Large-Scale Chromatin Structure–Function
Relationships during the Cell Cycle and Development:
Insights from Replication Timing

VISHNU DILEEP,1 JUAN CARLOS RIVERA-MULIA,1 JIAO SIMA,1 AND DAVID M. GILBERT1,2
1Department of Biological Science, Florida State University, Tallahassee, Florida 32306-4295
2Center for Genomics and Personalized Medicine, Florida State University, Tallahassee, Florida 32306-4295
Correspondence: gilbert@bio.fsu.edu

Chromosome architecture has received a lot of attention since the recent development of genome-scale methods to measure chromatin interactions (Hi-C), enabling the first sequence-based models of chromosome tertiary structure. A view has emerged of chromosomes as a string of structural units (topologically associating domains; TADs) whose boundaries persist through the cell cycle and development. TADs with similar chromatin states tend to aggregate, forming spatially segregated chromatin compartments. However, high-resolution Hi-C has revealed substructure within TADs (subTADs) that poses a challenge for models that attribute significance to structural units at any given scale. More than 20 years ago, the DNA replication field independently identified stable structural (and functional) units of chromosomes (replication foci) as well as spatially segregated chromatin compartments (early and late foci), but lacked the means to link these units to genomic map units. Genome-wide studies of replication timing (RT) have now merged these two disciplines by identifying individual units of replication regulation (replication domains; RDs) that correspond to TADs and are arranged in 3D to form spatiotemporally segregated subnuclear compartments. Furthermore, classifying RDs/TADs by their constitutive versus developmentally regulated RT has revealed distinct classes of chromatin organization, providing unexpected insight into the relationship between large-scale chromosome structure and function.

Shortly after J.H. Taylor synthesized tritiated thymidine and his pre–Messelson and Stahl demonstration of semi-conservative, antiparallel DNA replication (Taylor et al. 1957; Meselson and Stahl 1958), he discovered that plant and animal chromosomes replicate segmentally in a defined temporal sequence (Taylor 1958, 1960). The segments labeled during a brief pulse were too large to be accounted for by a single replicon, rather being more consistent with the coordinated initiation of replicon clusters within each segment. Later, visualization of purified DNA fibers confirmed the synchronous firing of origin clusters (Jackson and Pombo 1998; Berezney et al. 2000). Today, it is understood that these initiation sites are stochastically selected from a large set of potential initiation sites and each chromosomal domain uses a different cohort of sites (Lebofsky et al. 2006; Hamlin et al. 2008; Cayrou et al. 2011; Letessier et al. 2011; Besnard et al. 2012). Despite this flexibility in origin choice, a defined temporal order of replication (replication timing; RT) emerges (Bechhoefer and Rhind 2012). Indeed, it is the temporal order of replication, not the initiation sites, that is conserved among species (Aladjem et al. 2002; Farkash-Amar et al. 2008; Liachko et al. 2010; Ryba et al. 2010; Yaffe et al. 2010; Di Rienzi et al. 2012; Müller and Nieduszynski 2012; Xu et al. 2012).

Blocks of synchronously firing origins can also be visualized by genome-wide RT profiling as large synchronously replicated chromosome segments, and their boundaries can be identified by monitoring the coordinated switches in RT that occur across half of the genome during cell fate transitions, revealing 400–800 kb units of replication regulation, or “replication domains (RDs)” (Hiratani et al. 2008, 2010; Hansen et al. 2010; Ryba et al. 2010). RDs are also likely to be the equivalent of the punctate “replication foci” that are seen whenever cells are briefly labeled with detectable nucleotide analogs. These foci persist through multiple cell generations, suggesting they are stable chromosome units, and foci that replicate early or late during S phase reside in separate compartments of the nucleus and correlate with transcriptional activity within each domain (Nakamura et al. 1986; Nakayasu and Berezney 1989; O’Keefe et al. 1992; Sparvoli et al. 1994; Ferreira and Carmo-Fonseca 1997; Jackson and Pombo 1998; Ma et al. 1998; Dimitrova and Gilbert 1999; Sadoni et al. 2004). Recent findings have mapped this 3D organization onto the genome sequence, providing molecular alignment of large-scale chromatin folding with RDs and their time of firing during S phase, both enhancing our understanding of the significance and mechanism of the replication program and providing a functional proxy for the molecular maps of chromosome folding (Ryba et al. 2010; Yaffe et al. 2010; Dixon et al. 2012; Moindrot et al. 2012). These results suggest a unifying model of how structural compartmentalization of the genome in the eukaryotic nucleus can influence overall functional output. Here, we summarize the novel insights that genome-wide analyses of RT have contributed to our understanding of large-scale chromosome structure and function.
Measuring RT Genome-Wide

The most robust (rapid/high-dynamic-range) method for genome-wide RT profiling is to pulse-label cells with a nucleoside analog such as 5-bromo-2-deoxyuridine (BrdU) and purify early and late S-phase fractions on the basis of DNA content by flow cytometry (Gilbert 2010). BrdU-labeled DNA from each fraction is immunoprecipitated, amplified, and sequenced or hybridized to a microarray, and a ratio of enrichment of sequences in early fraction to late fraction is generated (E/L method) (Fig. 1A; Ryba et al. 2011a; Dileep et al. 2012). For nonlive cell samples (e.g., proliferating cells that were frozen or fixed), RT can be inferred by measuring relative copy number variation (CNV) between cells sorted in S phase versus G1 phase (Woodfine et al. 2004; Gilbert 2010). This circumvents the nucleoside analog incorporation and immunoprecipitation steps, but the maximum dynamic range of the data is decreased from 1000-fold or higher for the E/L method to twofold. It is even possible to glean RT from the slight CNV within a genomic sequence, providing that sufficient numbers of cells were in S phase and the sequencing is deep enough to overcome noise (Manukjan et al. 2013; Dimitriadou et al. 2014; Koren et al. 2014).

Interpreting RT Profiles (CTRs and TTRs)

RT profiles from multicellular organisms consist of broad, computationally defined constant timing regions (CTRs) of relatively synchronous initiation of replication, separated by timing transition regions (TTRs) (Fig. 1B; Farkash-Amar et al. 2008; Hiritani et al. 2008; Desprat et al. 2009; Ryba et al. 2010). CTRs usually correspond to several tandem RDs that replicate at the same time; their boundaries cannot be detected because forks emanating from each RD fuse quickly. Often however, other cell types will replicate them at sufficiently different times to create a TTR and by comparing RT profiles from many cell types, increasingly more RD boundaries can be identified (Pope et al. 2014). Hence, RDs and their boundaries are cell type–invariant, whereas CTRs reflect the developmental regulation of RT. Similarly, replication within TTRs can proceed passively through one or more RDs before fusing with a late replicon or can terminate within an RD by fusing with a fork emanating from a distal early RD. As a result, only the earliest replicating border of a TTR aligns with the boundary of an RD (Pope et al. 2014). It is still a matter of debate as to whether TTRs consist of a single unidirectional fork (Norie et al. 2005; Schultz et al. 2010) or sequential firing of origins resulting in the appearance of unidirectional replication (Guilbaud et al. 2011). However, sequences that can function as origins at their normal locations initiate poorly when inserted into a TTR (Guan et al. 2009), whereas conditions of slowed fork movement (replication stress) may activate dormant origins (Blow and Ge 2009) within the TTR to accelerate its replication. Hence, both interpretations are likely to be correct under the appropriate conditions.

Common Misinterpretations

Can RT profiles identify replication origins? Closer inspection of CTRs reveals a series of “earliest replicating

RT profiles from multicellular organisms consist of broad, computationally defined constant timing regions (CTRs) of relatively synchronous initiation of replication, separated by timing transition regions (TTRs) (Fig. 1B; Farkash-Amar et al. 2008; Hiritani et al. 2008; Desprat et al. 2009; Ryba et al. 2010). CTRs usually correspond to several tandem RDs that replicate at the same time; their boundaries cannot be detected because forks emanating from each RD fuse quickly. Often however, other cell types will replicate them at sufficiently different times to create a TTR and by comparing RT profiles from many cell types, increasingly more RD boundaries can be identified (Pope et al. 2014). Hence, RDs and their boundaries are cell type–invariant, whereas CTRs reflect the developmental regulation of RT. Similarly, replication within TTRs can proceed passively through one or more RDs before fusing with a late replicon or can terminate within an RD by fusing with a fork emanating from a distal early RD. As a result, only the earliest replicating border of a TTR aligns with the boundary of an RD (Pope et al. 2014). It is still a matter of debate as to whether TTRs consist of a single unidirectional fork (Norie et al. 2005; Schultz et al. 2010) or sequential firing of origins resulting in the appearance of unidirectional replication (Guilbaud et al. 2011). However, sequences that can function as origins at their normal locations initiate poorly when inserted into a TTR (Guan et al. 2009), whereas conditions of slowed fork movement (replication stress) may activate dormant origins (Blow and Ge 2009) within the TTR to accelerate its replication. Hence, both interpretations are likely to be correct under the appropriate conditions.
peaks,” descriptively referred to as “ripples” (Mukhopadhyay et al. 2014; Bartholdy et al. 2015), that are often referred to as active replication origins (Hause and Shen-dure 2014; Koren et al. 2014). However, the statistical reproducibility of ripples across several biological replicates must first be rigorously determined and even then, ripples likely result from heterogeneous firing of several adjacent low-efficiency origins in the population (Rhind and Gilbert 2013). Even the sharpest and most reproducible ripples fall in the range of several tens of kilobases, sufficient to harbor multiple potential replication origins (Rhind and Gilbert 2013). Importantly, no in-depth study of these features has been performed to confirm their identity from their behavior in RT profiles. Hence, although some ripples likely represent regions of frequent initiation, it is misleading to refer to these initiation regions as “replication origins.”

Resolution. Another common misinterpretation is that deeper sequencing and/or more extensive fractionation of the cell cycle will give increasingly higher resolution data. It is important to understand that the resolution of ensemble RT data is limited by the biology of DNA replication. Because replication forks move bidirectionally at nearly 2 kb/min at an average spacing of 120 kb (Cayrou et al. 2011), most replicons within any 400–800-kb RD will complete synthesis in 30–40 min (<10% of S phase). Perhaps more importantly, stochastic origin firing produces a different pattern of replication in each cell, so the averaging of thousands of cells further diminishes the

Figure 1. Measuring replication timing (RT) and its relationship to chromatin structure. (A) Repli-seq method. Cells pulse labeled with BrdU are separated into early and late S-phase fractions by flow cytometry. BrdU-substituted DNA from each fraction is immunoprecipitated with an anti-BrdU antibody and subject to next-generation sequencing. (B, Top) Exemplary RT profile of 50 Mb from hChr.10 in IMR90 fibroblasts, compared with Hi-C interaction compartments (Eigenvector display) and LaminB1 contact maps (DamID). (Bottom) 10 Mb containing large adjacent early and late synchronously replicating regions is expanded and compared with topologically associating domains (TADs) and Hi-C compartments. At the bottom is a schematic model; larger coordinatey replicated regions consist of multiple tandem early (green) or late (red) replicating domains that are off or on the periphery, respectively. Blue vertical lines are replication domain (RD) boundaries identified in IMR90 and gray dotted lines are RD boundaries found in other cell types. FACS, fluorescence-activated cell sorting.
signal from any given origin. Single-cell or single-molecule methods (Norio and Schildkraut 2001; Gilbert 2010; Schultz et al. 2010; Van der Aa et al. 2013) are needed to overcome these barriers.

**Neither RDs nor CTRs Align with Chromomeric Banding Patterns.** When labeled cells are chased into metaphase, a close correlation is consistently observed between banding patterns revealed by replication labeling and chromomeric R/G chromosome bands (Latt 1977; Holmquist et al. 1982; Craig and Bickmore 1993; Hoshi and Ushiki 2011). However, neither CTRs nor the much smaller RDs align to R/G bands (Fig. 1B). Presumably, this must reflect higher-order packaging of early and late replicating DNA into the metaphase chromosomes, an interesting phenomenon that deserves further study.

**RT IS A STABLE EPIGENETIC PROPERTY**

RT is a robust property resistant to most artificial perturbations of chromatin structure and transcriptional programs. Except a few localized changes, the RT program is virtually unaffected by knockout/knockdown or overexpression of many chromatin and transcription regulators, aging, and induced cellular senescence, and conditions of replication stress where origin choice is demonstrably altered population-wide (Hiratani et al. 2008; Yokochi et al. 2009; Pope et al. 2011; Ryba et al. 2012; Takebayashi et al. 2013; JC Rivera-Mulia, unpubl.). In fact, the only molecular perturbations that have been shown to cause widespread changes in RT were Rif1 (16% of genome affected) and Pol Theta (Cornacchia et al. 2012; Hayano et al. 2012; Fernandez-Vidal et al. 2014). Table 1 summarizes and ranks factors according to their effect on RT.

In contrast to the resilience of RT to artificial manipulations, cell fate changes reproducibly and extensively modify RT. RT profiles of different genetically polymorphic embryonic stem cell (ESC) lines were nearly identical. However, after differentiation of ESCs toward distinct lineages, a global reorganization of RT occurred that affected ∼50% of the genome and 20% of the genome between any given cell fate transition (Hiratani et al. 2008, 2010; Ryba et al. 2010; Rivera-Mulia et al. 2015). In addition, RT changes resulted in a significant increase in the average CTR size together with a drop in the number of domains, with small differentially replicated RDs consolidating into larger regions of tandem RDs that replicate at similar times during S phase (Hiratani et al. 2008; Ryba et al. 2010; Rivera-Mulia et al. 2015). In summary, RT is mitotically stable and a cell type–specific epigenetic property. The stability of RT has enabled us to generate unique RT “signatures” and “fingerprints” that can distinguish cell types (Ryba et al. 2011b; Rivera-Mulia et al. 2015) and even distinguish subtypes of cancers (Ryba et al. 2012).

**RT IS REGULATED AT THE LEVEL OF DEFINED CHROMOSOMAL UNITS**

Cytogenetic studies have long shown that DNA synthesis can be visualized as discrete foci that complete replication within 45–60 min and are stable structural units of 0.5–1 Mb (Nakamura et al. 1986; Nakayasu and Berezney 1989; O’Keefe et al. 1992; Jackson and Pombo 1998; Ma et al. 1998; Dimitrova and Gilbert 1999; Sadoni et al. 2004; Maya-Mendoza et al. 2010). Because DNA fiber studies show coordinate firing of multiple origins (“replicon clusters”) across hundreds of kilobases (Berezney et al. 2000), replication foci likely correspond to replicon clusters. Further, differentially labeling early and late replicating DNA showed that early and late foci reside in distinct spatially separated compartments of the nucleus (Sparvoli et al. 1994; Ferreira and Carmo-Fonseca 1997; Jackson and Pombo 1998).

More recently, RT comparison of distinct cell types showed that developmentally regulated RDs were uniform in size, on the order of 400–800 kb, defining a minimal unit of RT control (Hiratani et al. 2008, 2010; Rivera-Mulia et al. 2015), consistent with the size of replication foci. The much larger CTRs seen in RT profiles are also consistent with the finding that groups of adjacent replication foci replicate sequentially within 1–2 h of each other (Sporbert et al. 2002; Sadoni et al. 2004), which could explain many of the submegabase peaks and valleys (ripples) observed within CTRs.

**Table 1.** List of experimental perturbations or conditions that affect replication timing (RT), arranged according to the percentage of genome affected, from least affected (top) to most affected (bottom).

| Conditions | Reference |
|------------|-----------|
| Mother–daughter pairs | Ryba et al. 2012 |
| Individual variation | Pope et al. 2011, Ryba et al. 2012 |
| FSHD versus normal myoblasts | Pope et al. 2011 |
| Oncogene-induced presenescence | Unpublished |
| Suz12 null | Pope et al. 2014 |
| McCP2 null | Unpublished |
| G9a null | Yokochi et al. 2009 |
| H1c, H1d, H1e triple null | Unpublished |
| c-Myc, n-Myc double null | Unpublished |
| BAF53a null | Takebayashi et al. 2013 |
| CAPD3 knockdown | Unpublished |
| LaminB knockdown | Unpublished |
| Rb knockdown | Unpublished |
| HMGA1 knockdown | Unpublished |
| MyoD overexpress | Unpublished |
| Pax5 overexpress | Unpublished |
| CTCF null | Unpublished |
| Replicative presenescence/ Hayflick limit | Unpublished |
| Pluripotent cell growth conditions | Unpublished |
| Age | Unpublished |
| Progeria | Unpublished |
| BAF250a or Brg1 null | Takebayashi et al. 2013 |

FSHD, facioscapulohumeral dystrophy; CTCF, CCCTC-binding factor.
The discovery of genomic units of RT regulation lead us to propose a large-scale “beads on a string” model for the spatiotemporal regulation of DNA replication, which we refer to as the “RD model” (Hiratani et al. 2008). It is now important to test the equivalence of these units to those measured by different methods. First, a proper computational analysis of ripples is needed to decipher their reproducibility and to distinguish regions of high initiation potential within RDs. Second, advances in DNA fiber analyses could elucidate the distribution of origins and forks across the genome to reveal whether domains with definable boundaries indeed exist. Finally, combined replication labeling with fluorescence in situ hybridization (FISH) could link replication foci to chromosome map units directly.

**TADs EQUAL RDs: LINKING STRUCTURAL UNITS TO FUNCTIONAL UNITS**

A major breakthrough came with the development of methods to map chromatin 3D interactions genome-wide (Hi-C). The first Hi-C study (Lieberman-Aiden et al. 2009) reported that chromatin is organized into two spatially distinct, mutually exclusive compartments with boundaries separating interactions between alternate compartments. Surprisingly, we found a strong correlation between early and late RT to the A/B compartments. Surprisingly, we found a strong correlation between early and late RT to the A/B compartments, better than any other chromosomal property analyzed to date (Fig. 1B; Ryba et al. 2010). In fact, even the subtle variations in RT (ripples) match subtle variations in contact frequencies. Higher resolution Hi-C produced even higher correlations to RT and identified the existence of substructures in the interaction maps that were termed topologically associating domains (TADs) (Dixon et al. 2012; Nora et al. 2012; Rao et al. 2014). This raised the hypothesis that TADs might be RDs. By mapping RD boundaries in several different cell types, we found that indeed there is a “near one to one” correspondence of TAD boundaries and the early border of the TTR. Further analysis demarcated RD boundaries at the early replicating border of TTRs as a major point of chromatin state bifurcation (Pope et al. 2014).

Thus RT is regulated at the level of TADs; multiple TADs replicate at the same time to form CTRs and chromatin folds in such a way that CTRs that reside in the same interaction compartment replicate at similar times. Consistently, changes in subnuclear position correlate with changes in RT during development (Williams et al. 2006; Hiratani et al. 2008, 2010). Further, developmental changes in the RT of a domain are accompanied by the reestablishment of chromatin contacts between the domain and other regions of similar RT (Takebayashi et al. 2012; Denholtz et al. 2013). Consistently, the changes observed by artificial perturbations of chromatin structure—for example, RT alterations in Rif1 or BAF250a mutants and in leukemia (Table 1)—align with TAD/RD boundaries (Ryba et al. 2012; V Dileep, unpubl.). These observations suggest that cis-acting elements at the early replicating border of TTRs may act as boundaries to confine units of RT regulation. More direct evidence for the insulating effect of TTRs comes from the analysis of RT on a rearranged human chromosome in mouse cells, which found that when two regions that replicate at different times were juxtaposed, early replication spread into the late domains, but the spread was limited to the closest TTR (Pope et al. 2012).

**FORMATION OF TADs AND NUCLEAR COMPARTMENTS COINCIDE WITH ESTABLISHMENT OF RT**

Another compelling link between RT and the spatial compartmentalization of chromatin was that RT is established coincident with the global repositioning and anchorage of chromatin as the nucleus is reassembled during early G1 phase (Dimitrova and Gilbert 1999), at a point we termed the TDP (timing decision point). The observation that the interphase organization of the genome is disrupted during mitosis and reestablished in G1 phase (Naumova et al. 2013) suggested that reestablishment of TADs and inter-TAD contacts might coincide with the TDP. To test this hypothesis, we used multiplexed 4C-seq to study dynamic changes in chromatin organization during early G1. Both the establishment of TADs and their compartmentalization occurred within the same time frame as the TDP (Fig. 2; Dileep et al. 2015).

The concurrent reestablishment of both RT and chromatin organization at TDP suggests a model in which RT could be the functional consequence of interphase chromatin organization during early G1. In contrast, G2 phase is a period where RT determinants are lost but the interphase spatial organization of chromatin remains unchanged (Lu et al. 2010). In fact, 4C-seq in G2 phase revealed that the TAD structure and the inter-TAD contacts that establish the nuclear compartments are still preserved in G2 (Dileep et al. 2015). Hence, although chromatin contacts may be necessary to establish RT, those chromatin contacts are not sufficient to dictate the RT program in the G2 nuclei. Together, these results support a model where the formation of TADs and their segregation into distinct nuclear compartments serves as a scaffold that seeds the assembly of RDs that are enriched or depleted for factors that modulate the efficiency of origin firing such as Rif1 (Gilbert 2001; Dileep et al. 2015). The loss of timing program in G2 may be due to the dilution or relocation of these chromatin components during replication.

**RDs THAT SWITCH RT ARE DIFFERENT FROM CONSTITUTIVE RDs**

The genome consists of domains that replicate either early or late in all cell types (constitutive domains) or domains that can switch RT from early to late or late to early during the course of differentiation (developmental domains). Generally, more open and transcriptionally active early replicating chromatin is in the nuclear interior and closed transcriptionally silent late replicating chromatin is located closer to the nuclear periphery (Rhind
and Gilbert 2013). A surprise, however, was that many of these correlations that have dominated the literature for decades are applicable to constitutive but not developmental domains. Although constitutive domains are either strongly compartmentalized to the nuclear interior (constitutive early domains) or to the nuclear periphery (constitutive late domains), developmental regions are more plastic in their compartmentalization, regardless of early or late RT. A systematic analysis of several histone modifications and transcription factors showed that enrichment of these features was correlated to RT for constitutive domains, whereas developmental domains showed moderate enrichment that was less correlated to RT (Dileep et al. 2015). This finding is consistent with several other studies showing that developmental domains are distinct as measured by several global features of the chromatin as summarized in Table 2 (Besnard et al. 2012; Takebayashi et al. 2012; Lubelsky et al. 2014; Rivera-Mulia et al. 2015).

**RT AND GENE EXPRESSION: A COMPLEX RELATIONSHIP**

Despite the strong correlation between RT program and gene expression observed for decades (Goldman et al. 1984; Hatton et al. 1988; Gilbert 2002), mechanistic and causal relationships remain elusive. Early observations found that regions of the genome that are heterochromatic (Lima-de-Faria 1959; Taylor 1960; Baer 1965; Himes 1967) and/or less transcriptionally active (Hsu et al. 1964; Berlowitz 1965) replicate late during S phase. In contrast, DNA replicated early during S phase contains most active genes (Kleckewie and Stubblefield 1967; Pfeiffer and Tolmach 1968). Studies of individual gene RT showed that constitutively active genes replicate early, whereas tissue-specific genes replicate early only in cell types where they were active (Goldman et al. 1984; Gilbert 1986; Holmquist 1987; Selig et al. 1992; Ermakova et al. 1999; Simon et al. 2001). Further characterization identified developmentally regulated RT switches that encompass hundreds of kilobases that were coincident with transcriptional activation of genes within the affected domain (Hiratani et al. 2004, 2008; Perry et al. 2004).

With the advent of genome-wide studies, the global correlation between early replication and active gene expression has been confirmed in all multicellular organisms studied (Schübeler et al. 2002; MacAlpine et al. 2004; Woodfine et al. 2004; Huvet et al. 2007; Hiratani et al. 2008; Desprat et al. 2009; Hiratani and Gilbert 2009; Schweiger et al. 2009; Maric and Prioleau 2010; Lubelsky et al. 2014). Moreover, development changes in the RT program are closely coordinated with transcriptional competence (Zhou et al. 2002; Hiratani et al. 2008, 2010; Desprat et al. 2009; Schultz et al. 2010; Yue et al. 2014). However, the dynamic changes in RT and gene expression were so coincident that the order of events was not determined and causal relationships remained unsolved. In addition, closer inspection of the genome-wide studies showed that the relationship between transcription and RT is clearly indirect, because changes in gene expression can occur without alterations in RT and ~20% of

---

**Table 2.** Comparison of chromatin features between constitutive early and late (CE, CL) and developmental replication domains that are either early (DE) or late (DL) in the particular cell type analyzed for each respective chromatin feature

| Property                              | CE                   | CL       | DE                   | DL                   |
|---------------------------------------|----------------------|----------|----------------------|----------------------|
| Degree of compartmentalization       | High                 | High     | Low                  | Low                  |
| MNase sensitivity, origin density     | High                 | Low      | Low                  | Low                  |
| LaminB association                    | Low                  | High     | Moderately low       | Moderately high      |
| 42 trans-acting factors (e.g., CTCF, EZH2), chromatin marks: H3K27ac, H3K4me1, H3K4me2, H3K9ac, H4K20me1, H2az | Enriched             | Depleted | Moderately enriched  | Moderately depleted |
late replicating genes are expressed (Schübeler et al. 2002, 2004; Hiratani et al. 2008, 2010).

To investigate the intriguing liaisons between RT and gene expression, we recently analyzed their coordinated changes during cell fate commitment (Rivera-Mulia et al. 2015). By taking advantage of optimized methods for human embryonic stem cell differentiation (Schulz et al. 2012; Menendez et al. 2013; Rivera-Mulia et al. 2015) and analyzing multiple cell types and differentiation intermediates, we characterized the establishment of cell type-specific RT programs and differential gene expression. We observed highly coordinated kinetics of RT and transcriptional changes; however, our study also revealed that the strong correlation between early replication and active transcription is restricted to RT constitutive genes (i.e., genes that do not change their RT program), whereas the RT switching genes have a weaker correlation that is lost during differentiation. Deeper analysis showed distinct classes of genes, among those, some are expressed only when they replicate early (E-class), whereas others can be expressed when they replicate late (C-class) (Fig. 3A,B). Surprisingly, the majority of RT switching genes belong to the C-class category and are strongly expressed in one or more cell types while late replicating, demonstrating

Figure 3. Cell cycle and developmental regulation of replication domains (RDs): The RD model. (A) RDs correspond to topologically associating domains (TADs). Replication timing (RT) profiles of two hypothetical cell types illustrate an RD that changes RT (developmental RD) and RDs that replicate either early or late in all cell types (constitutive RD). Genes in constitutive early/late RDs are colored black. E-class and C-class genes are shown in green and blue, respectively. (B) Early replicating RDs are more open and positioned toward the interior and late replicating RDs are closed and positioned near nuclear periphery. Late RDs and their adjacent timing transition regions (TTRs) associate with the nuclear lamina to form lamina-associated domains (LADs). During differentiation, developmental RDs move toward or away from nuclear lamina depending on the direction of the RT switch. Expression of E-class genes is closely linked to RT and is expressed coordinately with their switch toward early replication, whereas C-class genes are less correlated to RT and are generally turned on before their switch to early replication. (C) Formation of TADs and their segregation into early/late compartments both coincide with establishment of RT during TDP.
that transcription is not sufficient for early replication. Additionally, a rare category of genes (L-class) contains RT switching genes that were expressed exclusively when replicated late. Interestingly, transcriptional activity of C-class genes is still coordinated with RT, despite their ability to be transcribed when replicated late. However, an increase in expression levels is commonly observed before changes to early replication, whereas changes to late RT are followed by down-regulation (Rivera-Mulia et al. 2015). These results suggest either that aspects of transcriptional control can indirectly influence RT or that different RDs are regulated by different mechanisms. Manipulative experiments are now imperative to dissect the mechanisms linking transcription and RT.

**CONCLUSION: THE RD MODEL**

Here, we finish with a unifying large-scale structure–function model that incorporates recent findings linking RT to chromatin 3D structure (Fig. 3). Our model has two main tenets. First, RT is regulated at the level of TADs that are stable across cell types (Pope et al. 2014; Dixon et al. 2015). Second, TADs that replicate at similar times frequently interact with each other, dividing the genome into two distinct spatially separated nuclear compartments. Consistently, early replication is associated with the nuclear interior, whereas late replication is observed near the nuclear periphery (Rhind and Gilbert 2013). Late replicating RDs together with TTRs associate with the nuclear lamina to form lamina-associated domains (Fig. 3B; Peric-Hupkes et al. 2010; Pope et al. 2014). Thus, the segregation of RDs/TADs into these early or late replicating compartments reflects each cell type’s unique RT program. This organization persists throughout interphase but is dismantled during mitosis (Naumova et al. 2013) and reestablished coincident with the establishment of RT early in G1 (Fig. 3C; Dileep et al. 2015). This emerging view of the genome recognizes RDs/TADs as epigenetic modules whose chromatin features are regulated during cell fate transitions. These features include RT, chromatin marks, chromatin accessibility, and 3D organization (Rhind and Gilbert 2013), which all affect gene regulation. Because disruption of TADs and misregulation of its properties could result in disease states (Ryba et al. 2012; Lupiñáez et al. 2015), a critical issue to be resolved now is the extent to which TAD structure is a cause or consequence of genome function. We predict that studies of RT using modern “domain engineering” methods will provide important insight into this question.

**ACKNOWLEDGMENTS**

Research in the Gilbert laboratory is supported by National Institutes of Health (NIH) grants GM083337 and GM085354.

**REFERENCES**

Aladjem MI, Rodewald LW, Lin CM, Bowman S, Cimbora DM, Brody LL, Epner EM, Grouin M, Wahl GM. 2002. Replication initiation patterns in the β-globin loci of totipotent and differentiated murine cells: Evidence for multiple initiation regions. *Mol Cell Biol* 22: 4421–4429.

Baer D. 1965. Asynchronous replication of DNA in a heterochromatic set of chromosomes in *Pseudococcus obscurus*. *Genetics* 52: 275–285.

Bartholdy B, Mukhopadhyay R, Lajugie J, Aladjem MI, Bouhassira EE. 2015. Allele-specific analysis of DNA replication origins in mammalian cells. *Nat Commun* 6: 7051.

Bechterow J, Rhind N. 2012. Replication timing and its emergence from stochastic processes. *Trends Genet* 28: 374–381.

Berezney R, Dubey DD, Huberman JA. 2000. Heterogeneity of eukaryotic replicons, replicon clusters, and replication foci. *Chromosoma* 108: 471–484.

Berlowitz L. 1965. Correlation of genetic activity, heterochromatinization, and RNA metabolism. *Proc Natl Acad Sci* 53: 67–73.

Besnard E, Babled A, Lapasset L, Millahvet O, Parrinello H, Dantec C, Marin JM, Lemaître JM. 2012. Unraveling cell type-specific and reprogrammable human replication origin signatures associated with G-quadruplex consensus motifs. *Nat Struct Mol Biol* 19: 837–844.

Blow JJ, Ge XQ. 2009. A model for DNA replication showing how dormant origins safeguard against replication fork failure. *EMBO Rep* 10: 406–412.

Cayrou C, Coulombe P, Vigneron A, Stanojicic S, Ganior O, Peiffer I, Rivals E, Puy A, Laurent-Chabalier S, Desprat R, et al. 2011. Genome-scale analysis of metazoan replication origins reveals their organization in specific but flexible sites defined by conserved features. *Genome Res* 21: 1438–1449.

Cornacchia D, Dileep V, Quivy JP, Foti R, Tili F, Santarella-Mellwig R, Antony C, Almouzni G, Gilbert DM, Buonomo SB. 2012. Mouse Rif1 is a key regulator of the replication-timing programme in mammalian cells. *EMBO J* 31: 3678–3690.

Craig JM, Bickmore WA. 1993. Chromosome bands—Flavours to savour. *Bioessays* 15: 349–354.

Denholtz M, Bonora G, Cronis C, Splinter E, de Laat W, Ernst J, Pellegrini M, Plath K. 2013. Long-range chromatin contacts in embryonic stem cells reveal a role for pluripotency factors and polycomb proteins in genome organization. *Cell Stem Cell* 13: 602–616.

Desprat R, Thierry-Mieg D, Lailler N, Lajugie J, Schildkraut C, Murray T, Urgaonkar S, Aladjem MI, Boukherroub R, Berezney R. 2015. Allele-specific analysis of DNA replication origins from stochastic processes. *Trends Genet* 31: 983–993.

Dimitrova DS, Gilbert DM. 1999. The spatial position and replication timing of the replication-timing establishment of the replication-timing program. *Mol Cell Biol* 19: 983–993.
Holmquist GP. 1987. Role of replication time in the control of tissue-specific gene expression. *Am J Hum Genet* 40: 151–173.

Holmquist G, Gray M, Porter T, Jordan J. 1982. Characterization of Gimsa dark- and light-band DNA. *Cell* 31: 121–129.

Hoshi O, Ushiki T. 2011. Replication banding patterns in human chromosomes detected using 5-ethyl-2'-deoxyuridine incorporation. *Acta Histochem Cytochem* 44: 233–237.

Hsu TC, Schmid W, Stubblefield E. 1964. DNA replication sequences in higher mammals. In *The role of chromosomes in development* (ed. Locke M), pp. 83–112. Academic Press, New York.

Huvet M, Nicolas S, Touchon M, Audit B, d’Aubenton-Carafa Y, Arneodo A, Thermenes C. 2007. Human gene organization driven by the coordination of replication and transcription. *Genome Res* 17: 1278–1285.

Jackson DA, Pombo A. 1998. Replication clusters are stable units of chromosome structure. Evidence that nuclear organization contributes to the efficient activation and propagation of S phase in human cells. *J Cell Biol* 140: 1285–1295.

Kleckner RR, Stubblefield E. 1967. RNA synthesis in relation to DNA replication in synchronized Chinese hamster cell cultures. *J Exp Zool* 165: 259–268.

Koren A, Handsaker RE, Karczewski KJ, Karlic R, Ghosh S, Polak P, Eggan K, McCarroll SA. 2014. Genetic variation in human DNA replication timing. *Cell* 159: 1015–1026.

Latt SA. 1977. Fluorescent probes of chromosome structure and replication. *Can J Genet Cytol* 19: 603–623.

Lebofsky R, Heilig R, Sonnleitner M, Weissenbach J, Bensimon A. 2006. DNA replication origin interference increases the spacing between initiation events in human cells. *Mol Biol Cell* 17: 5337–5345.

Letessier A, Millot GA, Koundrioukov S, Lachagné AM, Vogt N, Hansen RS, Malfoy B, Brison O, Debatisse M. 2011. Cell-type-specific replication initiation programs set fragility of the FRA3B fragile site. *Nature* 470: 120–123.

Liaichko I, Bhaskar A, Lee C, Chung SC, Tye BK, Keich U. 2010. A comprehensive genome-wide map of autonomously replicating sequences in a naïve genome. *PLoS Genet* 6: e1000946.

Lieberman-Aiden E, van Berkum NL, Williams L, Imakaev M, Ragoczy T, Telling A, Amit I, Lajoie BR, Sabo PJ, Dorschner MO, et al. 2009. Comprehensive mapping of long range interactions reveals folding principles of the human genome. *Science* 326: 289–293.

Lima-de-Faria A. 1959. Differential uptake of tritiated thymidine into hetero- and euchromatin in *Melanopus* and *Seacale*. *J Biophys Biochem Cytol* 5: 457–466.

Lu J, Li F, Murphy CS, Davidson MW, Gilbert DM. 2010. G2 phase chromatin lacks determinants of replication timing. *J Cell Biol* 189: 967–980.

Lubelsky Y, Prinz JA, DeNapoli L, Li Y, Belsky JA, MacAlpine DM. 2014. DNA replication and transcription programs respond to the same chromatin cues. *Genome Res* 24: 1102–1114.

Lupuñiez DG, Kraft K, Heinrich V, Kravitz P, Brancati F, Klopcic E, Horn D, Kayserili H, Opitz JM, Laxova R, et al. 2015. Disruptions of topological chromatin domains cause pathogenic rewiring of gene-enhancer interactions. *Cell* 161: 1012–1025.

Ma H, Samarabandu J, Devdhari RS, Acharya R, Cheng PC, Meng C, Berezney R. 1998. Spatial and temporal dynamics of DNA replication sites in mammalian cells. *J Cell Biol* 143: 1415–1425.

MacAlpine DM, Rodriguez HK, Bell SP. 2004. Coordination of replication and transcription along a *Drosophila* chromosome. *Genes Dev* 18: 3094–3105.

Manukjian G, Tauscher M, Steinemann D. 2013. Replication timing influences DNA copy number determination by array-CGH. *Biotechniques* 55: 231–232.

Maric C, Prioleau MN. 2010. Interplay between DNA replication and gene expression: A harmonious coexistence. *Curr Opin Cell Biol* 22: 277–283.
Maya-Mendoza A, Oliwares-Chauvet P, Shaw A, Jackson DA. 2010. S phase progression in human cells is dictated by the genome-wide distribution of DNA foci. *PLoS Genet* e1000900.

Menendez L, Kulik MJ, Page AT, Park SS, Lauderdale JD, Cunningham ML, Dalton S. 2013. Directed differentiation of human pluripotent cells to neural crest stem cells. *Nat Protoc* 8: 203–212.

Meselson M, Stahl FW. 1958. The replication of DNA in *Escherichia coli*. *Proc. Natl Acad Sci* 44: 671–682.

Monardova A, Axline B, Kleinsmith L, Baker A, Therms C, de Laat W, Bouvet P, Mongelard F, Arneodo A. 2012. 3D chromatin conformation correlates with replication timing and is conserved in resting cells. *Nucleic Acids Res* 40: 9470–9481.

Mukhopadhyay R, Lajugie J, Fourel N, Selzer A, Schizas M, Bartholdy B, Mar J, Lin CM, Martin MM, Ryan M, et al. 2014. Allele-specific genome-wide profiling in human primary erythroblasts reveal replication program organization. *PLoS Genet* 10: e1004319.

Müller CA, Nieduszynski CA. 2012. Conservation of replication timing reveals global and local regulation of replication origin activity. *Genome Res* 22: 1953–1962.

Nakamura H, Morita T, Sato C. 1986. Structural organization of replication domains during DNA synthetic phase in the mammalian cell line *HeLa*. *Exp Cell Res* 165: 177–187.

Nakayasu H, Berezney R. 1989. Mapping replicational sites in *Escherichia coli*. *Cell* 16: 269–274.

Norio P, Schildkraut CL. 2001. Visualization of DNA replication at individual replication foci in *Escherichia coli*. *Nature* 408: 381–385.

Norio P, Schildkraut CL. 2001. Visualization of DNA replication on individual Epstein–Barr virus episomes. *Science* 294: 2361–2364.

Norio P, Kosiarytrakul S, Yang Q, Guan Z, Brown NM, Thomas S, Riblet R, Schildkraut CL. 2005. Progressive activation of DNA replication initiation in large domains of the immunoglobulin heavy chain locus during B cell development. *Mol Cell* 20: 575–587.

O’Keeffe RT, Henderson SC, Spector DL. 1992. Dynamic organization of DNA replication in mammalian cell nuclei: Spatially and temporally defined replication of chromosome-specific α-satellite DNA sequences. *J Cell Biol* 116: 1095–1110.

Peric-Hupkes D, Meuleman W, Bouvet P, Merkel F, Arneodo A. 2012. 3D chromatin conformation correlates with replication timing and is conserved in resting cells. *Nucleic Acids Res* 40: 9470–9481.

Pfeiffer SE, Tolnach LJ. 1968. RNA synthesis in synchronously growing populations of HeLa S3 cells. I. Rate of total RNA synthesis and its relationship to DNA synthesis. *J Cell Physiol* 71: 77–93.

Pope BD, Tsumagari K, Battaglia D, Ryba T, Hiratani I, Ehrlich M, Gilbert DM. 2011. DNA replication timing is maintained genome-wide in primary human myoblasts independent of D4Z4 contraction in FSH muscular dystrophy. *PLoS One* 6: e27413.

Pope BD, Chandra T, Buckley Q, Hoare M, Ryba T, Wiseman FK, Kuta A, Wilson MD, Odom DT, Gilbert DM. 2012. Replication-timing boundaries facilitate cell-type and species-specific regulation of a rearranged human chromosome in mouse. *Hum Mol Genet* 21: 4162–4170.

Pope BD, Ryba T, Dileep V, Yue F, Wu W, Denas O, Vera DL, Wang Y, Hansen RS, Canfield TK, et al. 2014. Topologically associating domains are stable units of replication-timing regulation. *Nature* 515: 402–405.

Rad S, Hunley MH, Philipp E, Stamenova EK, Bockiov KD, Robinson JT, Sanborn AL, Machol I, Omer AD, Lander ES, et al. 2014. A 3D map of the human genome at kilobase resolution reveals principles of chromatin looping. *Cell* 159: 1665–1680.

Rhind N, Gilbert DM. 2013. DNA replication timing. *Cold Spring Harb Perspect Med* 3: 1–26.

Roca-Mula JC, Buckley Q, Sasaki T, Zimmerman J, Didier RA, Nazor K, Loring JF, Lian Z, Weissman S, Robins AJ, et al. 2015. Dynamic changes in replication timing and gene expression during lineage specification of human pluripotent stem cells. *Genome Res* 25: 1091–1103.

Ryba T, Hiratani I, Lu J, Itoh M, Kulik M, Zhang J, Dalton S, Gilbert DM, Schulz TC, Robins AJ, et al. 2010. Evolutionarily conserved replication timing profiles predict long-range chromatin interactions and distinguish closely related cell types. *Genome Res* 20: 761–770.

Ryba T, Battaglia D, Pope BD, Hiratani I, Gilbert DM. 2011a. Genome-scale analysis of replication timing: From bench to bioinformatics. *Nat Protoc* 6: 870–895.

Ryba T, Hiratani I, Sasaki T, Battaglia D, Kulik M, Zhang J, Dalton S, Gilbert DM. 2011b. Replication timing: A fingerprint for cell identity and pluripotency. *PLoS Comput Biol* 7: e1002225.

Ryba T, Battaglia D, Chang BH, Shirley JW, Buckley Q, Pope BD, Devidas M, Druker BJ, Gilbert DM. 2012. Abnormal developmental control of replication-timing domains in pediatric acute lymphoblastic leukemia. *Genome Res* 22: 1833–1844.

Sadoni N, Cardoso MC, Stelzer EH, Leonhardt H, Zink D. 2004. Stable chromosomal units determine the spatial and temporal organization of DNA replication. *J Cell Sci* 117: 5353–5365.

Schübeler D, Scalzo D, Kooperberg C, van Steensel B, Delrow J, Groudine M. 2002. Genome-wide DNA replication profile for *Drosophila melanogaster*: A link between transcription and replication timing. *Nat Genet* 32: 438–442.

Schübeler D, MacAlpine DM, Scalzo D, Wirbelauer C, Kooperberg C, van Leeuwen F, Gottschling DE, O’Neill LP, Turner BM, Delrow J, et al. 2004. The histone modification pattern of active genes revealed through genome-wide chromatin analysis of a higher eukaryote. *Genes Dev* 18: 1263–1271.

Schultz SS, Desbordes SC, Du Z, Kosiarytrakul S, Lipchina I, Studer L, Schildkraut CL. 2010. Single-molecule analysis reveals changes in the DNA replication program for the POU5F1 locus upon human embryonic stem cell differentiation. *Mol Cell Biol* 30: 4521–4534.

Schultz TC, Young HY, Aguilnick AD, Babin MJ, Baetge EE, Bang AG, Bhournik A, Cepa I, Cesario RM, Haakmeester C, Studer L, Schildkraut CL. 2010. Single-molecule analysis reveals changes in the DNA replication program for the POU5F1 locus upon human embryonic stem cell differentiation. *Mol Cell Biol* 30: 4521–4534.

Selig S, Okumura K, Ward DC, Cedar H. 1992. Delineation of DNA replication time zones by fluorescence in situ hybridization. *EMBO J* 11: 1217–1225.

Siminovitch L, Tenen D, Mostoslavsky R, Ficahn E, Lande L, Milot E, Gribnau J, Grosfeld F, Fraser P, Cedar H. 2001. Developmental regulation of DNA replication timing at the human β globin locus. *EMBO J* 20: 6150–6157.

Sparvoli E, Levi M, Rossi K, Zachei SJ, Hanekom EJ, de Ruyter J, Schuëbel D, 2009. Chromatin state marks cell-type- and gender-specific replication of the *Drosophila* genome. *Genes Dev* 23: 589–601.

Sparvoli E, Levi M, Rossi E. 2009. Replicon clusters may form adjacent origin clusters. *PLoS Genet* 5: e1000386.

Sparvoli E, Levi M, Rossi E. 1994. Replicon clusters may form adjacent origin clusters. *PLoS Genet* 5: e1000386.

Sparvoli E, Levi M, Rossi E. 1994. Replicon clusters may form adjacent origin clusters. *PLoS Genet* 5: e1000386.

Sparvoli E, Levi M, Rossi E. 1994. Replicon clusters may form adjacent origin clusters. *PLoS Genet* 5: e1000386.
tally regulated chromosomal domains reveals an unusual principle of chromatin folding. *Proc Natl Acad Sci* **109**: 12574–12579.

Takebayashi S, Lei I, Ryba T, Sasaki T, Dileep V, Battaglia D, Gao X, Fang P, Fan Y, Esteban MA, et al. 2013. Murine esBAF chromatin remodeling complex subunits BAF250a and Brg1 are necessary to maintain and reprogram pluripotency-specific replication timing of select replication domains. *Epigenetics Chromatin* **6**: 42.

Taylor JH. 1958. The mode of chromosome duplication in *Crepis capillaris*. *Exp Cell Res* **15**: 350–357.

Taylor JH. 1960. Asynchronous duplication of chromosomes in cultured cells of Chinese hamster. *J Biophys Biochem Cytol* **7**: 455–463.

Taylor JH, Woods PS, Hughes WL. 1957. The organization and duplication of chromosomes as revealed by autoradiographic studies using tritium-labeled thymidimee. *Proc Natl Acad Sci* **43**: 122–128.

Van der Aa N, Cheng J, Mateiu L, Zamani Esteki M, Kumar P, Dimitriadou E, Vanneste E, Moreau Y, Vermeesch JR, Voet T. 2013. Genome-wide copy number profiling of single cells in S-phase reveals DNA-replication domains. *Nucleic Acids Res* **41**: e66.

Williams RR, Azuara V, Perry P, Sauer S, Dvorkina M, Jørgensen H, Roix J, McQueen P, Misteli T, Merkenschlager M, et al. 2006. Neural induction promotes large-scale chromatin reorganisation of the Mash1 locus. *J Cell Sci* **119**: 132–140.

Woodfine K, Fiegler H, Beare DM, Collins JE, McCann OT, Young BD, Debernardi S, Mott R, Dunham I, Carter NP. 2004. Replication timing of the human genome. *Hum Mol Genet* **13**: 191–202.

Xu J, Yanagisawa Y, Tsankov AM, Hart C, Aoki K, Kommajosyula N, Steinmann KE, Bochicchio J, Russ C, Regev A, et al. 2012. Genome-wide identification and characterization of replication origins by deep sequencing. *Genome Biol* **13**: R27.

Yaffe E, Farkash-Amar S, Polten A, Yakhini Z, Tanay A, Simon I. 2010. Comparative analysis of DNA replication timing reveals conserved large-scale chromosomal architecture. *PLoS Genet* **6**: e1001011.

Yokochi T, Poduch K, Ryba T, Lu J, Hiratani I, Tachibana M, Shinkai Y, Gilbert DM. 2009. G9a selectively represses a class of late-replicating genes at the nuclear periphery. *Proc Natl Acad Sci* **106**: 19363–19368.

Yue F, Cheng Y, Breschi A, Vierstra J, Wu W, Ryba T, Sandstrom R, Ma Z, Davis C, Pope BD, et al. 2014. A comparative encyclopedia of DNA elements in the mouse genome. *Nature* **515**: 355–364.

Zhou J, Ermakova OV, Riblet R, Birshtein BK, Schildkraut CL. 2002. Replication and subnuclear location dynamics of the immunoglobulin heavy-chain locus in B-lineage cells. *Mol Cell Biol* **22**: 4876–4889.