A nucleosome-free region locally abrogates histone H1–dependent restriction of linker DNA accessibility in chromatin

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Eukaryotic genomes are packaged into linker–oligonucleosome assemblies, providing compaction of genomic DNA and contributing to gene regulation and genome integrity. To define minimal requirements for initial steps in the transition of compact, closed chromatin to a transcriptionally active, open state, we developed a model in vitro system containing a single, unique, “target” nucleosome in the center of a 25-nucleosome array and evaluated the accessibility of the linker DNA adjacent to this target nucleosome. We found that condensation of H1-lacking chromatin results in ~60-fold reduction in linker DNA accessibility and that mimics of acetylation within all four core histone tail domains of the target nucleosome synergize to increase accessibility ~3-fold. Notably, stoichiometric binding of histone H1 caused >2 orders of magnitude reduction in accessibility that was marginally diminished by histone acetylation mimics. Remarkably, a nucleosome-free region (NFR) in place of the target nucleosome completely abrogated H1-dependent restriction of linker accessibility in the immediate vicinity of the NFR. Our results suggest that linker DNA is as inaccessible as DNA within the nucleosome core in fully condensed, H1-containing chromatin. They further imply that an unrecognized function of NFRs in gene promoter regions is to locally abrogate the severe restriction of linker DNA accessibility imposed by H1s.

The genomes of eukaryotes are packaged into chromosome-sized oligonucleosome assemblies, which provide orderly compaction of genomic DNA and contribute to the regulation of transcription and other processes. These functions are due in part to the propensity of oligonucleosomes (referred to as nucleosome arrays) to spontaneous fold and condense into higher-order chromatin structures within the ionic environment of the nucleus (1—3). Such condensation requires inter-nucleosome interactions mediated through the core histone tail domains (4—8). Importantly, reconstituted arrays containing only core histones and defined DNA templates recapitulate the behavior of native chromatin with regard to salt-dependent folding and compaction (1, 3, 9). Oligonucleosome arrays lacking H1 exist as extended “beads on a string” structures (defined as primary chromatin structure (10)) in low ionic strength buffers in vitro but equilibrate between moderately compacted and fully folded secondary structures such as the chromatin fiber in 0.5–2 mM MgCl₂. In slightly higher concentrations of divalent salts (>2–3 mM MgCl₂), the arrays oligomerize (self-associate) into large globular tertiary structures that resemble the size, shape, and appearance of interphase chromosomes (3). Arrays containing H1 undergo similar transitions but at lower concentrations of multivalent or monovalent salts (1, 11). Therefore, the core histones are sufficient to direct formation of chromatin fibers and higher-order structures in vitro (1, 12).

Nucleosomes impose a severe restriction to the accessibility of DNA that is exploited by gene regulatory mechanisms. For example, pioneering experiments showed that even in the absence of activating conditions or upstream activating sequence elements appropriate repression of several yeast genes requires nucleosomes (13–15). In vitro experiments have demonstrated that the accessibility of trans-acting factors to DNA within the ~147-bp nucleosome core region is restricted 10³–10⁵-fold, depending on proximity to the nucleosome dyad, an extent of inhibition sufficient to elicit a regulatory effect (16, 17). The core histones compete with transcription factors for binding to cognate sites, thereby providing a thermodynamic barrier to binding that must be overcome by nucleosome remodeling, increased factor concentration, or increased binding activity (18).

In contrast, linker DNA, which extends between successive nucleosome core regions in nucleosome arrays, is not in tight association with the core histones and is nearly as accessible as naked DNA in mononucleosomes (19, 20). However, within condensed nucleosome arrays lacking linker histones, accessibility to linker DNA is reduced compared with naked DNA about 15–50-fold, consistent with a role of condensed higher-order chromatins in restricting access to cognate sites regardless of their position in chromatin (16, 21). In addition, linker histones (H1s) are present in similar abundance to the core histone, bind to the exterior of the nucleosome, and neutralize the charge of linker DNA to stabilize the salt-dependent folding of nucleosome arrays (22, 23). Despite the occurrence of a single canonical H1-binding site per nucleosome, a range of H1:nucleosome ratios has been observed in cells with significantly less than one H1 per nucleosome in more transcriptionally active and pluripotent cell types to ≥1 in quiescent cell types, suggesting a general role of H1s in gene repression (24, 25). However, although H1 drastically inhibits transcription of chromatin templates in vitro (26–28), H1 knockdowns and

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knockouts in vivo do not result in genome-wide increases in transcription, consistent with additional overlapping mechanisms of gene repression (29–31). Moreover, stable reduction or elimination of H1 results in reduced nucleosome spacing, which may also contribute to repression (30, 31). However, the precise effect of H1 on linker DNA accessibility in condensed chromatin and the extent to which histone acetylation or other chromatin modifications mitigate these effects have not been precisely quantified.

The activation of a silent gene locus is thought to involve initial binding of a transcription factor(s) that initiates chromatin remodeling events such as histone acetylation and displacement of H1s to open chromatin (32). This transition typically involves recruitment of histone acetyltransferases to target nucleosomes to direct local histone acetylation (32, 33). Importantly, acetylation of the core histone tail domains directly destabilizes higher-order condensed chromatin structures (11, 34, 35). In addition, active promoters have a nucleosome-free region (NFR)3 immediately upstream of the transcription start site (TSS), generated by a combination of the binding of general transcription factors, ATP-dependent nucleosome remodeling, and, in some cases, DNA sequences that exhibit reduced affinity for binding core histones (15, 36–41). The NFR and associated activities serve to order nucleosomes upstream and downstream of the TSS with the regularity of spacing decaying with distance from the TSS (37, 42, 43). Although the NFR allows space for binding of the preinitiation complex, the lack of a canonical nucleosome may disrupt stable folding/condensation of higher-order chromatin structure and therefore contribute to promoter accessibility (21, 44).

To better understand the effects of factors involved in locus activation on chromatin accessibility, we developed an in vitro model system wherein a single “designer” nucleosome is located in the center of an otherwise unmodified 25-nucleosome array (21). Independent reconstitution of the central target nucleosome allows installation of any single or combination of histone modifications or other alteration associated with locus activation. In initial experiments with this system, we showed that acetylation mimics within the histone H4 tail domain of the target nucleosome increased accessibility of the surrounding linker DNA about 2-fold, whereas acetylation mimics within the H3 tail had little effect but were able to synergize with H4 tail acetylation mimics to further increase accessibility. Here, we assess the effects of acetylation mimics in histones H2A and H2B, linker histones, and an NFR on linker DNA accessibility. We found that acetylation mimics within all four core histones increase accessibility beyond that when only H3 and H4 contain acetylation mimics. Importantly, we found that stoichiometric association of H1 with nucleosomes in the array results in a drastic (>100-fold) reduction in accessibility that is minimally abrogated by acetylation. However, installation of a NFR completely abolished the effect of H1.

Results

To investigate the effects of chromatin remodeling events that initiate opening closed chromatin, we generated 25-nucleosome arrays in which a single “target” nucleosome (T_{Nuc}) was ligated between two 12-mer nucleosome arrays (Fig. 1A). In previous work, the array templates contained tandem repeats of the Lytechinus 5S nucleosome positioning sequence (21, 45). However, nucleosome positioning on these arrays is degenerate (46). Although the 5S arrays recapitulate the somewhat loose positioning found in some regions of native chromatin, arrays based on the 601 nucleosome positioning sequence exhibit precisely positioned nucleosomes and have slightly altered salt-dependent folding properties in vitro compared with 5S arrays (5, 47). Indeed, the latter might better model selected regions of native chromatin in which nucleosome positioning is highly ordered.

To determine whether the precision of nucleosome positioning significantly affects accessibility in condensed chromatin, we compared accessibility within arrays based on the 601 sequence with prior results with 5S arrays (21). We first assessed the impediment to accessibility in maximally condensed nucleosome arrays containing unmodified core histones in 10 mM MgCl₂ (high-Mg2⁺) buffer. In such buffers, the 25-mer nucleosome arrays spontaneously condense and self-associate into large aggregates resembling the size and density of individual interphase chromosomes (1, 3). Indeed, >90% of the arrays are insoluble in 3 mM MgCl₂ buffer, indicating complete saturation of the template with nucleosomes (Fig. 1B and Ref. 1). A 4.5-kb naked DNA fragment containing a single DraIII site was included in every reaction as an internal control, allowing normalization for variations in enzyme activity (21). Digestion was initiated by addition of DraIII, and the ratio of the rates of disappearance of 25-mer DNA and 4.5-kb DNA bands (k relative (k_{rel})) was determined from plots of digestion kinetics (Fig. 1, C and D) with correction for inherent target-specific differences in cleavage rates (see “Experimental procedures”). Plots showed that the vast majority of both the DNA and chromatin digests contained a single kinetic component (Fig. 1D). Our analysis revealed that DraIII sites in the T_{Nuc} linker DNA were about 60-fold less accessible in the condensed chromatin compared with the naked DNA template (k_{rel} = 0.015 ± 0.001) (Table 1), commensurate with previous measurements for 601 arrays (16). Thus, the T_{Nuc} linker DNA is about 5-fold less accessible in the 601 arrays compared with 5S arrays (21), suggesting that precise nucleosome spacing generates a more stable and less accessible condensed chromatin structure. However, similar to the 5S nucleosome arrays (21), acetylation mimics within the H3 and H4 tail domains of the target nucleosome locally increased DNA accessibility about 1.5-fold within the 601-based arrays (Table 1).

In addition to the H3 and H4 tail domains, the N-terminal tail domains of H2A/H2B dimers also contribute to the stability of nucleosome packing in higher-order chromatin structures and are acetylated in association with gene expression (34, 48, 49). To determine whether acetylation of the H2A and H2B N-terminal tails in a single nucleosome can open chromatin structure alone or in combination with H3/H4 acetylation, we assembled 25-mer arrays in which the T_{Nuc} contained unmodified histones (WT), acetylation mimics (ac⁰⁰) only within H2A/H2B (H2Aac⁰⁰/H2Bac⁰⁰), or mimics within all four core histones (All-ac⁰⁰) (Fig. 2A). Comparison of k_{rel} values indicated that acetylation mimics within H2A/H2B resulted in an ~1.4-

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3The abbreviations used are: NFR, nucleosome-free region; TSS, transcription start site; T_{Nuc}, target nucleosome; k_{rel}, k relative; ac⁰⁰, acetylation mimics.
fold increase in the accessibility of T_{Nuc} linker DNA, comparable with that observed for H3acm/H4acm (Fig. 2B and Table 1). Importantly, installation of acetylation mimics in all four core histones resulted in significantly more enhancement in linker DNA accessibility (k_{rel}/H11006 = 0.10) than the predicted combined effects of the individual sets of mimics (1.5/0.21 = 0.1; p < 0.001), indicating that acetylation mimics within the H2A/H2B dimers and the H3/H4 tetramer of a single nucleosome cooperatively increase accessibility of the adjoining linker DNA.

Linker histones (H1s) bind to the exterior of nucleosomes and stabilize higher-order chromatin structures (50, 51). Thus, linker histones restrict the accessibility of DNA to trans-acting factors, and many (or all) H1 subtypes, including H1.0, are depleted in the vicinity of active promoters (52–57). However, quantitative determinations of the effect of linker histones on DNA accessibility in chromatin are lacking, hampered by the difficulty in precisely controlling the stoichiometry of H1 binding, as slight excesses of H1 can cause aggregation of chromatin samples, whereas undersaturation can lead to drastically different results compared with saturated conditions. To overcome these issues, we used Nap1 as an H1 chaperone to precisely control H1 deposition (58, 59). Incubation of increasing amounts of preformed H1–Nap1 complex results in a small but detectable reduction in the electrophoretic migration of 12-mer nucleosome arrays until a point where a uniquely migrating species is apparent over a range of H1–Nap1 concentrations expected to deposit one H1 per nucleosome (Fig. 3A, white line). Importantly, the H1-bound arrays are completely soluble in 0.5 mM MgCl$_2$ (see “Experimental procedures”), indicating that the arrays are not overloaded with H1 (Fig. 3B).

We next determined the extent to which H1 restricts access to the T_{Nuc} linker DNA. Note that in these experiments we directly compared rates of digestion as calculation of k_{rel} based on inclusion of a naked DNA control was not possible because linker histones bind cooperatively to naked DNA with affinities...
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![Diagram](Image)

Figure 2. Acetylation mimics in all core histone tail domains increase accessibility of central target nucleosome linker DNA within a condensed 25-mer array. A, location of Lys → Gln substitutions as acetylation mimics (acm) in each of the core histone N-terminal tail domains (blue lines). B, representative SDS-agarose gels showing the DraiIII digestion time course for 25-mer arrays with TNuc, containing the indicated modified histones. Digestions were carried out for the times indicated above the lanes, and then products were separated on 0.7% SDS-agarose gels and stained with ethidium bromide. Gels were quantified, and rates of digestions were determined as described under “Experimental procedures.” The relative rate of digestion for each construct, normalized to the WT TNuc, is shown below the gel. See also Table 1. Bands corresponding to the 5.3-kb 25-mer array template (array) and the 4.5-kb naked control DNA (DNA) are indicated.

only a few-fold less than to nucleosomes (60, 61). We first determined the effect of H1 binding to arrays in buffer containing 0.5 mM MgCl₂ where the arrays are only partially condensed and not self-associated (21). Remarkably, we found that stoichiometric association of H1 to arrays (one per nucleosome) resulted in an ∼50-fold reduction in accessibility of the WT T₅₄ucleosomal linker DNA (Fig. 3, C and E, and Table 2). Moreover, acetylation mimics within all four histones in the TNuc (All-acm) had only a small effect on the H1-dependent reduction in accessibility (Fig. 3, D and E, and Table 2). Importantly, titration of H1 into both WT and All-acm nucleosome arrays yielded a reduction in T₅₄ nucleosome linker DNA accessibility, as measured by DraIII digestion, until the same apparent point of saturation as indicated by the gel-shift binding assay (Fig. 3, compare F with A), further indicating that the arrays in the above experiments are saturated with H1 over a range of H1–Nap1 inputs. Moreover, substoichiometric ratios of H1:nucleosomes, such as are found in many cell types, result in significantly less restriction in access to linker DNA (see “Discussion”).

We next assessed the effect of H1 association on linker DNA accessibility when the arrays are fully condensed and self-associated (21). Remarkably, we found that binding of H1 to arrays in 10 mM MgCl₂ resulted in an even more drastic, ∼200-fold reduction in linker DNA accessibility compared with that observed in the low-Mg²⁺ buffer (Fig. 4, A and B, and Table 2). Given that accessibility of the linker DNA is already ∼60-fold inhibited in such buffers in the absence of H1, these results suggest that the restriction of linker DNA accessibility approaches that found within the nucleosome core region (see “Discussion”).

Active genes in metazoans contain an NFR upstream of the transcription start site that generates a gap in the ordered array of nucleosomes and leads to a 3–4-fold increase in linker DNA accessibility in H1-lacking nucleosome arrays (21). Because an NFR also results in the loss of a canonical H1-binding site on the nucleosome, we determined the effect of replacing the target nucleosome with an NFR on observed restriction of linker DNA accessibility by H1. Remarkably, in stark contrast to the effect of acetylation mimics, installation of a single NFR completely reversed the effect of H1 on DraIII accessibility for both partially and maximally condensed arrays (Fig. 4, C and D, and Table 2). Indeed, equivalent amounts of DraIII can be used to achieve the same extent of digestion in NFR arrays in the presence or absence of H1, in contrast to WT arrays. To determine whether the drastic effect of the NFR on H1 repression extended throughout the entire 25-mer arrays, we digested arrays with EcoRV, which cuts between every nucleosome in the flanking 12-mer arrays except for the T₅₄−linker DNA (Fig. 5A). We found that the effect of the NFR is largely localized to the vicinity of the TNuc as the remainder of the array exhibits roughly equivalent, inhibited rates of digestion with EcoRV in the presence of H1, regardless of whether the central position was a WT nucleosome or an NFR (Fig. 5B). These results indicate that H1 imposes a dominant constraint on DNA accessibility in native chromatin and suggests that the NFR, which lacks a canonical H1-binding site, locally counteracts H1-mediated condensation. These results are consistent with whole-genome mapping studies, which show that H1 is widely distributed across genomes except for in the immediate vicinity of NFRs of active genes (54–57).

Discussion

Our data show that chromatin condensation can drastically affect linker DNA accessibility in model nucleosome arrays. We observed a 60-fold reduction in linker DNA accessibility in condensed chromatin lacking H1 compared with naked DNA. Moreover, we found an additional 50–200-fold reduction in accessibility in an H1-containing over H1-lacking chromatin, depending on whether the chromatin was moderately or fully condensed, respectively. Interestingly, acetylation had a modest 2–4-fold effect in both the presence and absence of H1. Therefore, our results indicate that displacement of H1 either locally or globally will result in a much larger effect than acetylation on accessibility and binding of transcription factors and associated gene activation (Fig. 6).

Our results indicate that acetylation of all four core histone N-terminal tail domains of a single nucleosome contributes to opening chromatin. Indeed, acetylation mimics within either H3/H4 or H2A/H2B tails resulted in a modest but significant increase in accessibility, whereas mimics within all four tail domains in a single nucleosome cooperate to increase accessibility of linker DNA. These results indicate that acetylation within a single nucleosome brought about by an activating pioneer factor is sufficient to initiate a transition to an open, active chromatin structure (Fig. 6).
Previous work showed that installation of a single NFR results in a 3–4-fold increase in accessibility in fully condensed H1-lacking arrays (21). In contrast, we found that the NFR completely abrogates the large repressive effect of H1, resulting in 50- and 200-fold increases in accessibility in partially and fully condensed arrays, respectively. Our results suggest that the effect of the NFR is primarily local as little effect on H1-independent reduction in accessibility was observed when arrays were digested with EcoRV, which cuts between the nucleosomes in the regions of the nucleosome array flanking the T_{Nuc} (Fig. 5).

Importantly, whole-genome ChIP studies indicate that many H1 subtypes, including H1.0, the subtype used in the current studies, are drastically depleted in the immediate vicinity of the NFR of active genes but only modestly reduced upstream and downstream of the NFR (54–56), consistent with the idea that the lack of a single nucleosome creates a discontinuity in the H1-stabilized condensed chromatin structure (Fig. 6). Although it is difficult to determine from these data whether the reduction of H1 in the promoter region is due to the NFR or transcription factors that bind to the DNA, our data indicate that the lack of a single nucleosome and canonical H1-binding site plays a primary role in counteracting H1-dependent repression. (Note that although H1 may bind to the NFR, binding occurs with a lower affinity to naked DNA than to nucleosomes and is not expected to have an equivalent chromatin-condensing effect.) Thus, our data suggest that recruitment of chromatin remodeling factors that ultimately evict histones contributes a...
much larger enhancement to promoter accessibility than acetylation of the core histones per se, although acetylation may be required for recruitment or efficient function of such activities.

Many cell types contain less than one linker histone per nucleosome. For example, mouse embryonic stem cells contain about 0.5 H1s per nucleosome, which appears to contribute to maintaining a transcriptionally permissive environment in these pluripotent cells (25, 30, 62). Moreover, an increase in the H1:nucleosome ratio is observed in embryonic stem cell differentiation along with chromatin condensation (25). Interestingly, our data show that chromatin containing less than one H1 per nucleosome is much more accessible than when saturated with H1. Indeed, a reduction in H1 stoichiometry approximately from one to 0.5 per nucleosome resulted in an ~10-fold increase in accessibility of the T_{Nuc} linker DNA (Fig. 3F). These results provide strong support to the idea that a similarly reduced H1 content will result in a much more transcriptionally permissive chromatin environment.

Even in the absence of chromatin folding, nucleosomes impart a $10^3$–$10^5$-fold restriction to DNA sites within the ~147-bp nucleosome core region, depending on location with respect to the nucleosome dyad. Thus, the vast majority of cognate sites within the nucleosome core would remain unoccupied by trans-acting factors at concentrations typically found

Figure 4. The NFR overrides the H1-dependent decrease in linker DNA accessibility. A, digests of nucleosome arrays containing WT T_{Nuc} in 10 mM MgCl₂ in the absence and presence of H1. Units of DraIII used in each digestion are indicated above the gels. B, plot of data taken from gel shown in A. C and D, as in A and B except arrays contained an NFR in place of the T_{Nuc}.

Figure 5. The effect of the NFR on H1-dependent restriction of linker DNA accessibility does not extend to the entire array. A, schematic showing the 25-mer array, EcoRV sites (vertical arrows) and the T_{Nuc} (center; shaded). B, 25-mer arrays containing either a WT nucleosome or an NFR at the central (T_{Nuc}) position were incubated with or without H1 in buffer containing 10 mM MgCl₂ as indicated and digested with EcoRV (2 units) for the indicated times, and then products were separated on SDS-agarose gels.
within the nucleus (16). (Exceptions include pioneer transcription factors that have the ability to bind to the nucleosome surface (32, 33). However, whether chromatin condensation reduces binding of pioneer factors remains an open question.) Interestingly, compaction of nucleosome arrays causes only a modest 3–8-fold additional restriction on accessibility within the nucleosome core region (16). However, we found that although the linker DNA separating core regions remains relatively accessible in expanded chromatin containing only the core histones, complete condensation of the chromatin imparts an −60-fold impediment on linker DNA accessibility in the absence of H1 (Table 1 and see also Ref. 21), consistent with prior published work (16). Moreover, binding of H1 causes an additional 50–200-fold reduction in accessibility, resulting in an overall restriction in linker DNA accessibility approaching that found in the nucleosome core region (Figs. 3 and 4). These results suggest that linker DNA in a fully condensed H1-bound chromatin is not appreciably occupied by sequence-specific DNA-binding factors responsible for driving gene activation, similar to that in the nucleosome core region (16). The large dynamic range of this effect suggests that linker DNA accessibility represents a potential H1-dependent switch within chromatin.

Our data indicate that precisely spaced 601 nucleosome arrays form a more tightly condensed chromatin structure than the 5S arrays (21), which have a more heterogeneous nucleosome spacing. We observed a $k_{\text{rel}}$ of 0.015 ± 0.001 for the WT 601 arrays, about 5-fold less than the $k_{\text{rel}}$ observed for the 5S arrays (21) (Table 1). However, we note that the inherent rate of digestion of the ligated 601 array DNA template is 2-fold greater than that of the 5S templates (see “Experimental procedures” and Ref. 21). These differences are likely due to sequence-dependent DNA structure, which results in modest effects in the rate of two-dimensional diffusion and cognate site recognition by the restriction enzyme. It is possible that these effects are not relevant when the templates are assembled into nucleosomes; however, even if corrections for DNA sequence effects are ignored, the 601 arrays are ≈3-fold less accessible than the 5S arrays ($k_{\text{rel}}$ of 0.17 ± 0.02 versus 0.06 ± 0.01, respectively), indicating that arrays with more regularly spaced nucleosomes form a more stable and compact structure. These results are supported by EM of long 601 arrays (35) and folding studies (63) and suggest that more regular spacing of nucleosomes in vivo results in a more regular and stable condensed chromatin structure.

We found that the accessibility of linker DNA is severely restricted by stoichiometric binding of linker histones. The extent of this restriction likely depends on the linker H1 subtypes as they vary in their degree of condensation, length of DNA, and potentially the type of enzymes (length of recognition sequence) being used. It will be interesting in the future to test subtypes that have a range of chromatin-condensing capabilities (57).

**Experimental procedures**

**Expression and purification of core and linker histone proteins**

WT and mutant recombinant *Xenopus* core histones were expressed and purified as described previously (21). Briefly, coding sequences for *Xenopus* core histones H2A, H2B, H3, and H4 containing specific lysine-to-glutamine substitutions were obtained using the Stratagene QuickChange site-directed mutagenesis kit. Mutants were confirmed by Sanger sequencing. *Escherichia coli* BL21(DE3) cells were transformed with the pET3a expression plasmids harboring the WT or mutant core histone gene sequences. Cultures (3 ml of LB) containing ampicillin (100 μg/ml) were inoculated from single colonies and grown overnight, then 1 ml was used to inoculate 100 ml of LB, and the cultures were allowed to grow at 37 °C in a shaker incubator until 0.6 $A_{600}$. Expression of histones was induced by addition of 0.4 mM isopropyl β-D-galactopyranoside for 3 h at 37 °C. Cells were harvested, lysed, and sonicated, and H3/H4 tetramers and H2A/H2B dimers were purified as described (64). *Xenopus* H1.0 protein was expressed and purified as described (65). Briefly, *E. coli* BL21(DE3) cells transformed with the plasmid pET3aH1(0) were grown in 100 ml of LB containing ampicillin (50 μg/ml) until 0.6 $A_{600}$ expression was induced as described above, cells were harvested, and H1 was purified as described (65). Protein concentrations were determined by quantitative analysis of standards and samples run on 15% SDS-polyacrylamide gels stained with Coomassie Blue. The concentrations of standards were determined by amino acid analyses (66). Proteins were stored in a −80 °C freezer.

**Preparation of DNA templates**

A 204-bp DNA fragment containing the 601 nucleosome positioning sequence and asymmetric DraIII overhangs (DC template) was prepared as described (21). A complementary array DNA template containing 12 tandem repeats of a 207-bp 601 nucleosome positioning element was created by conversion of the Xbal site in the plasmid pSub601-12 to an asymmetric DraIII site to create plasmid PD6. A 2.5-kb DNA fragment containing the array template was generated by digestion of 200 μg of PD6 plasmid with 200 units of HindIII-HF (High Fidelity) and DraI enzymes in CutSmart buffer (New England Biolabs) in a 300-μl reaction for 6 h. The digested plasmid was incubated with 50 units of calf intestinal alkaline phosphatase at 37 °C for 30 min, extracted with phenol/chloroform, ethanol-precipitated, dried, resuspended in 200 μl of TE buffer (10 mM Tris-HCl, pH 8.0, 1 mM EDTA), and digested with 200 units of DraIII-HF at 37 °C for 2 h. The 2.5-kb band was isolated on a preparative 0.7% agarose gel, electroduted, precipitated, and resuspended in TE buffer. Note that the 2.5-kb 12 × 207 array template has a DraIII (3') overhang complementary to the DC template overhangs at one end and a dephosphorylated 5' terminus at the opposite (HindIII) end to block self-ligation. A −4.5-kb DNA fragment, derived by digestion of p12-5S-C1 with Sall and AlwN1 (21), was also prepared for use as an internal control in the DraIII digestion assays.

**Preparation of nucleosomes and nucleosome arrays**

Reconstitution of DC mononucleosomes and 12-mer nucleosome arrays was independently carried out via the salt dialysis method as described previously (21). Briefly, 25 μg of DC DNA or 12 × 207 array template was mixed with 12.5 μg each of H2A/H2B dimer and H3/H4 tetramer in 2 mM NaCl, TE
buffer containing 20 mM DTT; incubated on ice for 30 min; and then transferred into 6–8-kDa–cutoff Spectrapore tubing. The sample was dialyzed into 1.2 M NaCl, TE buffer in the cold room followed by addition of TE buffer every 90 min to bring the NaCl concentration to 1, 0.8, and 0.6 M. Finally, the mixture was dialyzed overnight into T_{10}F_{0.1}_v (10 mM Tris-HCl, pH 8.0, 0.1 mM EDTA) buffer. Mononucleosomes were purified by sedimentation through 7–20% sucrose gradients (10 ml) with ultracentrifugation at 197,868 × g for 18 h in a Beckman SW41 Ti rotor at 4 °C. Nucleosome fractions (~0.4 ml) were collected in BSA-treated 0.6-ml Eppendorf tubes, and 20 μl of each was analyzed on a 0.7% native agarose gel (0.5 TBE). Array reconstitutions were carried out identically except that dialysis was for 4 h in TE buffer, 1 M NaCl; 3 h in TE buffer, 0.75 M NaCl; and then overnight in T_{10}F_{0.1}_v buffer. The extent of saturation of the array template with nucleosomes was assessed by magnesium–dependent self-association assays (67).

Ligation of DC mononucleosomes and 12-mer arrays was carried out with empirically optimized ratios of components to maximize formation of 25-mer arrays (Fig. 1). The ligation reactions typically contained 50 ng of sucrose gradient–purified DC nucleosomes, 1.5 μg of 12-mer arrays, and 2 μl (800 units) of T4 DNA ligase in ligation buffer (0.2 μg of BSA, 1 mM ATP, 10 mM MgCl₂, 0.25 mM EDTA, 10 mM Tris-HCl, pH 8.0) incubated in a final volume of 50 μl for 30 min at room temperature and then overnight at 4 °C. The 25-mer arrays were subjected to buffer exchange into TEN buffer (10 mM Tris, 0.25 mM EDTA, 2.5 mM NaCl) via microfiltration (48). To generate arrays containing a central NFR, the naked 204-bp DC DNA (601 nucleosome positioning sequence) was ligated with the 12 × 207 arrays. After ligation, all nucleosome dyads in the 25-nucleosome arrays are separated by 207 bp. Self-association assays and DraIII digestion kinetics support minimal loss of nucleosomes through the ligation process (see below).

**Binding of linker histone to nucleosome arrays**

*Xenopus* Napl and H1 were mixed in 2:1 ratio in binding buffer containing 10 mM Tris, pH 8.0, 1 mM EDTA, and 50 mM NaCl (58). The H1–Napl complex was mixed with 12- or 25-mer nucleosome arrays in the ratios stated in the figure legends in binding buffer containing 10 mM Tris-HCl, pH 8.0, 0.1 mM EDTA, and 50 mM NaCl and incubated on ice for 1 h, and binding was assessed by loading arrays with 5% glycerol onto a 0.7% native agarose gel (0.5 TBE).

**Quantification of 25-mer array DNA digestion and identification of k_{rel} rates of cleavage**

Analytical DraIII digestions were performed with 25-mer arrays in which the central DC nucleosome contained either all native histones (WT) or combinations of H3/H4 tetramers and H2A/H2B dimers in which acetylatable lysines were swapped for glutamines (H4 K5Q,K8Q,K12Q,K16Q (H4acm); H3 K4Q,K9Q,K14Q,K18Q,K23Q,K27Q (H3acm); H2A K5Q,K9Q (H2Aacm); and H2B K5Q,K12Q,K15Q,K20Q,K24Q,K27Q (H2Bacm)) as stated in the text and figure legends. For digestions, 20 μl of each array preparation (~1 μg) was mixed with an equal amount of p12 DNA control in 80 μl of 1× New England Biolabs CutSmart digestion buffer (50 mM potassium acetate, 20 mM Tris acetate, 10 or 0.5 mM magnesium acetate (as indicated), 1 mM DTT, pH 7.9). Nine microliters of the solution was removed as an undigested control (t = 0 min) and added to 2 μl of 6× SDS gel loading solution (10 mM Tris-HCl, pH 8.0, 1 mM EDTA, 30% glycerol, 1% SDS, 0.25% bromphenol blue dye). The remaining 71 μl was mixed with DraIII-HF (units as indicated in the figure legends) followed by rapid mixing and further incubation at 37 °C. Nine microliters of digested product was removed at time points as indicated in the figures legends and immediately mixed with 2 μl of 6× SDS gel loading solution. Samples were run on 0.7% SDS-agarose gels, the gels were imaged on a Bio-Rad imager (Gel Doc™ XR+), and the 25-mer array DNA bands and p12 control DNA bands were quantified with the volume tools in Image Lab. The rates of digestion were determined by plotting ln(fraction uncut) versus time for both the nucleosome array and naked DNA bands, and the data were fitted to a simple y = mx + b equation in Microsoft Excel as described previously (21). To allow comparison of the absolute rates of cleavage for chromatin and DNA templates within each sample, we also determined the relative rates of cleavage for the naked 25-mer array template and the naked 4.5-kb control DNA templates. We found that the naked 25-mer template, containing two DraIII sites, was cleaved 8 times faster than the 4.5-kb control template, which contains a single DraIII site, indicating that there is an inherent difference in the overall probability of a first-hit single cleavage event of about 4-fold between the two DNAs. The parameter k_{rel} was thus calculated as Slope chromatin/8x slope DNA for each of the 25-mer arrays, correcting for the differences in inherent rates between the two templates. For quantification of rates of cleavage for H1-bound arrays, the experiment was set up as described above except that naked DNA was not used as an internal control as the extent of H1 binding to the naked template and the effect on the control digestion rate were impossible to determine. Note that as originally shown by Widom and co-worker (19), under the conditions used for the digestions, the reactions are first-order in enzyme concentration, allowing correction of rates obtained at different restriction enzyme concentrations. Moreover, the rates reflect the probability of site exposure (equilibrium) rather than the rate at which sites become exposed.

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