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Nucleosomes Shape DNA Polymorphism and Divergence

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Abstract
An estimated 80% of genomic DNA in eukaryotes is packaged as nucleosomes, which, together with the remaining interstitial linker regions, generate higher order chromatin structures [1]. Nucleosome sequences isolated from diverse organisms exhibit ~10 bp periodic variations in AA, TT and GC dinucleotide frequencies. These sequence elements generate intrinsically curved DNA and help establish the histone-DNA interface. We investigated an important unanswered question concerning the interplay between chromatin organization and genome evolution: do the DNA sequence preferences inherent to the highly conserved histone core exert detectable natural selection on genomic divergence and polymorphism? To address this hypothesis, we isolated nucleosomal DNA sequences from Drosophila melanogaster embryos and examined the underlying genomic variation within and between species. We found that divergence along the D. melanogaster lineage is periodic across nucleosome regions with base changes following preferred nucleotides, providing new evidence for systematic evolutionary forces in the generation and maintenance of nucleosome-associated dinucleotide periodicities. Further, Single Nucleotide Polymorphism (SNP) frequency spectra show striking periodicities across nucleosomal regions, paralleling divergence patterns. Preferred alleles occur at higher frequencies in natural populations, consistent with a central role for natural selection. These patterns are stronger for nucleosomes in introns than in intergenic regions, suggesting selection is stronger in transcribed regions where nucleosomes undergo more displacement, remodeling and functional modification. In addition, we observe a large-scale (~180 bp) periodic enrichment of AA/TT dinucleotides associated with nucleosome occupancy, while GC dinucleotide frequency peaks in linker regions. Divergence and polymorphism data also support a role for natural selection in the generation and maintenance of these super-nucleosomal patterns. Our results demonstrate that nucleosome-associated sequence periodicities are under selective pressure, implying that structural interactions between nucleosomes and DNA sequence shape sequence evolution, particularly in introns.

Introduction
Sequence-dependent differences in the physical properties of DNA influence its associations with the histone core, as well as the kinetics of nucleosome assembly and stability [2–9]. One of the most generalizable sequence affinities of the histone octamer is the periodic variation of dinucleotide frequencies across nucleosomal DNA. Alignments of nucleosomal sequences from diverse eukaryotes display a prominent ~10 bp periodic enrichment of AT-rich dinucleotides, along with an anti-correlated periodicity of GC-rich dinucleotides [8,10–14]. The ~10 bp spacing of AA/TT dinucleotides generates intrinsically curved DNA molecules with increased nucleosome binding affinity [5,8,14–17]. Peaks of AA/TT frequency are found specifically over positions where the minor groove bends interiorly, whereas GC dinucleotides peak where the major groove is facing the histone core. Structural data suggest that DNA shape, in particular the narrowing of the minor groove and the associated lowering of its electrostatic potential at AT-rich sequences facilitate contacts with key histone arginines [9,18,19]. GC dinucleotides contract the major groove, which also facilitates the tight winding of DNA around the core [9,20]. Although these broadly conserved dinucleotide patterns have been cited as evidence for a genomic “code” for nucleosome positioning [10], the role of sequence in nucleosome function remains contested and unresolved [6,8,21,22]. Correlation between in vitro and in vivo nucleosome maps in yeast may reflect the influence of the inherent sequence preferences of the histone core on nucleosome positioning [7,23]. However, strong experimental evidence suggests that trans-acting factors (e.g. RNA polymerase II, transcription factors and ATP-dependent remodelers) are central to establishing nucleosome positions along genomic DNA (translational positions), particularly in genic regions, with sequence providing a weaker contribution [7,8,21,24,25]. In cases where DNA sequence does impact translational nucleosome positions, its influence is largely attributed to GC content and anti-nucleosomal sequences, such as poly-dA/dT tracts, rather than dinucleotide patterns [5,8,25–27]. Dinucleotides are instead thought to play a distinct but integrally connected role in directing and preserving the ‘rotational positioning’ of nucleosomal DNA [8,9,20,28], which refers to the orientation of DNA relative to the core. Due to the structural constraints inherent to nucleosome formation, a given
Author Summary

In eukaryotic cells, the majority of DNA is packaged in nucleosomes comprised of ~147 bp of DNA wound tightly around the highly conserved histone octamer. Nucleosomal DNA from diverse organisms shows an anti-correlated ~10 bp periodicity of AT-rich and GC-rich dinucleotides. These sequence features influence DNA bending and shape, facilitating structural interactions. We asked whether natural selection mediated through the periodic sequence preferences of nucleosomes shapes the evolution of non-protein-coding regions of D. melanogaster by examining the inter- and intra-species genomic variation relative to these fundamental chromatin building blocks. The sequence changes across nucleosome-bound regions on the melanogaster lineage mirror the observed nucleosome dinucleotide periodicities. Importantly, we show that the frequencies of polymorphisms in natural populations vary across these regions, paralleling divergence, with higher frequencies of preferred alleles. These patterns are most evident for intronic regions and indicate that non-protein coding regions are evolving toward sequences that facilitate the canonical association with the histone core. This result is consistent with the hypothesis that interactions between DNA and the core have systematic impacts on function that are subject to natural selection and are not solely due to mutational bias. These ubiquitous interactions with the histone core partially account for the evolutionary constraint observed in unannotated genomic regions, and may drive broad changes in base composition.

Translational position in the genome will assemble with a particular rotational alignment. This determines which bases face the nucleosome interior and exterior, and also the positioning of the major and minor grooves relative to the core. Nucleosomes tend to occupy translational genomic positions which are offset by ~10 bp increments [13,28,29]. Thus, due to the helical structure of DNA, with ~10.4 bp per turn, the rotational orientation of DNA relative to the core is thought to be unchanged as nucleosomes assume new favored translational positions (Figure 1A). This 10 bp incremental movement leaves the exposure of sites at the surface unchanged [20,28], and is in agreement with the reported step size of many chromatin remodelers [30]. By influencing the rotational positioning of DNA relative to the histone core, nucleotide changes at particular nucleosome positions (or in flanking regions) could have diverse functional impact, for example on nucleosome assembly, stability, remodeling efficiency, RNA and DNA polymerase processivities and transcription factor binding site access. However, despite considerable evidence that dinucleotide patterns impact nucleosome positioning and dynamics in vitro, in vivo evidence of function has remained elusive.

One approach to discovering function is to look for evidence of natural selection in sequence polymorphism (variation within species) and divergence (variation between species). Individual mutations influencing histone-DNA interactions may have only slight, undetectable phenotypic effects in the laboratory; in contrast, the associated fitness consequences in large natural populations can strongly shape rates of divergence and levels of polymorphism over many generations [31]. Of course, strongly selected variants will go to fixation quickly and be maintained in very high frequency against the weaker force of mutational reversion. However, observations of extensive DNA sequence polymorphism and divergence throughout the genome, including nucleosomal sequences, indicate that such systematic selection is not dominating stochastic effects (mutation, genetic drift, and variation in selection coefficients) in the evolutionary dynamic.

Analysis of codon bias suggests that at equilibrium between selection, mutation and genetic drift, the ratio of the frequencies of two alternative synonymous codons throughout a single genome can be used to estimate the direction and magnitude of selection [32]. The action of natural selection can be inferred when synonymous codon pairs exhibit a strong “bias” towards one state relative to the other. This analysis extends to the distribution of polymorphic allele frequencies in genomes sampled from natural populations [33–35]. A similar approach can be applied to alternative nucleotides at particular positions within nucleosomal sequences. As the magnitude of selection increases, the expected frequency of preferred alleles increases. Consequently, the distribution of SNP frequencies (or “site frequency spectrum”) at a given nucleosome position (analogous to a synonymous SNP) is expected to shift towards relatively higher frequencies of preferred alleles.

If the observed dinucleotide patterns reflect selectively favored states, ancestrally unpreferred base pairs across nucleosomes should diverge towards the “preferred” state along species lineages. Further, if the “preferred” divergence patterns reflect the average impact of natural selection, then frequencies of polymorphisms in natural populations should be more skewed at sites experiencing stronger selection. “Unpreferred” variants, specifically substitutions or polymorphisms away from favored nucleotides, such as substitution of an ancestral A with a G at nucleosome positions which are systematically enriched for AA dinucleotides, should diverge more slowly and be rarer when polymorphic in the population. In contrast, “preferred” variants, such as substitution of an ancestral A with a G at positions of enriched for the GC dinucleotide, should diverge more rapidly and be more common when polymorphic.

At the lower resolution of an entire nucleosome and its nearby flanking regions, both divergence and polymorphism are observed to vary [36–39], but evidence of a role for natural selection in the underlying evolutionary dynamics remains sparse [40–43]. Studies of human SNPs [37,38] and divergence in humans, yeast and medaka [36,38,39,43] show that both expected heterozygosity and divergence between species are elevated near the central dyad and depressed in the adjacent linker regions, though these patterns appear to differ by substitutional pathway [38]. One possibility is that patterns of variation relative to nucleosomes derive from nucleosome-specific mutational biases. This could result from suppression of mutation by a protective aspect of nucleosome occupancy [44], or it could arise from an interaction between the histone core and DNA damage recognition or repair mechanisms [45–48]. Of course, natural selection mediated via DNA-nucleosome interactions may also strongly reshape the patterns of SNP variation and divergence between taxa [40–43]. Analysis of the site frequency spectrum promises to distinguish between these two alternatives.

The whole-nucleosome-resolution analyses considered above cannot leverage the specific structural predictions of dinucleotide interactions with the core and their strong mechanistic implications. Examination of polymorphism and divergence at each base pair position across the nucleosomal DNA opens a rich and precise view, as well as powerful tests of alternative mechanisms such as biased mutation and natural selection. We report the discovery of fine-scale periodicities in inter- and intra-species sequence variation relative to nucleosomes and discuss their implications for the role of natural selection mediated through nucleosome function. Our analysis of DNA sequence polymorphism and divergence across isolated nucleosomal fragments from D. melanogaster embryos...
reveals that nucleosomal sequences are diverging towards “preferred” nucleotides. Regions where the minor groove is interior are becoming more AT-rich, and regions where the major groove is interior are becoming more GC-rich along the melanogaster lineage. Using a new index for quantitating the frequency spectrum ($A_2$), we identify clear signals associated with natural selection, which parallel the observed periodicities in divergence. This selection is strongest in intronic regions, where nucleosome assembly and positioning are expected to have greater functional impacts. These findings support the hypothesis that the widely observed sequence affinities of the core octamer have functional consequences that are subject to natural selection. Given the dominant role of nucleosomes in the packaging of the genome and their conserved sequence preferences, their interactions may broadly shape the sequence of melanogaster and other genomes.

**Results**

**Periodicity in dinucleotide frequencies in *D. melanogaster***

To investigate the impact of nucleosomes on DNA sequence variation, we isolated nuclei from *D. melanogaster* embryos, performed Micrococcal nuclease (MNase) digestion, and used paired-end sequencing to position fragments on the genome (Figure S1). Previous studies in *Drosophila* identified a range of periodic dinucleotides in association with nucleosomes [10,11]. Our collections of 276,614 intergenic and 270,998 intronic autosomal 147 bp nucleosomal fragments (hereafter n147, Tables S1 and S2) cover 68.5% of the unique intergenic and intronic euchromatic autosomal genome and display a ~10 bp periodicity for many dinucleotide frequencies (Figures 1 and S2). In these and subsequent analyses, the 5'-3' sequence from bases -73 to -1 were joined to the reverse complement of bases 1 to 73, to reflect the dyad symmetry of the nucleosome (see Materials and Methods). AA, TT, and GC showed the strongest periodicity of WW and SS (where $W=A^1T$, $S=G^1C$) dinucleotide pairs, respectively (Figure 1B). These same dinucleotides show a distinct overrepresentation in the non-coding regions of the genome as a whole (Figure 1C). As noted in previous studies, AA and TT are similar periodic and occur where the minor groove is interior (at superhelix locations, SHL, ±(0.5); where i is 0, 1,...6). However, noticeable differences between the distributions are apparent. For example, the frequency of TT displays a distinctly smaller peak at ~SHL 4.5, and AA frequency displays a stronger drop at ~SHL 2 (Figure 1B). GC frequency across n147 regions is anti-correlated with AA/TT and is characterized by a prominent upward concavity (Figure 1B). These dinucleotide periodicities extend well beyond n147 edges into linker regions, consistent with the proposed translational step size of 10 bp.

Upon examination of the dinucleotide frequencies flanking aligned n147 regions, we discovered an additional large-scale pattern in AA/TT and GC dinucleotide frequencies (Figure 1D). This ~180 bp periodic variation in frequency tracks with overall nucleosome “occupancy” in the regions flanking the n147. Average AA/TT frequencies (Figure 1D) and overall A/T frequencies (Figure S3) are higher in regions of greater nucleosome “occupancy” and lower in putative “linker” regions. Thus, the AA/TT sequence features that facilitate nucleosome formation are enriched over regions with higher nucleosome “occupancy.” Conversely, GC frequency (and overall G/C frequencies, Figure S3) peaks at the periphery of more nucleosome-dense regions and in “linker” regions. These surprising “super-nucleosomal” periodicities extend the observed n147 patterns to flanking multi-nucleosomal arrays, and suggest a contribution of sequence to translational positioning. Consistent with chemical mapping of nucleosomes, this result suggests that the observed experimental correlation between MNase nucleosome “occupancy” and GC content [1,8,22,26,49,50] reflects differential recovery, rather than positional preference [23,51].

**Divergence along the melanogaster lineage mirrors periodic nucleosomal base preferences**

If variations in dinucleotide frequencies relative to nucleosomes result from accumulated sequence divergence, we expect substitution patterns to parallel the observed base preferences. However, the timescale(s) at which these patterns evolve is unknown. Lineage-specific or “polarized” divergence is the proportion of nucleotide sites that are different in melanogaster while identical in its sister taxa *simulans* (most recent common ancestor 2.5 MYA) and the proximate outgroup *yakuba* or * erecta*, 6–7 MYA, see Materials and Methods). Overall genomic divergence on the melanogaster lineage shows a marked excess of G→A (inferred ancestral G, derived A in melanogaster) and C→T (ancestral C, derived T) substitutions compared to A→G and T→C (Figure 2A). This is in agreement with earlier estimates of divergence on the melanogaster lineage [32,53] and with the observed two-fold greater mutation rate [34]. We next considered the average divergence at each site across n147 regions, normalized for underlying base frequencies. This analysis revealed a striking ~10 bp periodicity in transitions (GC→AT and AT→GC) for two estimates of divergence; per-n147 in Figure 2B is weighted by the redundancy in the n147 set, while per-site in Figure S4 weights each site equally. Rates for GC→AT and AT→GC are anti-correlated and track with underlying dinucleotide frequencies. Thus, ancestral GC bases are more likely to become AT in nucleosomal regions where AA/TT dinucleotides are in higher frequency, and AT bases are more likely to become G or C at sites where GC is enriched. GC→AT divergence also shows a marked curvature, with a peak at the dyad axis.

Given the substantial variation in individual substitution rates, we next examined specific pathways to determine their relative contributions. Of all pathways, G→A, C→T, A→G and T→C...
exhibited the most obvious periodicities in divergence (Figure 2C; see Discussion). In some cases, divergence patterns reflect the subtleties observed for dinucleotide frequency patterns. For example, C→T rates are less peaked at SHL 4.5, the location of the lowest peak in TT frequency (Figure 2C). C→T also shows a greater difference in rates between the n147 periphery and linker regions (compared to G→A). Interestingly, for each ancestral base, the periodicities of substitutions that do not change GC content,
appear weaker, perhaps due to both scaling and weaker signal to noise (Figure 2C). A subset of non-overlapping n147 regions showed similar patterns (Figure S5A).

When mapped onto the DNA from the nucleosome structure [55], peaks of intergenic G→A divergence clearly occur within regions where the minor groove is interior and in contact with key arginines of the histone core (Figure 2D). Note also the higher G→A divergence toward the central axis, as reflected by the downward concavity in Figure 2C. This is consistent with analyses of the impact of sequence variation on nucleosome structure, which identified this central region of H3/H4 interactions as most constrained [36]. Conversely, A→G substitution rates are highest in regions where the major groove is interior (Figure S5B). This pattern is consistent with established SS dinucleotide patterns [8,10–14] and the observation that GC rich sequences are disfavored for minor groove compression and favor narrowing of the major groove [9,18].

Divergence patterns should also reflect the observed nucleosome-scale periodicities in base and dinucleotide frequencies (Figures 1D and S3). To increase signal, we combined complementary substitutions, G→A and G→T (G→A:C→T) and A→G and T→C (A→G:T→C). Aligned n147 regions show substantially lower divergence rates than their immediate flanking sequences (Figure 2E). Rates drop to the local background within ~500 bp, following the skew of AA/TT dinucleotides (and overall AT content; Figures 1D and S3). In spite of this local variation in rates, due at least in part to MNase preferences (Figure S6), we observe a large-scale (180 bp) periodicity in G→A:C→T divergence surrounding intergenic n147 nucleosomal regions (Figure 2E). Introns showed a similar but weaker pattern, potentially due to the influence of flanking coding regions (Figure 2E). Any periodicity of the A→G:T→C divergence in flanking regions is less obvious (Figure S5C), at least partially due to a 50% lower rate of divergence and thus inherently weaker signal.

These large-scale patterns allow us to resolve general trends in divergence relative to nucleosome occupancy. We find that, on average, G→A:C→T changes along the melangaster lineage are fixed at higher rates across nucleosomes relative to linkers, mirroring underlying AA/TT dinucleotide frequencies. This is in apparent contrast to the report that the cytosine deamination mutational pathway (a major source of G→A:C→T transitions) and associated divergence is suppressed by nucleosome occupancy [44]. To clarify this discrepancy, we examined the interactions between divergence and “occupancy” of the n147 fragments, as estimated by depth of coverage by 142–152-bp nucleosomal fragments, n142-152. Indeed, we observe a negative correlation between this metric and all substitutional pathways (Table S3, Figure S7A). However, we note that n142-152 coverage is derived SNPs, such as G→GA (ancestral G and a derived, polymorphic A), exhibit clear periodicities that generally parallel divergence and nucleosomal dinucleotide frequencies (Figures S10, intergenic, and S11, intronic).

To systematically assess the periodicity in the frequency spectra we calculated a new index, Δₚ, (closely related to Tajima’s D [59]) across n147 regions for the Raleigh sample [65]. Where $\hat{p}$ is the frequency of a SNP in the sample and $\bar{\pi} = 2\hat{p}(1 - \hat{p})$ is the estimate of the heterozygosity, we define $\Delta_p$ as the average (per SNP) deviation in $\bar{\pi}$ from expectation under equilibrium between genetic drift and mutation to selectively equivalent alleles (see Materials and Methods). The “folding” of the frequency spectrum such that $\hat{p}$ is equivalent to $(1 - \hat{p})$ mitigates the impacts of errors in the inference of the ancestral state [66] and emphasizes variation in the midrange of $\hat{p}$. Weak positive selection is predicted to skew $\bar{\pi}$ toward higher values (more positive $\Delta_p$), while weak negative selection leads to more negative $\Delta_p$. Thus, systematic differences in selective forces at different positions across n147 regions should yield a pattern in $\Delta_p$ that parallels that observed for divergence. These patterns of $\Delta_p$ are superimposed on the observed genome-wide average negative skew [65] (Table S4) that can be attributed.
Figure 3. Periodicity in average SNP frequencies aligns with nucleosomal divergence and dinucleotide sequence preferences. (A) Smoothed average intergenic (black) and intronic (red) per-n147 frequency of A→G polymorphic sites in the Raleigh sample. (B) Average smoothed (black) and unsmoothed (grey) per-n147 Δε across n147 regions (±50 bp) for G→GA, C→CT, and A→AG polymorphic sites. Color-coded dinucleotide frequencies are included for reference. (C) Average smoothed n147 G→GA Δε for the Rwanda sample (orange) and across a non-overlapping subset of intergenic and intronic n147 regions (teal). Note that for both B and C the estimated divergence are outside the range of the plots near the edges of the n147. AA dinucleotide frequencies included for reference (green). (D) Combined intergenic and intronic (G→A;C→T; green) divergence and (G→A:C→CT; black) Δε surrounding intergenic n147 (±1 kb), n142-152 read depth in grey (axis in blue). Note that divergence and Δε, immediately flanking (±5 bp) the edges of the n147 are not plotted, since their true values are largely obscured by sequence bias of the MNase. (E) Average scaled intergenic and intronic per-n147 π, Δε and divergence for specific substitutions (see Table S4).

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Discussion

Histones are among the most ubiquitous and highly conserved eukaryotic proteins. Thus, it is not surprising that nucleosomal dinucleotide periodicities, which derive from key structural interactions between DNA sequence and the histone core, are shared widely across species. In spite of the near universality of these patterns among eukaryotes and decades of research, our understanding of their functional impact and evolutionary dynamics remain unsettled. In this work we examined genomic variation across regions defined by isolated nucleosomal DNA fragments. Our goal was to first determine if these regions showed interpretable variation in divergence between species, then to analyze population genomic variation for evidence of a role for...
natural selection in the generation and maintenance of nucleosome-associated sequence variation.

We find that divergence on the *melanogaster* lineage mirrors the sequence preferences of the histone core. This periodic variation in substitution rates across nucleosomal regions indicates that interior minor groove regions display more rapid substitution of AT for GC, and that AT base pairs in regions where the major groove faces inward are more likely to become GC rich. These striking patterns align directly with dinucleotide patterns that stabilize associations between DNA and the histone core, as documented in numerous biochemical and structural studies [5,8,15,18,19,56]. If nucleosome-bound regions are evolving toward the observed nucleosome sequence preferences, a key question is whether this is the result of mutational bias relative to the positioning of chromatin proteins, or whether it is the consequence of natural selection based on functional differences. The available depth of population data and our new index $\Delta_S$ allowed us to directly address this question. We find remarkable periodicities in $\Delta_S$ that parallel the observed patterns of divergence. The spectra of SNP frequencies across n147 regions are variable, with higher $\Delta_S$ when the inferred ancestral allele is unpreferred, and the derived allele is structurally favored. Therefore, we conclude that selection is, at least in part, driving the maintenance of nucleosome-associated sequence patterns on the *melanogaster* lineage.

If the fitness differences associated with histone-DNA interactions are largely arising from nucleosomal dynamics (assembly, disassembly, movement and modification) and rotational positioning of functional elements, then we can further hypothesize that transcribed (intronic) nucleosomal sequences should exhibit stronger periodicity than untranscribed (intergenic) nucleosomal sequences. Consistent with this hypothesis, correlations of lineage specific divergence and $\Delta_S$ with the relevant underlying dinucleotide frequencies are stronger for intronic sequences. Table 1 shows that in each case where a large difference between intergenic and intronic is apparent, it is the intronic that is larger. The two exceptions, G→A & AA and A→G & GC, are those where both correlations are among the highest. As might be expected given the longer timescale and greater number of variable sites, divergence correlates more strongly with dinucleotide frequencies than $\Delta_S$. Interestingly, Figures 2C and 3B show that these intronic vs. intergenic differences in correlation of divergence and $\Delta_S$ with dinucleotide patterns may be attributable to large deviations from expectation in specific regions of the nucleosomal sequences, while other regions follow the expected periodic patterns.

While natural selection is the most direct interpretation of these results, interactions between chromatin proteins and DNA damage and repair are well documented [44–48,73–76]. Contextually biased mutation (substitution) pathways could underlie the observed periodicities in nucleosomal divergence. However, Drosophila does not have a significant level of 5-methylcytosine [77], the deamination of which is thought to drive the strong contextual biases (NpCpG) in vertebrates [78]. Indeed, a recent genomic sequencing study of Drosophila mutation accumulation lines yielded no evidence for contextual biases [54]. Most importantly, such sequence-contextual as well as nucleosome-mediated biases in mutation rates are excluded as an explanation for the observed periodicities in the skew of the SNP frequency spectrum ($\Delta_S$), since strictly neutral mutations should display the same frequency distribution across the genome [31,35,59]. Support for a role of natural selection maintaining these periodicities is bolstered by the stronger periodicities in intronic nucleosomal sequences, where transcription-associated remodeling and disruption of nucleosome-DNA interactions are more likely to have functional impacts.

There is, however, one potential “selectively neutral” mechanism to explain the observed periodic patterns in $\Delta_S$. Biased gene conversion (BGC), a process where heteroduplex regions formed between homologs are repaired in a direction favoring one base, can create SNP dynamics analogous to those of directional selection [79]. BGC systematically favoring GC over AT has been observed in a few species and indirectly implicated in others by associations of local GC content with estimated rates of crossing over [80]. However, evidence for such an association is not observed in Drosophila [81]. Given that the magnitudes of average $\Delta_S$ and its periodicities for G→GA and C→CT SNPs are comparable to those of A→AG, any explanation of our results invoking BGC would have to involve multiple distinct gene conversional biases that depend on nucleosome position. While this is conceivable and worthy of further investigation, we conclude that the canonical GC-biased gene conversion is not a significant component of the evolutionary dynamics leading to these intricate nucleosomal patterns of polymorphism and divergence.

Whether these periodic patterns are the product of natural selection or BGC, the magnitude of the average force shaping the dynamics of nucleosomal SNPs must be small compared to that affecting the evolution of nonsynonymous variants. The shifts in G→A divergence between peaks and valleys in the n147 are ~0.001 against a background average of ~0.01, suggesting relatively weak constraint of 1 in 10 mutations. The nonsynonymous rate of divergence on the *melanogaster* lineage, ~0.006, is about one tenth of that for synonymous divergence corresponding to 9 out of 10 mutations being selected against [61]. Comparable conclusions could be drawn from the modest magnitudes of periodic fluctuation in expected heterozygosities and, indeed, in the widely observed periodicities in dinucleotide frequencies of nucleosomal sequences. Still, by virtue of its four-fold greater genomic footprint, the net selective impact of just the selection associated with such nucleosomal periodicities could approach the magnitude of non-synonymous variants. As is the case for coding sequence, differences in the relative (average) rates reflect the aggregate impact of selection that must vary substantially among nucleosomes, as well as among sites.

Table 1. Pearsons correlation of lineage specific (per-site) divergence or $\Delta_S$ with dinucleotide frequencies.

|                  | G→A & AA | C→T & TT | A→G & GC |
|------------------|----------|----------|----------|
| intronic divergence | 0.648    | 0.521    | 0.847    |
| intergenic $\Delta_S$ | 0.803    | 0.279    | 0.830    |
| intronic $\Delta_S$ | 0.499    | 0.531    | 0.559    |
| intergenic $\Delta_S$ | 0.317    | −0.059   | 0.093    |

![Image](https://www.plosgenetics.org/lookup/doi/10.1371/journal.pgen.1004557.t001)
Evidence of natural selection supporting nucleosome-associated sequence periodicities and the implication of their biological impact casts the potential functions of non-protein-coding regions in a new light. Substantial portions of Drosophila, human and other genomes appear to be under evolutionary constraint, yet lack any functional annotation [67,68]. Further, SNPs identified by genome-wide association studies (GWASs) of interesting human phenotypes often have mild attributable effects and map to unannotated intronic or intergenic regions, where mechanistic hypotheses concerning the impacts of such genomic variation are lacking. We demonstrate that at least part of the constraint in Drosophila arises from interactions between histone proteins and DNA sequence.

Our results suggest dinucleotide periodicities and the rotational positioning that they guide have significant biological consequences. Sequences affecting rotational positioning can influence the binding of transcriptional activators and participate in regulation of expression or gene splicing [25,28,82–84]. More generally, they impact nucleosome assembly and stability [2–7,9,17], properties that broadly impact chromatin dynamics and may influence higher order chromatin structures. Further, the observed large-scale periodicities in dinucleotide frequencies (and divergence and $\Delta_p$ patterns supporting them) demonstrate that sequences that facilitate rotational positioning are specifically enriched relative to adjacent nucleosomes. So, while periodic sequence patterns are considered more relevant to rotational positioning, they clearly interact with the translational positioning of arrayed nucleosomes in Drosophila. Going forward, deeper and more detailed population genomic analyses should provide a unique window into the complex in vivo interactions between DNA sequence and nucleosome function.

The significance of these periodic patterns of polymorphism and divergence is amplified in light of the substantial proportion of the eukaryotic genome packaged in nucleosomes (four-fold greater than that of coding sequence in Drosophila) and the broad conservation of dinucleotide interactions with the histone core. Indeed, no other DNA-protein interaction remotely approaches the genomic density or structural impact of nucleosomes. The striking periodic variation we observe relative to nucleosomes fundamentally changes expectations about divergence and SNP frequency, particularly in non-protein-coding regions. Our results point to a layer of evolutionary forces across entire genomes, emanating from the interactions of DNA sequence variation with the structure and function of the histone core.

Materials and Methods

Nuclear isolation and MNase digestion

Embryos were collected from population cages [85] over a 1 hr period and aged at 25°C for 2–3 hr. Staged embryos were dechorionated in 50% bleach for 2 minutes, washed extensively, and then homogenized on ice in SEC buffer (10 mM HEPES, 150 mM NaCl, 10 mM EDTA 10% glycerol, 1 mM DTT) with Protease Inhibitors (PI) (0.1 mM PMSF and 2X Roche EDTA-free Protease Inhibitor tablets). After lysis filtering and centrifugation, pelleted nuclei were resuspended in CIB (15 mM Tris pH 7.5, 60 mM KCl, 15 mM NaCl, 0.34 M Sucrose, 0.15 mM Spermine, 0.5 mM Spermidine)$\pm$PI and then repelleted. Centrifugation of nuclei in CIB was repeated 3 times, and the resultant pellet was flash frozen and stored at −80°C.

Pelleted frozen nuclei were resuspended in CIB$\pm$PI, and chromatin was digested with 0.5 U/ml Micrococcal nuclease (Sigma) for 37°C for 15 min. MNase treated nuclei were pelleted, resuspended in 0.1% NP-40 PBS$\pm$PI, and incubated at 4°C for 3 hrs to release (primarily) mononucleosomes. Nuclei were repelleted, and chromatin from the supernatant was phenol-chloroform extracted. Digestion was analyzed on an agarose gel.

Library construction and sequencing

Approximately 882 ng of DNA was used as starting material for paired-end sequencing library construction following the Illumina protocol (PE-102-1001). 10 µl of paired end adapter oligos were ligated to the end-repaired, A-tailed fragments in a 50 µl reaction. The adapter-ligation product was gel-purified to select molecules approximately 150–700 bp in length and re-suspended in 30 µl total volume. 1 µl of size-selected ligation product was used as template for 12 cycles of library enrichment PCR in a 50 µl reaction volume. The enriched library was purified using QIAAGEN MinElute columns and sequenced (2$\times$36 cycles) on one lane of a flow cell (FC42JBB) with an Illumina GAIIx running the Illumina software SCS v2.4.135/Pipeline v1.4.0.

Subsequently, 8 µl of the same size-selected ligation product was used as a template for 10 cycles of library enrichment PCR in a 100 µl reaction volume. To enrich for 147 bp fragments, the library was purified using QIAAGEN MinElute columns, then size selected on an agarose gel to recover fragments approximately 273 bp in length (as determined by an Agilent Bioanalyzer). This size-selected library was sequenced (2$\times$36 cycles) on four lanes of a flow cell (FC61BGN) with an Illumina GAIIx running the Illumina software SCS v2.5.38/Pipeline v1.5.0.

Read mapping and filtering

Reads that passed the Illumina pipeline’s quality filters were then aligned to the Berkeley Drosophila Genome Project’s Release 5 reference sequence [86] using Version 0.7.0 of the MAQ program [85]. Read pairs that mapped more than 1,000 bp apart and those for which the combined sum of the quality scores of mismatches exceeded 300 were filtered using the $\text{maq map} – a 1000 – e 300$ command. Otherwise, default $\text{maq}$ parameters were used.

147 bp Fragments in intronic and intergenic regions

Mapped paired end clones of length 147 bp were then filtered based on Release 5.16 FlyBase Annotation of the D. melanogaster genome and classified as intronic or intergenic. For classification, all bases, including the flanking $\pm 50$ bp, were required to map entirely to a contiguous intronic or intergenic region. Heterochromatic reads were removed using cytogenetically-defined boundaries [87]. Downstream analysis was carried out on 276,614 intergenic and 270,998 intronic autosomal 147 bp nucleosomal fragments, referred to as non-CpG nucleosomes. n147. These cover 61% and 79% respectively of the target nucleosomal fragments, referred to as intronic and intergenic n147. These cover 61% and 79% respectively of the target intergenic and intronic regions in the euchromatic autosomes (chr2 and chr3). The coordinates of the n147$\pm$50 bp flanking regions are in Tables S1 and S2. Average nucleosomal read depths, where represented, are a pileup representation of a set of similarly processed paired end clones with lengths ranging from 142–152 bp.

In calculating dinucleotide frequencies, divergence, $\pi$ and $\Delta_p$ across n147 regions (and, where relevant, $\pm$50 bp flanking), both positional and average calculations took the dyad symmetry into account. Substitutional pathways were switched at the dyad axis, such that positions $\pm$73 to $\sim$1 were joined to the reverse complement of bases 1–73. Where included, flanking regions ($\pm$50 bp) were treated similarly. For larger scale positional divergence and genomic averages, data from complementary substitutional pathways were combined.
Dinucleotide frequencies

The n147 dinucleotide frequencies are the averages (in the reference sequence) over all n147 fragments for each position. Genomic over/underrepresentations of dinucleotides were calculated by dividing the difference between observed and expected frequencies by the expected frequency. Estimates of expected intergenic and intronic dinucleotide frequencies were calculated based on underlying base frequencies. Observed frequencies were computed directly from the reference sequence.

Population genomic data

The sequences of the euchromatic portions of 36 D. melanogaster genomes from Raleigh, North Carolina were released by DPGP (http://www.dpgp.org/1K_50genomes.html - Reference_Release_1.0). The sequencing, alignment and assignment of estimated quality scores are described in Langley, et al., 2012 [65]. The sequences of the 22 D. melanogaster genomes from Rwanda, Africa were released by DPGP (http://www.dpgp.org/dpgp2/update_20Jan2012/dpgp2_v2_rg.ID5.nohets.fasq.bz2). The sequencing, alignment and assignment of estimated quality scores are described in Pool, et al., 2012 [71]. For both data sets, only bases with a minimum quality score of Q30 or greater were included in the analyses.

Divergence, frequency, π and Δπ

Calculations of divergence on the melanogaster lineage, frequency, expected heterozygosity (π) and the index of skew in the frequency spectrum (Δπ) were done only on sites that could be polarized using a multiple alignment of D. melanogaster, D. simulans, D. yakuba and D. erecta genomes [65], i.e., simulans and yakuba and/or erecta have the identical base. For consideration of the potential impact of ascertainment bias association with the isolation of nucleosomes, sites were subjected to a more stringent polarization (simulans, yakuba and erecta have the identical base) and calculations were done only on sites where the experimental genome had the ancestral allele. These statistics (divergence, π and Δπ) were estimated with two alternative weightings. The first, per-n147, gave weight to genomic sites proportional to their occurrence at nucleosome positions among the n147. Thus, in instances where a particular site was found multiple times in the n147 set, the divergences or SNPs at that site were given proportionally more weight. This in effect weights the signal by nucleosome site “occupancy.” The second, per-site, counted normalized the weighting such that each site (conserved, divergence or SNP) contributed equally, independent of its recurrence in the n147. The per-n147 estimates reflect those nucleosomal sequences that were readily isolated, while the per-site method treats each population genomic variant equally.

A third representation of the mapping of divergence and Δπ consists of random non-overlapping regions sampled from the n147. These analyses are presented to simply address whether the periodicities arise solely from a ~10 bp periodicity in the overlap of n147 fragments. For divergence, non-overlapping n147 sets (72,710 intronic and 72,859 intergenic sequences) were generated by random sampling of intergenic and intronic n147 without replacement. Newly drawn sequences were added to the non-overlapping subset only if they did not share any positions with prior sampled sequences. These non-overlapping subsets together cover 53% of the target intergenic and intronic regions in the euchromatin autosomes (chr2 and chr3) covered by the full n147. For non-overlapping Δπ, intergenic and intronic n147 (not including the flanking ±50 bp) were first filtered for only those nucleosomal regions containing the relevant SNP (taking dyad symmetry into account). This produced non-overlapping intergenic and intronic sets of ~90,000 regions each for G→GA and C→CT and ~55,000 regions for A→AG. These sets were then subjected to random sampling without replacement. A new n147 was added to the set if it did not overlap any already in the set. Non-overlapping G→GA and C→CT intronic and intergenic sets contained ~46,000 regions each and A→AG sets contained ~30,000.

All three of these methods for calculating divergence and Δπ yielded similarly periodic patterns reflecting the fact that while the genomic coverage of the n147 is not deep, it is also relatively uniform (cv 0.65) and the periodicities are not arising from a small subset of the n147 or interactions from overlapping n147 sequences.

Definition of divergence. Divergence was based on the species reference sequences (see above). Polarized divergence estimates were calculated as the number of specific substitutions (e.g. number of G→A) divided by the number of polarized sites inferred to have been the specific ancestral state (i.e., G in this case). For n147 average divergence, 5 bp at each end of the fragments were trimmed from the analysis to minimize the influence of sequence bias at the enzyme cleavage site (Figure S4).

Definition of frequency and π. All estimates of frequency and expected heterozygosity, π were calculated over sample sizes between n_min and n_max as

\[ \text{frequency} = \frac{1}{L} \sum_{i=n_{\min}}^{n_{\max}} \frac{\sum_{j=1}^{S} \chi_j}{L} \]

\[ \bar{\pi} = \frac{1}{L} \sum_{i=n_{\min}}^{n_{\max}} \left[ \frac{i}{i-1} \sum_{j=1}^{S} \chi_j (1 - \chi_j) \right] \]

where L is the total number of sites (bp) with sample size between n_min and n_max, x_i is the frequency of an allele at the jth of S sites with sample size i. These SNPs can be categorized simply by state (A, C, G or T) or also as derived from an inferred ancestral state (e.g., A in a G→GA SNP). For n147 average π, n147 regions were trimmed as described above for divergence.

Definition of Δπ

We require a sample-based index of the skew in the site frequency spectrum. Tajima [59] proffered the test statistic, D, a normalization of d, the difference between two estimates of the same population parameter, 4Nμ, where μ is the mutation rate to selectively neutral alleles and N is the population size. These estimates are

\[ k = L \bar{\pi} = \frac{n}{n-1} \sum_{i=1}^{S} 2x_i (1 - x_i) \]

and

\[ \bar{M} = \bar{\theta} L = \frac{S}{\sum_{j=1}^{S} (1/j)} \]

in a sample of size n, where x_i is the frequency of one of the two alleles at each of the S segregating sites in an arbitrary genomic segment of length L base pairs. Thus \( \bar{\pi} \) (expected heterozygosity) and \( \bar{\theta} \) are per site (base pair) estimates of 4Nμ.
But here we seek not a test statistic for a genomic segment, but an index of the same deviation, that can be aggregated across heterogeneously sampled data and compared across classes of genomic annotation.

To that end consider
\[
\Delta_s = \frac{k}{S} - \frac{1}{S} \sum_{i=1}^{a-1} \left( \frac{1}{n_i} \right)
\]
a simple rescaling of Tajima’s (little) \( \bar{d} \),
\[
d = S \Delta_s.
\]

Tajima [59] also presents the distribution of the proportion of \( S \) segregating sites with frequency \( i/n \) in the sample, \( G_s(i)/S \). \( G_s(i) \) can not be used to compute properties of a sample unless one can argue that the sites evolved independently and are sampled independently. If we choose a set of \( S \) segregating sites (assumed to be independently sampled from a population, i.e. no linkage disequilibrium), rather than a genomic segment, we have the expected heterozygosity in a sample of size \( n \),
\[
E\left( k | n, S \right) = \frac{1}{2} \sum_{i=1}^{n} G_s(i) \left[ 2 \left( \frac{i}{n} \right) \left( \frac{n-i}{n} \right) \right]^{-1} - \frac{1}{S} \sum_{i=1}^{a-1} \left( \frac{1}{n_i} \right)
\]
and so our index,
\[
\Delta_s = \frac{k}{S} - E\left( k | n, S \right) = \frac{k}{S} - \frac{1}{S} \sum_{i=1}^{a-1} \left( \frac{1}{n_i} \right)
\]
This index is thus a measure of the deviation of the population (expected) heterozygosity per segregating site from its predicted value under the assumptions of equilibrium between selectively neutral mutation and genetic drift in a Wright-Fisher population.

To estimate this deviation across sites with different sample sizes, we can calculate the weighted average, weighting by the reciprocal of the variance.

The variance of \( \Delta_s \) is
\[
Var[\Delta_s(n)] = \frac{1}{n} \sum_{i=1}^{n} \left\{ \frac{G_s(i)}{2S} \left[ \frac{2i(n-i)}{n(n-1)} \right] \right\}^2
\]
\[
= \frac{1}{n} \sum_{i=1}^{n} \left( \frac{1}{n_i} \right) \sum_{j=1}^{n-1} \left( \frac{1}{n} \right) \frac{2j^2(n-j)^2}{n^2(n-1)^2}
\]
Notice that this is the theoretical variance in \( \bar{d} \) at a single segregating site in a sample of size \( n \) under the assumptions of the neutral model (above).

If the frequencies at different sites are independent then we can estimate the sample variance of \( \Delta_s(n) \) for \( S \) sites with sampling depth \( n \), simply as \( Var[\Delta_s(n)]/S \).

Assuming again that these SNPs are sampled independently both within and over sample sizes (i.e., no linkage disequilibrium), the average \( \Delta_s \) can be estimated by the weighted average
\[
\Delta_s = \frac{\sum_{i=1}^{n_{\max}} \Delta_s(i) \left( \frac{S_i}{Var[\Delta_s(i)]} \right)}{\sum_{i=1}^{n_{\max}} \frac{S_i}{Var[\Delta_s(i)]}}
\]
where \( n_{\min} > 3 \) and \( n_{\max} \) is largest sample size.

\( \Delta_s \) for each position in the \( n_{147} \) regions (and its average across positions) was calculated over polarized sites with sampling sizes (\( n \)) between 32 and 34 in the Raleigh [65] and 18 to 21 for Rwandan data [71]. For \( n_{147} \) average \( \Delta_s \) the \( n_{147} \) regions were trimmed as described above for divergence. Data from complementary pathways were merged for larger scale positional \( \Delta_s \).

Correlations of divergence with GC frequency and nucleosome "occupancy"

\( n_{142}\)-\( 152 \) coverage of intergenic \( n_{147} \) regions was defined as the sum of the coverage of the region by the larger set of \( 142-152 \) bp nucleosomal fragments. This corresponds to what some authors call "occupancy." GC frequency of intergenic \( n_{147} \) regions was calculated based on the nucleotide frequencies in the \( D. melanogaster \) reference sequence from bases 6 to 141 of the \( n_{147} \) regions (to minimize the potential impact of MNase sequence bias). For density plots and Spearman’s \( \rho \), only \( n_{147} \) regions with at least 50 (out of 147) intergenic bases polarizable were included in the analyses. As above, the interior 5 bp on each end of \( n_{147} \) regions were trimmed from the analysis. For correlations between divergence and genomic intergenic GC frequencies, Spearman’s \( \rho \) were reported for non-overlapping 500 bp windows in which at least 166 intergenic bases were polarizable and no more than 250 bp in the reference were "N".

Plot details

Plots were generated using R [88]. Prior to plotting, all calculations were symmetrized around the dyad axis. \( n_{147} \) (\( \pm 50 \) bp) divergence and \( \Delta_s \) plots (Figure 1D, Figure 2B,C, Figure 3A) were smoothed using running average in a window of 5 bp (weights: 0.125, 0.250, 0.250, 0.250, 0.125). For large scale plots of dinucleotide frequency, divergence and \( \Delta_s \) (Figure 1C, Figure 2D, Figure 3B), flanking regions were smoothed using running average smoothing with a window of 50 bp of equal weights. In those plots, the central \( n_{147} \) regions were smoothed separately using a 50 bp window of equal weights. Regions of 5 bp upstream and downstream of the \( n_{147} \) edges were trimmed prior to smoothing for large-scale plots.

Structural mapping of divergence rates

To elucidate the distribution of the smoothed (as above) divergence on DNA from the structure of the nucleosome we colored-coded values of each base pair in a schematic rendering of the DNA strands of \( p h 1 k s 5 \) [53] using PyMOL [89]. Only the "top" turn of the DNA (base pairs 73 to -6) is shown. Bases 73 to 70 were rendered as grey, due to extreme values induced by MNase sequence bias.
Supporting Information

Figure S1  Paired end sequencing of MNase released mono-nucleosomal fragments produces a range of fragment sizes. (A) Electrophoretic separation of DNA isolated from digestions of embryonic chromatin with three dilutions of MNase (at top). (B) Distributions of sizes of sequenced nucleosomal fragments. The proportion of paired-end sequenced fragments of each size is plotted against the size (bp). The sizes of 8,124,351 fragments that mapped to the 96.6 Mb of euchromatic arms from a single GA2x lane of the initial library prepared from DNA isolated from the 1× digestion in panel A is shown in black. Almost completely superimposed are the distributions of sizes of fragments from four GA2x lanes of a size-selected fraction of that initial library (numbers: 7,892,551, 7,947,142, 7,871,834 and 7,825,097; red, blue, green and magenta respectively). Also shown in light green is the position of the n147 fraction used in the most of the analyses.

Figure S2  Dinucleotide frequencies across intergenic (black) and intronic (red) n147 regions display a ~10 bp periodicity. Dinucleotide frequencies are represented for bases 0 to ~75 across n147 and ~50 bp of flanking sequence.

Figure S3  Nucleotide frequencies surrounding intergenic and intronic n147 regions show a nucleosome-scale periodicity. Single nucleotide frequencies surrounding n147 regions (±600 bp).

Figure S4  Per-site divergence on the melanogaster lineage for the substitutional pathways displays ~10 bp periodicity, consistent with per-n147 results. Smoothed average per-site polarized divergence across intergenic and intronic n147 regions (±50 bp) for specific substitutions is shown. Indicated dinucleotide frequencies are plotted above for reference.

Figure S5  Polarized divergence on the melanogaster lineage is periodic across n147 regions. (A) Smoothed average polarized divergence for specific substitutions across regions defined by non-overlapping subsets of intergenic and intronic n147 (±50 bp). Intergenic dinucleotide frequencies are represented above for reference. (B) Smoothed intergenic A→G divergence mapped on to bases +73 to ~6 from the nucleosome structure [55]. Arginines that contact the minor groove are color coded by histone. (C) Smoothed average combined A→G;T→C polarized divergence surrounding intergenic (black) and intronic (red) n147 regions (±1 kb). Read depth of intronic (light grey) and intergenic (grey) n142-152 is represented below.

Figure S6  MNase cleavage sites show consistent sequence bias at the boundaries of intergenic and intronic n147 fragments. Average 5′-3′ nucleotide frequencies surrounding the MNase cleavage sites for intergenic and intronic n147 fragments (±50 bp).

Figure S7  Divergence of six substitutional pathways along the melanogaster lineage correlate with GC content and nucleosome “occupancy”. (A) Plots of the rate of divergence for intergenic n147 sequences for six substitutional pathways versus the log10 of coverage by n142-152 (“occupancy”). Correlation coefficients are reported in Table S3. (B) Plots of the rate of divergence for intergenic n147 sequences for six substitutional pathways versus their fraction GC. Correlation coefficients are reported in Table S3. Red lines show the linear regression fits, while the green represent the least-square fits to a third degree polynomial.

Figure S8  G→A:C→T divergence on the melanogaster lineage and AA/TT frequency varies with n147 GC content and nucleosome “occupancy.” (A) Average frequency of AA/TT dinucleotides (green) and rate of G→A:C→T divergence (orange) for intergenic n147 (and 1 kb flanking) regions binned by nucleosome “occupancy”, i.e., coverage of the n147 by the larger set of n142-152 nucleosomal fragments. n147 regions were ranked by coverage and separated into 5 bins (top is lowest and bottom is highest). Average divergence and AA/TT frequencies are shown for each bin. Intergenic n142-152 coverage is represented in grey (scale on blue axis). Although the axis limits for coverage vary, the scale of these axes is consistent. (B) Average G→A:C→T divergence across intergenic n147 regions for 5 bins based on n142-152 coverage (Q1 – lowest, Q5 – highest). (C) Average G→A:C→T divergence across intergenic n147 regions for 5 bins based on ranked n147 GC content (Q1 – lowest, Q5 – highest). (D) Scatter plot of intergenic n147 GC frequency against log10 n142-152 coverage. Spearman’s r is reported.

Figure S9  per-n147 SNP frequencies display a ~10 bp periodic variation across intergenic and intronic n147 regions. Smoothed average per-n147 frequency of A (A) or T (B) alleles for G-A or C-T polymorphic sites respectively in the Raleigh sample (purple). Frequencies after removal of singleton classes are plotted in blue.

Figure S10  Polarized per-n147 SNP frequencies across intergenic n147 regions show evidence of ~10 bp periodicities aligning with dinucleotide periodicities. (A) Average smoothed (black) and unsmoothed (grey) per-n147 frequencies of derived alleles for ancestrally G to G-A polymorphic, ancestrally C to C-T polymorphic and ancestrally A to A-G polymorphic sites in intergenic regions. (B) The same as A but with singleton classes (where only one allele of the derived or ancestral base was observed) removed.

Figure S11  Polarized per-n147 SNP frequencies across intronic n147 regions show evidence of ~10 bp periodicities aligning with dinucleotide periodicities. (A) Average smoothed (black) and unsmoothed (grey) per-n147 frequencies of derived alleles for ancestrally G to G-A polymorphic, ancestrally C to C-T polymorphic and ancestrally A to A-G polymorphic sites in introns. (B) The same as A but with singleton classes (where only one allele of the derived or ancestral base was observed) removed.

Figure S12  The weighting of estimates of Δr has little impact on the ~10 bp periodic patterns for per-site Δr and per-n147 Δr. Average smoothed (black) and unsmoothed (grey) per-site Δr across n147 regions (±50 bp) for G→GA, C→CT, and A→AG polymorphic sites. Color-coded dinucleotide frequencies are included for reference.

Figure S13  The patterns of per-n147 Δr across five independently sampled non-overlapping subsets of intergenic and intronic n147 sequences retain much of the ~10 bp periodic signal. Average smoothed (black) and unsmoothed (grey) Δr (Raleigh data) for G→GA, C→CT, and A→AG polymorphic sites across regions defined by non-overlapping subsets (see Materials and Methods).
Methods) of intronic and intergenic n147 (±50 bp). Frequencies of the indicated dinucleotides are plotted for reference. (EPS)

Figure S14  Per-n147 Δπ across intergenic and intronic n147 for Rwandan lines shows evidence of nucleosome-associated SNP periodicities in an independent sample. Average smoothed (black) and unsmoothed (grey) per-n147 Δπ for Rwandan lines for G→GA, C→GT, and A→AG polymorphic sites across regions defined by intronic and intergenic n147 (±50 bp). Frequencies of the indicated dinucleotides are plotted for reference. (EPS)

Figure S15  Ascertainment bias resulting from SNPs in the experimental strain does not explain the ~10 bp periodic patterns of Δπ. Average smoothed (black) and unsmoothed (grey) intronic per-n147 Δπ across n147 regions (±50 bp) for G→GA, C→CT, and A→AG polymorphic sites in the Raleigh sample where the experimental strain had the ancestral allele. (EPS)

Figure S16  A→G; T→C Δπ, in the regions flanking n147 sequences. Average combined A→G; T→C divergence (green) and A→AG; T→TC Δπ (black) surrounding intergenic and intronic n147 (±1 kb). Read depth of intronic or intergenic n142-152 is represented in grey below (axis label in blue). (EPS)

Figure S17  Genomic average values of divergence, π, and Δπ follow trends seen for n147 regions. Average scaled genome-wide π, Δπ and divergence for specific substitutions, partitioned by intergenic versus intronic (see Table S4). (EPS)

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Table S1  The genomic coordinates (Release 5) for intergenic autosomal 147 bp nucleosomal fragments ±50 flanking bp. (TXT)

Table S2  The genomic coordinates (Release 5) for intronic autosomal 147 bp nucleosomal fragments ±50 flanking bp. (TXT)

Table S3  Correlations between the rate of divergence of the indicated substitutional path and GC content or nucleosome “occupancy” (TXT).

Table S4  Estimates (per-n147) of π, Δπ and divergence for intronic and intergenic nucleosomal regions (n147), a non-overlapping subset of n147 and the euchromatic genome (see text). (TXT)

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Author Contributions

Conceived and designed the experiments: SAL GHK CHL. Performed the experiments: SAL CHL. Analyzed the data: SAL CHL. Contributed reagents/materials/analysis tools: SAL GHK CHL. Wrote the paper: SAL GHK CHL.
