Visualization of chromatin higher-order structures and dynamics in live cells

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Chromatin has highly organized structures in the nucleus, and these higher-order structures are proposed to regulate gene activities and cellular processes. Sequencing-based techniques, such as Hi-C, and fluorescent in situ hybridization (FISH) have revealed a spatial segregation of active and inactive compartments of chromatin, as well as the non-random positioning of chromosomes in the nucleus, respectively. However, regardless of their efficiency in capturing target genomic sites, these techniques are limited to fixed cells. Since chromatin has dynamic structures, live cell imaging techniques are highlighted for their ability to detect conformational changes in chromatin at a specific time point, or to track various arrangements of chromatin through long-term imaging. Given that the imaging approaches to study live cells are dramatically advanced, we recapitulate methods that are widely used to visualize the dynamics of higher-order chromatin structures. [BMB Reports 2021; 54(10): 489-496]

INTRODUCTION

Chromatin is hierarchically arranged in the cell nucleus (1). Accumulating evidence from current studies suggests that the organization of the three-dimensional (3D) genome is critical for regulating gene expressions and maintaining various cellular functions (2-8). To investigate large-scale chromatin organization, chromatin conformation capture (3C) methods have been widely used (9-11). Such sequencing-based techniques can efficiently identify contacts between chromatin domains and genomic loci by proximity ligation of cross-linked DNA fragments from fixed cells. Hi-C experiments recently revealed higher order chromatin organization that defines two types of segregated chromatin interactions—compartment A, and compartment B (Fig. 1) (10, 12). These two compartments have distinct characteristics. Compartment A is known to have open and accessible chromatin with higher transcription activity, while compartment B is associated with compactly arranged, closed chromatin with lower gene density. In addition to genomic compartmentalization, the topologically associating domain (TAD) was identified through Hi-C experiments, whose boundaries are associated with insulator binding protein CCCTC-binding factor (CTCF) and cohesion complex to form self-interacting chromatin regions at megabase scale (Fig. 1) (13).

Although sequencing-based techniques can identify long-range chromatin interactions and detect the spatial organization of chromosomes in the nucleus, these methods require millions of cells to calculate the averaged pattern of chromosome conformations. Since ensemble averaged data from the techniques underestimated cell-to-cell variability, the single-cell Hi-C method has been developed (14-18). However, regardless of the groundbreaking development of Hi-C analysis, what still remains as challenging is that sequencing-based approaches cannot detect dynamic chromatin movements, as their protocols rely on fixed cells. Chromatins are known to have dynamic properties, of which their structural changes or positioning can affect numerous nuclear activities, including transcription, replication, and DNA repair. Thus, live-cell imaging has become essential for understanding spatiotemporal chromatin movements and related functions as clustered regularly interspaced short palindromic repeats (CRISPR)-based imaging systems and their applications combined with fluorescence super-resolution microscopy have been widely used to elucidate unknown nuclear mechanisms.

In this review, we discuss the development of microscopy techniques and strategies for visualizing the real-time dynamics of genomic organization. The major nuclear structures that we focus on are chromosome territories (CTs), lamina-associated domains (LADs), and nucleolar-associated domains (NADs), of which their structures are closely related to gene-expression regulation and cell-type specificity.
Recent imaging methods for chromatin structures  
Tae Lim Park, et al.

**MAIN TEXT**

Visualization of chromosome territories
Genomic organization in the nucleus has been considered crucial for gene regulation and other nuclear processes. Followed by the advances in microscopic approaches, one key feature of nuclear architecture was revealed that chromosomes are not randomly positioned in the interphase nucleus, but instead occupy distinct nuclear spaces (1, 19-22). Such spatial segregation of individual chromosome is termed CT (Fig. 1). The most commonly used technique to study CTs is FISH (23, 24).

To perform FISH, probes that consist of oligonucleotides to label a genomic region of interest are hybridized to the target chromosome, and enabled to visualize each CT (25). To further investigate the spatial arrangement of chromosomes, 3D FISH has been developed to generate 3D image reconstruction that is accomplished by confocal microscopy that could obtain serially sectioned nuclei, and stack all the images, to ultimately provide the 3D locations of chromosomes (26, 27). With 3D FISH technique, a critical pattern of genomic arrangement was discovered. Human (HSA) chromosome 19 with high gene density was found in the nuclear interior, while HSA 18 with low gene density was located at the nuclear periphery (28). A similar test was performed on HSA 12; gene-dense and gene-poor sections on the same chromosome were separately labeled with two different probes, and the resulting 3D reconstruction image correlated with the previous finding. Regions with more genes tended to be positioned in the interior, while those with lower genes were found in the peripheral sites of the nucleus. These observations confirmed that gene density is closely related to the radial positions of chromosomes. Further studies also highlighted that among primates, the non-random positions of CTs are evolutionarily conserved, while they are significantly different in various cell types.

However, FISH-based techniques are limited to fixed cells, and involve DNA denaturation, which cannot provide information on the spatiotemporal changes of natural chromatin structures. To overcome the existing limitations, a number of approaches have been developed to achieve live-cell imaging, which allows the real-time dynamics of chromatin organization to be observed. The most well-known method applied for visualizing living cells is the clustered regularly CRISPR-based imaging system.

CRISPR/dCas9 systems for genome labeling and dCas9 orthologs for multi-color imaging in live cells
The CRISPR technique was initially introduced for genome engineering, in combination with single-guide RNA (sgRNA), which contains specific sequences targeting a genomic region of interest, and CRISPR-associated protein 9 (Cas9), which has
nuclease activity to cleave the targeted site, and induce DNA repair for genome editing (29-31). The mechanism that can recruit CRISPR/Cas9 complex to the specific genomic site via sgRNAs has made this technique suitable for genome imaging. To label the genome in live cells, Cas9 protein is modified to dead Cas9 (dCas9), which has deactivated nuclease function, and is fused with a fluorescent protein (32). Fluorescently labeled dCas9 complexed with CRISPR is recruited to the genomic region where sequences complementary to the specific sgRNA reside (Fig. 2B). The process required to select and generate appropriate sgRNAs is much simpler and easier than that for other methods, such as the transcription-activator-like effector nuclease (TALEN)-based technique, which involves specific protein productions for each targeting region. Thus, the CRISPR/Cas9 system is extensively utilized for tracking the dynamics of genomic structures.

Genome labeling with the CRISPR/Cas9 technique can also be used to visualize multiple genomic regions (33). There are several orthogonal Cas9 variants that have different protospacer adjacent motif (PAM) sequences. PAM is a short DNA sequence downstream of genomic site targeted by sgRNA, and such diversity and specificity in PAM enable multi-locus labeling with Cas9 orthologs. The most commonly used dCas9 is from Streptococcus pyogenes (SpyCas9) with the PAM sequence 5'-NGG-3'. Other known orthogonal Cas9 variants are from Staphylococcus aureus (SadCas), Neisseria meningitidis (NmdCas9), and Streptococcus thermophilus (St1dCas9), which recognize 5'-NNRRT-3', 5'-NNNGATT-3', and 5'-NNAGAAW-3', respectively. Recent studies tested whether these orthologs could be used in a combination to label interchromosomal and intra-chromosomal loci in live cells, and their results confirmed that each ortholog distinctively labeled the targeted genomic region with high specificity achieved by the unique PAM sequences.
having high SNR with high specificity.

In addition to CARGO-dCas9 system, an alternative strategy to improve SNR has recently been proposed, and is known as CRISPR LiveFISH (37). Unlike the conventional CRISPR/Cas9-based approach that uses fluorescent protein fused dCas9 for signal detection, gRNA tagged with a synthetic organic dye, cyanine 3 (Cy3), is used to track targeted regions. dCas9-EGFP and Cy3-gRNA targeting the same repetitive site in chromosome 3 were tested to compare their signal intensity and SNR, respectively. The results proved that both relative fluorescence and SNR were found to be much higher in the region labeled with Cy3-gRNA. Moreover, long-lasting expression of Cy3-gRNA was detected by forming more stable dCas9 complex. gRNAs known to be easily degraded in the cellular environment. However, Cy3-gRNA presents higher stability when compared to the conventional gRNA used with fluorescently labeled dCas9, enabling long-term live cell imaging. Another advantage of CRISPR LiveFISH is that this method overcomes a nucleolar accumulation of dCas9, a common phenomenon in which dCas9 proteins not bound to gRNAs are frequently enriched in the nucleoli. Since CRISPR LiveFISH only detects target-bound dCas9-Cy3-gRNA complex, it is applicable for spotting specific genomic sites without signals in the nucleoli.

**Mapping LADs and NADs with sequencing approaches**

Accumulated evidence from recent studies implies that the major structural components in the nucleus, nuclear membrane, and nucleolus could be considered as genome organizers, which affect gene expression regulatory processes (38, 39). The nuclear membrane is structured with double lipid bilayers composed of an outer nuclear membrane (ONM), and an inner nuclear membrane (INM). Below the INM, nuclear lamina is organized through a fibrillar meshwork of nuclear lamins, A-type and B-type lamins, which serve as a scaffold for the binding of various proteins, and chromatin domains as well. A-type lamins, which mainly include lamin A and C, are encoded by *LMNA*, while B-type lamins are classified as lamin B1 and B2, coded by *LMNB1* and *LMNB2*, respectively (40, 41). Another genome organizer, the nucleolus is the largest subcellular compartment in the nucleus, and is known as a site for ribosomal RNA (rRNA) synthesis and ribosome biogenesis. The nucleolus does not have a membrane, even though it is the largest sub-organelle inside the nucleus, while it consists of three layers forming a concentric arrangement: fibrillar center (FC), dense fibrillar component (DFC), and granular component (GC) (38). Numerous sequencing-based experimental approaches and studies with fluorescence imaging have revealed that the nuclear lamina and nucleolus represent the hubs for the chromatin organization, especially through interacting with heterochromatin domains.

Heterochromatin is spatially segregated into two main regions within the nucleus: nuclear periphery, and nucleolar periphery. The genomic regions that make molecular contacts with the nuclear lamina are termed lamina-associated domains (LADs) (42-45), while the other genomic sites that frequently interact with the nucleolus are termed nucleolus-associated domains (NADs) (46, 47).

LADs are found on all chromosomes, covering around 40% of the entire genome. They typically range in length from (100 kb to 1 Mb), with a median size of 0.5 Mb. The human LADs are characterized as consisting of genes in transcriptionally silent state, or with low expression levels, and overlap with relatively gene-poor chromatin domains. LADs are especially enriched in B compartments with repressive histone modifications, such as di- and tri-methylated histone H3 lysine 9 (H3K9me2 and H3K9me3) (43, 48). A recent study with mouse embryonic stem cells has classified three types of chromatin regions in terms of LADs: constitutive LADs (cLADs), facultative LADs (fLADs), and constitutive interLADs (cILADs). In detail, cLADs are defined as regions that remain in position during differentiation, while fLADs represent regions that interact with the nuclear lamina in a cell-type specific manner. cILADs are sites that are outside of LADs in any cell type (45, 49). Interestingly, cILADs with the lowest gene density are highly conserved in their genomic positions, suggesting that cILADs play a crucial role in forming a “structural backbone” of chromatin organization in the nucleus (50, 51).

Similarly, NADs have low gene density, have a repressive chromatin signature, and contain a high density of AT-rich sequence elements. NADs have a comparable size range of (0.1-10) Mb and median sequence length (749 kb) to those of LADs (46, 47). There are two distinct classes of NADs: Type I NADs, and Type II NADs. Type I NADs commonly associate with both the nucleolar periphery and nuclear lamina, and exhibit features of constitutive heterochromatin. In contrast to Type I NADs, Type II NADs interact with nucleoli, but not with nuclear lamina (52). However, compared to LADs, NADs have been less thoroughly studied for their biology. Altogether, LADs and NADs share the general characteristics of heterochromatin.

How could these heterochromatin domains be identified? Genome-wide mapping studies of LADs and NADs were performed on multiple eukaryotic species, including *Caenorhabditis elegans*, *Drosophila melanogaster*, mouse, and human cells via several distinct methods, such as DNA adenine methyltransferase identification (DamID), chromatin immunoprecipitation sequencing (ChIP-seq), and NAD-seq (42-45, 48, 53).

LADs were firstly identified by DamID, a proximity DNA labeling method. DamID works by fusing a bacterial adenine methyltransferase (Dam) to a protein of interest. To detect LADs, Dam is fused with lamin B1. Dam methylase recognizes GATC motifs in lamin B1-binding regions and methylates GATC sites, resulting in 6-methyl-adenine (mA). Because mA modification is not normally detected in eukaryotes, DamID achieves specific labeling of lamin-interacting regions. Those regions containing mA are isolated by cutting with the methyl-specific restriction enzyme, DpnI. These methylated fragments are selectively amplified and identified by DNA microarrays or high-
throughput sequencing (DamID-seq) (43). DamID has emerged as a comprehensive technique to profile genome-wide protein-DNA interactions, and construct a high-resolution map for protein interaction sites in the genome. Beside the DamID method, LADs have also been mapped with ChIP-seq of A- or B-type lamins using lamin-specific antibodies (54-56).

NADs have been analyzed genome-wide through a combination of fluorescence comparative genome hybridization (CGH) and high-throughput DNA sequencing with biochemically isolated nucleoli from nuclei, referred to as NAD-seq (46, 47).

In addition to sedimentation, which is one of the nucleolar isolation methods, it has been reported that nucleoli from Arabidopsis thaliana leaves can be purified by introducing chimeric m6A fragments that include the binding domain, and has no contact with the nuclear lamina, even if the interaction occurs simultaneously, or across long-range distances, within the nucleus (58). However, since H3K9me3-marked heterochromatin is resistant to sonication and the experimental processes to isolate nucleoli vary, sequencing based on purified nucleoli through sonication can be responsible for obtaining biased maps of NADs (59). These limitations led to the establishment of another novel method, called ‘Nucleolar-DamID’. By applying the existing DamID, Dam is fused with the histone, H2B, where in this adaptation, a nucleolar localization signal (NoLS, RKK) is inserted at the C-terminus (60). Through these sequencing approaches, the genome-wide maps and heterochromatic characteristics of LADs and NADs have been defined.

Visualization of LADs and NADs in live cells

Although sequencing approaches have improved our understanding of LADs and NADs, those methods still have several limitations, including the lack of coordination of spatial and temporal information. Real-time visualization in living cells has been considered a powerful method, as it enables observation of the spatial arrangement and dynamics of heterochromatin domains in the three-dimensional space of the nucleus.

The dynamics of LADs has been tracked in live single-cells using the DamID-derived imaging method known as ^\textsuperscript{\texttt{A}}-Tracer" (Fig. 3A). Dpn1 contains a domain that recognizes and selectively binds to m6A. Genetically encoded ^\textsuperscript{\texttt{A}}-Tracer is fused with enhanced green fluorescent protein (eGFP) in a truncated dpn1 fragment that includes the binding domain, and has no cleavage activity. Since m6A is a stable covalent modification, m6A-tagged LADs can be continuously tracked once DNA makes contact with the nuclear lamina, even if the interaction no longer occurs at the moment of imaging. Therefore, this "molecular contact memory" strategy can provide a microscopic "history tracking" of nuclear lamina-interacting LADs (61). Many studies that utilized ^\textsuperscript{\texttt{A}}-tracer or its modified system have demonstrated that during interphase, LADs have constrained mobility, while after mitosis, have appeared to be intrinsically stochastic (Fig. 3B). These observations suggest that LAD localization is generally not inherited. Interestingly, after mitosis, some of the nucleoplasmic LADs become closely associated with nucleoli (61, 62). These results are consistent with the genome-wide mapping studies in which NADs sequences partially overlap with LADs (46, 47).

Unlike the nuclear lamina, the nucleolus is a subnuclear membrane-less organelle (63), which makes it challenging to apply the DamID-derived tool to track NADs in live cells. This implicates that there seems to be no possible candidate molecules as scaffold to bind in and around the nucleoli, while there are the stable lamins under the nuclear envelope for mapping LADs. However, Nucleolar-DamID using Dam-fused and Nol5 inserted histone, which can bind DNA sequences near nucleoli without motif specificity, has recently been proposed (60), thereby applying it may allow insight to be gained into the dynamics of NADs, as has been done for LADs. In addition to Nucleolar-DamID, other creative technologies for visualizing NADs need to be established.

Live cell imaging is an attractive analytical methodology for understanding cellular structures and their dynamics. It can...
provide intuitive and dynamic information over time, which are possibly lost in snapshots from fixed cells. Heterochromatin is critical to the 3D organization of the interphase genome, thereby imaging it in real-time will provide answers to the fundamental question of which mechanisms are involved in the spatial genome organization.

**DISCUSSION**

Recently, higher-order structures of chromatin have become more appreciated for their roles in gene regulation and cellular processes. Unlike sequencing-based methods or FISH that rely on fixed cells, microscopic approaches have been developed, and have opened an era for studying chromatin dynamics in live cells. With an attempt to visualize chromatin organization in living cells, there are many methods proposed for fluorescently labeling specific genomic regions, or whole chromosomes. One of the most well-known methods is the CRISPR/Cas9 system, and its modified techniques, such as CARGO-dCas9 and CRISPR LiveFISH, have recently been introduced to obtain enhanced results from live cell imaging. Moreover, there is another way to label genomic sites in live cells, called the DamID-derived technique. This system uses Dam protein fused with the protein of interest to methylate adenines, and generates m6A modifications at specific regions of the genome. These modified DNA regions are recognized by catalytically inactivated DpnI enzyme fused with fluorescent protein, which allows stable tracking of genomic structures, such as LADs and NADs.

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**CONFLICTS OF INTEREST**

The authors have no conflicting interests.

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