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The nuclear matrix protein HNRNPU maintains 3D genome architecture globally in mouse hepatocytes

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Eukaryotic chromosomes are folded into higher-order conformations to coordinate genome functions. In addition to long-range chromatin loops, recent chromosome conformation capture (3C)-based studies have indicated higher levels of chromatin structures including compartments and topologically associating domains (TADs), which may serve as units of genome organization and functions. However, the molecular machinery underlying these hierarchically three-dimensional (3D) chromatin architectures remains poorly understood. Via high-throughput assays, including in situ Hi-C, DamID, ChIP-seq, and RNA-seq, we investigated roles of the Heterogeneous Nuclear Ribonucleoprotein U (HNRNPU), a nuclear matrix (NM)-associated protein, in 3D genome organization. Upon the depletion of HNRNPU in mouse hepatocytes, the coverage of lamina-associated domains (LADs) in the genome increases from 53.1% to 68.6%, and a global condensation of chromatin was observed. Furthermore, disruption of HNRNPU leads to compartment switching on 7.5% of the genome, decreases TAD boundary strengths at borders between A (active) and B (inactive) compartments, and reduces chromatin loop intensities. Long-range chromatin interactions between and within compartments or TADs are also significantly remodeled upon HNRNPU depletion. Intriguingly, HNRNPU mainly associates with active chromatin, and 80% of HNRNPU peaks coincide with the binding of CTCF or RAD21. Collectively, we demonstrated that HNRNPU functions as a major factor maintaining 3D chromatin architecture, suggesting important roles of NM-associated proteins in genome organization.

[Supplemental material is available for this article.]

In the eukaryotic cell nucleus, appropriate higher-order chromatin structures are critical to genome organization and functions (Cremer and Cremer 2010; Bickmore 2013; Dekker and Misteli 2015; Rowley and Corces 2016). It has been well established that long-range chromatin loops are important for connecting enhancers and promoters and for insulating chromatin domains (Tang et al. 2015; Ghirlando and Felsenfeld 2016; Hnisz et al. 2016). Furthermore, recent 3C-based experiments revealed that the genomes consist of multiple levels of self-associating chromatin structures including compartments (Lieberman-Aiden et al. 2009) and topologically associating domains (TADs) (Dixon et al. 2012; Nora et al. 2012; Sexton et al. 2012). There are two types of compartments: A and B, associated with active and inactive chromatin, respectively. The compartments are recently validated as relatively stable physical structures of individual cells as detected by super-resolution microscope (Wang et al. 2016b) and single cell Hi-C (Stevens et al. 2017). TADs are evolutionarily conserved and are relatively invariant among cell types (Dixon et al. 2015; Schmitt et al. 2016). It is increasingly evident that TADs may be functional units of the genome, involving biological processes such as transcriptional coregulation, DNA replication, and V(D)J recombination (Nora et al. 2012; Lucas et al. 2014; Pope et al. 2014; Symmons et al. 2014; Hu et al. 2015; Dekker and Mirny 2014). The binding of CTCF is enriched at the boundary of TADs in a convergent manner (Dixon et al. 2012; Ruan et al. 2014; Vietri Rudan et al. 2015). Deletion or inversion of CTCF binding sites disrupts the topology of chromatin loops (Guo et al. 2015; Narendra et al. 2015), and an extrusion model was proposed to explain the mechanism for loop formation mediated by CTCF and cohesin (Sanborn et al. 2015; Fudenberg et al. 2016). Furthermore, loss-of-function studies indicated that the depletion of CTCF or cohesin led to alterations of intra- and inter-TAD chromatin interactions (Seitan et al. 2013; Sotueva et al. 2013; Zuin et al. 2014). Other proteins such as ZNF143, SMARCA4, and RUNX1 are also found...
to play roles in the regulation of chromatin interactions (Heidari et al. 2014; Bailey et al. 2015; Barutcu et al. 2016a,b). However, the molecular machinery responsible for global and hierarchical chromatin organization remains poorly understood.

The nuclear matrix (NM) consists of the inner NM and peripheral NM, and the latter includes nuclear lamina and nuclear pore complexes (Linnemann and Kravetz 2009). It has long been speculated that the NM provides structural frameworks for topologically confined chromatin domains (Getzenberg et al. 1991), but the roles of the inner NM in global chromatin architecture such as compartments and TADs have not been investigated systematically. As one of the major NM-associated proteins, HNRNPU (also known as SAF-A) was recognized as an attachment factor linking specific DNA elements, scaffold/matrix attachment regions (S/MAR), to the NM (Gohring and Fackelmayer 1997). Purified HNRNPU proteins can bind DNA to form higher-ordered nucleic-acid-protein complexes in vitro (Fackelmayer et al. 1994). Moreover, HNRNPU is required for the localization of Xist RNA (Hasegawa et al. 2010), the key regulator of X Chromosome inactivation. Therefore, we hypothesized that HNRNPU could play a general role in regulating global 3D genome organization.

Results

HNRNPU is required for global nuclear organization

To test roles of HNRNPU in nuclear organization, we knocked down Hnnpu gene expression by RNA interference (RNAi) with two independent short hairpin RNAs (shRNAs) in AML12 cells (Fig. 1A), a mouse hepatocyte cell line in which the nuclear architecture is well preserved (McDonald et al. 2011; Fu et al. 2015). Based on immunofluorescence analysis, we observed strong signal reductions of the nucleolus marker C23 and a significant increase of heterochromatin foci numbers (marked by H3K9me3) in HNRNPU-depleted cells (Fig. 1B,C), we only used shU-1 for further high-throughput assays (labeled as “shU”). For genome-wide experiments, we conducted two replicates using AML12 cells treated with shU-1 containing lentivirus separately.

Disruption of HNRNPU alters gene expression and increases chromatin-lamina interactions

To reveal biological consequences of HNRNPU depletion, we examined gene expression by RNA-seq. We observed significant expression changes on 639 genes upon HNRNPU depletion. Among them, 500 (78%) genes were down-regulated and 139 genes (22%) were up-regulated (Supplemental Table S1; Supplemental Fig. S2A,B). Gene Ontology (GO) analysis shows that up-regulated genes are enriched in biological processes including cell adhesion and cell proliferation, and down-regulated genes associate with immune processes, oxidation-reduction processes, and triglyceride metabolic processes (Supplemental Fig. S2C).

To investigate spatial organization of the genome with HNRNPU depletion, we mapped chromatin-lamina interactions by DNA Adenine Methyltransferase Identification (DamID) assay with lamin B1-Dam fusion proteins. The data show that chromatin-lamina interactions are substantially remodeled upon HNRNPU depletion (Fig. 2A,B), and the genome coverage of lamin-associated chromatin domains (LADs) (Guelen et al. 2008) is increased from 53.1% to 68.6% (Fig. 2C; Supplemental Table S2). Importantly, the gain of LADs is significantly associated with the reduction of gene expression (Fig. 2D). Consistently, the electron microscope (EM) imaging displays a global relocalization of chromatin toward the nuclear membrane in HNRNPU-depleted cells (Fig. 2E). Thus, HNRNPU depletion results in spatial changes of chromatin toward the nuclear periphery, which associates with gene expression down-regulation.

HNRNPU regulates chromatin compartments

To investigate the impact of HNRNPU on chromatin interactions, we generated high-resolution interaction maps by an in situ Hi-C assay (Rao et al. 2014). We conducted two replicates of Hi-C experiments, and the results are highly consistent (Supplemental Fig. S3). Hi-C data provide information about multiscale chromatin interactions including compartments, TADs, and chromatin loops, and we first examined compartments. While contact maps of control and HNRNPU-depleted cells are similar at the chromosome level (Fig. 3A), interactions among compartments are changed globally (Fig. 3B). We applied different statistical methods to detect these differences and obtained similar results (Supplemental Fig. S4).

To take a closer look at compartment organization, we examined inter- and intra-compartment interactions. Consistent with the results of Figure 3B, inter-compartment interactions between A and B increase significantly, but those among the same type of compartments (A vs. A; B vs. B) decrease significantly (Fig. 3C). Further, we compared changes in inter-compartment interactions along distances. Interactions among the same type of compartments become significantly decreased if distances are >2 Mb. However, interactions between A and B compartments decrease significantly in all distance ranges (Supplemental Fig. S5). Moreover, intra-compartment interactions significantly increase in A compartments but decrease in B compartments (Fig. 3D). Of note, results obtained from replicates (rep1 and rep2) are almost identical, exhibiting high repeatability of our Hi-C experiments and analysis. These data suggested that the depletion of HNRNPU remodels long-range interactions at the compartment level.

We next examined compartment switching with Hnnpu knockdown. The numbers of switched bins (200 kb each) in two independent knockdown experiments are 1094 (8.5%) and 1099 (8.6%), but those between two controls and two knockdown samples are only 113 (0.9%) and 182 (1.4%), respectively (Supplemental Fig. S6A). The percentages of compartment switching upon HNRNPU depletion are significantly higher than those between replicates in the same condition (generalized linear model with a
binomial outcome, \( P < 2.2 \times 10^{-16} \). Additionally, switched compartments are highly consistent between the two replicates: A to B, 88% overlapping; B to A, 84% overlapping (Supplemental Fig. S6B). By considering consistent results from replicates only, percentages of the switched region are 4.9% for A to B, and 2.6% for B to A (Fig. 3E; Supplemental Table S3). There are 890 genes in compartments switched from A to B, and 747 genes in those shifted from B to A. The gene density of these switched compartments is between those of unchanged A and B compartments (Supplemental Fig. S6C).
Furthermore, the A-to-B compartment switching coincides with increases of chromatin-lamina interaction and decreases of gene expression, but B-to-A compartment switching displays opposite trends (Fig. 3F,G). The down-regulated genes paralleled with A-to-B switching are significantly enriched in GO terms related to liver functions, such as the P450 pathway and lipid transport (Supplemental Fig. S6D,E), suggesting a loss of cell identity resulted from changes of chromatin compartments.

HNRNPU regulates TAD boundaries and interactions

We next examined chromatin interactions at the TAD level. We calculated TAD boundaries using two replicates separately. The overlap between replicates in the same condition is higher than 95% (Supplemental Fig. S7A,B). The down-regulated genes paralleled with A-to-B switching are significantly enriched in GO terms related to liver functions, such as the P450 pathway and lipid transport (Supplemental Fig. S6D,E), suggesting a loss of cell identity resulted from changes of chromatin compartments.

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HNRNPU regulates chromatin loops

To reveal roles of HNRNPU in 3D chromatin organization at finer scales, we investigated changes of chromatin loops. We first merged all contacts from control and knockdown samples for a
Figure 3. HNRNPU regulates chromatin compartment organization. (A) Hi-C contact maps of Chromosome 19 in (left) shCtrl- and (right) shU-treated cells at 200-kb resolution. Compartments A (red) and B (blue) are shown by PC1 eigenvectors. (B) (Left) Log2 (shU/shCtrl) of Chromosome 19 binned at 200-kb resolution. (Right) Log2 (shU/shCtrl) of 40-kb bins (lower left) and averaged by compartments (upper right) in a 15.9-Mb region. (C,D) Statistics of interaction changes between (C) and within (D) compartments. (Top) Schematic view of regions examined. (Middle) Cumulative distribution of obs/exp values of interactions between compartments. P-values were obtained by two-factor ANOVA. (Bottom) Box plots showing the pairwise fold changes of interactions. Results from two replicates are presented. (E) Percentages of compartment switching between shCtrl- and shU-treated cells. (F) The relationships between compartment switching and the lamina association (based on lamin B1 DamID data). (G) Box plots showing expression changes with compartment switching. P-values were obtained by Wilcoxon rank-sum test.
more accurate loop detection and identified, in total, 7994 chromatin loops (Supplemental Table S5). The overall strengths of chromatin loops are reduced in the HNRNPU-depleted cells as shown by the aggregate signal from all loops (Fig. 5A). To test the statistical significance of loop changes upon HNRNPU depletion, we compared the intensity of loops (peaks) and “fold changes” of peaks relative to four neighboring regions (Fig 5B), as proposed previously (Rao et al. 2014). In the HNRNPU-depleted cells, the intensity of peaks (Fig. 5C) and their relative ratios to adjacent regions (Fig. 5D) are all significantly reduced. Then, we defined differential loops only if all the five values (peak intensity and ratios relative to four neighboring regions) change in the same direction. In total, we identified 4651 (58.2%) decreased and 680 (8.5%) increased loops upon HNRNPU knockdown (Fig. 5E). Two representative regions showing decreased chromatin loops are provided in Figure 5F.

HNRNPU associates with CTCF and RAD21

Next, we explored how HNRNPU affects chromatin organization. High-throughput mass spectrometry data suggested that HNRNPU might interact with CTCF (Hutchins et al. 2010). Our immunofluorescence data indicated that both CTCF and RAD21, a component of the cohesin complex, are partially colocalized with HNRNPU in the nucleus (Fig. 6A). Co-immunoprecipitation experiments further validated that HNRNPU, CTCF, and RAD21 could be associated (Fig. 6B).

Then, we mapped genome-wide chromatin binding of CTCF, RAD21, and HNRNPU in AML12 cells (Fig. 6C; Supplemental Tables S6, S7). For CTCF and RAD21, we used regular ChIP-seq; for HNRNPU, we applied bioChIP-seq, which is based on in vivo biotinylation of interested proteins by the BirA enzyme (He and Pu 2010). Our HNRNPU bioChIP-seq experiments were carefully controlled by examining expression and localization of tagged HNRNPU proteins (Supplemental Fig. S8). By comparing with the BirA-only control, we identified 5285 HNRNPU peaks. Strikingly, more than 80% of HNRNPU peaks overlap with CTCF or RAD21, and 59.6% of HNRNPU peaks overlap with both (Fig. 6D), further supporting the interaction among these three proteins. However, we still cannot determine whether the interaction is direct or indirect.

Furthermore, HNRNPU peaks are enriched at TAD boundaries (Supplemental Fig. S9A), and intensities of HNRNPU-associated boundaries are significantly higher than those not associated
with HNRNPU (Supplemental Fig. S9B). Interestingly, upon HNRNPU depletion, only boundaries bound by CTCF and RAD21 are significantly decreased in their strengths (Supplemental Fig. S9C).

To test whether HNRNPU affects the binding of CTCF and RAD21 on chromatin, we compared ChIP-seq data from control and knockdown cells (Supplemental Fig. S10A,B). While CTCF binding is almost unchanged, 3816 (9.8%) RAD21 peaks are significantly reduced in their binding strengths (Supplemental Fig. S10C,D). These decreased RAD21 peaks are highly associated with CTCF binding (Supplemental Fig. S10E). Concordantly, chromatin loops associated with decreased RAD21 binding are significantly weaker than those associated with invariant RAD21 peaks (Supplemental Fig. S10F,G), but proteins levels of CTCF and RAD21 remain unchanged upon HNRNPU depletion (Supplemental Fig. S10H).

HNRNPU associates with active chromatin

Finally, we examined the distribution and strength of HNRNPU binding relative to genes. Surprisingly, ~60% of HNRNPU peaks locate at promoters (Fig. 6E). Further, binding strengths of HNRNPU near transcription start sites (TSSs) are positively correlated with levels of gene expression (Fig. 6F). Consistently, 90% of HNRNPU peaks are within compartment A or non-LADs (Fig. 6G). Therefore, our data indicated that HNRNPU mainly associates with active chromatin.

Discussion

In conclusion, using mouse hepatocytes as the model, we demonstrate that HNRNPU functions as a major regulator of the 3D genome, regulating multiple levels of chromatin architectures, including LADs, compartments, TADs, and chromatin loops, partially through cooperating with CTCF and RAD21.

Although the concept of the NM has been disputed for decades (Pederson 2000; Nickerson 2001), the NM is still being hypothesized as a platform for 3D genome organization (Razin et al. 2014). Besides the HNRNPU data presented here, topoisomerase II beta, another NM-associated protein, also associates with cohesin and CTCF at TAD boundaries (Uuskula-Reimand et al. 2015).
2016), suggesting general roles of the NM and/or its associated proteins in 3D genome organization. As a potential mediator connecting chromatin and NM, HNRNPU mainly associates with A compartments, as indicated in Figure 6G. HNRNPU may maintain proper chromatin architecture by “stabilizing” chromatin to the inner NM. The depletion of HNRNPU may decrease the constraint on chromatin, therefore remodeling chromatin-chromatin and chromatin-lamina interactions (Supplemental Fig. S11).

During the review process of our manuscript, Nick Gilbert and colleagues reported that, primarily based on FISH, biochemistry, and mutagenesis, HNRNPU/SAF-A is involved in de-compaction of chromatin via oligomerization with chromatin-associated RNAs (caRNAs), mainly in gene-rich regions (Nozawa et al. 2017). In the current study, we showed that HNRNPU depletion leads to global chromatin condensation, an increase of LADs from 53.1% to 68.6% of the genome, compartment switching on 7.5% of the genome, and a significant decrease of TAD boundary strengths and chromatin loop intensities. Therefore, these two studies are highly complementary, demonstrating important roles of HNRNPU in the higher-order chromatin organization.

Furthermore, the Gilbert study demonstrated that the regulatory role of HNRNPU on chromatin depends on transcription and caRNAs (Nozawa et al. 2017). Our study indicated that HNRNPU strongly associates with active genes and with CTCF- and RAD21-bound regions. These observations suggested that caRNAs could be involved in the organization of higher-order

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Figure 6. HNRNPU associates with CTCF and RAD21. (A) Immunofluorescent staining displays partial colocalization of HNRNPU (red) and CTCF or RAD21 (green) in AML12 cells. Scale bar, 10 µm. (B) Endogenous co-immunoprecipitation with HNRNPU (left) or CTCF (right) antibodies, and blots with indicated antibodies. (C) ChIP-seq tracks of CTCF and RAD21; bioChIP-seq tracks of HNRNPU and BirA only (negative control) on a 1.3-Mb region on Chromosome 19. (D) Venn diagram showing overlaps between peaks of CTCF, RAD21, and HNRNPU. (E) Distribution of HNRNPU peaks relative to genes. (F) Average profiles of HNRNPU bioChIP-seq signal of genes at different expression levels (100% denotes the highest expression). (G) Distribution of HNRNPU peaks relative to compartments and LADs.
chromatin via cooperating with RNA binding proteins (such as HNRNPU), structural proteins (such as CTCF and cohesin), and the transcription machinery, presumably.

Additionally, as a protein binding most kinds of RNAs (Xiao et al. 2012), HNRNPU also interacts with long noncoding RNAs (lncRNAs) such as Xist (Chu et al. 2015; McHugh et al. 2015) and Fire (Hacisuleyman et al. 2014), both of which are known regulators of chromatin interactions at specific regions. Other lncRNAs, including LincGET and Blen1, are also found to be interacting with HNRNPU (Wang et al. 2016a; Mi et al. 2017). These studies suggest potential lncRNA-dependent roles of HNRNPU in genome organization of specific loci, which awaits further investigation.

Our data indicated that HNRNPU affects the strengths of more than 40% of TAD boundaries. Additionally, TAD boundaries are enriched for the binding of HNRNPU and other known 3D genome regulators including CTCF, RAD21, and SMARCA4 (Dixon et al. 2012; Sofueva et al. 2013; Barutcu et al. 2016b). Our study further indicated that HNRNPU proteins interact with CTCF and RAD21. Thus, these “boundary factors” may form macrocomplexes, directly or indirectly, to segregate TADs. Further studies are needed to identify components and functional roles of these potential macrocomplexes and to reveal their dynamics and dysfunction in development and diseases.

The HNRNPU-deficient mice displayed phenotypes of post-implantation lethality (Roshon and Ruley 2005) and impaired heart development as demonstrated by conditional knockout (Ye et al. 2015). Human de novo mutations or microdeletions of the Hnrnpu gene are associated with brain disorders (Epi4K Consortium et al. 2013; Gupta et al. 2014). The experimental observations in both humans and mice suggest that HNRNPU could regulate genome organization in vivo in development and diseases.

Methods

RNAi and Rescue

For RNAi experiments, shRNA plasmids were constructed by cloning the target sequences to the pLKO.1-puro vector: AGATCATGGCCAGGGATTTT (mouse-shU-1), TGCCCTGAAGAA GCAGAAATTT (mouse-shU-2), AGATCATGGCCGTTGATATTT (human-shU-1), GACCTGTACCTGCTCATTAA (human-shU-2), and ACTCGACATAGTATCTCA (negative control). For rescue experiments, mouse Hnrnpu cDNA was mutated for the targets of shU-1. AML12 cells were treated with shRNAs or expression vectors by lentivirus-based systems for 5 d.

Western Blotting

The cell lysates were blotted against primary antibodies as follows: anti-H3 (Abcam, ab1791), anti-H3K9me3 (Abclonal, A2360), anti-C23 (Proteintech, 10556-1-AP), anti-HNRNPU (Abcam, ab20666), and anti-RAD21 (Abcam, ab9922). The blots were visualized with peroxidase-coupled secondary antibodies.

Immunofluorescence

Immunofluorescence was conducted with the same antibodies used for Western blots except for HNRNPU (Abcam, ab10297). The slides were imaged with a confocal laser scanning microscopy (Leica, SP5).

Electron Microscopy Imaging

Cells were fixed with 2.5% glutaraldehyde and further processed and imaged by the EM facility at the Shanghai Institute of Biochemistry and Cell Biology (SIBCB).

DamID

DamID experiments were performed as described (Guelen et al. 2008). Two biological replicates were conducted for the control and Hnrnpu knockdown AML-12 cells. Microarray data were analyzed as previously described (Fu et al. 2015).

In situ Hi-C

The in situ Hi-C libraries were prepared as previously described (Rao et al. 2014). Two biological replicates were performed for both control and HNRNPU-depleted AML12 cells. The libraries were then sequenced via the Illumina HiSeq X Ten system. Detailed information about Hi-C data analysis is provided in the Supplemental Methods.

ChIP-seq and bioChIP-seq

ChIP experiments were performed as previously described (Wen et al. 2008) with antibodies against CTCF or RAD21. For HNRNPU, we applied bioChIP experiments as previously described (He and Pu 2010; Roux et al. 2012) with some modifications. Detailed information of bioChIP experiments and ChIP-seq data analysis are provided in the Supplemental Methods.

Data Access

The raw and processed DamID, RNA-seq, ChIP-seq, and Hi-C data sets from this study have been submitted to the NCBI Gene Expression Omnibus (GEO; http://www.ncbi.nlm.nih.gov/geo/) under accession number GSE95116.

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Author Contributions: H.F. and B.W. conceived and designed this study. H.F. performed most of the experiments except for DamID assay (by X.H.), Hi-C library preparation (Q.W.), Co-IP (J.W.), and DamID microarray experiments (L.Z. and F.Z.). P.L. analyzed the data, with the guidance of H.W., Y.L., and X.Z. L.C. and Q.-Q.T. provided some experimental support. B.W., H.F., and P.L. prepared the manuscript with input from all authors.

References


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