## **Replication Timing**

Replication timing refers to the order in which segments of DNA along the length of a chromosome are duplicated. DNA replication errors increase genetic instability, and may be a causative factor in diseases such as cancer and neuronal disorders. Replication in eukaryotes initiates from discrete genomic regions, termed origins, according to a strict, often tissue-specific, temporal program. The genetic program that controls activation of replication origins in mammalian cells has still not been elucidated.

Recent technology advancement can now allow measurements of replication timing genomewide. One such a technology is developed by Hiratani et al. (2008)(1). Briefly, cells were pulse labeled with BrdU and separated into early and late S-phase fractions by flow cytometry (Figure 1A). BrdU-substituted DNA from each fraction was immunoprecipitated with an anti-BrdU antibody, differentially labeled, and cohybridized to a whole-genome oligonucleotide microarray (Figure 1A). The ratio of the abundance of each probe in the early and late fraction ["replication timing ratio" = log2(Early/Late)] was then used to generate a replication-timing profile for the entire genome (Figure 1B). Biologists are often interested in partitioning the genome according to the obtained replication profiles to early replication domains, late replication domains and timing transition regions (TTR). The circular binary segmentation (CBS) algorithm (2) is often used to segment the genome(1;3;4). Early and late replication domains then are called as segments with log2(Early/Late)<0 and log2(Early/Late)>0, respectively (Figure 1B). The TTRs are called by first loess-smoothing the profile and identification of the regions with large positive and negative slopes(3) (Figure 1C). This method is not satisfactory both statistically and biologically. The replication domains and the transition regions should have no overlap and this cannot be guaranteed by this method. The choice of the slope to determine the TTRs is arbitrary (e.g. Ryba et al. 2010 used +/-68e-7 RT/bp) and the slope estimates could be influenced by the parameters used in loess smoothing. The aim of this project is to develop a new method that can improve the aforementioned method for detecting early/late replication domains and TTRs.

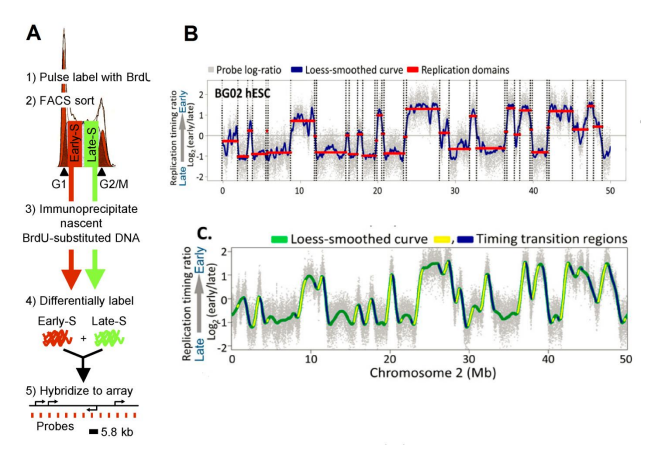


Figure 1 (A) Protocol for genome-wide replication timing analysis using oligonucleotide microarrays. (B) Replication timing profile across a 50-Mb segment of human chromosome 2. Data shown are the average of two replicate hybridizations (dye-swap) for hESC line BG02. DNA synthesized early vs. late during S phase was hybridized to an oligonucleotide microarray, and the log2 ratio of early/late signal for each probe (probe spacing 1.1 kb) across the genome was plotted on the y-axis vs. map position on the x-axis. (Gray dots) Rawdata. (Blue line) Loess-smoothed data. Replication domains (red lines) and

boundaries (dotted lines) were identified by circular binary segmentation (CBS algorithm). (C) Identification of timing transition regions (TTRs; blue and yellow highlight alternating TTRs) from loess-smoothed RT profiles. (Green) BG02 hESC.

## **Project Task**

- 1. Develop a new method for detecting early/late replication domains and TTRs. You need to show that your method works well or even works better than the aforementioned method.
- 2. Apply your method to the data in Ryba et al. 2010 (3) and perform data analysis to show if you can achieve similar conclusions about the replication domains and TTRs that you find. The data in Ryba et al. 2010 (3) can be downloaded from <a href="http://www.replicationdomain.org">http://www.replicationdomain.org</a>. Note: Since Ryba et al. 2010 (3) have a large amount of analyses, you do NOT need to perform every analysis in this paper, but you will be rewarded (specifically, in your final scores) if you perform more. If you have analysis results that do not agree with Ryba et al. 2010(3) and you can logically and convincingly explain why your results are correct, you will be significantly rewarded in your final score.

## Remark about the data

Suppose that you want to download the replication timing data for the human embryonic stem cell line (hESC) BG01. After you open <u>http://www.replicationdomain.org</u> in your browser, click *Database* (shown

as the following figure).

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Note that there are 6 columns in the table shown in the above. The 5<sup>th</sup> column (Data Type) tells you the data type. The rows with a Data Type value *RT* are replication timing data. The 6<sup>th</sup> column (Build) tells you which version of the reference genome the coordinate is based on. **Please use data under hg19**. If you search for *"bg01hESC"*, you can get something like the following.

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Contact Information	Download	BG01hESC_1_UGA_15473002CGH	RD_BG01hES1_Sm300_090127.txt	RT hg17
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hESC Lineage	Download	BG01 2 UGA 15707202CGH	RD BG01hES2 Sm300 090127.txt	RT hg38

You can download data at the row with hg19 by clicking the *Download* text (See the above figure).

## **Reference List**

- 1. Hiratani,I., Ryba,T., Itoh,M., Yokochi,T., Schwaiger,M., Chang,C.W., Lyou,Y., Townes,T.M., Schubeler,D. and Gilbert,D.M. (2008) Global reorganization of replication domains during embryonic stem cell differentiation. *PLoS Biol*, **6**, e245.
- 2. Venkatraman, E.S. and Olshen, A.B. (2007) A faster circular binary segmentation algorithm for the analysis of array CGH data. *Bioinformatics*, **23**, 657-663.
- 3. Ryba,T., Hiratani,I., Lu,J., Itoh,M., Kulik,M., Zhang,J., Schulz,T.C., Robins,A.J., Dalton,S. and Gilbert,D.M. (2010) Evolutionarily conserved replication timing profiles predict long-range chromatin interactions and distinguish closely related cell types. *Genome research*, **20**, 761-770.
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