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Differences in the Localization and Morphology of Chromosomes in the Human Nucleus

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Abstract. Using fluorescence in situ hybridization we show striking differences in nuclear position, chromosome morphology, and interactions with nuclear substructure for human chromosomes 18 and 19. Human chromosome 19 is shown to adopt a more internal position in the nucleus than chromosome 18 and to be more extensively associated with the nuclear matrix. The more peripheral localization of chromosome 18 is established early in the cell cycle and is maintained thereafter. We show that the preferential localization of chromosomes 18 and 19 in the nucleus is reflected in the orientation of translocation chromosomes in the nucleus. Lastly, we show that the inhibition of transcription can have gross, but reversible, effects on chromosome architecture. Our data demonstrate that the distribution of genomic sequences between chromosomes has implications for nuclear structure and we discuss our findings in relation to a model of the human nucleus that is functionally compartmentalized.

Key words: chromosome territories • genome organization • nuclear compartmentalization • transcription • translocations

The arrangement of human chromosomes on the mitotic spindle could reflect an ordered arrangement of chromosomes at interphase (Nagele et al., 1995). Whole chromosomes occupy discrete territories in the nucleus (Cremer et al., 1993 and references therein). Nevertheless, with the exception of the inactive X (Xi) that is positioned against the nuclear membrane, and the rDNA-containing chromosomes that are intrinsic to the nucleolus, there has been little evidence that other whole mammalian chromosomes regularly adopt defined addresses within the nucleus (Manuelidis, 1990).

Here, we investigate whether the two small human chromosomes 18 and 19 (HSA18 and HSA19) adopt different dispositions within the nucleus by fluorescence in situ hybridization (FISH). There are several reasons why these two chromosomes make for interesting comparisons: HSA18 and 19 are similarly sized DNA molecules, containing 85 M b and 67 M b of DNA, and so comprising 2.6 and 2% of the physical length of the genome, respectively (Morton, 1991), yet these two chromosomes have contrasting functional and structural characteristics. HSA19 contains a high density of CpG islands (Craig and Bickmore, 1994) and repeats of the Alu family (Korenberg and Rykowski, 1988), replicates most of its DNA in the early part of S phase (Dutrillaux et al., 1976), and has abundant hyperacetylated histone H4 in its chromatin (Jeppesen and Turner, 1993). HSA19 also has the highest observed/expected ratio of gene-based marker assignments of any human autosome (Cross et al., 1997; Deloukas et al., 1998) with 3.7% of such markers assigned to it. In contrast, HSA18 has far fewer gene assignments than expected for its size (Cross et al., 1997; Deloukas et al., 1998), with only 1.7% of gene-based markers mapped to it. It also has a low CpG island density, a high concentration of the L1 family of repeats, and a high proportion of late replicating DNA (Dutrillaux et al., 1976; Korenberg and Rykowski, 1988; Craig and Bickmore, 1994). Little hyperacetylated H4 is detected on this chromosome (Jeppesen and Turner, 1993).

We compare the nuclear position and chromosome morphology of human chromosomes 18 and 19 through the cell cycle and in different cell types and visualize the association of DNA from each chromosome with substructure of the nucleus. The effects of inhibiting transcription and histone deacetylation on nuclear organization are assessed.
Materials and Methods

Cell Culture, Cell Cycle Fractionation, and Preparation of Nuclear Halos

H human primary lymphocytes were stimulated with phytohemagglutinin; both lymphocytes and lymphoblasts (46XY) were grown in RPMI + 10% FCS. Human primary fibroblasts (less than passage 10) and HT1080 cells were grown in DMEM + 10% FCS. To identify cells in 5 phase, BrdU was added to cultures at 0.1 m. To irreversibly inhibit transcription, actinomycin D (AM) was added 2 h before harvest to 5 μg/ml. To reversibly inhibit RNA pol II cells were cultured with 50 μg/ml 5,6-dichloro-β-β-rifobouranoribenzimidazole (DRB) for 5 h (Haa and Ward, 1996). Inhibition was relieved by culturing the cells for a further 1 h in the absence of drug. To inhibit histone deacetylase activity, cells were cultured in the presence of 10 ng/ml Trichostatin A (TSA) for 2 h (Bickmore and Carothers, 1995).

Cells were fixed in 3:1 methanol/acetic acid (MAA) using standard procedures. Human lymphoblasts, swollen in 0.075 M KCl, were also cytogenetically analyzed on slides and fixed with 4% paraformaldehyde (pFA) made up in 120 mM KCl, 20 mM NaCl, 10 mM Tris HCl, pH 8.0, 0.5 mM EDTA, 0.1% Triton X-100. To preserve threedimensional (3D) nuclear structure, HT1080 cells and primary fibroblasts were grown on slides and fixed with 4% pFA for 10–20 min.

An exponential culture of lymphoblastoid cells was labeled with 0.1 mM BrdU for 45 min before harvest. 1.25 × 10^6 cells in 10 ml of PBS, 1% FCS, 0.3 mM EDTA, 0.1% glucose were loaded at 11 ml/min into a 5 ml elutriation chamber of a J E5.0 centrifugal elutriator (Beckman) at 2500 rpm. Fractions were taken at increasing flow rates up to a maximum of 60 ml/min; the last fraction was collected at 60 ml/min, 2000 rpm. Fractions were assayed by FACS® after staining with propidium iodide (PI), by nuclear size, and by their pattern of BrdU incorporation.

Interphase nuclei from lymphoblasts, prepared in polyamine buffer, were incubated in extraction buffer (Bickmore and Oghene, 1996) containing increasing concentrations of NaCl (0.5, 1.0, 1.2, and 1.8 M). Slides were then fixed twice in MAA before FISH.

Preparation of Chromosome Painting Probes

Chromosome paints for HSA18 and 19 were prepared in a variety of different ways and their specificity was confirmed by FISH to mitotic chromosomes. A chromosome 19 paint was produced by human-specific inter-Alu PCR from a Chinese hamster–human monochromosome 19 hybrid cell line GM10494A and labeled by nick translation. To test that this probe detected the entirety of HSA19, total DNA from the hybrid cell line was nick translated and a commercially available HSA19 paint, produced by both inter-Alu and inter-LI PCR (Oncor) was also tested. Chromosome paints prepared by the amplification of total DNA from microdissected HSA19p and q arms were also used (Guan et al., 1996).

A total chromosome 18 paint was prepared from Facs®-sorted chromosomes digested with Msel, and ligated to catch-linkers (3'-TAGCTTAACCCGTA GGCAAATCCG TTACCGGAGGG-3' [CH18-1] and 5'-CCATGTGATGGCTCCTTCGTC GCACTTAACCCGTA GGCAAATCCG TTAAGGAGGG-3' [CH18-2]), provided by S. Cross. CH18-2 was used as a primer for PCR in the presence of biotin- or digoxigenin-labeled dUTP for 35 cycles. Total DNA from the Chinese hamster–human somatic cell hybrid GM10110, a commercially available chromosome 18 paint (Oncor) and microdissected 18p and q arm paints (Guan et al., 1996) were also tested.

The area taken up by FISH signal with paints from different sources was assessed in 50 nuclei. No significant differences were found between samples of the same chromosome prepared in different ways but detected with the same fluorochrome. However, when signals from the same paints labeled with both biotin and digoxigenin and cohybridized were compared, Texas red (TR) signals occupied a larger proportion of the nuclear area than did FITC signals. Therefore, it was important that comparisons made were with the same fluorochrome.

FISH and Immunofluorescence

FISH on 3:1 fixed nuclei was as described previously (Bickmore and Carothers, 1995). Where flattened cells had been fixed in pFA for 15 min, slides were treated with 0.07 M NaOH–EDTA (2:5) for 5 min before denaturation. 3D-preserved cells were fixed in 4% pFA for 10 min, washed in PBS, and then permeabilized with 0.5% saponin, 0.5% Triton X-100 in PBS for 10 min at room temperature. The slides were incubated in 20% glycerol–PBS for 30 min and then subjected to ~5 repeated freeze-thaw cycles in liquid N2. Before hybridization these slides were treated with 0.1 M HCl for 5 min at room temperature and with 100 μg/ml RNase A at 37°C for 20 min. Slides were then denatured at 75°C in 70% formamide, 2× SSC, pH 7.0, for 3 min followed by 1 min in 50% formamide, 2× SSC, pH 7.0 (Kur et al., 1996). 150 ng of chromosome paint and 10–30 μg CotI DNA, or 200 ng labeled P1-BAC DNA's and 3 μg Cot-I DNA, were used per slide. Biotinylated probes were detected using fluorochrome-conjugated avidin (FITC or TR) (Vector), followed by biotinylated antidigoxigenin (Vector) and FITC-conjugated anti-sheep (Vector). To combine immunofluorescence with FISH in cells fixed with 4% pFA, after detection of the hybridization signals, slides were incubated for 1 h at room temperature, or at 4°C overnight, with a 1:10 dilution of rabbit anti-pH27/8 pH1B1-1 antibody (Dianova) and a monoclonal antibody I.CD43 (gift of B. Lane) directed against B-type lamin. These were detected with TR anti-rabbit and A.M.CA anti-mouse secondary antibodies (Vector). Slides were counterstained with either 1 μg/ml 4,6-diamidino-2-phenylindole (DA PI) or 0.2 μg/ml PI and examined either using a Zeiss Axioplan fluorescence microscope, equipped with a triple band-pass filter (Chroma B1000), or with a BioRad-MA-R 60 confocal laser scanning microscope fitted with an argon laser and a dual filter set (FITC and PI). For random selection of nuclei for analysis, images were taken of consecutive nuclei that presented in a spiral scan pattern from the center of the slide and which did not touch adjacent nuclei. Gray scale images from the A xiphan were collected with a CCD camera (Photometrics), pseudocolored, and merged using Digital Scientific SmartCapture extension to IPLab Spectrum.

Immunohistochemistry

U sing IPLab Spectrum software, scripts were written to analyze the data from flattened specimens. In the first, most applicable to the analysis of circular (spherical) nuclei, the DAPI image was segmented and the area and centroid coordinates calculated. The mean FITC and TR pixel intensities within the area of the DAPI segmented nucleus were calculated and subtracted from the FITC and TR images to remove background. A region of interest was then defined manually around the signal. The signal within this region was then segmented and the area and signal intensity weighted centroid coordinates calculated. The area of the signal was normalized for nuclear size by dividing by the nuclear area (see Table 1). The DAPI image was converted to binary form. U sing the coordinates of the signal weighted centroid as the center, an appropriately sized segment of circular (spherical) nuclei, the DAPI image was segmented and the area and centroid coordinates calculated. The mean FITC and TR pixel intensities within the area of the DAPI segmented nucleus were calculated and subtracted from the FITC and TR images to remove background. A region of interest was then defined manually around the signal. The signal within this region was then segmented and the area and signal intensity weighted centroid coordinates calculated. The area of the signal was normalized for nuclear size by dividing by the nuclear area (see Table 1). The DAPI image was converted to binary form. Using the coordinates of the signal weighted centroid as the center, an appropriately sized segment of circular (spherical) nuclei, the DAPI image was segmented and the area and centroid coordinates calculated. The mean FITC and TR pixel intensities within the area of the DAPI segmented nucleus were calculated and subtracted from the FITC and TR images to remove background. A region of interest was then defined manually around the signal. The signal within this region was then segmented and the area and signal intensity weighted centroid coordinates calculated. The area of the signal was normalized for nuclear size by dividing by the nuclear area (see Table 1). The DAPI image was converted to binary form. Using the coordinates of the signal weighted centroid as the center, an appropriately sized segment of circular (spherical) nuclei, the DAPI image was segmented and the area and centroid coordinates calculated. The mean FITC and TR pixel intensities within the area of the DAPI segmented nucleus were calculated and subtracted from the FITC and TR images to remove background. A region of interest was then defined manually around the signal. The signal within this region was then segmented and the area and signal intensity weighted centroid coordinates calculated. The area of the signal was normalized for nuclear size by dividing by the nuclear area (see Table 1). The DAPI image was converted to binary form.
above. The proportion of FISH signal and DAPI fluorescence was then calculated for each shell (see Fig. 1).

For confocal analysis nuclei were scanned and imaged using BioRad Comos software. Images from four high resolution scans were averaged using a Kalman electronic filter and 0.5 or 1 µm optical sections were taken. The positioning of chromosomes in relation to the nuclear periphery was assessed manually from the PI staining using display software in IP LabSpectrum. The area of hybridization signal in optical sections was measured as a proportion of the nuclear area as defined by the counterstain.

**Results**

**Chromosomes 18 and 19 Occupy Contrasting Positions within the Nucleus**

Chromosome paints for HSA 18 and 19 were used in FISH to two-dimensional (2D) preparations of nuclei, swollen in hypotonic, and fixed in MAA, from asynchronous cultures of two primary and two transformed human cell types (Fig. 1, a–d), and also to nuclei swollen in hypotonic and then fixed with 4% pFa (Fig. 1 e). In all of these cases HSA 18 and 19 were seen to adopt different positions within the nucleus, and when nuclei were cohybridized with differently labeled paints for HSA 18 and 19 (Fig. 1, a and b) signals for both chromosomes 18 appeared to be in a more peripheral location than those of HSA 19.

Using flattened specimens enabled the capture and analysis of images from large numbers of nuclei (~50 in each case) hybridized either with biotinylated HSA 18 or HSA 19 (Fig. 1) and subsequent statistical evaluation of the relative nuclear positions of these two chromosomes. In the circular (spherical) nuclei of lymphocytes and lymphoblasts the distances between both the edge, and the signal intensity weighted center of the chromosome territory, to the edge of the nucleus were measured, as was the distance between the center of the nucleus and the signal intensity weighted center of the chromosome territory (Table I). Signals for both chromosomes 18 were significantly closer to the nuclear periphery in lymphocytes and lymphoblasts than those of HSA 19 and, conversely, chromosomes 19 were located significantly closer to the center of the nuclear area than HSA 18s. These data were independent of whether total PCR amplified chromosome 19p and q arm DNA was used as a probe or whether the DNA was labeled by inter-Alu PCR (Table I).

The nuclei of fibroblasts and fibrosarcoma cells are ellipsoid, so the analyses in Table I were not appropriate. 2D nuclei from four cell types were examined by erosion. The nuclear area was divided into concentric shells (1–5) of equal area from the periphery of the nucleus to the center. The proportion of FISH signal and DAPI fluorescence was calculated for each shell. The absolute values of these are influenced by the geometric properties of the nucleus in each cell type and the definition of the nuclear periphery afforded by the DAPI counterstain, but a comparison of the proportion of the DAPI stain or hybridization signal from HSA 18 and 19 paints in each concentric nuclear shell (Fig. 1) indicates that chromosome 18 is more peripheral than HSA 19 within the diploid nuclei of all the cell types and in nuclei fixed with either MAA or pFa. The proportion of hybridization signal from HSA 18 in the outermost shell (shell 1) was always greater than, or similar to, the proportion of DAPI stain located there, whereas the proportion of HSA 19 located in this shell was always significantly less than the proportion of the DAPI stain. Conversely, in the most central shell (shell 5) the proportion of HSA 19 signal found there was always greater than that of the DAPI stain. The same relative positions of HSA 18 and 19 were also seen in fibrosarcoma cells carrying trisomies

<table>
<thead>
<tr>
<th>Table I. Relative Nuclear Positions of HSA18 and 19</th>
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<tr>
<td>Cell type</td>
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<tr>
<td>Lymphoblastoid</td>
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<tr>
<td>Interphase (4% pFa)</td>
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<tr>
<td>Interphase (MAA)</td>
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<tr>
<td>p arm</td>
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<tr>
<td>q arm</td>
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<td>G1</td>
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<td>Early S</td>
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<td>Late S</td>
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<td>G2</td>
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<tr>
<td>AMD</td>
</tr>
<tr>
<td>DRB</td>
</tr>
<tr>
<td>DRB release</td>
</tr>
<tr>
<td>TSA</td>
</tr>
<tr>
<td>Normal lymphocytes</td>
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<tr>
<td>t(18;19) lymphocytes</td>
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<tr>
<td>Normal chromosomes</td>
</tr>
<tr>
<td>Derived chromosomes</td>
</tr>
</tbody>
</table>

Measurements made (in pixels) from 50 2D nuclei for the distance between the edge of the chromosome territory to the edge of the nucleus, and between the center of gravity of the chromosome territory to the center of the nucleus, in lymphoblastoid cells fixed either with 4% pFa or MAA, and from both asynchronous cultures as well as from different cell cycle fractions separated by centrifugal elutriation. Cell cycle stage was assessed both by FACS analysis and by staining patterns with BrdU (see Fig. 3 b). Measurements were also made with 2D nuclei from lymphoblasts treated with inhibitors of transcription (AMD or DRB) and histone deacetylase inhibitors (TSA). Nuclei within standard cytogenetic preparations of normal lymphocytes and those from a t(18;19) were also measured. All distances were standardized by dividing by the square root of nuclear area (NA) as an estimate of the nuclear radius. All standard errors of the mean were ±0.01 unless otherwise indicated. The significance of the difference in position between HSA18 and 19 was assessed using a two-sample, two-tailed distribution Student’s t test. P < 0.001 in all cases. The significance of the difference in position between the derived and normal chromosomes in the t(18;19) was similarly assessed, but P < 0.059 for normal and derived 18s and 0.11 for the chromosomes 19.
 Extrapolation of data from 2D preparations assumes that the relative organization of the intact nucleus is not significantly perturbed. In Drosophila, a comparison of data collected from hypotonically treated and squashed nuclei, with those collected from 3D-preserved nuclei, has confirmed the validity of this assumption (Csink and Heni-
However, we still considered it important to confirm our observations in 3D human nuclei that had not been subject to hypotonic swelling. Also, signals that appear to be in the center of flattened nuclei might be located close to the periphery on either the top or bottom surface of the nucleus. Therefore, hybridization signals for HSA18 and 19 were examined in optical sections through the nuclei of primary fibroblasts and HT1080 cells that had been fixed with pFa, using confocal laser scanning microscopy (Fig. 2). In 94% of cells (n = 78 cells, 156 chromosomes), a substantial amount of the signals from both chromosomes 18 was coincident with the nuclear periphery, as defined from the DNA counterstain (e.g., Fig. 2 a). By contrast, in 16% of cells hybridized with an inter-Alu HSA19 paint (n = 56 cells, 112 chromosomes) and sectioned at 0.5-μm intervals, and in 20% of cells hybridized with total HSA19 DNA (n = 35 cells, 70 chromosomes) (1-μm sections) we could not detect any signal from either chromosome 19 adjacent to the nuclear periphery at this level of resolution (e.g., Fig. 2 d). Hence our observations in 3D-preserved cells, that had been fixed with pFa and not subjected to hypotonic swelling, are consistent with the conclusions of our analysis of 2D specimens. It was also interesting to note that in the z sections of 3D-preserved fibroblasts, 65.5% of chromosomes 18 were close to the lateral edge of confocal midsections (e.g., both chromosomes in Fig. 2 a). Proximity of chromosomes 18 to the upper or lower edges of the nucleus could be seen (note the green hybridization signals within the outer optical sections of Fig. 2 b). Where chromosomes 19 did appear close to the nuclear periphery this was often at the top or bottom surface of the nucleus.

The demands of preserving nuclear structure yet allowing hybridization to denatured target DNA are conflicting. However, it has been shown that in cells fixed with pFa, permeabilized, and then processed for FISH using the method described here, that centromeres remain in the same spatial position before and after the FISH procedure (Cremer et al., 1993; Kurz et al., 1996). To confirm that some components of nuclear compartments and the nuclear periphery were not disrupted by this procedure, FISH with HSA18 and 19 paints was combined with immunofluorescence for a B-type lamin, a component of the nuclear lamina (Gerace et al., 1978), and for pKi-67, a protein that localizes from satellite DNA to the nucleolus from mid-late G₁ (Fig. 3 a) (Bridger et al., 1998). Between 50 and 100 nuclei were examined and the majority of chromosomes 18, but not 19, were located close to the nuclear lamina, as detected with anti-lamin B.

**Chromosome 18 and 19 Localization Is Established Early in the Cell Cycle**

The pKi-i-67 antigen coats chromosomes in mitosis and for most of interphase (from mid G₁ through to G₂) it accumulates in the nucleolus. However, in very early G₁, pKi-i-67 is distributed in nuclear foci that correspond with blocks of heterochromatin (Bridger et al., 1998). In combined immunofluorescence-FISH of 3D preparations of human fibroblasts HSA18 was located close to the lamina both in cells with a nucleolar pattern of pKi-i-67 distribution and also in those cells with a pKi-i-67 staining pattern indicative of early G₁. At these same stages human chromosome 19 appeared to be in a more internal region of the nucleus (Fig. 3 a). Hence, the difference in subnuclear compartmentalization of HSA18 and 19 is established very early in the cell cycle, probably as cells exit telophase.

HSA18 and 19 territories were also examined in the 2D nuclei from exponentially growing human lymphoblasts, pulsed with BrdU, and fractionated by centrifugal elutria-
tion (Fig. 3 b). 50 nuclei from each cell cycle stage fraction were examined. The relative positions of HSA 18 and 19 were maintained throughout the cell cycle. Chromosome 18 was more peripheral than HSA 19 even in cells that, on the basis of their fractionation and pattern of BrdU staining, appeared to be in late S phase (Fig. 3 b and Table I). This suggests that the relative nuclear positions of chromosomes 18 and 19 are maintained before, during, and after chromosome replication.

**Human Chromosome 19 Occupies a Less Compact Chromosome Territory than Chromosome 18**

Paralleling their DNA content, chromosomes 18 at C-metaphase have a mean area 10% greater than chromosomes 19 (Table II). However, HSA 19 hybridization signal appeared to occupy a larger proportion of nuclear area in 2D preparations than did HSA 18 (Fig. 1). The normalized area of hybridization signals for both homologues of HSA 18 and 19 were measured in lymphocytes and lymphoblasts (Table II and Fig. 4 a). HSA 18 occupied a significantly smaller fraction of nuclear area than HSA 19, in flattened preparations of both MAA- and pFa-fixed nuclei, independent of the method of labeling of HSA 19. The differences in the proportion of nuclear area occupied by whole chromosome paints for HSA 18 and 19 were also seen when the signals from chromosome paints specific for the p or q arms of the chromosomes were compared (Table II and Fig. 5, a–d) (Guan et al., 1996). For example, although the q arm of the submetacentric chromosome 18 is estimated to contain 65 Mb of DNA, it occupied the same proportion of the nuclear area as the estimated 37 Mb of DNA in the q arm of metacentric chromosome 19 (Table II).

Variance in the proportion of nuclear area occupied by the two chromosomes 18 within a single nucleus was similar to that of chromosomes 18 in different nuclei (0.007 vs. 0.010, P < 0.079). However, for HSA 19 territories there was significantly less variance between homologues within the same nucleus (0.016) than between chromosomes in different nuclei (0.040, P < 0.001) indicating that chromosome 19 compaction may vary between cells that, for example, are at different stages of the cell cycle to a larger extent than chromosomes 18. In elutriated fractions of
lymphoblasts, areas of HSA 18 signals were always smaller than those of chromosome 19 and the proportion of nuclear area that they occupied decreased from G1 to G2. In contrast, HSA 19 occupied the largest proportion of nuclear area in early S phase (Table II).

Signals from chromosome 19 also appeared to be more dispersed and irregular than those of HSA 18 in both flattened nuclei (Fig. 1) and in optical sections of 3D nuclei (Fig. 2). We do not have the facilities necessary to directly compute the volumes of HSA 18 and 19 in 3D reconstructions. However, we analyzed the sum of the areas occupied by segmented hybridization signal from each chromosome as a proportion of the sum of nuclear areas in 1-μm sections through 10 nuclei (20 chromosomes 18 and 19 each). By this analysis, chromosomes 18 and 19 occupied 3.4 ± 0.3% and 5.0 ± 0.3%, respectively, of the summed nuclear areas of these 3D-preserved cells (Fig. 4b). The proportion of the summed nuclear areas occupied by signals from chromosomes 19 is significantly larger (P < 0.001) than that occupied by chromosome 18, consistent with the analyses performed on flattened specimens (Table II).

Transcription and Histone Acetylation Affect Territory Compaction but Not Nuclear Position

To analyze the effects of transcription on the territories of HSA 18 and 19, lymphoblasts were treated with AMD, an irreversible inhibitor of RNA pols I and II. To assess the drug’s effectiveness a sample of cells, before and after treatment, was examined by immunofluorescence with an antibody against snRNPs (data not shown) (Carmon-Fonseca et al., 1992; Bregman et al., 1995). In AMD-treated cells the chromosome 19 territory, but not that of HSA 18, occupied a significantly smaller proportion of the nuclear area than in untreated cells (Table II and Fig. 4a). The adenosine analogue DRB is a reversible and specific inhibitor of RNA polymerase II. The effect of DRB, and its removal, on lymphoblast and fibroblast cells was assessed by the redistribution of nuclear and nucleolar antigens by immunofluorescence (data not shown) (Scheer et al., 1984; B. Riddler et al., 1988). In the presence of D R B H SA 19 territories occupied a smaller proportion of the nuclear area in flattened specimens than those of HSA 18 and this compaction of HSA 19 was partially reversed after removal of DRB for 1 h (Table II and Fig. 4a). This was confirmed by analyzing the proportion of nuclear area occupied by signal from each territory summed across confocal 1-μm optical sections of 3D-preserved cells fixed in pFA. Whereas in untreated cells the summed chromosome 19 territory areas were 1.5 times larger than those of HSA 18, in DRB-treated cells chromosome 19 appeared to be only 0.33 times the size of chromosome 18 (n = 10 cells each) (Fig. 4b). No significant differences between the summed areas of chromosomes 18 within untreated and untreated cells were detected (P < 0.3), however, chromosomes 19 in D R B-treated cells appeared to be significantly smaller than those in untreated controls (P < 0.000).

By contrast, treatment of cultures with the histone deacetylase inhibitor TSA, to concentrations known to influence replication timing within lymphoblast cells (Bickmore and Carothers, 1995), enhanced the differences in HSA 18 and 19 areas (Table II and Fig. 4a). There was no change in the relative chromosome positions within the nucleus when transcription was inhibited with AMD or D R B, or in the presence of TSA (Table I).

The Orientation of Normal and Translocation Chromosomes

Subnuclear localization of specific human chromosomes may result from differences in DNA sequences distributed along the chromosome arms, or at the centromeres or telomer es of each chromosome. To determine whether both arms of chromosome 18 lie close to the nuclear periphery, and whether both 19p and q are equidistant from the center of the nucleus, paints specific for either 18p or q and 19p and q were cohybridized to flattened lymphoblast nuclei (Guan et al., 1996). Both 18p and q appeared in association with the edge of the nucleus (Fig. 5, a and b) and both 19p and q were found in a more central location (Fig. 5, c and d). Examination of the distances between the edge of the nucleus and the edge or intensity weighted signal center for the p and q arms, and between the center of the nucleus and the intensity weighted signal centers (Table I) confirmed that the relative subnuclear localization observed for HSA 18 and 19 reflects properties of both the p and q arms equally. To establish the orientation of chromosome 18 with respect to the nuclear periphery, HSA 18 paint was also cohybridized with P1 52M11 and BAC 75F20, specific for 18pter and 18qter, respectively (gift of J. Fantes). Most (63–65%) territories in flattened nuclei fixed in MAA had the p or q telomere at one or the other end of the territory (n = 98). In most other cases the telomere appeared to be on the surface of the territory facing the nuclear interior; localization to the surface of the terri-

<table>
<thead>
<tr>
<th>Cell type</th>
<th>Cell cycle stage or treatment</th>
<th>Proportion of occupied nuclear area by chromosome</th>
<th>Area HSA18/ HSA19</th>
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<tbody>
<tr>
<td>Lymphocytes</td>
<td>C-Metaphase</td>
<td>N/A</td>
<td>1.11 (P = 0.000)</td>
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<td></td>
<td>Interphase</td>
<td>5.0 ± 0.2</td>
<td>9.2 ± 0.3</td>
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<td></td>
<td>Chromosome 19</td>
<td>N/A</td>
<td>N/A</td>
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<tr>
<td></td>
<td>P + q arms</td>
<td>1.4 ± 0.5</td>
<td>2.5 ± 0.2</td>
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<td>Interphase (4% pFA)</td>
<td>5.0 ± 0.2</td>
<td>6.2 ± 0.3</td>
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<tr>
<td>Lymphoblasts</td>
<td>C-Metaphase</td>
<td>N/A</td>
<td>0.3</td>
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Figure 4. Dependence of territory sizes on transcription and histone deacetylation. (a) Frequency histograms of the proportion of nuclear area occupied by hybridization signals (detected with FITC) of HSA 18 (filled bars) and 19 (open bars) in 50 swollen and flattened MAA-fixed lymphoblastoid nuclei. (b) Histograms of the proportion of summed nuclear area occupied by hybridization signals (detected with FITC) of HSA 18 (filled bars) and 19 (open bars) through the confocal sections of 5–10 fibroblast nuclei fixed with 4% pFA. Vertical solid and dashed lines show mean proportionate areas for HSA 18 and 19, respectively. Cells were either untreated, or treated with AMD, DRB, or TSA before harvest. DRB-treated cells were also grown for 1 h in the absence of DRB (DRB release).
...position of the edge of the signal in relation to the edge or the center of the nucleus, in 50 t(18;19) nuclei, for both the normal (open bars) and derived (filled bars) chromosomes 18 and 19. There is no significant difference in the positions of derived and normal chromosomes (P < 0.059 for HSA18 and P < 0.110 for HSA19).

A re the positions of HSA18 and 19 within the nucleus therefore confined to the intact chromosomes or can they be conferred by subchromosomal regions? Chromosomes and nuclei in standard cytogenetic preparations from an asymptomatic individual with a balanced reciprocal trans-
location t(18;19)(p11;p13) were analyzed. We estimate that 20 M b of material from 18p (24% of HSA 18) and 15 M b of 19p (23% of HSA 19) have been exchanged in this translocation (Fig. 5 g). The nuclear disposition of both the normal and derived chromosomes was analyzed (Fig. 5, h and i). Although the derived chromosome 18 tended to be less peripheral than the normal 18, this difference was not statistically significant (Table I, P < 0.059), neither was the apparently more central location of the normal 19 over that of the derived 19 significant (P < 0.11) (Table I and Fig. 5 j). However, there was a striking difference in the relative orientations of the two derived chromosomes. The translocated portion of 19p was more central than the remainder of the derived 18 in almost 80% of nuclei. Conversely, nearly 90% of translocated 18ps were peripheral to the derived 19 material. Hence the characteristic subnuclear localization of HSA 18 and 19 can be conferred by 15–20-M b subchromosomal regions and is not dependent on the centromeric heterochromatin from each chromosome. As no viable cells are available from this t(18;19) individual we could not analyze the localization of the derived chromosomes in 3D-preserved cells.

**Chromosomes 18 and 19 Have Different Associations with Nuclear Substructure**

The nuclear scaffold and matrix are nuclear substructures that include the lamina, residual nucleoli, and a proteinaceous network pervading the nuclear volume and that are left after extraction of soluble nuclear proteins with the detergent-like salt lithium diiodosalicylate (LIS) or with high salt (Mirkovitch et al., 1984; Luderus et al., 1992).

Nuclei were extracted with increasing concentrations of NaCl before fixation in MAA and hybridization with HSA 18 and 19 paints (Fig. 6). At low salt concentrations (0.5 M) no substantial release of DNA from the nucleus was seen. At 1 M NaCl, residual nuclei surrounded by halos of released DNA were revealed by DAPI staining and FISH exposed striking differences in behavior of HSA 18 and 19. Most chromosome 18 DNA was released up to 3 μm away from the nuclear remnants, whereas the detectable chromosome 19 DNA remained in the center of the nucleus. A salt concentration increased up to 1.8 M, release of HSA 18 became more exaggerated, extending for 7 μm away from the confines of nuclei, within which chromosome 19 was still tightly restrained (Fig. 6). These differences in chromosome behavior were unchanged in nuclei from cells that had been treated with AMD to inhibit transcription.

**Discussion**

In this paper we have examined and compared the subnuclear localization of human chromosomes 18 and 19. In different primary and transformed human cells, and in both flattened specimens and in cells fixed to preserve 3D structure, we observed distinct dispositions toward either the interior or periphery of the nucleus for both the p and q arms of chromosomes 19 and 18, respectively (Figs. 1, 2, and 5 and Table I).

Specific parts of chromosomes may adopt different orientations during the cell cycle (Ferguson and Ward, 1992; Vourc’h et al., 1993; Brown et al., 1997; Csink and Henikoff, 1998; Li et al., 1998), but it was not known whether entire mammalian chromosomes move or change their state of condensation. Measurements of chromatim movements in vivo suggest that significant passive diffusion of chromatin could occur during the course of interphase but predict that whole chromosomes are largely constrained within a limited subregion of the nucleus (Marshall et al., 1997). Therefore, specific subnuclear localization of DNA segments might be established most effectively as cells exit mitosis and before interphase nuclear architecture is fully formed. The distinctive arrangements of HSA 18 and 19, that we have described here, are established early in the cell cycle and are maintained throughout interphase (Fig. 3). However, the difference in spatial localization between HSA 18 and 19 is at its smallest during late S phase (Table I). This may reflect some movement of HSA 18 to a more internal location accompanying the replication of its DNA (Li et al., 1998).
In addition to the differences between HSA 18 and 19 in subnuclear localization, we have also demonstrated differences in the proportion of nuclear area that these two human chromosomes occupy in 2D preparations. The larger proportion of nuclear area occupied by HSA 19, as compared with that of HSA 18, is in contrast to the larger physical size (in bp) of the latter chromosome (Morton, 1991) and its greater size in metaphase preparations (Table II). Since much of HSA 18 has the characteristics typical of G-band chromosome regions, whereas HSA 19 is more R-band-like in its properties (Dutrillaux et al., 1976; Korenberg and Ryzkowska, 1988; Jepessen and Turner, 1993; Craig and Bickmore, 1994), our data are consistent with the regional differences in chromatin compaction at the 0.1–1.5-M b level that have been recorded between G- and R-band regions of the human genome in nuclei prepared in similar ways to those described here (Yokota et al., 1997). The proportion of nuclear area occupied by HSA 19 reached a peak in early S phase. This might reflect HSA 19 chromatin decondensation preceding its replication (Csink and Henikoff, 1998) or merely the doubling of DNA content for this gene-rich chromosome before most of the rest of the genome.

The apparently larger area of HSA 19 within flattened nuclei is also seen in 3D-preserved nuclei (Fig. 4 b). While sophisticated computational algorithms are necessary to accurately compute chromosome volumes within the nuclear space (Eils et al., 1996; Visser et al., 1998), our simplified approach of adding together the chromosome signal area and nuclear area in series of optical sections taken through nuclei suggests that chromosome 19 may occupy a larger volume within the human nucleus than chromosomes 18 and hence be less condensed. Hybridization signals seen with paints for HSA 19 also appeared to us to have a more irregular and scattered character than those from HSA 18 (e.g., Fig. 1). The chromosome territory of the active X (Xa) is similarly more rutted in appearance than that of Xi (Eils et al., 1996); however, the volumes calculated for the territories occupied by Xa and Xi in optical sections are, in fact, the same despite the more compact appearance of Xa in flattened preparations (Eils et al., 1996). It remains to be determined whether the actual nuclear volumes occupied by HSA 18 and 19 are different from each other.

We have also shown here that transcription affects the topology of chromosome territories since the larger apparent size of HSA 19 is only seen in the presence of transcription by RNA polymerase II (Fig. 4), i.e., in untreated cells or in cells in which the inhibition of transcription by D R B has been relieved. In the absence of transcription (A M D or D R B treatments) HSA 19 occupies a compact territory similar in size, or smaller, than that of HSA 18 (Table II and Fig. 4). We do not see the gross disruption of chromosome territories in the presence of D R B that was reported previously (H aaf and Ward, 1996), except in a small minority of both treated and untreated cells that we believe are dying cells. We have no explanation for this discrepancy, but different cell types may have different tolerances and responses to the same concentrations of drugs.

The enhanced differences in areas of HSA 19 and 18 we recorded when histone deacetylation was inhibited with TSA (Table II and Fig. 4 a) suggest that levels of steady-state histone acetylation influence the gross architecture of chromosome territories. However, chromosome position within the nucleus is independent of transcription and histone acetylation activities (Table I).

More genomic sequences partitioning with the operationally defined nuclear matrix (M A R s) or nuclear scaffold (S A R s) derive from HSA 18 than from 19 (Craig et al., 1997). This, together with the proximity of HSA 18 to the lamina (Fig. 3 a), a component of the nuclear matrix, lead us to expect that there might be a tight association of this chromosome with the matrix. However, in Drosophila, sequences isolated as S A R s do not correspond with loci at the nuclear periphery (Marshall et al., 1996) and so the relationship between M A R s and S A R s and sequences that visibly remain inside of the residual nucleus after extraction, rather than in the surrounding halo of DNA loops, is not clear. Indeed, we saw very little retention of chromosome 18 DNA within nuclear matrices in contrast to the retention of chromosome 19 within the bounds of residual nuclei (Fig. 6). The degree of extension of chromosome 18 sequences varied between nuclei, but within individual nuclei the two homologues behaved similarly (Fig. 6). It has been reported that ~16 kb of inactive DNA can decondense to cover ~5 µm in extracted nuclei (Gerdes et al., 1994); therefore, the 85 M b of chromosomes 18 extruded from nuclei with high salt retain a substantial degree of higher order structure that is independent of interactions with the nucleus.

RNA is an important component of the nuclear matrix, and active genes associate with residual nuclei and not with the nuclear halo (Gerdes et al., 1994). However, retention of HSA 19 within the residual nucleus and release of HSA 18 was seen in the absence of transcription (A M D treatment). The more central location of chromosome 19 in the human nucleus may be mediated by substantive and transcription-independent association with, as yet unidentified, nuclear proteins that resist extraction from the nucleus with high salt. Jackson and Pombo (1998) have demonstrated that early replicating DNA is retained within the residual nucleus of salt-extracted human cells and, indeed, the bulk of HSA 19 replicates earlier in S phase than does HSA 18 (Dutrillaux et al., 1976).

Subnuclear localization of HSA 18 and 19 is not determined by the centromeres of the chromosomes since distinctive localization is retained by regions (~<20 M b) of the chromosome arms of HSA 18 and 19 that are translocated to the reciprocal-derived chromosome (Table I and Fig. 5, g–j). We also find no evidence that the telomeres of the chromosomes are attached at the nuclear periphery of human nuclei (Fig. 5, e and f) as has been observed in simpler eukaryotes (Hiraoka et al., 1990; Funabiki et al., 1993; Gotta et al., 1996). We conclude that differences in the overall composition of bulk chromosome 18 and 19 DNA sequences may play a direct role in the nuclear destiny of these two chromosomes and the genes placed upon them.

It has been argued that lack of phenotypic abnormalities in individuals with balanced translocations is evidence that spatial arrangement of different chromosomes in the nucleus is not functionally important. This was based on the assumption that such translocations disrupt the normal nuclear location of chromosome domains (H aaf and Schmid, 1991; Qumsiyeh, 1995). However, we have shown that this
is not necessarily the case to any significant degree (Table I and Fig. 5, g–j). The diffusion constraints on chromatin movement in vivo (Marshall et al., 1997) mean that the physical proximity of different chromosomes in interphase, and their interactions with nuclear substructure, may be important in determining the likelihood of any two chromosomes meeting and exchanging material in a translocation (Qumsiyeh, 1995). The most recurrent chromosome rearrangements in humans are Robertsonian translocations between the acrocentric rDNA-carrying chromosomes that are known to be physically close to one another in both interphase and metaphase (Kaplan et al., 1993). We surveyed two large databases cataloguing balanced translocations in humans (http://www.hgmp.mrc.ac.uk/local-data/Cad_Start.html and http://mendel.imag.fr/BAC/Transloc/carto/) and found that t(18;19) is indeed very rare in the human population when compared with other translocations among small chromosomes.

The segregation of different chromosomes of the karyotype with different functional characteristics, described here, is reminiscent of the extreme genome separation seen in plant hybrids (Letch et al., 1991). What are the biological consequences of this type of compartmentalization? The chromosomal and nuclear position of a gene can influence its activity (Brown et al., 1997; Andrilis et al., 1998) and the position of a gene within the nucleus can be dictated by the sequences it is joined to on the chromosome (Csink and Henikoff, 1998; Dernburg et al., 1996). The edge of the nucleus is also a place where genes are repressed in many eukaryotes (Bridger and Bickmore, 1998). Condensed heterochromatin and later replicating DNA tend to concentrate toward the nuclear periphery in many vertebrates (Rae and Franke, 1972; Fox et al., 1991; Kili et al., 1991; Ferreira et al., 1997), whereas early replicating DNA and poly(A) RNA partition toward the nuclear interior (Carter et al., 1993). At the level of individual loci some, but not all, active mammalian genes have been found predominantly within the nuclear interior, whereas inactive genes have been located at the nuclear and nucleolar peripheries (Xing et al., 1995). The number of gene-based markers that has been assigned to H SA 18 is small in comparison to those located on H SA 19 (Craig and Bickmore, 1994; Deloukas et al., 1998). Because of their location within the nuclear space, the relatively small number of genes located on human chromosome 18 might be habituated to very different types of transcriptional regulation to those on H SA 19. Flexibility of expression of exogenous genes placed into these two different chromosome environments may also differ. This might impose constraints on the chromosomal position of genes through evolution.

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