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Spatial organization of active and inactive genes and noncoding DNA within chromosome territories

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The position of genes within the nucleus has been correlated with their transcriptional activity. The interchromosome domain model of nuclear organization suggests that genes preferentially locate at the surface of chromosome territories. Conversely, high resolution analysis of chromatin fibers suggests that chromosome territories do not present accessibility barriers to transcription machinery.

To clarify the relationship between the organization of chromosome territories and gene expression, we have used fluorescence in situ hybridization to analyze the spatial organization of a contiguous ∼1 Mb stretch of the Wilms' tumor, aniridia, genitourinary anomalies, mental retardation syndrome region of the human genome and the syntenic region in the mouse. These regions contain constitutively expressed genes, genes with tissue-restricted patterns of expression, and substantial regions of intergenic DNA. We find that there is a spatial organization within territories that is conserved between mouse and humans: certain sequences do preferentially locate at the periphery of the chromosome territories in both species. However, we do not detect genes necessarily at the periphery of chromosome territories or at the surface of subchromosomal domains. Intraterritory organization is not different among cell types that express different combinations of the genes under study.

Our data demonstrate that transcription of both ubiquitous and tissue-restricted genes is not confined to the periphery of chromosome territories, suggesting that the basal transcription machinery and transcription factors can readily gain access to the chromosome interior.

Introduction

There is a well-established functional link between chromatin structure at the level of the nucleosome and gene expression (Jenuwein and Allis, 2001). However, the functional significance of higher-order chromatin structures in transcription remains unclear and has been examined at different levels; from large scale structures above the 30-nm fiber to whole chromosome territories (Mahy et al., 2000). It was thought that large scale chromatin organization above the 30-nm fiber might affect transcription by presenting an accessibility barrier to large protein complexes (Zirbel et al., 1993). Hence, it has been proposed that transcription and RNA processing might occur in a space between territories called the interchromosome domain (ICD)* compartment (for review see Cremer and Cremer, 2001). In support of this model, specific gene transcripts and components of the splicing machinery have been reported to concentrate at the border of chromosome territories (Zirbel et al., 1993). However, poly(A) RNA is not excluded from chromatin domains, and nascent RNA can be seen deep within chromosome territories (Abranches et al., 1998; Verschure et al., 1999). The idea of a compact chromosome territory that would act as a barrier to the transcription machinery is contradicted by studies of chromatin fibers at higher resolution by both light microscopy and EM (Belmont et al., 1999).

If transcription does occur close to the surface of chromosome territories, genes should preferentially be found there, and noncoding sequences should be more internal. The intraterritory positions of a small number of individual genes from scattered genomic locations have been examined. In one report, three coding regions of the human genome were all located in the periphery of their respective chromosome territories independent of transcriptional status, whereas a noncoding sequence was not (Kurz et al., 1996). An active gene has also been found in a more peripheral location...
within the active X chromosome (Xa) territory than its inactive counterpart in the inactive X chromosome (Xi) territory (Dietzel et al., 1999). The imprinted SNRPN genes are found at the periphery of both chromosome 15 homologues (Nogami et al., 2000). Over larger regions, it has been shown that the gene-rich major histocompatibility complex (MHC) lies on large chromatin loops that extend away from the surface of the bulk chromosome 6 territory that is detectable with a chromosome paint (Volpi et al., 2000). An extreme interpretation of these data is that most (active) genes lie on the surface of chromosome territories. However, with the exception of the Xi, both early and late replicating DNA that are usually equated with gene-rich and gene-poor domains, respectively, appear to be distributed throughout chromosome territories (Visser et al., 1998). Similarly, the most GC-rich fraction of the human genome (which has a high gene density) is also distributed throughout the volume of territories (Tajbakhsh et al., 2000).

Models of higher-order chromatin fiber and chromosome organization have implications for the spatial organization of genomic DNA in the nucleus both at the long range (chromosome band) level and at a more local megabase (Mb) level. To investigate the relationships between the organization of chromosome territories and gene expression, we have used fluorescence in situ hybridization (FISH) on both two-dimensional (2D) and three-dimensional (3D) samples to analyze the spatial organization of a contiguous 1 Mb stretch of the human genome and the syntenic region in the mouse. We have used cell types that express different repertoires of genes and their chromosome territories. All of the 11p13 loci studied, including expressed genes, have a mean position well within their chromosome territories. Therefore, there is sufficient spatial resolution at interphase to detect any significant differences in their intraterritory distribution.

Using reverse transcriptase (RT)-PCR, we confirmed the expression of RCN and PAXNEB in lymphoblastoid cells and primary fibroblasts. WT1 and PAX6 are not expressed (Fig. 1). Cosmids encompassing the human WAGR region were first hybridized together with a paint for 11p to MAA-
cells. Rather, it is more likely that small local changes in large scale chromatin fiber conformation accompany gene expression (Belmont et al., 1999). Our data may be compatible with the modified ICD model that suggests that this compartment penetrates into territories, ending at the surface of compact chromosomal subdomains of 0.3–0.45 μm diameter (Verschure et al., 1999; for review see Cremer and Cremer, 2001). Chromatin fibers containing transcriptionally active DNA may be then be decondensed at the surface of these sub-domains or extend into the interchromatin spaces (Verschuer et al., 1999). However, although we have found that a ubiquitously expressed gene is indeed located at the surface of, or outside of, such subdomains and although adjacent noncoding DNA is located within the compact subdomain, we find no clear correlation between chromosome territory subdomains and the expression of tissue-restricted genes.

Results
Active genes from 11p13 do not locate preferentially at the chromosome territory surface
It has been suggested that genes are preferentially located in the periphery of chromosome territories (Kurz et al., 1996; Dietzel et al., 1999; Volpi et al., 2000; Cremer and Cremer, 2001). However, the positions of only a small number of individual genes, usually from scattered chromosomal locations, have been examined. To determine if genes from a contiguous region of the human genome are located together at the chromosome periphery, we examined the intra-chromosomal organization of a megabase stretch of the human genome.

Within the distal ~1 Mb of human 11p13, there are four known genes. RCN (Kent et al., 1997) and PAXNEB (Kleinjan et al., 2002) are ubiquitously expressed, whereas the expression of WT1 and PAX6 is tissue restricted (Hastie, 1994; Xu et al., 1999). Gene order is conserved at the syntenic region on MMU2E, and there are large intergenic regions (~300 kb) between WT1 and RCN and between RCN and PAX6 in both man and mouse (Kent et al., 1997; Gawin et al., 1999). Gene order and spacing are conserved at the region of conserved synteny on mouse chromosome 2 (MMU2E) (P. Gautier, personal communication) (http://www.ensembl.org/Mus_musculus/; http://www.ncbi.nlm.nih.gov/Omim/Homology/human11.html). We have compared the organization of these genomic regions to a more distal region on human chromosome 11 (HSA11) (11p15) and its region of conserved synteny in the mouse MMU7.

Our data show that there is spatial organization within chromosome territories. However, genes from 11p13 or MMU2E do not preferentially localize at the periphery of their chromosome territories. All of the 11p13 loci studied, including expressed genes, have a mean position well within the bulk chromosome territory compared with a locus from 11p15 that locates at the territory edge. Tissue-restricted genes are not relocated to the territory periphery in expressing cells.

We conclude that in general there is no gross remodelling of chromosome territory organization to accommodate small changes in gene expression within mammalian
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Fixed lymphoblast nuclei (Fig. 2 a). Images from 50 randomly selected nuclei were analyzed per locus. The distance (μm) between the center of each locus signal to the nearest chromosome territory edge was calculated as described (see Materials and methods). For the expressed gene RCN, 60% of signals were ≥0.4 μm from the territory edge (Fig. 3 b). Similar figures (61 and 65%) were seen for the linked inactive gene PAX6 and the noncoding D11S324 locus. This was in contrast to a locus (D11S12) from a more distal chromosome band 11p15.4 for which the bulk of signals were at the territory edge and only 27% of signals were ≥0.4 μm from the territory edge (Fig. 2 b and Fig. 3 b). The mean distances of all the WAGR probes from the territory edge (0.45–0.65 μm) were significantly greater than that for D11S12 (Fig. 3 c) and other loci from 11p15 (unpublished data). WAGR probes were not found preferentially at the territory edge but neither were they in the center of the territory. The average distance between the territory centroid and edge was 1.15 μm. Because we found that territories detected in G2 cells were larger than those in G1 cells (unpublished data), we also normalized actual distances relative to the radius of a circle of equal area to the chromosome territory. Values of 0 and 1 then equate to theoretical positions at the territory edge or center, respectively (Fig. 3 c). Because territories are not circular, the actual position of the

Figure 1. Expression of WAGR locus genes in human cell lines. Transcripts from WAGR locus genes were amplified by RT-PCR. Amplification was specific to the RNA and not from contaminating DNA, since there was no productive amplification from control −RT samples. The ovarian carcinoma cell line COV434 expresses the WT1 gene; RCN is expressed in all four cell lines tested; PAX6 is expressed only in the lens epithelium-derived cell line CD5a; PAXNEB is expressed in all four cell lines. M, size markers in base pairs.

Figure 2. Visualizing intraterritory position of loci from human 11p13 and 11p15. FISH of selected cosmids (red) from the WAGR region (11p13) or 11p15 together with an 11p paint (green) to MAA-fixed FATO lymphoblast (a) and 1HD primary fibroblast (b) nuclei. cH11148 detects the ubiquitously expressed RCN gene. cB2.1 contains the WT1 gene that is not expressed in either of the two cell types. cA08102 is from the intergenic region between RCN and PAX6 (D11S323) (Fig. 3 a). D11S12 is an anonymous marker from 11p15.4. The last image in panel a shows the territory segmentation masks for the cA08102 image. (c) Three plane images through a 3D preserved pFa-fixed fibroblast nucleus at 1-μm intervals in the z direction after hybridization with probes to RCN and 11p. Views of an MAPaint reconstruction of the same nucleus after delineation of the nucleus, chromosome territory, and locus probe domain are shown at 0 and 90° rotation about the x axis. Nuclei were counterstained with DAPI (blue). Bars, 5 μm.
Figure 3. Intraterritory organization of the human WAGR region. (a) Position of genes within the distal 1 Mb of human 11p13 DNA relative to cosmids used in this study. Open boxes represent ubiquitously expressed genes. Filled boxes represent genes with tissue-restricted expression. Arrows indicate the direction of transcription. The bar represents 100 kb. (b) Histogram showing the distribution of signals relative to the territory edge (in µm) for expressed (RCN) and nonexpressed (PAX6) genes from 11p13, a noncoding region of 11p13 (D11S324), and a marker (D11S12) from 11p15 in MAA-fixed FATO lymphoblast nuclei. (c) Mean position (± SEM) of 11p13 and 11p15 probes within the 11p territory of MAA-fixed lymphoblast nuclei. Distance in µm is shown on the left, and distances normalized for territory size are shown on the right where values of 0 and 1 equate to positions at the territory edge or the theoretical territory center, respectively (n = 100). The average distance between territory centroid and nearest territory edge in these cells was 1.14 µm, and the normalized position of the measured territory center is 0.64. Intercepts for the graphs are thus placed at these values. (d) Mean position of 11p13 and 11p15 probes within the 11p territory of MAA-fixed fibroblast nuclei (●) or in fibroblast nuclei fixed in pFa to preserve 3D structure (▲).
territory centroid in lymphoblasts was at 0.65 ± 0.02 (Fig. 3 c). Using analysis of variance, there was no significant difference between the positions of any of the WAGR loci examined \( (p = 0.68) \), but WAGR probes were located further inside of the chromosome territory than D11S12 \( (p < 0.01) \) (Fig. 3 c).

 Territory organization may vary through the cell cycle, for example, before, during, or after replication. Cosmid signals often appear as single spots of hybridization before replication of the locus that they detect and as paired dots (Fig. 2 a) after replication (Bickmore and Carothers, 1995). We detected no significant difference \( (p = 0.41) \) between the intraterritory position of a locus before and after its replication. Similarly, the variance in locus position within the two territories of homologous chromosomes within a cell was not less than that between cells in an unsynchronized population, indicating that territory organization within homologous chromosome territories of nuclei is independent.

 Human primary fibroblasts were also subjected to the same analysis (Fig. 2 b). All of the 11p13 probes tested were located inside of the chromosome territories compared with D11S12 (Fig. 3 d).

 Analysis of 2D specimens allows for statistical analysis of large numbers of images. However, it was important to establish that data from these preparations was consistent with intraterritory organization in cells in which 3D nuclear architecture had been preserved (Kurz et al., 1996; Croft et al., 1999; Volpi et al., 2000). Using human HT1080 cells that stably express centromere-localized GFP-tagged CENP-A, we first determined that the spatial organization of centromere sequences was not perturbed during neither the fixation procedure with 4% paraformaldehyde (pFa) nor the subsequent FISH procedures. We took image stacks of centromere-localized GFP signal in the living cells (Fig. 4 a) and then in the same cells after pFa fixation (Fig. 4 b). Lastly, hybridization signal for an \( \alpha \)-satellite probe was analyzed in these same cells after 3D FISH (Fig. 4 c). The images from the three stages of the experiment were comparable. Signals from the detection of \( \alpha \) satellite by FISH were generally larger and more variable than those from GFP–CENPA (Fig. 4). This is because similar amounts of CENPA are assembled into the kinetochore of every chromosome, whereas the size of \( \alpha \)-satellite arrays varies amongst centromeres and extends beyond the kinetochore itself. In addition, the efficiency with which the \( \alpha \)-satellite probe detects each centromere will vary depending on the exact sequence of the arrays. We measured the distance separating pairs of centromeres \( (n = 10) \). The distance between two centromeres in living cells decreased by an average of 0.1 \( \mu m \) upon fixation with pFa. After 3D FISH, the distance between pairs of centromeres had increased by 0.18 \( \mu m \) compared with living cells. Hence, any nuclear reorganization brought about by the 3D FISH process is small and does not affect our conclusions.

 We then analyzed the intraterritory position of loci from the WAGR region in 3D preserved primary fibroblast nuclei (Fig. 2 c). Measurements were made on 15–18 randomly selected nuclei per locus. Probes from the human WAGR locus (Fig. 3 a) were on average positioned 0.4–0.5 \( \mu m \) away from the edge of the chromosome territory. These distances were generally a bit smaller than those measured in 2D samples, but the normalized position within the territory was similar between 2D and 3D specimens (Fig. 3 d). This suggests a general swelling of the chromosome territory in 2D preparations but no reorganization of sequences with respect to each other (Volpi et al., 2000). In 3D preparations, the 11p15.4 locus D11S12 was located at the territory periphery as in 2D preparations.

 Therefore, we conclude that there is no preference for genes from 11p13 to locate at the periphery of the 11p territory, as defined in FISH by a chromosome paint, compared with intergenic sequences from the same region. Ubiquitously expressed genes can be transcribed at a mean distance of 0.6 \( \mu m \) away from the visible territory edge. However, there is a reproducible spatial compartmentalization within the chromosome 11 territory: a locus D11S12 from 11p15.4 and other 11p15 loci (unpublished data) are located preferentially at the territory edge.

 Evolutionary conservation of chromosome architecture

 Gene order and spacing are conserved at the murine WAGR region of synteny on MMU2E (P. Gautier, personal communication; http://www.ensembl.org/Mus_musculus/; http://www.ncbi.nlm.nih.gov/Omim/Homology/human11.html) (Fig. 3 a and Fig. 5 b). To investigate whether this linear sequence conservation also extends to the spatial organization of the locus, we investigated the intraterritory position of genes spanning the murine WAGR locus. Bacterial artificial chromosomes (BACs) encompassing this region (Fig. 5 b) were hybridized together with a paint for MMU2 to MAA-fixed embryonic stem (ES) cell nuclei. Compared with the human WAGR locus on 11p (Fig. 3), genes and intergenic DNA from the mouse WAGR region are located even further (0.75–1 \( \mu m \)) away from the edge of the MMU2 territory (Fig. 5 c). This is due to the larger size of the MMU2 territory compared with HSA11p (200 versus 50 Mb) (http://www.ensembl.org/Mus_musculus/ and http://www.ensembl.org/Homo_sapiens/), and territory centroids were on average 1.58 \( \mu m \) away from the nearest territory edge. Indeed the positions of the human and mouse WAGR loci, normalized to take account of territory size,
are very similar (Fig. 5 c). On human 11p, sequences from 11p15 are at the territory periphery (Fig. 3). The mouse region of conserved synteny to human 11p15 is on MMU7 (http://www.ncbi.nlm.nih.gov/Omim/Homology/human11.html). DNA hybridizing to a BAC 245N5 from this region (Engemann et al., 2000) was also found to locate at the edge of the MMU7 territory (Fig. 5, a and c). Therefore, in addition to primary sequence conservation there is conservation of relative intraterritory organization between regions of the mouse and human genomes that are in conserved synteny, suggesting that this organization has functional significance.

Figure 5. Intraterritory organization in the mouse. (a) FISH of BACs 82K15 (Wt1) and 50J5 (Rcn) (red) from the WAGR region together with a paint for MMU2 (green) or BAC 245n5 (red) with an MMU7 paint (green) to MAA-fixed ES cell nuclei. Nuclei were counterstained with DAPI (blue). Note the yellow color of the (red) probe signals located within the (green) MMU2 chromosome territories and the red color of the MMU7 probe signal at the territory edge. Bar, 5 μm. (b) Map of the mouse region of conserved synteny to WAGR at human 11p13 on MMU2, relative to the BACs used as probes. (c) Mean position (± SEM) of MMU2 and seven probes within chromosome territories of MAA-fixed ES cell nuclei. Distance in μm is shown on the left, and distances normalized for territory size are shown on the right where values of 1 and 0 equate to positions at the theoretical territory center or periphery, respectively (n = 100). The mean distance between the MMU2 territory centroid and nearest territory edge was 1.48 μm in ES cells, and the normalized position of the territory centroid was 0.58. Graph intercepts are placed at these values.
Changes in gene expression do not require spatial reorganization of 11p13

Transcription of genes from the WAGR region from within the volume of the chromosome territory could be restricted to ubiquitously expressed genes that do not require specialized transcription factors for their expression. To examine whether expression of tissue-restricted genes could also take place away from the periphery of chromosome territories, the organization of the human WAGR locus was analyzed in nuclei from cell lines that also express tissue-restricted genes. RT-PCR analysis showed that CD5a, a lens epithelial cell line, expresses PAX6 and that COV434, an ovarian carcinoma cell line, expresses WT1 (Fig. 1). Immunofluorescence (WT1) and immunohistochemistry (PAX6) showed that all cells in the populations express the proteins (unpublished data).

The spatial organization of the WT1 and PAX6 genes was assessed in 3D-preserved CD5a and COV434 cells. PAX6 is not preferentially located at the periphery of the 11p territory in CD5a cells (Fig. 6 a); 84% of FISH signals were ≥0.4 μm away from the territory edge (Fig. 6 b). 75% of WT1 signals were ≥0.4 μm from the territory edge in COV434 cells (Fig. 6 b). Therefore, we find no evidence that tissue-specific genes are repositioned to the periphery of chromosome territories to facilitate transcription, and transcription factors for such genes must be able to access their targets inside of chromosome territories.

Genes and subchromosomal domains

The original ICD compartment model described chromosome territories as compact objects surrounded by a network of channels that connect to the nuclear pores (Zirbel et al., 1993; Kurz et al., 1996). However, the ICD compartment has been extended recently to include small channels penetrating chromosome territories, ending between 1-Mb domains of perhaps 0.3–0.45 μm diameter (Verschure et al., 1999; Cremer and Cremer, 2001). Actively transcribed loci are thought to be positioned at the surface of these compact subchromosomal domains rather than at the surface of entire chromosomes, depositing newly synthesized RNA directly into the extended interchromatin space (Verschure et al., 1999). This is more compatible with high resolution observation of large scale chromatin fibers of different compaction (Belmont et al., 1999). In accordance with this idea, in 3D preserved fibroblast nuclei we found a preference for the ubiquitously transcribed RCN gene to be positioned in areas of the chromosome territory unlabeled by FISH (50% of cases, n = 14). In a single image plane, the probe appears as a red signal, indicating no colo-
calization with the deconvolved green chromosome territory signal (Fig. 7 a). This contrasted strongly with a probe cAO8102, detecting noncoding intergenic DNA ∼300 kb distal of RCN (Fig. 3 a). No signals from this probe (n = 13) were found outside of the compact territory subdomains detected by FISH with a chromosome paint. Instead, the probe signal was frequently (61.5%, n = 13) positioned within a region of densely stained chromatin. Hence the probe signal appeared yellow due to its colocalization with the labeled chromosome subdomain (Fig. 7 b). Our data supports a correlation between the position of DNA relative to chromosome subdomains and gene expression (Verschure et al., 1999), at least in the case of housekeeping genes. However, the position of genes with a tissue-restricted expression pattern with respect to subdomains was less clear. At this level of analysis, the position of the WT1 and PAX6 genes within chromosome subdomains from fibroblasts, COV434, and CD5a cells did not appear to be correlated with their expression pattern (unpublished data). Therefore, there is not a consistent correlation between transcription and chromosome territory substructure defined by chromosome paint. Furthermore, unlabeled areas of a chromosome territory cannot be “holes” devoid of chromatin, since we find gene loci located in these regions. It is likely that unlabeled regions represent decondensed chromatin fibers within the territory that cannot be detected by FISH with chromosome paints rather than defined interchromatin channels.

Discussion

Transcription of genes can occur within the interior of a chromosome territory

Components of the splicing machinery and specific gene transcripts of an integrated human papilloma virus genome are often excluded from the interior of chromosome territories (Zirbel et al., 1993). Furthermore, three coding regions of the human genome have been observed at the periphery of chromosome territories, and a single nontranscribed sequence was randomly distributed or more internally positioned (Kurz et al., 1996). Within the context of the ICD compartment model of nuclear organization (Cremer and Cremer, 2001), an extreme interpretation of these data is that most or all genes lie on the periphery of chromosome territories and that the territory interior is filled with intergenic sequence (Zirbel et al., 1993; Kurz et al., 1996). However, here we have shown that both coding and noncoding sequences from 11p13 are similarly located inside of the HSA11p territory. Both ubiquitously expressed genes and genes with a tissue-restricted expression pattern can be transcribed from within the territory interior (Figs. 3 and 6). This argues against the concept of a compact chromosome territory and indicates that the basal transcription machinery and transcription factors must be able to readily access chromatin located within the interior of chromosome territories. Our data indicate that large scale chromatin remodelling to position genes on the surface of a territory is not required to facilitate transcription but rather that any changes to the chromatin fiber are likely to be local (Belmont et al., 1999).

Therefore, our data question the extent to which the organization of chromosomes within territories can contribute to control of gene expression. Replication has been shown to take place in foci located throughout the entire chromosome territory volume (Visser et al., 1998). This demonstrates that activity of macromolecular enzyme complexes takes place throughout chromosome territories and is not confined to the territory surface (Zirbel et al., 1993; Kurz et al., 1996). In addition, transcription sites have been visualized throughout the nucleus (Abranches et al., 1998; Verschure et al., 1999), and using FRAP nuclear proteins with diverse functions and distinct distribution patterns have been shown to diffuse rapidly throughout the entire nucleus (Phair and Misteli, 2000). The fact that the human WAGR locus is positioned inside the HSA11p chromosome territory rather than at its periphery is consistent with these observations.

Intraterritory organization is not random and is conserved

Although we did not find genes from 11p13 preferentially at the periphery of the HSA11 territory, we did find that territories are spatially organized. In contrast to 11p13 DNA, sequences from 11p15 were preferentially located at the surface of the HSA11 territory (Fig. 3).
We extended our study of the WAGR locus to the syntenic region in the mouse. This is the first time that the organization of murine chromosome territories has been considered. We found that sequences, including expressed genes, from the murine WAGR locus were located in a similar relative position inside of the territory of MMU2 as the human region is within the HSA11p territory (Fig. 5). Likewise, the region of MMU7 in conserved synteny with HSA11p15 is located at the periphery of the MMU7 territory (Fig. 5). Conservation of sequence organization within chromosome territories suggests that it has functional significance, but it remains to be determined what that is. The conservation of spatial organization within chromosome territories is interesting in light of the conservation of territory position within the nucleus that has been seen in vertebrates (Croft et al., 1999; Boyle et al., 2001; Habermann et al., 2001; Tanabe et al., 2002).

Small scale changes in the transcription profile of a region do not require territory reorganization

Three studies have indicated a role for active transcription in organizing gene position within a chromosome territory (Dietzel et al., 1999; Volpi et al., 2000; Williams et al., 2002). Volpi et al. (2000) noted a difference in the incidence of extrusion of chromatin containing the MHC locus from the surface of HSA6 in cell lines with different expression profiles of MHC genes. Inducing transcription from this region using IFNγ increased the incidence of chromatin looping (Volpi et al., 2000). A relationship between transcription status and position was observed for a single gene within the Xa and Xi territories. The SLC25A5 (previously ANT2) gene was more peripheral within the Xa chromosome from which it is actively transcribed than within the Xi territory in which it is silent (Dietzel et al., 1999).

Using cell lines that express the tissue-restricted WT1 or PAX6 genes (Fig. 1), we found no significant alteration of the intraterritory position adopted by the WAGR locus that correlated with increased gene expression. Genes that are inactive in lymphoblasts and fibroblasts were not relocated to the HSA11p territory surface when actively transcribed but rather remain within the chromosome territory in a position similar to those of neighboring ubiquitously active genes and noncoding sequences (Fig. 6).

Differences in intraterritory organization dependent on transcription status of associated genes were ascertained previously within the context of chromosome domains subject to large scale transcriptional differences. The MHC locus consists of families of genes with the same or a related function; therefore, all of the genes are subject to the same regulation kinetics (Volpi et al., 2000). Similarly, the human epidermal differentiation complex at 1q21 contains functionally related genes involved in keratinocyte differentiation. This region appears to be extended outside of the HSA1 territory in keratinocytes where the genes are highly expressed but not in lymphoblasts where they are silent (Williams et al., 2002). Transcriptional activation of a 90-Mb heterochromatic region containing 256 copies of the lac operator sequence has also been associated with large scale chromatin decondensation (Tumbar et al., 1999). Lastly, the Xi adopts a unique structure which differs from the Xa territory (Eils et al., 1996; Dietzel et al., 1999) and is generally devoid of sites of transcription (Vereschure et al., 1999). It is unclear how widely applicable observations made from study of X chromosome territory organization will be to the rest of the human karyotype.

Do levels of chromatin packaging within territories contribute to control of gene expression?

A high resolution chromosome painting study has suggested that locally compacted and unfolded regions within chromosome territories form distinct subdomains of 0.3–0.45 μm diameter and that chromatin is organized in such a way that transcriptionally active loci are at the surface of large scale chromatin fibers (Veschure et al., 1999). These data imply that limitations on protein accessibility within chromosome territories are likely to map to the periphery of large scale chromatin fibers or larger chromosome subdomains formed by the folding of these fibers. Transmission EM localization of uridine or BrUTP incorporation has shown heavy labeling at the edge of condensed large scale chromatin domains rather than at the surface of chromosomes itself (Fakan and Nobis, 1978; Wansink et al., 1996; Cmarko et al., 1999). Transcripts of 1α1 collagen located within the chromosome 17 territory similarly often coincide within holes in the chromosome paint (Clemson and Lawrence, 1996). Such observations have been used to extend the concept of the ICD (now interchromatin domain) to include channels penetrating into chromosome territories (Cremer and Cremer, 2001), ending in branches between 1 Mb and 100 kb chromatin loop domains. Transcription apparently occurring within chromosome territories is therefore actually occurring on the “surface” of chromosome subdomains (Veschure et al., 1999). In accordance with this idea, we found that a ubiquitously transcribed gene often colocalized with unlabeled or less intensely labeled areas of the chromosome territory, whereas the linked intergenic locus was positioned frequently in intensely labeled (compact) subdomains of the territory (Fig. 7). The fact that we found gene sequences themselves and not just their transcripts in apparent “holes” in the chromosome territory implies that these are not in fact channels devoid of chromatin but rather that they correspond to areas of decondensed chromatin fibers not detectable by chromosome painting. The positions of genes with a tissue-restricted expression pattern with respect to chromosome subdomains did not appear to correlate with their expression status. Therefore, the role of subchromosomal packaging in facilitating transcription remains open to debate and further investigation.

Materials and methods

Cell culture

Human FATO lymphoblasts (46XY) were grown in RPMI plus 10% FCS. Human 46XY primary fibroblasts (less than passage 12) and the human lens epithelium-derived cell line CD5A (a gift from A. Prescott, University of Dundee, Dundee, UK) were grown in DMEM plus 10% FCS. COV434 human ovarian (granulosa) carcinoma cells (Berg-Bakker et al., 1993) were grown in a 1:1 mixture of F10 and DMEM supplemented with 10% FCS. The murine ES cell line E14 (Hooper et al., 1987) was maintained on a gelatinized 0.1% in PBS surface in MEM/BSK21 medium supplemented with 0.23% sodium bicarbonate, 1X MEM nonessential amino acids, 2 mM glutamine, 1 mM sodium pyruvate, 50 μM β-mercaptoethanol (all Life Technologies), 10% FCS (Sigma-Aldrich), and 100 μM soluble DIA/LIF (Sigma-Aldrich).

Immunofluorescence and immunohistochemistry

WT1 protein was detected in COV434 cells by immunofluorescence using the C19 polyclonal antibody (Santa Cruz Biotechnology) that detects the COOH-terminal 19 amino acids of the protein and an FITC-conjugated
anti-rabbit secondary antibody (Jackson ImmunoResearch Laboratories). PAX6 protein was detected in CDSA cells (A. Seawright, personal communication) using AD1.5.6 and AD2.3.7 antibodies (1:1) raised against the NH2-terminal 206 amino acids of PAX6 (Engelkamp et al., 1999). Staining was visualized using Vectastain ABC Systems and NBT/BCIP substrate kits (Vector Laboratories).

**FISH**

A paint for human chromosome 11p (HSA11p) was labeled with biotin-16-DUTP by PCR amplification (Guan et al., 1996; Croft et al., 1999). Cosmids from 11p (Fantes et al., 1995) were labeled by nick translation with digoxigenin-11-DUTP. This labeling scheme was adopted because the 11p chromosome territories detectable with biotin-labeled paint were brighter and more reproducible than those seen using digoxigenin-labeled paints. 210 ng paint and 50 ng cosmids were used per slide in a volume of 12 μl together with 6 μg human Cot1 DNA (GIBCO BRL) as competitor.

**MMU2 paint** (Jentsch et al., 2001) was amplified and biotin labeled by PCR using the degenerate primer 5′-CCGCAGCTGAG(N)6TACACC-3′. Mouse BACs, 82K15, 98L11, 67L02, and 46F10 were selected from a 129 library (Genome Systems, Inc.) (D.A. Kleijnan, personal communication) and labeled with digoxigenin by nick translation. 200 ng paint were used together with 100 ng Bac DNA per slide in a volume of 12 μl with 14 μg mouse Cot1 DNA (GIBCO BRL) added as competitor.

For 2D analysis, cells were swollen in 0.5% trisodium citrate/0.25% KCl before fixation in methanol acetic acid (MAA; 3:1 vol/vol) using standard procedures. Slide hybridization was as described previously (Fantes et al., 1995; Croft et al., 1999). To preserve 3D nuclear structure, human primary fibroblasts were grown on slides and fixed with 4% pFA/PBS for 10 min before fixation in methanol acetic acid (MAA; 3:1 vol/vol) using standard procedures. After signal detection, the same cells were recloned on the slide, and image stacks were taken of the centromeric hybridization signals (Fig. 4).

Biotinylated paints were detected using fluorochrome-conjugated avidin (FITC or Texas red) (Vector Laboratories) followed by biotinylated antiavidin (Vector Laboratories) and a final layer of fluorochrome-conjugated avidin. Digoxigenin-labeled probes were detected with sequential layers of FITC-conjugated antidigoxigenin (BCL) and FITC-conjugated anti–sheep (Vector Laboratories). Slides were counterstained with 1 μg/ml DAPI. 2D slides were examined using a ZEISS Axioplan fluorescence microscope equipped with a triple band-pass filter (Chroma #83000). Gray scale images were collected with a cooled CCD camera (Princeton Instruments Pentamax) and analyzed using Adobe Photoshop software. For 2D analysis, a focus motor was used to select images at 0.5-μm intervals in the z direction from a ZEISS Axioplan using a Xillig CCD camera. 3D image stacks were analyzed using IPLab extensions and deconvolved using Hazebuster (Vaytek, Inc.). For random selection of nuclei for analysis, slides were scanned in a methodical manner, beginning at the top left hand corner, scanning to the right, then moving downwards and scanning the next row of nuclei from right to left, etc. Images of consecutive nuclei that had two chromosome territories in which locus signals were visible and that were not touching adjacent nuclei, since this interferes with segmentation (see below), were collected.

**Image analysis**

The following script was used to analyze the position of loci within chromosome territories in 2D. Nuclear area was calculated from the segmented DAPI image. Background was removed from FITC and Texas red images as described previously (Croft et al., 1999). Locus-specific hybridization signals were automatically segmented, and a region of interest was manually defined around one of the signals (each locus signal was treated independently; each territory contained either one or two, depending on whether the cell had been replicated). The centroid coordinates of the locus signal were calculated. Hybridization signal from the chromosome territory was then segmented interactively on the basis of pixel thresholding without knowledge of the locus signal, and a region of interest was manually defined to include all of the detectable territory. An example of a chromosome territory segmentation mask is shown in Fig. 2a. The area of the territory was calculated. A segmentation distance map was dilated out from the signal centroid and then eroded until a pixel with zero intensity from the territory signal was found. This was taken to be the nearest edge of the territory to the locus, and the radius of the disc was calculated representing the distance from the center of the locus to the nearest edge of the territory (μm). A similar procedure was used to determine the distance between the territory centroid and territory edge. Because actual territory size varied between different cell types and at different cell cycle stages, we normalized the locus to territory edge distance by dividing it by the radius of a circle of equal area to the territory. A value of 0.0 thus denotes a locus at the edge of a chromosome territory, and 1.0 denotes a locus at the theoretical center of that circle. Negative values indicate loci that appear to be located outside of the visible limits of the chromosome territory. In practice, values of 1.0 are not seen because territories are not circular, and the actual territory centroid had a mean value of 0.64 ± 0.02 along this theoretical radius for HSA11p and 0.58 ± 0.03 for MMU2. To assess the reproducibility of this method of analysis, the same set of 50 images were analyzed independently by two different observers. The actual distance (0.53 ± 0.05 and 0.58 ± 0.04 μm), and the normalized distance (0.31 ± 0.02 and 0.32 ± 0.02) of the PAX6 gene from the HSA11p territory surface in lymphoblasts was similar in both cases.

To analyze 3D images, signals from the chromosome territory and the cosmid or BAC probe were converted to gray scale images. Regions corresponding to the chromosome territory and probe were segmented manually in each plane using the program MAPaint (http://genex.hgu.mrc.ac.uk/Software/paint/). Chromosome domains were delineated by thresholding, which segments connected regions within the image that have the same signal intensities. The probe signal was delineated using the “paint ball” facility. The volume of the chromosome territory domain and the minimum distance between the center-of-mass of the probe domain and the surface of the territory were calculated. Spatial coordinates were normalized using the measured voxel size (x, y, z) to compensate for 0.5-μm spacing between consecutive image planes.

For visualization of the chromosome and probe domains in context, the nucleus was also segmented using the same “thresholding” facility. The AVS/Express visualization system (Advanced Visual Systems, Inc.) was used to produce a surface-rendered 3D view, and this was used to make an MPEG movie by rotation around the x axis (http://www.mpeg.org).

The significance of differences in the relative positions of different loci using 2D and 3D analysis techniques was tested by one-way analysis of variance using MINITAB software (release 12.21). The results of all analyses on WAGR showed that the component of variance between the intraterritory position of probes in different nuclei was not significantly greater than the variance between intraterritorial position within homologous chromosomes within a single nucleus. Therefore, the territory was taken as the basic unit of observation.

**RT-PCR**

5 μg RNA extracted from cells using Bio/RNA-XCell (Bio/Gene) were reverse transcribed at 42°C for 2 h using 20 U M-MuLV reverse transcriptase (BCL) in a volume of 20 μl containing 50 mM Tris, pH 8.3, 7.5 mM KCl, 3 mM MgCl2, 10 mM DTT, 20 U RNaase inhibitor (BCL), and 200 ng random hexanucleotides (BCL). Identical reactions without reverse transcriptase were performed as a control (−RT). The products of both reactions were then amplified with primer pairs to each of the four genes of the human WAGR locus in a total of 25 μl for the PANBNE gene and 50 μl for the other three genes. For WT1, D609 from exon 1 and D610 from exon 6 amplify a 686-bp fragment (Williamson, 1996). For RCN, primers G829 and G830 amplify a 795-bp fragment containing most of the coding region (Kent et al., 1997); PAX6 primers B251 (from exon 7) and C402 (exon13) amplify a 898-bp fragment (Hanson et al., 1993). For PANBNE, primers 023 (5′-CTCTTCTTCGACTGAGCAC-3′) from exon 7 and 041 (5′-TCCACAGGTTCAAGGCG-3′) from exon 10 amplify a 395-bp fragment using the following conditions: 94°C for 5 min and then (94°C for 45s, 56°C for 45 s, and 72°C for 45 s) × 10 (D.A. Kleijnan, personal communication).

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