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Direct visualization of cardiac transcription factories reveals regulatory principles of nuclear architecture during pathological remodeling

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Abstract

Heart failure is associated with hypertrophying of cardiomyocytes and changes in transcriptional activity. Studies from rapidly dividing cells in culture have suggested that transcription may be compartmentalized into factories within the nucleus, but this phenomenon has not been tested *in vivo* and the role of nuclear architecture in cardiac gene regulation is unknown. While alterations to transcription have been linked to disease, little is known about the regulation of the spatial organization of transcription and its properties in the pathological setting. In the present study, we investigate the structural features of endogenous transcription factories in the heart and determine the principles connecting chromatin structure to transcriptional regulation *in vivo*. Super-resolution imaging of endogenous RNA polymerase II clusters in neonatal and adult cardiomyocytes revealed distinct properties of transcription factories in response to pathological stress: neonatal nuclei demonstrated changes in number of clusters, with parallel increases in nuclear area, while the adult nuclei underwent changes in size and intensity of RNA polymerase II loci. Fluorescence *in situ* hybridization-based labeling of genes revealed locus-specific relationships between expression change and anatomical localization—with respect to nuclear periphery and heterochromatin regions, both sites associated with gene silencing—in the nuclei of cardiomyocytes in hearts (but not liver hepatocytes) of mice subjected to pathologic stimuli that induce heart failure. These findings demonstrate a role for chromatin organization and rearrangement of nuclear architecture for cell type-specific transcription *in vivo* during disease. RNA polymerase II ChIP and chromatin conformation capture studies in the same model system demonstrate formation and reorganization of distinct nuclear compartments regulating gene expression. These findings reveal locus-specific compartmentalization of stress-activated,

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housekeeping and silenced genes in the anatomical context of the endogenous nucleus, revealing basic principles of global chromatin structure and nuclear architecture in the regulation of gene expression in healthy and diseased conditions.

Keywords

transcriptional regulation; epigenomics; chromatin structure; heart disease

1. Introduction

Eukaryotic transcription requires coordination between DNA accessibility, transcription factor binding and recruitment of RNA polymerase machinery. The mechanisms of local gene regulation, as well as the global patterns of histone modifications associated with transcription, are increasingly well understood [1–3]—but how these processes are regulated within the three-dimensional environment of the nucleus remains to be determined.

The nucleus is arranged into structural and functional compartments that facilitate the cell type-specific utilization of genetic information [4, 5]. The genome itself is partitioned into chromosome territories [6, 7] and on the sub-chromosomal scale into topologically associating domains which themselves are associated with active and inactive compartments [7, 8]. These structural features can contribute to co-regulation at the transcriptional level, forming chromatin neighborhoods in which similar gene regulation regimes prevail. One such environment is a transcription factory, defined by enrichment with active RNA polymerases and permissiveness for transcription [9]. The nucleolus, which can be readily visualized with DAPI staining in many cell types, is a common example of a nuclear compartment that houses transcription factories for RNA polymerase I [10], which synthesizes ribosomal RNA. However, RNA polymerase II transcription factories, which produce mRNA, are less well understood with more undefined properties.

RNA polymerase II factories are hypothesized to allow for efficient organization of mRNA/ncRNA transcription, based on the observation that a portion of GFP-tagged RNA polymerase II molecules appears to be fixed within the nucleus [11] and labeling of nascent RNA in HeLa cells shows an increase in intensity rather than abundance of puncta over time [12, 13]. Molecular analyses have identified that transcription factories may bring together single or multiple co-regulated gene promoters with enhancer elements and RNA polymerase II [14]. Genes may alter their association with transcription factories upon activation [15] and while stoichiometric measurements of RNA polymerase II molecules support the notion of stable compartments [15], quantitative live-cell super-resolution imaging has suggested that polymerase molecules tend to act alone and demonstrate dynamic behavior, and do not cluster into super-molecular complexes [16].

Lacking from this body of work is a direct measurement of RNA polymerase II distribution and behavior *in vivo*. Furthermore, the dynamics of transcription have not been studied in post-mitotic cells. Lastly, most studies examine reporter genes and/or fusion protein constructs to understand endogenous chromatin dynamics during transcriptional activation:

the dynamics of endogenous RNA polymerase II localization and activity, in the context of endogenous genetic loci following disease relevant stimuli, has not been explored.

To address this paucity of knowledge, we examined transcription factory behavior in the setting of heart failure, a condition in which the heart fails to meet the circulatory demands of the body. The cardiomyocyte is a terminally differentiated, post-mitotic cell that serves as a valuable model to understand endogenous structural features that facilitate transcription in the interphase nucleus. During the development of heart failure, cardiomyocytes—the contractile muscle cells of the heart—hypertrophy (i.e. they increase in size, but not number) and exhibit a pathological shift in their gene expression profile. Levels of nascent transcripts increase in hypertrophic cardiomyocytes [17], as do levels of elongating RNA polymerase II in animal models and heart failure patients [18, 19], all supportive of global transcriptional activation. The present investigation used the cardiomyocyte as a model to test the general hypothesis that in terminally differentiated cells *in vivo*, RNA polymerase II factories are dynamic entities into which genes are actively recruited in a locus specific manner. Two models were used: neonatal myocytes, which retain some proliferative potential and yet respond to disease stimuli, and adult cardiomyocytes, which do not proliferate and are centrally involved in disease pathogenesis. Our findings demonstrate global changes in endogenous RNA polymerase II localization and activity, resulting in coordinated, nucleus-wide partitioning of the genome into compartments of distinct transcriptional output.

2. Materials and Methods

2.1 Cell Culture

Neonatal rat ventricular myocytes (NRVM) were isolated using enzymatic digestion, collected and plated with DMEM supplemented with 17% M199, 1× Penicillin-Streptomycin-Glutamine (Gibco), 10% Horse Serum (Gibco) and 5% Newborn Calf Serum (Gibco). For NRVM imaging, cells were plated on glass coverslips pre-coated with laminin (10µg/ml in PBS, Sigma). After 24hr plating, media was switched to serum-free (1% Penicillin/Streptomycin [Gibco] and 1:1000 ITS [BD] in DMEM). The next day cells were treated with 10µM phenylephrine for 48hr to induce hypertrophy.

2.2 Transcriptional Run-on Assay

Transcriptional run-on assay with 5' fluorouridine (5'FU, Sigma) was performed to label nascent RNA [20, 21]. NRVMs were treated in culture with 4mM 5'FU for the indicated times after which the cells were rinsed with 1×HEPEM (65mM PIPES, 30mM HEPES, 2mM MgCl₂-6H₂O, 10mM EGTA, pH 6.9). Cells were fixed and permeabilized (3.7% formaldehyde/1×HEPEM/0.5% Triton X-100), underwent a series of washes (1×HEPEM, PBS, 0.05% Tween20/PBS) and incubated with primary antibody (1:50 in PBS for BrdU, Sigma [1:100 for all other antibodies]). The cells were then washed with PBS and 0.05% Tween20/PBS, incubated with secondary antibody/DAPI/PBS (1:100) and mounted with Prolong Gold.

To examine RNA polymerase II/III activity, NRVMs were supplemented with 2µM CX-5461 (Selleckchem), an RNA polymerase I inhibitor [22], for 15min prior to the addition of 5'FU.

The addition of 4mM 5'FU also included CX-5461. To differentiate nucleolar versus nucleoplasmic transcription, cells were colabeled with nucleolin to mark the nucleolus. 5'FU intensity analyses were carried out using FIJI/ImageJ software. Mann-Whitney tests were used for statistical analyses for comparing control and hypertrophic cardiomyocytes. To test the specificity of RNA labeling, samples were treated with 0.2mg/ml RNase A (in 0.2% Tween20/PBS, Thermo Scientific) for 45min and rinsed with PBS prior to the incubation with primary antibody.

2.3 Cell and Nuclear Size Analyses

For cell/nuclear size measurements, NRVMs were fixed with formalin and then washed with PBS and permeabilized with 0.1% Triton X-100/PBS. Cells were stained with Phalloidin/DAPI/PBS, washed with PBS and mounted using Prolong Gold. Area and circularity measurements were performed using FIJI/ImageJ and statistical analyses were performed using Mann-Whitney test. For correlations, statistical significance was determined using Pearson correlation coefficient.

2.4 Animal Models

All animal procedures were performed in compliance with NIH Guide for the Care and Use of Laboratory Animals and approved by the UCLA Animal Research Committee. Adult male C57/Bl6 mice (8weeks of age, Jackson Lab) were subjected to transverse aortic constriction (TAC) surgery until the development of heart failure, as indicated by cardiac dimensions and functional parameters monitored by echocardiography. Analyses were performed on animals in the heart failure state (~6weeks post surgery). Whole hearts and corresponding liver tissue from the same mice were fixed with formalin and paraffin-embedded to be used for imaging. For a subset of samples, isolated cardiomyocytes and homogenized brain and liver tissues were used for RNA, protein and CHIP analyses.

2.5 Adult Mouse Cardiomyocyte Isolation

Adult mouse cardiomyocytes were isolated using a Langendorff system [23]. Mice were injected with heparin and then anesthetized with pentobarbital, after which hearts were excised and cannulated. Hearts were perfused with Tyrode's calcium-free solution (13mM NaCl, 0.54mM KCl, 0.06mM NaH₂PO₄, 0.1mM MgCl₂, 1mM HEPES, 10mM glucose, pH 7.37), digested with 0.1mg/ml Protease (Sigma)/0.7mg/ml Collagenase, Type 2 (Worthington) prepared in Tyrode's solution, then washed with KB solution (25mM KCl, 10mM KH₂PO₄, 2mM MgSO₄, 5mM HEPES, 20mM glucose, 20mM taurine, 5mM creatine, 100mM glutamic acid-potassium salt, 10mM aspartic acid, 0.5mM EGTA, pH 7.18). Atria and right ventricle were removed, and left ventricular myocytes were dissociated into solution. Cells were spun and washed with PBS. Cell pellets were distributed for subsequent RNA, protein and CHIP analyses.

2.6 Immunolabeling (Tissue)

Formalin-fixed paraffin embedded tissues were deparaffinized and rehydrated with serial washes (xylene, 100% ethanol, 95%, 70%, dH₂O, PBS), underwent heat-mediated antigen retrieval in 10mM sodium citrate/0.05% Tween20 buffer using vegetable steamer, then

allowed to cool to room temperature. For immunofluorescence, samples were washed with PBS, blocked with 5% BSA/PBS and then incubated with primary antibody (prepared with 2.5% BSA; 1:100 dilution). Tissue was then washed with 0.05% Tween20/PBS, incubated with secondary antibody (1:100 in PBS), washed with PBS and mounted using Prolong Gold.

2.7 Generation of DNA FISH Probes

For labeling of genes, we generated DNA FISH probes as previously published, using *hdfish.nl* [24]. Gene coordinates (mm10 mouse genome) were used to obtain a list of PCR primers to be used to generate DNA products of ~200bp that tile the region of interest. For shorter genes, coordinates were expanded upstream and downstream, such that the probe covered at least 20kb. The following genomic sites were targeted: *Nppa*, gene Chr4:148,000,745–148,002,067, primers Chr4:147,997,910–148,021,634 (64 primer sets); *Atp2a2*, gene Chr5:122,453,512–122,502,225, primers Chr5:122,453,757–122,499,765 (51 primer sets); *Gapdh*, gene Chr6:125,161,851–125,165,773, primers Chr6:125,148,520–125,176,210 (60 primer sets); *Nefl*, gene Chr14:68,083,883–68,087,737, primers Chr14:68,069,935–68,102,872 (62 primer sets). After semi-quantitative PCR, products of primers that successfully produced specific amplicons of ~200bp were pooled, fluorescently labeled with AlexaFluor 647 using Ulysis Nucleic Acid Labeling Kit (Molecular Probes) and underwent column purification to remove excess label (Biorad). Labeled probes were reconstituted at 2ng/μl in hybridization buffer (1.7xSSC [saline sodium citrate buffer], 70% formamide, 50mM phosphate buffer [Na₂HPO₄/NaH₂PO₄], 10% dextran sulfate, 5x Denhardt's solution, pH 7.5) and included 20ng/μl mouse cot-1 DNA (Invitrogen) and 8ng/μl salmon sperm DNA (Ambion).

2.8 3D-DNA Fluorescence In Situ Hybridization

Formalin-fixed, paraffin-embedded tissue sections were deparaffinized and rehydrated. Samples were heated in Pretreatment Reagent (Cytocell) for 20min using a vegetable steamer, allowed to cool and washed with 2xSSC. Tissue was denatured with 60% formamide/2xSSC (70C on heat block). Concurrently, FISH probes were denatured (75C, 7min) and incubated at 37C for at least 1hr to pre - anneal. After pre-annealing, probes were added to tissue, sealed with rubber cement and allowed to hybridize at 37C (at least 24hr). Sections were washed with 50% formamide/2xSSC (42C, 2min) and 2xSSC (4x2min) and then counterstained with DAPI and WGA (Molecular Probes), washed with PBS and mounted with Prolong Gold.

2.9 RNA and Protein Preparation

Isolated cardiomyocytes were resuspended in either Trizol (Ambion) for RNA or lysis buffer (50mM Tris pH 7.4/10mM EDTA/1% SDS/0.1mM PMSF/0.2mM sodium orthovanadate/0.1mM sodium fluoride/10mM sodium butyrate/protease inhibitor cocktail tablets [Roche]). RNA was isolated using chloroform extraction, and preparation of cDNA was carried out using iScript cDNA Synthesis kit (Biorad), followed by quantitative PCR. For protein, lysates were sonicated and samples were diluted in Laemmli buffer for westerns.

2.10 Chromatin Conformation Capture Analysis

HiC data (significant interactions with $q < 0.01$ at 40kb resolution) from isolated cardiomyocytes of sham and TAC mice was used to examine gene-gene contacts of differentially expressed genes [23]. There are 1566 differentially expressed genes in response to TAC surgery with $q < 0.05$ (622 downregulated and 944 upregulated). For each of these genes, the number of interactions with other differentially expressed genes was recorded and subsetted based on whether they displayed increases or decreases in expression. The fractions of the gene-gene interactions for lost interactions (unique to the sham state) and gained interactions (specific to TAC) that showed up- or down-regulation were analyzed.

2.11 Chromatin Immunoprecipitation

For chromatin immunoprecipitation (ChIP) experiments, isolated adult mouse cardiomyocytes were fixed with 1% formaldehyde and quenched with 125mM glycine. Cells were then lysed using a dounce homogenizer and sonicated using both probe and bath sonications to obtain DNA fragments averaging ~500bp. Sonicates from 6 animals were pooled, and ChIP was performed using the ChIP -IT High Sensitivity kit (Active Motif), with 30 μ g chromatin for each IP reaction and input. Quantitation was determined by quantitative PCR by first normalizing to input and then determining fold enrichment from IgG.

2.12 Microscopy

Confocal imaging was carried out using a Nikon A1R system using 60x objective (100x was used for DNA FISH images). For DNA FISH imaging, nuclei from tissue were assessed by scanning through the z axis and 2D images were acquired selecting the plane giving the strongest FISH signal. For RNA polymerase II imaging, a 2-color in-house built stimulated emission depletion (STED) microscope was used. Samples were visualized with Atto647N (Active Motif) and Oregon Green 488 (Life Technologies) fluorophore-conjugated secondary antibodies and imaged using a 100x objective. We used 635nm excitation and 750nm depletion lasers for Atto647 visualization and 485nm excitation and 592nm depletion lasers for Oregon Green 488 labels.

2.13 Image Analysis

For RNA polymerase II distance and intensity quantifications, 2D images were processed by the following: application of a Gaussian filter, subtracting background noise and removing objects less than 50nm (based on STED microscope resolution). Imaris software (Bitplane) was then used to designate polymerase spots (clusters) and measure closest distances of center of the spots with respect to each other (Spots to Spots Closest Distance tool) as well as with the nuclear periphery (by generating a surface and using distance transformation function to calculate spot distance to surface). To determine the nuclear distribution of RNA polymerases for each nucleus (to normalize for differences in nuclear area as a complementary approach), the minimum and maximum cluster distances to the nuclear periphery were used to determine the distance range. This range was divided by 5 to generate bins, and clusters were then assigned their corresponding bin based on their

distance to the periphery for analyses. Mean intensity values were also recorded. For the top and bottom most intense clusters, the spots were ranked by intensity for each nucleus and the top and bottom 20% values were used for subsequent analyses. The data plotted represents the average of means generated from individual nuclei.

For DNA FISH analyses, we carried out quantitation of nuclear gene positioning as described [25] to segment nuclei into 5 equal concentric areas using erosion image processing and determine localization of gene loci with respect to the nuclear periphery. For distance quantifications, Imaris software was used to create surfaces to demarcate the nucleus or heterochromatin (determined based on DAPI intensity). The closest distance from the center of the FISH signal to the nuclear periphery (inside of nuclear surface) or distance to heterochromatin (outside of heterochromatin surface) was calculated using the distance transformation function.

For statistical analyses of mean intensities or distances, a Mann-Whitney test was used to determine significance while a Chi-squared test was used to assess comparisons of distributions between control and treated groups.

2.14 Antibodies

ANF (Abcam ab91250); BrdU (Sigma B8434); Desmin (Sigma D1033); Gapdh (Millipore MAB374); H3K9me3 (Abcam ab8898); H3K27me3 (Abcam ab108245); IgG (Santa Cruz sc2027); Nefl (Abcam ab72997); Nucleolin (Abcam ab22758); RNA polymerase II Ser2P (Abcam ab5095); Serca2 (Abcam ab2861); secondary AlexaFluor antibodies (Life Technologies)

3. Results

3.1 Endogenous features of transcriptional activity in cardiomyocytes

Transcriptional activation is a feature of heart disease that contributes to increased abundance of pathological mRNA and protein—these factors in turn contribute to cell growth, a necessary step in disease progression [26, 27]. To understand the regulatory properties of the cardiomyocyte nucleus underpinning pathological gene expression, a transcriptional run-on assay using 5'fluorouridine (5'FU) incorporation [20] was used to obtain a direct readout of RNA polymerase activity. This assay allowed spatial and temporal measurement of nascent RNA transcripts, for example following treatment of neonatal rat ventricular myocytes (NRVM) with hypertrophic agonists (Figures 1A, S1A, S1B). Total nascent RNA content, measured by 5'FU intensity, increased by 42% in phenylephrine (PHE)-treated cardiomyocytes. RNA polymerase I mediated transcription also increased by 19% (measured by 5'FU intensity in nucleolar regions) (Figure 1B). Nucleolar area increased by 53% in these cells, as did the ratio of nucleolar to nuclear area, 0.05 in control to 0.07 after phenylephrine treatment ($p < 0.001$), without change in number of nucleoli per nucleus (Figures 1B, S1C). This assay measured the transcriptional activity of all polymerases; because of the high transcriptional activity in the nucleolus due to RNA polymerase I, the signal observed is contributed primarily by nascent rRNA (Figure S1B). To discriminate the role of other polymerases, RNA polymerase I activity was blocked with

CX-5461 [22], revealing increases in RNA polymerases II and III activity (up by 46%; Figures 1C, S1D, S1E). While previous reports demonstrated increases in transcription with hypertrophic stress in cardiac cells [17, 28, 29], these studies are the first to provide direct visualization of these changes in the anatomical context of the nucleus.

Because both nuclear and total cell areas increase after hypertrophic stimulus (Figures 1B, 1C), we next examined their relationship (Figure S1F). The adrenergic agonist phenylephrine, which is analogous to endogenous ligands upregulated *in vivo* during heart failure and is sufficient to induce cardiomyocyte hypertrophy and pathologic gene expression [30, 31], was employed and induced a more robust response in comparison to other hypertrophic stimuli such as isoproterenol (Figures S1I, S1J). Hypertrophic cardiomyocytes (phenylephrine-treated) showed both significant increases in cell and nuclear sizes by 58% and 8% respectively without consistent gross changes to nuclear shape, indicated by circularity (Figure S1G). To determine whether this increase was proportional, we calculated the ratio of nuclear to cell areas, which was found to decrease (nuclear area dropping from 12% to 8% of the cell area), demonstrating that cell volume increases to a greater degree than nuclear volume (Figure S1G). Cardiomyocytes do not divide and their hypertrophic growth is associated with disease. Recent studies have suggested that nuclear content (most cardiomyocytes are binucleated) correlates with functionality of the cells during regenerative processes in disease [32, 33]. Interestingly, the change in nuclear to cell area ratio was consistent when cells are segregated based on nuclei number (Figure 1D). While there was an upward trend, there was no statistically significant difference in the fraction of binucleated cells in control and phenylephrine-treated cell populations (Figure S1H). In control cells, the nuclear area versus cell area relationships between mononucleated and binucleated cells were significant and consistent, with slopes of 8.22 ($r=0.655$, $p<0.001$) and 7.321 ($r=0.506$, $p=0.023$); phenylephrine altered these relationships to 11.20 ($r=0.563$, $p<0.001$) and 4.691 ($r=0.233$, $p=0.272$), respectively (Figure 1E). While the relationship between cell and nuclear area was maintained—although different—in control and phenylephrine-treated mononucleated cells (significant correlations but different slopes), this relationship was lost in binucleated cells with phenylephrine treatment. While hypertrophy was associated with increases in both cell size and transcriptional activity, these variables showed little correlation with each other in either control or hypertrophic states (Figure S1K).

3.2 Developmental state differences in stress response of transcription factories

Although neonatal cells (NRVMs) exhibit transcriptional responses reminiscent of those seen in the adult heart during disease, neonatal and adult cells are quite different in terms of morphology, metabolism, myofilament function and—linking, and perhaps driving, these other differences—gene expression. No previous analyses have examined real-time transcription in cardiac cells, and to our knowledge, no characterization of the effects of developmental stage and terminal differentiation on the endogenous transcriptional environment of any cell lineage has been reported. To investigate how terminal differentiation influences transcription in the anatomical context of the nucleus, we directly measured RNA polymerase II localization. Previous studies using live cell imaging of RNA polymerase II have demonstrated a subpopulation of polymerases are transcriptionally

active, engaged in the formation of so-called transcription factories [11]. The punctate nature of the labeling patterns of 5'FU provided support for the organization of such transcription factories in cardiomyocytes. We next sought to characterize the properties of these transcription sites in healthy neonatal and adult cardiomyocytes, as well as those in the setting of disease. Because previous studies have reported the size of transcription factories to be <200nm (below the resolution of conventional microscopy) [9, 34], super-resolution stimulated emission depletion (STED) microscopy, which improves the resolution down to 50nm, was used to characterize features of cardiac transcription factories, as marked by actively elongating RNA polymerase II (phosphorylated at serine 2 of the C terminal domain) (Figures 2A, S2A, S2B). Hypertrophic NRVMs had a higher number of detected puncta (averaging 988 in control and 1400 in phenylephrine-treated nuclei) but were comparable to control cells when normalized to nuclear area (Figure 2B). The cluster size and average intensity of polymerase puncta also did not differ between control and phenylephrine-treated cells (Figures 2B, 2C). The intensity ranges, minimum and maximum values across nuclei, between groups were also comparable (Figure S2C). To examine the distribution of clusters within the nucleus for determination of spatial organization, the clusters were ranked based on closest distance to the nuclear periphery. The maximum and minimum distances were then used to determine the range of cluster distribution across each nucleus, which was used to generate 5 bins (Figure 2D); this segmentation generates bins of equal distances to the periphery. The fraction of total clusters per nucleus across each bin did not differ between groups ($p=0.999$), nor did the average mean intensity of the clusters ($p=0.861$; Figure 2E). These findings suggest that RNA polymerase II distribution is tightly controlled within the developmentally immature cardiomyocyte.

We next tested this phenomenon in adult cardiomyocytes *in situ* within heart tissue sections (Figure S2D). Adult mice were subjected to transverse aortic constriction surgery (TAC), which induces pressure overload to the heart and causes heart failure, marked by a reduction in left ventricular ejection fraction and increased cross-sectional cardiomyocyte area [31]. Heart tissue sections were used to examine cardiomyocyte nuclei from the left ventricle labeled for endogenous RNA polymerase II using STED imaging. Interestingly, unlike the neonatal cardiomyocytes treated with phenylephrine, the number of RNA polymerase II puncta in adult mouse cardiomyocytes from TAC animal heart sections did not show significant change when compared to heart tissue sections from mice subjected to sham control (731 in sham vs 938 in TAC nuclei), but the cluster size increased with heart failure (Figures 2F, 2G, S3G). Furthermore, the mean intensity of the puncta also increased (Figures 2F, 2H, S3I). Again, when examining the cluster properties across the nucleus, the fraction of the clusters across each bin remained consistent ($p=0.989$) with a uniform increase in intensity across nuclei after TAC stress ($p<0.001$; Figures 2I, S3J, S3K, S3L). These findings reveal the behavior of endogenous transcription factories in adult myocytes (which appear to target all RNA polymerase clusters, rather than a discrete subset), highlighting previously unrecognized differences in transcription factory function.

To further investigate this observation, we examined the spatial organization of the RNA polymerase II puncta with respect to the nuclear envelope and with respect to each other. The distribution of all of the clusters as measured to the nuclear envelope increased in NRVMs, from 1625nm to 2036nm (Figures 3A, S3A), as did the absolute distances for the

most intense clusters (top 20% of clusters, as ranked per nucleus; Figures 3B, S3B, S3C). The most intense puncta were more centralized in the nucleus, and this pattern was unchanged with phenylephrine treatment (Figures S3B, S3C). Spacing between them, measured as the average distance to a spot's closest neighbor, did not differ in NRVMs (139.6nm vs 140.3nm; Figure 3C). Transcription factories were also measured by 5'FU incorporation (Figures 3D, S3D): while the number of 5'FU clusters increased with phenylephrine treatment, the density of clusters (number of puncta normalized to area) was unchanged between control and treated nuclei (Figure 3E). The distance of a 5'FU spot to the nuclear periphery increased without change to spacing between 5'FU clusters (Figures 3F, S3E, S3F), consistent with observations made with active RNA polymerase II. To confirm this observation, the distance of each RNA polymerase II cluster to the closest 5'FU was measured and found to be unchanged (134.9nm in control and 133.2nm in hypertrophic cells; Figure 3G). Additionally, we also examined colocalization between RNA polymerase II and 5'FU signals and found no changes in average correlation coefficient, determined by intensities for each channel (0.206 in control vs 0.199 in hypertrophy, $p=0.697$).

Adult cardiomyocytes analyzed from diseased hearts showed no change in distance of clusters to the nuclear periphery compared with sham hearts (Figures 3H, S3H). On the other hand, the spacing between clusters increased during disease, from 123.5nm to 125.8nm (Figures 2F, 3I). This increased distance was pronounced in the cohort of most intense puncta, where the increase was 3.2%, from 115.2nm to 118.8nm (Figure 3J). These analyses reveal distinct features between developing and mature cardiomyocytes in the response of endogenous transcription clusters to stress. In the neonatal cardiomyocyte, there is an increase in the number of RNA polymerase II factories without a change in individual cluster intensity, maintaining the global organization and spacing between clusters in response to hypertrophic stress. In contrast, adult cardiac nuclei exhibit recruitment and accumulation of active RNA polymerase II (indicated by increases in size and intensity of the clusters without a change in number).

3.3 Nuclear positioning of genes is a mechanism for differential gene expression

It has been suggested that physical positioning of genes within the nucleus influences expression, with central localization associated with expression and peripheral localization associated with silencing [35, 36]. Whether this phenomenon is true for endogenous gene loci, and in the setting of adult terminally differentiated cells, has never been examined. To assess this hypothesis, DNA fluorescence *in situ* hybridization (FISH) was used to map endogenous gene loci within the cardiac nucleus in tissue sections from healthy and diseased mouse hearts (Figure 4A) [24]. Nuclei were segmented into concentric annuli of equal areas (or bins) by erosion processing, thereby giving the measured gene locus an equal probability of falling in any one of the analyzed areas [25]. This approach served to normalize for variability in nuclear size and shape. *Atp2a2*, a downregulated gene in the failing heart [26], underwent a significant change in nuclear positioning, shifting towards the nuclear periphery (bin 1, 0.5% to 4.4% of measured loci) and away from the center (bin 5, 46.3% to 36.7%) (Figures 4B, S4). A second gene, *Nppa*, whose expression increases with disease, exhibited the antithetical pattern of change in distribution, shifting away from the periphery (7.5% to 1.1% of loci; Figures 4C, S4) [26]. These changes in gene localization were specific and not

reflective of wholesale chromatin reorganization: the control genes *Gapdh* (housekeeping gene that remains actively expressed in both states) and neuronal *Nefl* (a neuronal gene, which remains silent in healthy and diseased hearts [37]) did not show changes in distribution (Figures 4D, 4E, S4). Interestingly, genes that *can be expressed* in the heart (*Atp2a2*, *Nppa* and *Gapdh*) tended to exhibit an overall localization pattern favoring the center of the nucleus, whereas *Nefl*, which is *never expressed* in the heart, lacks preferential distribution and is uniformly distributed (Figure 4). When examining the absolute distances (rather than relative distances, as was done in Figure 4) of gene loci to the nuclear envelope, the same trend of localization was observed, although the effect was muted (differences in distance were minimal and not significant for *Atp2a2* and *Nppa*; the differences were significant for *Gapdh* and *Nefl*; Figure 5A). We attribute this effect to variation in three-dimensional nuclear morphology. As a further control, we examined gene localization in the liver. This choice is notable because the liver is centrally involved in heart failure [38], yet these target genes undergo altered expression in the heart (and not the liver), allowing us to test the hypothesis that systemic stresses that impact multiple organs and cells have cell type-specific effects due to prevailing chromatin environments in the given cell. The data indicate that expression and positioning of these genes (*Atp2a2*, *Nppa*, *Gapdh* and *Nefl*) were unaffected in liver (Figures S4, S5) supporting a model in which cell type-specific chromatin enables distinct spatial organization of transcription.

3.4 Heterochromatin surfaces are structural features of cell type-specific nuclear architecture and gene expression

Adult mouse cardiomyocyte nuclei exhibit a unique nuclear architecture visible with DAPI staining: a large heterochromatic feature is commonly observed at the center of the nucleus, in addition to a ring of heterochromatin at the periphery. Thus, there is the possibility that gene silencing may also be occurring at this centrally located heterochromatin island (Figure 5B). To test this possibility, we performed a complementary analysis in which the distance of a gene locus was measured relative to the nearest region of constitutive heterochromatin. We observed that the aforementioned DAPI patterns closely overlapped with H3K9me3, a mark for constitutive heterochromatin, and not H3K27me3 (a facultative heterochromatin mark that is found across the nucleus), and thus measured the distances between a FISH locus for an individual gene and the nearest intense DAPI surface (Figures 5B, 5C). The *Atp2a2* loci exhibited a significant decrease in average distance to heterochromatin surfaces (182nm in sham and 128nm in diseased cells) whereas the same distances for *Nppa* increased from 392nm to 453nm (Figure 5D), in agreement with the changes in gene expression and nuclear localization described above. Furthermore, the number of *Atp2a2* loci that colocalize with DAPI (that is, those with a distance equal to 0nm) also increased (35% to 45% of loci) in disease. *Gapdh* displays the opposite behavior, moving away from constitutive heterochromatin surfaces, again in agreement with changes in expression and the measurements in total nuclear orientation.

3.5 Role of chromatin features to establish nuclear architecture and transcription factories

To investigate the role of chromatin features to establish gene expression microenvironments, we examined chromatin immunoprecipitation-DNA sequencing (ChIP-

seq) data for RNA polymerase II enrichment from healthy and diseased hearts [39], confirming behavior of specific genes by ChIP-PCR in isolated adult cardiomyocytes (Figure S6). Activation of *Nppa* expression in disease was associated with enrichment of active RNA polymerase II at its promoter, whereas the converse was observed with *Atp2a2*, which is downregulated and showed reduced occupancy. *Gapdh* (constitutively expressed) had substantial enrichment of RNA polymerase II along its gene body, whereas *Nefl* (not expressed) had no enrichment. Our results from isolated cardiomyocytes were comparable to results from whole hearts (which are composed of a heterogeneous cell population); the qualitative differences may be explained by the contributions from non-cardiac cells in the ChIP-seq experiment. To test the involvement of specific chromatin domains determined by sequencing, data from chromosome conformation capture studies (Hi-C) performed in the same model [23] were used to investigate interactions amongst genomic regions constituting a unique transcription factory. For each up- or down-regulated gene, the corresponding interactions with other differentially expressed genes were examined. Of the other genes interacting with differentially expressed genes, the majority of them also change expression in the same direction: that is, upregulated genes tend to have a greater fraction of interactions with other upregulated genes and vice versa (Figure 5E). This was a dynamic process: upregulated genes gain interactions with other differentially expressed genes, 72.7% of which are also upregulated, whereas 65.8% of gained interactions of downregulated genes are also downregulated. These results indicate that genes that are co-regulated tend to congregate with each other in nuclear environments tailored for transcription and demonstrate a mechanism for coordinated transcription during stress that involves physical reorientation of genomic regions to designated sites for transcription.

4. Discussion

Our data support a model in which stress-induced changes in global transcription are mediated by recruitment or loss of RNA polymerase II molecules at transcription sites in the adult mouse cardiomyocyte nucleus, with corresponding movement of differentially expressed genes to and from these factories (Figure 6). During diseases like cancer and heart failure, there is a global shift in DNA packaging towards a more plastic phenotype as evinced by changes in histone isoforms and modifications [31, 40, 41]. Super-resolution imaging of histone H3 proteins previously provided evidence of global structural reorganization of chromatin domains in hypertrophic cardiomyocytes [30]. Recently, chromatin capture studies in adult cardiomyocytes have demonstrated changes to genomic interactions in the setting of heart failure [23] as well as cardiac development [42]. How these global arrangements affect transcriptional output is unclear and the spatial contributions to gene expression have not been examined.

In various cell models—but to our knowledge, not tested in mammalian tissues—it has been shown that gene silencing can occur at the nuclear envelope, such that recruitment represses expression and release from the lamina towards the center of the nucleus promotes expression. Imaging approaches have utilized reporter genes to complement sequencing-based DamID analyses, which have mapped genomic interactions with the nuclear lamina [35], and shown gene silencing occurs at the nuclear envelope, with anchoring mechanisms dependent on histone methylation and histone deacetylase proteins [43–45].

Our results also point to a clear role for distinct nuclear orientation of genes, with respect to nuclear periphery and chromatin microenvironments, that positions genes in the appropriate physical space for distinct transcriptional responses. Additionally, our findings support distinct mechanisms of transcriptional regulation at different stages of development and in response to pathological stress during disease—these features may be reflective of altered chromatin plasticity during cellular maturation. These novel features of transcription are reflective of cell type-specific nuclear architecture, insofar as the systemic stresses of disease did not affect gene positioning (or expression) in the liver. This study provides support for the compartmentalization of gene expression *in vivo* and quantifies the relationship between gene expression and chromatin architecture.

Consistent with earlier studies of cellular hypertrophy [17], we observed increased RNA production in hypertrophic cardiomyocytes, extending previous observations by (i) quantifying the relative contributions of different RNA polymerase isoforms and (ii) determining the anatomical compartments of the nucleus responsible for transcription in distinct developmental stages. We analyzed the relationship between global transcription and cell hypertrophy and characterized the structural organization of cardiac transcription in the mature cardiomyocyte nucleus. Increases in nuclear size are correlated with cell size in a state specific manner (ratio of nuclear to cell size areas differs between control and hypertrophic states) and is dependent on nuclear content: the correlation is lost in binucleated cells in hypertrophy, although it is unknown if these cells underwent mitosis without cytokinesis as a result of phenylephrine treatment. While there was an increase in the number of binucleated cells, it was not statistically significant, suggesting nucleation is independent of regulation of cell size. Interestingly, a direct correlation between transcript abundance, measured by total 5'FU intensity, and cell area was not observed in either control or hypertrophic states. Nevertheless, these findings suggest that mononuclear and binuclear cells exhibit distinct transcriptional programs that may contribute to the different functional capabilities of these cells [32]. In a time-course study of mouse skeletal muscle hypertrophy, an inverse relationship was observed between transcription and cell size at the peak of transcriptional activation in myofibers, whereas no association between transcriptional activity and cell size was found in control conditions [46]. It is intriguing to speculate that our measurements reflect a new state in which the cells have stopped responding to the phenylephrine and reached a steady cell size—future experiments would be required to test whether the relationship between transcription and cell size was different at a more acute time point after hypertrophic stimulation.

Characterization of the spatial organization of active RNA polymerase II clusters revealed different mechanisms of transcriptional remodeling. In hypertrophic neonatal cardiomyocytes, new RNA polymerase II/5'FU clusters formed, ostensibly to accommodate increases in transcriptional demand as the nucleus gets larger. On the other hand, adult cardiac transcription factories physically expanded in size, indicated by increases in area and intensity of RNA polymerase II puncta. This observation suggests recruitment of polymerase molecules to established factories in adult cardiomyocytes, in agreement with previous evidence from sequencing studies that indicates RNA polymerase II elongation increases with disease [18, 19]. Single molecule imaging of RNA polymerase II reveal that the majority of polymerases exist as individual molecules, are dynamic and form transient

interactions at the basal state without perturbation [16, 47]. However, RNA polymerase clusters have increased lifetime and enhanced clustering upon transcriptional activation induced by serum stimulation [47, 48], which is correlated with increased mRNA production [49]. These support that the pathological stress induced by TAC surgery affects the stabilization and properties of RNA polymerase II clusters in the cardiomyocytes. The significant changes observed with the pressure overload model of disease *in vivo*, in comparison to phenylephrine treatment in the neonatal cells, could be reflective of the severity of stress as well as varied genome plasticity in these different developmental states. During differentiation to parietal endoderm, changes in transcription rates (based on labeling of nascent RNAs) are associated with changing nuclear volumes, maintaining the activity and distribution of transcription factories across the nucleus [50]. This prior observation is supported by our data on RNA polymerase II patterning in the neonatal cell model of hypertrophy, whereas size and intensity of the polymerase clusters are enhanced in the adult cardiomyocyte. The neonatal cardiomyocyte is still maturing and has residual proliferative potential (both characteristics that are absent in the adult cardiomyocyte): these features may be the result of a nuclear architecture that maintains greater potential for dynamism in the formation of transcription factories and the reorganization of genes. In the heart, targeting of TFIIB, important for the recruitment of RNA polymerase II to disease-associated genes, can prevent pathological hypertrophy [51]. Our studies extend this observation to map the physical orientation of these transcriptional events within the nucleus.

These experiments represent the first demonstration of the endogenous localization of distinct cohorts of genes in the adult cardiomyocyte nucleus, quantifying how this localization contributes to gene expression during disease. Genes expressed in the heart were more centrally located, whereas those not expressed were randomly distributed. Surprisingly, the liver also displays similar basal localization patterns to those observed in the heart, although unlike the heart, these distributions were unaffected in an *in vivo* model of heart failure. These patterns could be explained by a number of reasons: (i) the gene locus is situated near large heterochromatin segments on the linear chromosome. In general, centromeres and telomeres converge at constitutive heterochromatin foci found centrally (chromocenters) or at the periphery. The distance of the genes on the linear chromosome to these domains could restrict the region of localization in three dimensions; (ii) the associating genomic regions in the basal setting could also determine where the gene is found, a *guilt by association* model. For example, if *Nppa* is localized in a permissive chromatin environment in the basal state, this could enable its more ready activation even in a state where it is off (a type of structural transcriptional poisoning); (iii) chromatin structure is largely determined by developmental stage, such that once chromosome territories and nuclear architecture have been established on a large scale, nuanced reorganization (such as that observed with pressure overload hypertrophy *in vivo*) may be sufficient to alter association of genes with sites of transcription and/or heterochromatin regions. Unlike stem cell to cardiomyocyte differentiation, the changes in positioning with respect to the nuclear lamina are subtler during disease progression and point to complementary influences by local epigenetic mechanisms.

Genes with similar transcriptional behavior were found to preferentially interact in the context of the endogenous nucleus. Strikingly, 70% of the gene-gene interactions were

between genes of increased expression, supporting the model wherein chromatin dynamics bring together coregulated genes in three dimensions in the cardiomyocyte. Our findings also demonstrate that a gene's local environment is important for "priming" the locus for transcriptional activation or repression, which can then be regulated by transcription factors. Analysis of transcription factories targeted by TNF α signaling showed evidence of hierarchical regulation, such that disruption of the dominant gene (by cleaving the gene to affect gene looping) had an outsized effect on the expression of other genes in the transcription factory as contrasted with disruption of more subordinate genes (which had minimal or no effect on transcription at the factory) [52]. Another approach disrupted gene activity through knockdown of regulatory ER α transcription factor, which indirectly affected the expression of interacting promoters at the shared transcription factory [14]. Identification of the specific genomic components of cardiac transcription factories will be key to understanding the features—including shared epigenetic regulation or binding of a common set of transcription factors—that recruit and stabilize genes and regulatory elements. The mechanisms that drive shifts in gene positioning are still not well understood because most studies either measure one or a few chromatin marks with great specificity in terms of genome binding (i.e. via ChIP -seq) but with little or no data on where these marks and their associated genes localize within the anatomical context of the nucleus (as is provided by studies like those in the current paper) or relative to each other (e.g. using chromatin capture technologies).

What are the factors involved in the structural dynamics observed in this study?

Decondensation of chromatin is sufficient to move gene loci away from the nuclear periphery, independent of transcription [25]. H3K9 methylation is an important signal for the recruitment and anchoring of heterochromatic loci to the nuclear lamina [53]. HDAC3 has been demonstrated to be important in keeping loci at the nuclear lamina in fibroblasts [43]. More recently, HDAC3 was shown to be a regulator of cardiac differentiation; loss of HDAC3 was sufficient for the release of cardiac genes resulting in premature cardiomyocyte differentiation [54]. In the cardiomyocyte, overexpression of histone deacetylase, HDAC4, has been shown to decrease the expression of a subset of genes, which was also linked to their displacement from the nuclear pores [55]. These observations provide support for the importance of three dimensional physical orientation, as regulated by histone-modifying enzymes, to determine regulation of gene expression.

Pathological transcription during heart failure exhibits at least two distinct behaviors: upregulation through pause-release of RNA polymerase II (employed for housekeeping genes), and *de novo* recruitment of RNA polymerase II (employed for disease-associated loci) [39]. A new frontier is now to understand how these categories of genes are physically localized in the nucleus using imaging approaches. The present investigation sets the foundation to search for cell type-specific chromatin features that have evolved for bespoke transcriptional requirements of different cells and to understand how spatial organization of genes regulates their function in health and disease.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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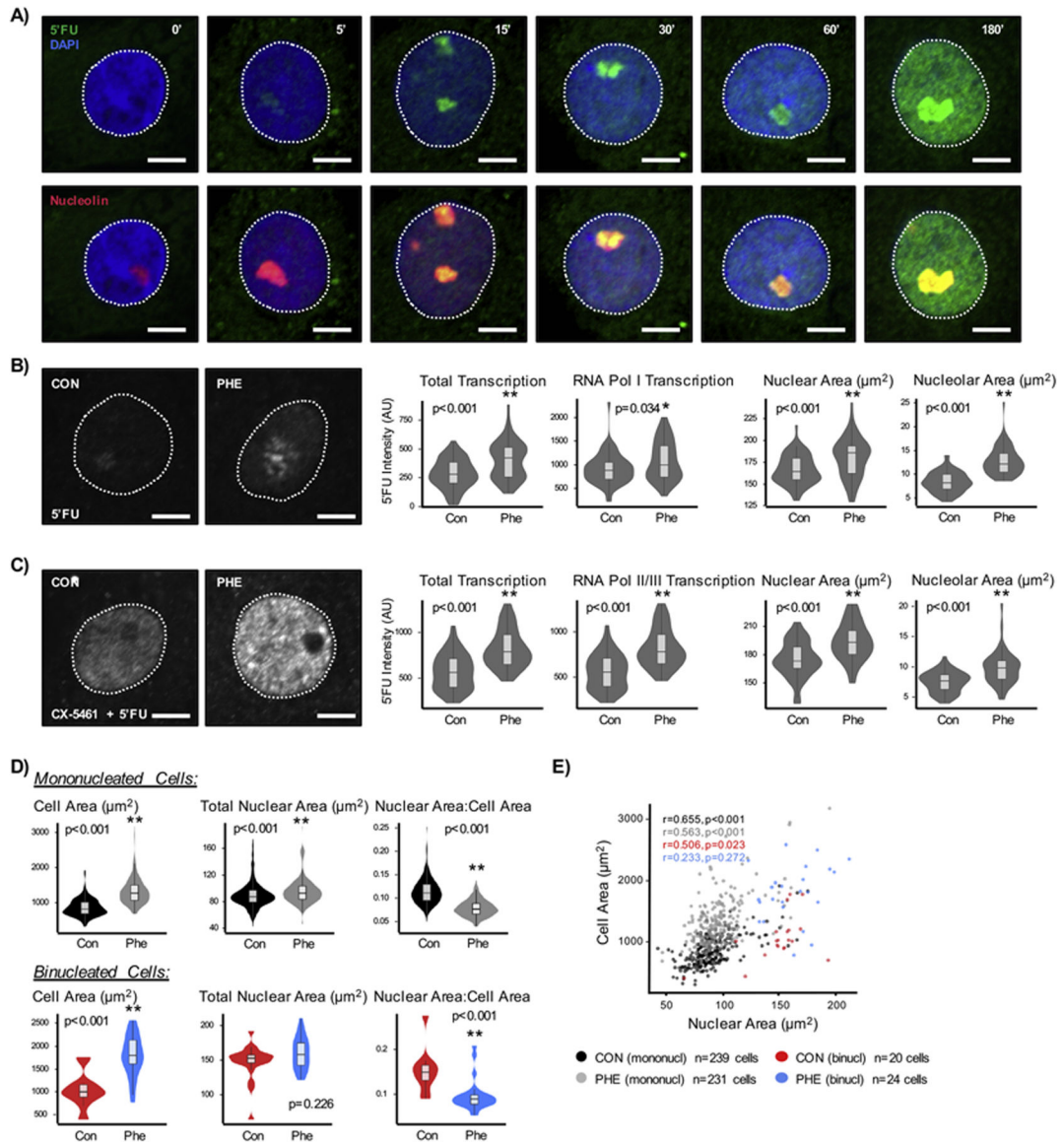


Figure 1. Endogenous transcriptional activity in cardiomyocytes following pathological stimulation.

A) Neonatal rat ventricular cardiomyocytes (NRVM) were treated with 5' fluorouridine (4mM) for the indicated times. There is a large enrichment of signal in the nucleolus (marked by nucleolin) that increases with time. The nucleolus is occupied by RNA polymerase I molecules, which carry out ribosomal RNA transcription and quench the 5' fluorouridine due to high transcriptional rates; signal is detected in the nucleolus within 5min of 5' fluorouridine treatment. **B)** NRVMs were treated with phenylephrine (10 μM , 48hr) to induce hypertrophy and treated with 5' fluorouridine to assess transcriptional activity. Representative images of nascent transcripts in nuclei are shown (left). 5' Fluorouridine intensities were measured across the entire nucleus (total transcription) or within nucleolus (marked by nucleolin, RNA Pol I transcription) (center), with distribution of nuclear and nucleolar areas (right). Average nucleoli number is not affected with treatment ($p=0.803$). Control: $n=60$ nuclei; Phenylephrine: $n=58$ nuclei. **C)** RNA polymerase

II and III activities were measured by inhibiting RNA polymerase I activity with CX-5461 (2 μ M). We observe 45% and 46% increases in total and nucleoplasmic 5'fluorouridine intensities, respectively. *Control: n=50 nuclei; Phenylephrine: n=47 nuclei. Data is representative of >3 biological replicates.* **D)** The relationship between cell and nuclear areas were analyzed in hypertrophying cardiomyocytes; measurements were subsetted based on nuclei number. The increase in cell and nuclear areas, as well as reduction in nuclear:cell area ratio, was consistent in the mononucleated cell population. Binucleated cell populations had larger cell areas in hypertrophic cardiomyocytes, but total nuclear area did not differ between control and hypertrophic states. **E)** The nuclear versus cell sizes were plotted for the different cell populations. Mononucleated cells show significant correlation between nuclear versus cell sizes (control: $r=0.655$, $p<0.001$; phenylephrine: $r=0.563$, $p<0.001$). On the other hand, binucleated cells lost this association when treated with phenylephrine (control: $r=0.506$, $p=0.023$; phenylephrine: $r=0.233$, $p=0.272$). *Scale bar = 5 μ m. * $p<0.05$; ** $p<0.001$ [Mann-Whitney].*

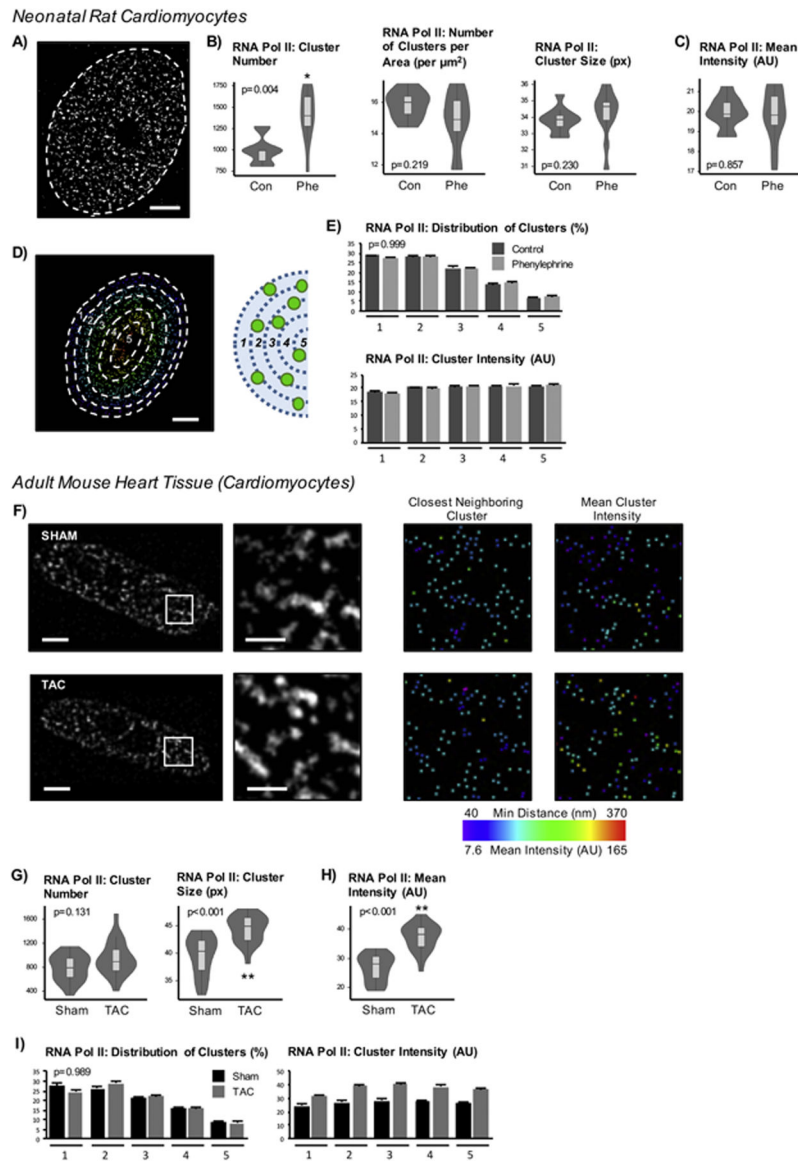
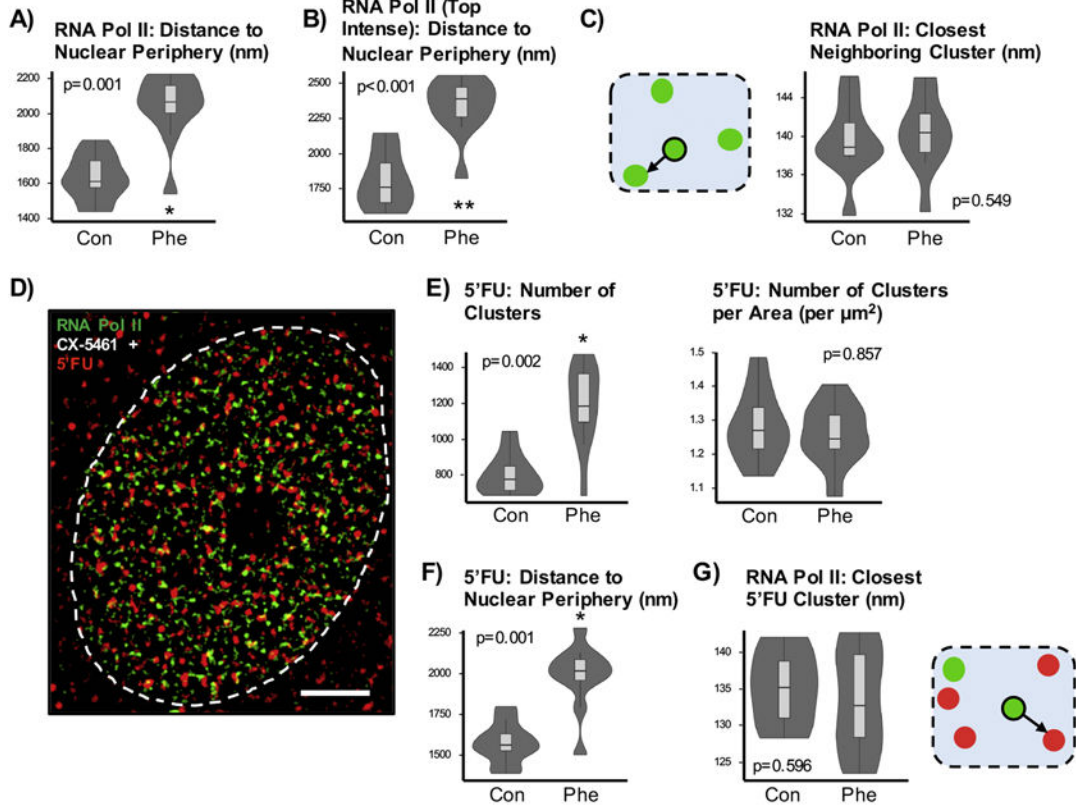


Figure 2. High transcriptional activity correlates with increases in cluster number in neonatal myocytes and increases in mean intensity and size in adult cardiomyocytes.

A) NRVMs were labeled with RNA polymerase II for STED imaging of cardiac transcription factories which were quantified using Imaris (non-nuclear signal was subtracted as background). *Scale bar = 2μm.* **B)** The number of clusters increases while the cluster density (spot number normalized to nuclear area) or size indicate no difference. **C)** Mean cluster intensity also does not differ. Together, these data suggest that increases in transcriptional activity are due to recruitment and activation of polymerases in the NRVM nucleus after agonist treatment. **D)** To examine the spatial distribution of puncta, the nucleus was divided into 5 bins based on the maximum and minimum distances to the periphery for each nucleus and then assessed cluster number and cluster intensity at each bin (*1* is closest to and *5* is furthest from periphery). *Scale bar = 2μm.* **E)** The distribution of percent of total clusters per nucleus found within each bin is plotted (*top*) along with the mean intensities of the clusters (*bottom*). Treatment does not affect intensity distribution ($p=0.861$, two-way

ANOVA). **F)** RNA polymerase II labeling was performed in adult mouse heart tissue sections in animals subjected to sham or TAC surgery, and cardiomyocyte nuclei from the fixed tissue sections were used for analyses (nuclei were selected based on co-labeling with desmin to ensure non-cardiomyocyte nuclei were not incorporated into analyses). Representative nuclei of RNA polymerase II-labeled nuclear are shown with respective magnifications and examples of subsequent cluster analyses. *Scale bar = 3 μ m (1 μ m for magnification).* **G)** RNA polymerase II total cluster number is not affected, but the mean cluster area does increase after pressure overload stress. **H)** Mean intensity of total active RNA polymerase II puncta is higher in nuclei from failing hearts. **I)** The distribution of spots across the nucleus does not differ between sham or TAC mice while the increase in intensity occurs uniformly across the nucleus as indicated by average of mean cluster intensity at each bin ($p < 0.001$, two-way ANOVA). *Control: n=10 nuclei; Phenylephrine: n=11 nuclei. Sham: n=21 nuclei; TAC: n=20 nuclei. (Imaging experiments, with different sets of control/phenylephrine -treated cells and sham/TAC hearts, were repeated >3 times. Representative dataset of analyses is presented here.) * $p < 0.05$; ** $p < 0.001$ [Mann-Whitney; for cluster distribution, Chi-squared]. Error bars represent SEM.*

Neonatal Rat Cardiomyocytes



Adult Mouse Heart Tissue (Cardiomyocytes)

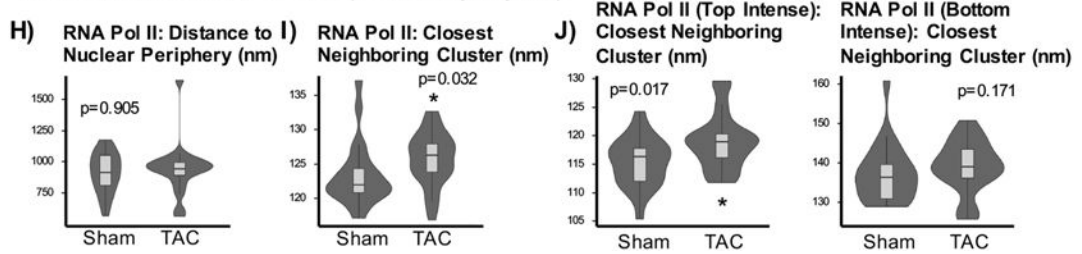


Figure 3. Distinct spatial and intensity properties of RNA polymerase II factories in neonatal and adult cardiomyocytes.

A) In neonatal cardiomyocytes, the distances from the center of each cluster to the nuclear periphery increases. **B)** For each nucleus, the clusters were ranked by mean intensity and the top 20% were used for analysis. The absolute distance to the periphery of the top-ranked clusters increases (highly intense clusters found more centrally). **C)** To assess anatomical distribution of cardiac transcription factories, we analyzed the distances to the closest neighboring spot. The cartoon illustrates RNA polymerase II clusters in green. For each spot, the distance to the nearest neighbor is recorded. The mean spacing between RNA polymerase II clusters remains similar between control and hypertrophic NRVMs. **D)** Analyses integrating 5' fluorouridine measurements, a direct readout of RNA polymerase activity, were performed. Neonatal myocytes were treated with CX-5461 and then supplemented with 5' fluorouridine for 30min. *Scale bar = 2 μm.* **E)** Like RNA polymerase II, the absolute number of 5' fluorouridine clusters increases though not when normalized to

nuclear area. **F)** The distance of 5' fluorouridine clusters to the nuclear periphery increases, consistent with measurement patterns of RNA polymerase II. **G)** The distance of closest neighboring 5' fluorouridine (*red*) with respect to RNA polymerase II cluster (*green*) does not significantly change, although the trend is towards a decrease. **H)** In adult cardiomyocytes from sham and pressure overload sections, the distance of clusters to the nuclear envelope does not change after TAC surgery. **I)** Unlike in neonatal cardiomyocytes, the average distances between closest neighboring RNA polymerase II clusters show change in spacing in the adult cardiomyocyte in the disease setting (123.5nm to 125.8nm). **J)** Examination of top intense clusters also shows increased spacing between close RNA polymerase II neighbors (115.2nm to 118.8nm) while the least intense clusters do not show changes in spacing between neighboring puncta. *Control: n=10 nuclei; Phenylephrine: n=11 nuclei. Sham: n=21 nuclei; TAC: n=20 nuclei. * p<0.05; ** p<0.001 [Mann - Whitney].*

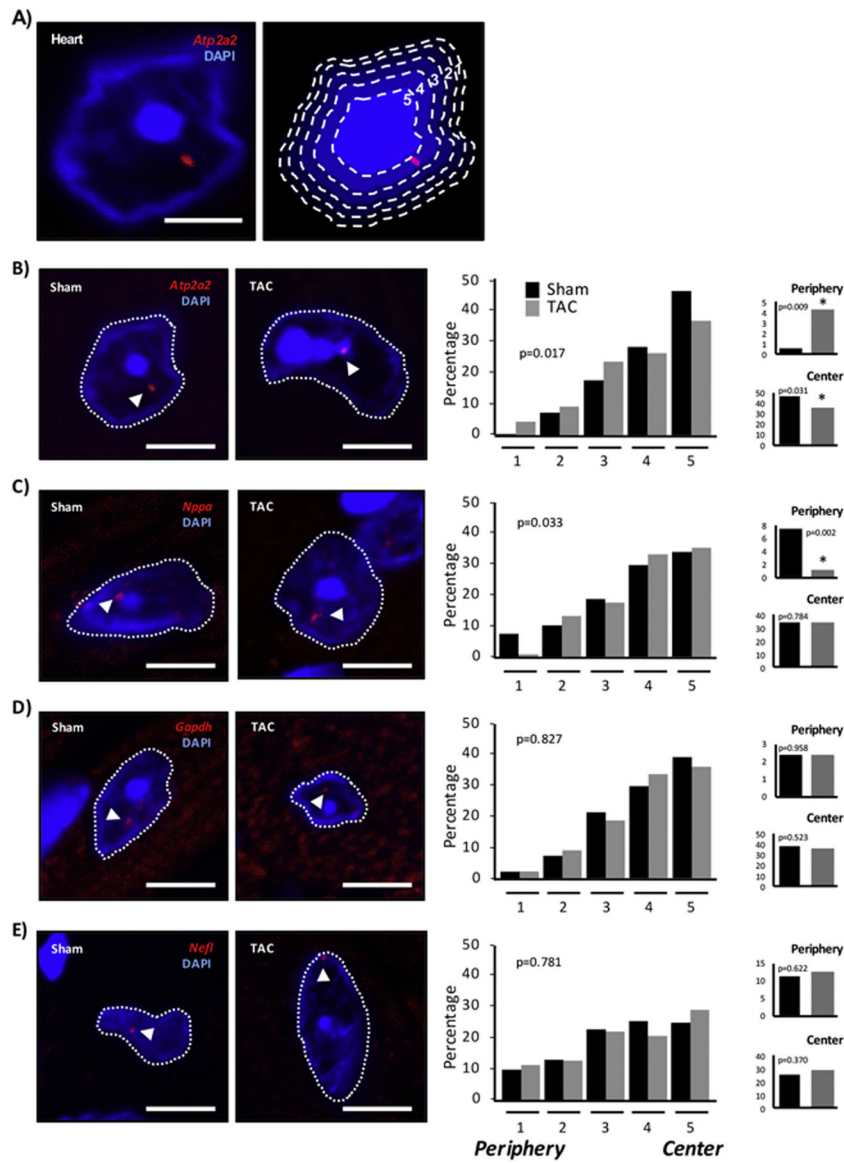


Figure 4. Cardiac gene localization is associated with differential expression in the failing heart. A) DNA FISH measurements to the nuclear periphery were normalized for shape and size between nuclei by using DAPI to bin the nucleus into 5 equal area annuli using erosion processing, allowing for an equal probability of finding a locus in any given bin [25]. Loci were assigned a bin location (1 near the periphery to 5 at the center) for analysis. Scale bar = 3 μ m. B-E) DNA FISH was performed in adult mice, either control or those that had developed heart failure. Heart tissue sections were labeled for the following genes *Atp2a2* (B), *Nppa* (C), *Gapdh* (D) and *Nefl* (E). The distribution of nuclear positioning is shown, with bins 1 (Periphery) and 5 (Center) highlighted on the right. Quantitation is compiled from different labeling experiments from 2–3 different hearts per treatment. *Atp2a2*: Sham $n=167$ nuclei/205 loci, TAC $n=220/297$; *Nppa*: Sham $n=179/213$, TAC $n=145/188$; *Gapdh*: Sham $n=178/215$, TAC $n=177/208$; *Nefl*: Sham $n=152/192$, TAC $n=177/224$. Scale bar = 5 μ m. * $p<0.05$ [Chi-squared].

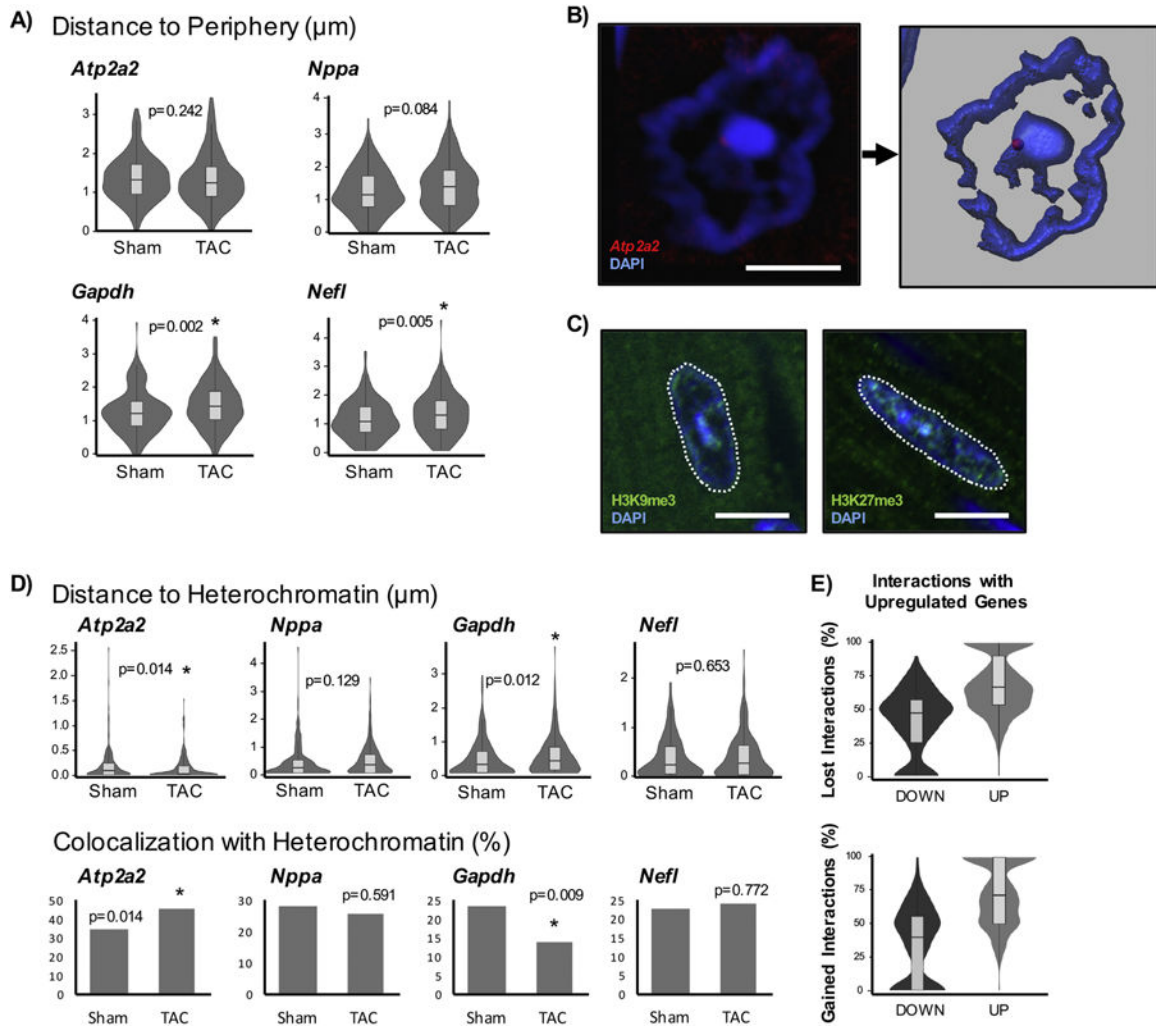


Figure 5. Localization of genes with respect to nuclear heterochromatin domains associate with patterns of differential expression.

A) In addition to performing binned assessments of radial positioning of genes, the absolute distances to the nuclear periphery were examined (note that these are absolute distances and do not account for differences in nuclear shape or area). **B)** To measure the distance to constitutive heterochromatin, a DAPI intensity threshold was used in Imaris to create a surface. The closest distance of the FISH spot to DAPI surface was measured. If loci were found inside the surface (within the heterochromatin), they were assigned a distance of 0 μm . *Scale bar = 5 μm .* **C)** The intense DAPI signal corresponds to constitutive heterochromatin, marked by H3K9me3, and is separate from facultative heterochromatin, indicated by H3K27me3. H3K9me3 colocalizes well with DAPI while H3K27me3 is found throughout the nucleus. *Scale bar = 10 μm .* **D)** The distribution of absolute distances to heterochromatin are shown on *top*. The fractions of loci that colocalize (have distances of 0 μm and associate directly with heterochromatin), are shown on *bottom*. *Atp2a2: Sham n=205 loci, TAC n=298; Nppa: Sham n=216, TAC n=190; Gapdh: Sham n=226, TAC n=212; Nefl: Sham n=192, TAC n=227.* **E)** HiC data was used to examine gene-gene interactions from isolated sham and TAC cardiomyocytes [23]. The lost and gained interactions for significant

differentially expressed genes (DOWN or UP) with other differentially expressed genes are described. Genes that are upregulated associate with other upregulated genes while those that are downregulated show the inverse relationship, providing support for genes with similar expression behavior tend to congregate together. * $p < 0.05$ [Mann-Whitney; for co-localization, Chi-squared].

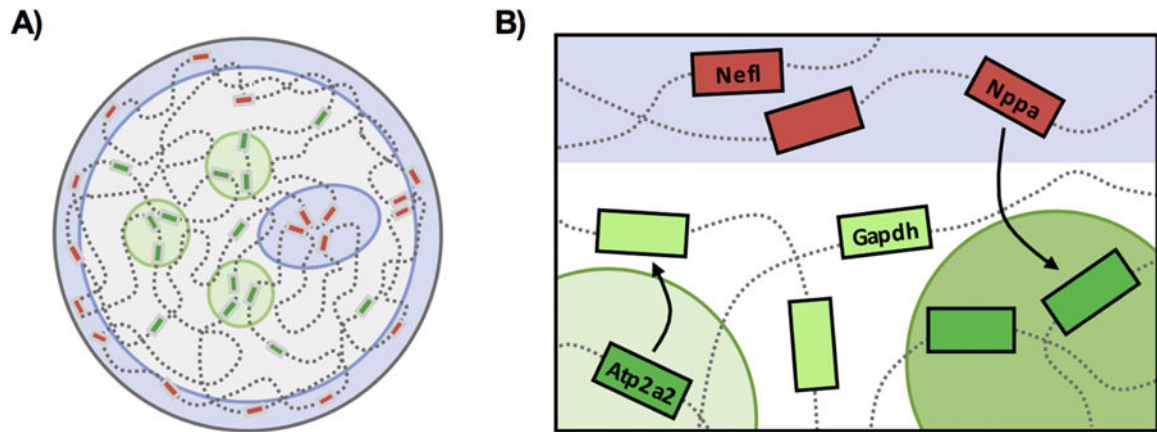


Figure 6. Gene expression is regulated by spatial positioning at the nuclear scale.

A) The nucleus is compartmentalized into active and inactive compartments. Inactive regions (*red boxes*) are found in heterochromatic regions (*blue*), which are heavily enriched at the nuclear lamina and centrally at chromocenters while highly transcribed regions (*green boxes*) are found at transcription factories (*green*). The cardiac nucleus contains stable RNA polymerase II transcription compartments enriched with active genes while genes outside of these sites in the nucleoplasm are expressed at low levels or poised for activation. Upon pathological stress, strength of transcription factory activity increases, without significant changes to number, through the recruitment of elongating RNA polymerase II molecules. **B)** The differential changes in gene expression are mediated by changes in gene positioning with respect to active (transcription factories, *green*) and inactive (heterochromatin, *blue*) compartments. Increases in gene expression correlate with the concentration of upregulated genes and RNA polymerase II molecules at discrete transcription sites.