Spatial Relationship between Transcription Sites and Chromosome Territories

Pernette J. Verschure, Ineke van der Kraan, Erik M.M. Manders, and Roel van Driel

E.C. Slater Instituut, BioCentrum Amsterdam, University of Amsterdam, 1018 TV Amsterdam, The Netherlands

Abstract. We have investigated the spatial relationship between transcription sites and chromosome territories in the interphase nucleus of human female fibroblasts. Immunolabeling of nascent RNA was combined with visualization of chromosome territories by fluorescent in situ hybridization (FISH). Transcription sites were found scattered throughout the territory of one of the two X chromosomes, most likely the active X chromosome, and that of both territories of chromosome 19. The other X chromosome territory, probably the inactive X chromosome, was devoid of transcription sites. A distinct substructure was observed in interphase chromosome territories. Intensely labeled subchromosomal domains are surrounded by less strongly labeled areas. The intensely labeled domains had a diameter in the range of 300–450 nm and were sometimes interconnected, forming thread-like structures. Similar large scale chromatin structures were observed in HeLa cells expressing green fluorescent protein (GFP)-tagged histone H2B. Strikingly, nascent RNA was almost exclusively found in the interchromatin areas in chromosome territories and in between strongly GFP-labeled chromatin domains. These observations support a model in which transcriptionally active chromatin in chromosome territories is markedly compartmentalized. Active loci are located predominantly at or near the surface of compact chromatin domains, depositing newly synthesized RNA directly into the interchromatin space.

Key words: chromosome territory • transcription sites • nuclear organization • chromatin structure • fluorescent in situ hybridization

The dynamic structure of the interphase nucleus is an important factor in the control of gene expression. Many nuclear functions, such as replication, transcription, and RNA processing, occupy well-defined domains in the interphase nucleus (Spector, 1993; Wansink et al., 1994b; van Driel et al., 1995; de Jong et al., 1996; Jackson, 1997; Lamond and Earnshaw, 1998). Mammalian chromosomes in the interphase nucleus are localized in distinct territories (Lichter et al., 1988; Pinkel et al., 1988; Cremer et al., 1993). The spatial organization of nuclear domains with respect to chromosome territories has become a focus of interest in the analysis of functional architecture of the interphase nucleus. For instance, Zirbel et al. (1993) have shown that components of the splicing machinery, as well as specific gene transcripts of an integrated human papilloma virus genome, are often excluded from the interior of chromosome territories. Furthermore, several coding genes were found positioned preferentially at the periphery of chromosome territories. A apparently, this configuration was independent of the transcriptional activity of the genes. Noncoding genomic DNA was found more randomly positioned in chromosome territories (Kurz et al., 1996). Since these observations are related to only a small set of genomic sequences, further studies are needed to explore whether nuclear compartmentalization with respect to the structural organization of interphase chromosomes is of functional significance for gene control.

The aim of the present study is to elucidate whether transcription sites have a distinct spatial distribution in relation to the organization of chromosome territories and chromatin. Detection of nascent RNA with the use of 5-bromouridine 5'-triphosphate (BrUTP) by in vivo and in vitro labeling procedures was developed by Wansink et al. (1993) and Jackson et al. (1993). These studies have shown that transcription sites in cultured mammalian cells are concentrated in numerous domains that occur scattered throughout the nucleoplasm. In the present study, detection of nascent RNA was combined with labeling of chromosome territories by fluorescence in situ hybridization (FISH), using chromosome-specific DNA whole library.

Address correspondence to P.J. Verschure, E.C. Slater Instituut, BioCentrum Amsterdam, University of Amsterdam, Plantage Muidergracht 12, 1018 TV Amsterdam, The Netherlands. Tel.: 020-5255151. Fax: 020-5255124. E-mail: a311pjvx@chem.uva.nl

1. Abbreviations used in this paper: 3-D, three dimensional; BrUTP, 5-bromouridine 5'-triphosphate; FISH, fluorescent in situ hybridization; GFP, green fluorescent protein; Xa, active X chromosome; Xi, inactive X chromosome.

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probes. Labeling of nascent rRNA in combination with visualization of X chromosome and chromosome 19 territories was analyzed in diploid human female fibroblasts. In mammals, one X chromosome is transcriptionally inactivated during early development, while the other remains active (Lyon, 1961; Gartler and Goldman, 1994; Rastan, 1994; H eard et al., 1997). The two X chromosome territories in somatic female cells nuclei provide an excellent model system to study the distribution of transcription sites in the active X chromosome (Xa), using the inactive X chromosome (Xi) as an internal control.

Our results show that transcription sites are scattered throughout the chromosome territories of Xa and chromosome 19, whereas Xi is devoid of transcription sites. We observe that interphase chromosome territories show a distinct substructure, consisting of domains that were intensely labeled in the FISH procedure, surrounded by less intensely labeled areas. Strikingly, newly synthesized and nascent rRNA was preferentially present in interchromatin areas within chromosome territories. These interchromatin areas contain little or no DNA. Since in situ hybridization involves relatively harsh conditions, we also analyzed the relationship between chromatin and transcription sites by investigating the spatial relationship between GFP-tagged histone H2B and transcription sites. Nascent rRNA was present in between intensely labeled chromatin domains, similar to that observed after performing FISH. These results indicate that there is a marked compartmentalization of transcriptionally active and inactive chromatin in chromosome territories.

Materials and Methods

Cell Culture

Human primary fibroblasts (kindly provided by Dr. L.H.F. Mullenders, University of Leiden, The Netherlands) with a normal female karyotype (46, XX) were grown at 37°C under a 2.5% CO₂ atmosphere in Ham’s F-10 (GIBCO BRL), supplemented with 15% (wt/vol) heat-inactivated FCS (Boehringer Mannheim Corp.), 2 mM L-glutamine (GIBCO BRL), 100 IU/ml penicillin and 100 μg/ml streptomycin (GIBCO BRL). Cells were grown to 50–70% confluency before use. Preannealed probes gave the same result in the FISH procedures as using a large excess of Cot-1 DNA. The probes have been shown to label their respective metaphase chromosomes completely and specifically, showing the specificity of the DNA library probes (Carter et al., 1990; Telenius et al., 1992). In some experiments, we specifically labeled only repetitive sequences. In that case, we used human Cot-1 DNA that was directly conjugated to FITC or Cy3 (kindly provided by Dr. J.C.A.G. Wiegant, University of Leiden, The Netherlands; 20 μg/ml).

Immunofluorescent Labeling of Acetylated Histone H4, Centromeres and PML Bodies

For immunofluorescent labeling, cells were fixed for 10 min at 4°C in 2% (wt/vol) formaldehyde in PBS. After fixation, cells were permeabilized with 0.5% (wt/vol) Triton X-100 in PBS for 5 min and incubated with PBS containing 100 mM glycine for 10 min. Subsequently, cells were incubated for 1 h at 37°C with an antibody recognizing acetylated histone H4, an antibody recognizing centromeres, or an antibody recognizing PML. Rabbit polyclonal antibodies (R232 and R252) against acetylated histone H4 (kindly provided by Dr. B. Turner, University of Birmingham, UK) were used. R232 polyclonal antibody recognizes histone H4 acetylated at lysine 8, and antibody R252 histone H4 acetylated at lysine 16 (Turner and Fellows, 1989, Turner et al., 1989). Human autoimmune serum H3 recognizes centromeres (kindly provided by Dr. W.J. Van Vennovo, K aholieke Universiteit Nijmegen, The Netherlands). To label PML protein in PML antibodies, we used the mouse mAb 5E10 (Stuurman et al., 1992; K oken et al., 1994). R232 and R252 antibodies were diluted 1:1,000, anticer-tromere 1:50,000, and 5E10:13 in PBS containing 0.1 μg/ml herring sperm DNA. A fer several washes in PBS, cells were incubated with appropriate secondary antibodies, using either Cy3- or FITC-conjugated donkey anti-rabbit antibody, goat anti-human antibody, or donkey anti-mouse antibody (Jackson Immunoresearch Laboratories, Inc.). Secondary antibodies were diluted in PBS containing 0.1 μg/ml herring sperm DNA and incubations were performed for 1 h at room temperature. As a control, cells were incubated in the absence of primary antibody. Subsequently, cells were washed in PBS and used for the in situ hybridization procedure.

Chromosome-specific Probes

Painting of the human X chromosome and of chromosome 19 was achieved using chromosome-specific biotin-labeled DNA library probes (Cambio). Probes were denatured at 85°C for 5 min. To suppress labeling of repetitive sequences, which are often not chromatin-specific, denatured probe samples (85°C, 5 min) were self-annealed (2-3 μg/ml, 60-100 min, 37°C) before use. Preannealed probes gave the same result in the FISH procedures as using a large excess of Cot-1 DNA. The probes have been shown to label their respective metaphase chromosomes completely and specifically, showing the specificity of the DNA library probes (Carter et al., 1990; Telenius et al., 1992). In some experiments, we specifically labeled only repetitive sequences. In that case, we used human Cot-1 DNA that was directly conjugated to FITC or Cy3 (kindly provided by Dr. J.C.A.G. Wiegant, University of Leiden, The Netherlands; 20 μg/ml).

In Situ Hybridization Procedure

The FISH procedure was an adaptation of protocols described by Lichter et al. (1988), Cremer et al. (1988, 1993), and Kurz et al. (1996). Fixation, pretreatment, and denaturation procedures were subjected to several modifications. The following protocol was found to produce optimal preservation of intranuclear organization of transcription sites and H4 acetylated histones. Cells were rinsed with PBS and fixed for 10 min at 4°C in 4% (wt/vol) formaldehyde diluted in PBS. To facilitate probe penetration, cells were treated with 0.1 M HCl for 10 min and subsequently with a mixture of 0.5% Triton and 0.5% (wt/vol) saponin in PBS for 10 min. After washing in PBS, genomic DNA was denatured by incubating cells in 2× SSC containing 70% formamide at 73°C for 4 min, followed by incubation in 2× SSC containing 50% formamide for 1 min. Hybridization with the biotin-tagged probe was allowed to proceed overnight at 42°C. Posthybridization washes with 2× SSC containing 50% formamide and subse-
Slides were kept at 4°C in the absence of probe. Slides were mounted in Vectashield (Brunschwig). As a control, cells were incubated in hybridization buffer with 4× sperm DNA. The wash steps after antibody incubations were performed with 0.4× SSC, pH 7.4. TWEEN 20. A fluorescent labeling, DNA staining was performed with 0.4 μg/ml Hoechst 33258 (Sigma Chemical Co.) in PBS, or with 0.5 μM Sytox green (Molecular Probes) in 25 mM Hepes buffer, pH 7.4. As a control, cells were incubated in hybridization buffer in the absence of probe. Slides were mounted in Vectashield (Brunschwig). Slides were kept at 4°C until evaluation and analyzed within 24 h.

Confocal Scanning Laser Microscopy

Images were recorded with a Leica or LSM 510 Zeiss confocal laser scanning microscope both equipped with a 100×/1.23 NA oil immersion lens. With the Leica microscope, a dual-wavelength argon ion laser was used, whereas the Zeiss confocal microscope used an argon laser at 488 nm in combination with a helium neon laser at 543 nm to excite green and red fluorochromes simultaneously at 488 nm and 514 nm, respectively. Emitted fluorescence was detected in the Leica confocal microscope using a 525 D F 10 bandpass filter for FITC and a 550-nm longpass filter for Cy3, whereas in the Zeiss microscope a 505-530-bandpass filter for green signal and a 560-nm longpass filter for the red signal was used. Pairs of images were collected simultaneously in the green and red channels. Three-dimensional (3-D) images were scanned as 512 × 512 × 32 voxel images (sampling rate 49-nm lateral and 208-nm axial).

Image Processing

Images were corrected for optical cross-talk (Manders et al., 1992). Image analysis was performed using SCIL-IMAGE software (Ten Kate et al., 1990; van Balen et al., 1994). Images were subjected to a 3-D image restoration procedure to correct for diffraction-induced distortion, using the Huygens System 2 (Scientific Volume Imaging BV; Van der Voort and Straster, 1985). The image restoration procedure uses a measured point spread function, which was obtained at precisely the same conditions as the image. To this end, 200-nm fluorescent beads (FluoSpheres; Molecular Probes, Inc.) were imaged. The 3-D image restoration procedure significantly improved the quality of the 3-D images by removing Poisson noise and by deblurring the images.

Results

Structural Preservation during In Situ Hybridization

We have developed a modified chromosome painting protocol with improved signal to background ratio, based on the chromosome painting procedures described by Lichter et al. (1988), Cremer (1993), and K urz et al. (1996). It was important to establish that the relatively harsh FISH procedure did not alter nuclear structure in general, and chromosome structure in particular. To investigate the degree of structural preservation, we have carried out four tests. First, we analyzed whether the FISH procedure induced an altered spatial distribution of transcription sites. Second, we investigated the effect of the FISH procedure on the distribution of DNA stained with Sytox green. Third, we analyzed the spatial distribution of acetylated histone H4 before and after the chromosome painting procedure. Histone acetylation is generally correlated with gene activity and reduced acetylation levels with gene silencing (H ebbes et al., 1988; Braunstein et al., 1993). Fourth, we measured in one and the same cell nucleus before and after the FISH procedure the spatial distribution of centromeres and of PML bodies (Stuurman et al., 1992).

Visual inspection shows that the spatial distribution of nascent RNA did not change in any recognizable way due to the FISH procedure (Fig. 1, A and B). Staining of DNA with Sytox green with or without chromosome painting showed that the FISH procedure also did not induce major changes in DNA distribution (Fig. 1, C and D). After the FISH procedure, the DNA staining pattern seemed slightly more blurred and the overall DNA labeling appeared more intense. Fig. 1, E and F, show immunofluorescent labeling of histone H4 acetylated at lysine 8, before and after FISH labeling. The punctate distribution of acetylated H4 throughout the nucleoplasm did not significantly change after chromosome painting. Only the dif-
fuse, low intensity component in the labeling was somewhat diminished after FISH, making the intense punctate granular labeling more prominent. This may be due to extraction of some histone protein during FISH labeling. These results show that the FISH procedure does not result in major rearrangements in DNA, chromatin, and nascent RNA.

Fig. 2 shows the spatial distribution of PML bodies (Fig. 2, A–C) and of centromeres (Fig. 2, D–F) imaged in the same cell before and after FISH. The corresponding optical sections before and after the FISH procedure (PML bodies, Fig. 2, A and B, and centromeres, D and E, respectively) show that nuclear organization is well-preserved during the FISH procedure. Both the integrity and the spatial distribution of the nuclear bodies and of the centromeres are essentially unchanged. The small changes in spatial distribution (see overlay in Fig. 2, C and F) can be attributed largely to a slight tilting of the nucleus relative to the substratum, due to the FISH procedure.

Together, these results establish that the FISH procedure does not induce significant changes in the spatial distribution of subnuclear structures at the light microscopy level. In particular, we did not observe any collapse, aggregation, unfolding, or other major rearrangement of chromatin in the nucleus. We conclude that, at the resolution level of the light microscope, nuclear structure is not affected by the chromosome painting procedure.

Substructure of Chromosome Territories

A striking feature that emerges from our confocal images after FISH labeling is that chromosome territories display a distinct substructure (Fig. 3). Chromosome territories do not appear as compact objects. Rather, they show a modulated intensity distribution inside the territory. Since the chromosome-specific library-probes label the complete metaphase chromosome, it is unlikely that considerable parts of the interphase chromosome remain unlabeled (Carter et al., 1990; Telenius et al., 1992). The chromosome territories contained strongly labeled chromosomal subdomains, surrounded by less intensely labeled areas. The strongly labeled subchromosomal structures have a diameter in the range of 300–450 nm (Figs. 3 and 4). Intensely labeled parts of a chromosome often seemed interconnected, forming thread-like, folded structures. A similar, distinct substructure in nuclei stained for DNA with Sytox green was observed (Fig. 1, C and D) and in nuclei of HeLa cells expressing GFP-histone H2B (see Visualization of Chromatin Using GFP-histone H2B; Fig. 5), suggesting a reticular organization of chromatin. It is tempting to suggest that, at least locally, the chromatin fiber that constitutes the chromosome can be followed in an optical section.

Images were subjected to 3-D image restoration to correct for diffraction-induced distortions. The advantage of such image restoration is that it significantly improves the quality of the 3-D images. It is conceivable that the restoration procedure preferentially enhances certain spatial frequencies. To verify this, we compared the unprocessed individual consecutive sections through a chromosome territory (Fig. 3, B and D) to the same sections after 3-D image restoration (Fig. 3, C and E). Careful comparison
shows that essentially all details of a chromosome territory in
the processed image are precisely matched by features that
can be found in the preprocessed image. Apparent slight dif-
f erences are most likely due to the fact that the 3-D restored
images contain information from optical sections above and
below those that are shown, whereas the unprocessed images
do not. Very low intensity signal may be lost by the image
restoration procedure, probably because it cannot be dis-
 criminated from noise in the restoration process. These re-
 sults show that no artificial structures in chromosome territo-
 ries are created by the 3-D restoration algorithm.

To investigate whether the less intensely labeled areas
in chromosome territories contained repetitive DNA,
which is suppressed during FISH labeling, we performed
hybridization with a probe specific for highly repetitive
human Cot-1 DNA, in dual labeling with an X chromosome
probe (data not shown). Hybridization of metaphase
chromosomes with the Cot-1 probe revealed intense label-
ing predominantly in the centromeric region, which is
known to contain a high concentration of repetitive DNA.
Simultaneous in situ hybridization with the X chromo-
some probe and the probe recognizing highly repetitive
DNA showed that repetitive DNA does not occur concen-
trated in the areas in the territory that are less intensely la-
beled with the chromosome-specific probe in the FISH
procedure. We conclude that the less intensely labeled ar-
 eas in chromosome territories constitute a compartment
that contains little or no DNA.
Figure 4. Distribution of transcription sites in relation to territories of the two X chromosomes and chromosome 19 in female primary fibroblasts. Labeling of nascent RNA by incorporation of BrUTP (green) in permeabilized cells (run-on-transcription) is combined with FISH labeling of X chromosome territories (A–C) and of chromosome 19 territories (D). In all nuclei analyzed, transcription sites occur as defined spots throughout one of the two X chromosome territories (most likely Xa; B) and in both chromosome 19 territories (D). A, Almost no transcription sites are observed in the other X chromosome territory (most likely Xi; C). Nascent RNA preferentially accumulates between the chromosomal subdomains in the areas with little or no FISH label. A, Single optical section through the center of a cell nucleus with labeled Xa and Xi chromosome territories (red) and transcription sites (green). Bar, 0.875 μm. B, Four consecutive
Transcription Sites throughout Chromosome Territories

To establish the 3-D distribution of transcription sites in relation to chromosome territories, chromosome painting was combined with the visualization of transcription sites. Fluorescent labeling of nascent RNA shows that transcription occurs scattered throughout the nucleoplasm (Jackson et al., 1993; Wansink et al., 1993, 1994a), often with exception of nucleoli, due to inaccessibility of the anti-Bru antibody (Wansink et al., 1993). The 3-D images show that transcription sites occur throughout the territory of one of two X chromosomes (Fig. 4, A and B), whereas almost no transcription sites are found in the other X chromosome (Fig. 4, A and C). Most likely, the X chromosome territory containing transcription sites represents the Xa and the territory devoid of transcription is the Xi. We also analyzed the distribution of transcription sites with respect to the autosomal chromosome 19. Chromosome 19 replicates early in S phase and is relatively gene rich, as indicated by the number of CpG islands (Craig and Bickmore, 1994). Transcription sites were found throughout the chromosome 19 territories (Fig. 4 D), similar to that observed for Xa. These results show that transcription sites are present throughout chromosome territories with exception of the Xi chromosome territory. Importantly, no striking differences in the substructure of the territories of the Xa, Xi, or chromosome 19 can be seen. This implies that the substructure does not depend on differences in overall transcriptional activity, or on differences between sex chromosomes and autosomal chromosomes.

Relationship between Transcription Sites and Chromosomal Substructure

We have analyzed the relationship between the spatial distribution of transcription sites and the substructure of chromosome territories. Strikingly, transcription sites (green signal in Fig. 4 A, and B III and IV) in Xa, as well as in chromosome 19 (Fig. 4 D, III and IV), were mainly found between the intensely labeled chromosomal subdomains (red signal in Fig. 4 A, B III and IV, and D III and IV) and almost never overlapped with them. Line scans were made to obtain more quantitative information about the spatial relationship between transcription sites and chromosome substructure. Fig. 4 E shows typical examples of line scans through Xa (Fig. 4 E, 1 and 2) and Xi (Fig. 4 E, 3) and chromosome 19 (Fig. 4 E, 4) territories. The position of the scans are marked in Fig. 4, B III and C IV. The line scans through the territory of putative Xa and of chromosome 19 territories confirm that transcription sites (indicated by arrowheads) are often localized near the surface of the intensely labeled chromosomal subdomains (Fig. 4 E, 1–3). The transcription sites marked with an asterisk have their 3-D center of gravity exactly on the scanned line. These transcription sites clearly show that nascent and newly synthesized RNA accumulates near a chromosomal subdomain, generally not overlapping with it.

Visualization of Chromatin Using GFP-histone H2B

Despite our efforts (see Structural Preservation during In Situ Hybridization) we cannot fully rule out that the FISH procedure induces local rearrangement of chromatin and/or nascent RNA. Therefore, we have used an alternative procedure to label chromatin, using cells that stably express GFP-tagged histone H2B (Kanda et al., 1998). Since essentially all histone protein is incorporated in chromatin, the distribution of GFP–H2B faithfully represents that of chromatin (Kanda et al., 1998). Obviously, this procedure does not allow visualization of individual chromosome territories. However, it does allow analysis of the chromatin distribution in nuclei under in vivo conditions. Intense GFP-labeling is seen particularly at the nuclear periphery and around nucleoli, suggesting a compact, heterochromatin-like local structure. In addition, an irregular, apparently reticular labeling of chromatin is seen throughout the nucleoplasm, with exception of the nucleoli. It is important to note that, like after FISH labeling, intensely labeled areas are observed with a diameter in the range of 300–450 nm (Fig. 5, B and E). Visualizing transcription sites in cells expressing GFP–H2B showed strikingly little overlap between transcription sites (red signal in Fig. 5, C, F, and both enlargements) and chromatin (green signal in Fig. 5, C, F, and both enlargements). Line scans confirm these observations (Fig. 5 G, 1 and 2). This observation is fully in agreement with the results after FISH labeling of chromatin in chromosome territories (see Relationship between Transcription Sites and Chromosomal Substructure). These results confirm that FISH does not affect nuclear structure at the light microscopy level and under-
score our observations showing that chromosome territories have a distinct substructure and that transcription sites are located near the surface of intensely labeled subchromosomal domains.

**Discussion**

Two major principles of organization of the interphase nucleus can be distinguished. First, many nuclear components and processes occur compartmentalized in well-defined domains in the interphase nucleus (Spector et al., 1993; Wansink et al., 1994b; Van Driel et al., 1995; De Jong et al., 1996; Lamond and Earnshaw, 1998; Mistelli and Spector, 1998). Second, individual chromosomes occupy discrete chromosome territories that are apparently not intruded by chromatin from other chromosomes (Lichter et al., 1988; Pinkel et al., 1988; Cremer et al., 1993). The dynamic structure of the interphase nucleus in general, and that of chromosomes and higher-order chromatin in particular, are important elements in the control of gene expression. Here, we investigate the spatial relationship between transcription sites and chromosome territories in the interphase nucleus. Chromosome territories were labeled by FISH, using chromosome-specific DNA whole library probes (Lichter et al., 1988; Jauch et al., 1990). We have analyzed in female primary fibroblasts the localization of transcription sites in relation to the territories of chromosomes X and 19. In addition, we have examined the spatial relationship between total chromatin in nuclei of HeLa cells that express GFP-tagged histone H2B.

*Figure 5. Distribution of transcription sites in relation to chromatin in HeLa cells expressing GFP tagged histone H2B. Transcription sites were visualized after in vivo incorporation of BrUTP into nascent RNA. Nascent RNA accumulated preferentially in nuclear domains that contain little or no GFP-labeled chromatin. A and D, Single optical sections of two different nuclei showing the distribution of transcription sites. B and E, Same optical as shown in A and D, respectively, showing the distribution of GFP-histone H2B. C and F, Overlay of A and B, and D and E, respectively, showing the spatial relationship between transcription sites (red) and chromatin domains (green). Magnified areas in C and F (white squares) are shown in the inserts at the right. The images show that nascent RNA is predominantly found in areas that contain little or no GFP-labeled chromatin. All images shown are subjected to 3-D image restoration. G, Line scans through compact GFP-histone H2B domains and transcription sites. The dotted curve represents the local intensity of the nascent RNA signal, the continuous line shows the intensity distribution of the GFP-histone H2B signal. The position of the line scan is indicated by a horizontal bar with a number in C (1) and F (2). Bars, 0.6 μm.*
(Kanda et al., 1998) and transcription sites. Sites of transcription were visualized by fluorescent labeling of nascent RNA (Wansink et al., 1993; Jackson et al., 1993).

Our results show that transcription sites are present throughout chromosome 19 (a gene-rich chromosome) and, as expected, in only one of the two X chromosome territories (most likely the Xa). The other X chromosome is devoid of transcription sites (most likely the Xi) and acts as an internal control. FISH labeling of chromosomes reveals a distinct substructure of interphase chromosome territories. Territories consist of areas that are intensely labeled, probably reflecting compact chromatin, and domains that are unlabeled, containing little or no DNA. An analysis of HeLa cells that express GFP-H2B supports this conclusion. Although no individual chromosomes can be distinguished, a similar pattern of intensely labeled domains and areas with little or no GFP signal is observed. These results show that chromosome territories are relatively open structures, rather than compact lumps of chromatin. Strikingly, newly synthesized and nascent RNA accumulates specifically in the interchromatin domains, indicating a distinct compartmentalization of transcriptionally active and inactive chromatin within territories. Each of these conclusions is discussed in detail below.

**Structural Preservation during In Situ Hybridization**

The relatively harsh FISH procedure may alter the structure of the nucleus and of chromosomes, even after formaldehyde fixation. Therefore, we have thoroughly analyzed the degree of structural preservation during the chromosome painting procedure. Recently, the spatial distribution of acetylated histones in interphase nuclei has been used as a sensitive marker to visualize possible alterations in chromatin structures that are induced by FISH labeling at the light microscopy resolution level (Hendzel and Bazett-Jones, 1997). We show that our FISH procedure does not alter the spatial distribution of acetylated histone H4 in fibroblast nuclei. The same is true for the distribution of DNA, stained with Sytox green. These results indicate that the FISH protocol has no major effect on the structure of interphase chromosome territories at the light microscopy resolution level.

This is in agreement with observations of Robinett et al. (1996), who demonstrated that the interphase chromosome structure is well-preserved after in situ hybridization in formaldehyde-fixed cells. In further support of this conclusion, we found that the chromosome painting procedure does not affect the spatial distribution of transcription sites. In addition, the 3-D distributions of centromeres and of PML bodies, compared in one and the same nucleus before and after carrying out the FISH protocol, was hardly affected by the FISH procedure. This is in agreement with other studies demonstrating that the distribution of other subnuclear structures, i.e., nuclear speckles and kinetochores, are not affected by the FISH procedure (Zirbel et al., 1993; Kurz et al., 1996; Hendzel and Bazett-Jones, 1997). Finally, in cells expressing histone H2B-GFP, we observe that chromatin is organized in compact domains surrounded by nonchromatin material, very similar to the substructure of chromosome territories observed after FISH labeling. We conclude that the chromosome painting procedure does not significantly alter the structure of the nucleus and of interphase chromosomes at the light microscopical resolution level.

**Chromosomal Subdomains**

Our results show that chromosome territories have a well-defined substructure and consist of distinct subchromosomal chromatid domains. In optical sections, chromosomes appear as clusters of such domains with a diameter in the range of 300-450 nm. Domains are sometimes interconnected, forming thread-like, folded structures. Consistently, a similar substructure is observed in nuclei after DNA staining with Sytox green. Similar types of spots and threads have been shown by Brakenhoff et al. (1985) in optical sections of mithramycin-labeled mouse neuroblastoma 2A cell nuclei. Chromosomal substructure is not only found in territories of chromosome 19 and Xa, but also in Xa territories, indicating that the presence of chromosomal subdomains does not depend on transcriptional activity.

It may be argued that the compartmentalization in subchromosomal domains that are intensely labeled and domains that appear to contain little or no DNA is an artifact. It cannot be ruled out that the chromosome-specific whole library probe that we used gives incomplete labeling. However, the probe libraries label metaphase chromosomes homogeneously over their complete length (Carter et al., 1990; Telenius et al., 1992). Alternatively, nonlabeled areas may be rich in repetitive sequences, the labeling of which is suppressed in the FISH procedure. However, dual labeling in situ hybridization with a fluorescently labeled repetitive Cot-1 DNA probe and the X chromosome-specific probe library shows that highly repetitive sequences do not occur in the less intensely labeled domains in chromosome territories (data not shown). Finally, the presumed chromosomal substructure could be an artifact due to the 3-D image restoration protocol. However, careful comparison of optical sections before and after image restoration shows that essentially all intensity modulations in the restored image are also present in the unrestored optical section, albeit in a more blurred state. If chromatin is visualized in a different manner, i.e., in cells expressing GFP-histone H2B, a very similar distribution of strongly labeled domains and areas containing little or no chromatin is observed, compared with the substructure in chromosome territories after FISH labeling. From this, we conclude that chromosome territories have a distinct substructure and are only partially filled with compact chromatin.

In earlier studies in which chromosome painting is used, territories are described as compact objects without a well-defined substructure (Lichter et al., 1988; Cremer et al., 1993; Zirbel et al., 1993; Eils et al., 1996; Kurz et al., 1996). However, close visual inspection of published images in these publications does show distinct intensity modulations in labeled territories, supporting the idea that chromosome territories do have a substructure. Understanding higher-order levels of chromatin structures above that of nucleosomes and 30-nm wide solenoid fibers is limited (Woodcock and Horowitz, 1997; Bustamante et al., 1997). It is evident that additional levels of organization exist between the 30-nm chromatin fiber and the 700-μm wide...
dense sister chromatids of metaphase chromosomes (Manuelidis and Chen, 1990; Manuelidis, 1990, 1997; Belmont and Bruce, 1994; Woodcock and Horowitz, 1997). Higher-order structures, such as 100–130-nm chromonema fibers, have been described by Belmont and Bruce (1994) and Robinett et al. (1996), and 240-nm fibers have been observed by Manuelidis (1990). Recently, Zink et al. (1998) showed that chromosome territories in living cells have a similar substructure. A analysis of cell cycle dynamics of a heterochromatic chromosomal region in vivo reveals that various levels of large-scale chromatin organization can be visualized by light microscopy and EM (Li et al., 1998). EM studies have shown that there are two major, intermingled compartments in the nucleus, one containing compact chromatin, whereas the other is mainly filled with proteins and RNA (Belmont and Bruce, 1994; Fakan, 1994; Woodcock and Horowitz, 1995, 1997). Our observations are in line with these studies, showing chromatin domains with comparable dimension. It is tempting to speculate that the tightly packed, irregularly shaped substructures represent the chromosome fiber that is folded in a highly convoluted way inside the territory.

**Functional Compartmentalization of Chromosome Territories**

We find that with the use of FISH labeling, newly synthesized nascent RNA accumulates in the interchromatin space inside and around chromosome territories. The same result was obtained using cells expressing GFP-tagged histone H2B. This indicates that the observed distribution is not due to labeling artifacts. It may be argued that antibodies recognizing BrU-containing RNA cannot penetrate into the chromosomal subdomains and, therefore, bind only to peripheral RNA. However, this is unlikely since, for instance, anti-DNA antibodies readily label DNA in interphase nuclei (Hendzel et al., 1998) and antihistone antibodies can be used to visualize chromatin in situ (Turner and Fellows, 1989; Turner et al., 1989). Our results indicate that transcription sites are located predominantly at the surface of chromosomal subdomains representing compact chromatin.

These results are in agreement with those of A branches et al. (1998), who showed that transcription occurs throughout wheat nuclei chromosomes in rapidly dividing root cells. Earlier studies, using tritiated RNA precursors and high resolution autoradiography, have shown that perichromatin fibrils, the ultrastructural in situ form of nascent transcripts, are located in the interchromatin space near the surface of compact chromatin domains (Fakan and Puvion, 1980; Fakan, 1994; Puvion and Puvion-Dutilleul, 1996). This has been confirmed in more recent studies, showing that BrU-immunogold-labeled nascent RNA coincides with perichromatin fibrils (Wansink et al., 1996, CM arco et al., 1999). Nascent or newly synthesized RNA is not found inside compact chromatin domains, confirming that transcriptionally active chromatin is exclusively located at or near the surface of compact chromatin domains. Our interpretation is in line with the model for the haploid interphase chromatid structure proposed by Manuelidis et al. (1990), stating that transcriptional active chromatin occurs on locally decondensed chromatin fibers that extend from a compact chromatin fiber.

It has been postulated that the interphase nucleus contains an interchromosomal domain (ICD) space, i.e., the space between the chromosome territories (Cremer et al., 1993; Zirbel et al., 1993). The ICD space is thought to be an interconnected reticular nuclear compartment in which transcription, RNA processing, and RNA transport take place. Our results extend this model by showing that the interchromosomal compartment between chromosome territories is continuous with the interchromatin space inside the territories. In fact, there is no sharp distinction between the surface of a territory and that of compact subchromosomal domains. This reconciles the at-first-sight conflicting results of Kurz et al. (1996) and our observations. They showed that coding DNA is found preferentially at the periphery of chromosome territories, whereas a noncoding genomic locus was found predominantly in the interior of the territory.

**Emerging Model**

The following picture is emerging. The chromosome fiber, which may be compacted or locally unfolded to some degree, follows an irregular, convoluted path inside a chromosome territory, very much as suggested by Belmont et al. (1989), Belmont and Bruce (1994), and Robinett et al. (1996). Large-scale chromatin folding is strictly organized...
in such a way that transcriptionally active DNA is at the surface of the chromosomal fiber (Fig. 6). It is not known whether this compartmentalization of active chromatin is static or dynamic, i.e., whether all actively transcribed genes and those poised for transcription are in the active compartment, or whether a gene, if activated, is moved from the interior of a subchromosomal domain to its surface. Aiso, whether transcriptionally active loci loop out into the interchromatin space remains to be established. If looping out occurs, it will be only locally and over relatively short distances, since perichromatin fibrils are localized close to the surface of compact chromatin domains. This compartmentalization of active and inactive chromatin allows direct deposition of newly synthesized RNA into the interchromatin space, which contains the molecular machineries for packaging, processing, and transport of RNA. Since the interchromatin space inside territories and the interchromosomal domain between chromosome territories is continuous, such nuclear organization would allow transport of RNA molecules directly to the nuclear pores or to other parts of the nucleus.

This model leads to a number of important questions: what is the molecular basis for the strict compartmentalization of active and inactive chromatin; do proteins that are involved in, e.g., epigenetic silencing and activation, induce large scale remodeling of chromatin in terms of this compartmentalization; does constitutive heterochromatin play a structural role in this higher-order chromatin organization; or is transcriptional activity itself responsible for the observed compartmentalization? Presently, we are using EM approaches to obtain insight in this matter.

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