Nuclear Organization of Mammalian Genomes: Polar Chromosome Territories Build Up Functionally Distinct Higher Order Compartments

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Abstract. We investigated the nuclear higher order compartmentalization of chromatin according to its replication timing (Ferreira et al., 1997) and the relations of this compartmentalization to chromosome structure and the spatial organization of transcription. Our aim was to provide a comprehensive and integrated view on the relations between chromosome structure and functional nuclear architecture. Using different mammalian cell types, we show that distinct higher order compartments whose DNA displays a specific replication timing are stably maintained during all interphase stages. The organizational principle is clonally inherited. We directly demonstrate the presence of polar chromosome territories that align to build up higher order compartments, as previously suggested (Ferreira et al., 1997). Polar chromosome territories display a specific orientation of early and late replicating subregions that correspond to R- or G/C-bands of mitotic chromosomes. Higher order compartments containing G/C-bands replicating during the second half of the S phase display no transcriptional activity detectable by BrUTP pulse labeling and show no evidence of transcriptional competence. Transcriptionally competent and active chromatin is confined to a coherent compartment within the nuclear interior that comprises early replicating R-band sequences. As a whole, the data provide an integrated view on chromosome structure, nuclear higher order compartmentalization, and their relation to the spatial organization of functional nuclear processes.

Key words: nuclear architecture • genome organization • chromosome structure • nuclear compartments • polar chromosome territories

Since it became evident that cell nuclei are compartmentalized organelles (for review see Spector, 1993; Cremer et al., 1995; van Driel et al., 1995; Strouboulis and Wolffe, 1996; Singer and Green, 1997; Lamond and Earnshaw, 1998), one of the major goals in cell biology has been to understand their functional organization. In particular, the functional organization of genomes within cell nuclei is a topic of long lasting and controversial discussions (Blobel, 1985; Hutchinson and Weintraub, 1985; Hochstrasser and Sedat, 1987; Cremer et al., 1988, 1995; Manuelidis, 1990; Belmont and Bruce, 1994; De Boni, 1994; Berezney et al., 1995; Kurz et al., 1996; Marshall et al., 1997a; Bridger and Bickmore, 1998). Although it has become evident that cell nuclei are compartmentalized organelles (for review see Spector, 1993; Cremer et al., 1995; van Driel et al., 1995; Strouboulis and Wolffe, 1996; Singer and Green, 1997; Lamond and Earnshaw, 1998), one of the major goals in cell biology has been to understand their functional organization. In particular, the functional organization of genomes within cell nuclei is a topic of long lasting and controversial discussions (Blobel, 1985; Hutchinson and Weintraub, 1985; Hochstrasser and Sedat, 1987; Cremer et al., 1988, 1995; Manuelidis, 1990; Belmont and Bruce, 1994; De Boni, 1994; Berezney et al., 1995; Kurz et al., 1996; Marshall et al., 1997a; Bridger and Bickmore, 1998). Although it has become evident that cell nuclei are compartmentalized within the level of whole chromosome territories in animal as well as in plant nuclei (Manuelidis, 1985; Schardin et al., 1985; Cremer et al., 1988; Lichter et al., 1988; Leitch et al., 1990), it was difficult to understand the internal structure of chromosome territories and the relation of their organization to presumptive higher order functional compartments within nuclei (for review see Cremer et al., 1995; van Driel et al., 1995; Bridger and Bickmore, 1998).

Studies on mitotic chromosomes have indicated a functionally significant compartmentalization of mammalian genomes, strongly related to the well known banding patterns and their specific isochore compositions. A correlation of isochores (for review see Bernardi, 1995) from GC poor and GC rich families with G- and R-bands (Bernardi et al., 1985, 1989) was demonstrated (Saccone et al., 1993; Federico et al., 1998). Since GC poor and GC rich isochores have low and high gene concentrations, respectively (Cuny et al., 1981; Saccone et al., 1996; Federico et al., 1998), these results also gave an estimate of gene distribution on chromosomal bands.

Two further pieces of evidence suggest that the classical-
banded structure observed only on mitotic chromosomes is relevant for genome organization with regard to important nuclear functions like transcription and replication. First, constitutively expressed, housekeeping genes reside almost exclusively within the R-bands, whereas G-bands harbor predominantly tissue-specific genes (Craig and Bickmore, 1993, 1994). Second, R-band DNA replicates first during S phase, whereas G-band DNA replicates thereafter, during the second half of S phase (Kim et al., 1975; Dutrillaux et al., 1976; Camargo and Cervenka, 1982). Recent studies indicate that the structures of mitotic chromosomes and nuclear chromosome territories are closely related and that the different bands of mitotic chromosomes are present as distinct domains regarded as subchromosomal foci (SF) within chromosome territories (Zink et al., 1998, 1999).

Moreover, there are several lines of evidence indicating, on the DNA level, a close relationship between SF and replication foci (RF) observed during S phase (Meng and Berezney, 1991; Sparvoll et al., 1994; Berezney, 1995; Jackson and Pombo, 1998; Ma et al., 1998; Zink et al., 1998, 1999). RF contain the actively replicated DNA, the nascent DNA, and all factors necessary for replication (Leonhardt and Cardoso, 1995). RF are organized into higher order patterns within the nucleus, and studies on fixed cells indicate that there are characteristic patterns for different stages of S phase (Nakayasu and Berezney, 1989; O’Keefe et al., 1992; Ferreira et al., 1997). Recent data indicate that higher order nuclear compartments, comprising DNA with a specific replication timing revealed by specific nuclear patterns of foci, are not established during S phase but directly after mitosis at late telophase/early G1 (Ferreira et al., 1997). However, the functional significance of these higher order compartments and their stability during interphase was not clear.

As a whole, the data described above suggest a close relationship between chromosome structure during mitosis and interphase, functional nuclear processes like replication and transcription, and nuclear higher order compartmentalization. However, because of the lack of direct evidence for these relations it was difficult to obtain an integrated view of mammalian genome functional architecture. Therefore, it was important to study the different parameters all in relation to one another. Our aim was to enable for an integration by relating chromosome structure during mitosis and interphase to nuclear higher order compartments characterized by a specific replication timing and to the spatial organization of transcription. We confirm that mammalian genomes are organized into higher order nuclear compartments that harbor DNA sequences with a specific replication timing (Ferreira et al., 1997). We demonstrate that higher order compartments established immediately after mitosis are stably maintained during all interphase stages. The organizational principle is clonally inherited. We provide direct evidence that higher order compartments are built up by the alignment of polar chromosome territories, as suggested by Ferreira et al. (1997). Polar chromosome territories display clusters of early or late replicating DNA that correspond to R- and G/C-bands of mitotic chromosomes, respectively, at distinct subterritorial positions. The R-band sequences of distinct chromosomes cluster within the nuclear interior to give rise to an early replicating compartment that is transcriptionally competent and active. The G/C-band sequences organize transcriptionally incompetent and inactive late replicating compartments in the perinuclear and perinucleolar regions.

Materials and Methods

Cell Culture

Cells were grown at 37°C in an atmosphere of 5% CO₂ in RPMI supplemented with 10% FCS (HeLa S6 cells [provided by W.W. Franke, DKFZ, Heidelberg], female human diploid fibroblasts [H v provided by Professor J. M. urken, LMU, Munich], SH-EP-N14 human neuroblastoma cells [derivative of SH-EP-cells [Ross et al., 1983], stably transfected with an N-Myc expression vector [Wenzel et al., 1991]) and CHO cells (provided by S. Müller, LMU, Munich). Primary human lymphocytes and C2C12 mouse myoblasts (provided by M.C. Cardoso, MDC, Berlin) were cultured in the presence of 20% FCS. The media were supplemented with antibiotics (100 µg/ml penicillin and 100 µg/ml streptomycin). 4 µg/ml phytohemagglutinin was added to cultures of primary lymphocytes. For synchronization in early S phase, 200 µM mimosin was added to the culture medium for 14–16 h. The block was released by adding fresh medium after washing the cells with PBS. For microinjection and microcopy of fixed cells, cells were grown on coverslips and fixed after replication labeling/nascent RNA labeling in PBS/3.7% formaldehyde for 10 min. Fixed cells were stored in PBS at 4°C and always kept wet during all of the following immunostaining/situ hybridization procedures. For following the clonal inheritance of replication labeling patterns in living cells, the cells were cultured, microinjected, and imaged in cell culture dishes with a grid-ded coverslip (cellocate, Eppendorf-Netheler-Hinz GmbH) inserted into the bottom.

Replication Labeling

Replication labeling was performed with exponentially growing cultures or cultures synchronized in early S phase. Replication labeling involving a single BrdU (bromodeoxyuridine) or Cy3-dUTP pulse (Cy3-dUTP was microinjected) or iododeoxyuridine (IdU) (idothymidyridine; IdU) double pulse labeling was performed as described in (Zink et al., 1998) according to the time schedules described in the results. FITC-dUTP (Boehringer) was microinjected at a concentration of 100 µM diluted in CMF-PBS (PBS without Ca²⁺ and Mg²⁺). If synchronized cultures were used, the block was released either after microinjection or before BrdU or IdU was supplemented to the medium to obtain early S phase patterns. To obtain later S phase patterns, the cells were replication-labeled 2–9 h after release. Cells were fixed at the time points indicated in the results either during or after S phase. If necessary, detection of incorporated nucleotides was performed as described in (Zink et al., 1998, 1999) (in the present study BrdU was only detected with a mouse anti-BrdU antibody and an anti-mouse TRITC-conjugated secondary antibody, see BrdU detection). No detection procedures were necessary for visualizing incorporated fluorescent nucleotides (Cy3-dUTP, FITC-dUTP).

Replication Labeling in Combination with Nascent RNA Labeling

Synchronized or unsynchronized cells from cultures of HeLa S6 or CHO cells were replication-labeled by microinjection of FITC-dUTP and grown after replication labeling for 14–19 h. Subsequently, replication-labeled cells were microinjected with 5-bromouridine-S-triphosphate (BrUTP, Sigma Chemical Co., 50 µg/ml in CMF-PBS). A 1 h microinjection, cells were fixed. For immunodetection of BrUTP, preparations were blocked for 30 min in PBS, 0.3% Triton X-100, 0.2% Tween 20, and 3% BSA, and subsequently incubated with a monoclonal anti-BrdU antibody (Becton and Dickinson, recognizes also BrUTP) diluted in blocking solution for 1 h. After washing the preparations three times for 5 min with...
Replication Labeling in Combination with Immunostaining Against Hyperacetylated Histone H4

Cells were replication-labeled either with Cy3-dUTP or BrdU and fixed. Fixed preparations were permeabilized and blocked in blocking solution (PBS, 0.2% Triton X-100, 0.2% TWEEN 20, 5% BSA) at room temperature for 1 h. Subsequently, cells were incubated with rabbit antiserum R 232/8 diluted 1:1,500 in blocking solution for 1 h at room temperature. R 232/8 is a high titer rabbit antiserum that is specific for histone H4 acetylated at lysine 8. It recognizes more highly acetylated H4 isoforms (mainly di- and triacetylated isoforms). Cells were washed three times for 10 min with PBS and 0.2% Triton X-100. A further washes, cells were incubated with an FITC-conjugated goat anti-rabbit antibody (Dianova) diluted in blocking solution. Cells were washed three times for 10 min with PBS and mounted for microscopy if they were replication-labeled with Cy3-dUTP. In the case when cells were replication-labeled with BrdU, they were postfixed with PBS, 3.7% formaldehyde for 10 min. A further fixation, DNA was denatured for BrdU detection by incubating the preparations for 10 min in 2 M HCl (for BrdU detection procedure see above).

Preparation of the H3 Isochore Probe for In Situ Hybridization

DNA from an H3 isochore fraction of human DNA (equivalent to one previously described, Saccone et al., 1996) was amplified and labeled by DOP-PCR (Telenius et al., 1992) with biotin-16-dUTP (Boehringer). Fragment length was checked by gel electrophoresis after DNA digestion (3 μg/ml for ~30 min at 15°C). 200 ng (for metaphase spreads) or 400 ng (for interphase nuclei) of labeled DNA was precipitated with 30 μl Cot-1 DNA (1 mg/ml; GIBCO BRL) and 5 μl of salmon testis DNA (12 mg/ml; GIBCO BRL). The dry pellet was dissolved in 10μl hybridization solution (50% formamide, 10% dextran sulfate, 1× SSC). 200 (for metaphase spreads) or 400 ng Cot-1 DNA (1 mg/ml; GIBCO BRL) and 5 μl of salmon testis DNA (12 mg/ml; GIBCO BRL). The dry pellet was dissolved in 10μl hybridization solution (50% formamide, 10% dextran sulfate, 1× SSC). Before hybridization, the probe DNA was denatured in hybridization solution for 6 min at 75°C and preannealed for 20 min at 37°C.

In Situ Hybridization and Probe Detection

Metaphase spreads of SH-EP-N14 cells and primary lymphocytes were prepared according to standard protocols. Immunostaining against hyperacetylated Histone H4 was performed as described in Eils et al. (1996).

Results

Higher Order Compartments Comprising DNA with a Defined Replication Timing Are Present during All Interphase Stages

Previous studies suggested a nuclear higher order compartmentalization of mammalian genomes according to the replication timing of DNA sequences (Ferreira et al., 1997). The nuclear compartments appear to be established at late telophase/early G1 (Ferreira et al., 1997). However, the stability of nuclear compartments established after mitosis during subsequent interphase stages and the stability of their inheritance was not clear. We were interested in the question of whether compartments comprising DNA sequences with a similar replication timing are indeed reproducibly established after mitosis and stably maintained during all subsequent interphase stages. In this case, a specific S phase pattern produced by DNA pulse labeling during replication should be observed at all other interphase stages and in all daughter cell nuclei.

To prove this prediction, we pulse-labeled DNA of different cell lines and primary cells derived from man, mouse, and hamster during the S phase. Studies were performed with the following cell types: primary human diploid fibroblasts (HDFs), Hela S6 cells, human neuroblastoma cells (SH-EP N14), CHO cells, and C2C12 mouse myoblasts. Pulse labeling was performed with BrdU or Cy3-dUTP. Cells were fixed 30 min after supplementing with the modified nucleotides to obtain typical S phase patterns (Fig. 1, left, a, c, e, g, and i).

Regarding S phase patterns, we mainly followed the classification of O’Keeffe et al. (1992) and defined five different types of pattern reflecting an ordered time sequence during S phase progression (type I beginning and type V end of S phase): type I (Fig. 1 a) displays hundreds of small foci (~300 nm in diameter) distributed throughout the DNA within the nuclear interior. DNA sequences located at the nuclear and nucleolar peripheries and minor accumulations of late replicating chromatin within the nuclear interior (see type IV and V patterns) are not labeled. Nucleoli are excluded in all types of patterns. The nuclear space occupied by the type I pattern will be regarded as the interior compartment in the following text.

The type II pattern (Fig. 1 c) shows some labeling of DNA located at nuclear and nucleolar peripheries. The interior compartment is still partially labeled, but fewer foci are observed here compared with the type I pattern and unlabeled regions appear within the interior compartment. The type III pattern (Fig. 1 e) displays heavy labeling of nuclear and nucleolar peripheries (excluded from the type I pattern), whereas the interior compartment is almost devoid of label. The compartments labeled by the type III pattern will be regarded as the peripheral compartments.

Type IV and V (Fig. 1, g and i) patterns are characterized by a few large (~800 nm in diameter) foci that are located within the nuclear interior as well as in the peripheral compartments. Type V displays less labeling of the peripheral compartments than type IV. The totality of late replicating chromatin accumulations within a nucleus labeled by type IV and V patterns will be regarded as the...
late replicating compartments in the following text. All cell lines examined displayed type I–V patterns.

These results confirmed for all cell types used the finding that DNA sequences with a defined replication timing occupy during S phase specific nuclear areas (Nakayasu and Berezney, 1989; O’Keefe et al., 1992). This way, higher order nuclear compartments are established, comprising DNA with a similar replication timing (e.g., the interior compartment comprises the early replicating DNA sequences). To investigate the stability of these compart-

Figure 1. S phase replication labeling patterns are preserved. Cells ([a and b] H\textsubscript{v}, human diploid fibroblasts; [c and d] SH-EP N14 human neuroblastoma cells; [e and f] HeLa cells; [g and h] CH\textsubscript{O} cells; and [i and j] C2C12 mouse myoblasts) were replication-labeled with BrdU (a–d, i, and j; 30-min pulses) or Cy3-dUTP (e–h). Cells were fixed immediately after BrdU labeling or 30 min after microinjection of Cy3-dUTP to obtain labeled S phase cells (left, a, c, e, g, and i). S phase cells display the typical replication labeling patterns (indicated on the left, for classification see text): (a) type I, (c) type II, (e) type III, (g) type IV, and (i) type V. Similarly labeled cells were grown for 1–5 d after labeling and fixed after this growth period (right, b, d, f, h, and j). The presence of similar patterns (types indicated on the right) several days after S phase labeling suggests that DNA replicating with a particular timing during S phase always occupies similar nuclear positions. Arrowheads in a mark the nucleoli that are always unlabeled. Arrowheads in c–e mark perinucleolar label. Arrows in b and d indicate unlabeled chromosome territories. The presence of unlabeled territories demonstrates that cells do not display similar patterns because they stopped cycling, but rather that they went through at least two mitoses after labeling (compare Fig. 3). Note also the two very close nuclei in f displaying similar type III patterns, suggesting that similar patterns were conserved in sister nuclei after cell division (compare Fig. 4, a–c). All pictures display single light optical sections through midnuclear planes except i and j, which show epifluorescence images. Bar in a is similar for all images.
ments during cell growth and distinct cell cycle stages, we pulse-labeled cells during the S phase as described above. In contrast to the previous experiment, cells were not fixed immediately but after a growth period of 1–5 d. The labeling patterns after this growth period were compared with typical S phase patterns (Fig. 1).

The patterns we observed 1–5 d after labeling (Fig. 1, right, b, d, f, h, and j) were similar to typical S phase patterns (Fig. 1). The distribution of numbers and sizes of foci was the same as that seen in cells fixed during S phase. 200 cells were examined for each cell type and no labeling patterns were observed that could not be classified according to the five types of patterns outlined above. Therefore, we extended the classification of the five types of patterns also to non-S phase cells.

The only difference to typical S phase patterns observed in cells grown for 1–5 d was the appearance of unlabeled nuclear areas corresponding to unlabeled chromosome territories (Fig. 1, b and d). The appearance of unlabeled chromosome territories is due to the random segregation of labeled and unlabeled chromatids beginning at the second mitosis after replication labeling (Taylor, 1984; Zink et al., 1998) (see also Fig. 3). This phenomenon confirmed that the cells went through at least two mitoses after labeling, and excluded the possibility that cells maintained the patterns because they stopped cycling. As the patterns were often similar in neighboring cells (Fig. 1 f) the data suggested that nuclear compartments comprising DNA with a similar replication timing were clonally inherited.

To exclude the possibility that cells displayed labeling patterns reminiscent of S phase patterns because of the fact that cells entered S phase again, exponentially growing CHO cells were pulse-labeled with Cy3-dUTP to obtain typical replication labeling patterns. Cy3-labeled cells were grown for 19 h, pulse-labeled with BrdU for 30 min, and fixed immediately (see scheme depicted in Fig. 2). BrdU labeling revealed whether cells displaying typical type I–V Cy3-patterns had entered S phase again. As the cultures grew exponentially, some daughter cells of Cy3-dUTP replication-labeled cells were in S phase at the time point of fixation (Fig. 2 c), whereas others were not. Nevertheless, all cells displayed the typical type I–V Cy3 labeling patterns (60 cells examined for each cell type), indicating that the corresponding nuclear compartmentalization is present during all interphase stages. Similar results were obtained with HeLa S6 cells.

**Double labeling scheme and results for CHO cells**

![Double labeling scheme and results for CHO cells](image)

**Figure 2.** Genome compartmentalization is similar during all interphase stages. The applied double labeling procedure is schematically drawn at the top. Typical replication labeling patterns were obtained by incorporation of Cy3-dUTP into the DNA of the S phase cells of exponentially growing CHO cultures. 19 h after Cy3-dUTP microinjection, cultures were replication-labeled with BrdU and fixed after 30 min. Cells in S phase at the time point of fixation (BrdU-labeled) can be distinguished from G1 and G2 cells (not BrdU-labeled). The DAPI staining is shown in a (a-c, same field of cells is imaged). Mitotic stages of exponentially growing cultures are indicated by arrowheads. Cy3-labeled cells are depicted in b. Note the two pairs of cells with similar labeling intensities and labeling patterns (upper pair, type I; lower pair, type II; blurred label is due to epifluorescent imaging). Cells of one pair are likely sister cells (compare Fig. 4). The right cells of each pair were in S phase at the time point of fixation (arrows) as indicated by the BrdU label depicted in c. Cy3 labeling patterns are similar in S phase and non-S phase cells.
Nuclear Genome Compartmentalization Is Stably Inherited through Mitoses

To confirm clonal inheritance of nuclear genome compartmentalization, single cells from HeLa S6 cultures exponentially growing on gridded coverslips were replication-labeled with Cy3-dUTP. Cells were not fixed but imaged each day to follow the inheritance of the initial labeling patterns in developing clones. Unfortunately, the motility of the cells usually did not allow the unequivocal identification of cells belonging to one clone (although usually neighboring cells and fields of cells were observed displaying similar patterns). However, this was possible in the case depicted in Fig. 3 where, indeed, all daughter and granddaughter cells displayed the initial type II labeling pattern.

To further confirm the inheritance of patterns HeLa S6 cells were synchronized with mimosin in early S phase. Immediately after release of the mimosin, block cells were pulse-labeled for 0.5 h with IdU, chased for 9.5 h, and pulse-labeled for 0.5 h with CldU (Fig. 4, labeling scheme).
Figure 4. Establishment of higher order nuclear compartments after mitosis and the underlying chromosomal structure. HeLa S6 cells synchronized in early S phase were labeled for 30 min with IdU and 9.5 h later for 30 min with CldU (see labeling scheme at the top). Therefore, initially labeled cells simultaneously displayed a type I pattern labeled by IdU and a pattern typical for the second half of S phase labeled by CldU. Daughter cells of the initially labeled cells were fixed 14 (a–c) or 69 h (d–f), respectively, after the CldU pulse and analyzed by confocal microscopy. Identical nuclear planes were imaged regarding TRITC (IdU detection, depicted in red) and FITC (CldU detection, depicted in green) fluorescence. The corresponding merged TRITC and FITC signals (colocalizing signals appear yellow) are shown in a–d. e and f display only the TRITC (e) or FITC (f) signals of the merged image in d (midnuclear plane). The two cells depicted in a–c display the typical morphology of early G1 cells shortly after mitosis. Both G1 cells display an IdU (red) type I pattern and a CldU (green) type III pattern present in their mother cell. Perinucleolar labeling is indicated by an arrowhead in c. After 69 h (d–f), initially labeled cells went through at least two mitoses as indicated by the presence of single-labeled chromosome territories (double-labeled patches within the nucleus depicted in d). Double-labeled chromosome territories reveal that the nuclear type I (IdU, red) and type III patterns are due to a reproducibly distinct distribution of IdU- and CldU-labeled DNA within single chromosome territories. CldU-labeled DNA is concentrated at subchromosomal positions near the nuclear and nucleolar peripheries, whereas IdU-labeled DNA is located at subchromosomal positions between these compartments.
We previously established (Fauth, 1998) that 93% of cells (n = 1,000) displayed a type I pattern if they were labeled within the first hour after release. 50% of cells (n = 1,000) displayed a type III pattern and 18% a type IV or V pattern if they were labeled 10 h after release. Therefore, most cells after IdU/CldU double labeling according to the scheme outlined above, display a type I pattern (labeled with IdU) as well as type III-V pattern (labeled with CldU).

Cells were fixed 14 or 69 h after the CldU pulse, respectively (Fig. 4, labeling scheme). Many cells fixed 14 h after CldU labeling went through mitosis. After cell division, daughter cells (39 cells examined) displayed the same nuclear compartmentalization as their mother cells after S phase labeling as revealed by the double labeling patterns. For example, cells with the typical morphology of early G1 cells displayed a type I IdU pattern in combination with a CldU type III pattern (Fig. 4, a-c). These data show that nuclear compartments comprising DNA sequences with a similar replication timing are immediately established after mitosis according to the pattern present in the previous interphase.

**Polar Chromosome Territories Build Up Higher Order Nuclear Compartments**

Similar compartmentalization patterns (IdU type I patterns with CldU type III-V patterns) were also present after at least two mitoses (indicated by the presence of unlabeled chromosome territories) 69 h after initial IdU/CldU labeling. Fig. 4, d-f, shows a typical example of the 46 nuclei examined. A single replication-labeled chromosome territories can be observed the data also indicate how single chromosome territories, composed of DNA replicating at distinct time points during S phase, contribute to higher order nuclear compartments. Single territories display a polar organization with DNA replicating early during S phase (IdU-labeled) clustered at subterritorial positions located within the nuclear interior and DNA replicating at later S phase stages (CldU-labeled) clustered at subterritorial positions located at nuclear or nucleolar peripheries. A alignment of polar territories gives rise to higher order nuclear compartments.

**Functional Nuclear Compartmentalization of Mammalian Genomes Is Related to the Organization Of Mitotic Chromosomes**

R-bands of mitotic chromosomes comprise DNA replicating early during S phase and G- or C-bands harbor DNA replicating at later S phase stages (Kim et al., 1975; Dutil-lox et al., 1976; Camargo and Cervenka, 1982). Therefore, one would expect that DNA located within these distinct bands of mitotic chromosomes corresponds to the observed early and late replicating DNA at different subterritorial positions and contributes distinctly to the different higher order compartments. It has been previously shown that clustering of R- and G-band sequences at distinct subchromosomal positions leads to the formation of polar interphase chromosome territories (Zink et al., 1999).

To confirm the contribution of DNA belonging to specific chromosomal bands to distinct higher order nuclear compartments, we performed in situ hybridization in combination with Cy3-dUTP replication labeling. A probe for in situ hybridization we used the H3 isochores fraction of human DNA (Saccone et al., 1996). The H3 fraction usually hybridizes with a subset of R-bands on human mitotic chromosomes (Saccone et al., 1996). However, when we tested the specificity of hybridization on metaphase spreads of human chromosomes prepared from cultured primary lymphocytes (n = 20; Fig. 5, a and b) as well as SH-EP N14 human neuroblastoma cells (n = 20; data not shown) we obtained almost a complete R-banding pattern although different R-bands were stained with variable intensity (Fig. 5 b). Even though the probe hybridized specifically to the whole set of R-bands, which was contrary to previous results (see Discussion), the result was highly reproducible under the conditions we used.

A probe under the conditions we used was an excellent and specific marker for R-band DNA, we hybridized it to replication-labeled nuclei of HeLa S6 and SH-EP N14 cells (in this case it was not possible to include hamster or mouse cells as the probe is specific for human DNA). The hybridization signal was spread all over the interior compartment (Fig. 6, a+d), but was excluded from the peripheral compartments (Fig. 6 e) and the late replicating compartment (Fig. 6 f). The data confirm that R-band DNA builds up the interior compartment, and reveals a clear correlation between the organization of mammalian genomes during interphase and the banded organization of mitotic chromosomes (see Fig. 10).

**Hyperacetylated Isoforms of Histone H4 Are Enriched within the Interior Compartment**

A probe for R-bands of mitotic chromosomes comprise most of the genes expressed during interphase, one would expect the transcriptionally competent chromatin within the interior compartment comprising the R-band sequences. Hyperacetylated isoforms of histone H4 are a characteristic feature of transcriptionally competent chromatin (Jeppesen, 1996; Hassig and Schreiber, 1997; Utley et al., 1998). To investigate whether transcriptionally competent chromatin is indeed confined to specific nuclear compartments, we stained replication-labeled nuclei with antisera R 232/8 specifically recognizing hyperacetylated isoforms of histone H4 (see Materials and Methods).

Fig. 7 summarizes the results obtained for female HDFs (100 nuclei examined) and mouse myoblasts (70 nuclei examined). Unsynchronized, exponentially growing cultures were fixed 27 h after BrdU pulse labeling and immunostained. Comparison of BrdU labeling patterns and the nuclear distribution of hyperacetylated histone H4 isoforms revealed a clear correlation. Hyperacetylated histone H4 isoforms were detected throughout the whole interior compartment labeled by the type I pattern (Fig. 7, a-c and e), but were excluded from the peripheral and late replicating compartments (Fig. 7, f-h). Also, in female HDFs, a heterochromatic structure that presumably is the Barr body was not stained by antisera R 232/8 (Fig. 7, b and d). This is in agreement with previous data demonstrating a lack of histone H4 acetylation for mammalian inactive X chromosomes during mitosis (Jeppesen and Turner, 1993).
As the transcriptional competence of genome compartments is an important feature with regard to their functional characteristics, we extended the analysis to CHO and HeLa S6 cells. Unsynchronized, exponentially growing cultures of HeLa or CHO cells were fixed 1 or 23 h after replication labeling with Cy3-dUTP to confirm similarity of results between S phase and non-S phase cells. Cy3-dUTP labeling excludes an influence of artificial changes of chromatin structure induced by DNA denaturation (Zink, D., unpublished results) that is necessary for BrdU detection on the results.

Cy3-dUTP–labeled CHO nuclei immunostained with R232/8 antiserum were examined and examples are shown in Fig. 8, a–i. For both groups (fixation after 1 h \(n = 28\) or 23 h \(n = 30\)) we obtained similar results. There was the same correlation between nuclear compartments revealed by replication labeling and the nuclear distribution of hyperacetylated histone H4 isoforms as observed for HDFs or C2C12 cells (compare Figs. 7 and 8). Hyperacetylated isoforms of histone H4 are confined to the interior compartment labeled by the type I pattern (Fig. 8, a–c). Type III–V patterns do not overlap with regions immunostained by antiserum R 232/8 (Fig. 8, d–i). Similar results were also obtained with HeLa S6 cells, fixed after 1 h \(n = 20\) or 23 h \(n = 20\) and double-labeled with Cy3-dUTP and antiserum R 232/8 (Fig. 8, j–l).

**Transcriptional Activity Is Compartmentalized within Mammalian Interphase Nuclei**

The results suggested that not only transcriptional competence, but also that the process of transcription itself also might be subject of higher order nuclear compartmentalization. Nuclear higher order compartments were visual-
ized within HeLa S6 and CHO nuclei by replication labeling with FITC-dUTP (synchronized and unsynchronized cultures used). On the next day, FITC-labeled cells were microinjected with BrUTP (incorporated into nascent RNA; Wansink et al., 1993) and fixed after 10 min of microinjection. Immunostaining of incorporated BrUTP revealed only the interior compartment as the transcriptionally active compartment. (Fig. 9, a–c). No considerable BrUTP labeling was visible within the peripheral compartments (Fig. 9, d–f). There was also no overlap between type IV
and V FITC patterns and BrUTP incorporation. Similar results were obtained for HeLa and CHO cells. Therefore, the interior compartment comprising the early replicating R-band sequences harbors the transcriptionally competent as well as the actively transcribed chromatin. The data are summarized in Fig. 10.

**Discussion**

In this study, we characterized the functional compartmentalization of mammalian genomes during interphase. The data demonstrated that a specific pattern of spatial genome compartmentalization was present during all interphase stages and was clonally inherited. Distinct
compartments comprised DNA sequences belonging to distinct chromosomal bands during mitosis. R-band sequences built up a coherent compartment within the nuclear interior, whereas G/C-band sequences localized to compartments at the nuclear and nucleolar peripheries as well as to minor internal compartments. The early replicating compartment comprising the R-band sequences harbored the transcriptionally competent chromatin. Detectable nascent RNA synthesis was confined to the interior compartment. Fig. 10 summarizes the correlations between functional higher order nuclear genome architecture and chromosome organization during mitosis and interphase.

The functionally different higher order compartments were built up through the alignment of polar chromosome territories with clusters of early replicating DNA directed towards the nuclear interior and clusters of later replicating DNA located at the nuclear or nucleolar peripheries. We demonstrated recently that early replicating R- or later replicating G/C-bands of mitotic chromosomes are retained as distinct chromosomal domains termed subchromosomal foci (SF) within interphase chromosome territories (Zink et al., 1998, 1999). Within chromosome territories of cycling (G1) cells, we observed a clustering of R-SF or G/C-SF at distinct sites of the territories (Bornfleth et al., 1999; Zink et al., 1999). Therefore, the emerging picture is as follows: distinct bands (in the megabase pair size range) alternating on mitotic chromosomes are retained as distinct domains during interphase (SF), but are reorganized within the territories. The distinct SF cluster within a territory and, therefore, give rise to a polar organization of this structure in the size range of several tens to hundreds megabase pairs (Morton, 1991). A alignment of these polar territories creates higher order functional genome compartments (size range: gigabase pairs) within mammalian cell nuclei (see also Ferreira et al., 1997; La- mond and Earnshaw, 1998).

The results and conclusions of the present study are in agreement with the present understanding of chromosome organization on the one hand, and a growing body of evi-
Figure 9. Transcriptional activity is confined to the interior compartment. Transcriptional activity within distinct genome compartments was investigated by BrUTP pulse labeling (10 min) of cells replication-labeled on the previous day. BrUTP was detected with TRITC fluorescence, whereas replication labeling was performed with FITC-dUTP. HeLa cells (a–c) and CHO cells (d–f) were investigated. Identical nuclear planes were imaged with regard to FITC detection (b and e, depicted in green) and TRITC detection (c and f, depicted in red). Merges of corresponding FITC and TRITC signals are depicted in a and d (colocalizing signals appear yellow). Transcriptional activity, indicated by BrUTP incorporation into nascent RNA (c and f, and, a and d, red) is confined to the interior compartment (a and b, green, colocalizing with the type I pattern). The TRITC signal in the upper left regions of a and c is due to an adjacent BrUTP-labeled nucleus that is not replication-labeled. d–f depict the exclusion of RNA synthesis from the peripheral compartments (d and e, green). Arrows in f show examples of perinuclear and perinucleolar regions strongly labeled by the type III replication labeling pattern (d and e, green) that display no TRITC signal (no RNA synthesis).

However, the in situ hybridization data do not rule out that these sequences also contribute to other compartments. We present cannot definitely exclude that a lack of signal in unlabeled compartments is due to accessibility problems. However, we think this is unlikely as we characterized R-band chromatin not only by sequence identity, but also by its replication timing and its content of hyperacetylated histone H4. Early replicating sequences do not obviously contribute to peripheral and late replicating compartments that are also not enriched in hyperacetylated histone H4. As both features are characteristic for R-band chromatin (Camargo and Cervenka, 1982; Jeppe sen and Turner, 1993), together the data indicate that R-band chromatin does not make a major contribution to peripheral and late replicating compartments.

Although we did not use specific DNA probes to localize G- and C-band DNA within the nucleus, a variety of data indicate that the corresponding sequences localize in the peripheral and late replicating compartments. First, many studies describe the detection of C-band sequences at the corresponding nuclear positions (Rae and Franke, 1972; M anuelidis and Borden, 1988; O’Keefe et al., 1992), whereas the localization of G-band sequences at these nu-
clear positions was suggested by replication labeling studies (Ferreira et al., 1997). G/C-band sequences are known to replicate during the second half of S phase (Kim et al., 1975; Dutrillaux et al., 1976; Camargo and Cervenka, 1982; O'Keefe et al., 1992; Ferreira et al., 1997). As our results show that DNA within the interior compartment replicates early in S phase, G- or C-band sequences cannot make a major contribution to this compartment and, therefore, have to be confined to the nuclear compartments comprising the transcriptionally inactive parts of the genome. This is in agreement with the fact that G-band DNA displays a low density of genes that are mostly not expressed and the fact that C-bands comprise highly repetitive sequences (Goldman et al., 1984; Bickmore and Summer, 1989; Craig and Bickmore, 1993).

The data are in agreement with many studies indicating that expressed sequences are generally located within the nuclear interior, whereas repressed sequences locate towards the nuclear periphery often in close association with constitutive heterochromatin (C-bands of mammalian chromosomes comprise the constitutive heterochromatin) (Hoehn and M arkin, 1973; M athog et al., 1984; E lmont et al., 1986; M anuelidis and Brown, 1988; M amilos and Borden, 1988; Csink and Henikoff, 1996; M ailet et al., 1996; Brown et al., 1997; Imai et al., 1997; A ndulis et al., 1998). However, these studies were performed with a variety of eukaryotic taxa as different as yeast, flies, and mammals. Although the principle of compartmentalization with repressed sequences at perinuclear positions might be general, experimental data indicate that the type of DNA sequences and chromosomal structures located within the different nuclear compartments, as well as the mechanisms mediating genome compartmentalization...
tion, might be different in distinct taxa (R abl, 1885; H och-strasser et al., 1986; Haaf and Schmid, 1991; M anuelidis and B orden, 1988; Palladino, 1993).

With regard to the specific localization of chromatin observed in mammals, previous analyses indicated that the process of transcription might not be responsible (Ferreira et al., 1997) for nuclear genome compartmentalization in mammals. The fact that the nuclear compartmentalization of mammalian genomes is strongly related to the banding patterns of mitotic chromosomes, and that the nuclear compartmentalization is established immediately after mitosis imply that chromatin belonging to the different bands is targeted to distinct nuclear positions when the nucleus reconstitutes. This process might take place independently for each chromosome (Ferreira et al., 1997). In this case, DNA or chromatin belonging to distinct bands of a chromosome has to be specifically recognized. Recognition could take place at the level of DNA sequence as R- and G/C-bands display a distinct isochose composition (Bernardi, 1995). However, we think that unlikely as the active (Xa) and inactive (Xi) X chromosomes of female mammals display a similar DNA sequence, but are affected by the nuclear genome compartmentalization regarding transcriptional competence and activity (Xi is usually located at the nuclear periphery; Hoehn and Martin, 1973; Belmont et al., 1986). As there are specific modifications of R- or G/C-band chromatin (J epsen and Turner, 1993; H aaf, 1995) that affect also Xa and Xi (Xi is exceptional in the sense that chromatin modifications affect almost the whole chromosome rather than its distinct bands), specific recognition at this level seems possible.

Although some chromatin domains might display dynamic relocalizations according to changes in their functional states, the stable maintenance of replication labeling patterns indicates that positional changes of chromosomes or chromosomal regions might be exceptional events within the nucleus. This is in full agreement with recent studies of DNA dynamics within nuclei of living mammalian cells (R obinett et al., 1996; Shelby et al., 1996; A bney et al., 1997; M arshall et al., 1997b; Zink and C remer, 1998; Z ink et al., 1998). Nevertheless, rare large-scale movements of chromosomes or parts of them were observed in living cell studies (Shelby et al., 1996; Zink and Cremer, 1998; Z ink et al., 1998). Such movements are compatible with the observed, stable compartmentalization of functionally defined chromatin, as long as they occur within or between corresponding compartments. Whereas the intra-nuclear location of the compartments is fixed, at least within similar cell types, there may be a degree of flexibility in the positioning of individual DNA sequences within the compartments. For example, if a centromeric C-band domain moves during interphase from the nucleolar to the nuclear periphery, the overall compartmentalization is not disturbed. In this regard, the observed stable compartmentalization is also compatible with the dynamic repositioning of nuclear domains observed in fixed cell studies (for review see De Boni, 1994).

We thank Irina Solovej for advice regarding H 3 isochose probe hybridization, R oel van D riel and P ernette V erschure for advice in BrUTP labeling, and R obert M artin for help in arrangement of images.

This work was supported by grants from the Wellcome Trust (045030/Z/ 95 to B.M. Turner) and the D eutsche Forschungsgemeinschaft (Z I 560/2-1).

Submitted: 6 May 1999
Revised: 29 July 1999
Accepted: 3 August 1999

References


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