Spatial and Temporal Dynamics of DNA Replication Sites in Mammalian Cells

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Abstract. Fluorescence microscopic analysis of newly replicated DNA has revealed discrete granular sites of replication (RS). The average size and number of replication sites from early to mid S-phase suggest that each RS contains numerous replicons clustered together. We are using fluorescence laser scanning confocal microscopy in conjunction with multidimensional image analysis to gain more precise information about RS and their spatial-temporal dynamics. Using a newly improved imaging segmentation program, we report an average of ~1,100 RS after a 5-min pulse labeling of 3T3 mouse fibroblast cells in early S-phase. Pulse-chase-pulse double labeling experiments reveal that RS take ~45 min to complete replication. Appropriate calculations suggest that each RS contains an average of 1 mbp of DNA or ~6 average-sized replicons. Double pulse–double chase experiments demonstrate that the DNA sequences replicated at individual RS are precisely maintained temporally and spatially as the cell progresses through the cell cycle and into subsequent generations. By labeling replicated DNA at the G1/S borders for two consecutive cell generations, we show that the DNA synthesized at early S-phase is replicated at the same time and sites in the next round of replication.

Key words: replication sites • replication timing • cell nucleus • chromosomes • computer image segmentation

In pioneering autoradiographic studies, it was determined that regions of euchromatin in the mammalian cell nucleus replicated early in S-phase and regions of heterochromatin replicate later in S-phase (Hay and Revel, 1966; Milner, 1969; Williams and Ockey, 1970; Ockey, 1972; Huberman et al., 1973; Comings and Okada, 1973; Fakan and Hancock, 1974; Sparvoli et al., 1976; Smith et al., 1984). Immunofluorescence microscopy of incorporated biotin labeled dUTP (Nakayasu and Berezney, 1989; Berezney, 1991) or 5-bromo-2'-deoxyuridine (BrdU; Nakamura et al., 1986; Nakayasu and Berezney, 1989; Vogel et al., 1989; Fox et al., 1991; Kill et al., 1991; O’Keefe et al., 1992) further revealed discrete granular sites termed replication sites (RS) that vary in number and location at different times of the S-phase in the mammalian nucleus. Analysis of the temporal and spatial order of RS has defined three major morphological patterns termed type I, type II, and type III (Nakayasu and Berezney, 1989). These three distinct replication patterns correspond to DNA synthesis in early to mid S-phase, mid to late S-phase, and very late S-phase, respectively. In cells of type I, DNA replication proceeds predominately over the extranucleolar euchromatic regions. Type II RS are concentrated over the perinucleolar and perinuclear heterochromatic regions. Type III RS correspond to DNA replication at satellite DNA heterochromatic regions (Nakayasu and Berezney, 1989; Kill et al., 1991). Other studies have demonstrated further division of the replication sites into five different intranuclear staining patterns (van Dierendonck et al., 1989; Neri et al., 1992; O’Keefe et al., 1992).

It is known that eukaryotic chromosomal DNA is divided into hundreds to thousands of independent subunits of replication termed replicons (Hand, 1978). Moreover DNA sequences are replicated at precise times within the S-phase of eukaryotic cells (Goldmen et al., 1984; Hatton...
et al., 1988), a phenomenon termed replication timing (RT). Previous studies concluded that a gene’s position in the chromosome, rather than its sequence, determines the timing of replication (Calza et al., 1984). Thus, changes in replication timing after gene rearrangement may be mediated by changes in the proximity of the affected gene to sites that control the temporal order of replication during S-phase (Calza et al., 1984; Hatton et al., 1988; Dhar et al., 1989). These studies provided the first indication that replication cluster synthesis may be temporally and spatially regulated along the chromosomal DNA.

Previous studies from our laboratory and others showed that the arrangement of replicated DNA into discrete RS-like structures persist throughout the cell cycle and subsequent daughter cells (Sparvoli et al., 1994; Berezney et al., 1995a,b; Jackson and Pombo., 1998). These results suggest that the organization of replicating DNA at distinct RS represents a fundamental feature of the higher order arrangement of chromatin in the cell nucleus. One important factor in this higher order arrangement may be the nuclear matrix. Previous studies demonstrated a remarkable preservation of RS and their S-phase–specific patterns after extraction for nuclear matrix (Nakayasu and Berezney, 1989; Neri et al., 1992; Berezney et al., 1995b).

In this study, we use fluorescence laser scanning confocal microscopy coupled to multidimensional image analysis to investigate the arrangement of individual replication sites in synchronized mammalian cells. The average number of replication sites in early S-phase and their average lifetime have been determined with great precision. Our study of the number of replication sites in early S-phase and their average lifetime have been determined with great precision. Our study of the replication timing of the in vivo replicon cluster. Further, we study the spatial and temporal dynamic of early and later replicated DNA throughout the cell cycle, and replicating timing of the early S-phase replicated DNA at two consecutive G1/S borders. We report a striking maintenance of the spatial arrangement and timing of replication sites from one cell generation to another.

Materials and Methods

Cell Culture and Synchronization

Mouse 3T3 fibroblasts (American Type Culture Collection [ATCC], Rockville, MD) were grown as monolayers in DME (GIBCO BRL, Gaithersburg, MD) plus 10% FCS (GIBCO BRL). Kangaroo kidney PtK1 cells (ATCC) were grown in MEM (GIBCO BRL) plus 10% FBS (Summit Biotechnology) supplemented with sodium pyruvate acid (GIBCO BRL) and MEM nonessential amino acids (GIBCO BRL). Cells were synchronized by serum starvation as previously reported (Nakayasu and Berezney, 1989). In brief, cells were cultured in 0.5% FCS for 3T3 cells, or 0.5% FBS for PtK1 cells for 72 h, and were released from the nonproliferating G0 phase by the addition of 10% serum into the culture medium. Cells were pulsed for early replication site labeling 10 h after serum addition.

To synchronize cells at the G1/S border, mouse 3T3 fibroblasts were serum deprived as described above. The cells were then cultured with 10% serum in the presence of 2 μg/ml aphidicolin (Cordeiro-Stone and Kaufman, 1985; Sorscher and Cordeiro-Stone, 1991; Brylawski et al., 1993) for 20 h. After aphidicolin release and an additional 10 h growth, the cells were blocked at the G1/S border in the second cell cycle by adding 2 μg/ml aphidicolin for 20 h.

Replication Site and Immunolabeling

Single Pulse or Pulse–Chase Experiments. Mouse 3T3 fibroblasts, either exponentially growing or synchronized in early S-phase by serum deprivation were pulsed with BrdU (10 μM) for 2-30 min. For chromosome spreads, PtK1 cells were synchronized by serum deprivation 10 h after serum readption (early S-phase), the cells were labeled with BrdU (10 μM) for either 2 or 15 min, chased for 10 h in fresh medium and processed for chromosomes (see Preparation of Chromosome Spreads). RS labeling was performed according to the instructions of the S-Bromo-2-Deoxy-Uridine Labeling and Detection Kit I (Boehringer Mannheim Corp., Indianapolis, IN) using reaction with mouse anti-BrdU RS FITC conjugated and antiamouse IgG (both 1:10 dilution; Boehringer Mannheim Corp.) for 30 min at 37°C. Chromosomes were counterstained with 0.025 μg/ml propidium iodide (PI).

Pulse-Chase-Pulse Experiments. In the double labeling (pulse-chase-pulse) experiments, synchronized 3T3 cells in early S-phase were labeled for 2 min with CldU (10 μM), chased for up to 10 h, and pulsed again for 5 min with IdU (10 μM).

Double replication site labeling were performed according to the method developed by Aten et al., (Aten et al., 1992) and modified in our laboratory (Berezney et al., 1995b). In brief, cells were fixed in 70% ethanol for at least 20 min in −20°C. The halogenated deoxyuridines incorporated DNA was denatured with 4 N HCl for 30 min at room temperature. Antibody reactions were carried out successively with rat anti-BrdU (1:10 dilution; Sera-Lab, Crawley Down, UK) and mouse anti-BrdU monoclonal antibodies (1:50 dilution) for 30 min at 37°C. FITC (Jackson Immunoresearch Laboratories, Inc., West Grove, PA) and Texas red (GIBCO BRL) secondary antibodies (both diluted 50-fold with PBS-0.02% Tween 20), respectively, were used for detection. High salt buffer (29.2 g/liter NaCl; 4.44 g/liter Tris-HCl, pH 7.5; and 0.5% Tween 20) was used to remove low affinity cross-reactive anti-BrdU antibodies.

Double Pulse–Double Chase Experiments. In the double pulse–double chase experiments, synchronized 3T3 cells in early S-phase were pulsed for 2 min with CldU, chased for 2 h and pulsed again for 5 min with IdU. The cells were then chased a second time for 1–48 h before fixation. Double labeling of RS was performed as described above in Pulse-Chase-Pulse Experiments.

For RS and chromosome three color labeling experiments, chromosomes were counterstained with PI (0.025 μg/ml) and displayed in pseudocolor blue. The secondary antibody against the first pulsed CldU was FITC conjugated (green color). The secondary antibody against the second pulsed IdU was Cy-5 conjugated goat anti-mouse IgG (diluted 50-fold with PBS-0.02% Tween 20), Jackson ImmunoResearch Laboratories, Inc.) and displayed in pseudocolor red.

Labeling Early Replication Site at Two Consecutive G1/S Borders. 3T3 cells synchronized at G1/S in the first cell cycle were pulsed for 30 min with CldU (10 μM) immediately after the release from aphidicolin, chased to the second G1/S border and pulsed again for 30 min with IdU (10 μM). Cell fixation, DNA denaturation, and double RS labeling was performed as described above in Pulse-Chase-Pulse Experiments.

Preparation of Chromosome Spreads

After the pulse-chase protocol for RS labeling, kangaroo kidney PtK1 cells were treated with 0.015 μg/ml colcemid (Sigma Chemical Co., St. Louis, MO) for 2 h at 37°C (Trask, 1991). The cell pellet was then incubated at 37°C for 17 min with prewarmed 75 mM KCl followed by fixation in three changes of fresh 3.1 methanol/acetic acid (10 min for the first fixation). Cells in suspension were dropped onto clear glass coverslips and incubated in ddH₂O at 4°C for at least 12 h in a humid environment to promote spreading of chromosomes. Slides were air dried overnight at room temperature. Chromosome spreads were hardened by incubated for 4 h at 37°C, and then fixed in 70% ethanol in glycine buffer, 50 mM, pH 2.0, for at least 20 min at −20°C.

Confocal Microscopy

All samples were mounted in SlowFade medium (Molecular Probes, Inc., Eugene, OR) and examined by laser scanning confocal microscopy. The confocal imaging was performed on a three channel laser scanning confocal imaging system (MRC-1024; Bio-Rad Laboratories, Hercules, CA) equipped with a Nikon Optiphot 2 microscope, a Nikon 60×/1.4 NA objective and an argon/krypton laser used to excite FITC (488 nm), Texas red or PI (568) and Cy-5 (649 nm). In some cases we used an Olympus GB200 laser scanning confocal microscope system equipped with a plan apo 60×, 1.4 NA objective and an argon laser (λ = 514 nm) to excite green and red fluorescence simultaneously. Optical sections of 512 pixels × 512 pixels × 8 bits/pixel (MRC-1024) or 1536 pixels × 1536 pixels ×
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24 bits/pixel (Olympus GB200) were collected through the samples at 0.5-μm intervals. 3-D images were reconstructed and pseudo-colored using Sterecon or VMLSM software.

**Results**

**Measuring Replication Sites in the Mammalian Cell Nucleus**

We have combined double labeling techniques, laser scanning confocal microscopy, and a newly developed image segmentation method to investigate the three-dimensional spatial and temporal properties of in vivo DNA RS in the mammalian cell nucleus. In the first series of experiments, we labeled mouse 3T3 fibroblasts with BrdU for 2–30 min. Then we analyzed the confocal images from cells engaged in early S-phase (type I; Nakayasu and Berezney, 1989) DNA replication with a spot based segmentation method (Samarabandu et al., et al., 1995). This is distinct from the commonly used threshold based methods for segmentation and enabled us to segment virtually all RS visible in the original images despite the wide range of intensities and the close proximity of many of the individual RS (Fig. 1 B). In contrast, application of a commonly used threshold-based algorithm (Samarabandu et al., 1991) resulted in very significant inaccuracies due largely to many small groups of closely positioned RS being segmented as single sites (Fig. 1 C, arrowheads) and the elimination of more weak staining RS during the thresholding (Fig. 1 C, arrow). Indeed, the total number of measured sites averaged less than one-half of that detected with our spot based segmentation in both individual optical sections (e.g., 495 versus 189 contours in Fig. 1, B and C, respectively) and in the overall nuclear volume (1,856 versus 788 sites for the image shown in Fig. 1).

Using these approaches, we have determined that the early S-phase (type I) replication patterns labeled after a 5-min BrdU pulse, contain an average of 1,080 individual sites per nucleus with an average diameter in the X-Y plane of 0.46 μm ± 0.01 (n > 12,000). Moreover, neither the X-Y diameter nor the overall volume of a RS, which represent the average of >12,000 sites for each pulse period, significantly changed between 2–15-min pulses with only a slight increase after a 30-min pulse (Fig. 2 A). There

| Pulse Time (min) | X-Y Diameter Per Site (μm) | Volume Per Site (μm³) | Density Per Site (Pixel Intensity) |
|------------------|---------------------------|----------------------|----------------------------------|
| 2                | 0.44                      | 0.16                 | 53                               |
| 5                | 0.46                      | 0.19                 | 86                               |
| 15               | 0.46                      | 0.18                 | 219                              |
| 30               | 0.48                      | 0.22                 | 293                              |

**Figure 2.** Quantitative image analysis of replication sites. (A) Volumes and labeling intensities of RS after two 30-min pulses. (B) Distribution of RS volumes after two 30-min pulses. Over 12,000 individual sites were measured for each pulse time.

![Image A](https://example.com/image1.png) ![Image B](https://example.com/image2.png) ![Image C](https://example.com/image3.png)

**Figure 1.** Computer image analysis of individual replication sites. Replication sites were single labeled for 5 min with BrdU in unsynchronized 3T3 fibroblast cells. (A) Confocal image of replication sites in 3T3 cells in a typical 0.5-μm midplane section. (B) Contours of replication sites in the optical section shown in A after the spot based segmentation method. 495 RS were counted. (C) Contours of replication sites for A using the threshold based segmentation method. 189 RS were counted. Yellow arrow indicates one cluster of weak RS that were imaged well in B but missed in C. Yellow arrowheads indicate closely spaced RS that were imaged as individual RS in B, but grouped into a single contour in C. Bar, 5 μm.

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was also a striking similarity in the volume distribution of the total population of RS for each pulse time (Fig. 2 B). In contrast, the relative intensity of the RS populations increase proportionally with pulse time (Fig. 2 A).

**Kinetic Analysis of Replication Site Lifetimes**

A series of pulse-chase-pulse experiments were performed to directly determine the average time to complete replication at individual RS. As described in Materials and Methods, synchronized 3T3 cells were pulsed in early S-phase with CldU (2 min) followed by a variable chase time (5–120 min) and a second pulse with IdU for 5 min. The strategy of this experiment is to measure the progressive decrease in overlap of early replication sites (first pulse, green color) with later replication sites (second pulse, red color). Thus the zero chase control experiment (simultaneous labeling with CldU and IdU) shows that all the labeled replication sites overlap (Fig. 3, 100% yellow sites). However, with increasing chase time between pulses there should be a progressive decrease in the number of labeled yellow sites (early and later replication at the same site) and a corresponding increase in the number of separately labeled green sites (sites labeled only with the first pulse) and red sites (sites labeled only with the second pulse). The kinetics of this progressive decay in the distribution of double labeled (Fig. 3, yellow) sites compared with singly labeled sites is a characteristic property of the average time it takes to complete replication at each site. As shown in Fig. 3, most of the replication sites are labeled as mixed yellow sites after a 5-min chase but this decreases to a small percentage after a 30-min pulse. By 1 h, all the sites are separated into red and green.

Next, we quantitated the progressive decay in mixed yellow labeled sites by direct counting of >10,000 RS for each chase time and calculating the percentage of yellow sites in the total population of yellow (early and late replication) plus green sites (early replication only). This was performed manually by direct comparison of individual sites of the separate channel and merged images. This enabled a more rigorous determination of overlap at individual sites than the simpler method of counting yellow sites in the merged images. The later method depends on a good balance of signals between the two channels whereas our method directly determines overlap without the need to visualize a yellow color. The results shown in Fig. 4, perfectly match the theoretical values obtained by assuming a replication lifetime of 45 min per site.

**Estimating the Amount of DNA and the Number of Replicons Per Replication Site**

The data generated in this study can then be used to estimate the average amount of DNA encompassed by each

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*Figure 3. Double labeling experiments in early S-phase. Replication sites in synchronized 3T3 fibroblasts were first labeled for 2 min with CldU (FITC secondary antibody, green sites), chased for 0–60 min and, pulsed again for 5 min with IdU (Texas red secondary antibody, red sites). (A) Simultaneously pulsed; chased for (B) 5 min; (C) 15 min; (D) 30 min; (E) 1 h; (F) 2 h. Bars, 5 μm.*
RS. Assuming that an average of 1,000 new replication sites are activated every 45 min during a typical 8-h S-phase, we estimate \( \approx 10,000 \) RS in the 3T3 cell. Since the 3T3 cell nucleus has \( \approx 10,000 \) mbp of DNA, each RS contains an average of 1 mbp. With a bidirectional replicative fork rate of \( \approx 3.4 \) kb (Jackson and Pombo, 1998), only \( \approx 150 \) kb of DNA can be replicated in 45 min and it would take \( \approx 300 \) min to replicate 1 mbp of DNA. Therefore, the average 1 mbp RS must be composed of at least six replicons that replicate in a relatively synchronous manner.

**Replication Sites Persist throughout the Cell Cycle**

Increasing the chase time up to 6 h in the pulse-chase-pulse experiments enabled simultaneous visualization of replication sites labeled in early S- versus mid to late S-phase. We observed type II RS around the nuclear and nucleolar periphery (Fig. 5 A, red RS) and type III RS at typical mouse centromeric heterochromatin regions (Fig. 5 B, red RS). Longer chase times (e.g., 10 h) enabled us to visualize early S-phase replication sites in the G2 phase where later S-phase replication sites are absent.

To study the fate of early S-phase RS in mitosis and the next cell generation, we labeled PTK1 cells with BrdU in early S-phase and chased the labeled cells into mitosis and the next cell generation. As previously reported by our group and others (Sparvari et al., 1994; Berezney et al., 1995b; Jackson and Pombo, 1998; Zink et al., 1998), the RS appear to be maintained during mitosis and into future cell generations (data not shown). Because it is difficult to see specific RS in the highly condensed mitotic spindle structure, we further analyzed chromosome spreads after these labeling protocols. As shown in Fig. 6, individual RS are easily resolved with this technique and are arranged along the chromosomes in a pattern reminiscent of chromosome bands. Remarkably this band-like pattern of individual RS along the chromosomes was even apparent at the earliest pulse period investigated (Fig. 6, A, C, D, and F, 2 min pulse). Previous studies have also reported band-like arrangement of BrdU labeled replication sites on mitotic chromosomes (Sparvoli et al., 1994; Ferreira et al., 1997). The doublets or sister chromatid replication pairs characteristic of RS labeling on mitotic chromosomes (Ferreira et al., 1997; Jackson and Pombo, 1998) were also observed (Fig. 6, D and J, arrows). However, the limited degree of swelling in these chromosome spreads compromised the efficiency of identifying sister chromatid replication pairs.

**Figure 5.** Double labeling experiments in early S-phase versus later S-phase and the G2-phase. Replication sites in synchronized 3T3 fibroblasts were first labeled for 2 min with CldU (FITC secondary antibody, green sites), chased for 4–10 h and pulsed again for 5 min with IdU (Texas red secondary antibody, red sites). (A) 4-h chase (mid S-phase, type II for red RS); (B) 6-h chase (late S-phase, type III for red RS); (C) 10-h chase (G2-phase, there are no red RS). Bars, 5 μm.
DNA from Replication Sites Labeled in One Cell Generation Does Not Significantly Mix with Later Replication Sites in Future Cell Generations

It was previously demonstrated (Figs. 3 and 4) that early replicated DNA is completely distinguishable from later replicated DNA by occupying separate RS after a 1- or 2-h chase. Therefore, we designed double pulse–double chase experiments (see Materials and Methods) to determine whether the DNA sequences replicated at individual RS maintain their separation or mix together as the cells proceed through subsequent cell generations. After the first 2-h chase between early and later labeling of RS, the cells are secondarily chased into late S- or G2-phase (Fig. 7 A), M-phase (Fig. 7, D–F) and into subsequent cell generations (Fig. 7, B and C). The results demonstrate that temporally distinct early S-phase RS (Fig. 7, green) and later replicated sites (red) are totally maintained in spatially distinct domains throughout the S-phase and in future cell generations. As a control, the replication sites are pulsed with CldU and IdU simultaneously in early S-phase and chased through the same time span. Yellow RS are maintained completely throughout this extensive chase period (data not shown). Separation of the early and later labeled RS is especially evident in chromosome spreads that reveal repeating band-like clusters of green versus red sites (Fig. 7, D and F, arrows) and confirm the previous observations by Ferreira et al. (1997) who used longer pulse labeling. We also observe a significant decrease in the number of replication sites in the third generation cells (Fig. 7 C) compared with first and second generation cells (Fig. 7, A and B), which is consistent with semiconservative replication (Sparvoli et al., 1994; Ferreira et al., 1997; Jackson and Pombo, 1998).

DNA Replicated in Early S-Phase Is Replicated at the Same Time and Site in the Next Round of Replication

The previous results demonstrate that the DNA replicated at individual RS in early S-phase is spatially maintained at distinct sites in mitotic chromosomes and in subsequent cell generations (Fig. 7). We next addressed whether this spatially distinct group of RS replicate at the same time in the next cell generation. For these experiments, mouse 3T3 cells were synchronized at the G1/S border with aphidicolin for two successive cell generations and pulsed with CldU and IdU, respectively, immediately after release from the G1/S border (see Materials and Methods and Fig. 8 B for details). Under these conditions, we achieved a high degree of synchrony in early S-phase with >99% of the S-phase cells (507 counted) having type I early S-phase replication patterns immediately after release and >90% of the cells entering S-phase within 2 h of releasing the aphidicolin block (data not shown). The results shown in Fig. 8 A demonstrate complete overlap of the RS labeled at two successive early S-phases. We conclude that DNA synthesized in early S-phase is replicated at the same time and at the same site in the next round of replication.
Discussion

Fluorescence microscopic analysis of newly replicated DNA has revealed discrete granular sites of replication (Nakamura et al., 1986; Nakayasu and Berezney, 1989; Mills et al., 1989; Berezney, 1991; Kill et al., 1991; Fox et al., 1991; O'Keefe et al., 1992). Moreover, the pioneering studies of Nakamura et al. (1986) showed that increasing times of BrdU in vivo labeling of DNA RS leads to progressively more elongated structures and to increased difficulty in resolving individual sites. We are using fluorescence laser scanning confocal microscopy in conjunction with a spot-based segmentation algorithm (Samarabandu et al., 1995) to address these issues and to gain more precise spatial information about individual RS and their potential organization into higher order domains.

We report an average of ~1,100 replication sites after a 5-min pulse of 3T3 mouse fibroblasts synchronized in early S-phase. Similar results were obtained in exponentially dividing cells where selection for measurement was based solely on visible type I patterns (data not shown). Our results, therefore, suggest that ~1,000 RS are active at any time in early S-phase. This number is several fold higher than previously reported values (100–300) using both epifluorescence and laser scanning confocal microscopy (Nakamura et al., 1986; Mills et al., 1989; Nakayasu and Berezney, 1989; Fox et al., 1991; Nakayasu and Berezney, 1989; Fox et al., 1991; Kill et al., 1991; Hozak et al., 1993; Hassan and Cook, 1993) as well as when the very same samples were analyzed by a widely adopted threshold-based segmentation algorithm (Samarabandu, 1991; see Fig. 1). Direct visual comparison demonstrates that our newly developed spot-based segmentation method (Samarabandu et al., 1995) is extremely accurate and sensitive in distinguishing individual sites over a broad range of intensities and in close spatial apposition (Fig. 1). In contrast, threshold-based methods result in nondetection of more weakly stained sites and the merging of closely positioned individual sites into one contour (see Fig. 1). Recently, Jackson and Pombo (1998) have applied a direct
RS is composed of at least six average-sized replicons. The overall population average is heterogeneous, ranging in size from 100–150 kb in mammalian cells. Thus each mbp RS is composed of at least six average-sized replicons.

Jackson and Pombo (1998) have come to similar conclusions based on DNA fiber analysis of replicon clusters size: the RS observed in the cell nucleus directly correspond to replicon cluster observed with DNA fiber analysis.

Previous studies have shown that postreplicated DNA maintains a discrete RS pattern throughout the cell cycle and into subsequent cell generations (Sparvoli et al., 1994; Berezney et al., 1995a,b; Jackson and Pombo, 1998; Zink et al., 1998). This has led to the view that DNA arranged in domains of approximately mbp size may be a fundamental feature of higher order arrangement of chromatin in the cell nucleus (Berezney et al., 1995a,b). An important question is whether the DNA that composes the individual RS is also maintained in these sites as the cell progresses through the cell cycle and from one cell generation to another. In other words, whereas individual RS may be reflecting a fundamental feature of higher order chromatin organization, the actual DNA sequences that compose individual sites may fluctuate. To address this we labeled the replicating DNA in early S-phase in two colors with a 2-h chase between the brief pulses. We then chased the spatially separate red and green sites through S-phase, mitosis and into the next two cell generations (Fig. 7). If the DNA in individual RS is promiscuous, then we should observe mixing of the separate green and red sites through S-phase, mitosis and into subsequent cell generations (Sparvoli et al., 1994; Berezney et al., 1995a,b; Jackson and Pombo, 1998; Zink et al., 1998). We conclude that the DNA at individual RS is maintained with a high degree of fidelity through at least several cell generations.

There is growing evidence that DNA replication in the eucaryotic cell proceeds in a precisely choreographed manner whereby specific groups of replicons are active at different times in S-phase. Studies of specific gene se-
quences have shown that most active genes replicate in early S-phase, whereas inactive ones tend to replicate in late S-phase (Goldman et al., 1984; Hatton et al., 1988). The higher order arrangement of chromatin into ~1-mbp domains of replication (RS) may represent the spatial basis for this replication timing. In pioneering studies, Schildkraut and his associates demonstrated that the position of a gene on a chromosome determines its replication timing (Calza et al., 1984; Dhar et al., 1988, 1989; Hatton et al., 1988). Specific timing was measured for up to several hundred kilobases in several multi-gene families (Brown et al., 1987; Dhar et al., 1988, 1989; Hatton et al., 1988). More recently, Selig et al. (1992) identified replication time (control) zones exceeding 500 kb within and flanking the cystic fibrosis gene on human chromosome 7 and suggested that these replication zones correspond to basic units of chromosome structure for both DNA replication and transcriptional regulation. Consistent with this view, replication control zones that range from several hundred kilobases to >1 mbp have been identified in the delayed replication of the FMR1 gene associated with fragile X syndrome (Hansen et al., 1993) and in the imprinted human Igf2 gene domain (Kitsberg et al., 1993).

Based on these findings and relationships, we suggest that replication time (control) zones correspond to individual RS, which are, in turn, composed of a cluster of replicons that are subject to replication timing control. To test this concept experimentally, we labeled RS at two consecutive S-phases after release from the G1/S borders. The results strikingly demonstrate complete colocalization of the RS (Fig 8). In other words, the large subset of RS at this discrete time in very early S-phase (~1,000 sites) behave as a collection of replication time zones and lead to the replication of the same subset of RS from one cell generation to another. However, due to the limited resolution of our approach, we can not rule out at this time a degree of fluctuation among the specific DNA sequences at individual sites. Future studies designed to map DNA sequences at individual replication sites will provide a more rigorous test. Moreover, Jackson and Pombo (1998) have come to similar conclusions using DNA fiber analysis after double labeling. This enables direct visualization of overlap at the level of individual replicons.

Whereas these studies are an important advance in understanding the relationship of chromatin organization and function in the cell nucleus, they do not directly address the nature of this higher order arrangement of chromatin. Indeed, our knowledge of how chromatin is arranged in the nucleus beyond the arrangement of nucleosomes into a 30-nm solenoid or zigzag structure (McGhee et al., 1980; McGhee et al., 1983; Butler, 1984; Pienta and Coffey, 1984; Thomas, 1984; Sen et al., 1986; Horowitz et al., 1994) is still in its infancy. However, direct ultrastructural studies of chromatin inside the cell nucleus have suggested a higher level of organization that is consistent with the ~1-mbp sites that we can identify with replication labeling (Lau and Arrighi, 1981; Mullinger and Johnson, 1983; Belmont et al., 1989; Manuelidis, 1990). The relationship of functional domains of replicon clusters to chromosomes bands has attracted particular attention. Indeed, the discrete patterns of R-bands versus G/Q bands observed in early versus later S-phase has lead to the proposal that individual bands correspond to replicon clusters (Klevecz and Keniston, 1975; Latt, 1975; Stubblefield, 1975; Lau and Arrighi, 1981; Holmquist et al., 1982).

Our results, however, suggest that chromosome bands may be composed of several RS (see Figs. 6 and 7) and are consistent with previous findings (Sparvari et al., 1994; Ferreira et al., 1997). Thus the equivalent of a chromosome band in the interphase nucleus may be a small cluster of several RS. Recently we have used quantitative image analysis to demonstrate that a large percentage of individual RS are indeed arranged in clusters of higher order domains or zones in the nucleus (Wei et al., 1998). Moreover, these higher order replication zones are spatially segregated from analogous zones containing clusters of transcription sites (Wei et al., 1998). Distinct clusters of RS have also been observed after BrdU labeling of RS followed by chase times of up to 10 cell generations and are believed to represent chromosome territories or territorial sub-regions (Jackson and Pombo, 1998; Zink et al., 1998).

The relationship of the replication and transcription zones elucidated by Wei et al. (1998) to chromosome territories remains to be elucidated.

Numerous findings point to the nuclear matrix as a key player in organizing the higher order domain structure of chromatin (Berezney and Jeon, 1995). Chromatin domains are believed to be attached to the nuclear matrix and chromosome scaffold in repeating 50–200-kb loops (Vogelstein et al., 1980; Goldberg et al., 1983; Nelson and Coffey, 1987; Chai and Sandberg, 1988; Laemmli et al., 1992; Roberge et al., 1992; Hiraoka et al., 1993). These loop domains may provide the structural basis for both domains of transcription and individual replicons (Vogelstein et al., 1980; Jackson and Cook, 1995). Models have been proposed that describe the replication of DNA on the nuclear matrix as a cluster of replicons in loop form (Dijkwel et al., 1979; Pardoll et al., 1980; Berezney and Buchholtz, 1981; Berezney, 1984; Cook, 1991). Consistent with these biochemical findings and models, the RS and the various patterns of RS characteristic of different stages in S-phase are strikingly maintained after extraction of cells for nuclear matrix (Berezney and Nakayasu, 1989; Neri et al., 1992).

These results suggest that the matrix may be involved in chromatin organization at levels even higher than the 100-kb loop domains such as the 1 mbp RS elucidated in this study. However, it is important to recognize that the precise role(s) of the nuclear matrix in chromatin organization remain(s) to be defined. For example, it can be argued that nuclear matrix organization may be regulated by chromosome territory organization rather than vice versa (Cremer et al., 1995). In this regard, we recently found that the integrity of chromosome territories is dependent on the corresponding integrity of the nuclear matrix (Ma, H., and R. Berezney, manuscript submitted for publication).

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