The prenucleosome, a stable conformational isomer of the nucleosome

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Chromatin comprises nucleosomes as well as nonnucleosomal histone–DNA particles. Prenucleosomes are rapidly formed histone–DNA particles that can be converted into canonical nucleosomes by a motor protein such as ACF. Here we show that the prenucleosome is a stable conformational isomer of the nucleosome. It consists of a histone octamer associated with ~80 base pair (bp) of DNA, which is located at a position that corresponds to the central 80 bp of a nucleosome core particle. Monomeric prenucleosomes with free flanking DNA do not spontaneously fold into nucleosomes but can be converted into canonical nucleosomes by an ATP-driven motor protein such as ACF or Chd1. In addition, histone H3K56, which is located at the DNA entry and exit points of a canonical nucleosome, is specifically acetylated by p300 in prenucleosomes relative to nucleosomes. Prenucleosomes assembled in vitro exhibit properties that are strikingly similar to those of nonnucleosomal histone–DNA particles in the upstream region of active promoters in vivo. These findings suggest that the prenucleosome, the only known stable conformational isomer of the nucleosome, is related to nonnucleosomal histone–DNA species in the cell.

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function as a more dynamic [less static and repressive] form of histone-bound DNA than canonical nucleosomes.

It was therefore important to gain a better understanding of the prenucleosome. Does it contain a histone octamer or some other combination of the histones? How much DNA is associated with a prenucleosome, and where is the DNA located relative to the histone octamer? Why do prenucleosomes not supercoil DNA like nucleosomes? In previous work, we assembled prenucleosomes onto relaxed plasmid DNA (Torigoe et al. 2011). While this approach has been useful for the characterization of the function of prenucleosomes in the chromatin assembly process, the resulting templates were heterogeneous in terms of the number and locations of the prenucleosomes. Hence, with that system, it was not possible to determine the composition and structure of the prenucleosome.

In this study, we took a new experimental approach to the analysis of prenucleosomes and determined that the prenucleosome is a stable conformational isomer of the nucleosome. We further found that the properties of prenucleosomes assembled in vitro are remarkably similar to those of nonnucleosomal histone–DNA particles. This partial wrapping of DNA around the histones in prenucleosomes corresponds to less than one turn of the DNA around the histone octamer in a nucleosome. This partial wrapping of DNA around the histones in prenucleosomes may have been due to the presence of some canonical nucleosomes, and the resulting samples were subjected to psoralen cross-linking and denaturing electron microscopy. Bubble sizes were measured in ImageJ and converted from nanometers to nucleotides. A total of 4623 prenucleosome bubbles and 5013 nucleosome bubbles was measured in four independent experiments. The plot displays the distribution of bubble sizes as the average ± standard deviation (n = 4) of 10-nt bins. The individual data points are placed at the center of the 10-nt bins. (C) Comparison of the psoralen bubble sizes observed in vitro and in vivo. The data from prenucleosomes and nucleosomes in vitro [shown in B] and at the active versus repressed PHO5 promoters in vivo in Saccharomyces cerevisiae [Brown et al. 2013]. The plot shows the distribution of bubble sizes as the average of 10-nt bins.

The measurement of the bubble sizes revealed a peak at 140–150 nucleotides [nt] with canonical nucleosomes, as expected. With prenucleosomes, however, we observed a peak at ~70–80 nt [Fig. 1B] as well as larger bubbles that may have been due to the presence of some canonical nucleosomes in the predominantly prenucleosomal samples [as in Torigoe et al. 2011]. These findings suggest that prenucleosomes associate with ~70–80 nt of DNA, which corresponds to less than one turn of the DNA around the histone octamer in a nucleosome. This partial wrapping of DNA around the histones in prenucleosomes...
(relative to that in nucleosomes) might explain the lack of DNA supercoiling that is observed during the formation of prenucleosomes [Torigoe et al. 2011].

These studies additionally enabled us to compare the properties of prenucleosomes assembled in vitro and nonnucleosomal chromatin particles observed in vivo. Remarkably, the biochemical data on prenucleosomes versus nucleosomes (Fig. 1B) exhibit a close resemblance to the psoralen bubble size distribution observed in vivo with the active versus repressed PHO5 promoter in yeast [Brown et al. 2013]. In the in vivo analysis, a peak bubble size of 70–80 bp of DNA was seen at the activated PHO5 promoter, whereas a peak of 140–150 bp was observed at the repressed PHO5 promoter. To demonstrate the correlation, we directly compared the in vitro data (this study) and in vivo data [Fig. 1C, Brown et al. 2013]. These data suggest that the active PHO5 promoter contains both prenucleosomes [or prenucleosome-like particles] and nucleosomes, whereas the repressed PHO5 promoter contains mostly canonical nucleosomes. Thus, although prenucleosomes were initially identified in the analysis of chromatin assembly, these findings indicate that they may also be present at the promoter regions of active genes.

**Figure 2.** Rapid and efficient formation of mono-prenucleosomes with 80-bp DNA fragments. (A) The NAP1-mediated formation of mono-prenucleosomes with an 80-bp genomic DNA fragment occurs rapidly and requires all four core histones. Histone–NAP1 complexes were combined with an 80-bp DNA fragment (an 80-bp segment in the coding sequence of the *Drosophila melanogaster* ISWI gene; henceforth termed the “80-bp genomic DNA”). The samples were incubated at room temperature for 30 sec and then subjected to native (nondenaturing) 5% polyacrylamide gel electrophoresis. The DNA was visualized by staining with ethidium bromide. One octamer equivalent of all four core histones per DNA would be a 1:1 octamer:DNA ratio. Note that one octamer equivalent of H2A+H2B+H3+H4 has the same amount of H3 and H4 as one octamer equivalent of H3–H4. (B) Mono-prenucleosomes can be formed with the central 80 bp of the 601 nucleosome positioning sequence. Mono-prenucleosomes were formed and analyzed as in A, with all four core histones along with either the 80-bp genomic DNA or the central 80 bp of the 601 sequence. (C) Mono-prenucleosomes appear to be the thermodynamically most stable arrangement of the four core histones and 80 bp of DNA at 50 mM NaCl. Mono-prenucleosomes were formed with the dNLP histone chaperone as well as by salt dialysis of the four core histones with the 80-bp genomic DNA fragment. For comparison, H3–H4 monotetrasomes were also generated in parallel by salt dialysis with H3–H4. The histones were used at an octamer equivalent:DNA ratio of 1.0. (D) Mono-tetrasomes can be converted into prenucleosomes by the addition of H2A–H2B. Monotetrasomes were formed by salt dialysis with the 80 bp of genomic DNA as in C. Next, NAP1–H2A–H2B complexes were added as indicated. The samples were incubated for 30 sec at room temperature and then subjected to native 5% polyacrylamide gel electrophoresis. As a reference, a mono-prenucleosome formed by salt dialysis as in C was included ["Mono-prenuc"].
this work, we refer to these nucleoprotein complexes as monomer prenucleosomes or “mono-prenucleosomes.”

We then tested whether mono-prenucleosomes can be assembled with different DNA segments. In Figure 2A, we used an 80-bp stretch of DNA from the coding sequence of the *Drosophila melanogaster* ISWI gene (sequence given in Supplemental Table S1); we refer to this fragment as the “80-bp genomic DNA.” We also used the central 80 bp of the 601 nucleosome positioning sequence (for sequence, see Supplemental Table S1; Lowary and Widom 1998); we term this fragment the “central 80 bp of 601 sequence.” As shown in Figure 2B, mono-prenucleosomes can be formed efficiently with either the 80-bp genomic DNA or the central 80 bp of 601 sequence. These two DNA fragments are used throughout this study.

Next, we sought to determine whether mono-prenucleosomes are generally formed under different reaction conditions. To this end, we tested a different core histone chaperone, *Drosophila* nucleoplasm-in-like protein (dNLP) (Ito et al. 1996) as well as salt dialysis techniques (Stein 1989) for the reconstitution of mono-prenucleosomes. These experiments revealed that mono-prenucleosomes are efficiently formed with not only NAP1 but also dNLP and by salt dialysis in the absence of histone chaperones (Fig. 2C). Thus, in a minimal purified reaction consisting of only the four core histones and 80 bp of DNA, mono-prenucleosomes can be formed by salt dialysis under standard conditions used for nucleosome reconstitution, in which the histones and DNA are combined in 1 M NaCl, and the samples are slowly dialyzed in a stepwise manner to 0.05 M NaCl (Stein 1989). These results indicate that the mono-prenucleosome is a thermodynamically stable arrangement of the four core histones and 80 bp of DNA.

It should also be noted that mono-prenucleosome reconstitution requires a histone chaperone, such as NAP1 or dNLP, or initial high-salt conditions that prevent the formation of histone–DNA aggregates. If core histones are added directly to free DNA in low-salt buffer (for example, with ≤100 mM NaCl), insoluble histone–DNA aggregates are formed that do not enter the gel (Fig. 2C).

Comparison of assembly by salt dialysis with all four core histones or equimolar amounts of histones H3–H4 reveals the efficient formation of mono-prenucleosomes with the four core histones and inefficient formation of mono-tetrasomes with histones H3–H4 (Fig. 2C). Moreover, we observed that mono-tetrasomes can be efficiently converted into mono-prenucleosomes by the addition of NAP1–H2A–H2B complexes (Fig. 2D). These findings reveal that mono-tetrasomes are distinct from mono-prenucleosomes.

Analysis of the composition and salt lability of mono-prenucleosomes

We next analyzed the composition of mono-prenucleosomes by sucrose gradient sedimentation analysis. In these experiments, mono-prenucleosomes were reconstituted by either NAP1 deposition or salt dialysis and then subjected to 10%–30% sucrose gradient sedimentation. The presence of mono-prenucleosomes was detected by native gel electrophoresis of the nucleoprotein complexes as in Figure 2, and the histones were detected by SDS–polyacrylamide gel electrophoresis and silver staining. These experiments revealed that mono-prenucleosomes cosediment with all four core histones (Fig. 3A).

![Figure 3](https://example.com/figure3.png)

**Figure 3.** Mono-prenucleosomes contain all four core histones and are distinct from hexasomes. [*A*] Sucrose gradient sedimentation analysis reveals that mono-prenucleosomes contain all four core histones. Mono-prenucleosomes were prepared by either NAP1-mediated deposition ([left panels] or salt dialysis ([right panels]) and then subjected to 10%–30% [w/v] ([left to right]) sucrose gradient sedimentation in a Beckman SW41 rotor (32,000 rpm for 18 h at 4°C). The arrows indicate the direction of sedimentation. [*Top panels*] The presence of mono-prenucleosomes was detected by native polyacrylamide gel electrophoresis and ethidium bromide staining of the DNA. [*Bottom panels*] The protein composition was analyzed by SDS–polyacrylamide gel electrophoresis and silver staining. The top two fractions and the bottom fraction did not contain histones (for example, see Supplemental Fig. S2A) and are not included. The sedimentation of the free core histones relative to prenucleosomal histones is shown in Supplemental Figure S2A. [*B*] Mono-prenucleosomes contain two copies of H2A and thus appear to contain a core histone octamer rather than hexamer. Mono-prenucleosomes were reconstituted with recombinant core histones onto the 80-bp genomic DNA by NAP1-mediated histone deposition. The H2A species were a combination of wild-type H2A and Strept-H2A at a 3:1 ratio of H2A:Strept-H2A. Prenucleosomes containing Strept-H2A were pulled down with streptavidin beads and then analyzed by Western blot with antibodies against histone H2A. An H2A Western blot and silver-stained SDS gel are also shown for the input samples. The Western blots were detected and quantitated by using 32P-labeled protein A.
As a control, histones in mono-prenucleosomes sediment faster than free histones (Supplemental Fig. S2A). In addition, it can be seen that the sedimentation rate of NAP1-assembled mono-prenucleosomes is the same as that of salt dialysis-reconstituted mono-prenucleosomes [Fig. 3A]. Throughout this study, we found that mono-prenucleosomes that are prepared by either method have the same biochemical properties.

Although the four core histones can be seen in mono-prenucleosomes [Fig. 3A], it was possible that the histones in mono-prenucleosomes exist as a hexamer (with two copies of H3 and H4 and only one copy of H2A and H2B) rather than as an octamer (with two copies each of H2A, H2B, H3, and H4) [for example, see Arimura et al. 2012]. To test the hexamer hypothesis, we reconstituted mono-prenucleosomes with a 3:1 ratio of wild-type H2A to Strep-tagged H2A along with the other three core histones at a 1:1:1:1 ratio of total H2A [wild-type H2A + Strep-H2A]:H2B:H3:H4. We then pulled down the mono-prenucleosomes containing Strep-tagged H2A and examined whether wild-type H2A coprecipitates with Strep-H2A by Western blot with antibodies against H2A (Fig. 3B). If there were a hexamer of histones in a mono-prenucleosome, then wild-type H2A would not coprecipitate with the Strep-H2A because of the single copy of H2A in a hexamer. However, these experiments revealed the presence of two copies of H2A in mono-prenucleosomes, as there was a ratio of approximately 1.0:1.4 of wild-type H2A to Strep-H2A. This ratio is nearly identical to the theoretical expectation of 1:0.1:1.3 for an octamer [given the 3:1 ratio of wild-type H2A to Strep-tagged H2A prior to precipitation]. These results therefore support the conclusion that a mono-prenucleosome comprises 80 bp of DNA and a histone octamer.

We additionally tested the salt lability of mono-prenucleosomes relative to mononucleosomes. In these experiments, we prepared mono-prenucleosomes and mononucleosomes by salt dialysis [with 80-bp and 146-bp DNA fragments, respectively]; adjusted the final NaCl concentrations to 0.1 M, 0.3 M, 0.8 M, or 2.0 M; and then subjected the samples to sucrose gradient sedimentation at the same salt concentrations [Supplemental Fig. S2B]. As a reference, mononucleosomes partially disassemble at 0.8 M NaCl, as seen with native chromatin (for example, see Germond et al. 1976), and are completely disassembled at 2.0 M NaCl. In contrast, mono-prenucleosomes begin to disassemble at 0.3 M NaCl and are substantially disassembled at 0.8 M NaCl. Thus, consistent with the fewer histone–DNA contacts in prenucleosomes relative to nucleosomes, mono-prenucleosomes are more salt labile than canonical nucleosomes.

**Mono-prenucleosomes can be converted into canonical nucleosomes by ACF**

A key property of prenucleosomes is their ability to be converted into canonical nucleosomes by ACF [Torigoe et al. 2011]. Therefore, we tested whether mono-prenucleosomes can be converted into canonical nucleosomes. To address this question, we formed poly-prenucleosomes by head-to-tail ligation of the 80 bp of DNA in mono-prenucleosomes to free DNA. Next, we incubated the poly-prenucleosomes with ACF and ATP and then digested the reaction products extensively with MNase, which converts nucleosome arrays into core particles that contain ~147 bp of DNA [Fig. 4A]. This experiment revealed that ACF is able to convert poly-prenucleosomes into canonical nucleosomes, as assessed by the generation of the ~147-bp DNA species that is diagnostic of core particles. In the absence of ACF, we did not detect 147-bp DNA fragments upon digestion of the poly-prenucleosomes with MNase. Thus, the conversion of poly-prenucleosomes to polynucleosomes is dependent on ACF. In addition, the ability of mono-prenucleosomes to be converted into canonical nucleosomes further supports the conclusion that mono-prenucleosomes contain a complete histone octamer.

In related experiments, we observed the conversion of mono-prenucleosomes to mononucleosomes by ligation of free DNA tails [85 bp] onto mono-prenucleosomes followed by the addition of ACF [Fig. 4B]. As seen with the ligated poly-prenucleosomes [Fig. 4A] as well as prenucleosomes formed by deposition of histones onto relaxed plasmid DNA [Torigoe et al. 2011], the conversion of prenucleosomes to canonical nucleosomes is dependent on the ACF motor protein. These experiments, along with the results in Supplemental Figure S3, A and B, further show that prenucleosomes formed with NAP1 or by salt dialysis can be converted into nucleosomes by ACF and that this process can occur with the 80-bp genomic DNA fragment or the central 80 bp of 601 DNA.

It is particularly notable that the ligation of free DNA to mono-prenucleosomes does not result in the wrapping of the DNA around the histones to give a canonical nucleosome. We hypothesize that the histone octamer is slightly unfolded or expanded in prenucleosomes relative to nucleosomes and that this alternate conformation of the histones in the prenucleosome does not enable facile wrapping of the DNA into a nucleosome. This postulated alternate conformation of the octamer in prenucleosomes could be due to charge repulsion between the histones that occurs in the absence of the extra histone–DNA contacts in canonical nucleosomes.

We also found that the Chd1 motor protein can be used in place of ACF for the conversion of ligated mono-prenucleosomes to nucleosomes [Supplemental Fig. S3C]. Hence, mono-prenucleosomes can be converted to nucleosomes in a variety of different conditions. Most importantly, these experiments reveal that mono-prenucleosomes are functionally active as precursors to canonical nucleosomes.

**MNase analysis indicates that prenucleosomes contain ~80 bp of DNA**

To complement the results from the psoralen cross-linking and electron microscopy experiments [Fig. 1], we used the mono-prenucleosome system to re-examine the length of DNA that is closely associated with prenucleosomes. For these experiments, we employed MNase as a
probe of histone–DNA interactions. In our studies of the ACF-mediated conversion of prenucleosomes to nucleosomes [Fig. 4], we observed that MNase digestion of prenucleosomes with ligated free DNA (in the absence of ACF) yields a set of bands of ~80 bp. We mapped the ends of these MNase digestion products by primer extension analysis and found that the majority of the DNA fragments ranged from ~78 to 85 bp in length and correlated with the central 80-bp 601 DNA that was used in the assembly of the mono-prenucleosomes (Fig. 5). The boundaries of MNase digestion of prenucleosomes are not as distinct as those seen with MNase digestion of canonical nucleosomes. Nevertheless, these results provide independent confirmation of the estimates of 70–80 bp of DNA per prenucleosome based on electron microscopy of prenucleosomes in relaxed plasmid DNA (Fig. 1). Collectively, these results indicate that prenucleosomes are closely associated with ~80 bp of DNA.

We also examined whether the DNA in a prenucleosome is accessible to restriction enzymes. To address this question, we assembled mono-prenucleosomes onto a variant of the 80-bp genomic DNA fragment that contains an EcoRV site (the center of the restriction site is 25/55 bp from the DNA ends) and an XhoI site (the center of the site is 46/34 bp from the ends). The addition of EcoRV or XhoI with mono-prenucleosomes versus naked DNA controls revealed that the packaging of DNA into mono-prenucleosomes blocks the access of the DNA to restriction enzymes [Supplemental Fig. S4], as seen with canonical nucleosomes. This restriction enzyme accessibility assay could be useful in the analysis of the properties of prenucleosomes.

The central region of the DNA in a mono-prenucleosome is located at approximately the same position as the analogous stretch of DNA in a core particle

We next sought to determine the location of the 80 bp of DNA in the mono-prenucleosome. To this end, we employed the histone-directed DNA cleavage method of Flaus et al. [1996] with the use of N-(1,10-phenanthroline-5-yl)iodoacetamide (OP) as the histone-modifying reagent [Brogaard et al. 2012; Henikoff et al. 2014]. In these experiments, we used wild-type core histones as well as histone octamers containing the H4S47C or H2BT87C mutant histones. In particular, it should be noted that H4S47C in a nucleosome is located near the dyad (Flaus et al. 1996). Alkylation of the cysteine sulfhydryl group by OP results in a covalent linkage between an o-phenanthroline moiety and the cysteine residue. In the presence of Cu(II)
and hydrogen peroxide, this o-phenanthroline group mediates the generation of hydroxyl radicals that cleave the nearby DNA. We thus alkylated wild-type and mutant [H4S47C or H2BT87C] histones with OP, reconstituted mono-prenucleosomes with the modified histones and the central 80 bp of 601 DNA, and then examined the cleavage of DNA upon addition of Cu(II) and hydrogen peroxide. These experiments revealed distinct cleavage sites near the middle of the 601 DNA sequence with the modified H4S47C histones but not with the wild-type or H2BT87C histones (Fig. 6). The DNA cleavage was also dependent on the addition of Cu(II). Strikingly, with the H4S47C histones, the sites of DNA cleavage of the central 80-bp 601 fragment in mono-prenucleosomes are identical to those seen in canonical nucleosomes with the full 147-bp 601 DNA fragment [Henikoff et al. 2014]; that is, the central 80 bp of 601 sequence in the mono-prenucleosome is located at the position analogous to the same DNA segment in a canonical nucleosome with the full 147-bp 601 DNA. We also observed a related DNA cleavage pattern of the 80-bp genomic DNA with the H4S47C histones [Supplemental Fig. S5], but the results were less distinct, possibly due to some heterogeneity in the interaction of the 80-bp genomic DNA to prenucleosomes relative to that of the well-positioned central 80 bp of 601 DNA. Hence, these findings indicate that the central region of the DNA in a mono-prenucleosome is located at approximately the same position as the analogous stretch of DNA in a core particle. It is also relevant to note that the H2BT87C residue, which is located at the opposite side of the dyad in a canonical nucleosome [for example, see Ferreira et al. 2007], was included as a probe in case the prenucleosomal DNA was located asymmetrically on the histone octamer. With its symmetric location, the prenucleosomal DNA would be cleaved by the OP-modified H2BT87C residue ~4–6 nt from the end (if the prenucleosome had the same histone–DNA contacts as a nucleosome) and yield small fragments that could not be clearly resolved. Thus, the absence of distinct cleavage sites with H2BT87C is not definitive but is consistent with the proposed central location of the DNA in the prenucleosome.

Figure 5. MNase digestion analysis reveals that prenucleosomes are associated with ~80 bp of DNA. Mono-prenucleosomes were reconstituted by NAP1-mediated deposition onto the central 80 bp of the 601 DNA (with two 5-nt overhangs) and ligated to two free 80-bp DNA fragments (each containing a single 5-nt overhang) to give mono-prenucleosomes that are flanked by 85-bp DNA extensions, as indicated in the diagram. The samples were digested with MNase, and the 5' ends of the resulting DNA fragments were mapped by primer extension analysis. The 5' ends of the primers corresponded to the ends of the central 80 bp of the 601 sequences; thus, the majority of the MNase-digested fragments ranged in size from ~78 to 85 bp. The control lanes show the ends of the unligated [single white dots] and ligated [double white dots] DNA fragments prior to MNase digestion.

p300 specifically acetylates H3K56 in prenucleosomes relative to nucleosomes

We further investigated the properties of prenucleosomes by subjecting prenucleosomes and nucleosomes to acetylation by purified p300 and then analyzing the resulting histones by mass spectrometry. Among the possible candidates for prenucleosome-specific acetylation by p300, H3K56 is the only amino acid residue that was found to exhibit this property [Fig. 7A]. Acetylation of H3K56 by p300 occurs with prenucleosomes but not with NAP1–histone complexes or nucleosomes. In addition, the acetylation at H3K56 was confirmed by the parallel analysis of H3K56A, with which acetylation was not detected.

In metazoans, H3K56 is acetylated by CBP/p300 proteins [Das et al. 2009]. H3K56 is located at the DNA entry and exit points of the nucleosome [for example, see Masumoto et al. 2005; Xu et al. 2005]. Hence, the greater accessibility of p300 to H3K56 in prenucleosomes relative to nucleosomes is consistent with the location of prenucleosomal DNA in the region that corresponds to the central 80 bp of the nucleosome (Fig. 6). H3K56ac has been found to be involved in chromatin assembly during DNA replication and repair [for example, see Masumoto et al. 2005; Han et al. 2007; Chen et al. 2008; Li et al. 2008]. In addition, genome-wide chromatin immunoprecipitation (ChIP) experiments in yeast, *Drosophila*, and human cells have shown that H3K56ac is highly enriched at active promoters as well as enhancers [for example, see Lo et al. 2011; Venkatesh et al. 2012; Skalska et al. 2015]. Moreover, in *Drosophila*, the increase in H3K56 acetylation at promoters and enhancers by Notch activation was found to occur rapidly by a mechanism that could not be clearly resolved. Thus, the absence of distinct cleavage sites with H2BT87C is not definitive but is consistent with the proposed central location of the DNA in the prenucleosome.
that requires CBP acetyltransferase activity but not transcriptional elongation (Skalska et al. 2015). These findings suggest that pre-existing H3K56 can be acetylated by CBP. Thus, the specific acetylation of H3K56 in prenucleosomes relative to nucleosomes provides another link between the properties of prenucleosomes and dynamic chromatin in cells. It is also possible that the presence of H3K56ac at promoters and enhancers may reflect the occurrence of prenucleosomes or prenucleosome-like particles. In the future, it should be interesting and informative to examine the relation between H3K56 acetylation and prenucleosome function in greater detail.

Figure 6. Mapping of the histone–DNA contacts in a mono-prenucleosome. Core histones containing the wild-type or the indicated mutant histones were modified with OP, which links an o-phenanthroline moiety onto the histones via alkylation of the thiol group on cysteine residues. The resulting derivatized histones were reconstituted by salt dialysis into mono-prenucleosomes with the central 80 bp of 601 DNA sequence that is 32P-labeled at the 5′ end. The hydroxyl radical cleavage reactions were initiated by the addition of Cu(II), hydrogen peroxide, and mercapto-propionic acid. The cleavage products were purified and analyzed by electrophoresis on a 10% polyacrylamide–urea gel. The Maxam-Gilbert G+A ladder was used to identify the OP cleavage products, which are indicated at the right side of the autoradiogram. In a canonical nucleosome, H4S47C is located near the dyad.

Figure 7. The prenucleosome, a conformational isomer of the nucleosome. (A) p300 specifically acetylates histone H3K56 in prenucleosomes relative to nucleosomes. Chromatin assembly reactions with ACF (Fyodorov and Kadonaga 2003; Torigoe et al. 2011) were performed with relaxed circular plasmid DNA in the presence of acetyl-CoA. ATP (or UTP as the −ATP control), DNA, and p300 were included as indicated. In addition, as a test for acetylation at H3K56, we performed parallel reactions with the mutant histone H3K56A, which cannot be acetylated at H3 residue 56. The resulting samples were then subjected to Western blot analysis with H3K56ac-specific antibodies (Millipore, catalog no. 07-677). As a reference, the blot was stripped and reprobed with anti-total H3 antibodies (Abcam, catalog no. AB1791). (B) Prenucleosomes comprise a core histone octamer and 80 bp of DNA at a location that is analogous to that of the central 80 bp of the core particle. H3K56 is accessible to p300 in a prenucleosome but not a nucleosome. Prenucleosomes can be converted into canonical nucleosomes by an ATP-driven motor protein such as ACF or Chd1. (C) Prenucleosomes or prenucleosome-related particles may be present in the upstream region of active promoters. (D) Model for the productive dynamic interconversion between prenucleosomes and nucleosomes. Prenucleosomes can be formed by the deposition of histones onto DNA and converted into nucleosomes by an ATP-driven motor protein such as ACF or Chd1. Nucleosomes can be disrupted by the action of enzymes such as polymerases as well as some ATP-driven chromatin remodeling factors. The resulting free histones are bound by the chaperones and then reassembled into prenucleosomes. It is not known whether a canonical nucleosome can be directly converted into a prenucleosome.
Discussion

The prenucleosome is a stable conformational isomer of the nucleosome

The prenucleosome was initially identified as a nonnucleosomal histone–DNA complex that is a precursor to the nucleosome in the assembly of chromatin in vitro (Torigoe et al. 2011). However, the ability to study prenucleosomes was limited by the heterogeneity of the prenucleosome samples that were assembled onto plasmid DNA templates. Hence, we lacked a fundamental understanding of the composition, structure, and organization of prenucleosomes.

In this study, we observed by psoralen cross-linking and electron microscopy that prenucleosomes appear to be associated with ~70–80 bp of DNA (Fig. 1). This finding led to the development of the mono-prenucleosome system that involves the assembly of mono-prenucleosomes from the four core histones and an 80-bp DNA fragment (Fig. 2). Prenucleosomes contain a histone octamer and are distinct from species such as tetrasomes or hexasomes that contain less than a complete octamer (Figs. 2, 3). Moreover, mono-tetrasomes can be converted into mono-prenucleosomes with no apparent accumulation of stable hexasome species (Fig. 2D). It is particularly notable that mono-prenucleosomes can be formed by the deposition of histones by NAP1 or dNLP as well as salt dialysis (Fig. 2C). Importantly, mono-prenucleosomes are functionally active, as they can be ligated to free [naked] DNA and then converted into canonical nucleosomes by a motor protein such as ACF or Chd1 (Fig. 4; Supplemental Fig. S3).

By using histone-directed DNA cleavage methodology (Flaus et al. 1996, Brogaard et al. 2012,Henikoff et al. 2014), we mapped the location of the 80 bp of DNA relative to the histone octamer in mono-prenucleosomes (Fig. 6; Supplemental Fig. S5). These experiments revealed that the central region of the 80 bp of DNA in a mono-prenucleosome is at the location analogous to the central region of the DNA near the dyad in a canonical nucleosome. We additionally mapped the ends of the DNA fragments generated by MNase digestion of prenucleosomes and found that the amount of DNA that is closely associated with prenucleosomes is ~80 bp [Fig. 5]. This is similar to the prenucleosomal DNA length that was independently estimated by psoralen cross-linking and electron microscopy (Fig. 1). The partial wrapping of DNA relative to that in a nucleosome is likely to be responsible for the lack of DNA supercoiling by prenucleosomes as well as the ability of p300 to acetylate H3K56 in prenucleosomes but not in canonical nucleosomes (Fig. 7A).

These findings indicate that the prenucleosome is a stable conformational isomer of the nucleosome (Fig. 7B). Moreover, no other histone–DNA particle was observed to be formed as efficiently and rapidly as prenucleosomes in the presence of the four core histones. Because there are probably only a small number of stable alternate conformations of the nucleosome, prenucleosomes may share a common fundamental structure with native nonnucleosomal particles such as those present at active chromatin throughout the genome.

As briefly discussed above, it is interesting to note that mono-prenucleosomes that are ligated to free DNA do not fold into canonical nucleosomes. The prenucleosomal histones appear to be in a different conformation than the histone octamer in a canonical nucleosome, possibly due to charge repulsion between the histones because of the reduced histone–DNA contacts in prenucleosomes relative to nucleosomes.

Based on the ability of prenucleosomes to be formed rapidly and then converted into canonical nucleosomes by a motor protein such as ACF or Chd1, we imagine that prenucleosomes are generated and assembled into nucleosomes during processes in which nucleosomes are disrupted, such as DNA replication, transcription, and repair. For instance, prenucleosomes have the same properties as histone–DNA complexes [in which H2B and H3 can be detected] at DNA replication forks that resemble nucleosomes but are formed much more rapidly than canonical nucleosomes [for example, see McKnight and Miller 1977; McKnight et al. 1978; Worcel et al. 1978]. In addition, some factors might be able to convert canonical nucleosomes directly into prenucleosomes, but such activities have not yet been identified.

Prenucleosomes appear to be related to nonnucleosomal histone–DNA complexes at active promoters

In our psoralen cross-linking and electron microscopy analysis of prenucleosomes versus nucleosomes, we observed a striking similarity between the distributions of psoralen bubble sizes with prenucleosomes versus nucleosomes compared with those obtained with active versus repressed promoters in vivo in yeast (Fig. 1C). These findings suggest that the “nucleosome-depleted regions” [NDRs; also termed “nucleosome-free regions” (NFRs)] that are located immediately upstream of the transcription start site of active genes contain prenucleosomes or prenucleosome-like particles (Fig. 7C).

In further support of this hypothesis, methidiumpropyl-EDTA sequencing [MPE-seq] analyses in mouse embryonic stem cells revealed that subnucleosome-sized chromatin fragments [including those containing 50–100 bp of DNA] are located specifically in the upstream promoter region of active genes [Ishii et al. 2015]. Notably, these noncanonical chromatin particles contain histones H2A and H3 by ChIP-seq [ChIP combined with deep sequencing] analysis. Moreover, the degree of enrichment of the subnucleosome-sized, histone-containing particles in the upstream promoter region correlates with the level of gene transcription, as assessed by RNA sequencing [RNA-seq] [Ishii et al. 2015]. Hence, these 50- to 100-bp-sized histone-containing particles that are present at active promoters have features that are similar to those of prenucleosomes.

In addition, the ability of prenucleosomes, but not nucleosomes, to be acetylated at H3K56 by p300 [Fig. 7A] may be responsible, at least in part, for the observed enrichment of H3K56ac at active promoters and enhancers [for example, see Lo et al. 2011; Venkatesh et al. 2012, Skalska et al. 2015]. p300 is associated with
transcriptional enhancers (for example, see Heinzelman et al. 2007, Visel et al. 2009), and it is possible that the enrichment of H3K56ac at enhancers is due to p300-mediated acetylation of prenucleosomes. In promoter regions, the averaged peak of H3K56 acetylation has been observed to flank the NDR at about –250 bp or about +250 bp relative to the +1 transcription start site. The absence of an H3K56ac peak precisely at the NDR could be due to the increased fragmentation of the sensitive DNA at the NDR (relative to the DNA in bulk chromatin) during the sonication of the chromatin. To clarify this issue, the generation of chromatin fragments for H3K56ac ChIP might optimally be performed with mild DNA cleaving reagents such as methidiumpropyl-EDTA-Fe(II) [MPE-Fe(II)] or low concentrations of MNase as in Ishii et al. (2015).

Prenucleosomes may also be related to “fragile” nucleosomes, which are MNase-sensitive nucleosomes that have been seen in yeast promoters (for example, see Weiner et al. 2010; Xi et al. 2011; Knight et al. 2014; Kubik et al. 2015). In addition, in HeLa [human] cells, salt-labile nucleosomes containing histones H2A.Z and H3.3 have been found at active promoters (Jin et al. 2009). Thus, the NDRs of active promoters appear to contain prenucleosomes or prenucleosome-related species (Fig. 7C).

Why might prenucleosomes or prenucleosome-like particles be present in the NDRs of active promoters? Because prenucleosomes interact with only ~80 bp of DNA, they would be more easily altered or disrupted than canonical nucleosomes. Also, if disrupted or displaced, prenucleosomes could be rapidly reassembled. In these respects, prenucleosomes appear to be compatible with the function of transcription factors. An alternate but related viewpoint is that prenucleosomes are intermediates in the dynamic process of nucleosome disassembly and reassembly at active promoters (Fig. 7D; see also Brown et al. 2013). It is even possible that prenucleosomes enhance transcription, such as in the establishment of the optimal structure of the active promoter. This notion is supported by the observation that the intensities of the H2B and H3 ChIP signals associated with the subnucleosome-sized DNA fragments correlate with transcriptional activity as measured by RNA-seq [Ishii et al. 2015]. In the future, it will be interesting and important to investigate the potential role of prenucleosomes or prenucleosome-like structures at active promoters.

Conclusion and perspective

Chromatin in the eukaryotic nucleus is multidimensional. There are covalent modifications of the histones, histone variants, ATP-driven chromatin remodeling factors, nonhistone chromosomal proteins, and nonnucleosomal chromatin particles. Notably, each of these dimensions of chromatin affects gene expression. The prenucleosome is the only known stable conformer of the nucleosome and the only distinct nonnucleosomal histone–DNA particle that has been observed to be rapidly and efficiently formed on DNA in the presence of the four core histones, as in the nucleus. (For instance, as seen in Figure 2, C and D, H3–H4 tetrasomes are inefficiently formed and rapidly converted into prenucleosomes in the presence of H2A–H2B.) It thus seems likely that many nonnucleosomal particles in the cell are prenucleosomes or prenucleosome-related particles. An additional attractive feature of this hypothesis is the ability of prenucleosomes to be converted into nucleosomes by ATP-dependent motor proteins such as ACF or Chd1. Hence, in this model, there is a productive dynamic interconversion between prenucleosomes and nucleosomes (Fig. 7D). It is our hope that, in the future, the new knowledge of nonnucleosomal components of chromatin will contribute to an integrated understanding of the dynamic structure and function of our genome.

Materials and methods

Reagents and methodology

DNA oligonucleotides were synthesized by Integrated DNA Technologies. The sequences of the oligonucleotides are in Supplemental Table S1. D. melanogaster NAP1 and ACF complexes were synthesized in S9 cells by using baculovirus expression vectors and purified as described previously [Fyodorov and Kadonaga 2003]. D. melanogaster dNLP was synthesized in Escherichia coli and purified by the method of Ito et al. [1996]. D. melanogaster Chd1 was synthesized in E. coli and purified as described [Torigoe et al. 2013]. Native core histones were purified from D. melanogaster embryos that were collected from 0 to 12 h after egg deposition [Fyodorov and Levenstein 2002]. Recombinant D. melanogaster histones were synthesized in E. coli and purified by the method of Luger et al. [1999]. Human p300 protein was purified and used as described by Kraus and Kadonaga [1998]. All experiments were performed independently at least twice to ensure the reproducibility of the data.

Reconstitution of mono-prenucleosomes

Mono-prenucleosomes were prepared by either histone chaperone-mediated deposition or salt dialysis. In chaperone-mediated reconstitution reactions, the core histones were incubated with either NAP1 [at a 5:1 mass ratio of NAP1:core histones] or dNLP [at a 10:1 mass ratio of dNLP:core histones] for 20 min on ice in a volume of 10 µL in the following buffer: 10 mM K-HEPES [pH 7.6], 0.1 M KCl, 0.1 mM EDTA, 1 mM DTT, and 70 µg/mL bovine serum albumin. Eight microliters of HEG buffer [25 mM K-HEPES at pH 7.6, 0.1 M KCl, 0.1 mM EDTA, 10% [v/v] glycerol] was then added. Next, 2 µL of the preannealed DNA oligonucleotides [in 10 mM Tris-HCl at pH 7.5, 0.1 mM EDTA, 0.05 M NaCl] was added to the histone chaperone mixture to give a final volume of 20 µL. The samples were mixed immediately by gentle vortexing, incubated for 30 sec, and analyzed by native 5% polyacrylamide gel electrophoresis.

Salt dialysis reconstitution of mononucleosomes, mono-prenucleosomes, and mono-tetrasomes was performed by the method of Stein [1989]. In a standard reaction, the DNA fragment [as indicated for each experiment] (50 pmol, 6 µL in TE buffer: 10 mM Tris-HCl at pH 7.5, 1 mM EDTA) was combined with 140 µL of TE buffer containing 1.07 M NaCl and 0.011% [v/v] NP-40, and core histones [50 pmol, 4 µL in 10 mM K-HEPES at pH 7.6, 0.1 M KCl, 1 mM DTT, 10% [v/v] glycerol] were added to give final concentrations of 1.0 M NaCl and 0.01% [v/v] NP-40 in a total volume of 150 µL. The resulting histone–DNA mixture was
subjected to dialysis at room temperature in a ThermoFisher Slide-A-Lyzer dialysis cassette [molecular weight cutoff of 3500 Da] for 2 h against TE containing 0.8 M NaCl for 1.5 h against TE containing 0.6 M NaCl and 2 h against TE containing 0.05 M NaCl. The resulting products were analyzed by native 5% polyacrylamide gel electrophoresis. Mono-prenucleosomes that were reconstituted by salt dialysis were stable for at least a few weeks at 4°C. For storage, prenucleosomes were dialyzed overnight at 4°C against histone storage buffer (10 mM K-HEPES at pH 7.6, 0.1 mM EDTA, 0.1 M KCl, 10% [v/v] glycerol, 1 mM DTT).

**Ligation of mono-prenucleosomes to free DNA and assembly into canonical nucleosomes by ACB**

Mono-prenucleosomes were ligated to free (naked) DNA by using methods similar to those employed for the ligation of nucleosomes to free DNA (Clark and Felsenfeld 1992; Stein et al. 2002). Typically, mono-prenucleosomes (85 pmol) of DNA with sticky ends) and the flanking DNA strands (85 pmol each, and each with a single sticky end that is complementary to one end of the prenucleosomal DNA) were combined in 10 mM K-HEPES (pH 8.0), 1 mM EGTA, 1 mM DTT, 0.5 mM ATP, an ATP regeneration system [3 mM phosphoenolpyruvate, 20 U/µL pyruvate kinase], 2.5 mM MgCl₂, 12.5 µg of bovine serum albumin, and 3000 U of T4 DNA ligase (New England Biolabs) in a volume of 138 µL. HEG (82 µL) was added to give a final volume of 220 µL, and the ligation was carried out overnight at 16°C. In the assembly of prenucleosomes into nucleosomes, the ligated prenucleosomes (20 pmol, 52 µL directly from the ligation reaction) were combined with a solution of 20 mM ATP and 5 µL of 33 mM MgCl₂, 1.0 µL of 700 nM ACB (to a final ACB concentration of 6 nM), and 17 µL of HEG buffer to a final volume of 70 µL. The reaction was carried out for 1.5 h at 27°C. The samples were analyzed by MNase digestion as described by Torigoe et al. (2011).

**Trimethylpsoralen cross-linking**

Cross-linking was performed essentially as described [Sogo and Tomha 1989; Brown et al. 2015]. The specific conditions were as follows: A 175-µL sample of chromatin assembly reaction (Fyodorov and Kadonaga 2003; Torigoe et al. 2011) was diluted with 125 µL of dilution buffer [15 mM K-HEPES at pH 7.6, 100 mM KCI, 5 mM MgCl₂, 0.1 mM EDTA, 6.6% [v/v] glycerol, 1% [w/v] polyvinyl alcohol [average molecular weight 10,000], 1% [w/v] polyethylene glycol 8000, 20 µg/mL bovine serum albumin] and transferred to a single well of a 24-well plate. The 24-well plate containing all samples was placed on an ice-water slurry and positioned 5 cm away from five 366-nm 15 W UV bulbs in a well plate containing all samples was placed on an ice-water slurry and positioned 5 cm away from five 366-nm 15 W UV bulbs and transferred to a single well of a 24-well plate. The 24-well plate containing all samples was placed on an ice-water slurry and positioned 5 cm away from five 366-nm 15 W UV bulbs and transferred to a single well of a 24-well plate.

**Sample preparation for electron microscopy**

DNA [1–3 µL in TEN] was denatured in 70% [v/v] deionized formamide (Sigma) and 0.5 M glyoxal (Sigma) in a total volume of 13 µL for 30 min at 37°C. Samples were immediately placed on ice and diluted with 5 µL of TEN. Benzalkonium chloride (Sigma) was added to 0.001% [w/v] to facilitate spreading of the DNA. A 10-cm petri dish with a mica ramp resting on the rim was filled with distilled water. Graphite powder was lightly dusted on the surface of the water. A portion (5 µL) of the denatured DNA sample was run down the mica ramp and spread on top of the water, which pushed back the graphite dusting. DNA near the mica border or mica ramp was picked up by using carbon-coated copper grids (Electron Microscopy Sciences) pretreated with 30 µg/mL ethidium bromide. Grids were stained with 0.5 mM uranyl acetate, washed with 100% ethanol, and air-dried. Following staining, they were secured to a rotary tilt stage in a 208C high-vacuum turbo carbon coater equipped with a metal evaporation accessory (Cressington) and shadowed at an angle of 3° with platinum-palladium (80:20). Electron Microscopy Sciences) until a thickness monitor (Cressington, MTM-10) reported 100 nm of metal deposition on the sensor.

**Electron microscopy**

Images were taken on a JEOL 1230 electron microscope at 120 keV at 20,000x magnification and were processed and analyzed in ImageJ. Chromatin assembly reactions were performed four separate times, and totals of 380 and 376 molecules were analyzed from chromatin assembly reactions without and with the addition of ACB, respectively.

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