Structure and Epigenetic Regulation of Chromatin Fibers

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In eukaryotic cells, the accessibility of DNA is dependent on the packing density of chromatin fibers. Genomic DNA firstly wraps around a histone octamer to form a nucleosome, which is connected by linker DNA to form the primary chromatin structure, the “beads-on-a-string” nucleosomal array. Nucleosomes in the array are further compacted by linker histone H1 to form a 30-nm chromatin fiber—typically regarded as the secondary structure of chromatin. Chromatin fibers are further organized into other higher-levels of chromatin structures, but so far details of these structural levels still remain obscure. The three-dimensional (3D) organization of genomic DNA plays a critical role in regulating DNA-related biological processes, such as gene transcription and DNA replication, repair, and recombination. Elucidating the structure and dynamics of chromatin fibers in molecular details is the key to understanding the epigenetic regulation of gene expression by different chromatin factors.

HIGH-RESOLUTION STRUCTURE OF 30-NM CHROMATIN FIBERS

It is still a puzzle how genomic DNA is hierarchically organized in eukaryotic cells. From a structural point of view, the DNA double-helix structure discovered by Watson and Crick is surely the most important milestone in molecular biology (Watson and Crick 1953). After more than 40 years, the high-resolution structure of the nucleosome core particle (NCP) has been defined by crystal X-ray studies (Luger et al. 1997), which undoubtedly reveals the structural details of histone–histone and histone–DNA interactions within nucleosomes (Fig. 1). Other high-resolution structures of NCPs containing core histones from different species or with a different DNA sequence, different histone variants, or different nucleosome-binding proteins/peptides have been resolved subsequently to investigate the regulation of nucleosome structure as reviewed previously (Cutter and Hayes 2015; McGinty and Tan 2015; Zhu and Li 2016).

Afterward, the manner of how a “beads-on-a-string” nucleosomal array folds into a condensed 30-nm chromatin fiber remains to be determined. Two basic classes of structural models had been proposed previously based on the studies of native 30-nm fibers in nuclei or isolated from nuclei (Finch and Klug 1976; Langmore and Paulson 1983; Widom and Klug 1985; Gerchman and Ramakrishnan 1987; Ghirlando and Felsenfeld 2008). One is the solenoid model, in which nucleosomes are arranged linearly in a one-start solenoid-type helix with bend linker DNA, and the other is the zigzag model, in which nucleosomes zigzag back and forth in a two-start stack of nucleosomes connected by a relatively straight DNA linker (Woodcock et al. 1984; Widom and Klug 1985; Williams et al. 1986). To discriminate between these two structural models, the detailed structure of chromatin fibers needs to be resolved. However, the heterogeneous properties of nucleosomes in native chromatin with different DNA sequences/linker lengths and different histone compositions/modifications make this difficult. The reconstitution of chromatin fibers in vitro using regular tandem repeats of unique nucleosome-positioning DNA sequences and purified histone proteins greatly improves the reproducibility and uniformity for structural analysis (Dorigo et al. 2004). Using this system, Richmond and colleagues first resolved the crystal X-ray structure of chromatin fibers reconstituted by tetranucleosomal arrays with a 20-bp linker DNA at a high concentration of Mg2+ (120 mM). The structure resolved at ∼9 Å resolution reveals two stacks of nucleosomes connected by straight linker DNA, which agrees with the zigzag model (Schalch et al. 2005). Within each nucleosome stack, strong interactions between the H2B–helix α1/αC and the adjacent H2A–helix α2 stabi-
lize the chromatin fibers (Fig. 1). However, the crystal structure was determined for a tetranucleosomal array, which is too short to form a solenoid structure, and with a very short nucleosome repeat length (NRL; 167 bp) in the absence of H1, which is uncommon in nature.

Recently, we had determined the 3D cryo-electron microscopy (EM) structures at ∼11 Å resolution of longer chromatin fibers reconstituted in vitro from 12-nucleosomal arrays with two different NRLs (177 bp and 187 bp), which reveal a left-handed double helix twisted with the repeating tetranucleosomal structural units (Song et al. 2014). The structures constitute the largest fragments of chromatin resolved at this high resolution so far and provide new insights into the nucleosome arrangement in helical structure of 30-nm chromatin fibers. The four nucleosomes within the structural unit zigzag back and forth to form two stacks of two nucleosome cores connected by straight linker DNA (Schalch et al. 2005; Song et al. 2014). Strong interactions between the H2B–helix α1/αC and the adjacent H2A–helix α2 stabilize each nucleosome stack of tetranucleosomal units. The tetranucleosomal units are twisted against each other in a left-handed manner to form the final double-helical structure of chromatin fiber (Song et al. 2014). Between the structural units, the H1–H1 interaction and the interactions of H4 amino-terminal tail with the H2A/H2B acidic patch between the nucleosomal interfaces play central roles.

The linker histone has been considered to play an essential role in the formation of compact chromatin fiber (Thoma et al. 1979; Allan et al. 1980; Bates and Thomas 1981; Thomas 1999). But the molecular details of how linker histones bind to nucleosome and compact chromatin fibers remain to be determined. Our cryo-EM structures, for the first time, clearly address the location and the role of H1 in the formation of the 30-nm chromatin fiber (Song et al. 2014). H1 directly interacts with both the dyad and the entry/exit nucleosomal DNA in a three-contact mode. The binding of H1, which locks the outer DNA wraps of nucleosomes at the enter/exit point as previously discussed (van Holde and Zlatanova 1996; Syed et al. 2010), enhances the stability of the outer nucleosomal wrap with the increase of ∼20 kJ/mol free energy and accelerates the folding/unfolding rate of the outer wrap.

Figure 1. The hierarchical organization of chromatin fibers: 147 base pairs (bp) of DNA wrapped around a histone octamer (with one histone H3/H4 tetramer and two H2A/H2B dimers) ∼1.7 times in a left-handed manner to form nucleosome, the basic repeating unit of chromatin (Luger et al. 1997). Nucleosomes interact with each other to form the tetranucleosomal structural units, within which the four nucleosomes zigzag back and forth to form two stacks of two nucleosome cores connected by straight linker DNA (Schalch et al. 2005; Song et al. 2014). Strong interactions between the H2B–helix α1/αC and the adjacent H2A–helix α2 stabilize each nucleosome stack of tetranucleosomal units. The tetranucleosomal units are twisted against each other in a left-handed manner to form the final double-helical structure of chromatin fiber (Song et al. 2014). Between the structural units, the H1–H1 interaction and the interactions of H4 amino-terminal tail with the H2A/H2B acidic patch between the nucleosomal interfaces play central roles in interaction and twisting between the structural units (Fig. 1). In comparison to the closely stacked nucleosomes within the tetranucleosomal structural unit, the apparently formed gaps between the structural units may provide a platform for histone modifications or other architectural proteins to modulate the internucleosomal surface interactions in the regulation of higher-order chromatin structures.
In addition, the H1 locates asymmetrically in each nucleosome core with its bulk of globular domain pointed outside the structural units, which allows the self-association of H1 by its globular domain between tetranucleosomal units and imparts an additional twist between each structural unit. However, the 11 Å resolution of the resolved cryo-EM structures cannot give the structural details of H1 at the atomic levels. Higher resolution structures of chromatin fibers still need to be determined to solve these problems.

SINGLE-MOLECULE STUDY ON THE DYNAMICS OF CHROMATIN FIBERS

The 30-nm chromatin fiber has been shown to be the first level of the transcriptionally dormant platform (Li et al. 2010). Thus, the structural transition between the 30-nm chromatin fiber and the nucleosomal array plays a critical role in regulating the accessibility of the DNA template (Li et al. 2010; Li and Reinberg 2011). Static conformations obtained by our cryo-EM structures of 30-nm chromatin fibers in vitro cannot provide much detailed information for such a dynamic process. In addition, because of the highly dynamic and heterogeneous properties of the chromatin fiber, it is technically challenging to study the dynamics of the chromatin fibers. Recently, several single-molecule techniques have been applied to investigate the dynamics of nucleosome/chromatin structures (Cui and Bustamante 2000; Pope et al. 2005; Kruithof et al. 2009). Mechanical manipulations of single mononucleosome show that a nucleosome in the absence of linker histone unravels in two major stages: the outer nucleosomal DNA unwrapped at ~3 pN and the inner turn unwrapped at ~10 pN (Brower-Toland et al. 2002; Hall et al. 2009; Bintu et al. 2012). For the mechanical decompaction of higher-order chromatin fibers, Bustamante and colleagues observed a distinct structural transition between “compacted” and “extended” states for the isolated native chicken erythrocyte chromatin fibers at medium ionic strengths (~40 mM NaCl) at 5–6 pN, which was attributed to the disruption of internucleosomal interactions for stabilizing the higher-order structures of chromatin fibers (Cui and Bustamante 2000). Van Noort and colleagues used magnetic tweezers to investigate the mechanical stretching of single reconstituted chromatin fibers and also observed the dynamic unfolding behavior of chromatin fibers at low forces (~0 pN) (Kruithof et al. 2009; Meng et al. 2015). They found that the chromatin fibers with 197-bp NRL stretch like a Hookian spring, which supports a solenoid topology with a high nucleosome–nucleosome stacking energy. Using magnetic tweezers (Fig. 2A), we investigated the hierarchical organization and dynamics of 30-nm chromatin fibers reconstituted in vitro in the presence of H1, whose high-resolution 3D cryo-EM structures

Figure 2. Mechanical unfolding of 30-nm chromatin fibers by single-molecular magnetic tweezers. (A) Schematic setup of the magnetic tweezers used in chromatin fiber studies (not to scale). (B) Comparison of two typical force–extension curves of a chromatin fiber with H1 (blue curve) and without H1 (orange curve) in HE buffer. Three major distinct stages as labeled can be recognized in the force-extension curve of the chromatin fiber with H1. The inset shows the details of Stage I at low forces (~8 pN). (C) Stepwise folding/unfolding dynamics of tetranucleosomal units with two alternative pathways at 3.5 pN for the 177-bp nucleosome repeat length (NRL)’s chromatin fiber with H1. (D) Model for the dynamic organization of chromatin fibers. The left-handed double-helical chromatin fiber unfolds to a “tetranucleosomes-on-a-string” extended structure; then the tetranucleosomal unit unfolds to a complete open nucleosomal array in one or two steps. (Adapted, with permission, from Li et al. 2016.)
have been resolved recently (Song et al. 2014; Li et al. 2016). We showed that mechanical unfolding of a 30-nm fiber is a multistep process (Fig. 2B). Under increasing tensile force applied, the compacted chromatin fiber first unfolds to a “tetranucleosomes-on-a-string” extended structure by disrupting the internucleosome interactions between tetranucleosomal units. The tetranucleosomal unit exists as a stable structural intermediate of the 30-nm chromatin fiber and further unfolds to a more extended “beads-on-a-string” conformation by disrupting the nucleosome–nucleosome interactions within the tetranucleosomal unit in a two-step, three-transition-state process at force of ~3.5 pN, which is characteristic of three transitions with the step sizes of 1L, 2L, and 3L (in which L is the linker length between the adjacent nucleosomes), respectively (Fig. 2C,D). Further increasing force causes nucleosome unwrapping in two major stages, including the one-step disruption of the outer DNA turn and the one-step unwrapping of the inner turn, consistent with previous investigations (Brower-Toland et al. 2002; Hall et al. 2009; Bintu et al. 2012; Li et al. 2016). Importantly, similar dynamic processes can be observed in the chromatin fibers assembled not only on regular tandem repeat of the 601 DNA sequence but also on the scrambled (nonrepetitive) DNA sequence, suggesting that the existence of tetranucleosomal units is not dependent on DNA sequence. Moreover, the low energy required to disrupt the interaction between two tetranucleosomal units (~1.8 kBT), which is comparable to the thermal fluctuations, suggests that chromatin fibers may undergo spontaneously rapid folding/unfolding dynamics between a compact regular 30-nm chromatin fiber and an extended “tetranucleosomes-on-a-string” at physiological conditions in vivo. Thus, it is likely that the chromatin fibers in vivo probably adopt an irregular “tetranucleosomes-on-a-string” structure, which combines fully folded regular zigzag tetranucleosomal clusters/stacks with partially unfolded “nucleosomal arrays” or “nucleosomal clutches” regions (Collepardo-Guevara and Schlick 2011).

**Figure 3.** Epigenetic regulation of chromatin fibers by chromatin factors. (A) Schematic model of 30-nm chromatin fibers with the three important interaction interfaces as indicated by arrows, including the DNA–histone interaction interfaces within nucleosome (1), internucleosome interaction interfaces within tetranucleosomal unit (2), and interaction interfaces between tetranucleosomal units (3). (B) The different chromatin factors potentially regulate the interaction interfaces of 30-nm chromatin fibers.

**EPIGENETIC REGULATION OF CHROMATIN FIBERS BY CHROMATIN FACTORS**

In the past 20 years, many chromatin factors, including chromatin modifications, histone chaperones, histone variants, chromatin remodelers, and chromatin architectural proteins, have been shown to be involved in regulating chromatin dynamics (Luger and Hansen 2005; Hake and Allis 2006; Greaves et al. 2007). Deciphering the structure and dynamics of the 30-nm chromatin fibers in molecular details is essential for understanding such regulations. Our cryo-EM structures clearly imply the presence of three important interaction interfaces in the 30-nm fiber—namely, DNA–histone interaction interfaces within the nucleosome, internucleosome interaction interfaces within the tetranucleosomal unit, and interaction interfaces between tetranucleosomal units, which can be potentially regulated by different chromatin factors (Fig. 3). The newly developed in vitro single-molecule techniques and in vivo genomic approaches enable us to investigate the epigenetic regulations of 30-nm fibers by different chromatin factors (Hsieh et al. 2015; Li et al. 2016; Risca et al. 2017).

Previously, FACT, a conserved histone chaperone for H2A–H2B dimers, has been shown to destabilize nucleosomes for RNA polymerase progression on chromatin templates (Orphanides et al. 1998, 1999; Belotserkovskaya et al. 2003) and maintain chromatin structure in vivo during DNA transcription, replication, and repair (Saunders et al. 2003; Fujimoto et al. 2012; Formosa 2013; McCullough et al. 2015). However, the molecular mechanisms of how FACT remodels chromatin remain largely unclear. As discussed above, the nucleosomal stacks within tetranucleosomal unit are mainly stabilized by the interactions between the H2B–helix α1/αC and the adjacent H2A–helix α2 (Song et al. 2014). A recent structural study revealed that the recognition of H2A–H2B heterodimer by FACT is mainly mediated by the interactions between the U-turn motif of Spt16M and the amino-terminal α1 helix of H2B (Hondele et al. 2013), suggesting that
FACT may remodel chromatin fibers via interfering these interfacial interactions. Indeed, our single-molecule measurements showed that FACT can remodel higher-order chromatin structure by destabilizing the tetranucleosomal unit (Li et al. 2016). More importantly, genomic analyses showed that FACT destabilizes the N/N+2 Micro-C interactions (tetranucleosomal motifs) to facilitate gene transcription in yeast. Distinct domains of FACT have been shown to interact with H2A/H2B or H3/H4 separately (Stuwe et al. 2008; Zhang et al. 2015; Tsanaka et al. 2016), suggesting that these distinct domains of FACT might be responsible for their different activities in remodeling the nucleosome and/or the tetranucleosomal unit. It will be of great interest to generate a series of FACT mutants to resolve the distinctive activity in remodeling the nucleosome and/or the tetranucleosome using single-molecule measurements. Beside FACT, several other chromatin remodelers (such as INO80, which recognizes the H2B αC helix) (Tosi et al. 2013), or certain repressive chromatin factors (such as PRC1, which has been shown to compact three to four nucleosomes) (Francis et al. 2004) may also fulfill their biological functions via modulating the dynamics or stability of tetranucleosomal units. The tetranucleosomal unit may provide an additional level of gene regulation beyond the nucleosome. It will be of great interest to identify these factors and decipher their regulatory interactions with the tetranucleosomal units.

The incorporation of histone variants has been shown to regulate the structure and dynamics of chromatin fibers and create architecturally distinct chromatin states that play diverse functions in genome-associated biological processes. Among them, histone variant H3.3 has been largely considered as a mark of transcriptionally activated genes and has been deposited into transcribed genes, promoters, and gene-regulatory elements (Schwartz and Ahmad 2005; Jin et al. 2009; Goldberg et al. 2010). However, a few recent studies showed that H3.3 was also incorporated at regions of the genome that are typically thought to be relatively transcriptionally inactive, such as the telomere and pericentric heterochromatin in mouse embryonic stem cells (van der Heijden et al. 2007; Goldberg et al. 2010; Wong et al. 2010). H3.3 differs from canonical H3 at only four amino acid residues, three are hidden inside the NCP in region 87–90 and one residue, Ser31, is exposed outside of the NCP. We and others have shown that Ala87 and Gly90 are the principal determinants of human H3.3 specificity in DAXX recognition (Elsässer et al. 2012; Liu et al. 2012). We also investigated how H3.3 regulates the nucleosome/chromatin dynamics and gene transcription (Fig. 4; Chen et al. 2013). Our FRET and magnetic tweezers experiments showed that H3.3 has little effect on the stability of nucleosomes, which was consistent with previous findings (Flaus et al. 2004; Thakar et al. 2009), but it greatly impairs the compaction of chromatin by analytical ultracentrifugation (AUC) and EM analyses (Chen et al. 2013). Interestingly, we also showed that the histone variant H2A.Z not only stabilizes the nucleosome but also facilitates compaction of nucleosomal arrays into a featured “ladder-lier” chromatin fiber. Although the incorporation of H3.3 does not affect the stabilization effect of H2A.Z on nucleosomes, it counteracts H2A.Z-mediated chromatin compaction (Fig. 4A,B). Residues 89 and 90 of H3.3 were mainly responsible for the counteractivity of H2A.Z-mediated chromatin compaction. Moreover, we found that H3.3 could antagonize the inhibitory effects of H2A.Z on chromatin transcription in vitro by RNA Pol II, which partially resulted from the counteractivity of compaction by H2A.Z. Our results suggest that H3.3 may play a dominant role in regulation of the dynamics of higher-ordered chromatin and transcriptional activity in the chromatin context. To this end, we further analyzed the dynamic depositions and/or replacement of H2A.Z and H3.3 and the corresponding structural changes of chromatin at the enhancer and promoter regions of RAR/RXR targeted genes during gene activation by all-trans-retinoid acid (tRA) induction. Our results (Fig. 4C) revealed that deposition of H2A.Z results in compaction of chromatin at promoter regions, which inhibits the transcription of the Cyp26a1 and Hoxa1 genes before tRA induction, whereas the incorporation of H3.3 at enhancer regions decorates the chromatin architecture to a relatively open conformation, allowing the recognition and binding of transcriptional activators, which subsequently recruits the ATP-dependent chromatin remodeling complexes and/or histone-modifying enzymes to remodel the nucleosome architecture at promoter regions upon gene induction, as previously reported (Li et al. 2010). Our results provide new insights into the molecular mechanism of how histone variants function cooperatively to establish featured chromatin structures at enhancer and promoter regions to prime inducible genes for rapidly activation in response to environmental stimulation.

The centromere-specific histone H3 variant (CenH3 [CENP-A in humans]), is an epigenetic factor essential for the centromere identity and function (Allshire and Karpen 2008). In human cells, CENP-A is specifically recognized and deposited into centromeres by its chaperone and assembly factor HJURP (Dunleavy et al. 2009; Foltz et al. 2009). Our structural and biochemical analysis showed that other than the CMTD of CENP-A, previously identified as the exclusive region responsible for HJURP binding (Black et al. 2004; Bassett et al. 2012), the residue Ser68 also plays essential roles for HJURP binding (Hu et al. 2011). We further showed that the dynamic phosphorylation/dephosphorylation of Ser68 in CENP-A, which is mediated by the Cdk1/Cyclin B and PP1α complex, temporally controls the HJURP-mediated assembly of CENP-A into centromeric regions (Yu et al. 2015). CENP-N, an indispensable member of CCAN (the constitutive centromere-associated network), was identified as the first “reader” of epigenetic marks present in the CENP-A-containing chromatin (Carroll et al. 2009). CENP-N is dynamically recruited to centromeres/kinetochore during the cell cycle (Hellwig et al. 2011) and plays an essential role in the assembly of kinetochore complex at centromeres and faithful segregation of sister chromatids during cell division (Foltz et al. 2006; Carroll et al. 2009). By using biochemical, biophysical, and cell-based assays, we showed that the RG loop, the two CENP-A-specific residues Arg80 and Gly81 in loop1 located at the lateral surface of CENP-A
nucleosome (Tachiwana et al. 2011), not only provides the recognition site for the binding of CENP-N to the CENP-A nucleosome, but also facilitates the folding of CENP-A arrays into a compact “ladder-like” chromatin structure in the presence of MgCl₂, as revealed by AUC and EM analyses (Fig. 5A–C; Fang et al. 2015). Interestingly, we found that the formation of a compact “ladder-like” structure of CENP-A chromatin inhibits the binding and recruitment of CENP-N through concealing/hiding the RG loop within chromatin fiber. More importantly, our fluorescence resonance energy transfer (FRET) analysis and SNAP imaging showed that upon G1/S phase transition, centromeric chromatin switches from the compact to an open state because of the dilution of CENP-A nucleosome during DNA replication, which enables the now-exposed RG loop to recruit CENP-N at the S phase (Fig. 5D).

Figure 4. Histone variants H3.3 and H2A.Z regulate chromatin structures cooperatively for gene activation. (A) Sedimentation coefficient distribution plots for the canonical, H2A.Z-, H3.3-, and double-variant H2A.Z/H3.3-containing nucleosomal arrays at 0, 1.0, and 1.5 mM MgCl₂. (B) Negatively stained electron microscopic (EM) images of the canonical, H2A.Z-, H3.3-, and double-variant H2A.Z/H3.3-containing chromatin fibers in 1.0 mM MgCl₂. Scale bar, 100 nm. (C) The model for the dynamic regulation of H2A.Z and H3.3 on chromatin structures at the enhancer and promoter regions during the gene activation process by the addition of trans-retinoic acid (tRA). (Adapted, with permission, from Chen et al. 2013.)
ORGANIZATION OF CHROMATIN FIBERS IN NUCLEUS

The 30-nm chromatin fiber has long been thought to be the first level of the hierarchical chromatin compaction, but the existence of the 30-nm fiber in vivo still remains controversial. Studies of purified or reconstituted chromatin have provided evidence supporting the existence of longitudinally compacted 30-nm chromatin fibers; however, until now high-resolution imaging of chromatin in living cells had not been possible (Elstov et al. 2008; Fussner et al. 2011). Early cryo-EM studies have shown that a 30-nm fiber is indeed the most predominant form of starfish sperm and nucleated chicken erythrocyte chromatin (Horowitz et al. 1994; Woodcock 1994; Scheffer et al. 2011). Using cryo-EM tomography of vitreous sections, Frangakis and colleagues showed that the most predominant form of chromatin in chicken erythrocyte nuclei is indeed a 30-nm fiber arranged in a two-start helix formation with \( \sim 6.7 \) nucleosomes per turn, in which the nucleosomes are juxtaposed face to face (Scheffer et al. 2011). In addition, the stacked nucleosomes were shown to be shifted off their superhelical axes, with an axial translation of \( \sim 3.4 \) nm and an azimuthal rotation of \( \sim 54° \), very consistent with the high-resolution structure of in vitro reconstituted chromatin fibers (Schalch et al. 2005; Song et al. 2014). In addition, Rando and colleagues recently developed a novel Hi-C-based method named ‘‘Micro-C’’ to probe nucleosome organization at nucleosome resolution in yeast (Hsieh et al. 2015). Despite the lack of periodicity...
in their data set, they showed that N/N+2 and N/N+1 nucleosome pairs are similarly abundant in the whole genome, which suggests the wide existence of tetraneu-
coleosomal folding motifs in yeast genome. Similarly, using the EM-assisted nucleosome interaction capture (EMANIC) cross-linking experiments in combination with mesoscale modeling of chromatin fibers, Schlick and colleagues showed a dominant relaxed two-start zigzag organization rather than longitudinal compaction associated with the 30-nm fiber (Grigoryev et al. 2009). Most recently, combining the ionizing radiation-induced spatially correlated cleavage of DNA with deep-sequencing technique (RICC-seq), Greenleaf and colleagues provided the first genome-wide map of the chromatin secondary structure in living human cells at the one- to three-nucleosome (50- to 500-bp) scale (Risca et al. 2017). Unbiased analysis of RICC-seq signals in intact interphase nuclei reveals that RICC-seq fragmentation patterns in H3K9me3- and H3K27me3-marked heterochromatin regions are consistent with variable longitudinal chromatin compaction of two-start helical fibers with face-to-face stacked alternating nucleosomes in tri- or tetranucleosome units, as seen in the structure of reconstituted chromatin fiber (Song et al. 2014).

This overwhelming body of experimental evidences from EM and genomic studies strongly support that two-start helical fibers with stacked alternating nucleosomes are an important mechanism for generating chromatin compaction both in vitro and in vivo. However, cryo-EM (Gan et al. 2013), X-ray scattering (Nishino et al. 2012; Maeshima et al. 2014), and electron spectroscopy imaging (ESI) studies (Bazett-Jones et al. 2008; Fussner et al. 2011b, 2012) of the nucleus do not support the existence of the regular 30-nm fiber within intact cells (Eltsov et al. 2008; Fussner et al. 2011a,b; Nishino et al. 2012). Because of the poor contrast of DNA in vitreous ice and the limited 3D sampling volume of ESI, it is technically difficult to identify chromatin unambiguously and to reconstruct 3D organization of chromatin through large nuclear volume in cryo-EM experiments. Most recently, O’Shea and colleagues developed a novel ChromEMT technique by combining electron microscopy tomography (EMT) with a labeling method, which explores a fluorescent dye (DRAQ5) that binds to DNA and enhances the contrast of DNA, enabling chromatin to be visualized with OsO4 in EM (Ou et al. 2017). Using this technique, they only observed a flexible and disordered granular chroma-
tin chain with diameters between 5 and 24 nm, but no regular higher-order chromatin fibers in human interphase and mitotic cells in situ. However, a few important issues still need to be clarified in the technique. The first is whether DRAQ5 also binds to other type nucleic acids in situ, such as RNA (particularly double-stranded RNA [dsRNA]). It is unclear how to distinguish chromatin fibers from other nucleic acid–protein particles, which are abundant and distributed widely in nuclei. Second, no evidence in vitro or in vivo showed that DRAQ5 can bind equally to free DNA and chromatin DNA. It is likely that the DNA in compacted chromatin fragments may not be accessible for staining by the dye. In addition, the path of DNA in the chromatin fiber cannot be directly observed in vivo by the ChromEMT technique under such low resolution. As discussed above, we and others showed that chromatin fibers mainly adopt a two-started helical conformation with alternating nucleosomes stacked face to face in vitro and in vivo, in which the two nucleosomal stacks are connected by linker DNA and distanced from each by \(-7.5–10\) nm (Bednar et al. 1998; Song et al. 2014; Li et al. 2016). In this context, when the linker DNA cannot be discriminated, only two separated nucleosomal stacks with a diameter of \(-10\) nm can be actually observed in ChromEMT. Moreover, extensive fixation by chemical cross-linking and dehydration by ethanol during sample preparation in ChromEMT may damage the ultrastructure and 3D organization of chromatin fiber. Therefore, it is still a big challenge to preserve the natural 3D conformation of chromatin fiber in ChromEMT or cryo-EM tomography.

PERSPECTIVES AND CONCLUSION

Our 3D cryo-EM structures at 11 Å resolution have provided the fundamental structural features of the elusive 30-nm chromatin fiber and a solid foundation for understanding the basic principle of chromatin compaction, whereas higher-resolution structures of chromatin fiber are needed to uncover much more structural details for the nucleosome–nucleosome, nucleosome–H1, and H1–H1 interactions in chromatin fibers. In addition, our cryo-
EM structures clearly imply the presence of three important interaction interfaces in the 30-nm fiber, whereas it still remains unclear how these interfaces are regulated by different chromatin factors. Regarding the variation of NRLs in vivo, reconstituted chromatin fibers with a combina-
tion of different NRLs will also be good candidates for single-molecule and cryo-EM studies in the future. These further studies will not only enhance our understanding of the diversity of chromatin structures in vivo but also provide structural basis for how different combina-
tions of DNA sequences, NRLs, histone variants, chroma-
tin modifications, and chromatin architectural proteins can be coordinated to precisely regulate the biological function of genomic DNA in the nucleus.

It is still a puzzle as to whether the structural results from the in vitro studies can represent the actual structure of chromatin fibers in situ. Therefore, the 3D organization of chromatin fibers in the intact nuclei needs to be further studied by using newly developed techniques. Well-char-
acterized chromatin fibers reconstituted in vitro, including compacted 30-nm fibers and open nucleosomal arrays, would be perfect structural references for analyzing the 3D organization of chromatin fibers in situ in these stud-
ies. The combination of cryo-EM with super-resolution fluorescence imaging techniques has been recently developed to visualize and quantify the ultrastructure of cryopreserved cells (Chang et al. 2014; Liu et al. 2015). The combination of genomic approaches (such as micro-C and RICC-seq) and CRISPR (clustered regularly interspaced short palindromic repeat)-based imaging techniques may
enable us to probe the ultrastructure and 3D organization of chromatin fiber at defined genomic regions in the intact nuclei. Undoubtedly, more structural detail for the 3D organization of chromatin fibers in situ can be obtained by the application of these advanced imaging techniques in the future.

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