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Methods

High-resolution nucleosome mapping reveals transcription-dependent promoter packaging

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Genome-wide mapping of nucleosomes has revealed a great deal about the relationships between chromatin structure and control of gene expression, and has led to mechanistic hypotheses regarding the rules by which chromatin structure is established. High-throughput sequencing has recently become the technology of choice for chromatin mapping studies, yet analysis of these experiments is still in its infancy. Here, we introduce a pipeline for analyzing deep sequencing maps of chromatin structure and apply it to data from *S. cerevisiae*. We analyze a digestion series where nucleosomes are isolated from under- and overdigested chromatin. We find that certain classes of nucleosomes are unusually susceptible or resistant to overdigestion, with promoter nucleosomes easily digested and mid-coding region nucleosomes being quite stable. We find evidence for highly sensitive nucleosomes located within “nucleosome-free regions,” suggesting that these regions are not always completely naked but instead are likely associated with easily digested nucleosomes. Finally, since RNA polymerase is the dominant energy-consuming machine that operates on the chromatin template, we analyze changes in chromatin structure when RNA polymerase is inactivated via a temperature-sensitive mutation. We find evidence that RNA polymerase plays a role in nucleosome eviction at promoters and is also responsible for retrograde shifts in nucleosomes during transcription. Loss of RNA polymerase results in a relaxation of chromatin structure to more closely match in vitro nucleosome positioning preferences. Together, these results provide analytical tools and experimental guidance for nucleosome mapping experiments, and help disentangle the interlinked processes of transcription and chromatin packaging.

[Supplemental material is available online at <http://www.genome.org>. The microarray data and the sequencing data from this study have been submitted to the NCBI Gene Expression Omnibus (<http://www.ncbi.nlm.nih.gov/geo>) under accession nos. GSE18629 and GSE18530.]

Eukaryotic DNA is packaged in nucleosomes, composed of 147 bp of DNA wrapped ~1.7 turns around an octamer of histone proteins (Luger et al. 1997; Kornberg and Lorch 1999). Nucleosomes influence the expression of a huge fraction of yeast genes (Wyrick et al. 1999) and the precise positioning of nucleosomes relative to underlying DNA, controls access to protein-binding sites, and thereby affects regulatory programs (Stunkel et al. 1997; Lomvardas and Thanos 2002; Lam et al. 2008; Radman-Livaja and Rando 2009). In yeast, transcription start sites (TSSs) are generally positioned just within the +1 nucleosome, downstream of a nucleosome-depleted region generally referred to as the nucleosome-free region (NFR) (Yuan et al. 2005; Albert et al. 2007; Mavrich et al. 2008a). Nucleosome positioning is different between promoter types; TATA-less promoters tend to be “housekeeping” genes and are characterized by a canonical NFR upstream of their TSS; conversely, TATA-containing genes tend to be stress-responsive, noisily expressed, are more responsive to genetic perturbation of chromatin remodeling complexes, and their promoters are often at least partly occupied by nucleosomes (Basehoar et al. 2004; Ioshikhes et al.

2006; Newman et al. 2006; Field et al. 2008; Tirosch and Barkai 2008; Choi and Kim 2009).

While nucleosomes can package almost any sequence, work over the past few decades has established sequence rules that influence this packaging. Most importantly, rigid poly(A) tracts in DNA are unfavorable for nucleosome assembly and can direct formation of NFRs in vitro (Kunkel and Martinson 1981; Drew and Travers 1985; Iyer and Struhl 1995; Sekinger et al. 2005; Yuan et al. 2005; Ioshikhes et al. 2006; Kaplan et al. 2008; Yuan and Liu 2008; Segal and Widom 2009). This was recently confirmed globally in an in vitro reconstitution study, emphasizing a major role for nucleosome-excluding sequences in “programming” promoter architecture (Kaplan et al. 2008; Zhang et al. 2009). Much of the remainder of in vivo chromatin structure is proposed to result from “statistical positioning,” the idea that packing as many nucleosomes as possible in a short stretch of the genome will result in positioned nucleosomes (Kornberg and Stryer 1988; Yuan et al. 2005; Mavrich et al. 2008a). While some aspects of yeast chromatin structure can, in principle, be predicted based on sequence and packing rules, a number of protein machines, such as the ATP-dependent chromatin remodeling complexes, regulate transcription by moving nucleosomes (Workman and Kingston 1998; Clapier and Cairns 2009). Perhaps the most widespread chromatin-perturbing complex is RNA polymerase, whose passage disrupts histone–DNA contacts, and at very high transcription rates results in nucleosome eviction (Studitsky et al. 1994, 1997; Lee et al. 2004;

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Schwabish and Struhl 2004; Bondarenko et al. 2006; Dion et al. 2007; Field et al. 2008).

The advent of high-throughput sequencing technology has enabled rapid genome-wide analysis of nucleosome positioning at high resolution and has been used to map nucleosomes in yeast, worms, flies, medaka, and humans (Field et al. 2008; Mavrich et al. 2008a,b; Schones et al. 2008; Shivaswamy et al. 2008; Valouev et al. 2008; Sasaki et al. 2009). Yet, analytical and experimental methods for deep-sequencing analysis of chromatin are still in their infancy and mostly focus on averaged nucleosome occupancy levels at genomic loci, effectively transforming deep-sequencing data into an analog of tiling array data.

Here, we describe a novel analytical method for identifying nucleosome positions from Illumina sequencing data and automatically estimate nucleosome position, occupancy, and length from *S. cerevisiae* data. Using parameters extracted from these nucleosome calls allows us to identify groups of functionally related genes with significantly high or low values of a given parameter. Experimentally, we examine the impact of nuclease titration levels on this assay and identify properties of yeast chromatin that are influenced by the level of digestion as well as invariant properties. Finally, we explore the role of RNA polymerase in chromatin structure through the use of a temperature-sensitive mutant in the gene encoding the large subunit of RNA polymerase II (Pol II), *RPO21* (also known as *RPB1*). Comparing nucleosome positions and properties before and after Pol II inactivation, we confirm the role of RNA polymerase in nucleosome eviction at promoters, and find a surprising role in retrograde movement of nucleosomes over genes. By quantitatively analyzing changes in nucleosome positioning after Pol II shutoff we find that different classes of genes are subject to distinct perturbations by RNA polymerase. Finally, we confirm a role for RNA polymerase in perturbing nucleosomes from their thermodynamically favored positions.

Results

Identifying nucleosome positions by template filtering

The application of deep sequencing to mononucleosomal DNA results in million of reads from both ends of the mononucleosomal DNA segments. Published methods for calling nucleosome positions from Illumina data typically involve extending each single-end short sequenced read to the expected segment length of ~140 bp and then examining the coverage of different genomic loci by the accumulated extended segments. These methods clearly highlight nucleosome-depleted vs. nucleosome-occupied regions, for example, in averaged gene alignments (Albert et al. 2007; Field et al. 2008; Shivaswamy et al. 2008). More elaborate approaches identify nucleosome positions by identifying the center of these inferred segments and estimating the occupancy at different center locations (Shivaswamy et al. 2008). These methods do not account for the possibility that nucleosomes might occupy different positions in subpopulations of cells and assume uniform nucleosome lengths (i.e., no variability in digestion level at different nucleosomes).

To overcome these issues, we developed a method for calling nucleosome positions, occupancy, and length using template filtering (Turin 1960). This method is based on the observation that sequencing the ends of a nucleosome will result in an expected pattern of offset forward (F) and reverse (R) strand reads at the two ends (Fig. 1A). Due to variability in exact nucleosome position from cell to cell and variability in the extent of MNase digestion at each end of a nucleosome, the peak of reads at each nucleosome

end will form a distribution of variable width. Our method is based on identifying occupancy templates of forward and reverse reads that are typical of nucleosomes. The method then uses a fast procedure to identify locations where the forward and reverse read distributions correlate with a series of model templates (Fig. 1B; Supplemental Fig. S1). The method examines a range of distances between forward and reverse templates to capture over- and underdigestion of the ~147-bp nucleosomal DNA, and determines nucleosome length by choosing an F-to-R offset that maximizes correlation to F and R templates (Fig. 1C,D).

To further elaborate the method, we aimed to identify the common read distributions from the data. In other words, what are the typical distributions of end reads for the nucleosomes in an experimental data set? We first applied our method using a Gaussian-shaped template to create a preliminary map of nucleosome positions. We next examined read distributions at the ends of these initial nucleosome predictions, clustered read patterns to identify typical behaviors, and selected seven representative templates for use in our scans (Fig. 1B; Supplemental Fig. S1). Six of the templates exhibited several peaks, potentially indicating nucleosomes with variable ends resulting from subpopulations and/or variable nuclease digestion. For example, nucleosomes with ends that match template 2 have sets of ends separated by 10 bp (Supplemental Fig. S1), suggesting that the first stage in overdigestion of nucleosomes by micrococcal nuclease is to cut one helical turn further into the nucleosomal DNA.

Our method fits the sequencing data to the best-correlated templates with varying distances between F and R templates. Local maxima in the correlation spectrum (Fig. 1C) are identified as potential nucleosome calls (Fig. 1E,F). To assemble the final set of nucleosomes, we use a greedy approach to choose the best-correlating template and distance per nucleosome (Methods). Occupancy is determined for each nucleosome by the number of reads contributing to a given nucleosome call. To account for different sequencing yields, we normalize nucleosome occupancies to a mean value of 1.

To evaluate the quality of our method's nucleosome calls, we used the nucleosome positions, templates, and occupancy calls to regenerate the sequenced reads that account for our called nucleosomes. This simulated data set is sampled with the same number of reads as the original sequencing run. These simulated reads closely match the original sequencing data, and the small difference (residual) between the measured and reconstructed data indicate that our nucleosome calls account for 88% of the sequencing reads, indicating that the small number of extracted parameters capture the majority of the experimental sequencing data (Supplemental Fig. S2).

We also examined data from seven additional previously published deep-sequencing datasets (Kaplan et al. 2008; Shivaswamy et al. 2008). We find that the same templates are common in all datasets (Supplemental Fig. S3A), confirming the generality of our approach. However, we did find that the occurrence frequency of different templates differed between datasets, which we ascribe to differences in numbers of reads as well as differences in digestion between different MNase preparations (Supplemental Fig. S3B; see below). Positions of nucleosome calls were generally concordant between datasets (Supplemental Figs. S4, S5), with the major difference occurring in -1 nucleosome calls, which we also identify below as being dependent on digestion levels.

Finally, we compared our nucleosome calling method to an alternative calling algorithm based on identifying peaks in data where forward and reverse reads have been shifted a

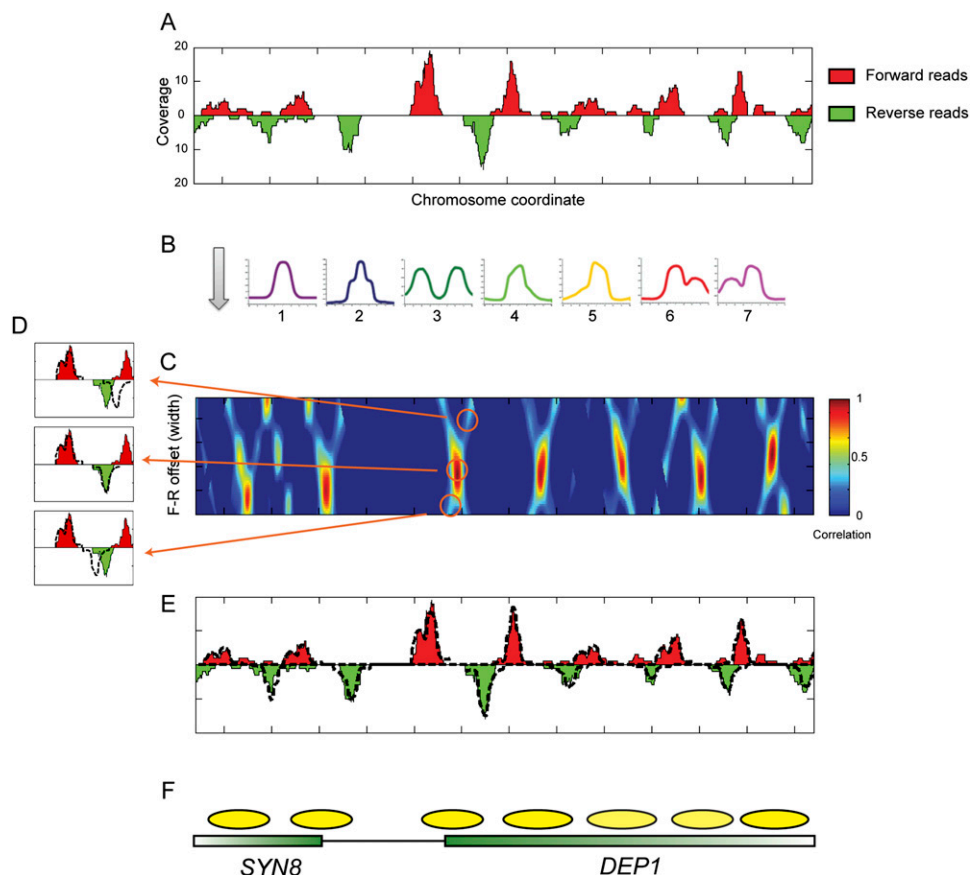


Figure 1. Template filtering overview. (A) Deep sequencing data for a typical stretch of the yeast genome. Coverage by forward-strand sequencing reads are shown as red peaks, whereas coverage by reverse-strand sequencing reads are shown as inverted green peaks. (B) Templates. Forward and reverse-strand read distributions are cross-correlated with each of the seven templates shown. (C) Correlation coefficient heat map of template 1 for forward and reverse templates at varying center positions (*x*-axis) and distances (*y*-axis). (D) Examples of templates spaced too far apart (*top*), at the optimal distance (*middle*), or too close together (*bottom*). Dotted lines indicate template outlines being compared with the underlying data. (E) Read distributions explained by the optimal template matches are shown as dotted lines for the region in A. (F) Schematic of nucleosome calls and underlying gene annotations.

half-nucleosome width toward one another (Shivaswamy et al. 2008). Nucleosome calls were highly concordant for both methods (Supplemental Fig. S5A), particularly for clearly well-positioned nucleosomes (those that match Template 1 in our approach). Examination of regions where nucleosome calls differed revealed that our method fails in regions where only one end of a nucleosome generates reads, whereas our method better captures data from “fuzzy” regions such as mid-coding regions (Supplemental Fig. S5B,C), thereby better capturing nucleosome occupancy over such regions.

By examining distributions of end reads, we found that nucleosomes at different locations vary in their digestion patterns. Over- and underdigestion of DNA can be due to many possible factors, including properties of the DNA sequence (e.g., sequence composition, bendability, etc.) and properties of the nucleosome (e.g., histone modification state). To investigate these two factors we tested whether the digestion template at nucleosome ends is associated with specific sequence properties and/or specific modification annotation. Consistent with previous reports of MNase sequence preference (Dingwall et al. 1981; Horz and Altenburger 1981), we find that different templates are associated with a clear sequence preference for location of A/T dinucleotides (Supplemental Fig. S6B). On the other hand, we also observed that mul-

timodal templates such as template 3 were enriched at locations previously described as “fuzzy” or delocalized, such as mid-coding regions and over promoters of stress-responsive genes (Supplemental Fig. S6C,D). These results suggest that both nucleosomal subpopulations (i.e., delocalization) as well as sequence composition are significant sources of variability in patterns of nucleosome reads. While these two factors cannot be completely disentangled when using MNase to map nucleosomes, we note that the use of several templates provides an automated means for taking MNase sequence biases into account.

Nucleosome positioning in growing yeast

Genome-wide maps of nucleosomes in actively growing yeast have been the subject of a rage of recent studies (Yuan et al. 2005; Albert et al. 2007; Lee et al. 2007; Field et al. 2008; Mavrich et al. 2008a; Shivaswamy et al. 2008). To further evaluate our methodology, we carried out deep sequencing of mononucleosomal DNA from actively growing yeast, and applied our method to generate a map of nucleosome locations (Supplemental Table S1). Our method automatically extracts features of interest, such as NFR width, nucleosome length (which is not available using previous methods),

nucleosome spacing, and occupancy. To systematically examine these features we used a compendium of experimental gene annotations that we previously collected (Dion et al. 2007; Wapinski et al. 2007) and compared these against multiple nucleosome attributes associated with each gene (e.g., +1 nucleosome occupancy, spacing between the +1 and +3 nucleosome, etc.). Using the Kolmogorov-Smirnov (KS) test (Fig. 2A) we discovered attributes whose distribution in specific gene groups was significantly (false discovery rate [FDR] < 0.05) different from the background (Fig. 2B; Supplemental Table S2).

This analysis highlights previously described features of yeast chromatin as well as novel ones. We recapitulate the dichotomy between promoter packaging of growth (TFIID-regulated, TATA-less, high-expression growth genes) and stress (SAGA-regulated,

TATA, noisy expression, rapid histone replacement) genes (Fig. 2B) (Ioshikhes et al. 2006; Newman et al. 2006; Albert et al. 2007; Dion et al. 2007; Choi and Kim 2008, 2009; Field et al. 2008; Tirosh and Barkai 2008). For example, canonical growth genes, such as those encoding ribosome proteins, are enriched with wider NFRs and often also exhibit lower nucleosomal occupancy of coding regions, whereas stress genes, such as heat shock-induced genes, show the opposite behavior. This dichotomy is clearly the major feature that separates classes of genes in yeast, but our analysis also shows finer distinctions beyond this dichotomy. In general, the second distinguishing axis beyond NFR width is coding region nucleosome occupancy, which further can be separated into occupancy of 5' nucleosomes such as the +1 nucleosome, and occupancy of mid-CDS nucleosomes located distal from either end of the gene. For

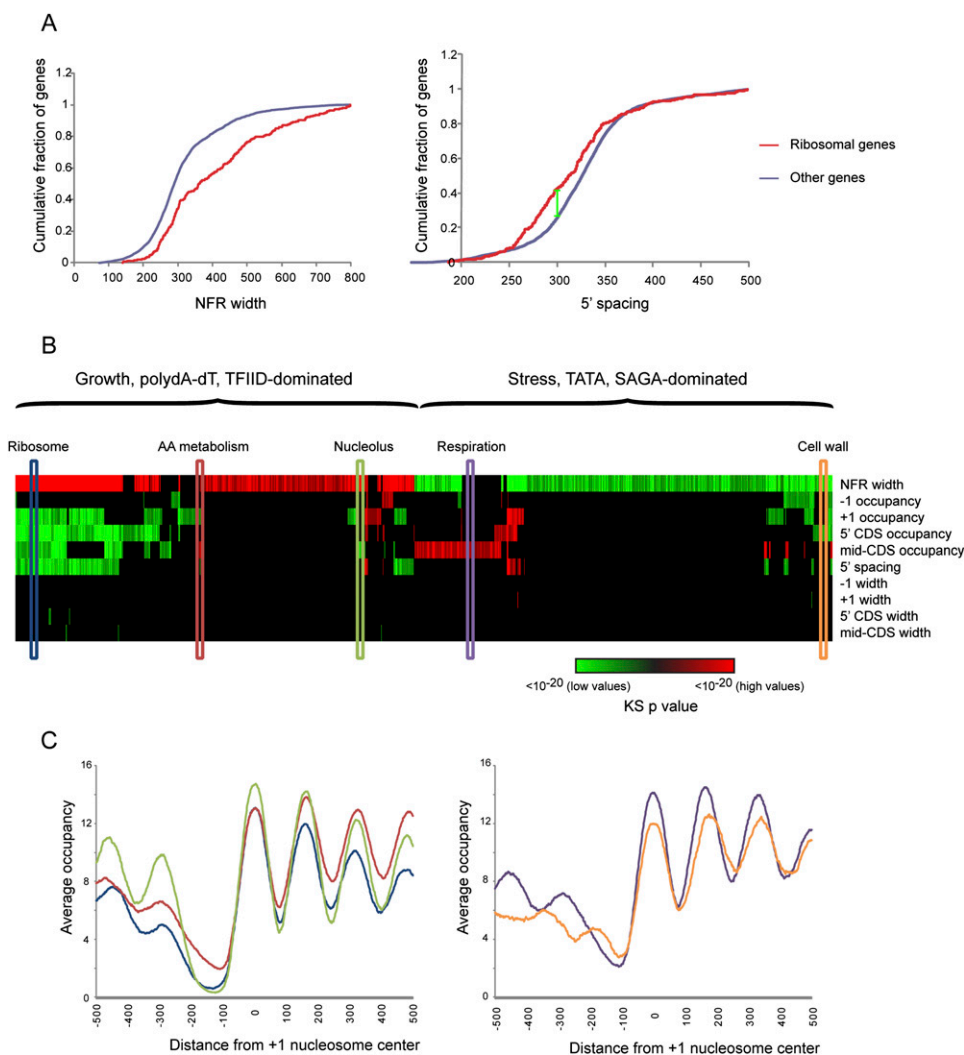


Figure 2. Different promoter types are differently packaged. (A) Cumulative distribution function (CDF) plots for two significant Kolmogorov-Smirnov (KS) enrichments. The gene set of 270 ribosomal genes is enriched for long NFRs (*left*), and close +1 to +3 nucleosome spacing (*right*). For example, 45% of ribosomal genes have 5' nucleosome spacing of <300 bp (green line), whereas only 25% of all genes have this spacing. (B) Enrichment of high or low values of various parameters for a set of promoter types. Various parameters such as +1 nucleosome occupancy (listed at *right*) were extracted for all yeast promoters. Each of the gene sets previously gathered (Dion et al. 2007; Wapinski et al. 2007) was tested (using KS test) for significantly high or low values of the various parameters. Significant (FDR < 0.05) high values are shown in red, and significant low values are in green. Colors represent \log_{10} of the P -value of the KS enrichment (saturated at $P < 1 \times 10^{-20}$). Colored elongated boxes indicate gene classes evaluated in C. (C) Averages for five promoter types as indicated in B, aligned according to +1 nucleosome center. The y -axis represents average normalized nucleosome occupancy, in number of reads per million mappable reads.

example, within the “growth” class of genes, ribosomal and metabolic genes can be roughly distinguished by the occupancy of coding-region nucleosomes (Fig. 2B,C). While these distinctions appear subtle, it is important to note that they are of similar magnitude to physiologically relevant (Ihmels et al. 2005) changes observed in the chromatin packaging of mitochondrial ribosomal genes between *S. cerevisiae* and *C. albicans* (Field et al. 2009), suggesting that such small differences can, indeed, play roles in transcriptional control.

To explore the role of transcription in shaping chromatin architecture we analyzed the relationship between Pol II enrichment (Methods) at a given gene and the various chromatin parameters described above (Supplemental Fig. S7). NFR width was positively correlated with transcription rate, while +1 occupancy, mid-CDS occupancy, and +1 to +3 spacing were slightly anti-correlated with transcription rate. To determine the extent to which gene set enrichments from Figure 2A were driven by transcription level, we corrected each gene’s chromatin parameters to account for transcription rate (Methods) and repeated the KS enrichment analysis from Figure 2A (Supplemental Fig. S7D). The majority of enrichments from Figure 2B repeated after correcting for polymerase abundance (699 of 1001 retained, 302 lost, 42 gained; see Supplemental Table S2), indicating that different regulatory mechanisms are linked to different chromatin architecture of gene sets.

Analysis of nuclease titration levels

A striking aspect of our analysis is that the majority of nucleosomes were matched using templates with multiple ends, suggesting that the majority of nucleosome ends are partially trimmed during a typical MNase digestion. Furthermore, in comparing our data to published datasets we found global variation in the distribution of nucleosome widths between datasets (Supplemental Fig. S3B), as well as in the relative occupancy of various nucleosome classes (i.e., genome-wide averages of +1 occupancy vs. mid-CDS occupancy differs between our data and that of Shivaswamy et al. (2008) (data not shown). Indeed, a recent study using MNase titrations followed by q-PCR identified variation in quantitative MNase susceptibility across nucleosomes associated with *GAL* genes (Bryant et al. 2008). We therefore sought to more thoroughly explore the influence of digestion levels in nucleosome positioning and occupancy. We have previously reported little change in nucleosome maps as measured by tiling microarray when mononucleosomal DNA is isolated from an early digestion step with only ~40% mononucleosomal DNA (Yuan et al. 2005). However, dynamic range compression by microarrays might hide changes in a relative abundance of nucleosomes, and we were unable to obtain enough DNA for microarray analysis from less-digested (<40% mononucleosome) titration steps. Moreover, small changes in nucleosome segment lengths are difficult to detect with tiling microarrays.

Since limited digestion with trypsin has proven a valuable structural probe for proteins, we were also interested in whether the same might be true of limited nuclease digestion of chromatin. We therefore carried out a titration of micrococcal nuclease, and gel-purified mononucleosomal DNA from three different titration levels—underdigested (~15% mononucleosomes, “BY2”), typical digestion (~80% mononucleosomes, “BY10”), and overdigested (only mononucleosomal DNA, “BY15”) (Fig. 3A). We used Template Filtering to call nucleosome positions in our titration data. As expected, nucleosome length was correlated with digestion level

(Fig. 3B), with increasing digestion leading to shorter and shorter nucleosomes, presumably due to “chewing” of nucleosome ends by MNase.

Inspection of nucleosome maps for the three digestion levels revealed extensive similarities between the three maps. However, notable changes occur, particularly between underdigested and typical digestion (Fig. 3C). To globally assess differences between the different titration steps, we aligned genes by transcriptional start site (Fig. 3D) or stop codon (Fig. 3E) and averaged data from all genes at the three different titration levels. At the 5’ ends of genes we found an anticorrelation between +1 nucleosome occupancy and digestion level as expected. +1 nucleosomes are most abundant in underdigested chromatin and least abundant in overdigested chromatin. This is seen both in TSS-aligned averages of all genes (Fig. 3D), as well as in systematic analysis of changes in relative nucleosome occupancy calls (Supplemental Fig. S8).

The loss of promoter-proximal nucleosomes during digestion was even more pronounced for –1 nucleosomes, where underdigested chromatin (BY2) showed high levels of the –1 (and a resulting decrease in the width of the average NFR), whereas this nucleosome was less abundant or completely missing in BY10 and BY15 (Fig. 3C, red bar). A similar effect was observed at the 3’ NFR (Fig. 3E). The presence of an easily digested nucleosome over the NFR was most commonly observed at the long NFRs associated with highly expressed genes (red bar), and can easily be seen in averaged data for genes with the highest levels of RNA polymerase (Fig. 3F). Thus, consistent with recent studies in *Drosophila* (Henikoff et al. 2009) and human (Jin et al. 2009) cells, we find that at least some of the “nucleosome-free” region seen in typical nucleosome mapping studies corresponds to a loosely bound (as determined by salt extraction in those studies), easily digested (seen here) nucleosome. Overall, nucleosome occupancy levels are most even in BY10, which is the digestion level we typically use for nucleosome mapping (Yuan et al. 2005) and histone modification mapping (Liu et al. 2005).

Together, the results of the titration series suggest the presence of easily digested nucleosomes or other protein complexes at the promoters of highly expressed genes and point toward the necessity of knowing the extent of digestion when comparing nucleosome maps from different labs or different experiments.

The role of RNA polymerase in chromatin structure

A number of recent studies have claimed that intrinsic affinity of various genomic sequences for the histone octamer accounts for much, or most, of the chromatin structure observed in vivo in yeast (Ioshikhes et al. 2006; Segal et al. 2006; Peckham et al. 2007; Field et al. 2008; Kaplan et al. 2008; Yuan and Liu 2008). However, experimental determination of nucleosome positioning after in vitro reconstitution revealed that intrinsic preferences can almost entirely be ascribed to the role of poly(dA/dT) in excluding nucleosomes, with little additional translational positioning information encoded in the genome (Kaplan et al. 2008; Zhang et al. 2009). The huge discrepancy between in vitro sequence preferences and in vivo nucleosome positioning is likely to result from the action of numerous factors, most notably protein complexes in vivo that move nucleosomes from their preferred positions, such as the ATP-dependent chromatin remodeler Isw2 (Whitehouse and Tsukiyama 2006; Whitehouse et al. 2007). Almost certainly the most widespread of these *trans*-acting factors is Pol II, as ~2/3 of the yeast genome codes for proteins, and combining distributions of RNA abundance (Yassour et al. 2009) with typical absolute

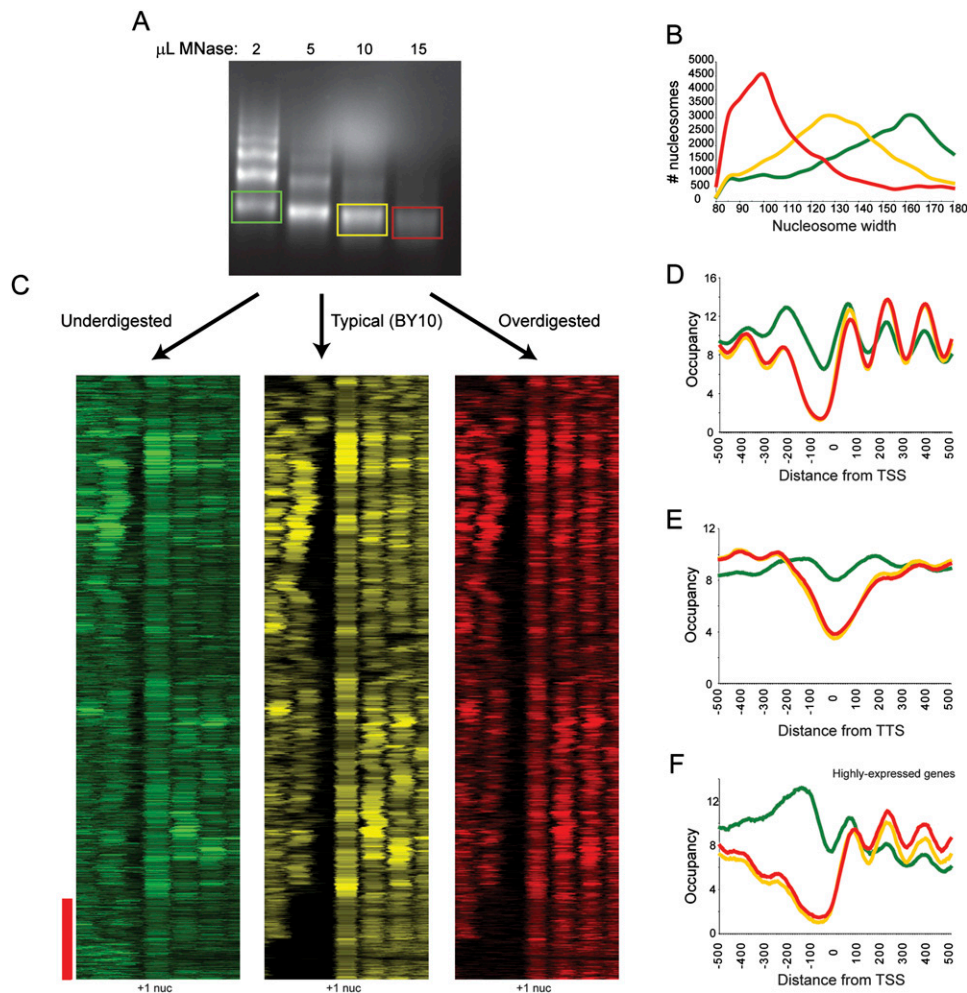


Figure 3. Effects of MNase level on chromatin structure. (A) Mononucleosomal DNA was isolated from ladders from three different MNase titration levels, and sequenced by Illumina sequencing. (B) Data from titration series was subjected to template filtering to generate nucleosome calls. Width distributions for nucleosomes from the three titration steps are plotted. Green, yellow, and red correspond to under-, mid-, and over-digested chromatin, respectively. (C) Data for under- (green), mid- (yellow), and over- (red) digested chromatin is shown in cluster view. Genes are aligned using BY10 +1 nucleosome center (indicated); all three clusters have genes ordered by clustering for BY10 data. Red bar indicates genes with wide NFRs in mid- and over-digested chromatin (largely highly expressed genes such as ribosomal genes), which are partially filled in underdigested chromatin. (D) TSS-aligned nucleosome occupancy data for all genes. (E) Stop-codon-aligned nucleosome occupancy for all genes. (F) As in D, but only for genes with Pol II ChIP occupancy >1, top 7% of genes.

mRNA abundances (Iyer and Struhl 1996) and half-lives (Wang et al. 2002) indicates that well over half of the yeast genome is likely transcribed at least once during a given cell cycle. The passage of RNA polymerase disrupts DNA–histone contacts (Wasylyk and Chambon 1980), leading us to ask how RNA polymerase globally affects chromatin structure in vivo.

We examined the effects of RNA polymerase on chromatin structure by using the *rpb1-1* yeast strain, which contains a temperature-sensitive allele of the gene encoding the large subunit of Pol II (Nonet et al. 1987). To identify both the early, as well as longer term effects of RNA polymerase deactivation, we performed MNase-seq at 0, 20, and 120 min after shifting these cells from 25°C to 37°C. Interestingly, we found that despite minimal change in cell density, increasing amounts of MNase were required to generate similar nucleosome ladders (Supplemental Fig. S9A–C) as the time course progressed, indicating a global role for RNA polymerase in increasing overall chromatin accessibility. Consistent

with the increased MNase required, we found that nucleosome length decreased over the time course (Supplemental Fig. S9D). Importantly, none of our conclusions below are affected by the titration level, as the inferred effects of polymerase loss do not match effects of overdigestion (see below).

Averaging genes by TSS for the three time points reveals two effects of polymerase loss on nucleosome positioning (Fig. 4A,B; Supplemental Fig. S10). First, NFR width decreases over time, largely because of gains in nucleosome occupancy at the –1 position (Venters and Pugh 2009). This was not a digestion artifact at later time points, as the nucleosome length distribution at 120 min was consistent with MNase overdigestion (Supplemental Fig. S9D), whereas increased –1 nucleosome occupancy is a general property of underdigested chromatin (Fig. 2). Second, coding-region nucleosomes shift downstream over time. This latter observation is interesting, as it is consistent with predictions from biochemical studies: The process of RNA polymerase transiting a nucleosome in

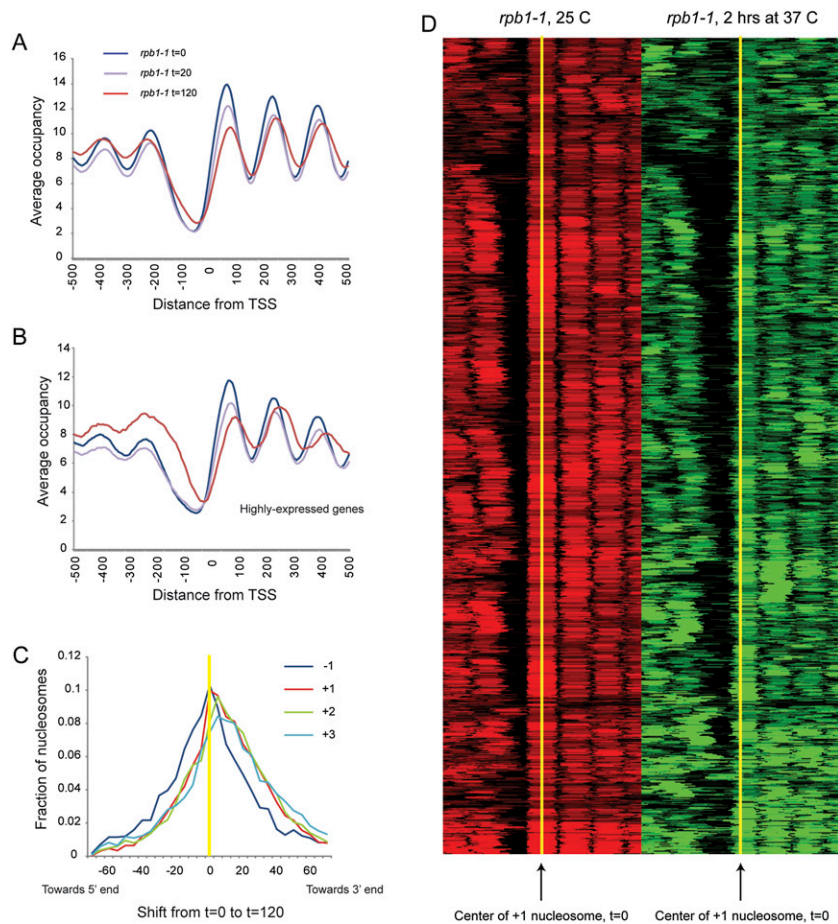


Figure 4. Effects of RNA polymerase on chromatin structure. (A) Nucleosomes were isolated from *rpb1-1* yeast grown at 25°C, and shifted to 37°C for 20 or 120 min. Data are presented in TSS-aligned average. (B) As in A, but for highly expressed genes. (C) Nucleosomes over genes shift downstream upon Pol II loss. For each indicated nucleosome type (−1, +1, +2, +3) we plot the distribution of center-to-center distances between the nucleosome calls at 0 and 120 min after Pol II inactivation. We find that 43% of −1 nucleosomes, 59% of +1 nucleosomes, 61% of +2 nucleosomes, and 60% of +3 nucleosomes shift away from the NFR. (D) Global view of +1 nucleosome shifts during Pol II inactivation. Nucleosome calls for all promoters with a downstream +1 nucleosome shift are shown as a heatmap, aligned by the center of the +1 nucleosome (yellow) before Pol II inactivation (red). After 2 h of Pol II inactivation (green), downstream shifts of these 59% of +1 nucleosomes are apparent.

vitro results in transient dissociation of 5' DNA from the octamer, followed by recapture of upstream DNA on the octamer surface, resulting in a predicted retrograde shift of the octamer relative to polymerase movement (Studitsky et al. 1994, 1997).

We investigated nucleosome position shifts by plotting the direction of nucleosome shift between pairs of time points during the temperature shift (Fig. 4C,D). Nucleosome shifts over coding regions were clearly biased in the downstream direction, consistent with the averaged view in Figure 4A. Importantly, −1 nucleosome shifts were not biased in either direction. This small (~10 bp on average) downstream shift is specific, as comparisons between other pairs of chromatin datasets (such as pre- and post-heat shock) show distributions of nucleosome shifts centered on zero (Supplemental Fig. S11). To determine whether nucleosome shifts preferentially occurred over particular classes of genes, we tested gene classes for significant deviation from the average shift of the −1, +1, +2, or +3 nucleosome by the KS statistic. Again, very highly expressed gene classes, such as those encoding ribosomal proteins or amino acid metabolism genes, exhibited more dramatic nucle-

osome shifts from 20 to 120 min (data not shown). Highly expressed genes are generally down-regulated during heat stress (Gasch et al. 2000), but nucleosome shifts at ribosomal genes are unlikely to be a consequence of heat shock-induced changes—reanalysis of the heat shock data from Shivaswamy et al. (2008) do not recapitulate the shifts we observe here (Supplemental Fig. S11A).

Why do nucleosomes shift after loss of polymerase? As noted above, RNA polymerase is a major factor in nucleosome movement and eviction in vivo, and the in vivo nucleosome positions over highly expressed genes deviate from in vitro nucleosome preferences to a greater extent than they do over poorly expressed genes. Thus, we compared our data with the in vitro data from Kaplan et al. (2008), reasoning that loss of RNA polymerase might allow chromatin to relax to more closely match local thermodynamic minima. We noticed that nucleosomes at many promoters, indeed, more closely match in vitro preferences after 2 h of polymerase inactivation (Fig. 5A).

To generalize this result, we calculated the correlation between in vitro nucleosome assembly data (Kaplan et al. 2008) and in vivo nucleosome positioning for 1-kb windows at the 5' ends of genes (Fig. 5B). At $t = 0$, correlation coefficients centered around 0.3, consistent with the ability of in vitro assembly to highlight NFRs (Sekinger et al. 2005; Kaplan et al. 2008; Zhang et al. 2009). After 2 h of polymerase inactivation, the distribution of correlations shifted to a higher value of ~0.5, indicating that RNA polymerase does help maintain nucleosomes in thermodynamically unfavored

locations in vivo. Of course, much of this is due to the above-mentioned role of transcription in −1 nucleosome eviction (Fig. 5C). We also asked whether lateral repositioning of nucleosomes upon polymerase loss resulted in relaxation to thermodynamically preferred positions. Strikingly, distances between +1 nucleosomes and the nearest in vitro occupancy peak decreased after polymerase inactivation (Fig. 5D; Supplemental Fig. S12), consistent with the hypothesis that both nucleosome eviction and sliding by RNA polymerase antagonize the thermodynamically preferred chromatin state.

Discussion

Genome-wide mapping of nucleosome positions in *S. cerevisiae* has been a tremendously productive method for illuminating the principles underlying chromatin structure and function. Deep sequencing methods have multiple advantages over tiling microarrays for genomic localization studies such as nucleosome mapping studies, including single-nucleotide resolution, expanded

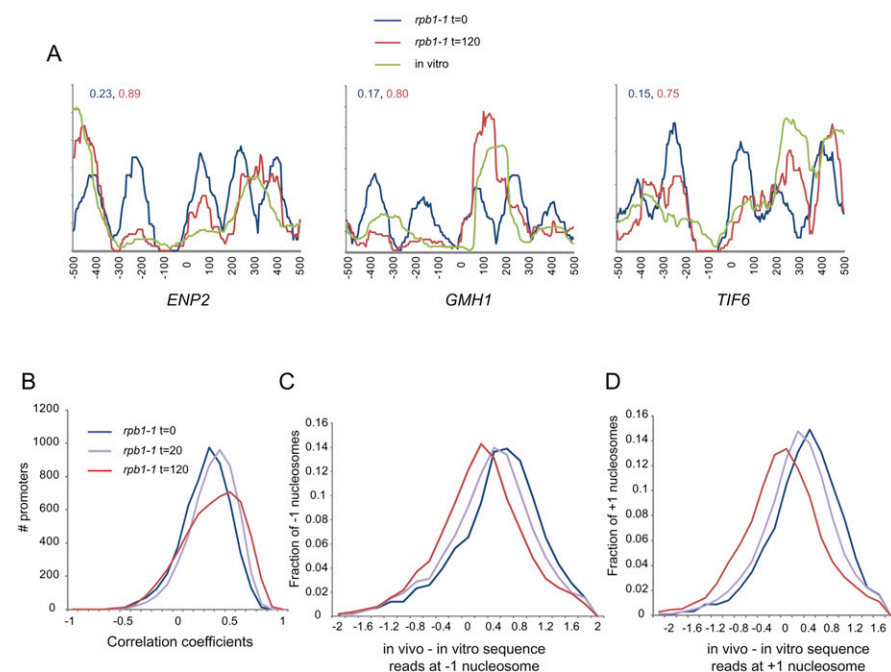


Figure 5. Nucleosomes relax toward in vitro preferred locations after Pol II loss. (A) Three examples of promoters where data from Pol II inactivation matches in vitro nucleosome assembly data better than data from before Pol II inactivation. Shown are extended read coverage along 1000 bp centered on TSS. Numbers shown in the inset are correlations between in vitro coverage and $t=0$ (blue) and $t=120$ (red) in vivo coverage. (B) Promoter chromatin architecture globally shifts toward in vitro preferences as polymerase is inactivated. Extended read coverage along the 1 kb centered on the TSS was extracted for all genes, and correlation coefficients were calculated to equivalent data for in vitro nucleosome reconstitution experiments (Kaplan et al. 2008). Histograms show a global shift of promoters toward the in vitro nucleosome pattern. (C) Normalized occupancy of -1 nucleosome better matches in vitro data after polymerase loss. For all -1 nucleosomes (called at $t=0$ or at $t=120$), the difference between in vivo normalized occupancy and in vitro normalized occupancy at the center of the in vivo nucleosome were calculated and presented as a histogram. (D) As in C, but for all $+1$ nucleosomes.

dynamic range, and nearly whole-genome coverage. Here, we present a novel method for analyzing deep sequencing data for chromatin maps. Our method automatically extracts nucleosome position, occupancy, and width, and accounts for variability in end digestion by MNase.

Analysis of chromatin packaging across the yeast genome confirmed previously described aspects of yeast chromatin structure, including widespread 5' and 3' nucleosome-depleted regions, a dichotomy between stress and growth genes reflected in NFR width, increased nucleosome fuzziness distal to the NFR, and a subtle anticorrelation between coding-region nucleosome occupancy and transcription. We also identified finer distinctions between certain classes within the major stress/growth branches. Most interestingly, we found that $+1$ to $+3$ nucleosome spacing was significantly shorter over ribosomal genes than over other gene types (see below).

We also analyzed data from an MNase titration series, as different laboratories isolate nucleosomes from different MNase digestion levels (see, for example, Shivaswamy et al. 2008). Analysis of data from underdigested chromatin revealed the abundant presence of nucleosome-sized peaks in the "NFR." A recent analysis of *Drosophila* chromatin also identified nucleosomes in NFRs in a low-salt extraction from underdigested chromatin (Henikoff et al. 2009), and similar results hold in human cells (Jin et al. 2009). Does this material correspond to easily digested nucleosomes or to DNA protected from MNase by other proteins such as transcription

factors? Two lines of evidence support the former hypothesis. One, the equivalent material in *Drosophila* is associated with the histone variant H3.3, indicating the presence of histones at these locations (Henikoff et al. 2009). Second, we only find nucleosomes filling in NFRs that are larger than 140 bp, suggesting that these are bona fide nucleosomes. Interestingly, analysis of sequence motifs that exhibit occupancy differences include the CGCG motif recently shown to be bound by the Rsc3/30 subunits of the RSC chromatin remodeling complex (Badis et al. 2008), suggesting that the easily digested -1 peak might correspond to a nuclease-accessible RSC-remodeled nucleosome state.

As many features of yeast chromatin correlate with transcription rate, we mapped nucleosomes before and after inactivation of Pol II. We find that NFRs become shorter and shallower upon loss of Pol II, particularly at highly expressed genes, consistent with a previously described role for RNA polymerase in eviction of -1 nucleosomes (Venters and Pugh 2009). We also found a surprising general role for RNA polymerase in nucleosome sliding—nucleosomes over coding regions generally shifted away from the NFR upon loss of polymerase (Fig. 4). These results are consistent with the predictions from biochemical studies—in vitro, RNA polymerase is capable of transiting a nucleosome without evicting

histones, apparently by invading a nucleosome edge and then propagating a bubble of DNA around the octamer surface (Studitsky et al. 1994, 1997; Hodges et al. 2009). This retrograde nucleosome movement may play a role in the stereotyped $+1$ nucleosome positioning in vivo, which is not explained by intrinsic thermodynamic preferences as measured by in vitro nucleosome assembly (Kaplan et al. 2008; Zhang et al. 2009). We speculate that after assembly of a newly replicated DNA into nucleosomes, polymerase passage could be responsible for nucleosome shifts toward the NFR until the $+1$ nucleosome is as close to either poly(A)s or to the preinitiation complex as physically possible. This retrograde nucleosome movement may also play a role in the surprisingly tight packing of nucleosomes over highly transcribed ribosomal genes and, indeed, loss of polymerase results of relaxation of the $+1$ to $+3$ spacing at these genes (Supplemental Figs. S10, S13).

This interpretation must be tempered by dynamic studies in yeast, which indicate that some nucleosomes (particularly $+1$ nucleosomes) are rapidly exchanged during G_1 (Dion et al. 2007; Jamai et al. 2007; Rufiange et al. 2007)—how is it that translational effects of transcription are observed on nucleosomes, given that nucleosomes are often rapidly exchanged (in some cases many times per cell cycle)? We consider two of many possible ways to reconcile these results. First, given that current locus-specific dynamic exchange measurements rely on transcriptional activation of tagged histones, there is a lower bound (~ 15 – 30 min) for the fastest exchange rates measurable. However, if the fastest exchange

rates are indeed on the order of ~15 min, then for many genes a high proportion of cells in a population will have had polymerase pass through the gene since the last histone turnover cycle, resulting in a detectable polymerase-driven retrograde shift in the population measurement despite ongoing nucleosome replacement. Second, we do not currently know the extent of correlation in the dynamics of adjacent nucleosomes at the single-gene level. In other words, at a highly transcribed gene with high levels of histone replacement throughout the gene body, does +1 eviction affect +2 eviction? If these do not always co-occur, then retrograde shifts of surrounding nucleosomes could provide a local “memory” of prior polymerase passage, such that a leftward-shifted +2 would constrain the replacement location for a replaced +1 nucleosome.

Finally, our results bear on the relationship between thermodynamic sequence preferences and *in vivo* chromatin structure. Dramatic claims have been made regarding the extent to which genomic sequence dictates the positioning of nucleosomes in the cell (Segal et al. 2006; Kaplan et al. 2008), although most studies find the effects of sequence on chromatin architecture to be modest (Ioshikhes et al. 2006; Peckham et al. 2007; Yuan and Liu 2008). While *in vitro* chromatin assembly correlates well with *in vivo* nucleosome positions (Kaplan et al. 2008; Zhang et al. 2009), this almost entirely results from the depletion of nucleosomes over poly(A) and related sequences (Kunkel and Martinson 1981; Drew and Travers 1985; Iyer and Struhl 1995; Sekinger et al. 2005). Our results confirm the expected role for RNA polymerase in movement of nucleosomes away from thermodynamically preferred positions.

Together, these results further emphasize the role for RNA polymerase in shaping the chromatin landscape of the genome and point toward the difficulty in disentangling cause and effect in the relationship between chromatin and transcription.

Methods

Nucleosome isolation

Yeast culture, fixation, and MNase titrations were carried out as previously described (Yuan et al. 2005). For *rpb1-1* temperature shifts, cells were grown to an OD of 0.6 in YPD at 25°C, then culture aliquots were immediately shifted to 37°C by addition of an equal volume of YPD at 49°C. After recovery of digested DNA, mononucleosomal was gel-purified and subjected to Illumina sequencing as described in Shivaswamy et al. (2008).

Template filtering algorithm

Using a sliding window across the genome, we cross-correlate each position with a pair of templates, one matching the forward reads and one the reverse reads. We enumerate all 7×7 possible combinations of forward and reverse templates. We repeat this scan with different spacing between both templates to capture over- and underdigestion of the ~146-bp nucleosomal DNA fragments. As a result, we obtain a correlation “heat map” for each pair of templates containing the correlation coefficient for each center position and width. Next, we search for local maxima points within this “heat map”; each maxima point is a potential nucleosome at a given position with a specific width. Finally, to assemble the final set of nucleosomes, we are using a greedy approach to select the best assignment of nucleosomes under overlapping constraints. Potential nucleosomes are sorted according to correlation score and occupancy, and are then selected to the final set allowing maximum overlap of 40% between adjacent nucleosomes.

Selecting representative templates

To generate a variety of templates that represents the prototypical distributions of reads at nucleosome ends, we first applied our method using a Gaussian-shaped template and obtained a preliminary map of nucleosome predictions. We aligned all predicted nucleosome ends and created a matrix of read patterns using a window of 80 bp flanking the edges. Next, we clustered this matrix using k-mean clustering and selected seven representative templates that capture the range of template shapes observed.

Correcting chromatin parameters to account for Pol II enrichment in KS tests

We represent each gene as a vector of chromatin properties (i.e., +1 nucleosome occupancy, NFR width, mid-CDS occupancy, etc). Using a compendium of experimental gene annotations we previously collected (Dion et al. 2007; Wapinski et al. 2007), we compared the distribution of each chromatin parameter for each gene set vs. the background. We discovered 1001 gene sets with at least one enriched chromatin property. To correct the genes’ properties vectors for RNA polymerase levels, we plotted RNA polymerase occupancy (measured by microarray as in Steinmetz et al. 2006; T Kim, S Buratowski, A Novogrodski, M Yassour, N Friedman, and O Rando, in prep.) vs. each chromatin property and calculated the LOESS curve for each property (Supplemental Fig. S7A–C). We then subtracted the smoothed LOESS curves from each gene, obtaining new chromatin properties vectors that represent the distance of each property from the LOESS curve. We repeated the K-S enrichment with these new vectors.

Source code

Source code is available at <http://compbio.cs.huji.ac.il/NucPosition>.

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Author contributions: O.J.R. designed the experiments in discussion with N.F., and A.H. carried them out. A.W. and M.Y. developed the analysis methodology and analyzed the data. O.J.R., A.W., and N.F. wrote the manuscript.

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