

High-Resolution Profiling of *Drosophila* Replication Start Sites Reveals a DNA Shape and Chromatin Signature of Metazoan Origins

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SUMMARY

At every cell cycle, faithful inheritance of metazoan genomes requires the concerted activation of thousands of DNA replication origins. However, the genetic and chromatin features defining metazoan replication start sites remain largely unknown. Here, we delineate the origin repertoire of the *Drosophila* genome at high resolution. We address the role of origin-proximal G-quadruplexes and suggest that they transiently stall replication forks *in vivo*. We dissect the chromatin configuration of replication origins and identify a rich spatial organization of chromatin features at initiation sites. DNA shape and chromatin configurations, not strict sequence motifs, mark and predict origins in higher eukaryotes. We further examine the link between transcription and origin firing and reveal that modulation of origin activity across cell types is intimately linked to cell-type-specific transcriptional programs. Our study unravels conserved origin features and provides unique insights into the relationship among DNA topology, chromatin, transcription, and replication initiation across metazoa.

INTRODUCTION

Maintenance of cellular identity critically relies on the faithful transmission of the parental genome through DNA replication and a reestablishment of the epigenome (Alabert and Groth, 2012). Perturbation of this finely orchestrated process poses a major threat to genome stability, thus linking aberrant DNA replication to several human diseases (Zeman and Cimprich, 2014).

In the circular chromosome of bacteria and archaea, DNA replication starts from a single locus termed replication origin (Mott and Berger, 2007). In contrast, eukaryotic DNA replication requires the concerted activation of thousands of replication origins (Leonard and Méchali, 2013). The firing of a eukaryotic origin is preceded by the orderly recruitment of protein factors

to potential initiation sites. In G1 phase, the origin recognition complex (ORC) binds replication origins and, along with the help of Cdc6 and Cdt1, nucleates the pre-replication complex (pre-RC) through the loading of an inactive form of the minichromosome maintenance (MCM) helicase. At the onset of S-phase, Dbf4-dependent kinase (DDK) and cyclin-dependent kinases (CDKs) catalyze sequential phosphorylation events, which recruit initiation factors. These in turn stimulate MCM activity, complete replisome assembly, and trigger the initiation of DNA synthesis, a process referred to as origin firing (Masai et al., 2010). Whereas 60 years of genetic and biochemical dissection have elucidated much of the activation cascade underlying origin firing, the mechanisms that target replisomes to replication origins remain poorly understood, raising the question of which sequence and chromatin features define origins *in vivo*. A comprehensive answer to this question is missing, partly because, prior to the genomic era, only a handful of origins were precisely mapped (Leonard and Méchali, 2013), and partly because isolation and characterization of transient replication intermediates are experimentally challenging (reviewed in Gilbert, 2010). The isolation of small nascent leading strands (SNSs, Bielinsky and Gerbi, 1998) currently is considered the most reliable method to map replication origins (Leonard and Méchali, 2013).

Recent high-throughput approaches coupled SNS purification with tiling arrays (SNS-ChIP) (Sequeira-Mendes et al., 2009; Cayrou et al., 2011) or next-generation sequencing (NGS, SNS-seq) (Besnard et al., 2012; Picard et al., 2014), enabling systematic origin mapping genome-wide. Particularly, origin profiling in four human cell types (Besnard et al., 2012) identified thousands of active origins and suggested that cell-type-specific origin-usage signatures are responsible for the observed plasticity of replication programs. However, despite considerable efforts, these works fell short in identifying a eukaryotic consensus sequence and converged to associate G-rich elements and G-quadruplexes (G4s) (Maizels and Gray, 2013) to a variable fraction of initiation sites, suggesting that features beyond nucleotide sequence define metazoan replication origins. Cooperation between genetic elements and epigenetic features is therefore likely to be key for origin function, but chromatin configurations of replication start sites remain largely unexplored.

In this study, we delineate the origin repertoire of the *Drosophila melanogaster* genome at an unprecedented resolution. We examine the role of origin-proximal G4s and provide evidence that these DNA secondary structures act as replication fork barriers in vivo. We carefully dissect the chromatin configuration of replication start sites, and demonstrate that specific DNA shape and chromatin configurations, as opposed to strict sequence specificity, mark and accurately predict replication origins in higher eukaryotes. Finally, our study reveals that differential origin usage across cell types is tightly connected to cell-type-specific transcriptional programs, thus providing a means to couple chromatin processes crucial for the maintenance of cellular identity.

RESULTS

High-Resolution Mapping of *Drosophila* Replication Origins

Upon origin firing, two nascent leading strands extend from a short RNA primer and emanate bidirectionally from the origin. SNS-seq aims at selectively isolating these covalent RNA-DNA hybrids, whose 5' ends define the site of replication initiation. In the SNS purification protocol, origin-proximal SNSs are first size-separated from Okazaki fragments and then enriched by lambda-exonuclease (Lexo) digestion of non-RNA-primed DNA. As this 5' to 3' processive nuclease exhibits very weak activity on ribonucleotides, SNSs are protected from digestion while contaminating DNA species are degraded. However, even in rapidly dividing cells, SNSs account for only ~0.002% of total genomic DNA (Gilbert, 2012). Accuracy and resolution of origin detection, therefore, critically depend on efficient degradation of contaminant, unreplicated DNA. Moreover, in the absence of other DNA species, the relative abundance of SNSs from all origins firing throughout S-phase is expected to reflect their firing efficiency within a cell population, thus allowing estimates of aggregated firing probabilities (Gilbert, 2010). Treatment with Lexo has proven essential in eliminating contamination and previous work enriched for SNS through two or three rounds of Lexo digestion (Cayrou et al., 2011; Besnard et al., 2012; Picard et al., 2014).

Here we adopted an enhanced sensitivity SNS purification protocol (Cayrou et al., 2011; see [Experimental Procedures](#)) to map active replication origins genome-wide in two *Drosophila* cell lines, the late embryo-derived S2 and the neuronal-derived Bg3 cells, whose epigenomes have been profiled extensively by the modENCODE project (Celniker et al., 2009). For each cell type, we obtained highly pure SNS preparations from two biological replicates by subjecting size-selected genomic DNA to up to five rounds of Lexo digestion (Figure S1B, inset). High-coverage, saturating deep sequencing of these SNSs yielded a total of 119 and 251 million reads aligning to the *Drosophila* genome for S2 and Bg3 cells, respectively (Figure S1A). This led us to identify 7,268 and 8,212 high-confidence replication origins in S2 and Bg3 cells (Figure 1A), respectively, whose replication start sites (RSSs) were defined as the summit of highly resolved origin peaks (Figures S1G–S1I). Furthermore, position and firing efficiency of a subset of origins were confirmed by qPCR (Figure S1B; Table S1).

Notably, 73%–81% and 69%–75% of origin peaks were independently detected within S2 and Bg3 biological replicates, respectively (Figures S1D and S1E). These values not only compare favorably to previous studies, but also exceed the technical reproducibility of recently published SNS-seq data in human K562 cells (Picard et al., 2014). Moreover, SNS-seq signals exhibited a nearly perfect correlation ($r = 0.98$) across biological replicates when computed in the union of all origin peaks from a given cell type (Figure S1F). To verify the accuracy of our origin mapping, we compared our results with previously published datasets. S2 and Bg3 origins covered a total of 6.9 and 6.4 Mb, respectively, a considerably smaller fraction of the *Drosophila* genome than the 27.3 Mb spanned by 6,184 origins previously identified in *Drosophila* Kc cells (Cayrou et al., 2011; Figures 1A, 1E, and S1G). In addition, <14% of S2 and Bg3 origin peaks sufficed to recall >72% of modENCODE early-origin regions (EORs) (Eaton et al., 2011; Figure 1B), which were mapped by BrdU immunoprecipitation from G1/S-synchronized cells and resulted in broad initiation zones. Our data finely resolved the composition of initiation zones exhibiting significantly higher origin scores than regions exclusively identified by modENCODE (Figures 1C and 1E for an example), suggesting that the remaining low-confidence events contain dormant origins that do not fire in an unperturbed S-phase (Zeman and Cimprich, 2014). Taken together, these results demonstrate the high quality and unmatched resolution of our origin mapping, which markedly expands the origin repertoire of the *Drosophila* genome to 14,005 distinct genomic loci.

Modulation of Origin Activity, Not Site Selection, Defines Cell-Type-Specific Replication Programs

Next, we analyzed the overlap among S2, Bg3, and Kc origins. We found 16%–20% of origin peaks common to all three cell types (constitutive origins) and 35%–45% of origin sites activated by at least two cell types. This overlap is significantly larger than expected by chance ($p < 0.001$, Figure 1A) and indicates a preferred origin localization across cell types. Previous work reported that cell-type-specific origins were on average poorly used (Besnard et al., 2012). However, whether firing of these sites was restricted to the cell type of detection or whether these origins are also marginally used in other cell types remained unclear. To address this question, we estimated firing efficiencies by integrating SNS-seq signals within all detected origin peaks and in ten matched background sets (see [Experimental Procedures](#)). In line with previous reports (Besnard et al., 2012), we found that constitutive origins exhibited on average the highest efficiency values across the entire origin repertoire (Figure 1D). However, cell-type-specific origin peaks were not only characterized by low efficiency in the cell type of detection, but they also yielded SNS-seq signals well above background in other cell types (Figure 1D). This result indicates that virtually all origins we identified fire in each *Drosophila* cell type but with characteristic frequencies. Cell-type-specific origins, therefore, could rather be termed cell-type-preferred origins, thus reflecting cell-type-specific preferences for low-efficiency origins. Our results strongly suggest that modulation of origin activity, not the selection of origin sites, is likely to define cell-type-specific replication programs in *Drosophila*.

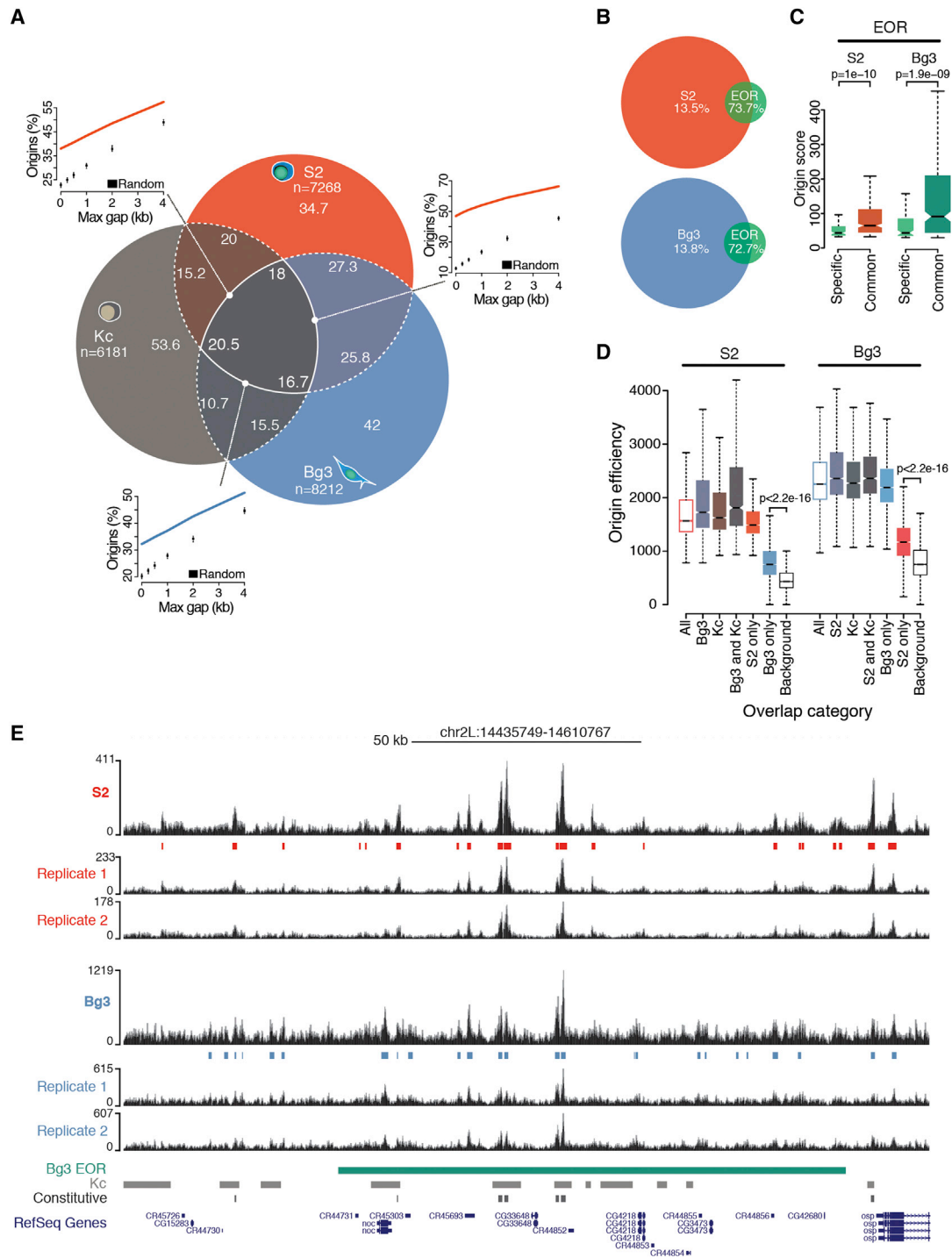


Figure 1. High-Resolution Mapping of the *Drosophila* Origin Repertoire

(A) Percentage overlaps of origin peaks identified in S2, Bg3, and Kc (Cayrou et al., 2011) *Drosophila* cells and comparison of observed pairwise overlaps (lines) with random expectations (boxplots) are shown (n, total number of origin peaks).

(B) Percentage overlaps of S2 and Bg3 origin peaks and modENCODE EORs are shown.

(C) Origin scores of EORs overlapping with S2 and Bg3 origins (common) or solely identified by modENCODE (specific) are shown.

(D) Efficiency of S2 and Bg3 origins partitioned and color-coded according to (A). Background estimates are shown.

(E) A representative snapshot shows the SNS-seq coverage in S2 and Bg3 cells from two biological replicates and detected origin peaks. A single EOR (green) spans most of this 175-kb genomic region. Kc (gray) and constitutive (black) origins are also shown (p values are from Wilcoxon rank-sum test). See also Figure S1.

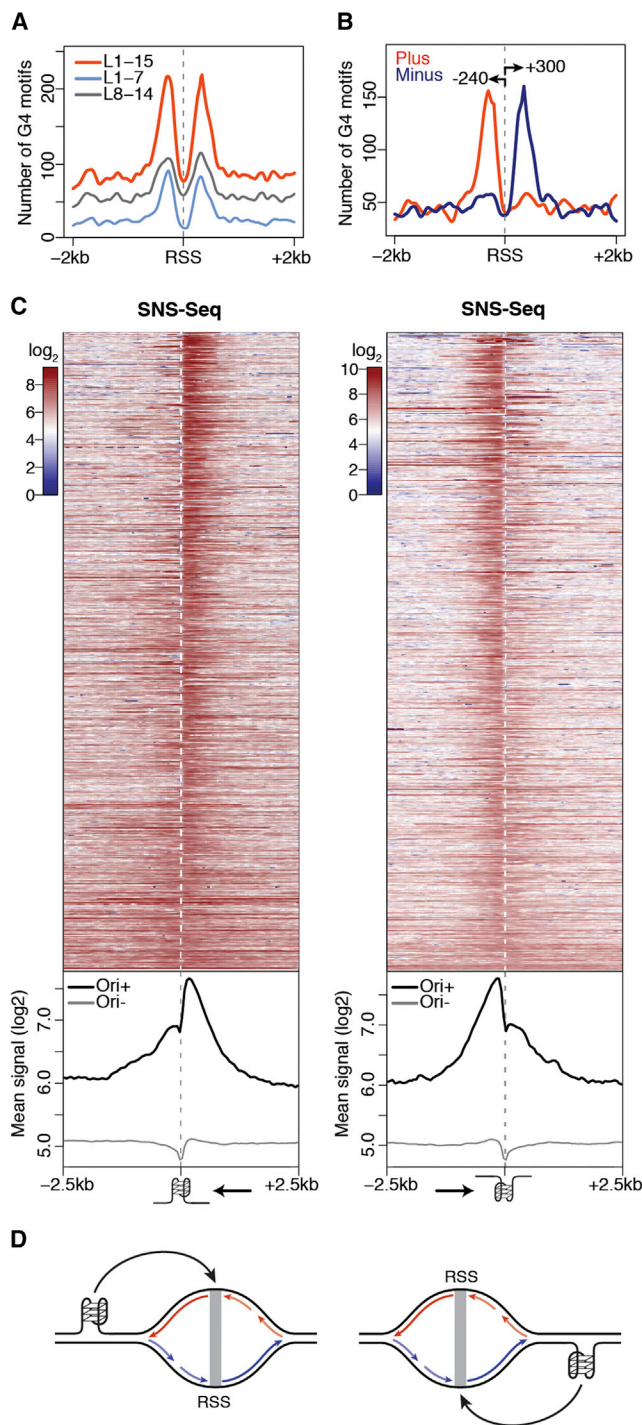


Figure 2. A G4 Signature at S2 Replication Origins

(A) Spatial distribution of G4 motifs within ± 2 kb of S2 RSSs is shown. (B) Same as (A) for strand-specific annotation of G4 L1–15 motifs. Arrows indicate peak distances (base pairs) from the RSS. (C) S2 SNS-seq signals within ± 2.5 kb of origin-associated G4 L1–15 motifs occurring on the plus (left) and minus (right) strands, ranked by coefficient of variation. Bottom shows the average of the signals above (Ori+) and at origin-negative (Ori-) G4 motifs. Arrows indicate the direction of the leading strand facing the G4. (D) Model describes how origin-proximal G4 motifs could orient (black arrows) replication forks. Leading strands (long arrows) and Okazaki fragments (short arrows) replicating the plus (red) and minus (blue) strands are indicated. See also Figure S2.

Origin-Proximal G4s Act as Transient Replication Fork Barriers In Vivo

Recent work investigated the role of G4s in DNA replication (Valton et al., 2014; Castillo Bosch et al., 2014), yet the potential contribution of G4 structures to replication initiation remained controversial. Here we examined the association between G4s and origins genome-wide.

We predicted G4 occurrences in the *Drosophila* genome and used the loop size (L) to define nested classes of G4 motifs. We found a significant association between origins and predicted G4s as compared to random expectation ($p < 0.001$, Figure S2A), with 9% and 22% of S2 origins overlapping 7% and 5% of L1–7 and L1–15 G4 motifs, respectively. G4-associated origins were more efficient than G4-negative origins (Figure S2B). This result is not mediated by colocalization with transcription start sites (TSSs, Figure S2C), which do not significantly associate with origins in *Drosophila* ($p = 0.81$, Figure S2D; Cayrou et al., 2011). We then examined the position and orientation of G4 motifs with respect to S2 origins. The alignment of G4-associated origins relative to their RSSs revealed a strong positional preference (Figure 2A), with strand-specific occurrences of G4s peaking at 240–300 bp from RSSs and largely restricted to their flanking regions (Figures 2B, S2E, and S2F).

Next, we quantified the spatial distribution of SNS-seq signals within a 5-kb region centered on each G4 L1–15 motif and partitioned G4 occurrences by strand and association with S2 origins. We observed a skewed distribution of SNSs at origin-proximal G4s, with most of the signal contributed by RSSs located downstream and upstream of G4 motifs mapping to the plus and the minus strands, respectively (Figure 2C). However, we noted that the SNS signal was not only asymmetrically distributed at these G4 sites, but it also sharply dropped exactly at the G4 position (Figure 2C). Thus, we reasoned that our data might capture G4-proximal replication fork-stalling events and that, if this is the case, only synthesis of the leading strand replicating the G4 template should be affected.

To test this hypothesis, we set out to indirectly monitor the progression of replication forks emanating from origin-associated G4s by purifying, barcoding, and deep sequencing SNSs of increasingly larger sizes (Figure 3A). Two sequencing libraries were prepared for each of three gradient fractions, for a total of 298 million reads aligning to the *Drosophila* genome (Figure S3A). Fractions 4 (shortest DNA molecules), 5 (intermediate), and 6 (largest) independently identified 4,505, 6,448, and 5,814 origin peaks, respectively, most of which (>81%) overlapped in two or more fractions (Figures S3B and 3B). The SNS enrichment of a subset of origin peaks was further confirmed by qPCR and negatively correlated with SNS sizes (Figure S1C; Table S1). SNS-seq signals in the union of origin peaks were highly correlated across fractions ($r = 0.94$ – 0.98 , Figure S3C). Moreover, >70% of origins detected by all fractions ($n = 3246$) overlapped S2 origins previously

(D) Model describes how origin-proximal G4 motifs could orient (black arrows) replication forks. Leading strands (long arrows) and Okazaki fragments (short arrows) replicating the plus (red) and minus (blue) strands are indicated. See also Figure S2.

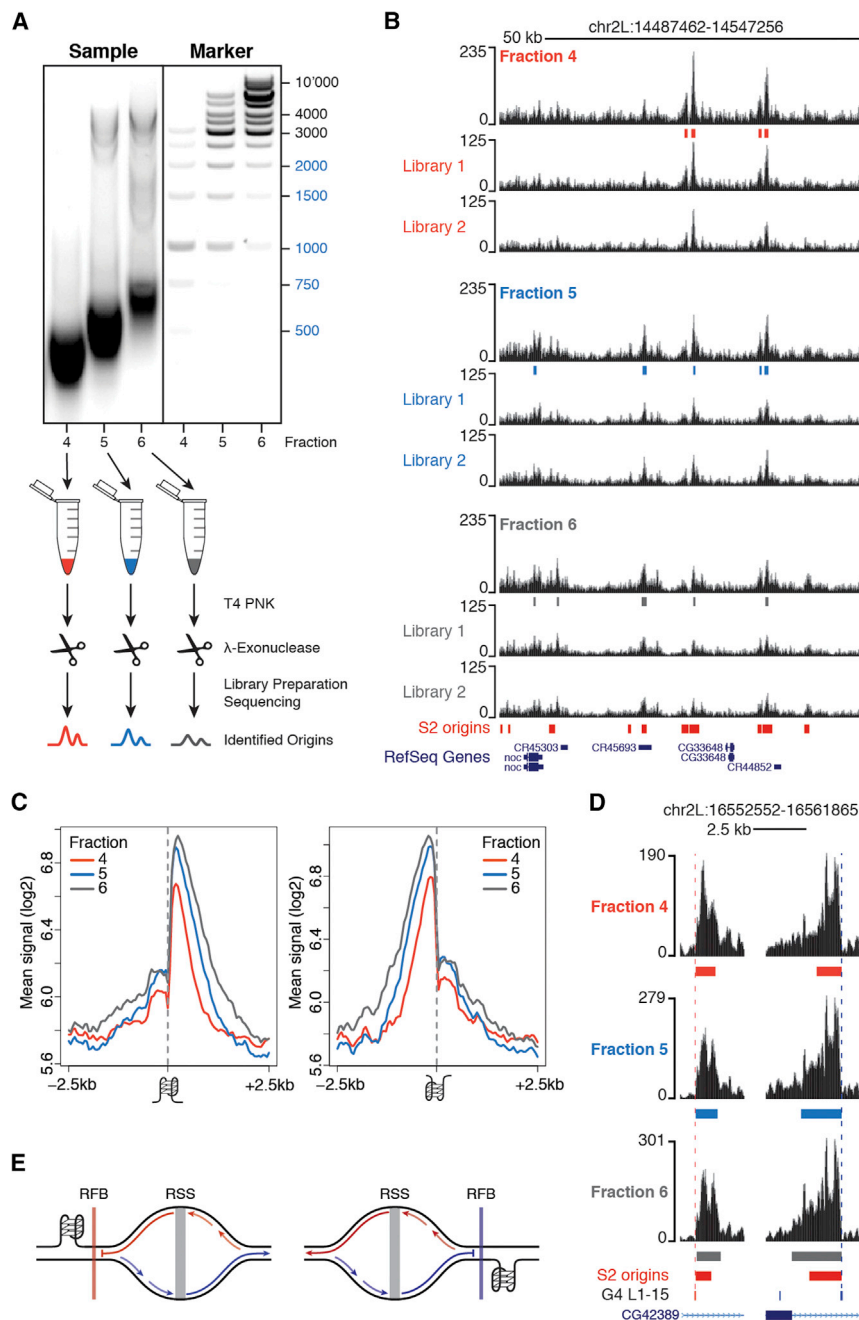


Figure 3. Origin-Proximal G4s Stall Replication Forks In Vivo

(A) An outline of the experimental strategy used to indirectly monitor replication fork progression at origin-associated G4s. Fractions 4–6, corresponding to marker lanes 4–6, were individually purified and subjected to two sequential rounds of T4 PNK phosphorylation and Lexo digestion. Two sequencing libraries were prepared for each sample and origin peaks were called on their union.

(B) A representative snapshot of the single-fraction SNS-seq coverage. Origin peaks identified in each fraction and S2 origin peaks from standard SNS-seq experiments are shown.

(C) Average single-fraction SNS-seq signal within ± 2.5 kb of origin-associated G4 L1–15 motifs occurring on the plus (left) and minus (right) strands is shown.

(D) Two representative G4 motifs occurring on opposite strands are shown.

(E) Model describes how origin-proximal G4 motifs could act as replication fork barriers. Origin-proximal G4s pause the synthesis of the nascent leading strands replicating the G4 template. See also Figure S3.

A DNA Shape Signature of Metazoan Replication Origins

With the high resolution of our data, it was possible to revisit the sequence characteristics of replication origins that have been elusive in lower resolution studies. We started by examining the local DNA sequence composition at RSSs. Interestingly, nucleosome-repelling AAAA polynucleotides and AA dinucleotides (Kaplan et al., 2009; Tillo and Hughes, 2009) were symmetrically distributed around RSSs, with depletion of these elements marking both the RSS and two proximal sites localized within 50 bp (Figure 4A). Moreover, depletion of poly(A) stretches at flanking regions was accompanied by features characteristic of nucleosome container sites, such as a central core of GC-rich sequences and a moderate decrease in AT content (Figure 4B; Tillo and Hughes,

2009). In contrast, RSSs reside at the global minimum in GC content and at a local maximum in AT content (Figure 4B), suggesting an enrichment of TpA base-pair steps, which are characterized by the weakest base-stacking interactions among all possible dinucleotides (Rohs et al., 2009). Interestingly, nucleosome containers similarly marked RSSs of human HeLa replication origins (Figures S4A and S4B), thus indicating that this feature is conserved across higher eukaryotes. Taken together, these results strongly suggest an increase in conformational flexibility of RSSs and immediately adjacent regions.

identified in separate biological replicates (Figure S1D), thus demonstrating the high sensitivity of our approach. These data allowed us to quantify the relative contribution of individual fractions to the SNS profile previously observed at G4 motifs (Figure 2C). Strikingly, we found that, while leading strands traveling away from the G4 motif extend normally, DNA synthesis on the strand replicating the G4 template is blocked at the G4 site (Figures 3C and 3D for an example). This finding suggests that most origin-proximal G4s are folded at the time of origin activation and function as replication fork barriers in vivo (Figure 3E).

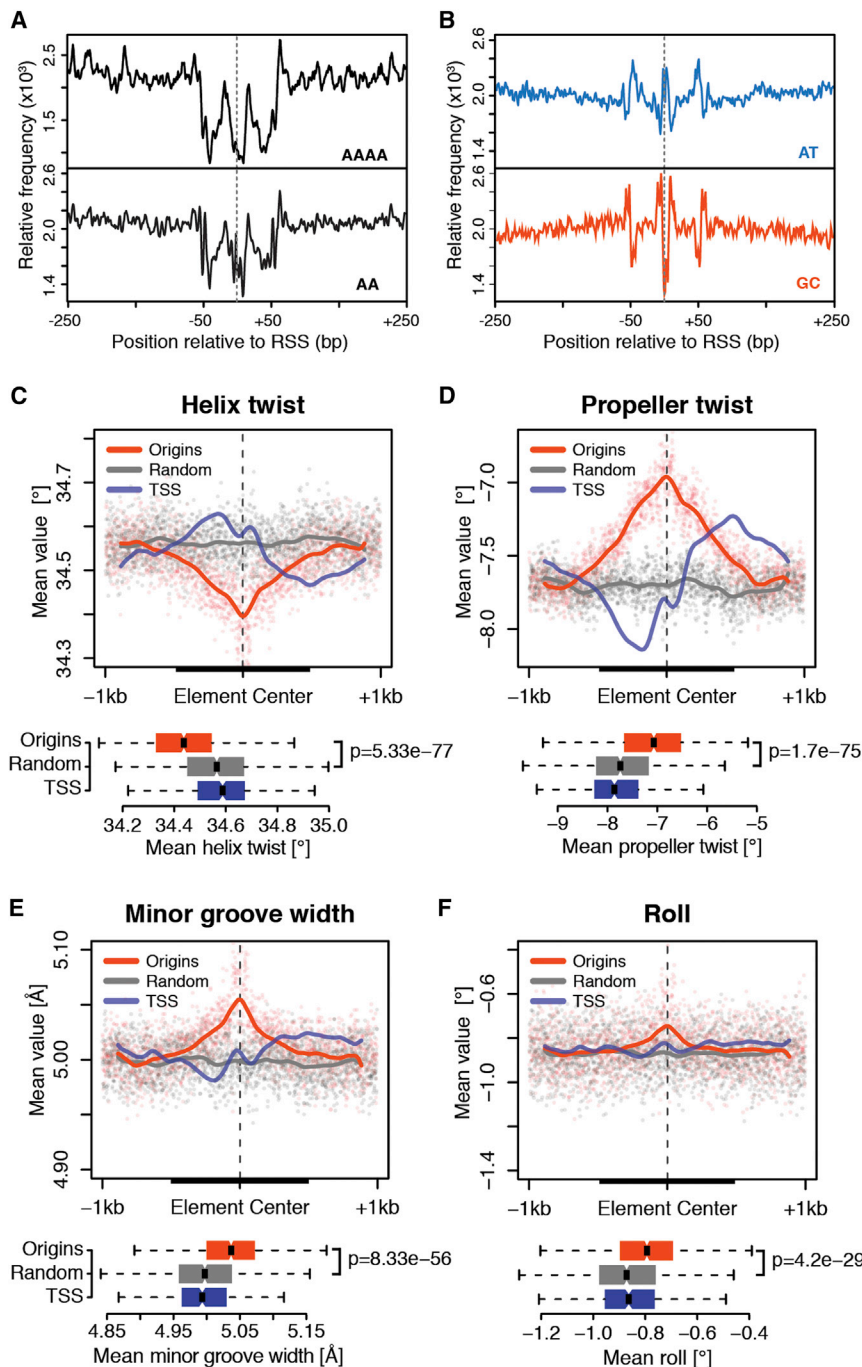


Figure 4. Specific DNA Shape Features Mark Metazoan Replication Origins

(A) Relative frequency of AAAAAA polynucleotides and AA dinucleotides within ± 250 bp of S2 RSSs is shown.

(B) Same as (A) is shown for AT and GC dinucleotides.

(C–F) Average of DNA shape features within ± 1 kb of RSSs for constitutive *Drosophila* origins, background regions, and TSSs. The latter were extended while preserving orientation. Solid lines are Loess fitted curves from single-nucleotide resolution shape predictions (dots). Boxplots of average feature values within 500-bp windows (thick black lines) are shown (bottom) (p values are from Wilcoxon rank-sum test). See also Figure S4.

twist (Figure 4D) suggests a reduction in inter-base pair hydrogen bonds in the major groove, which is the main stabilizing force for the formation of rigid poly(A) elements (Rohs et al., 2009). This, in turn, has been correlated previously with widening of the minor groove at the corresponding positions (Figure 4E; Hancock et al., 2013). Moreover, the local increase in roll (Figure 4F) suggests an enrichment in pyrimidine-purine base-pair steps, such as TpA dinucleotides, thus generating weak stacking interactions that enhance local flexibility of RSSs (Rohs et al., 2010). Together, these data provide compelling evidence that degenerate sequence features dictate a conserved DNA structure that is likely to play a key role in origin function.

The Chromatin Composition of *Drosophila* Replication Origins

Previous studies noted increased chromatin accessibility at sites of early replication in *Drosophila* (Bell et al., 2010; MacAlpine et al., 2010) and identified DNase I-hypersensitive sites (DHSs) as a determinant of replication initiation in human cells (Gindin et al., 2014). In line with these studies, S2 origins were significantly associated with DHSs ($p < 0.001$, Figure 5A), and their firing efficiency posi-

To further test this hypothesis, we generated high-throughput predictions of DNA shape features at origins (Zhou et al., 2013). Strikingly, a specific DNA shape signature common to *Drosophila* (Figures 4C–4F) and human (Figures S4C–S4F) origins emerged, characterized by reduced helix twist and increased propeller twist, minor groove width, and roll. A decrease in helix twist (Figure 4C) indicates helical unwinding, which renders bending and other DNA deformations energetically more favorable (Chen et al., 2013). The increase in propeller

tively correlated with local chromatin accessibility (Figure 5B). Moreover, averaging of DNase-seq signals across 5-kb windows centered on RSSs revealed a strong enrichment for DNase I-digested fragments at origins as compared to randomized genomic regions (Figure 5C). At first glance, these results are incompatible with a nucleosome container signature at initiation sites, and this apparent contradiction prompted us to examine the spatial distribution of DNase-seq signals across RSSs. To our surprise, we found a striking difference between chromatin

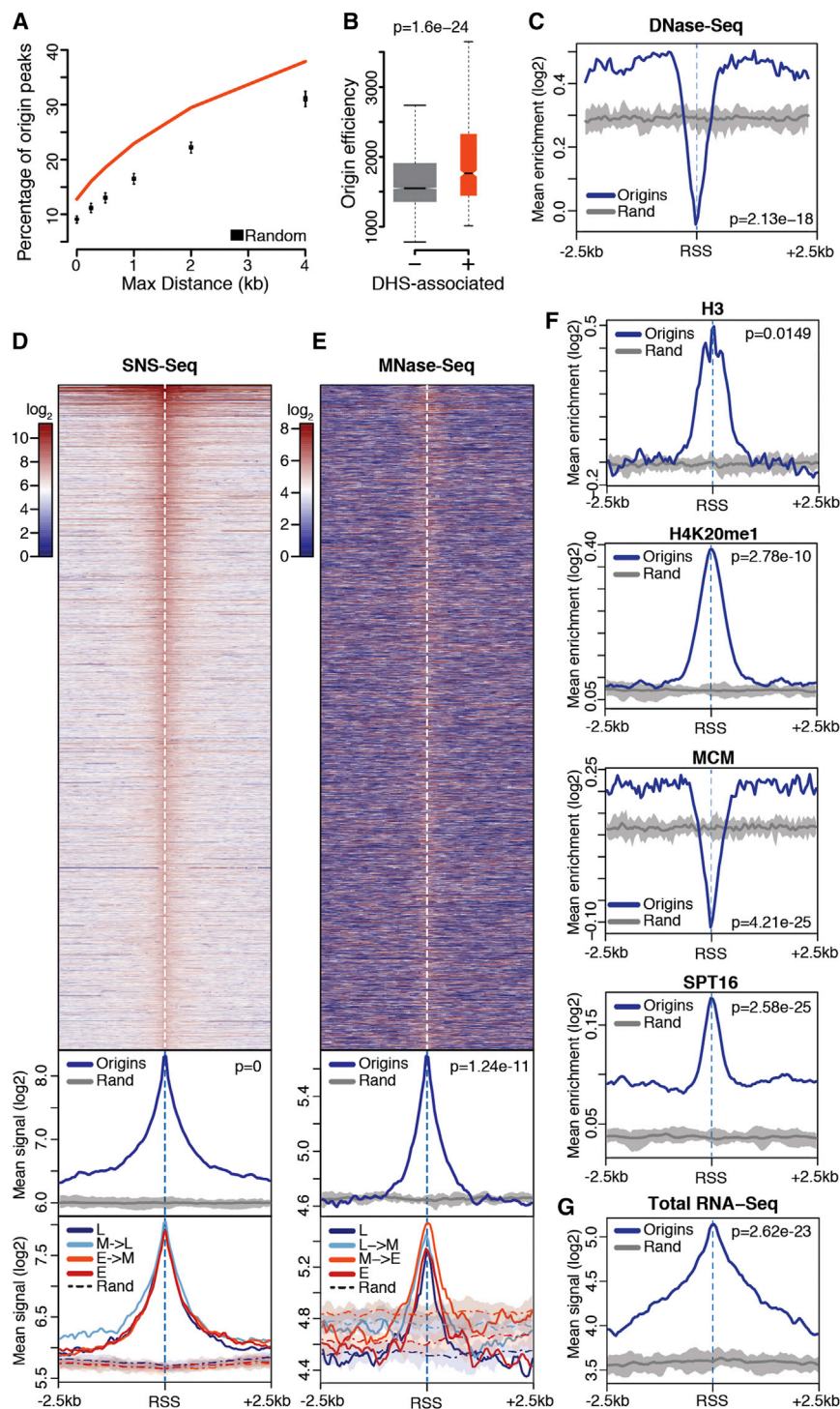


Figure 5. The Chromatin Composition of *Drosophila* Replication Origins

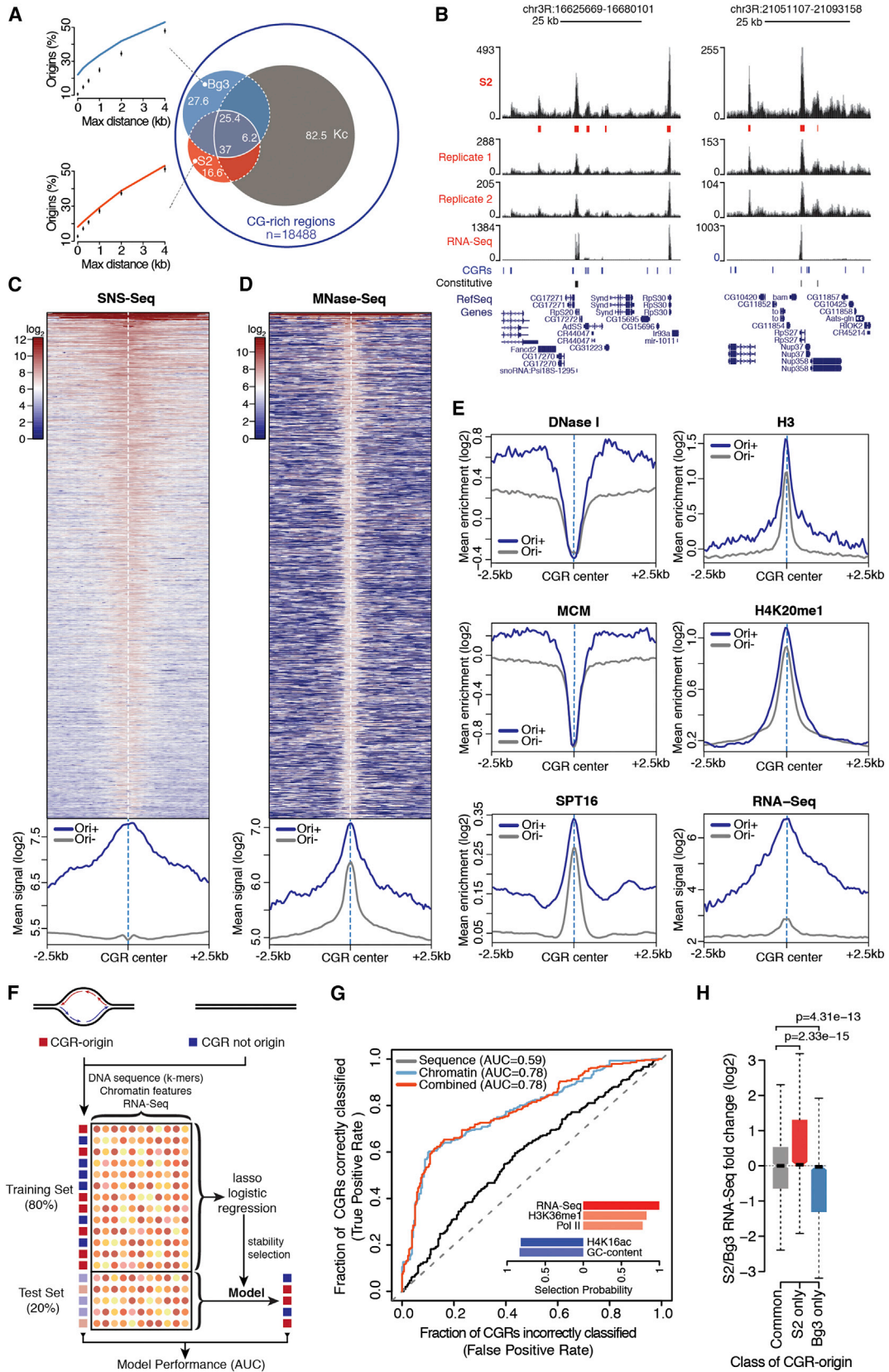
(A) Percentage overlap of S2 origin peaks with DHSs and random expectation are shown. (B) Efficiency of S2 origins localizing within (+) or outside (–) DHSs is shown. (C) Average DNase-seq enrichment within ± 2.5 kb of S2 RSSs and within ten sets of randomized genomic regions (Rand). The thick gray line traces average background values. (D) Spatial distribution of SNS-seq signal within ± 2.5 kb of S2 RSSs (top), metaprofiles comparing origins with ten sets of randomized genomic regions (middle), and further partitioning of the signal above in four timing classes (L, late S-phase; M, mid; E, early) based on replication timing quartiles (bottom) are shown. (E) Same as (D) is shown for MNase-seq. (F and G) Same as (C) is shown for the indicated features (p values are from Wilcoxon rank-sum test). See also [Figure S5](#).

To test this hypothesis, we set out to survey the chromatin landscape of replication origins at an unprecedented resolution. First, we compiled a comprehensive representation of the chromatin landscape of S2 cells comprising 85 chromatin features profiled by the modENCODE project (Celniker et al., 2009) or independent studies (Table S2). Second, we analyzed the spatial distribution of each feature within 5-kb windows centered on inferred RSSs or in ten sets of matched control regions at 50-bp resolution. The potentially confounding contribution of TSS-associated chromatin features was limited by excluding origin-TSSs from the analysis. Third, as chromatin features do not uniformly distribute across replication-timing compartments, we partitioned origins and control regions in four timing classes (from early to late replicating) based on replication-timing quartiles. This allowed us to probe for evidence of timing-specific chromatin signatures by directly comparing matched timing classes.

As expected, origins were strongly enriched for SNS compared to control regions, with SNS-seq signals sharply peaking at the inferred RSS positions (Figure 5D). However, nearly no difference

accessibility of RSSs and their flanking regions. Indeed, while the latter exhibit clear features of open chromatin, a sharp reduction in DNase I-digested fragments was seen at the RSS (Figure 5C). These results reconcile our observations and led us to posit that a rich, spatially organized chromatin configuration marks eukaryotic origins.

in the average SNS-seq signal was observed across timing classes, indicating that firing efficiency does not correlate with replication timing at the single origin level. Next, we focused on direct and indirect measurements of nucleosome occupancy. Interestingly, while the genomic regions flanking the RSSs exhibit background MNase-seq signals (Figure 5E) and histone H3/H4



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enrichments (Figures 5F and S5A), RSSs correspond to positions of high nucleosome occupancy, a feature that is shared across timing classes. Moreover, we found that nucleosomes at RSSs are decorated by histone modifications, such as lysine mono-methylation and acetylation. H3K9me1, H3K23me1, and H4K20me1, a PR-Set7-dependent histone modification that promotes loading of the pre-RC at origins (Tardat et al., 2010), sharply peaked at RSSs (Figures 5F and S5B). Intriguingly, enrichment of these marks was not limited to early replicating regions (Figure S5B), suggesting that initiation sites could be invariably bookmarked by these modifications. In contrast, pre-RC binding was restricted to the accessible chromatin regions flanking RSSs (Figures 5F and S5C), in line with the concept that ORC-mediated pre-RC nucleation requires direct contact with the DNA template (Masai et al., 2010) and occurs adjacent to RSSs (Lombraña et al., 2013).

Rapid nucleosome turnover has emerged as a distinguishing feature of both promoters and origins (Deal et al., 2010), and binding of chromatin remodelers was previously correlated with early replication timing in *Drosophila* (Eaton et al., 2011; Comoglio and Paro, 2014). In agreement, members of different remodeling complexes and the histone chaperone Spt16, a core component of the facilitates chromatin transcription (FACT) complex, were markedly enriched at origins (Figures 5F and S5D), with the highest values throughout early replicating, active chromatin. However, while the overall enrichment of these proteins largely varied across timing classes, RSSs were invariably marked by the highest occupancy and exhibited similar enrichments relative to flanking regions in each timing class (Figure S5E). Finally, as the correlation between transcription and origin firing remains elusive, we leveraged our high-resolution data to examine local transcriptional outputs at origins. Strikingly, analysis of the spatial distribution of both total and poly(A)⁺ RNA-seq signals at origins revealed conspicuous transcription at RSSs, once again, irrespective of their replication timing (Figures 5G and S5F).

Chromatin Landscape and Transcriptional Output Predict Origin Activity of CG-Rich Regions

Several CpG islands (CGIs) in mammals and CG-rich regions (CGRs) in *Drosophila* share the potential to initiate DNA replication (Cayrou et al., 2012a; Besnard et al., 2012). Indeed, 18% of S2 and 22% of Bg3 origin peaks were significantly associated with CGRs ($p < 0.001$, Figure 6A). However, only 6%–37% of origin CGRs are highly efficient, constitutive origins (Figures 6A

and 6B), raising the question of which features might favor or prevent replication initiation at these sites. Here we asked whether the local chromatin context at CGRs correlates with their firing potential.

We started by contrasting the chromatin landscape of origin CGRs active in S2 cells (Figure 6C) with that of origin-negative CGRs. Intriguingly, contrary to our expectation, we found that chromatin was overall similarly configured across CGRs, irrespective of origin activity (Figures 6D, 6E, and S6A). A few noteworthy features, therefore, likely render the chromatin configuration of origin CGRs compatible with efficient origin firing: (1) higher chromatin accessibility and higher pre-RC loading proximal to the CGR center (a proxy for the RSS), (2) higher nucleosome occupancy, and (3) markedly higher levels of Spt16 throughout the entire origin region that sharply peaked at the CGR center (Figure 6E). Moreover, while origin-negative CGRs were on average poorly transcribed, origin CGRs were strongly enriched for RNA-seq reads (Figure 6E).

These findings led us to test whether chromatin configurations and transcriptional outputs could predict the firing potential of CGRs. To this purpose, we trained binary classifiers based on lasso logistic regression (Tibshirani, 1996), using DNA sequence content (k -mers, $k \leq 4$), chromatin feature enrichments, and RNA-seq signal at CGRs as predictors. A test set of CGRs that was not previously seen by the models was used to evaluate performances (Figure 6F, see Experimental Procedures). Interestingly, while the DNA sequence content of CGRs was a poor predictor of origin activity (area under the receiver operating characteristic curve, AUC = 0.59), chromatin features and transcription were able to accurately classify CGRs (AUC = 0.78). Moreover, a more-complex model combining genetic and epigenetic features did not perform better than epigenetic features alone (AUC = 0.78), indicating that these two sets of features are highly redundant. Next, we unbiasedly assessed the importance of individual predictors by estimating feature selection probabilities with bootstrap-lasso (Comoglio and Paro, 2014). Intuitively, the more a feature is required for accurate predictions, the higher its selection probability. Our analysis identified RNA-seq, H3K36me1, and RNA polymerase II (Pol II) as top-ranked, positive predictors of origin firing at CGRs. In contrast, local GC content and H4K16ac were stably selected, negative predictors of origin activity (Figure 6G, inset). Further analysis of the H4K16ac distribution at CGRs revealed a striking contrast between early- and late-replicating origin CGRs, which were depleted and enriched for this mark, respectively (Figure S6B).

Figure 6. Origin Activity of CGRs Is Predicted by Chromatin Landscape and Transcriptional Output

- (A) Percentage overlaps of origin peaks associated with CGRs in S2, Bg3, and Kc (Cayrou et al., 2011) cells, and comparison of the observed overlap between S2 and Bg3 origin peaks and CGRs (lines) with random expectation (boxplots) are shown (n, total number of CGRs).
- (B) Two representative snapshots of the S2 SNS-seq coverage from two biological replicates, origin peaks, and poly(A)⁺ RNA-seq coverage across several CGRs. Constitutive origins (black) are also shown.
- (C) Spatial distribution of SNS-seq signal within ± 2.5 kb of S2 origin CGR midpoints (top) and metaprofiles (bottom) comparing origin CGRs (Ori+) with origin-negative CGRs (Ori-) are shown.
- (D) Same as (C) is shown for MNase-seq.
- (E) Same as bottom of (C) is shown for the indicated features.
- (F) An outline of the modeling strategy used to classify CGRs is shown.
- (G) ROC curves and AUC values for lasso models trained on the indicated sets of features. The inset shows selection probabilities of the top-ranked features selected by bootstrap-lasso. Bars are color-coded according to coefficient signs (positive, red; negative, blue) and absolute value of coefficient Z scores.
- (H) S2/Bg3 RNA-seq fold changes for the indicated classes of origin CGRs are shown (p values are from Wilcoxon rank-sum test). See also Figure S6.

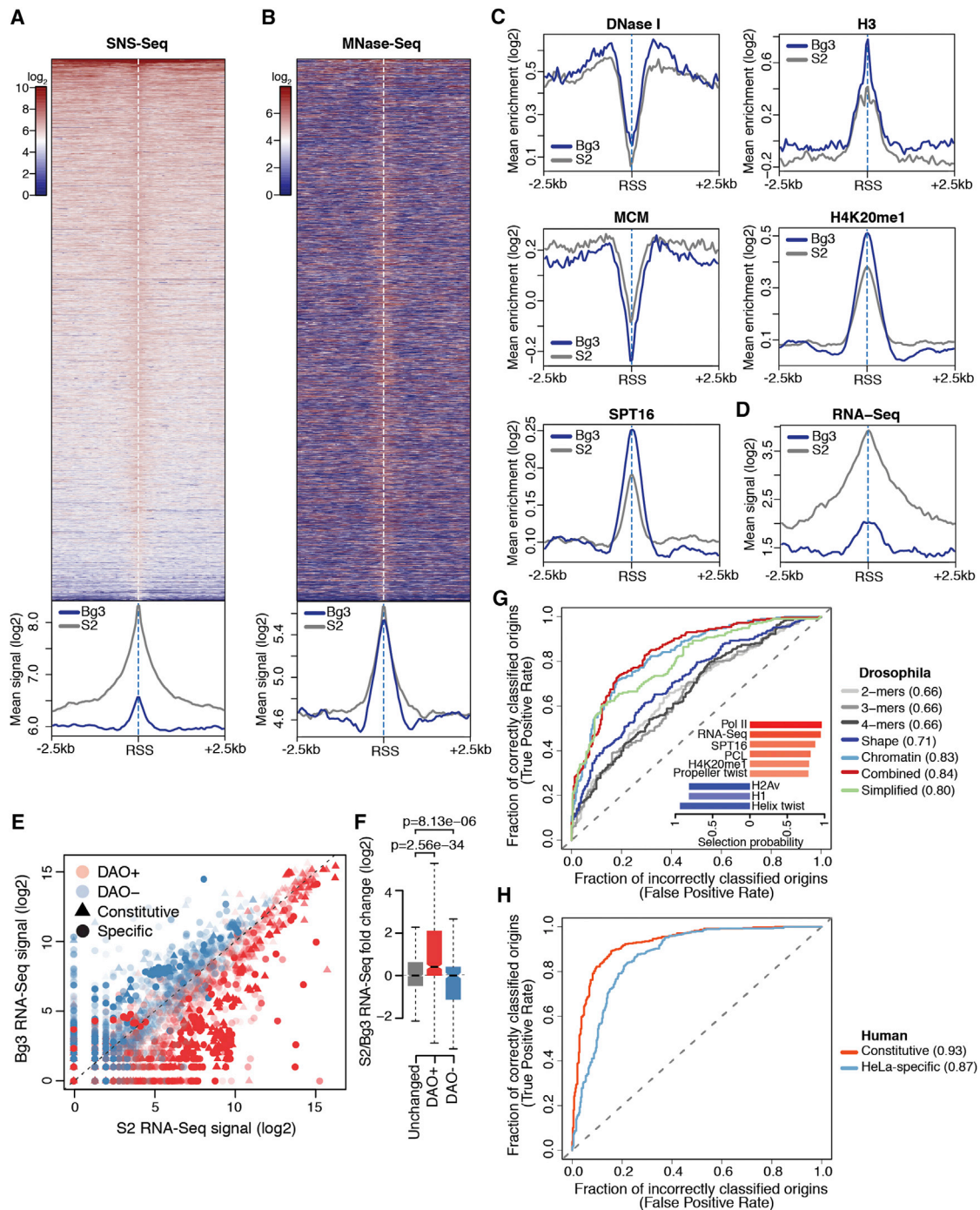


Figure 7. Differential Origin Activity Mirrors Differences in Cell-Type-Specific Transcriptional Programs

(A) Spatial distribution of S2 SNS-seq signal within ± 2.5 kb of RSSs of origin peaks solely identified in Bg3 cells (top) and metaprofiles (bottom) comparing all S2 origin peaks with these sites are shown.

(B) Same as (A) is shown for MNase-seq.

(C and D) Same as bottom of (A) is shown for the indicated features.

(E) Scatter plot shows S2 and Bg3 RNA-seq signals at DAOs that were more efficiently used by S2 (DAO+) or Bg3 (DAO-) cells (triangles, constitutive origins; circles, origin peaks solely detected in one cell type). Opacity reflects the statistical significance of differential origin activity and is proportional to $-\log_{10}$ -transformed adjusted p values.

(F) S2/Bg3 RNA-seq fold changes of equally activated origins (unchanged) and of differentially activated ones are shown (p values are from Wilcoxon rank-sum test).

(legend continued on next page)

As the RNA-seq signal was stably selected by all models, we investigated whether differential expression of origin CGRs could explain differential usage of these sites across cell types. To this end, we computed the S2/Bg3 RNA-seq fold change of origin CGRs that efficiently fired in both cell types or that were efficiently activated only in one of the two. Intriguingly, while the former were on average similarly transcribed in S2 and Bg3 cells, transcription of the latter was significantly upregulated in the cell type of efficient activation (Figure 6H). Taken together, these results establish a tight coupling between transcription and origin firing at origin CGRs.

Differential Origin Activity Mirrors Differences in Cell-Type-Specific Transcriptional Programs

Bg3-preferred origins exhibited poor, yet highly significant, firing efficiency in S2 cells (Figure 1D). Indeed, analysis of S2 SNS-seq signals at these sites indicated that virtually all Bg3-preferred origins also fire in S2 cells (Figure 7A). Therefore, we reasoned that a systematic comparison between these low-efficiency sites and S2 origin peaks could shed light on poorly understood epigenetic determinants of origin firing.

Dissection of the chromatin configuration of these two origin sets revealed an enrichment of H3K9me3, Su(var)3-9, and increased binding of insulator proteins (Figure S7A) at Bg3-preferred origins in S2 cells compared to S2 origin peaks, suggesting that these sites are preferentially embedded within constitutive heterochromatin in S2 cells. However, despite their heterochromatic localization, the distinctive chromatin signature previously identified at S2 origin peaks invariably marked Bg3-preferred origins in the S2 epigenome (Figures 7B and 7C). In fact, Bg3-preferred origins shared several features with efficient origins, including accessible flanking regions as well as high H4K20me1, Spt16, and chromatin remodeler levels (Figures 7C and S7B). Conversely, a markedly lower transcriptional output distinguished Bg3-preferred origins from S2 origin peaks (Figure 7D), suggesting that a chromatin environment less permissive to transcription might suffice to prevent efficient origin firing irrespective of local chromatin cues. These results, along with convergence of transcription and replication programs at CGRs, prompted us to test whether differential origin activity across cell types could similarly mirror differences in the cell-type-specific transcriptional outputs genome-wide. To this end, we identified 5,917 differentially activated origins (DAOs, see Experimental Procedures) between S2 and Bg3 cells. Notably, DAOs encompassed 38% of constitutive origins and >49% of origins efficiently firing only in one cell type at a very stringent significance threshold (adjusted p value $\leq 1e-5$). When S2 and Bg3 RNA-seq signals were compared at DAOs, a clear-cut correlation emerged between differential origin activity and transcriptional output (Figure 7E). Indeed, while equally activated origins were similarly transcribed in S2 and Bg3 cells (Figures 7F and S7C), DAOs that were more efficiently used by S2 cells were, in turn, signif-

icantly more transcribed in this cell type than in Bg3, with the opposite trend being observed at DAOs more efficiently used by Bg3 cells (Figure 7F).

DNA Shape and Epigenetic Features Accurately Predict Active Origins in the *Drosophila* and Human Genomes

Next, we asked whether the identified genetic and epigenetic characteristics of replication origins could discriminate active replication initiation sites from the rest of the *Drosophila* genome. To this end, we trained lasso origin classifiers on DNA *k*-mers and shape features, chromatin feature enrichments, and transcriptional output, using the same learning scheme of CGR classifiers (Figure 6F).

Intriguingly, DNA shape features alone not only exhibited a moderately high predictive power (AUC = 0.71), but they also outperformed DNA sequence content (AUC = 0.66) in classifying constitutive origins (Figure 7G). Moreover, chromatin feature enrichments and transcription were able to generate accurate origin predictions (AUC = 0.83). Further inclusion of *k*-mers only marginally improved model performances (AUC = 0.84) (Figure 7G). Indeed, no *k*-mer was consistently selected in this more-complex model. Conversely, a simplified origin classifier solely based on the nine most stably selected features (selection probability ≥ 0.8), including RNA-seq signal, six chromatin features, and two DNA shape features (Figure 7G, inset), generated remarkably accurate predictions (AUC = 0.80, Figure 7G). These results led us to examine whether local DNA shape and transcription at origins could similarly predict active origins in the human genome. Strikingly, a classifier solely based on helix twist, propeller twist, and RNA-seq signal was not only able to discriminate active origins from the rest of the human genome, but it also outperformed the *Drosophila* origin classifier at both constitutive (AUC = 0.93) and HeLa-specific (AUC = 0.87) origin peaks (Figure 7H). Interestingly, model performances correlated with the DNA shape profiles observed within these origin sets, with poorly efficient origins exhibiting a less-pronounced DNA shape signature at RSSs (Figure S4).

DISCUSSION

Origin specification in higher eukaryotes involves mechanisms other than simple replicator-initiator interactions (Leonard and Méchali, 2013), yet the genetic and epigenetic features that specify replication origins in vivo remain enigmatic. ORC is the first replication factor to bind origins but it lacks a sequence-specific binding motif (Gilbert, 2010). Interestingly, studies have suggested that DNA topology rather than sequence motifs might mediate nucleation of the pre-RC at origins. Mainly, *Drosophila* ORC exhibits much higher affinity for negatively supercoiled DNA than for linear or relaxed DNA in vitro (Remus et al., 2004), and human ORC binds preferentially to G4-forming RNA and single-stranded DNA (Hoshina et al., 2013). Moreover, these findings are reminiscent of observations in bacteria where

(G) ROC curves and AUC values for origin classifiers trained on the indicated set of features in *Drosophila*. The inset shows selection probabilities of the top-ranked features selected by bootstrap-lasso and used to train the simplified model. Bars are color-coded according to coefficient signs (positive, red; negative, blue) and absolute values of coefficient z-scores.

(H) Same as (G) is shown for constitutive and HeLa-specific human origins. See also Figure S7.

negatively supercoiled DNA coordinates replication initiation (Mott and Berger, 2007).

By high-throughput predictions of DNA shape, our study demonstrates that a specific DNA topology, characterized by increased conformational flexibility of RSSs, marks both *Drosophila* and human replication origins. This relaxed DNA conformation likely serves two important functions. First, positioning of nucleosomes at the RSS, previously observed at efficient mammalian origin CGIs (Lombrana et al., 2013), is likely to be weaker in this region, thus favoring nucleosome displacement. Formation of atypical nucleosomes at RSSs could also contribute to this process. Second, melting of the double helix at the RSS is energetically assisted, further facilitating initiation. Interestingly, while enhanced DNA flexibility similarly marks *Drosophila* (Figure 4) and human (Figure S4) origins, the sequence composition of RSSs is remarkably different between these two organisms. Indeed, while *Drosophila* RSSs are locally AT-rich, high GC content is found at human RSSs (Figure S4B). Intriguingly, the increased GC content of human RSSs might reflect a role for cytosine methylation of CpG dinucleotides, which is widespread in mammals but not *Drosophila* (Takayama et al., 2014), in granting conformational flexibility to human origin sequences, thus functionally replacing TpA base-pair steps (Lazarovici et al., 2013). Together, these findings indicate that by integrating over degenerate sequence features, DNA shape appears to represent a universally conserved origin bookmark.

A confounding feature of metazoan origins, however, is that mechanisms of origin specification are likely not sufficient in defining origin activity. Origin firing is intrinsically stochastic at the single-cell level (Bechhoefer and Rhind, 2012; Cayrou et al., 2011), yet aggregated firing probabilities within a replicon could be modulated by extrinsic factors. A major question in the field is which *cis*- and *trans*-acting factors influence these probabilities. G4s and noncoding RNAs (Ge and Lin, 2014) are likely candidates. G4s have been proposed to orient replication forks and enhance the efficiency of origin firing at the chicken β^A promoter (Valton et al., 2014), but they are not sufficient for origin activation. Our data suggest that origin-associated G4s determine the precise position of replication initiation at a subset of origins (Figure 2D). However, single-fraction SNS-seq experiments indicated that synthesis of the leading strand pauses at origin-associated G4 motifs *in vivo* (Figure 3), despite the presence of Pif1 and other G4-unwinding helicases (Maizels and Gray, 2013). Replication fork stalling therefore might be responsible for the observed accumulation of SNSs at G4-associated origins (Valton et al., 2014). Moreover, it provides an alternative explanation to the repeatedly observed, yet enigmatic, higher firing efficiency of these sites (Figure S2B; Besnard et al., 2012; Valton et al., 2014). Our finding is unlikely to be a technical artifact. First, SNS purifications underwent up to five rounds of Lexo digestion. Second, SNSs were absent from non-dividing cells and degraded upon RNase or alkali treatment (Cayrou et al., 2011, 2012b). Third, recent work based on independent assays revealed transient replication fork stalling at exogenous G4 sequences *in vitro* (Castillo Bosch et al., 2014).

Whereas most studies have focused on the relation between chromatin features and replication timing (Bell et al., 2010; Eaton et al., 2011; Gindin et al., 2014), little is known about the chro-

matin configuration of metazoan origins. Moreover, conflicting evidence has been reported on the role of transcription in origin firing (Sequeira-Mendes et al., 2009; Martin et al., 2011; Lubelsky et al., 2014). An answer to this question likely depends on the genomic scale at which conclusions are drawn. Here we show that a specific chromatin configuration similarly marks efficient and poorly used origins, irrespective of replication timing (Figures 5, 6, and 7). However, differential transcription at RSSs discriminates active from inactive origin CGRs, explaining differential usage of these sites (Figure 6), and reflects differential origin activity across cell types (Figure 7). These findings reinforce the convergence of transcriptional and replication programs at replication origins, and they support a model in which DNA shape and chromatin features primarily define origin localization. Transcriptional output at RSSs, in contrast, likely contributes to modulate aggregated firing probabilities, which could be directly reflected in the number of MCM molecules recruited at the origin site (Bechhoefer and Rhind, 2012).

While our study does not establish a causal link between transcription and origin firing, we anticipate that the genome-wide datasets reported here will facilitate mechanistic studies of the interplay between transcription and origin function in higher eukaryotes.

EXPERIMENTAL PROCEDURES

Cells and Cell Culture

Drosophila S2-DRSC cells and ML-DmBG3-c2 cells were cultured at 25°C in 145-mm plates (Greiner) at a density of 1.5×10^6 cells/ml in Schneider's Insect Cell Medium (Sigma S0146) and Shields and Sang M3 insect medium (Sigma S3652) with 10 μ g/ml insulin (Sigma), respectively, both supplemented with 10% fetal bovine serum (FBS, Pansera ES).

SNS Purification

SNS fragments were isolated essentially as described in Cayrou et al. (2011), with the following modifications. Adjustments were made according to SW-41 Ti rotor specifications (Beckman Coulter) and centrifugation was carried out at 4°C and 26,700 rpm for 21 hr. DNA fragments of 0.5–2.5 kb were collected, purified, and subjected to four/five rounds of T4 PNK (Fermentas) phosphorylation and lambda Exonuclease (Fermentas) digestion. SNSs were then prepared for sequencing by digesting the RNA-primer, second-strand synthesis (NEBNext mRNA Second Strand Synthesis) and purified with AMPure XP beads (Beckman Coulter). This procedure ensures that RNA species do not contribute to SNS-seq signals. For a complete description of the experimental procedures, see the Supplemental Experimental Procedures.

Data Analysis and Statistical Learning

S2 and Bg3 SNS-seq reads were aligned to the dm3 *Drosophila* reference genome using Bowtie2 (Langmead and Salzberg, 2012). Human SNS-seq data (Besnard et al., 2012) were mapped to the hg19 human reference genome. Origin peaks were called using MACS (Zhang et al., 2008). The coverage function was used to define the position of the RSS within each origin peak, and read counts normalized to the peak length in kilobases were used as a proxy for origin efficiency. G4 motif occurrences in the *Drosophila* genome were predicted with QuadParser (Huppert and Balasubramanian, 2005). DNA shape features were obtained from high-throughput predictions (Zhou et al., 2013). CGRs were predicted according to Gardiner-Garden and Frommer (1987). Feature scoring was performed essentially as described previously (Comoglio and Paro, 2014). Lasso logistic regression models were trained with 10-fold cross-validation. Feature selection probabilities were computed with bootstrap-lasso (Comoglio and Paro, 2014). Differential origin activity analysis was carried out using DESeq2 (Love et al., 2014). For a complete description of

the algorithms and analysis procedures, see the [Supplemental Experimental Procedures](#).

ACCESSION NUMBERS

The accession number for the SNS-seq data reported in this paper is GEO:GSE65692.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, seven figures, and two tables and can be found with this article online at <http://dx.doi.org/10.1016/j.celrep.2015.03.070>.

AUTHOR CONTRIBUTIONS

F.C., T.S., C.B., and R.P. designed the study. F.C. carried out most of the experimental work with the help of T.S. and V.S. F.C. carried out computational and statistical analyses and wrote the manuscript with the help of T.S., R.P., and R.R., who also contributed to the DNA shape analysis.

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