Genome-wide depletion of replication initiation events in highly transcribed regions

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This report investigates the mechanisms by which mammalian cells coordinate DNA replication with transcription and chromatin assembly. In yeast, DNA replication initiates within nucleosome-free regions, but studies in mammalian cells have not revealed a similar relationship. Here, we have used genome-wide massively parallel sequencing to map replication initiation events, thereby creating a database of all replication initiation sites within nonrepetitive DNA in two human cell lines. Mining this database revealed that genomic regions transcribed at moderate levels were generally associated with high replication initiation frequency. In genomic regions with high rates of transcription, very few replication initiation events were detected. High-resolution mapping of replication initiation sites showed that replication initiation events were absent from transcription start sites but were highly enriched in adjacent, downstream sequences. Methylation of CpG sequences strongly affected the location of replication initiation events, whereas histone modifications had minimal effects. These observations suggest that high levels of transcription interfere with formation of pre-replication protein complexes. Data presented here identify replication initiation sites throughout the genome, providing a foundation for further analyses of DNA–replication dynamics and cell-cycle progression.

[Supplemental material is available for this article.]

DNA replication is tightly orchestrated to duplicate the entire genome precisely once each somatic cell cycle. In eukaryotes, replication initiates at multiple sites on each chromosome in a temporally regulated and often tissue-specific manner. Approximately 30 replication initiation sites in the human genome have been identified and characterized in some detail (Aladjem et al. 2006; Hamlin et al. 2008). Additional replication initiation sites have been identified using high-throughput targeted microarray analyses (Cadoret et al. 2008; Karnani et al. 2009; Gilbert 2010). Identified replication initiation sites in human cells do not share clear consensus sequences. In contrast, replication initiation sites exhibit sequence similarities in single-celled eukaryotes such as yeast. In yeast, consensus replication initiation yeast sequences are characterized by a unique, asymmetric AT-rich sequence, leading to a distinct pattern of nucleosome positioning (Eaton et al. 2010). However, even in yeast, replication is limited to a subset of potential initiation sites exhibiting the consensus sequence, and in many instances, potential yeast replication origins do not initiate replication. In metazoans, strong DNA sequence similarity is not observed among replication initiation sites, suggesting that primary DNA sequence is not the sole determinant of replication initiation competence (Aladjem 2007; Mechali 2010). Replication initiation sites in metazoa share

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several DNA sequence motifs, including AT-rich sequences, matrix attachment sites, and asymmetric purine:pyrimidine sequences (Aladjem 2007). None of these motifs, however, is absolutely essential for replication initiation. Sequence variation within known replication initiation sites suggests that the regulation of site localization and the timing of DNA replication initiation may involve epigenetic mechanisms.

DNA replication may be regulated epigenetically to allow transcription and replication to proceed in a coordinated fashion. In mammalian cells, mapping studies suggest that replication initiation events are enriched in transcription factor binding sites (Cadoret et al. 2008; Karnani et al. 2009; Gilbert 2010; Valenzuela et al. 2011) and CpG islands (Antequera 2004; Gomez and Brockdorff 2004). Consistent with this observation, distal sequences involved in transcription are required for replication initiation at a number of loci, including a region 40 kb upstream of the human betaglobin (HBB) replication origin (Aladjem et al. 1995), the promoter of the Chinese hamster Dhfr locus (Kalejta et al. 1998), and an enhancer of the Th2 locus (Hayashida et al. 2006). In several biological contexts, however, transcription appears to suppress replication initiation. This occurs during differentiation of Xenopus (Hyrien et al. 1995) and Sciara (Lunyak et al. 2002) embryos, at the Chinese hamster Dhfr locus (Saha et al. 2004), and at the murine HoxB locus (Gregoire et al. 2006). Conversely, at the human IGH locus (Zhou et al. 2002) replication initiates in an expanded region after the onset of transcription; and on human (Gray et al. 2007) and murine (Rowntree and Lee 2006) X chromosomes, replication

initiates from distinct origins regardless of transcriptional status. Finally, at the human *MYC* locus, transcription factor binding seems to activate initiation of DNA replication, whereas high transcription rates suppress it (Ghosh et al. 2004).

To investigate the relationship between replication initiation and transcription, we measured replication density and transcriptional activity in nonrepetitive genomes of two human cancer cell lines, which have different patterns of gene expression. Replication initiation events were more frequent and clustered near regions of moderate transcription. In contrast, low levels of replication initiation were found in genomic regions either devoid of transcription, or with very high levels of transcription. High-resolution analyses of initiation event distribution also suggested that replication initiation sites were enriched near promoters, but excluded from transcription start sites (TSSs). Finally, replication tended to initiate in genomic regions containing methylated CpG sequences, which are frequently associated with low rates of transcription

Results

Visualization of replication initiation sites

To map replication initiation at a genome-wide scale, short, RNAprimed nascent DNA was isolated from two human cell lines (myeloid erythroleukemia K562 cells and breast cancer MCF7 cells) (Bielinsky and Gerbi 1998; Wang et al. 2004) and analyzed using massively parallel sequencing. The frequency of initiation events at the individual genomic regions was measured as the ratio between reads obtained from a nascent strand preparation and reads obtained from a corresponding control genomic DNA preparation. Reads were calculated as reads per kilobase per million mapped reads (RPKM). The number of reads and aligned reads from K562 and MCF7 cells are shown in the Supplemental Methods (Supplemental Table 1). Results were visualized using the Integrative Genome Viewer 1.4.1 (Broad Institute), and data validated by analyzing initiation at well-characterized replication initiation sites (Fig. 1; Supplemental Fig. 1).

We first analyzed replication profiles at several, well-characterized genomic loci. At these loci, replication has been mapped several times previously by several independent research groups using methods that rely on nonoverlapping principles. Figure 1A shows the replication initiation ratio (nascent strands vs. genomic RPKM) at the MYC locus on human chromosome 8. At this locus, DNA replication initiates from several sites within and around the promoter and within the first exon of the gene (Ghosh et al. 2004). As expected, our analysis showed that the replication initiation ratio was high in this region, and several replication initiation peaks were observed, consistent with previous observations (e.g., Ghosh et al. 2004). Similarly, replication initiation ratios near the HBB gene (chromosome 11) were highest near the promoter (Fig. 1B), consistent with previous data (Kitsberg et al. 1993; Aladjem et al. 1995). Finally, Figure 1C shows the replication profile at the HPRT1 locus on the X chromosome. Consistent with previous observations (Cohen et al. 2004), replication preferentially initiated near the promoter region.

Although replication initiation sites seemed clustered in some loci, we also identified very large genomic regions that were devoid of replication initiation events. For example, replication initiation sites were entirely absent from the *CTCF* gene, a region that spans \sim 120 kb on chromosome 16 (position 67564943–67704608) (Fig. 1D). Irregular clustering of replication initiation events such as this was evident throughout the genome.

Replication initiation events and transcriptional activity

To compare replicative and transcriptional activities, data from Affymetrix Human Genome U133 Plus 2.0 microarrays (Shankavaram et al. 2007) were used to determine gene expression values for K562 and MCF7 cells. Replication initiation data for chromosome regions within 3 kb of known genes were then aligned with the gene expression data. Figure 2, A and B shows that in MCF7 cells, regions with very low transcriptional activity were not enriched in replication initiation events, and that replication density generally increased with gene expression. Peak initiation activity coincided with moderate expression levels. Replication initiation events were again less frequent in genes with high levels of transcriptional activity. Similar results were obtained with K562 cells (Fig. 2C,D). In addition, no replication initiation rate differences were found when 5' and 3' flanking gene regions were compared or when chromosomal regions between two genes transcribed either in the same or opposite directions were compared (data not shown). In general, our data suggest that replication initiation events are more frequent in moderately transcribed regions and less frequent in regions exhibiting either high or low levels of transcription.

Replication initiation and TSSs

The relationship between replication initiation and distance from a TSS was examined for all predicted genes in K562 and MCF7 cells. Replication initiation events were generally enriched in regions within 10 kb of a TSS (Supplemental Fig. 1B). However, higherresolution analyses showed that in MCF7 cells, the replication initiation ratio decreased several hundred base pairs upstream of the TSS, and peaked \sim 500 bp downstream from the TSS (Fig. 3A). As RPKM values around the TSS were generally elevated in control samples (genomic DNA), the decreased RPKM in nascent strands versus the genomic control was particularly striking (for details, see Supplemental Information). The replication initiation pattern at the TSS indicates that replication initiation events are excluded from regions that nucleate transcriptional complexes. To evaluate whether the depletion of initiation events at the TSS was associated with transcription activity, the analysis was repeated using four groups of genes with different levels of expression. For the purpose of this analysis, log₂ GCRMA normalized expression values of <2.3, 2.3–5.3, 5.3–8.5, or >8.5 were classified as very low, low, medium, or high, respectively. For genes with very low and low levels of transcriptional activity, initiation events were not excluded from the TSS, and instead, a preference for initiation near the TSS was detected (Fig. 3B). In contrast, a marked reduction in initiation activity was observed at the TSS in genes with medium and high levels of transcription. Similar results were observed in nascent strands from K562 cells (Fig. 3C,D). These observations suggest that the depletion of replication initiation events from regions adjacent to the TSS are directly associated with transcriptional activity, and possibly the assembly of transcription initiation complexes.

Replication initiation activity in discordantly expressed genes

We next tested whether genomic regions that exhibit discordant levels of gene expression between K562 and MCF7 cells exhibit disparate frequencies of replication initiation events. We calculated the replication initiation activity of discordantly expressed genes, defined as genes that exhibit (1) high expression in MCF7 cells and low expression in K562 cells (MCF7_H:K562_L), or (2) low expression in MCF7 cells and high expression in K562 cells (MCF7_L:K562_H). High and low levels of gene expression were defined as >6.3 and Α







Figure 2. Replication and transcription. (*A*,*B*) Replication enrichment ratios (nascent strand [NS] versus genomic control RPKM) for all identified genes in MCF7 cells plotted against \log_2 GCRMA normalized gene expression. Genes on the *x*-axis were binned according to gene expression, with each bin containing 897 genes. The first column, however, represents a combination of the first five bins, which includes 4485 low-expressing genes that did not show significant differences in \log_2 GCRMA normalized gene expression. (*A*) The mean enrichment ratio for each bin is plotted against gene expression. (*B*) Replication enrichment ratios calculated as in *A*, showing the distribution of enrichment ratio values as a box plot. (*C*,*D*) Replication enrichment ratios calculated as in *A* and *B* for K562 cells plotted as a histogram of mean values (*C*) or as a box plot (*D*). For mean value histograms (*A*,*C*), asterisks represent statistically significant (*P* < 0.001) divergence from the central bin, which is marked with an arrow. For box plots (*B*,*D*), boxes indicate distributions of the average enrichment ratio of the entire genome.

<4.3 log₂ GCRMA values, respectively. Genes that exhibited similar GCRMA values were defined as concordant genes. As shown in Table 1, a substantial fraction (58.5%) of genes that exhibited high expression in MCF7 cells and low expression in K562 cells had a higher frequency of initiation in K562 cells. Similarly, 54.5% of genes that exhibited high expression in MCF7 exhibited a higher frequency of initiation in K562 and low expression in MCF7 cells. In contrast, in concordantly expressed genes that exhibited similar levels of expression in both cell lines, 51.1% of the genes exhibited a higher frequency of initiation in MCF7.

We next tested whether discordantly expressed genes exhibited TSS replication initiation patterns typical of their expression group. Discordantly expressed genes expressed at high levels in MCF7 cells (and therefore low levels in K562 cells) exhibited TSS exclusion patterns in MCF7 cells that resembled those for all MCF7 highly expressed genes (Fig. 4A). In contrast, those same genes did not exhibit TSS exclusion in K562 cells (Fig. 4B). Similarly, genes that exhibited low expression levels in MCF7 cells (and therefore high K562) did not show marked TSS exclusion in MCF7 cells (Fig. 4C) but did show marked TSS exclusion in K562 cells (Fig. 4D).

Replication initiation and chromatin modifications

With all replication initiation events precisely mapped for these two cell lines, we next asked whether replication initiation events were associated with particular modifications and properties that affect chromatin conformation. Associations between replication initiation frequency and chromatin modifications in K562 cells were analyzed using the UCSC database of epigenetic modifications. As shown in Figure 5, the highest replication initiation ratios were associated with methylated CpG sequences and DNase hypersensitivity. Other features, such as unmethylated CpGs, CTCF binding sites, regions encoding miRNA transcripts, and RNA Polymerase II binding sites were also enriched, albeit at lower levels. Replication initiation events were more frequent in regions that exhibited both methylated CpGs and DNase hypersensitivity (Supplemental Fig. 2A), but this did not represent a striking synergistic effect. CpG methylation in combination with CTCF binding sites did not enhance the initiation ratio, and CTCF binding sites in combination with other chromatin modifications did not enhance the initiation ratio (Supplemental Fig. 2). Consistent with previous reports, higher replication density was associated with binding



Figure 3. Replication initiation depletion at the transcription start site (TSS) in transcribed genes. (*A*) Average replication enrichment ratio in MCF7 cells (calculated as in Fig. 2) plotted against distance from the TSS for all known genes. (*B*) Distribution of replication enrichment ratios in MCF7 cells for groups of genes that exhibit different levels of expression. Levels include the following: very low (log₂ GCRMA <2.3), low (log₂ GCRMA 2.3–5.3), medium (log₂ GCRMA 5.3–8.5), and high (log₂ GCRMA >8.5). (*C*,*D*) The same analyses are shown for K562 cells.

sites for the JUN transcription factor (Cadoret et al. 2008), but this enrichment was minimal. Histone modifications such as methylation of H3K4 or acetylation of H3K9 and H3K27 exhibited minor but statistically significant enrichment in replication initiation.

CpG methylation, replication, and gene expression

To further explore the effect of CpG methylation on replication initiation and its relationship with gene expression, we plotted replication initiation ratios and gene expression values against the frequency of DNA methylation. As shown in Figure 6, the frequency of replication initiation events generally increased with

Discussion

Results from our study suggest that replication initiation is a dynamic, flexible process that is sensitive to both chromatin conformation and transcriptional activity. We found that replication initiation sites were nonrandomly distributed throughout the genome, with clusters of sites in some regions and other regions completely devoid of sites (replication initiation sites were absent in regions up to several hundred kilobases). Replication initiation events were enriched in open chromatin and moderately transcribed regions. Although replication density was highest immediately adjacent to (and downstream from) TSS, the TSS themselves were characterized by low replication density. Methylated CpG tracks were particularly enriched in replication initiation events. Finally, replication initiation events were excluded from tightly condensed chromatin, as well as from heavily transcribed chromatin.

The replication initiation sites identified in our study are in agreement with previous reports, which examined replication initiation events at individual loci (for reviews, see Aladjem et al. 2006; Hamlin et al. 2008). These previous maps were generated using various methodologies that rely on nonoverlapping principles. Overall, we observed replication initiation enrichment within genic regions, which presumably represent open chromatin. The enrichment of replication initiation sites in open chromatin is concordant with the notion that in budding yeast, initiation events occur at an AT-rich consensus (Newlon and Theis 1993; Aladjem and Fanning 2004) or at several near-consensus sequences (Theis and Newlon 2001). Also, in fission yeast, AT-rich sequences are the primary de-

terminants of replicator activity (Dai et al. 2005). Notably, only a subset of AT-rich consensus sequences bind the origin recognition complex and initiate replication, and selected consensus sequences exhibit a unique asymmetric nucleosome positioning pattern (Eaton et al. 2010). In our current study, human replication initiation sites were remarkably enriched immediately downstream from TSSs, colocalizing with nucleosome-free and variant nucleosome regions (Jin et al. 2009). It should be noted that these observations are in agreement with data from the ENCODE project, which has mapped replication initiation sites in $\sim 1\%$ of the human genome (ENCODE Project Consortium 2007; Karnani et al. 2009). Our results are similar to those obtained in a microarray-based study

Definition	Expression value in K562 (log ₂ GCRMA)	Expression value in MCF7 (log ₂ GCRMA)	% Higher replication enrichment ratio in K562 ^a	% Higher replication enrichment ratio in MCF7 ^a
MCF7 H:K562 L	<4.3	>6.3	58.8	40.8
MCF7_L:K562_H	>6.3	<4.3	44.4	54.5
Concordant	4.3< log ₂ GCRMA <6.3; <	4.3 or >6.3 in both	51.1	48.9

Table 1. Replication enrichment values for genes exhibiting discordant and concordant expression

^aPlease note that values do not add to 100% because some genes exhibited equal replication enrichment ratios in both cell lines.

reporting enrichment of replication initiation in transcription factor binding sites (Cadoret et al. 2008); although unlike that study, we have not observed a very high enrichment for JUN binding sites. Our studies are also consistent with other microarray-based studies reporting the enrichment of replication initiation events in regions of decondensed chromatin associated with histone modifications such as H3K4Me3 (Karnani et al. 2009; Valenzuela et al. 2011), and RNA Pol II (Karnani et al. 2009; Valenzuela et al. 2011). Here we have shown that while transcription factor binding sites were somewhat enriched near replication initiation events, transcriptional activity and CpG methylation appeared to have the strongest impact on replication density. The effect of transcriptional activity and CpG methylation was more critical than factors such as DNase I hypersensitivity, transcription factor binding, and unmethylated CpGs.

In our analysis, replication initiation events clustered near transcriptionally active regions, as has been reported previously (Cadoret et al. 2008; Karnani et al. 2009; Valenzuela et al. 2011). We also observed, however, that initiation events were less frequent in regions that exhibited very high levels of transcription. Hence, differences in gene expression may explain the low concordance between two previous microarray-based studies that mapped replication initiation events (Cadoret et al. 2008; Karnani et al. 2009). In addition, while those previous studies



Figure 4. Replication initiation in discordantly expressed genes. We calculated replication enrichment ratios (nascent strands RPKM vs. control genomic DNA RPKM) for genes whose level of expression differed significantly between K562 and MCF7 cell types. Genes with log₂ GCRMA values <4.3 in MCF7 and >6.3 in K562 were considered MCF7-low and K562-high (MCF7_L:K562_H); genes with log₂ GCRMA values <4.3 in K562 and >6.3 in MCF7 were considered MCF7-low and K562-high (MCF7_L:K562_H); genes with log₂ GCRMA values <4.3 in K562 and >6.3 in MCF7 were considered MCF7-high and K562-low (MCF7_H:K562_L). (A) Distribution of nascent strand enrichment ratios in MCF7 cells for MCF7_H:K562_L genes. For comparison, enrichment ratio plots for MCF7_H and MCF7_L genes are shown. (B) Distribution of enrichment ratios in K562 cells for MCF7_H:K562_L genes. For comparison, enrichment ratio plots for K562_H and K562_L genes are shown. (C) Distribution of nascent strand enrichment ratios in MCF7_Cells for MCF7_L:K562_H genes. For comparison, enrichment ratio plots for K562_H and K562_L genes are shown. (C) Distribution of nascent strand enrichment ratios in MCF7_Cells for MCF7_L:K562_H genes. For comparison, enrichment ratio plots for K562_H and K562_L genes are shown. (D) Distribution of nascent strand enrichment ratio plots for MCF7_L:K562_H genes. For comparison, enrichment ratio plots for K562_L genes are shown. (D) Distribution of enrichment ratio plots for MCF7_L:K562_H genes. For comparison, enrichment ratio plots for K562_H and K562_L genes are shown. (D) Distribution of enrichment ratio plots for K562_H genes. For comparison, enrichment ratio plots for K562_H and K562_L genes are shown. (D) Distribution of enrichment ratio plots for K562_H genes. For comparison, enrichment ratio plots for K562_H and K562_L genes are shown.



Chromatin Feature

Figure 5. Chromatin modifications and replication initiation events. The average nascent strand versus genomic DNA ratio (calculated as described for Fig. 2) is shown for genomic regions that contain the indicated chromatin modification features. Modified regions were identified using data from the UCSC Genome Browser (Table 3). Mean NS/control RPKM ratios (dot), and corresponding SDs (error bars), are shown. The horizontal line indicates a matched random control. Chromatin modification features are sorted according to the level of enrichment for replication initiation frequency. Statistically significant deviations from replication ratios of the entire genome (P < 0.001) were observed for all modifications except c-Jun and SIRT6. For intersections between chromatin features (regions that exhibit combinations of chromatin modifications), please see Supplemental Information.

(Cadoret et al. 2008; Karnani et al. 2009; Valenzuela et al. 2011) noted that replication initiation events are enriched in the vicinity of the TSS, our high-resolution data demonstrated that initiation events were excluded from the TSS but enriched immediately downstream from the TSS. Our data also showed that high levels of transcriptional activity were incompatible with the initiation of DNA replication. The depletion of replication initiation events at the location of transcription initiation complexes is similar to the situation in budding yeast, where replication initiates almost exclusively between genes (Raghuraman et al. 2001). These data imply that a high density of transcription initiation complexes may interfere with the formation of replication initiation complexes.

Previous studies have noted a preference for replication initiation in CpG islands (Delgado et al. 1998), and it has been suggested that CpG islands are associated with promoters that can also function as replication origins (Antequera 2004). Analyses of strandspecific mutational asymmetries in long, intergenic regions also suggest that CpG sequences serve as origins for bidirectional replication (Touchon et al. 2004; Polak and Arndt 2009). In murine embryonic stem cells, 0.4% of all promoters contain TSSs that are also associated with replication initiation (Sequeira-Mendes et al. 2009), further suggesting a preference for replication initiation events in CpG-rich sequences. These suggestions are consistent with our analyses using high-resolution sequencing, as we have detected enrichment of replication initiation events immediately downstream from (but not including) the TSS, and this enrichment was particularly associated with CpG methylated genomic regions. Our combined results are consistent with the hypothesis that promoters and replicator sequences have coevolved to ensure the coordination between replication and transcription (Delgado et al. 1998; Antequera and Bird 1999; Sequeira-Mendes et al. 2009), and suggest that frequent transcription initiation events prevent the assembly of pre-replication complexes on chromatin. Importantly, methylated CpG tracks, which we found associated with a high frequency of initiation events, typically mark transcriptionally inactive regions of the genome (Weber et al. 2007). As such, these regions may be inclined to nucleate pre-replication and pre-initiation complexes.

While the present study does not address the issue of replication timing, the relationship among replication initiation sites, replication timing, and transcription is interesting. For example, replication timing is influenced by and correlated with changes in chromatin condensation, as DNase I hypersensitivity is an effective marker for early replicating regions (Hansen et al. 2009). In our current study, the location of initiation events was affected by transcription, whereas others have shown that replication timing correlates poorly with transcription (in studies involving loci with mono-allelic expression). In agreement, a high-resolution study of replication timing revealed that replication timing correlates with gene density in both pluripotent and differentiating human embryonic stem cells and that changes in replication timing sometimes, but not always, reflect gene expression (Desprat et al. 2009). Cellular differentiation influences replication timing over large genomic regions (400-800 kb) in human and mouse cells, and chromatin domains that replicate concomitantly are often located in distinct nuclear compartments (Ryba et al. 2010). Combined with our studies, current observations are consistent with the notion that chromatin accessibility and nuclear compartmentalization primarily affect DNA replication timing, whereas local metabolic processes (such as transcription) primarily affect the location of replication initiation events.

Methods

Cell culture

Cells were obtained from the Developmental Therapeutics Program at the National Cancer Institute. For replication analyses,



Figure 6. Effect of CpG methylation on the frequency of replication initiation events and gene expression. The level of CpG methylation for MCF7 cells (A) and K562 cells (B) is plotted against gene expression levels (log₂ GCRMA, gray histograms) and replication enrichment ratio (nascent strands vs. genomic control, red histograms). For box plots of the data , please see Supplemental Information.

K562 erythroleukemia cells and MCF7 breast cancer cells were grown in Dulbecco's modified Eagle medium supplemented with 10% fetal calf serum and antibiotics, as needed. K562 cells (2.22 \times 10^8) were harvested after 48 h of growth, and 9.36×10^8 MCF7 cells were grown until 85% confluent and then harvested for the

CGCAGGCTTTTTGTAGTGAG

CTGCCAAACATGTGCAGGTA

(Invitrogen). Briefly, nascent strand DNA was incubated for 1 h at 37°C, ethanol precipitated, and resuspended in 30 µL sterile water. Double-stranded nascent DNA (1 µg) was sequenced using the Illumnia (Solexa) genome analyzer II. Sheared genomic DNA was sequenced as a control.

Table 2.	List of	primers	used for	validation

Primer

LCR nonorigin

CTCF nonorigin

Poll 2 nonorigin

Lamin B2

RI TPR

Poll 2

Forward Reverse Probe Previously characterized replication initiation sites GGTGAAGGCTCATGGCAAGA Human beta-globin AAAGGTGCCCTTGAGGTTGTC CCTTTAGTGATGGCCTGGCTCACCTG GGATCCACTTGCCCAGTGTT TCTCAGCAGGGTTCAGGAAGA TCCTTAGTTCCTACCTTCGACCTTGATCCTCCTT TGGGACCCTGCCCTTTTT CGTGACGAAGAGTCAGCT TTCTAGTGAGCCTCCGAC CACACCTGAGGACCCCAATAAC TCACCCTGGATAAGCCTGCGTCGG CCCTGGTCTCTTCTTGTGTATCC Lamin B2 non origin Newly predicted replication initiation sites for validation GGATTGATCCTGGGATTCCT TCCCCACCATTTTAGTCGAG TGCACGGTCTATGGTATTGTG CGGGGTGGAGGAGGTTTC GGACGGTAGAGGGAGACAAA

ACCAGTTCATCCGGACTCAG

ACACACCCAGTGCTTTCCTC

CTCTAGGTGTCACGATGGGG CATCAAGAGAGTCCAGTTCGG CCTCTGAGATGAGTGGGAGC

nascent strand abundance assay, or as described previously (Shankavaram et al. 2007). For gene expression analysis, cell growth and harvesting were performed as described previously (Shankavaram et al. 2007). In brief, cells were thawed and placed in RPMI-1640 (Lonza) with 5% fetal calf serum (Atlantic Biologicals Corporation) and 2 mM glutamine (Invitrogen Corporation). Cells were harvested at \sim 80% confluence, as assessed by phase microscopy for attached cells, or at ${\sim}0.5\,{\times}$ 10⁶ cells/mL for suspended cells. Total RNA was extracted using the Qiagen RNeasy Midi kit.

Nascent strand abundance assay

Genomic and nascent strand DNA was isolated from K562 and MCF7 cells as previously described (Wang et al. 2004), with some modifications.

Nascent DNA was loaded onto a neutral 5%-30% sucrose gradient in TNE. Gradients were centrifuged for 20 h at 21,000 rpm in an SW40 rotor at 25°C. Fractions containing 400-600 bp DNA fragments were pooled, dialyzed against TE, and precipitated. The sample was divided into two aliquots, and plasmid DNA was added to one aliquot as a carrier or positive control. The sample was extracted with phenol-chloroform, ethanol precipitated, and resuspended in 22 µL DEPCtreated water. DNA was digested overnight at 37°C in 32 µL lambda exonuclease buffer and 2 µL lambda exonuclease. The sample was heated for 10 min at 95°C, ethanol precipitated, and resuspended in 50 µL sterile water. RT-PCR was carried out on an aliquot of the sample, using primers for known replication initiation sites. Nascent strands were random-primed using the DNA polymerase I Klenow fragment and the DNA Prime Labeling System

Table 3. Data sources for histone	and DNA modifications	
Label (Fig. 5; Supplemental Fig. 3)	Tracks	URL
Methyl CpG; Unmethyl CpG (K562 and MCF7)	EncodeHaibMethylRrbsK562Hudsonalpha (Note: split into methylated >90% reads/	http://genome.ucsc.edu/cgi-bin/hgTrackUi?hgt_tSearch=Search&g=wgEncodeHaibMethylRrbs
DNase HS	unmetriylated <10% reads ENCODE UW Digital Dhase I Peaks (EDR 0 5%) - 1 et (in K562 rells)	http://genome.ucsc.edu/cgi-bin/hgTrackUi?hgsid=148574667&c=chrX&g=wgEncodeUwDnaseSeq
CTCF binding	ENCODE Open Chromatin, UT ChIP-seq Peaks	http://genome.ucsc.edu/cgi-bin/hgTrackUi?hgsid=148574667&c=chrX&g=wgEncodeChromatinMap
RNA RNA Pol II binding	ENCODE Den Chromatin, UT ChIP-seq Peaks	http://genome.ucsc.edu/cgi-bin/hgTrackUi?hgsid=148574667&c=chrX&g=wgEncodeChromatinMap
H3K27Ac	(NVA FOI II III NJOZ CEIIJ) ENCODE Histone Mods, Broad ChIP-seq Peaks	http://genome.ucsc.edu/cgi-bin/hgTrackUi?hgsid=148574667&c=chrX&g=wgEncodeChromatinMap
H3K4Me3	(H3K9ac, K362) ENCODE Histone Mods, Broad ChIP-seq Peaks	http://genome.ucsc.edu/cgi-bin/hgTrackUi?hgsid=148574667&c=chrX&g=wgEncodeChromatinMap
MYC	(H3N4me3, N302) ENCODE TFBS, Yale/UCD/Harvard ChIP-seq Peaks	http://genome.ucsc.edu/cgi-bin/hgTrackUi?hgsid=148574667&c=chrX&g=wgEncodeYaleChIPseq
JUN	(inyc in KJood Cells) ENCODE TFBS, Yale/UCD/Harvard ChIP-seq Peaks	http://genome.ucsc.edu/cgi-bin/hgTrackUi?hgsid=148574667&c=chrX&g=wgEncodeYaleChIPseq
SIRT6	(c-Jun in K-SOC Cells) ENCODE TFBS, Yale/UCD/Harvard ChIP-seq Peaks (SIRT6 in K562 cells)	http://genome.ucsc.edu/cgi-bin/hgTrackUi?hgsid=148574667&c=chrX&g=wgEncodeYaleChIPseq
Source: UCSC Human Genome Brows	er March 2006 (NCBI 36/hg18) and Feb 2009 (GRCh	37/hg19) regulation tracks.

Nascent strands and fragmented genomic control strands were sequenced using the Illumina GA II platform. The resultant sequences were aligned to the hg19 (NCBI Build 37) human reference genome using Bowtie (Langmead et al. 2009). Alignments were converted to .bam and .tdf format using SAMtools (Li et al. 2009) and igvtools for visualization in Broad's Integrative Genomics Viewer (http://www.broadinstitute.org/igv) (Robinson et al. 2011).

Reads per kilobase per million aligned reads (RPKM) values were calculated for each sample using 100-base genomic bins (Mortazavi et al. 2008). A Gaussian smoothing algorithm was applied to the bin values. To correct for sequencing biases (Dohm et al. 2008; Harismendy et al. 2009) and copy number variation, an enrichment ratio was defined as the ratio of nascent strand RPKM to control RPKM and was calculated for each 100-base bin.

In addition, a .bed format file of the regions with significant replication initiation activity was prepared from the bin-level enrichment ratios. The threshold of enrichment ratios for significant replication initiation activity was selected using a Metropolis Monte Carlo simulation (Metropolis et al. 1953) to find a ratio that identified peaks with a specified target false-discovery rate (FDR) level. The empirical FDR value (Pepke et al. 2009) of a given enrichment ratio threshold was calculated by inverting test and control samples to find the ratio of false (inverted) to true (normal) peaks at the threshold. Initiation at select locations was validated with RT-PCR using primers listed in Table 2.

Transcript expression profiling and data sources

K562 and MCF7 mRNA samples were labeled and processed at Gene Logic using the 54,674 probe-set Human Genome U133 Plus 2.0 Array microarrays (Affymetrix). Each expression profile was done in triplicate. Signal intensities were determined using robust multi-array average (RMA) (Wu et al. 2003), with GC correction (Wu et al. 2003), which is based on perfect match probe intensities that have been background corrected. Intensities were transformed to logarithm base 2. Expression values were GCRMA normalized and log₂ transformed. NCBI Gene was used to determine the genomic position of genes, including TSS (NCBI Gene [http://www. ncbi.nlm.nih.gov/gene]). Analysis was restricted to genes with at least one transcript sequence record in RefSeq or GenBank (20,063 genes). GEO record for transcription data is GSE5720.

Chromatin modification data were obtained from the UCSC Genome Browser (Rhead et al. 2010). In cases where features were not available in hg19 coordinates, their coordinates were converted to those in hg19 by extracting feature sequences from the original build and realigning them to hg19. Data sources are listed in Table 3. For a more detailed description of the analysis process, please see the Supplemental Data Analysis.

Data access

Replication initiation data reported in this article have been submitted to the NCBI Gene Expression Omnibus (GEO) (http://www. ncbi.nlm.nih.gov/geo/) under accession no. GSE28911.

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