA to identify ribosomes and probes targeting the Cre mRNA sequence to identify transduced cells in 6-month-old mice. While HA-CRE signal was nuclear due to the nuclear localization signal in CRE proteins, the Cre mRNA signal was expected to be both nuclear and cytoplasmic and showed a perinuclear accumulation. In both control and Rpsa-knockout mice, Cre-negative cells maintained a typical striated pattern when probed for 18S rRNA, indicative of normal ribosomal localization (8) (Figure 7). However, in Rpsa-knockout mice, Cre-positive cells exhibited a loss of this striation pattern, showing a disruption in the normal sarcomeric localization of ribosomes. RPSA loss disrupted ribosome localization but did not eliminate ribosomes, explaining why localized translation was lost in the OPP assay while protein translation was not completely abrogated. We also examined mRNA localization using dual smFISH with probes directed at polyA to identify mRNAs and probes targeting the Cre mRNA (Supplemental Figure 5). In control cardiomyocytes, mRNAs displayed the expected cross-striated distribution (7) (Supplemental Figure 5). Due to the small size of the Rpsa-knockout cardiomyocytes, we could not reliably quantify and determine if the mRNA distribution in these cells was intact but in some knockout cardiomyocytes we could observe a cross-striated pattern.

The N-terminal domain (amino acids 1-209) of RPSA is globular highly conserved region that shares high resemblance to the prokaryotic ribosomal proteins S2 (RPS2) family (10). This region was shown to be sufficient for ribosomal binding. The C-terminal domain (amino acids 210-295) is absent in prokaryotes and has an intrinsically disordered structure. We therefore wanted to verify if the C-terminal domain of RPSA was sufficient for localization to the sarcomere. We expressed HA-tagged mCherry-RP-SA-C-terminal fragment₂₁₀₋₂₉₅ in mice using AAVs. A staining of cardiac sections from these mice with anti-HA and anti-ACTN2 antibodies showed that the C-terminus of RPSA was sufficient to localize the construct at the same location as endogenous RPSA and ribosomes, on both sides of the Z-line (Supplemental Figure 6, A and B). A biochemical study showed that RPSA directly binds tubulin with high affinity (13), suggesting this interaction was mediated by the C-terminal domain of RPSA. Our protein quantification showed that knockdown of Rpsa resulted in reduced tubulin levels in cardiomyocytes (Figure 6, K and L), but staining of microtubules in Rpsa-knockdown cardiomyocytes showed that,

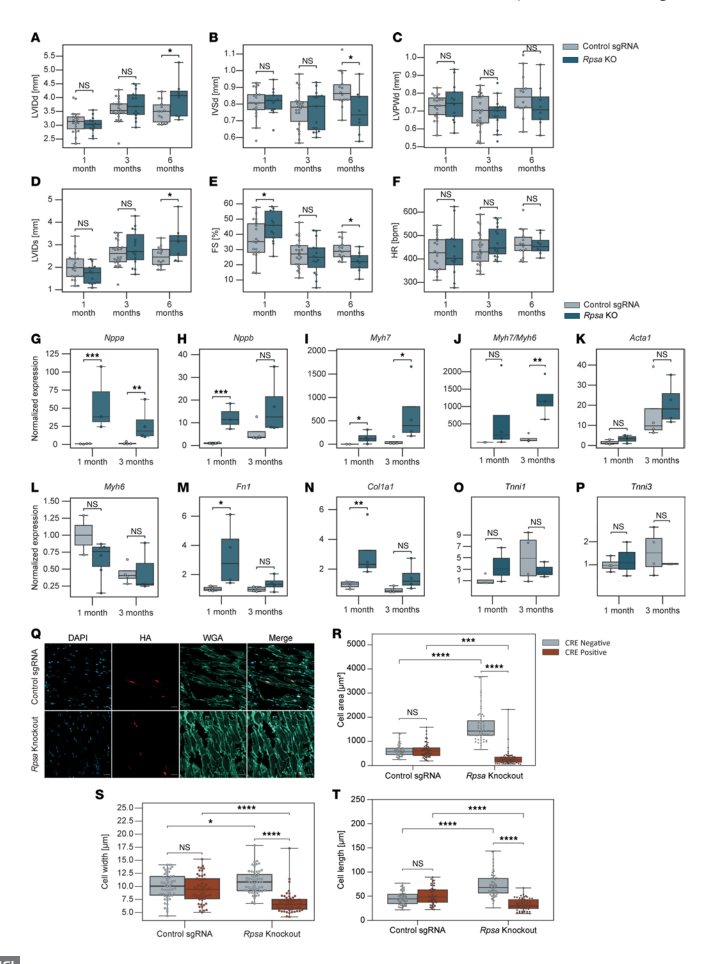


Figure 5. Cardiomyocyte cell autonomous loss of Rpsa results in cardiac dysfunction and cardiomyocyte atrophy. (A-F) Echocardiographic assessment of mice with Lox-stop-Lox-CAS9 transduced with AAVs encoding for sgRNAs targeting Rpsa or control sgRNAs and troponin promoter-driven HA-CRE expression to knockout Rpsa specifically in cardiomyocytes. Analysis shows Rpsa knockout results in cardiac dilation with increased LVIDd, decreased IVSd and LVPWd, and decreased contractile function with increased LVIDs and decreased FS percentage. HR was unchanged. n = 8, 12, 13, 14, 15, 19, and 22 mice per experimental group; mean of 3 measurements per mouse. (G-P) Gene expression analysis by RT-qPCR shows an increase in markers of cardiac stress and hypertrophy (Nppa, Nppb, Myh7, Myh7/Myh6 ratio) as well as in markers of fibrosis Fn1 and Col1a1 in Rpsa-knockout hearts. Tnni1 and Tnni3 gene expression had no statistically significant alteration. Expression data are normalized to Gapdh. n = 2, 3, and 4 mice per age and experimental group; 2–3 technical replicates per mouse. (Q-T) Representative images and quantitative analysis in Rpsa-knockout hearts showing decreased cardiomyocyte width, length, and area in targeted HA-CRE-positive cardiomyocytes, indicating atrophy, and an increase in nontargeted HA-CRE-negative cardiomyocytes, indicating hypertrophy. Sections were stained with WGA (light green), HA-tag antibody (red) to identify nuclear HA-CRE, and DAPI (blue) for nuclei. n =51, 53, 54, 57, 58, and 60 cells from 2-3 mice per group. Scale bar: 20 μm. * $P \le 0.05$, ** $P \le 0.01$, *** $P \le 0.001$, **** $P \le 0.0001$, by Student's t test (A-P) or 1-way ANOVA test (R-T). Data are presented as individual values, with box plots displaying the median with 25th and 75th percentiles.

although the levels of microtubules appeared lower, the overall distribution patterns did not change (Supplemental Figure 6C). In contrast, the disruption of the microtubular network in cultured adult rat ventricular cardiomyocytes (ARVMs) by either colchicine or nocodazole resulted in loss of RPSA sarcomeric localization and perinuclear accumulation of RPSA (Supplemental Figure 6, D-F). These observations support the role of microtubules in RPSA localization to the sarcomere.

Finally, we examined the distribution and expression of RPSA in adult hypertrophic *Mybpc3*_{c.2373InsG} mice, which are homozygous for the hypertrophic cardiomyopathy Dutch founder mutation (14). Immunostaining showed that RPSA was positioned on both sides of the Z-line in both hypertrophic and in wild-type cardiomyocytes (Supplemental Figure 7, A and B). Expression analysis on the other hand showed a strong trend toward downregulation of *Rpsa* in the hypertrophic hearts (Supplemental Figure 7C).

Discussion

Here, we used proximity ligation to unravel the Z-line proteome and identified RPSA as a potential localization candidate. We show that RPSA is required for localization of ribosomes, for the localization of translation, and for sarcomere maintenance in cardiomyocytes in vivo. The loss of RPSA causes the ribosomes and translation to lose their sarcomeric localization, eventually leading to dilated cardiomyopathy. In genetic mosaic mice the untransduced cells undergo compensatory hypertrophy in response to the atrophy of the *Rpsa*-knockout cardiomyocytes. RPSA is known to bind the microtubular system (13), and we also show here that the C-terminus extraribosomal domain of RPSA is sufficient for localization to the Z-line of the sarcomere and that disruption of the microtubules results in RPSA collapse around the nucleus.

We used Cypher (LDB3) as our bait for the spatial proteomic analysis. This protein interacts through its N-terminal PDZ domain with the Z-line protein α -actinin (15), and BioID2 was tethered

to its C-terminal part. We showed that the bait is localized to the Z-line, and our proteomic analysis identified multiple know Z-line proteins, including α -actinin, other PDZ-LIM proteins, such as Enigma (Pdlim7) and Enigma Homologue (Pdlim5), Palladin, and Myozenin, confirming that we specifically enriched for Z-line proteins. We also identified multiple proteins associated with ribosomes and protein translation, providing further evidence that this region is an important protein translation hub. A previous study used proximity labeling experiment to unravel the Z-line proteome (16). As compared with our study, this study used human iPS-derived cardiomyocytes, which are less structurally mature than neonatal rat cardiomyocytes, a different bait, α-actinin instead of Cypher, and a different control — no BioID as opposed to untethered BioID here. While several ribosomal proteins were identified in this study, RPSA was not among the significant hits, and interestingly, IGF2BP2, a major hit, was not identified in our study. The width of the Z-line measured by electron microscopy is 130 nm (17), and the exact location of the C-terminus of Cypher within the structure is not known. Given the uncertainty regarding the exact localization of Cypher C-terminus and that BioID has a practical labeling radius estimated at approximately 10 nm (18), it is difficult to determine precisely where the ribosomal and translation nanodomain resides inside or near the Z-line. Our coimmunostaining of RPSA and α-actinin suggests that ribosomes or at least RPSA mostly reside on both sides of the Z-line at the lateral ends of the I-band. We speculate therefore that we mostly isolated proteins from a nanodomain located on both sides of the of the Z-line while the α-actinin-BioID study identified a domain that is more at the center of the of the Z-line.

There is controversy over the location and functions of RPSA. RPSA was separately identified as a nonintegrin laminin receptor with a 67-kDa molecular weight from muscle cell membranes (19) and as a component of the 40S ribosome with a 32- to 37-kDa molecular weight (20) and was called 37/67-kDa laminin receptor (LAMR). It was reported to reside in the plasma membrane, the ribosomes, the cytosol, and the nucleus (10). There are many unanswered questions regarding the cellular biology of this protein, about its ability to bind laminin, and about the function of its intrinsically disordered C-terminal region (10). While RPSA's N-terminal domain (amino acids 1-209) is homologous to the bacterial small ribosomal protein RPS2, its C-terminal domain (amino acids 210-295) is not found in prokaryotes. It was therefore proposed that the N-terminal domain binds ribosomes while the C-terminal domain of RPSA functions in laminin binding. Nevertheless, it was also proposed that the C-terminal domain of RPSA participates in ribosome binding (21).

It is also unclear how RPSA functions as a ribosomal protein. It is highly abundant, with 6–8 copies per ribosome and does not appear to be a constitutive component of ribosomes (21–23). We found that in cardiomyocytes RPSA is localized to both sides of the Z-line and near the intercalated discs. This location coincides with our previously documented location of ribosomes (7, 8) and, together with the proteomic analysis here, which identified other ribosomal protein at this site, suggests that RPSA is predominantly associated with ribosomes near the Z-lines of cardiomyocytes. Our polysome profiling showed that RPSA was associated with monosomes and polysomes, but it is unclear whether all cardiomyocytes' ribosomes contain RPSA

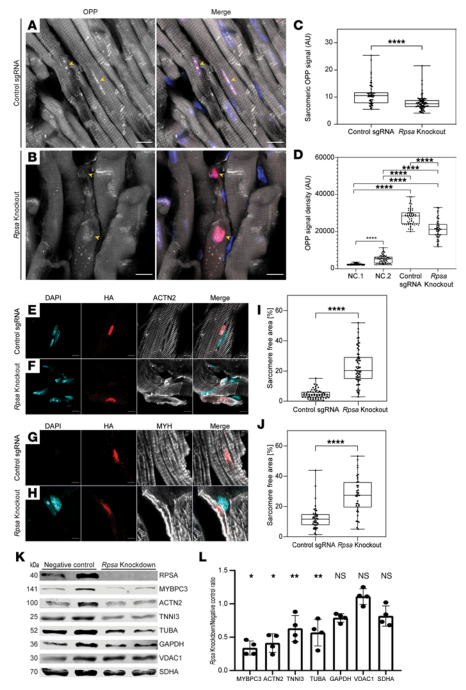


Figure 6. *Rpsa* knockout disrupts local sarcomeric protein translation and sarcomere integrity in vivo. (A–D) OPP-injected mosaic *Rpsa*-knockout and control mice were stained for OPP (white) and HA-CRE (red) to label nascent proteins and targeted cardiomyocytes, respectively. Representative images show OPP localized sarcomeric translation pattern in control mice (A). In *Rpsa*-knockout mice, the localized sarcomeric translation is lost only in targeted cardiomyocytes (B). Yellow arrowheads mark targeted nuclei. DAPI (blue). Scale bar: 10 μm. (C) Line-scan quantification in targeted cardiomyocytes confirms the loss of localized sarcomeric translation in *Rpsa*-knockout cardiomyocytes. n = 2 mice per group; n = 57 and 88 cardiomyocytes. (D) The translational activity in *Rpsa*-knockout cardiomyocytes, assessed by OPP cytoplasmic integrated density, is reduced yet higher than the negative control samples (NC.1- no OPP or NC.2- no Click labeling). n = 2 mice per group; n = 63, 64, 70, and 75 cardiomyocytes. (E–J) Representative images and analysis showing disrupted ACTN2 and MYH staining in *Rpsa*-knockout cardiomyocytes, quantified by the increased percentage of the sarcomere-free area. n = 2 mice for control sgRNA, n = 3 mice for the *Rpsa*-knockout group, n = 56 and 64 (I), n = 39 and 42 (J) cardiomyocytes. (K and L) Western blot and densitometry analysis after *Rpsa* siRNA-mediated knockdown in NRVMs showing significantly reduced sarcomeric protein levels (MYBPC3, ACTN2, TNNI3) and α -tubulin (TUBA) but not in the nonsarcomeric proteins (GAPDH, VDAC1, SDHA). n = 4 biological samples from 3 independent experiments. * $P \le 0.05$, ** $P \le 0.01$, **** $P \le 0.0001$, by 1-way ANOVA test (D) or Student's t test (C, I, J, and L). Data are presented as individual values, with box plots displaying the median with 25th and 75th percentiles, and bar graph (mean ± SD).

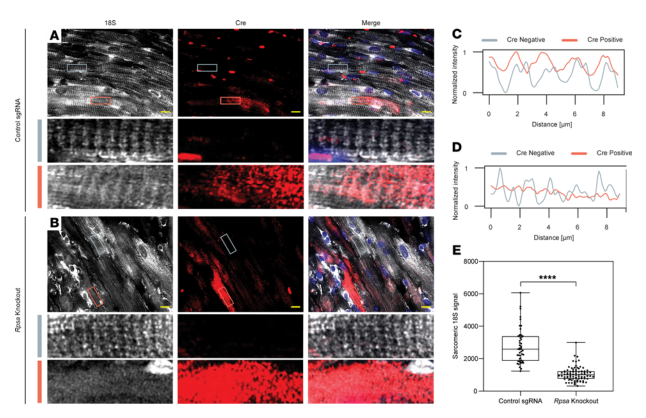


Figure 7. *Rpsa* knockout disrupts sarcomeric ribosome localization in vivo. (**A** and **B**) Dual smFISH with probes for ribosomal 18S RNA (white) and *Cre* mRNA (red) was used to visualize ribosomes and targeted cardiomyocytes, respectively. Representative images show sarcomeric cross-striated pattern of ribosomes localization in both targeted and untargeted cardiomyocytes in negative control mice (**A**). In *Rpsa*-knockout mice, the sarcomeric localization of ribosomes is lost in targeted *Cre*-positive cardiomyocytes, while untargeted cells are normal (**B**). High-magnification images (magnified 6.5 times the original image) of the boxed areas showing untargeted cytoplasmic *Cre* mRNA-negative cell (gray rectangle) and targeted cytoplasmic *Cre* mRNA-positive cell (light red rectangle) for both the negative control (**A**) and *Rpsa*-knockout (**B**) hearts. Scale bar: 10 μm. (**C** and **D**) The intensity of ribosomal 18S RNA FISH signal along the cardiomyocytes in the boxed areas shows sarcomeric periodicity in both the targeted (light red line) and untargeted (gray line) cardiomyocytes in negative control mice (**C**). In *Rpsa*-knockout mice (**D**), the sarcomeric periodicity of ribosomes is lost in the targeted *Cre*-positive cardiomyocyte (light red line), while the untargeted cell has normal periodicity (gray line). (**E**) Line-scan quantification confirms the loss of localized sarcomeric 18S signal in *Rpsa*-knockout cardiomyocytes. *n* = 2 mice per experimental group; *n* = 49 and 83 cells. ****P ≤ 0.0001, by Student's *t* test. Data are presented as individual values, with box plot displaying the median with 25th and 75th percentiles.

or use it for localization. We showed that the sarcomeric localization of ribosomes was lost after deletion of RPSA in vivo. In cardiomyocytes and muscle cells, the bulk of the transcriptome is located along the Z-line (7, 8), including transcripts for nonsarcomeric proteins (9). We examined mRNA localization in Rpsa-knockout cardiomyocytes using dual smFISH for polyA. The small size and relatively weak signal of dual-color FISH made it difficult to determine whether the cross-striated localization of mRNA was still intact in these knockout cardiomyocytes. However, we and others have previously shown that the sarcomeric mRNA distribution does not depend on ribosomes and translation (7, 9). The untethering of ribosomes from the Z-line should increase the distance between ribosomes and mRNAs, resulting in a decrease in protein translation. Indeed, we show that localized translation in cardiomyocytes was lost after knockdown of Rpsa in cultured cardiomyocytes or by CRISPR knockout in vivo with a loss of sarcomeric proteins.

Although knockout of *Rpsa* resulted in cardiomyocyte atrophy, we could identify knockout cardiomyocytes, even several months after RPSA ablation, and could show that these cardiomyocytes contained ribosomes and exhibited translation, albeit reduced and unlocalized. We also did not observe a significant decline in the

numbers of knockout cardiomyocytes between 1 and 6 months of age or evidence for apoptosis. Considering the limited half-lives of proteins in the heart (24), these results strongly suggest that RPSA is not required for ribosome biogenesis or protein translation in general. The loss of sarcomeric but not of mitochondrial proteins after *Rpsa* knockdown also supports this view. However, we cannot rule out that RPSA has additional ribosomal functions in addition to localization. This question is further complicated by the fact that ribosomes are responsible for the translation of ribosomal proteins and, therefore, the untethering of ribosomes from the Z-lines and from the localized mRNAs is also expected to reduce the translation of ribosomal proteins, leading to further reduction in translation. Our data also suggest that the atrophy and functional loss of approximately 15% of cardiomyocytes is sufficient for the development of heart failure.

Proteomic pull-down in fibroblast showed that RPSA interacts with cytoskeletal elements in addition to its interactions with multiple ribosomal proteins (25). In particular, elements of the microtubular system, including tubulin and Dynein cytoplasmic 1 intermediate chain 2, a molecular motor protein responsible for transport of proteins along microtubules, were identified. A biochemical study

showed that recombinant RPSA, but not the bacterial ortholog rps2, directly binds both tubulin and actin with high affinity, suggesting that the C-terminal domain of RPSA, which is absent in bacteria, mediates this interaction (13). This study also suggested that ablation of the microtubular system results in altered distribution of ribosomes in cultured fibroblasts. We have previously shown that the sarcomeric Z-line association of ribosomes and translation is lost upon ablation of the microtubular system (7, 8). Together with our observations that the C-terminal domain of RPSA is sufficient for binding to the Z-line, and that RPSA loses its sarcomeric localization after disruption of the microtubular system, these data imply that the ability of RPSA to bind to the microtubular system by its C-terminal domain and its ability to bind ribosomes are responsible for Z-line localization of ribosomes and translation in cardiomyocytes. We cannot rule out, however, that the localization is mediated by binding of RPSA to additional proteins.

A spontaneously occurring mutation in Rpsa in a mouse line resulted in arrhythmogenic right ventricular cardiomyopathy, which was ascribed to nuclear functions of RPSA and binding to heterochromatin protein 1, suggesting that this gene may have a specific function in the heart (26). We show here that mice with the homozygous Dutch Mybpc3 mutation, which results in complete loss of MYBPC3, have a strong trend for reduced expression of Rpsa. Further research is needed to determine how such cardiac disease-associated Rpsa reductions contribute to disease progression. The homozygous Rpsa-knockout mice die at an early embryonic age, and therefore the necessity for this gene could not be studied in vivo (10). In this study, we used a cardiomyocyte-specific genetic mosaic knockout of Rpsa to investigate the cardiomyocyte cell-autonomous effects of RPSA loss in postnatal cells within the context of normally functioning myocardium, although the mice eventually succumbed to cardiomyopathy. A knockout of Rpsa in all cardiomyocytes would have likely resulted in profound cardiac dysfunction and would have made the dissection of the direct effects of RPSA deletion the from secondary effects difficult (27).

The role of RPSA in tethering ribosomes to microtubules has been proposed before (13), but the highly localized and cross-striated distribution of ribosomes and translation in cardiomyocytes allowed us to examine and demonstrate the necessity of RPSA for localized translation in vivo. It is likely, however, that RPSA is required for localized translation in many cell types. Multiple mechanisms were shown to be responsible for mRNA localization in different cell types and cellular locations (28), and therefore, it is possible, or even likely, that additional mechanisms and elements are responsible for ribosome and translation localization beyond RPSA in muscle cell and in additional cell types.

Methods

Sex as a biological variable. Our study examined male and female animals equally, and similar findings are reported for both.

sgRNAs. See Supplemental Table 2 for details.

Antibodies. See Supplemental Table 3 for details.

Plasmids. pAAV2/9n (Addgene plasmid 112865; http://n2t. net/addgene:112865; RRID: Addgene_112865) and pAdDeltaF6 (Addgene plasmid 112867; http://n2t.net/addgene:112867; RRID: Addgene_112867) were a gift from James M. Wilson (Perelman School of Medicine at the University of Pennsylvania, Philadelphia, PA, USA).

mycBioID2-pBABE-puro was a gift from Kyle Roux (University of South Dakota Sanford School of Medicine, Vermillion, SD, USA) (Addgene plasmid 80900; http://n2t.net/addgene:80900; RRID: Addgene_80900).

Isolation of cardiomyocytes and tissue culture. NRVMs were isolated using the Worthington Biochemical Neonatal Cardiomyocyte Isolation system (catalog LK003300) according to the manufacturer's instructions. Cardiomyocytes were separated using Percoll gradients. Cytosine β -D-arabinofuranoside (ARA-C, Sigma-Aldrich, C1768) was added to the culture medium for a final concentration of 3 μ M to inhibit the proliferation of fibroblasts. Isolation of ARVMs was performed according to the previously published protocol (29).

Production of adenoviruses and AAVs. Adenoviruses were generated as previously described with adenovirus expression plasmid containing the insert (Chypher-BioID2 or BioID2) (7). The production of AAV followed a previously published protocol (30). HEK293 cells were transfected with the pAAV plasmid, along with a helper plasmid (pAdDeltaF6) and a packaging plasmid (pAAV2/9n). For injection of AAV to mice, 50 μL containing approximately 3 \times 10 11 viral genomes were injected into the intraperitoneal cavity of 3- to 5-day-old mouse pups.

Microscopy. Samples were imaged using a Zeiss LSM 900 with Airyscan2 superresolution system, attached to an Axio Observer 7 inverted microscope. Acquisition was performed with Plan-Apochromat 20×/0.8 M27 and 63×/1.4 oil DIC M27 objectives from Zeiss. Airyscan images were processed with the ZEN software using default parameters.

For wide-field imaging, slides were imaged with Axio Observer inverted fluorescent microscope (Zeiss) using an X-cite metal-halide light source and a high-resolution camera (Hamamatsu Orca R2), with an X63/1.4NA objective (Olympus).

Microtubule ablation. The cell culture medium was replaced with a medium containing either Nocodazole or Colchicine at a final concentration of 10 μ M. After a 24-hour incubation period, the cells were fixed and stained.

Image analysis. For automated image analysis of OPP and immunofluorescence signals in NRVMs a custom tool based on the Sarcomere Texture Analysis method (11) was developed. Briefly, cells were segmented from multichannel immunofluorescence images, and algorithms were used to quantify 2 aspects of the sarcomeric structure, namely homogeneity and colocalization of different markers. To quantify the sarcomeric signal of OPP or 18S in sections, lines were drawn using Fiji software along the long axis of cardiomyocytes. We measured the mean peak-to-valley intensity along the line. We used 2 lines per cell, and the average value from these 2 lines was used as the value for the cell. To quantify the signal density, we used 2 different regions of interest of fixed size in the cytoplasm and calculated the integrated density of signal using Fiji. The average value from these regions was used as the value for the cell. To quantify the sarcomere-free area, we manually measured and divided the area devoid of sarcomeric signal in the cytoplasm by the total area of the stained cardiomyocyte in the sections using Fiji.

Immunofluorescence staining. Immunofluorescence staining was performed on isolated cardiomyocytes and frozen heart tissue sections. For immunofluorescence combined with OPP assay, immunostaining was performed first, followed by the OPP assay as described below. For WGA staining, sections underwent incubation with FITC-conjugated WGA for 1 hour at room temperature (diluted 1:10 in PBS ×1) prior to the permeabilization step.

TUNEL assay. The assessment of apoptosis was conducted using the Click-iT TUNEL Alexa Fluor 488 imaging assay (Thermo Fisher Scientific) according to the manufacturer's instructions. For the positive control, sections were treated with DNase I-containing solution for 30 minutes at room temperature prior to conducting the TUNEL assay.

Rodents. Mice and rats were maintained in a pathogen-free facility, with temperature and humidity control, and a 12-hour-light cycle, with ad libitum water and food supply. Male and female animals were equally used. The constitutive Rosa26-Cas9 knockin on B6J (RRID:IMSR_JAX:026179) and the conditional Rosa26-LSL-Cas9 knockin on B6J mice (RRID:IMSR_JAX:026175) were obtained from The Jackson Laboratory. The *Mybpc3* mutant mice (MYBPC3c.2373InsG) were generated in C75BL6/J background by J. van der Velden Laboratory. Wistar rats were obtained from the Technion animal facility.

Single-molecule FISH. Stellaris RNA FISH probes (Biosearch Technologies) were used, tagged with either 5-TAMRA or Quasar-670 fluorophores. smFISH was performed according to the manufacturer's protocol, as previously described (7). Slides were imaged with Axio Observer inverted fluorescent microscope (Zeiss, see *Microscopy*). Exposure times for smFISH signals were between 800 and 3000 milliseconds. Images were acquired as a full-thickness *Z*-stack with a 0.24 μ m section thickness. Images were imported into the ImageJ software, a Laplacian of Gaussian filter was applied to the smFISH channels using the Log3D plugin, and a maximum intensity merge of the *Z*-stack was performed.

RT-qPCR. For cell cultures, RNA was isolated using the Nucle-oSpin RNA (Macherey-Nagel, 740955) according to the manufacturer's instructions. For isolation of RNA from heart tissues, atrial tissue was removed, and a Bullet Blender homogenizer (Next Advance) was used to homogenize the whole ventricular tissue prior to the standard protocol of RNA extraction. All studies were performed with a standard curve, technical duplicates or triplicates, and controls (negative control and no reverse transcriptase control).

Polysome profiling. Whole hearts with atria removed were dissected into small pieces and homogenized with a Bullet Blender homogenizer (Next Advance) containing polysome extraction buffer (20 mM Tris-HCl pH 7.4, 200 mM KCl, 10 mM MgCl₂, 1 mM DTT, 0.15 mg/mL Cycloheximide [Sigma Aldrich, 0180-5G], 1× Protease inhibitors cocktail). The homogenates were centrifuged at 17,000g for 12 minutes at 4°C. Triton X-100 was added to a final concentration of 1%, followed by 30 minutes of incubation at 4°C. The samples were centrifuged again, and the supernatant was transferred to a prechilled new tube. Linear sucrose gradients of 10%-50% were prepared in SW41 centrifugation tubes (Beckman Coulter, Ultra-Clear 344059), using the Gradient Station (BioComp). Gradient solutions 1 (GS1) and 2 (GS2) were prepared with RNase/DNase-free UltraPure water and filtered with a 0.22 μm filter, consisting of 20 mM Tris-HCl pH 8, 200 mM KCl, 10 mM MgCl₂, 0.2 mg/ml CHX, and 10% or 50% w/v RNase-free sucrose, respectively. SW41 centrifugation tubes were filled with GS1, followed by GS2 layered at the bottom of the tube. The linear gradient was formed using the tilted methodology, with the Gradient Station Maker (BioComp). 700 μL of each lysate was loaded on top of the gradients and centrifuged at 4°C for 150 minutes at 21,000 g. Gradients were immediately fractionated using the Gradient Station (BioComp Instruments), and 24 × 450 µL fractions were collected while monitoring the absorbance at 260 nm continuously. Proteins from individual fractions were extracted by adding freshly made Trichloroacetic acid (Carlo Erba, 411527) to a final concentration of 10% v/v, and allowed to precipitate overnight at -20°C. Samples were then pelleted at 17,000g for 15 minutes at 4°C, supernatant was discarded, and the pellet was washed in ethanol and air dried for 15 minutes. The pellets were resuspended in Laemmli buffer, heated at 95°C for 5 minutes, and loaded on SDS-PAGE gels for subsequent Western blot analysis.

BioID proximity labeling assay. NRVMs were transduced using adenoviruses encoding either 3x-HA-Cypher1s-mycBioID2 (bait construct) or mycBioID2 (control construct) and incubated for 48 hours to allow for expression of the encoded constructs. Culture medium was supplemented with 50 μ M biotin, and the cells were incubated for 18 hours to allow for labeling. Subsequently cells were scraped, and pellet was then frozen in liquid nitrogen.

Pull-down of biotinylated proteins. Pull-down procedure was based on a previously published protocol (31). The supernatant was incubated with Streptavidin magnetic beads (EMD Millipore). Beads were washed twice with wash buffer 1 (2% SDS), once with wash buffer 2 (0.1% deoxycholic acid, 1% Triton X-100, 1 mM EDTA, 500 mM NaCl, 50 mM HEPES), once with wash buffer 3 (250 mM lithium chloride, 10 mM Tris-HCl, 1 mM EDTA, 0.5% NP-40, 0.5% deoxycholic acid), and 3 times with 50 mM Tris-HCl. Washed beads were resuspended in 50 mM ammonium bicarbonate.

Mass spectrometry. The proteins on the beads were reduced in 3 mM DTT, 100 mM ammonium bicarbonate, and 8 M urea; modified with 10 mM iodoacetamide in 100 mM ammonium bicarbonate and 8 M urea (room temperature 30 minutes in the dark); and digested in 2 M urea, 25 mM ammonium bicarbonate with modified trypsin (Promega), overnight at 37°C in a 1:50 (M/M) enzyme-to-substrate ratio. The tryptic peptides were desalted using homemade C18 stage tip, dried, and resuspended in 0.1% formic acid. The peptides were resolved by reverse-phase chromatography on 0.075 × 300 mm fused silica capillaries (J&W) packed with Reprosil reversed-phase material (Dr Maisch). The peptides were eluted with linear 60 minutes gradient of 5 to 28%, a 15 minutes gradient of 28 to 95%, and 15 minutes at 95% acetonitrile with 0.1% formic acid in water at flow rates of 0.15 μL/min. Mass spectrometry was performed with a Q Exactive HF mass spectrometer (Thermo Fisher Scientific) in a positive mode (m/z300-1,800, resolution 60,000 for MS1 and 15,000 for MS2); using repetitively full MS scan followed by high collision induces dissociation (at 27 normalized collision energy) of the 18 most dominant ions (>1 charges) selected from the first MS scan. A dynamic exclusion list was enabled with an exclusion duration of 20 seconds.

The mass spectrometry data were analyzed using MaxQuant software 1.5.2.8 for peak picking and identification using the Andromeda search engine, searching against the rat proteome from the Uniprot database with mass tolerance of 6 ppm for the precursor masses and 20 ppm for the fragment ions. Oxidation on methionine and protein N-terminus acetylation were accepted as variable modifications and carbamidomethyl on cysteine was accepted as static modifications. Peptide- and protein-level false discovery rates were filtered to 1% using the target-decoy strategy. The mass spectrometry proteomics data were deposited to the ProteomeXchange Consortium via the PRIDE (32) partner repository with the data set identifier PXDO46385.

Proteomic analysis. Proteomic analysis was performed using MaxQuant and Perseus with log₂ of LFQ values. Proteins that were identified by fewer than 2 distinct peptides were removed. In addition, a protein was considered for the data set and for further

analysis if it had a valid value in 2 or more replicates in at least 1 of the groups (control or experiment). Imputation of missing values was performed using Perseus by random numbers drawn from a normal distribution. We used a cutoff of \log_2 fold change ≥ 1 and t test P < 0.05 for significantly enriched proteins. The R Enhanced-Volcano package (https://bioconductor.org/packages/devel/bioc/vignettes/EnhancedVolcano/inst/doc/EnhancedVolcano.html) was used to draw volcano plots. The ShinyGO 0.77 tool was used for pathway enrichment (33). STRING was used for visualization (34).

OPP assay. Analysis of the location of protein synthesis was performed using OPP and the Click-iT Plus OPP Alexa Fluor 488 Protein Synthesis Assay Kit (Thermo Fischer Scientific), as previously described (7). For in vivo protein synthesis assay, mice were injected intraperitoneally with OPP (0.049 mg/g). Mice were sacrificed 1 hour after OPP injection. The heart was then removed, frozen, and cryosectioned.

Statistics. Statistical methods are described in each of the figure legends and include 1-way ANOVA and unpaired, 2-tailed Student's *t* test. *P* values of less than 0.05 were considered significant.

Study approval. All procedures and protocols were approved in advance by the Technion animal care and use committee (IACUC) and were performed according to IACUC guidelines.

Data availability. The mass spectrometry proteomics data were deposited in the ProteomeXchange (https://www.ebi.ac.uk/pride/archive/projects/PXD046385), with the data set identifier PXD046385. Values for all data points in graphs are reported in the Supporting Data Values file.

Author contributions

IK conceived of and conceptualized the study, designed experiments, wrote the manuscript, and interpreted the data with intellectual input from all authors. RH, OS, TZ, and LHC conceptualized and designed experiments, conducted and analyzed experiments, and contributed to the review and editing of the manuscript. RH and OS are co-first authors, equally contributed to this work, and the author order was agreed upon based on workload distribution. IE developed tools and performed analysis. JVDV contributed resources and contributed to the review and editing of the manuscript. AS developed tools for the analysis of the data and contributed to the review and editing of the manuscript.

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