Internucleosomal Interactions Mediated by Histone Tails Allow Distant Communication in Chromatin

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Background: Gene expression is regulated by DNA elements that often lie far apart along genomic sequences. Action across long distances on chromatin is a hallmark of eukaryotic transcriptional regulation. Although chromatin structure per se can support long-range interactions, the mechanisms of efficient communication between widely spaced DNA modules in chromatin remain a mystery. The molecular simulations described herein suggest that transient binary inter-nucleosomal interactions can mediate distant communication in chromatin. Electrostatic interactions between the N-terminal tails of the core histones and DNA enhance the computed probability of juxtaposition of sites that lie far apart along the DNA sequence. Experimental analysis of the rates of communication in chromatin constructs confirms that long-distance communication occurs efficiently and independently of distance on tail-containing, but not on tailless, chromatin. Taken together, our data suggest that internucleosomal interactions involving the histone tails are essential for highly efficient, long-range communication between regulatory elements and their targets in eukaryotic genomes.

The action of protein molecules over large distances on DNA requires special facilitating mechanisms. Indeed, sequences separated by more than a thousand base pairs (1 kb) along linear DNA do not communicate efficiently in vitro (1, 2). Prokaryotic proteins bound to more closely spaced regulatory elements can take advantage of the natural looping properties of DNA in forming the long-range contacts needed to control gene expression. The protein-mediated DNA loops implicated in the expression of eukaryotic genes, however, are much longer than the closed (~0.5 kb) DNA structures formed most easily in solution (3–5). Most eukaryotic genes are regulated by short DNA sequences called transcriptional enhancers that, after binding of proteins, can activate transcription over distances up to hundreds of kb. The action of these enhancers involves the direct association of the proteins bound at the enhancer site and at its target (promoter) site, with accompanying formation of a large loop of the intervening chromatin-covered DNA (6–8).

Although both kinetic and equilibrium aspects of enhancer-promoter communication over long distances on histone-free DNA have been extensively investigated (3, 9–12), the mechanisms of communication in chromatin remain poorly understood. New experimental approaches that yield quantitative data on the rate of enhancer-promoter communication on nucleosome-bound DNA constructs in vitro (13, 14) provide unique information for deciphering the mechanisms of eukaryotic gene expression in chromatin. The levels of RNA detected upon association of regulatory proteins that are bound to the ends of these constructs reveal surprising new information about the intervening DNA. In particular, the presence of nucleosomes along the DNA strongly facilitates distant communication between the bacterial nitrogen regulatory protein C (NtrC)3 and the RNA polymerase-α54 assembly (a convenient prokaryotic system for monitoring long-range association) and suggests that the observed enhancement of gene expression is more than simple DNA compaction (15).

Understanding long-range communication in chromatin requires knowledge of the contributions of the nucleosomes and the interspersed DNA linkers to the enhancer-promoter search. Early molecular simulations (16, 17) focused on a few of...
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the many facets of the complex protein-DNA assembly that determine the degree of chromatin compaction, such as the interplay between the length of linker DNA and the entry-exit angle of the DNA wrapped on the histone octamer. More recent studies (18–20) have begun to look at the effects of other structural features, such as the histone tails (effective electrostatic charge, tail modifications, etc.), on the equilibrium configuration and dynamics of short fragments of chromatin. None of the aforementioned work has looked at the contribution of chromatin structure to long-range enhancer-promoter interactions or explored the chromatin-folding problem from a multiscale perspective.

In this study, to identify the features of chromatin that are important for long-range communication, we have performed Monte Carlo simulations of different coarse-grained representations of chromatin and explored the relative likelihood of enhancer-promoter contacts in nucleosome-bound and protein-free DNA chains. Our computations suggest that transient electrostatic internucleosomal interactions mediated by the N-terminal tails of the core histones could strongly facilitate enhancer-promoter communication between widely separated sites on DNA and contribute to enhancer action. Our measurements of enhancer-promoter communication on physiologically relevant, saturated arrays of regularly spaced and precisely positioned nucleosomes, obtained with the techniques noted above, corroborate the predicted looping propensities. The fully saturated arrays support highly efficient, enhancer-promoter communication over DNA distances from 0.7 to at least 4.5 kb and further show that the core histone N-terminal tails become progressively more important as the distance between elements is increased. Taken together, the data suggest that transient, internucleosomal electrostatic interactions mediated by the histone tails give rise to highly efficient, long-range communication between regulatory elements and their targets in eukaryotic chromatin.

**EXPERIMENTAL PROCEDURES**

**Bead-spring Model**—In the bead-spring model of chromatin, the spring is modeled by a finitely extensible nonlinear elastic potential, which captures the resistance of chromatin to stretching. An energy cost to bending is introduced to capture the bending rigidity of chromatin, and phantom beads are introduced so that the springs and beads are self-avoiding and the model polymer is non-self-intersecting.

Modeling the screened electrostatic interaction between nucleosomes requires some experimentation. We argue that the histone tails, because of their flexibility, are available for temporary electrostatic interactions with nucleosomes. The possibility of nucleosome-nucleosome interaction has been raised in this and many other studies (21–23). Such interactions have been detected, and participating molecular surfaces have been identified experimentally (24, 25). In this picture, the histone tails are either extended and available for electrostatic interaction with other nucleosomes or wrapped on the nucleosome surface, and unable to mediate interactions with other nucleosomes. We have simplified this picture further and introduced a single effective tail at each nucleosome that is in two discrete states, available or unavailable, that it switches between stochastically.

We ignored the orientational dependence of nucleosomal interactions, which should not matter in a coarse-grained description. The screened attractive potential between beads available for bonding is a Gaussian well potential, which is harmonic in bead separation within the screening length and assigned a depth (attained when beads are touching) of a few $k_B T$, where $k_B$ is the Boltzmann constant, and $T$ is the temperature in Kelvin. The interaction state of a histone tail was modeled by an internal variable, which is either available or unavailable. This internal variable encountered a uniform potential barrier to flipping. We tuned the height of this barrier and the depth of the potential well for comparisons in our simulations. We simulated a ring of 200 beads using standard Metropolis-Monte Carlo methods. All beads were identical, and the data generated were the set of temporary bonds formed between beads at fixed sampling time. We collected the order of $5 \times 10^4$ accepted Monte Carlo steps for each bead, where each step is 5% of the bead size.

**Mesoscale Chromatin Model**—Our mesoscale treatment of chromatin represents DNA in terms of the set of rigid-body base-pair-step parameters that specify the orientation and displacement of successive base pairs. These details, excluded in earlier models of fluctuating chromatin fibers (17, 18), are needed to keep precise account of the spatial disposition of interacting proteins. The protein-free linker DNA is governed by an ideal elastic potential that allows for deformations in local structure consistent with the solution properties of DNA (26), and the nucleosomes are treated as rigid bodies. The atoms of proteins are stored in the frames of the nucleosomal base pairs and thereby included in the model (27). Each nucleosome also carries a superhelical reference frame, which is used in the detection of excluded volume and the characterization of multinucleosomal arrangements. The enhancer-binding protein NtrC was represented as a cylinder (12.4 nm diameter $\times$ 4 nm high), and the RNA polymerase-octet assembly, on the promoter, was represented as a prolate ellipsoid (15 $\times$ 11 $\times$ 11.5 nm) consistent with the observed dimensions (28, 29) (supplemental Fig. S1).

The electrostatic interactions that govern nucleosomal formation and interaction (30–32) were treated approximately in terms of charge clusters identified on the basis of the spatial locations and chemical identities of protein and DNA atoms in the core-particle structure. Our treatment ignored the charge-compensated cationic and anionic residues in the protein interior and placed the negative charges on DNA into groups comparable in size with those identified for protein side groups (supplemental Fig. S2). The positions of charges on DNA and bound nucleosomes thus varied with change in linker conformation. The charges on protein also moved in accordance with a Gaussian distribution based on the positions of the amino acid atoms comprising the clusters. These small moves very crudely mimicks the dynamics of the histone tails, which adopt fairly defined structures in solution (33, 34). The model thus included structural features more realistic than those in typical physics-based models (17) but simpler and easier to decipher and con-
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FIGURE 1. Nucleosome positions and occupancies on saturated 601 arrays. A, schematic diagram of the pYP05 template. The 601 nucleosome-positioning sequences are indicated by wide arrows. The glmAp2 promoter and the enhancer are positioned over 0.7–4.5-kb distances from each other (2.5 kb in the image). The first 601 nucleosome-positioning sequence (601 NPS) is truncated by 35 bp from the 3’ end and overlaps with the 176-nucleotide transcript (dashed arrow). B, schematic map of the arrays containing different numbers N of 601 nucleosome-positioning sequences (NPS) between the glmAp2 promoter and the enhancer (E). AluI sites for restriction endonucleases are located close to the middle of the positioning sequences, and Scal sites are in the middle of the 30-bp DNA linkers. C, experimental approach: a restriction enzyme sensitivity assay with primer extension. Chromatin was assembled on a negatively supercoiled pYP05 plasmid and digested with an excess of restriction enzymes AluI or Scal. Purified DNA was next digested with EcoRI restriction endonuclease to create one DNA end and then subjected to primer extension using a promoter-proximal end-labeled primer. D, characterization of the 601-nucleosomal arrays assembled on pYP05 using the restriction digestion sensitivity assay. Analysis of primer end-labeled DNA by denaturing PAGE was performed. Undigested (U) and fully digested (D) DNA fragments are indicated. Note that the increase in the chromatin assembly level results in almost quantitative protection of the templates from AluI, but not Scal restriction enzyme. M denotes theMspI digest of pBR322 plasmid.

trol at the local level than those fitted to all-atom treatments of electrostatic properties (18).

We simulated the chromatin system, given its complexity, in multiple stages (see supplemental Methods). The dense packing of nucleosomes on the system made the computations challenging in that internucleosomal contacts often trap the system in small, deep energy wells. Samples of 10^6–10^8 configurations were generated to access the available configuration space.

We computed the distances and angles that specify the relative spatial disposition of the proteins at the ends of the simulated chains and a loop-closure probability term analogous to the classic Jacobson-Stockmayer cyclization factor (35). The latter quantity depends upon the fraction of configurations that meet the selected chain-closure criteria (27), here the probability that the distance between the bound proteins terminated in the desired range.

**Plasmids, Proteins, and Chromatin Assembly**—Plasmids were constructed with the approaches given in supplemental Table S1. All templates for in vitro transcription used in this work are derivatives of the pLY10 plasmid, which has been described before (2) and used in previous studies (15). The pYP05 plasmid (Fig. 1A) containing an NtrC-dependent enhancer and a glmAp2 promoter separated by a spacer DNA of 0.7–4.5 kb, including specific numbers of strong 601 nucleosome-positioning sequences, was cloned from pLY10 in several consecutive steps. In the first nucleosome-positioning sequence (NPS), 35 bp were truncated from the promoter-distant end and replaced by the bacteriophage T7 transcription terminator site, thereby limiting the length of the transcript by 176 nucleotides (Fig. 1B). The lengths of the spacers between all NPSs within the pYP05 plasmid were the same (29 bp). The plasmids were constructed using PCR amplification of the desired fragments followed by restriction digestion of the fragments and the target vectors and ligation.

Established methods were used to purify all transcriptional machinery proteins (12), to prepare H1/H5-depleted chicken erythrocyte donor chromatin (36), to prepare tailless donor chromatin (by digestion of long H1/H5-depleted chromatin with trypsin) (37), and to reconstitute chromatin on supercoiled templates in vitro (36). Nucleosome positioning within the assembled arrays was verified using a restriction enzyme sensitivity assay. After chromatin assembly, the samples were digested with an excess of restriction enzymes, either AluI or Scal. Purified DNA was then digested with EcoRI restriction endonuclease to set one DNA end and subjected to primer extension with Tag DNA polymerase using a radioactively labeled primer localized immediately upstream of the promoter.

**Transcription**—Conditions for in vitro transcription were optimized for maximal utilization of the chromatin templates. Transcription was conducted as described previously (13). In short, all templates were linearized, the closed initiation complexes were formed, and the enhancer-binding protein (NtrC) was prebound to DNA in the absence of ATP. Next, ATP was added for a short time (1 min) to initiate enhancer-promoter communication and formation of the open initiation complex, which is competent to begin elongation. A mixture of all four NTPs (containing [α-32P]GTP and heparin) was then added to the reaction to start transcript elongation and to limit it to a single round. The reaction was continued for 15 min, and labeled RNA was purified and analyzed by denaturing PAGE. The gel was dried, exposed, and scanned on a Cyclone imager (PerkinElmer Life Sciences). The data were analyzed using the...
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**RESULTS**

**Internucleosomal Interactions Increase Long-range Chromatin Looping**—Many models of chromatin, incorporating varying degrees of structural detail, have been explored for different goals (23, 38–41). Here we focused on the effects of electrostatic interactions between nucleosomes, likely mediated by flexible histone tails, in enhancer-promoter communication. The existing models are insufficient for our purposes as we explain below. We initially modeled chromatin minimally as a bead-spring polymer, where the beads represent a few nucleosomes and the spring connecting them represents the linker DNA and overall flexibility of packed nucleosomes. We also introduced a discrete variable for each bead, which indicates whether the flexible histone tails are available for interaction with other beads. Two nearby beads that are available for interaction could form a temporary “bond.” Two key parameters of this model are the free energy barrier to the tails becoming available and the free energy loss when the temporary bond is broken.

The key quantity of interest is the contact probability distribution as a function of the separation of beads. For a self-avoiding, semiflexible polymer, this distribution exhibits a maximum for separations on the order of the persistence length. Below the persistence length, contacts are unlikely because of the cost of bending the polymer, and above this reference point, the contact frequency falls off as a power law of the separation. The typical contact probability is peaked around a separation determined by the choice of persistence length, and that peak is excluded from this plot. As expected, the tail of the distribution falls off as a power law for the noninteracting case (blue line), but has a much slower fall-off for our interaction model (dotted and solid red lines correspond to two different interaction strengths). A, number of contacts against simulation time (arbitrary units). The timeline plot (simulation steps) shows the number of transient bonds formed in a window of a typical run. Notice the occasional peaks in the number of contacts, where the value fluctuates significantly above the mean. The side panel shows the frequencies of the contacts.

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At intermediate interaction strength (1.25 or 1.5 $k_B T$, Fig. 2A), the probability distribution displays a distinct profile with a much slower fall-off at longer separation of beads when compared with that of the noninteracting case. This flattened tail of the distribution implies enhancement of long-range communication between nucleosomes, mediated by histone-tail interactions, and is not simply an overall enhancement of contact frequency. The time evolution of the number of contacts (Fig. 2B) shows that the system intermittently goes through states with a relatively high number of contacts. We conclude that the interactions lead to a novel mechanism of long-range action, which avoids a collapsed state but enhances long-range interactions.

**Histone Tails Introduce Intrinsic Compaction and Curvature That Enhance Long-range Chromatin Interactions**—A closer look at chromatin structure at the level of specifically positioned nucleosomes and fluctuating linker DNA segments shows how the presence of the histone tails enhances the likelihood of long-range intramolecular contacts when compared with that in both free DNA of the same chain length and chromatin fragments modeled with tailless nucleosomes in the same positions. We simulated saturated arrays of regularly spaced nucleosomes with spacings known to form higher-order chromatin structures in vitro (42, 43), i.e. 147-bp nucleosomal DNA segments separated by 30-bp linkers. We treated DNA at the base-pair level and incorporated, in the bound nucleosomes, the precise DNA pathway and the positions of charged amino acid atoms found in the currently best-resolved core-particle structure (44). We let the elastic properties of the linker DNA, the steric overlap of nucleosome cylinders, and the electrostatic interactions of the DNA polyanion and histone polycations guide the spatial arrangements of the mesoscale assembly (see “Experimental Procedures”).

The interactions of the histone tails have a profound effect on the configurations of the simulated chains. The charges on the surfaces of modeled intact nucleosomes shift the distributions of both the end-to-end distance and the radius of gyration toward lower values than those of the corresponding DNA with bound tailless nucleosomes (Fig. 3A, supplemental Fig. S3). Both nucleosome-bound systems are globally stiffer and substantially more compact than free DNA and thus more prone to looping mediated by other proteins. For example, the simulated likelihood of close contact between a cylindrically shaped NtrC...
activator protein complex located on an enhancer site at the 3′ end of the DNA and a prolate ellipsoidal RNA polymerase-σ54 complex bound at a promoter site on the 5′ end is more than an order of magnitude greater for duplexes with 4–25 intervening nucleosomes than for ideal, protein-free DNA of the same chain length (Fig. 3B). Moreover, the computed enhancement of looping for fragments with nucleosomes that contain intact histone tails is 1.3–3.0 times greater than that for the corresponding tailless arrays. Remarkably, the positive impact of the histone tails on the efficiency of looping in chromatin becomes larger as the size of the loop increases (Fig. 3B). In other words, in the presence of histone tails, the likelihood of chromatin looping is less distance-dependent, the same result found at longer chain lengths with the bead-spring polymer model. The combined data suggest that the histone tails are essential for distant communication within a chromatin fiber.

The detailed depiction of individual nucleosomes and linker base pairs incorporated in the mesoscale representation of chromosomal DNA reveals an intrinsic compaction and curvature in the fragments bearing intact nucleosomes when compared with those that bind tailless nucleosomes (Fig. 3C, supplemental Figs. S1 and S4). This difference in chain configuration underlies the increased levels of end-to-end contact detected in the simulations of nucleosomal arrays.

The changes in average global structure reflect slight differences in the local arrangements of nucleosomes, which arise, in turn, from the modeled interactions of the histone tails. These interactions affect both the number and the type of intramolecular contacts within the simulated chromatin fragments. For example, there are more close contacts of nucleosomes in the modeled chromatin systems with histone tails than in those without tails (Fig. 4A, supplemental Fig. S5). There are also more such contacts in the middle than at the ends of the nucleosome-bearing DNA. The three nucleosomes at either end of the chains make substantially fewer contacts on average with other nucleosomes than those in the middle. The central tail-bearing nucleosomes make as many as eight and on average 2.6 contacts with other nucleosomes in the simulated structures. These contacts tend to occur at close range, involving the three sequentially closest nucleosomes, i.e. nucleosomes in positions ±1, ±2, or ±3 with respect to that of the nucleosome of interest (Fig. 4B). Moreover, there is a much greater likelihood that a nucleosome in these chains will contact a nucleosome at position ±2, the nucleosome known to be spatially closest in a perfect zigzag model of chromatin, than at positions ±1 or ±3 (supplemental Fig. S6). By contrast, the nucleosomes in the middle of the chains that bind tailless histones contact 1.7 other
nucleosomes on average and associate with sequentially close nucleosomes with slightly different propensities. These subtle changes in number and type of contacts lead to the differences in mean internucleosomal structure that give rise to the overall compaction and curvature noted above. For example, both the angle between the cylindrical axes of nearby nucleosomes and the displacement of the centers of the nucleosomes tend to be greater when the tails are removed (Fig. 4C). The few longer-range contacts captured in the simulations, such as the \((n, n + 24)\) contact illustrated on a 50-nucleosome construct in Fig. 5, occur in concert with numerous shorter-range contacts and appreciable bending of the intervening DNA. The formation of such interactions is much less frequent in chains with tailless histones than in those with intact nucleosomes (Fig. 4B).

Histone N-tails Mediate Long-range Enhancer-Promoter Communication in Vitro—To evaluate the primary prediction of the computational studies (the effect of histone tails on distant communication in chromatin), we constructed a series of \(601_{177 \times N}\) plasmid derivatives with different numbers of 601 nucleosome-positioning sequences (45) between enhancer and promoter sites \((n = 4, 7, 13, \text{ or } 25)\) (Fig. 1, A and B). Nucleosomal arrays were reconstituted by transfer methods by use of continuous dialysis from 1 m NaCl (14) with intact or tailless core histones (Fig. 1C). Each 177-bp long nucleosomal repeat contained 147 bp of nucleosome-covered DNA and a 30-bp

![FIGURE 4. Histone tails affect contacts and spatial arrangements of nucleosomes in simulated chromatin fragments. A, color-coded mosaic of the numbers and locations of close (≤ 15 nm) internucleosomal contacts detected in simulated chains bearing 25 evenly spaced nucleosomes with intact (left) or tailless (right) histones. The positions of nucleosomes are denoted by the numbers at the bottom of the images, and the number of contacts (\#) are denoted by the scale on the right. The total number of contacts per simulated configuration, plotted to the right of each row, fluctuates around an average value of 60 for intact (red line) and 38 for tailless (blue line) nucleosomes. See supplemental Fig. S5 for mosaics of 4-, 7-, and 13-mers. B, color-coded map of the frequency of close contacts between all pairs of nucleosomes in simulated chains binding 50 regularly spaced nucleosomes. The color coding is based on a logarithmic scale to pinpoint the locations of rare, longer-range interactions, e.g. \(-5\) denotes a frequency of \(10^{-5}\). Note the distinctly different frequencies (colors) for the intact (+ tails) nucleosomes plotted below the diagonal versus those for the tailless (− tails) nucleosomes plotted above the diagonal. See supplemental Fig. S6 for frequency counts. C, representative configurations of four-nucleosome chromatin fragments in simulated chains with intact (top) and tailless (bottom) histones. The DNA is shown in pink, with the phosphorus atoms on complementary strands depicted by ribbons and the bases drawn as stick figures. The histones of successive nucleosomes are shown alternately in green and silver; the blue spheres depict charge cluster sites on the histone tails (see supplemental Fig. S2).](https://www.jbc.org/content/287/24/20253)

![FIGURE 5. Long-range contacts of nucleosome-decorated DNA occur in concert with numerous short-range interactions and appreciable bending of intervening chain. Simulated hairpin configuration of chromosomal DNA bearing 50 intact, evenly spaced nucleosomes is shown. In this example, one pair of nucleosomes, at positions 18 and 42, comes within 21 nm of each other, and almost every nucleosome pair separated by 1–3 intervening nucleosomes lies within 20 nm of each other. The NtrC assembly bound to the enhancer at the 3’ end of the chain is represented by a yellow cylinder, and the polymerase-α\(^+\) complex bound to the promoter at the 5’ end is represented by a gold prolate ellipsoid. The blue spheres depict charged clusters on the histone tails. Images were created with Mathematica™ and rendered with Blender.](https://www.jbc.org/content/287/24/20253)
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spacer between successive positioning sites. These constructs place the promoter and enhancer on the same or opposite sides of a fiber with the 6- and 12-nucleosome repeats per turn deduced from chemical cross-linking and low-resolution structural studies (46, 47) (Fig. 6A).

We determined the rate of enhancer-promoter communication in the 601_{177×N} nucleosomal arrays using the recently developed approach, outlined in Fig. 1C, which allows for the assembly of saturated nucleosomal arrays and quantitative characterization of the resulting chromatin system (13, 14). The DNA plasmids (Fig. 1B) contain, at one end, the NtrC-dependent enhancer, which can strongly activate the glnAp2 promoter placed at the other end (2, 12, 48).

Restriction digestion sensitivity assays (Fig. 1, B–D) show that the chromatin samples contain saturated nucleosomal arrays. The Alu restriction sites positioned within nucleosomes are fully protected from digestion by the core histones, whereas the Scal sites positioned on the linker DNA are fully sensitive to restriction enzymes (Fig. 1, B–D). A different version of the restriction enzyme sensitivity assay shows that the majority of nucleosomes assemble specifically on the high-affinity nucleosome-positioning sites, and not at random along the plasmid template nor on the enhancer and/or promoter (14). Taken together, the data suggest that saturated, defined nucleosomal arrays that contain specifically positioned nucleosomes are formed during assembly. Notably, nucleosome-saturated 601_{177×N} arrays spontaneously form chromatin fibers in buffers similar in composition to the transcription buffer used in this study (42, 43).

Evaluation of the relative rates of enhancer-promoter communication entailed linearization of the chromatin templates followed by a single-round transcription assay (Fig. 6, B and C). We previously found that enhancer-promoter communication is the rate-limiting step in the entire transcription reaction and that the yield of transcripts is directly proportional to the level of communication on both DNA and chromatin templates (2, 15). Furthermore, in the case of histone-free DNA, the experimental results quantitatively mirrored the expected end-to-end properties of flexible DNA chains (12).

The assembly of saturated chromatin arrays of different lengths with intact core histones resulted in strong and comparable activation of transcription on all 601_{177×N} templates (Fig. 6, C and D). Thus chromatin structure can support efficient enhancer-promoter communication over distances from 0.7 to at least 4.5 kb in vitro. As expected, the extent of activation by the assembly of intact chromatin became progressively lower as the length of the DNA spacer was increased, possibly due to the progressively increasing average spatial separation of the enhancer and promoter. The magnitude of activation between 0.7 and 4.5 kb, however, differed by only ∼1.5-fold, an indication that intact chromatin can maintain enhancer-promoter communication that is almost distance-independent in the constructs considered here. If the chromatin constructs were rigid fibers with the assumed nucleosomal repeat, the stimulatory effect of chromatin assembly on enhancer-dependent transcription would strongly and periodically depend on the number of nucleosomes on the intervening DNA (Fig. 6A). No strong periodic differences in the efficiency of activation were detected (Fig. 6D). The data do not support a rigid fiber model and suggest that the constructs are intrinsically dynamic structures that can support long-range communication in chromatin.

In striking contrast to enhancer-promoter communication in intact chromatin, the level of communication in tailless chromatin strongly decreases as enhancer-promoter spacing increases (Fig. 6D). Thus the rate of enhancer-promoter communication over 0.7 kb decreased only by ∼20% after removal of histone tails, but that over 4.5 kb dropped ∼4-fold. Taken together, the data suggest (in agreement with the predictions of our computations) that: (i) histone tails are essential for enhancer-promoter communication over large distances in saturated arrays of nucleosomes that form chromatin fibers and (ii) histone tails mediate long-range enhancer-promoter communication in chromatin.
DISCUSSION

Our composite findings show that internucleosomal interactions involving the histone tails can account for long-distance regulatory communication along chromatin. Our study of the bead-spring model, which represents chromatin at a length scale corresponding to many hundreds of nucleosomes, indicates that chromatin interactions induced by histone-tail contacts give rise to transient collapsed structures that nucleate long-range communication (Fig. 2). In addition, we have developed a mesoscale model of chromatin to decipher the effects of nucleosomal fine structure and linker DNA deformability on long-range polymeric interactions. Our Monte Carlo simulations of chains decorated with 4–50 evenly spaced nucleosomes reveal a natural compaction and curvature induced by the histone tails that enhance the likelihood of intranuclear loop formation and the concomitant association of bound, widely spaced proteins (Figs. 3–5). Our in vitro experimental system allows for analysis of enhancer-promoter communication over different distances on physiologically relevant, saturated arrays of precisely positioned nucleosomes that form higher-order chromatin structures (Fig. 1). Using this system, we have shown (Fig. 6) that: (i) efficient enhancer-promoter communication in chromatin occurs over distances from 0.7 to at least 4.5 kb and only weakly depends on the assumed rotational positioning of the communicating DNA elements within the assembled fiber; (ii) the assembled fiber is a dynamic structure that supports efficient communication over a variety of enhancer-promoter distances; and (iii) histone N-terminal tails are essential for long-range communication in chromatin.

Our experiments with nucleosomal arrays of different lengths suggest that at least two types of deformation contribute to the overall mobility of our chromatin constructs. Precise alignment of the enhancer and promoter over a short distance from each other (4–7 nucleosomes) is impossible without changes in the local structure of the fiber. To account for efficient short-range communication, we propose that nucleosomes on neighboring gyres of the fiber can move relative to each other. Such nucleosome repositioning within the fiber could be accomplished by spontaneous partial uncoiling of nucleosomal DNA from the histone octamer (49). Communication over larger distances (10–25 nucleosomes) cannot occur only by this mechanism because extensive uncoiling of nucleosomal DNA is unlikely. Therefore communication over large distances must involve bending of the chromatin fiber as a whole (Fig. 5) and take advantage of the deformability of the DNA linkers and the electrostatic interactions of nucleosomes. Such features would also occur in shorter chains.

As noted above, our experimental design was based on chromatin fiber models with 6- and 12-nucleosome repeats deduced from histone cross-linking and low-resolution structural studies (42, 43). Neither model accounts for our findings. For example, if the six-nucleosome repeat applied, the construct with seven nucleosomes would have produced much higher levels of RNA than that with four nucleosomes. That is, enhancer and promoter sites separated by seven nucleosomes would lie on roughly the same face of such a fiber and thus come into more frequent contact than sites with four intervening nucleosomes, which would lie on nearly opposite faces of the fiber. On the other hand, a four-nucleosome spacer may not be sufficiently long to form an organized fiber. Similarly, if the fiber model with a 12-nucleosome repeat were correct, the construct with 13 nucleosomes would have produced much higher levels of RNA than a similarly organized structure with seven nucleosomes (Fig. 6A).

Determination of the nucleosome repeat of constructs like ours in solution requires the preparation and analysis of the long-range interactions on an extensive series of new templates. The monotonic decay of transcript levels found here (Figs. 3B and 6D) suggests that the enhancer and promoter lie on the same face of the examined constructs. Preliminary simulations based on the current mesoscale model reveal an oscillatory pattern of contact frequencies indicative of a structural repeat shorter than six nucleosomes. The constructs described here lie near the peaks in the predicted oscillations, and simulated arrays of intermediate lengths with the same nucleosomal spacing contribute to both peaks and valleys in the profiles. The differences in compaction between the predicted structures and the published images of similar nucleosome constructs (42, 43) may reflect the different conditions of our experiments and computations (low divalent salt) when compared with those needed to visualize compacted fibers (100 mM MgCl₂ or the addition of linker histones).

Our simulations add a new perspective on the looping of chromatin not considered in previous research. The likelihood of looping chromatin into closed structures of the sizes detected in cells (∼100 nucleosomes) is often treated analytically (50, 51), and only recently has it been estimated in direct computations (52). The transient binary interactions introduced in our coarse-grained model allow for isolated long-range internucleosomal contacts not considered in previous work. Our mesoscale treatment of nucleosome-decorated DNA is consistent with previous simulations, which take explicit account of the positively charged histone proteins and the DNA polyelectrolyte backbone (20, 53). For example, the average number of internucleosomal contacts per core particle found here (Fig. 4A) is comparable with the values found in Langevin dynamics simulations of interacting charged beads (used to approximate nucleosomes) in the presence of low levels of Mg²⁺ (20), and the reduction in contact number found in the absence of the charged histone tails is similar to the loss of contacts determined upon reduction of bead charge. Our explicit treatment of nucleosomal and linker DNA, however, generates different distributions of nucleosome-nucleosome contacts from those reported in simulations that represent the linker by a few beads and confine the histone-bound DNA to an idealized pathway (53), i.e. a preponderance of contacts with nucleosomes at positions ±2 here (Fig. 4) versus roughly equivalent numbers of contacts at ±1 and ±2 in related simulations with short linkers (3–4 beads, each spanning ∼9 bp steps). Like many others (50, 54–56), we find that chromatin is a highly efficient communication device and a highly flexible polymer. Our computed estimates of enhancer-promoter contacts in chromatin versus free DNA, however, exceed the measured levels by roughly 4-fold (note the different scales in Fig. 3B versus Fig. 6D). As noted above, communication in chromatin could
be limited by the probability of enhancer-promoter juxtaposition, such as found on relaxed and supercoiled DNA (12). This idea is consistent with data showing that the rates of communication on supercoiled DNA and chromatin are comparable (15).

Finally, our studies suggest that efficient communication over a large distance (more than 1.5 kb) critically and selectively depends on the presence of intact tails on the core histones. This observation implies that communication over a distance could constitute a separate regulated step during execution of various intranuclear DNA-targeted processes. Specifically, histone tails are subject to intensive modifications (57); many of these modifications (e.g., acetylation or phosphorylation) change the charges of histone tails, destabilize higher-order chromatin structure (e.g, histone H4 acetylation at position 16 (H4K16Ac) (58, 59)), and most likely affect chromatin dynamics. We propose that in addition to their other important functions, some histone tail modifications could affect the rate of enhancer-promoter communication, perhaps in a regulated way. In particular, H4K16Ac is associated with active chromatin (60) and could facilitate distant communication in chromatin.

Acknowledgment—We thank Tim Richmond for the plasmid containing 601,177×12 nucleosome-positioning sequences.

REFERENCES

Histone Tails Help to Activate Genes on Chromatin

.. figure:: Histone_Tails_Help_to_Activate_Genes_on_Chromatin.png

Histone tails help to activate genes on chromatin.
SUPPORTING INFORMATION

Internucleosomal Interactions Mediated by Histone Tails Allow Distant Communication in Chromatin

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FIGURE S1. Histone tails reduce the distance between regulatory proteins bound at the ends of simulated chromatin fragments. Representative three-dimensional arrangements of fluctuating chromatin chains bearing a 3′-terminal enhancer site with bound NtrC, a 5′-terminal promoter site with bound RNA polymerase, and 4, 7, 13, or 25 evenly spaced nucleosomes with intact (left) or tailless (right) histones. The DNA and protein are depicted and rendered as in Fig. 5. The enhancer-promoter separations correspond to the simulated mean values.
FIGURE S2. Simulated electrostatic interactions of nucleosomes and linker DNA depend upon the positions and magnitudes of histone and DNA charge centers. Charge centers identified by K-means clustering (1) on the currently best-resolved nucleosome core-particle structure (Protein Data Bank file 1kx5) (2). The DNA backbones are displayed as cyan ribbons and the histone proteins as gray ribbons. The cluster centers, which represent 16 groups of closely spaced charges from the cationic amino-acid nitrogens on the histone tails, are color-coded in red and the 18 groups from amino acids at the surface of the folded histone core in blue. The anionic clusters on DNA, which combine the charges of three successive base-paired units, are depicted in magenta. The model ignores amino acid charges within 2.3 nm of the nucleosomal axis, given that this region has a net charge close to zero and the charges within this limit tend associate as ion pairs (3).
FIGURE S3. Histone tails introduce a natural compaction in simulated nucleosome-decorated DNA chains. Simulated distributions of the end-to-end distances (left) and radii of gyration (right) of DNA chains binding (top) 4, (middle) 7, and (bottom) 25 regularly spaced, fully intact or tailless nucleosomes (red and blue curves, respectively) compared to those of protein-free DNA chains (inset, black curves) of the same length. See legend to Fig. 3A.
FIGURE S4. Histone tails induce greater curvature in simulated nucleosome-decorated DNA chains. Variation of average curvature $\kappa$ with contour length $\ell$ along the fiber axes of ensembles of simulated chromatin fragments bearing (left) 13, (center) 25, or (right) 50 nucleosomes evenly spaced at 177-bp increments, i.e., 147-bp nucleosomes and 30-bp linkers. Snapshots of representative configurations and the fiber axes (center lines), obtained from the average positions of the centers of three successive nucleosomes, are shown below each plot. The DNA is depicted as a colorless tube and the histone cores by alternating green and silver cylinders. The center lines of the fibers made up of intact nucleosomes are depicted in red and those made up of tailless nucleosomes in blue. The same color-coding is used in the plots of curvature, including the mean curvature of the fibers as a whole (dashed lines). Note the approximately constant curvature of chains with different lengths and the greater curvature and shorter contours of the chains with intact vs. tailless nucleosomes. Images generated as in Fig. 5.
FIGURE S5. Histone tails increase internucleosomal contacts in simulated chromatin fibers. Color-coded mosaic of the numbers and locations of close (≤15 nm) internucleosomal contacts detected in simulations of chromatin chains bearing (left) 4, (center) 7, and (right) 13 evenly spaced, intact or tailless nucleosomes. See legend to Fig. 4A.
FIGURE S6. Histone tails affect the types of internucleosomal contacts in simulated chromatin fibers. Total frequency of contact between all nucleosomes separated by the same sequential interval $m$ in chromatin fibers binding 50 regularly spaced, intact (native) or tailless nucleosomes. $W(m)$ is the sum of the frequencies of contact for all pairs of nucleosomes $n$ and $n+m$ for each $m$ between 1 and 49, where $n$ is the sequential position of the nucleosome in the array, i.e., the sum of the probabilities along the diagonals in Fig. 4B. Red bars: native histones; blue bars: tailless histones.
Table S1. Cloning approaches used in this study.

<table>
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<tr>
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<th>Oligonucleotides/DNA fragments used *</th>
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<th>Cloned into</th>
<th>Restriction enzymes</th>
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<td>pYP01</td>
<td>5´- aactaccgattaaaAGCTTgcgcgaacacact</td>
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*Restriction sites are in capital letters.
†See (4).
‡See (5).
SUPPORTING METHODS

Mesoscale simulations

We simulate the chromatin system, given its complexity, in multiple stages. The system is initiated by positioning nucleosomes along an otherwise naturally straight DNA chain. The starting states of the linker DNA and the enhancer and promoter sequences at the ends of the chain correspond to the equilibrium rest state of an ideal, perfectly straight double helix with 10.5 base pairs per turn and 3.6 nm pitch, corresponding to ~34.3° twist and 0.34-nm rise at each base-pair step. The nucleosomal DNA is assigned the spatial pathway found in the currently best-resolved nucleosome structure (2). Trial moves that generate a new set of rigid-body parameters (shift, slide, rise, tilt, roll, twist) (6) for a randomly selected base-pair step on an arbitrary DNA linker are performed using Gaussian sampling and an ideal elastic potential (7). The resulting configuration is generated and the excluded volume, i.e., pairwise steric overlap of cylinders embedded on each the nucleosome core particle, is evaluated for the proposed move. The axis of the cylinder is the line from which the 147 base-pair centers are minimally displaced (8) and the radius is large enough to include the nucleosomal DNA. Electrostatic interactions are determined if the system is free of such contacts. The acceptance or rejection of a trial move is based on the difference in the electrostatic energies of the proposed and current states using the standard Metropolis criterion (9). Once a linker move is accepted, the positions of the charged clusters in histones are varied at random with Gaussian sampling. The repositioning is based on the observed spatial dispersion of the charge-bearing nitrogen atoms from the cationic amino acids that comprise each cluster (specifically, the coordinates of the arginine N<sup>ε</sup> and lysine N<sup>γ</sup> atoms in the nucleosome core-particle structure (2)). The configurations reported herein were found using a Debye-Hückel potential with a screening length of 0.83 nm (consistent with the ionic composition of the transcription buffer detailed in the companion article (10)) and a dielectric constant of 80. The configuration of the DNA on the two (promoter, enhancer) ends of the chromatin system is sampled following successful linker-DNA and histone-tail moves. DNA structure is varied, interactions are computed, and trial moves are accepted with the same procedure used to sample linker DNA. In contrast to earlier simulations of enhancer-promoter communication (11), which ignore the interactions of NtrC and polymerase, the simplified protein models described in the text are 'bound' to the
accepted nucleosome-decorated configurations, i.e., offset by an appropriate distance from the central base pairs of the enhancer and promoter sequences (see Fig. S1), and used in the computation of the looping probabilities. Replica-exchange methods (12-14) are used to enhance the efficiency of Monte-Carlo simulations, i.e., the system is assigned different temperatures and simulated in parallel on different processors. The choice of potential and sampling methodology do not affect our general conclusions about the role of the histone tails in long-range communication on chromatin. Complete details will be presented in a forthcoming publication (NC, AVC, GZ & WKO, in preparation).

REFERENCES


