Targeted Deficiency of the Transcriptional Activator Hnf1α Alters Subnuclear Positioning of Its Genomic Targets

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Abstract

DNA binding transcriptional activators play a central role in gene-selective regulation. In part, this is mediated by targeting local covalent modifications of histone tails. Transcriptional regulation has also been associated with the positioning of genes within the nucleus. We have now examined the role of a transcriptional activator in regulating the positioning of target genes. This was carried out with primary β-cells and hepatocytes freshly isolated from mice lacking Hnf1α, an activator encoded by the most frequently mutated gene in human monogenic diabetes (MODY3). We show that in Hnf1α−/− cells inactive endogenous Hnf1α-target genes exhibit increased trimethylated histone H3-Lys27 and reduced methylated H3-Lys4. Inactive Hnf1α-targets in Hnf1α−/− cells are also preferentially located in peripheral subnuclear domains enriched in trimethylated H3-Lys27, whereas active targets in wild-type cells are positioned in more central domains enriched in methylated H3-Lys4 and RNA polymerase II. We demonstrate that this differential positioning involves the decondensation of target chromatin, and show that it is spatially restricted rather than a reflection of non-specific changes in the nuclear organization of Hnf1a-deficient cells. This study, therefore, provides genetic evidence that a single transcriptional activator can influence the subnuclear location of its endogenous genomic targets in primary cells, and links activator-dependent changes in local chromatin structure to the spatial organization of the genome. We have also revealed a defect in subnuclear gene positioning in a model of a human transcription factor disease.

Introduction

The recognition of nucleotide sequences in the vicinity of genes by DNA binding factors is central to the regulation of gene-specific transcription [1]. The mechanism by which DNA binding transactivators lead to gene activation is in part dependent on their ability to promote the remodeling of chromatin structure and the covalent modification of nucleosomal histone tails [1–3].

Numerous studies have linked different covalent histone modifications with the transcriptional state of gene loci [4]. Amongst these, the methylation of H3-Lys4 at gene promoters has been linked to gene activity [5,6], whereas transcriptional silencing correlates with increased methylation of H3-Lys9 or H3-Lys27 [7–10]. For example, trimethylated histone H3 Lysine 9 (H3-Lys9me3) is enriched at pericentromeric repeats forming constitutive heterochromatin [9,11], while trimethylated H3-Lys27 (H3-Lys27me3) has been linked to other forms of inactive chromatin, including chromosome X facultative heterochromatin, imprinted loci, and Polycomb-mediated silencing of homeobox gene clusters [7,8,10].

It has recently become apparent that the positioning of gene loci within the three dimensional structure of the nucleus may provide a further level of regulation [reviewed in [2,12,13]]. Gene activation has been linked to selective looping of loci away from chromosome territories [14,15], and appears to be associated with an increased likelihood that a locus intermingles with heterologous chromosome territories [16]. Transcribing genes have also been shown to colocalize with nuclear domains that are visibly enriched in RNA polymerase II [17,18]. Other observations revealed that active loci localize in the nuclear interior, whereas inactive genes have been found to be preferentially positioned at the nuclear periphery [19,20]. Moreover, repositioning to centromeric regions has been shown for several hematopoietic genes during differentiation-related silencing, and correlates with mutation-induced silencing of the brown locus in Drosophila [21–24].

The precise relationships between gene positioning and transcriptional regulation, however, are not understood. Some studies suggest that gene compartmentalization may play a decisive regulatory role. An example is the demonstration that artificial recruitment of genes to the nuclear lamina results in transcriptional repression in certain, though not all, experimental settings [25,26]. On the other hand, several studies show that gene-rich regions tend to locate outside of their respective chromosome territories or occupy more central
nuclear positions [27–30]. Therefore, the extent to which a gene’s subnuclear position in a given cell type depends on its gene-specific transcriptional activity or on the regional organization of the chromatin varies [37–40]. However, in this process, we have employed a genetic approach. We used mice lacking Hnf1α, a transcription factor gene that is mutated in an inherited form of diabetes. We studied genes that are directly bound by Hnf1α, as well as various control genomic regions, and determined their position in nuclear space in liver and insulin-producing β-cells. The results showed that the absence of Hnf1α causes local changes in the chromatin of target genes. At the same time, it modifies the position of target genes in nuclear space. The findings of this study lead us to propose a model whereby transcription factor dependent local chromatin modifications are linked to subnuclear gene positioning. They also revealed abnormal subnuclear positioning in a model of a human transcription factor disease.

Author Summary

All cells in an organism share a common genome, yet distinct subsets of genes are transcribed in different cells. Selectivity of gene transcription is largely determined by transcription factors that bind to target genes and promote local changes in chromatin. Such changes are thought to be instrumental for transcription. Emerging evidence indicates that the position of genes in the three-dimensional structure of the nucleus may also be important in transcriptional regulation. However, the role of transcription factors in gene positioning, and its possible relationship with chromatin modifications, is poorly understood. To examine this, we employed a genetic approach. We used mice lacking Hnf1α, a transcription factor gene that is mutated in an inherited form of diabetes. We studied genes that are directly bound by Hnf1α, as well as various control genomic regions, and determined their position in nuclear space in liver and insulin-producing β-cells. The results showed that the absence of Hnf1α causes local changes in the chromatin of target genes. At the same time, it modifies the position of target genes in nuclear space. The findings of this study lead us to propose a model whereby transcription factor dependent local chromatin modifications are linked to subnuclear gene positioning. They also revealed abnormal subnuclear positioning in a model of a human transcription factor disease.

Results

Hnf1α Alters Histone Methylation Patterns and Chromatin Compaction of Its Target Genes

Earlier studies showed that Hnf1α-dependent transcription is dependent on the recruitment of histone acetyltransferases and the local acetylation of nucleosomal histones [40,41]. We have now examined the methylation state of histone H3 in target genes. For this analysis we selected the most profoundly downregulated genes identified in expression profiling experiments of Hnf1α−/− hepatocytes (Afh, Cyp2j5 and Pah) and islets (Kis12), all of which are specifically downregulated in their respective Hnf1α-deficient cell-types (Figure 1A). The four genes contain evolutionary conserved high-affinity Hnf1 binding sites in their promoter regions, and were experimentally shown to be directly bound by Hnf1α (Figure 1B and not shown).

As shown in Figure 1C–D, dimethylated H3-Lys4 (H3-Lys4me2) was decreased in the 5′ region of such genes in hepatocytes from Hnf1α-deficient mice, while no changes were observed in control genes.

H3-Lys9me3, an established repressive mark associated with constitutive heterochromatin [9], was not increased in the 5′ region of these genes in Hnf1α−/− hepatocytes (Figure 1E), but was readily detected in minor satellite positive control sequences (data not shown).

In contrast, methylated H3-Lys27 was increased in Hnf1α-dependent targets in Hnf1α−/− hepatocytes to a similar extent as in two constitutively silenced genes known to be enriched in this repressive mark (Nanog and Hoxa9), whereas no changes were observed in non Hnf1α-dependent control genes (Figure 1F–H and not shown). Increased methylated H3-Lys27 was primarily the trimethylated form, as it was detected with selective antisera for H3-Lys27me2,3 and H3-Lys27me3, but not H3-Lys27me2 (Figure 1F–H and not shown). Interestingly, increased H3-Lys27me3 was spread throughout the Cyp2j5 locus, rather than being circumscribed to discrete segments (Figure 1H). Dimethylated H3-Lys9, another histone mark previously associated with facultative heterochromatin, was also increased by 3.5 to 5-fold in inactive Hnf1α-targets in Hnf1α−/− cells (data not shown).

We also examined the consequences of Hnf1α-deficiency on target chromatin condensation. General DNase I sensitivity studies revealed reduced degradation of Cyp2j5 chromatin in Hnf1α−/− vs. Hnf1α+/− hepatocytes, whereas no differences were observed between genotypes for the control gene Actb (Figure 1I). Thus, in direct Hnf1α target genes that are inactive due to Hnf1α-deficiency, there is a switch from an active chromatin conformation enriched in methylated H3-Lys4, to a more compacted state enriched in trimethylated H3-Lys27.

Different Isoforms of Methylated Histone H3 Are Distributed Non-Randomly in Nuclear Space

To explore possible relationships between Hnf1α-dependent gene activity, site-specific histone modifications, and nuclear organization, we first assessed subnuclear distributions of histone modifications in primary hepatocytes and pancreatic islet-cells. Both of these cell types are largely quiescent under normal cellular mitotic conditions, and we therefore examined the nuclear occupancy of subnuclear compartments containing the euchromatin marks H3-Lys4me2 and H3-Lys9me3, as well as the heterochromatin mark H3-Lys27me3.

The Histone Code in Nuclear Space

Hnf1α-mediated changes in histone methylation patterns were assessed using immunofluorescence microscopy and quantitative histone mark enrichment analysis. The results showed that Hnf1α deficiency leads to decreased dimethylated H3-Lys4 (H3-Lys4me2) and increased trimethylated H3-Lys27 (H3-Lys27me3) in target genes, as compared to control genes. These changes are thought to be mediated by Hnf1α-dependent chromatin remodeling events. In contrast, Hnf1α deficiency leads to increased trimethylated H3-Lys9 (H3-Lys9me3) in response to DNA damage, as compared to control genes. These changes are thought to be mediated by Hnf1α-dependent transcriptional repression events. The results provide novel insights into the function of Hnf1α in regulating gene expression and cellular identity in hepatocytes and pancreatic islet-cells.
Figure 1. Inactive Hnf1α targets in Hnf1α−/− cells exhibit local enrichment of H3-Lys27me3, decreased H3-Lys4me2, and reduced DNAse I sensitivity. (A) RNA expression of tissue-specific Hnf1α targets (Afm, Cyp2j5, Pah, Kif12) and control genes (Tbp, Actb) in Hnf1α−/− and Hnf1α+/− liver and islets. (B) ChIP analysis of Hnf1α occupancy in the promoter region of tissue-specific targets and a control gene (Nanog) in Hnf1α−/− and Hnf1α+/− hepatocytes (Afm, Cyp2j5 and Pah, black and white bars respectively) and MIN6 beta-cells (Kif12, black bars). Results are normalized by Tbp enrichment. *p<0.05 and **p<0.01 relative to Nanog. (C–H) ChIP analysis of histone modifications in Hnf1α-targets and control genes in Hnf1α−/− and Hnf1α+/− hepatocytes. For all genes the 5′ flanking regions are analyzed, except in D,G, where the entire Cyp2j5 locus is analyzed. Blue horizontal lines indicate amplicon positions, grey boxes are exons, red lines depict computationally predicted high-affinity HNF1 binding sites, and an arrow indicates the transcription start site. Graphs depict mean ± SEM of the ratio of the percent input immunoprecipitated with anti-methyl specific H3 relative to anti-H3 antibodies in 3 independent experiments. Black bars represent Hnf1α+/− and white bars Hnf1α−/− hepatocytes. *p<0.05 and **p<0.01 relative to Hnf1α+/− hepatocytes. (I) General DNAse sensitivity of Cyp2j5. Representative PCR analysis of Cyp2j5 and Actb 5′ flanking and coding sequence regions after digestion of Hnf1α+/+ and Hnf1α−/− nuclei with increasing amounts of DNAse I. Results show reduced DNAse I sensitivity in Hnf1α−/− nuclei in the Cyp2j5 region, but not in the Actb control locus.

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conditions. The results showed that H3-Lys4me2-rich subnuclear regions displayed a high degree of colocalization with regions that are enriched in RNA polymerase II phosphorylated on serine 5 of the C-terminal repeat, the predominant polymerase form in the transcriptional initiation complex [42] (hereafter referred to as RNA polymerase II) (Figure 2A). In sharp contrast, gene-silencing marks H3-Lys9me3 and Lys27me3 were more abundant in regions that were not enriched in RNA polymerase II (Figure 2A). These subnuclear distributions were independent of the fixation and processing methods used, and were observed with different H3-Lys27me3 antibodies (Figure S1A,B). Furthermore, the H3-Lys27me3 immunostaining pattern was distinct from that of Histone H3 and other modifications including H3-Lys4me2, H3-Lys27me1, H3-Lys27me2, H3-Lys9me3, as well as the DNA stain TO-PRO-3, indicating that it does not merely reflect chromatin density (Figure S1C–F, Figure S2, and not shown).

We next examined the radial distribution of histone modifications. H3-Lys27me3 was markedly enriched whereas H3-Lys4me2 displayed relative depletion in the immediate vicinity of the inner nuclear membrane, as shown by co-immunostaining of Lamin A/C (Figure 2B). Erosion analyses using non-thresholded images furthermore revealed markedly different radial enrichment patterns for RNA polymerase II, H3-Lys4me2, and H3-Lys27me3 (Figure 2C). Thus, RNA polymerase II and H3-Lys4me2 were significantly depleted in peripheral nuclear zones compared to more interior nuclear regions (Figure 2C, ANOVA p values $5.4 \times 10^{-10}$ and $8.9 \times 10^{-23}$). In contrast, H3-Lys27me3 was significantly enriched in the outermost zones, compared to more internal regions (Figure 2C, ANOVA p value $8.7 \times 10^{-10}$).

These results are largely consistent with recent studies describing distinct nuclear patterns of histone modifications in cultured cell lines [43], but extend it by showing that H3-Lys4me2 exhibits preferential colocalization with RNA polymerase II in central nuclear domains, while H3-Lys27me3 is particularly abundant in peripheral domains lacking enrichment in RNA polymerase II, H3-Lys4me2, or H3-Lys9me3.

Hnf1α-Regulated Gene Loci Display Differential Association with Distinct Subnuclear Domains in Hnf1α+/− versus Hnf1α−/− Cells

Immunofluorescence analysis indicated that Hnf1α is clearly enriched in H3-Lys4me2- and RNA polymerase II-rich, H3-Lys27me3-poor subnuclear domains, suggesting that there might be a subnuclear compartmentalization of Hnf1α function (Figure S3). We therefore tested if Hnf1α promotes not only changes in

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**Figure 2.** Methylated histone H3 marks exhibit a non-random subnuclear distribution. (A) Dual immunofluorescence confocal analysis of domains enriched in H3-Lys4me2, H3-Lys9me3, and H3-Lys27me3 (red) compared with RNA polymerase II (green) in interphase wild-type hepatocyte nuclei. Adjacent Venn diagrams display percentages of colocalization (mean ± SEM from 20 nuclei) of signals exceeding the 75th percentile of nuclear signal intensity in wild-type nuclei. (B) Triple immunofluorescence of RNA polymerase II (red), lamin A/C (green) and either H3-Lys4me2 or H3-Lys27me3 (blue) in hepatocytes. Insets below show peripheral nuclear segments at higher magnification. (C) Erosion analysis of the nuclear distribution of RNA polymerase II, H3-Lys4me2, or H3-Lys27me3. Nuclei were subdivided into 5 concentric zones, and total nuclear fluorescence intensities were determined for each zone (mean ± SEM). The values were normalized to the relative nuclear areas occupied by the different zones, so that a value of 1 was obtained if the percentage is as expected in case of unbiased distribution. At least 20 nuclei were analyzed in each case. Significance values for the comparison between the 5 zones for each of the 3 epitopes were obtained by ANOVA.

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site-specific histone modifications, but also in the subnuclear positioning of its targets relative to histone modification domains. To address this question we performed DNA immuno-FISH experiments and non-thresholded images were analyzed to determine the enrichment of defined histone modifications and RNA polymerase II at Hnf1α-dependent loci in control vs. null-mutant nuclei (Figure 3 and Figure S4). Importantly, the compartmentalization of histone marks was conserved after the immuno-FISH procedure, and the spatial patterns of histone modifications were unaltered in Hnf1α−/− cells (Figure S1A and Figure S5).

In several studies it has been observed that gene silencing is associated with relocation to constitutive heterochromatin domains enriched in satellite repeat sequences [21–24,44]. In mice, H3-Lys9me3 is enriched at pericentromeric regions [11]. In agreement, this was also observed under the conditions used here, where pericentromeric regions clustering at chromocenters were highlighted by TO-PRO-3 staining (Figure S2). However, inactive Hnf1α-targets in mutant hepatocytes (Cyp2j5) and islets (Kif12) were not positioned in domains that are enriched in H3-Lys9me3 or TO-PRO-3 as compared to wild-type cells (Figure 3J, Figure S2D, and not shown). Furthermore, the distance of Cyp2j5 to H3-Lys9me3-rich chromocenters, and the frequency with which the two were in contact, did not differ in wild-type vs. null-mutant cells (0.94 ± 0.10 vs. 0.88 ± 0.09 μm, and 10% vs. 7.7%, respectively). Thus, Hnf1α-deficiency does not result in repositioning of inactive Hnf1α-targets to pericentromeric heterochromatin clustering at chromocenters enriched in H3-Lys9me3.

In sharp contrast, Cyp2j5 alleles in activator-deficient cells were positioned in nuclear domains that are relatively enriched in H3-Lys27me3 (Figure 3K). Analogous results were observed for the pancreatic islet Hnf1α-dependent gene Kif12 (Figure S4I). These observations were specific for Hnf1α-dependent loci because they were not observed in 4 control loci in hepatocytes (Hnf1b, Ly9, Actb and Nanog) (Figure 3F and Figure S6) or one control locus in islet-cells (Figure S4E). Furthermore, silent Cyp2j5 and Kif12 loci in Hnf1α-deficient cells were located in subnuclear domains with decreased H3-Lys4me2 and RNA polymerase II (Figure 3LM and Figure S4JK).

Simultaneous imaging of two protein marks at each locus allowed us to more accurately assess the extent to which loci were differentially positioned in domains enriched in distinct marks. We found that the average ratio of non-thresholded H3-Lys27me3/ubiquitinated H3-Lys4me2 was 2.6 and 3.3 times more frequently in Hnf1α−/− vs. Hnf1α+/+ cells, respectively (Figure 3P and Figure S4N). This finding did not reflect just deocclusion from RNA polymerase II-rich domains, as Cyp2j5 and Kif12 in null-mutant cells were not more frequently in RNA polymerase II-poor/H3-Lys27me3-poor domains (Figure 3P and Figure S4N), nor in RNA polymerase II-poor/H3-Lys9me3-rich or RNA polymerase II-poor/TO-PRO-3-rich domains (data not shown). The results remained significant using the median (50th percentile) of nuclear epitope intensity as an alternate threshold to define epitope enrichment (2.6-fold and 2.2-fold increased presence of Cyp2j5 and Kif12 in H3-Lys27me3-rich/RNA polymerase II-poor domains in Hnf1α−/− vs. Hnf1α+/+ cells, respectively; Fisher’s exact test, p<0.01). Differences were again not observed in four control genes using similar criteria (Figure 3O, Figure S4M, and not shown).

In concordance with the preferential nuclear compartmentalization of Hnf1α in RNA polymerase II- and H3-Lys4me2-rich domains (Figure S3), target loci were also preferentially localized in Hnf1α-rich domains in hepatocytes, in contrast to the inactive control locus Ly9 (Figure S7, and not shown). However, we found no evidence that this preferential localization reflected the existence of an activator-specific subnuclear domain, because an Hnf1α-independent active control gene (Actb) exhibited a similar subnuclear compartmentalization with Hnf1α as Cyp2j5 (Figure S7).

We also compared the radial positioning of Hnf1α-dependent loci by erosion analysis in wild-type vs. mutant hepatocytes and islet cells, respectively. In contrast to the unchanged radial positioning of the control locus Ly9, significantly increased percentages of Cyp2j5 and Kif12 alleles localized in the most peripheral nuclear zone, where H3-Lys27me3 is mostly enriched, in mutant nuclei compared to wild-type (p = 0.002 and p = 0.01, respectively) (Figure 4). Conversely, a significant decrease in the number of Kif12 loci in mutant cells was observed in the interior shell 3 (p = 0.04) (Figure 4).

Thus, in the presence of Hnf1α its direct target genes Cyp2j5 and Kif12 are positioned in more central nuclear domains enriched in RNA polymerase II and H3-Lys4me2, whereas in the absence of Hnf1α inactive targets are positioned in more peripheral, H3-Lys27me3-rich domains. Interestingly, these subnuclear histone modification enrichment patterns parallel those observed locally in Hnf1α-target nucleosomes.

Spatial Resolution of Hnf1α-Dependent Positioning

Altered positioning of Hnf1α targets in null mutant cells could represent a localized activator-dependent phenomenon, or a more global effect of Hnf1α-deficiency on the configuration of nuclear structures. To address the mechanisms involved, we performed two-color DNA FISH using contiguous BAC probes mapping to sites adjacent to the Cyp2j5 locus (Figure 5). Despite their proximity, signals from adjacent clones could be clearly separated by dual FISH analysis in a substantial number of nuclei (Figure 5A), thus enabling us to test how genomic regions in the vicinity of Cyp2j5 were positioned relative to subnuclear domains in wild-type and mutant cells. To assist the interpretation of results, we first analyzed the gene content in these regions. We noted that there were two additional Hnf1α-dependent genes immediately centromeric to Cyp2j5, while an extensive telomeric region was completely devoid of any experimentally defined spliced transcripts (Figure 5B, Table S2). Parallel ImmunoFISH studies showed that unlike Cyp2j5, the adjacent regions 12L1, 68H9, and 114C9 were not differentially distributed with respect to nuclear RNA polymerase II or histone marks domains in Hnf1α−/− vs. Hnf1α+/+ cells (Figure 5C,F and G). Nonetheless, the region marked by clone 263F12 that is in immediate proximity to the Hnf1α-dependent gene (Cyp2j6) did show differential positioning similar to Cyp2j5 (Figure 5E, Table S2). These findings indicated that Hnf1α-dependent positioning of Cyp2j5 into histone modification/RNA polymerase II subnuclear domains is a locally restricted phenomenon, encompassing a somewhat extended domain of up to 300 Kbp containing at least two additional coordinately regulated Hnf1α-dependent genes.

We next sought to determine if Hnf1α-dependent positioning of the Cyp2j5 locus can be elicited relative to adjacent genomic regions, thus providing reference points that are independent of histone mark and RNA polymerase II spatial distributions. We used two-color DNA FISH to measure the distance of Cyp2j5 to
adjacent loci in wild-type vs. null-mutant cells. We found that the distance between Cyp2j5 and the two telomeric clones 68H9 and 114C9 was significantly increased in wild-type compared with Hnf1a+/2 hepatocytes (0.36 ± 0.02 vs. 0.28 ± 0.02 μm, and 0.46 ± 0.02 vs. 0.36 ± 0.02 μm, respectively, Mann-Whitney test p < 0.001; Figure 5J,K). The distance between Cyp2j5 and the most proximal centromeric 263F12 region was not affected by Hnf1a-deficiency (in keeping with the lack of differences in RNA polymerase II/K27me3 colocalization studies), but for the more distal clone 12L1 it was decreased from 0.40 ± 0.03 μm in wild-type cells to 0.29 ± 0.02 μm in Hnf1a+/2 cells (Mann-Whitney test p < 0.001; Figure 5H–I). Accordingly, the percentage of non-overlapping loci, which was systematically defined as those located at >0.4 μm center to center distance, was higher in wild-type vs. null mutant cells for Cyp2j5-68H9 (39 vs. 17%, Fisher’s exact test p < 0.05), Cyp2j5-114C9 (57 vs. 42%, p < 0.05), and Cyp2j5-12L1 (35 vs. 24%, p < 0.05) comparisons (Figure 5H–K). In contrast, the distances separating 68H9 and 114C9, which do not contain Hnf1α-dependent genes, do not differ between control and null-mutant cells (Figure 5L). Thus, Cyp2j5 showed altered Hnf1α-dependent positioning relative to neighboring centromeric and telomeric chromosomal regions.

Figure 3. Hnf1α-dependent Cyp2j5 activity correlates with differential positioning in RNA polymerase II and histone code domains. (A–D) Representative confocal immuno-FISH analysis in Hnf1α+/+ and Hnf1α−/− hepatocytes of the Cyp2j5 locus (red) with RNA polymerase II (RNA Pol II, green) and either H3-Lys4me2 (A,B) or H3-Lys27me3 (C,D) (blue). The framed regions containing Cyp2j5 FISH signals are shown at higher magnification on the right of each panel with omission of blue or green channels (E–N) Quantitative analysis of histone marks and RNA polymerase II in Cyp2j5 (J–N) and Ly9 control (E–I) loci in Hnf1α+/+ and Hnf1α−/− hepatocytes. For each condition, non-thresholded fluorescence intensities of histone marks and RNA polymerase II were measured at 70–200 FISH signals, and each value was divided by its nuclear median intensity in the same channel. The graphs thus depict the average of such normalized signal values ± SEM, except in I.N which shows mean ± SEM of H3-Lys27me3/RNA polymerase II ratios (mK27/PoIi) calculated for each allele. (O–P) Classification of Cyp2j5 (P) and Ly9 control (O) alleles into 4 categories according to the simultaneous enrichment (+) or non-enrichment (−) of RNA polymerase II (RNA Pol II) and H3-Lys27me3 (K27me3) in Hnf1α+/+ (black bars) and Hnf1α−/− (white bars) hepatocytes. Each allele was scored as enriched (+) or non-enriched (−) based on whether or not the signal intensity exceeded the 75th percentile of nuclear signals. Alternate thresholds such as the nuclear median yielded comparably significant results (see text). Results are expressed as % of all alleles for each genotype. *p < 0.05 and **p < 0.01 relative to Hnf1α+/+ cells using Mann-Whitney or Fisher’s exact test as appropriate.
Figure 5. Spatial resolution of Hnf1α-dependent repositioning. (A) Two-color DNA FISH detection of adjacent loci Cyp2j5 (red) and 68H9 (green) in Hnf1α+/+ hepatocytes. (B) Schematic representation of the relative positions and distances (Kb) of BACs located centromeric (12L1 and 263F12) and telomeric (68H9 and 114C9) to Cyp2j5. Hnf1α-dependent genes in the region are drawn schematically in red. Note that no spliced transcript has been mapped to the region encompassed by BACs 68H9 and 114C9. (C–G) Classification of 12L1 (C), 263F12 (D), Cyp2j5 (E), 68H9 (F), and 114C9 (G) alleles into 4 categories according to the simultaneous enrichment (+) or non-enrichment (−) of RNA polymerase II (RNA Pol II) and H3-Lys27me3 (K27me3) in Hnf1α+/+ (black bars) and Hnf1α−/− (white bars) hepatocytes as described in Figure 3O–P. (H–L) Comparison of distances
between the indicated BAC clone FISH signals in Hnf1α+/+ and Hnf1α−/− hepatocytes. For each comparison, the upper panels show the mean±SEM interlocus (signal center to center) distances in μm, with significance values calculated with the Mann-Whitney test. Lower panels show the percentage of non overlapping alleles, defined as those with signal center distances exceeding 0.4 μm, with significance values assessed with Fisher’s exact test. *p<0.05, **p<0.01 relative to Hnf1α+/+ cells. More than 100 nuclei were analyzed in each case.
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We further assessed Hnf1α-dependent positioning of Cyp2j5 with respect to its chromosomal territory. We observed that Hnf1α-deficiency did not affect the position of the nearby 114C9 genomic region relative to its chromosomal territory, whereas Cyp2j5 alleles less frequently extended away from their territory surface in Hnf1α−/− cells versus wild-type cells (Figure S8). Collectively, these findings reveal the existence of Hnf1α-dependent, spatially restricted positioning of a target locus relative to chromosomal reference landmarks and subnuclear RNA polymerase II/histone modification domains. The analysis of distances between adjacent regions and relative to the chromosome territory furthermore indicates that Hnf1α-dependent positioning involves chromatin decondensation of the Cyp2j5 locus.

Discussion

Genetic Evidence for Activator-Dependent Gene Positioning

We have used a genetic model to show that a transcriptional activator regulates the subnuclear positioning of its direct endogenous targets in primary differentiated cells. We documented Hnf1α-dependent differential gene positioning with respect to: a) subnuclear regions enriched in H3-Lys27me3, H3-Lys4me2, and phosphoserine-5 RNA polymerase II (Figure 3), b) radial nuclear zones (Figure 4), c) genomic regions adjacent to an Hnf1α-dependent gene (Figure 5), and d) chromosomal territories (Figure S8). The analysis of four control loci in trans allowed us to conclude that the observed Hnf1α-dependent spatial changes are specific. Experiments comparing the position of an Hnf1α-dependent locus to adjacent chromosomal regions and its chromosomal territory further demonstrated specificity, and revealed that changes were locus-selective and did not reflect broad chromosomal reconfigurations.

Although numerous studies have shown a relationship between gene transcription and subnuclear positioning, several variables that are only indirectly related to gene transcription, such as regional gene density or nucleotide composition, also appear to impact the subnuclear location of genomic regions, independent of their actual transcriptional activity [27–29,45]. Our new findings demonstrate that transactivator-dependent functions are dominant over such variables in the regulation of subnuclear gene positioning.

Earlier reports have linked the function of sequence-specific DNA binding proteins such as Ikaros and NF-E2p18 with the repositioning of endogenous loci [21,46,47]. In such examples, repressor-mediated repositioning of silenced loci to pericentromeric compartments was observed during developmentally regulated gene-silencing processes. This clearly represents a different situation compared to the current analysis where gene inactivity results from the sheer lack of an activator and gives rise to a different pattern of subnuclear positioning that does not involve association with chromocenters.

Previous evidence supporting the role of transactivators in gene positioning comes from studies of transgenes. Some of these studies took advantage of a lac repressor-VP16 acidic activation domain fusion protein, which was shown to cause repositioning of targeted multicopy loci away from the nuclear periphery [31,32]. Another study has analyzed transgenes with intact or mutated transactivator binding sites and showed that intact sites prevent association of transgenes with pericentromeric heterochromatin [48]. The role of transactivators in the positioning of endogenous loci, however, has not been directly assessed. One study showed that the deletion of a 24 Kb endogenous genomic region containing the β-globin locus control region results in gene silencing and increased perinuclear localization of the endogenous locus [49]. These effects were probably due to activator functions because the deleted region contained multiple binding sites for essential transcription factors. Nevertheless, it could not be excluded that structural changes due to deletion of an extended genomic segment also affected nuclear positioning by transactivator independent mechanisms. Our results provide genetic evidence in primary cells that positioning of endogenous genes can be dependent on a single transactivator. Together with previous studies, this suggests that the regulation of the subnuclear location of target gene loci might be a general function of sequence-specific DNA binding transcriptional regulators.

Gene Positioning Relative to Subnuclear Domains

Earlier studies describing correlations between gene silencing and perinuclear positioning were based on the comparisons of different cell types or developmental stages [19,20,50,51]. Such studies can theoretically be confounded by cell-specific differences in global spatial chromosomal arrangements [39]. It is thus important that peripheral positioning is now elicited in a model where transcriptional inactivity is ascribed to the selective absence of a direct transactivator.

Previous studies have also shown that genes are preferentially transcribed in nuclear subdomains enriched in RNA polymerase II [17,52,53]. This has led to models postulating that active loci loop into domains with high local RNA polymerase II concentrations [17,54]. Our findings confirm that gene activity is associated with localization to phosphoserine-5 RNA polymerase II domains in primary cells, and furthermore demonstrate that association with such domains is linked to the function of a transcriptional activator. Importantly, the new results extend our understanding of this phenomenon by showing that relocation does not only occur with respect to domains enriched in RNA polymerase II, but also involves repositioning amongst compartments that differ in the composition of histone modifications known to be critically involved in transcriptional regulation, and that such domains display distinct radial distributions. Our integrated analysis of a transactivator-deficient model thus suggests that transcription-related gene positioning with respect to RNA polymerase II foci, distinct radial nuclear zones, and domains enriched in specific histone modifications might reflect different experimental measurements of a single biological phenomenon.

Relationships between Hnf1α-Dependent Locus Specific Chromatin Changes and Subnuclear Positioning

Together with previous findings, our data shows that binding of Hnf1α to target loci promotes local histone tail hyperacetylation, methylation of H3-Lys4, and chromatin decondensation, while preventing methylation at H3-Lys27 [38,40,41]. H3-Lys27 methylation thus appears to represent a default state, consistent with genetic studies showing that the H3-Lys4-specific methyltransferase Trithorax suppresses default gene silencing mediated
the histone modification enrichment pattern of Hnf1 and relative to chromosomal territories. Our findings also show that locus that is reflected by changes in distances between adjacent loci chromatin decompaction, but also the decondensation of the Lys4me-rich/H3-Lys27me-poor domains in a possible mechanism. Histone modifications may be partly instrumental in gene positioning. Although similar measurements of locus positioning relative to histone modification domains have not been carried out before, two studies previously showed that treatment with histone deacetylase inhibitors causes repositioning of inactive genes away from the nuclear periphery [20,56]. Local histone modifications could affect compartmentalization of gene loci by regulating interactions with the nuclear lamina [56] and could also affect mobility, since acetylated histones have been previously shown to increase chromatin fiber flexibility [57]. Taken together, these findings support the proposal that local Hnf1-dependent chromatin decompaction and histone modifications might result in augmented mobility and loop formation, thus increasing the likelihood of accessing and establishing dynamic interactions with components of transcriptionally active nuclear regions (Figure 6). Local activator-dependent changes in chromatin structure may thus play a role in regulating the spatial organization of the genome.

Emerging evidence indicates that gene transcription is an integrated process involving multiple levels of regulation [2,3]. The data presented here link the in vivo function of an activator to different levels of regulation, namely the binding to specific target sequences, the local modification of target chromatin, and the positioning of targets in distinct subnuclear domains. This demonstration is provided in a genetic model of human diabetes, indicating cellular defects at multiple regulatory levels in a human transcriptional disease. Thus, our findings provide not only new insights into the complexity of trans-activator functions and transcriptional regulation, but are also important for understanding mechanisms underlying human disease.

**Material and Methods**

**Cell Preparation**

Hepatocytes and pancreatic islets were isolated from 4–6 week-old Hnf1α+/− and Hnf1α−/− mice [36] by local perfusion of the organ with collagenase for digestion and subsequent isolation of the cells as described [34,40]. For immuno-staining and FISH studies, after isolation islets were gently dissociated for 2 min in pre-warmed trypsin solution. Cells were processed for chromatin, RNA, immunofluorescence, and FISH analysis immediately after isolation.

**RNA Extraction and RT-PCR**

RNA isolation, reverse transcription and PCR were carried out as described [34].

**General DNAse I Sensitive Assay**

Isolated hepatocytes (30–40×10^6) were resuspended in 10 mL NI buffer (15 mM Tris-HCl pH7.5, 300 mM sucrose, 15 mM NaCl, 60 mM KCl, 4 mM MgCl2 and 0.5 mM DTT), and 10 mL of NI buffer supplemented with 1% NP40 was added for 10 min incubation in ice. Nuclei were collected at 500g for 3 min, washed in 4 mL NI buffer, resuspended in a final volume of 700 µL NI buffer and distributed in 100 µL aliquots for DNAse I digestion. To each suspension of 100 µL NI buffer, either 0, 10, 20, 30, 40, 50 or 80 µg DNAse I was added for 10 min on ice. The reaction was stopped with Nuclei Lysis Solution from Wizard Genomic DNA purification kit (Promega), and DNA was extracted as indicated by the manufacturer. DNAs were resuspended in 20 µL DNA rehydration solution and 1 µL was used for PCR amplification. Oligonucleotides are presented in Table S1.

**Chromatin Immunoprecipitation**

Approximately 2×10^6 isolated hepatocytes or MIN6 beta-cells were used per immunoprecipitation as described [40,58], with modifications. Immunoprecipitations were carried out before, two studies previously showed that treatment with histone deacetylase inhibitors causes repositioning of inactive genes away from the nuclear periphery [20,56]. Local histone modifications could affect compartmentalization of gene loci by regulating interactions with the nuclear lamina [56] and could also affect mobility, since acetylated histones have been previously shown to increase chromatin fiber flexibility [57]. Taken together, these findings support the proposal that local Hnf1-dependent chromatin decompaction and histone modifications might result in augmented mobility and loop formation, thus increasing the likelihood of accessing and establishing dynamic interactions with components of transcriptionally active nuclear regions (Figure 6). Local activator-dependent changes in chromatin structure may thus play a role in regulating the spatial organization of the genome.

**Figure 6. Summary model. Hnf1α+/−.** In the absence of Hnf1α, target nucleosomes exhibit increased trimethylated H3-Lys27, and are more likely to be located in peripheral nuclear domains with condensed chromatin enriched in methylated H3-Lys27. Hnf1α+/−: In wild-type cells, Hnf1α binding recruits complexes that lead to site-specific histone acetylation, H3-Lys4 methylation, and chromatin remodeling, while preventing trimethylation of H3-Lys27. This chromatin configuration is associated with relocalization to transcriptionally active, more centrally located nuclear subregions enriched in RNA polymerase II and H3-Lys4me2. We propose that activator-dependent local chromatin changes may be instrumental in gene positioning.

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by methylated H3-Lys27 [55]. Concomitant with local chromatin changes, Hnf1α binding also causes the recruitment of targets to predominantly central subdomains that are enriched in phosphoserine-5 RNA polymerase II and concordant histone modifications (see model in Figure 6).

How Hnf1α controls subnuclear positioning of its targets remains to be clarified. Treatment with the RNA polymerase II inhibitors α-amanitin and 5,6-dichlorobenzimidazole riboside (DRB) does not alter the preferential positioning of Kif12 in H3-Lys4me-rich/H3-Lys27me-poor domains in a β-cell line with normal Hnf1α expression (Figure S9). This suggests that Kif12 compartmentalization is not solely dependent on ongoing transcriptional activity per se, and points to the involvement of other activator-dependent functions.

Changes in local chromatin structure represent another potential mechanism. Our results showed that Hnf1α regulates not only local chromatin decompaction, but also the decondensation of the Cyp2j5 locus that is reflected by changes in distances between adjacent loci and relative to chromosomal territories. Our findings also show that the histone modification enrichment pattern of Hnf1α-dependent genes in nuclear domains with which they associate coincides with the local post-translational histone modification profile. This raises the possibility that histone modifications may be partly instrumental in gene positioning. Although similar measurements of locus positioning relative to histone modification domains have not been carried out before, two studies previously showed that treatment with histone deacetylase inhibitors causes repositioning of inactive genes away from the nuclear periphery [20,56]. Local histone modifications could affect compartmentalization of gene loci by regulating interactions with the nuclear lamina [56] and could also affect mobility, since acetylated histones have been previously shown to increase chromatin fiber flexibility [57]. Taken together, these findings support the proposal that local Hnf1α-dependent chromatin decompaction and histone modifications might result in augmented mobility and loop formation, thus increasing the likelihood of accessing and establishing dynamic interactions with components of transcriptionally active nuclear regions (Figure 6). Local activator-dependent changes in chromatin structure may thus play a role in regulating the spatial organization of the genome.

Emerging evidence indicates that gene transcription is an integrated process involving multiple levels of regulation [2,3]. The data presented here link the in vivo function of an activator to different levels of regulation, namely the binding to specific target sequences, the local modification of target chromatin, and the positioning of targets in distinct subnuclear domains. This demonstration is provided in a genetic model of human diabetes, indicating cellular defects at multiple regulatory levels in a human transcriptional disease. Thus, our findings provide not only new insights into the complexity of trans-activator functions and transcriptional regulation, but are also important for understanding mechanisms underlying human disease.
4°C with 7.5 μg rabbit anti-HNF1 (H-205) (Santa Cruz, sc8986), 2 μg rabbit anti-H3-Lys4me2 (Upstate, 07-030), 2 μg mouse anti-H3-Lys27me2,3 [38] (D. Reinberg, University of Medicine and Dentistry of New Jersey), 10 μg rabbit anti-H3-Lys27me3 [11] (T. Jenuwein, The Vienna Biocenter and Upstate, 07-449), and 20 μg rabbit anti-H3-Lys9me3 [11] (T. Jenuwein and Upstate, 07-442). For anti-H3-Lys4me2 and H3-Lys27me2,3, 1% Triton was added to the antibody binding solution. For anti-H3-Lys27me2,3, 3 μg rabbit anti-mouse IgG (Sigma) was added for a further 3 h incubation at 4°C. Immune complexes were collected by adsorption to protein A+G Sepharose (Amersham). Beads were washed and eluted as described, except for the anti-H3-Lys9me3 immunoprecipitation that was with 250 mM NaCl. Purified immunoprecipitated DNA was analyzed in duplicate by SYBR green real-time PCR, and compared to a standard curve generated with serial dilutions of input chromatin DNA. Oligonucleotides are shown in Table S1.

Immunofluorescence

5,10^4 isolated hepatocytes or islet-cells were lightly cytospun and fixed at room temperature for 5 min in freshly prepared 4% paraformaldehyde. In control experiments immunostaining was carried out as described [60] except that antibody retrieval was not employed. Primary antibodies were used with the following dilutions: rabbit anti-H3 (Abcam, Ab1791) (1/500), rabbit anti-H3-Lys4me2 (Upstate, 07-030) (1/500), rabbit anti-H3-Lys9me3 [11] (T. Jenuwein, and Upstate, 07-442) (1/500), rabbit anti-H3-Lys9me2 [11] (T. Jenuwein, and Upstate, 07-442) (1/100), mouse anti-H3-Lys27me3 (Abcam, Ab 6002) (1/100), rabbit anti-H3-Lys27me2,3 [11] (T. Jenuwein, and Upstate, 07-442) (1/100), mouse anti-H3-Lys4me2 (Upstate, 07-030) (1/100), rabbit anti-H3-Lys9me3 (T. Jenuwein, and Upstate, 07-442) (1/100), mouse anti-H3-Lys9me2 (Upstate, 07-442) (1/100), rabbit anti-H3-Lys9me2 (Abcam, Ab 5408) (1/1000), goat anti-Lamin A/C N18 (Santa Cruz) (1/200), mouse anti-HNF1α (Transduction Laboratories, H69220) (1/50), and rabbit anti-HNF1 (H-205) (Santa Cruz, sc8986) (1/100). The specificity of methylated H3-Lys27 stainings was verified by co-staining with two different highly specific antibodies directed against the same epitope and using alternate fixation (methanol at −20°C for 10 min). The specificity of anti-HNF1α staining patterns was verified using Hnf1α−/− cells and alternate fixation procedures (Figure S3).

DNA Immuno-FISH

We used purified BAC DNAs (Table S2) for labeling with Dig-nick translation or BioNick kits (Roche). Immuno-FISH was based on modifications of the protocol described by Brown et al [21]. Cells were immunostained essentially as described above except that nuclei were fixed in 4% paraformaldehyde for 15 min and heated in a microwave in 10 mM citrate buffer, pH 6, for 5 min before permeabilization. Immunostained nuclei were then post-fixed in 4% paraformaldehyde for 15 min, denatured in NaOH 0.1M in PBS, pH 13 for 110 sec, and washed in cold PBS and 2× SSC. One μL digoxigenin-labeled probe in 14 μL hybridization buffer (50% formamide, 2× SSC, 125 μg/mL Cot-1 and 10% dextran sulphate) and 1 μg mouse Cot-1 (Invitrogen) were denatured for 5 min at 90°C. Probes were hybridized overnight at 37°C, and sequentially washed in 2×SSC, 1×SSC, PBS-triton 0.2%, and PBS for 5 min at room temp. Slides were then sequentially incubated with Sheep anti-digoxigenin antibody (Roche) (1/300) for 3 h, and donkey anti-sheep Cy3 antibody (Jackson Immunoresearch) (1/200) for 2 h at room temp., with washes after each step. Cells were mounted with ProLong Antifade (Amersham).

Two-Color DNA FISH

Cells were fixed in 4% paraformaldehyde for 15 min at room temperature, washed in PBS and permeabilized for 30 min in PBS-0.5% Triton X-100. Cells were then heated in 10 mM citrate buffer, pH 6 for 5 min and post-fixed in 4% paraformaldehyde, washed in PBS, and incubated in 2× SSC. One μL digoxigenin-labeled probe and one μL biotin-labeled probe were added to 14 μL hybridization buffer as described above. Both probes and cells were simultaneously heated at 90°C for 5 min to denature DNA, and hybridized and washed essentially as in the immuno-FISH protocol. The digoxigenin-labeled probe was detected as in the immuno-FISH procedure, and the biotin-labeled probe was detected with AF488-streptavidin (Molecular Probes)(1/500). Cells were washed and counterstained with TO-PRO-3 (Molecular Probes)(1/50,000) and mounted with ProLong Antifade (Amersham).

Chromosome Painting

Cells were fixed and permeabilized as described above. After permeabilization, hepatocytes were incubated with 100 μg/mL RNase A at 37°C for 30 min. Nuclei were then denatured at 74°C in 2×SSC-70% formamide for 3 min followed by 1 min in 2×SSC-50% formamide. Ten μL of chromosome 4 biotin-labeled probe (Cambo) was denatured at 70°C for 10 min in the supplied buffer (Cambo) and pre-annealed for 20 min at RT. Subsequently 1 μL of either Cyp2j5 or 114C9 digoxigenin-labeled probe denatured at 90°C for 5 min was added for overnight hybridization at 37°C. After sequential washes of 5 min at 45°C in 2×SSC-50% formaldehyde, 1×SSC, PBS-0.2% Triton and PBS, biotin and digoxigenin-labeled probes were detected and processed for confocal image acquisition as described above.

Image Collection

Confocal images for each fluorochrome were acquired sequentially at room temperature with a Leica TCS SL laser scanning confocal spectral microscope, using a 63× oil immersion objective lens (NA 1.32). Focal Check Fluorescent microspheres (Molecular Probes) were used before image capture to align laser lines. Non-saturated, unprocessed images were further analyzed with ImageJ. Contrast-stretch and gamma adjustments were made using Photoshop (Adobe) only for display.

Colocalization Analysis

This analysis was carried out to determine colocalization between the most intense nuclear signals of each epitope. Ten to twenty nuclei were assessed in each double immunofluorescence experiment, and pixels with values exceeding the 75th percentile in each channel were selected for further analysis. The rationale for this threshold is that nuclear RNA polymerase II and H3-Lys27me3 signal intensities do not adhere to a normal distribution, and the 75th percentile enabled separation of visually evident RNA polymerase II and H3-Lys27me3-enriched domains from most remaining nuclear signals. Signals filtered in this manner were used to calculate Manders’ coefficient of colocalization using the appropriate ImageJ plug-in [Wayne Rasband and Tony Collins, www.uhresearch.ca/wef]. Manders’ coefficient calculates, for each channel, the proportion of colocalizing pixels respect to the summed up intensities of all pixels in the nucleus. Comparable results were obtained by subtracting pixels lower than the 50th percentile in each channel, and then applying a modified
Mander’s coefficient using the Colocalization threshold plugin, that first calculates an automated threshold (%colocalized pixels above threshold): RNA polymerase II vs. H3-Lys9me3: 4%.
RNA polymerase II vs. H3-Lys27me3: 19%; RNA polymerase II vs. H3-Lys4me2: 73%; H3-Lys27me3 vs. H3-Lys4me2: 22.5%.
H3-Lys27me3 vs. H3-Lys9me3: 31%).

Erosion Analysis
Erosion analyses were performed with single light optical sections and mid-nuclear planes were analyzed. Radial positioning of FISH signals was analyzed as described [20]. With respect to immunostaining data erosion analyses were performed as follows.
Based on the DNA counterstaining signal, the nuclear plane was subdivided into five concentric zones, each having a thickness of 20% of the nuclear radius. The numbers of pixels in each zone were counted and the grey value of each pixel was determined separately for each channel. Each grey value (I) was multiplied by its frequency N (I×N) and the sum of all values obtained for a given zone was determined (Σ I×N). The result obtained for each zone was divided by the sum of results obtained for all nuclear zones in order to determine the percentage of total nuclear fluorescence intensity in each nuclear zone. The values obtained were normalized to the relative nuclear areas occupied by the different zones. Thus a value of 1 was obtained if the percentage of the total nuclear fluorescence intensity corresponded to the percentage of the total nuclear area occupied by a given zone. It should be noted that this procedure did not involve any thresholding.

Immuo-FISH Image Analysis
For each condition, typically 100 alleles (range 70–200) from at least two independent experiments were analyzed blindly in unprocessed images to quantify signal intensities of histone marks and RNA polymerase II. In H3-Lys27me3 experiments Barr bodies were avoided. The 9-pixel area containing the brightest and most central pixels of each FISH signal was selected by inspection of single color images, and the average signal for each channel in this area was obtained using RGB Measure (ImageJ). Each non-thresholded immunofluorescence signal at a FISH-detected locus was divided by the median value of the entire nucleus in the same cell to correct for cell to cell and inter-assay technical variability. The mean intensity for each channel was also calculated from a broad cytoplasmic area in every stack and used to subtract non-specific background from both FISH and nuclear signals. This background value was similar to the non-specific nuclear signal defined as those with signal center distances exceeding 0.4 μm from the center of adjacent FISH signals defined by the 9-pixel square area unprocessed images to quantify the distance (in μm) between the center of the locus-specific FISH signal and the nearest chromosome surface was then measured as described for the two-color DNA FISH analysis. One hundred alleles from each genotype were classified as being located either within a territory and >0.4 μm from the edge, outside and >0.4 μm from the edge, or in contact if they were <0.4 μm from the edge.

Statistical Analysis
A two-tailed Student’s t-test was used for comparison of ChIP values. Mann-Whitney test was used for comparisons of immuno-FISH and allele distance values, which did not adhere to a normal distribution. ANOVA was used for erosion analysis. Fisher’s exact test was used for comparison of qualitative two-color DNA FISH, chromosomal territory, and immuno-FISH results.

Supporting Information
Figure S1 Immunofluorescence analysis of H3-Lys27me3 in hepatocytes. (A) Dual immunofluorescence analysis of RNA polymerase II (green) and H3-Lys27me3 (red) in hepatocytes after the immuno-FISH procedure showing that this process does not alter staining patterns. (B–F) Dual immunofluorescence analysis of hepatocytes with anti-H3-Lys27me3 (green) and either an alternate anti-Lys27me3 antibody (B), or anti-H3-Lys4me2 (C), anti-Lys9me2 (D), anti-Lys27me2 (E) or anti-Lys9me3 (F) antibodies (red). Colocalization analyses are depicted as Venn diagrams on the right side of immunolocalization images and were performed as described in the legend of Figure 2. Found at: doi:10.1371/journal.pgen.1000079.s001 (7.62 MB TIF)

Figure S2 Analysis of the correlation between TO-PRO-3 density and histone modification patterns and gene positioning. (A–C) Immunofluorescence analysis of H3-Lys4me2 (A), H3-Lys9me3 (B), and H3-Lys27me3 (C) enrichment (red) compared with the DNA marker TO-PRO-3 (blue). Colocalization was analyzed and depicted with Venn diagrams as described in Figure 2. (D). Quantitative analysis of H3-Lys27me3 and TO-PRO-3 enrichment in G2/M phase and H3-Lys1a+/− or H3-Lys1a−/− hepatocytes. Non-thresholded signal intensities of methylated histone marks or TO-PRO-3 were measured at 70-200 FISH alleles. To correct for cell to cell variability each value was divided by its nuclear median value, and is referred to as the normalized signal in the graphs. The graphs depict mean±SEM values. **p<0.01 relative to H3-Lys1a+/− cells. Found at: doi:10.1371/journal.pgen.1000079.s002 (4.07 MB TIF)

Figure S3 Subnuclear distribution of Hnf1a in histone code and RNA polymerase II domains. (A–B) Dual confocal immunofluorescence analysis of Hnf1a (red) and H3-Lys4me2 (A) or H3-Lys27me3 (B) (green) in interphase hepatocyte nuclei, (C–E) Dual immunofluorescence analysis of Hnf1a (red) and RNA polymerase II (green) in control hepatocytes fixed with 4% paraformaldehyde (C) or methanol (D) and in Hnf1a−/− hepatocytes fixed with 4% paraformaldehyde (E). Colocalization analysis was performed and Venn diagrams were arranged as described in the legend of Figure 2. Found at: doi:10.1371/journal.pgen.1000079.s003 (4.12 MB TIF)

Figure S4 Hnf1a-dependent Klf12 activity in islet-cells correlates with differential positioning in RNA polymerase II and histone code domains. (A–D) Representative confocal immuno-FISH analysis in Hnf1a+/− and Hnf1a−/− islet-cells of the Klf12 locus.
Classification of alleles from H3-Lys27me3/RNA polymerase II ratios (mK27/Pol II). (M,N) a

with RNA polymerase II in interphase

enrichment (Lys27me3 in signal values channel. The graphs thus depict the average of such intensities were measured at 100–200 FISH signals and each value was divided by the nuclear median intensity in the same channel. The graphs thus depict the average of such normalized signal values±SEM, except in HLL which shows mean±SEM of H3-Lys4me2/RNA polymerase II ratios (mK27/Pol II). (M,N) Classification of alleles from Lys9 control (M) and Kif12 (N) in 4 categories according to the simultaneous enrichment (+) or non-enrichment (−) of RNA polymerase II (RNA Pol II) and H3-Lys27me3 (K27me3) in Hfla1a+/− (black bars) and Hfla1a−/− (white bars) ilets as described in Figure 3O–P. Results are expressed as % of alleles for each genotype. *P<0.01 relative to Hfla1a+/+ cells using Mann-Whitney or Fisher’s exact test as appropriate.

Found at: doi:10.1371/journal.pgen.1000079.s005 (0.61 MB TIF)

Figure S5 Histone modification/RNA Polymerase II colocalization and radial distribution patterns are similar in Hfla1a+/+ and Hfla1a−/− hepatocytes. (A–C) Venn diagrams showing the colocalization of H3-Lys4me2 (A), H3-Lys9me3 (B) and H3-Lys27me3 (C) with RNA polymerase II in interphase Hfla1a+/+ and Hfla1a−/− hepatocytes. The mean±SEM percentage of colocalizing pixels of 20 nuclei is shown below. (D–G) Erosion analysis of the nuclear distribution of RNA polymerase II (D, E), and H3-Lys27me3 (F, G) in interphase Hfla1a+/+ and Hfla1a−/− hepatocytes. Erosion analyses were performed as described in the legend of Figure 2.

Found at: doi:10.1371/journal.pgen.1000079.s006 (0.61 MB TIF)

Figure S6 Quantitative analysis of histone marks and RNA polymerase II at control genes in Hfla1a+/+ and Hfla1a−/− hepatocytes by DNA immuno-FISH. Non-thresholded signal intensities were measured at 100-200 FISH signals and corrected by the nuclear median value (normalized signal) exactly as described in the legend of Figure 3. The graphs depict mean±SEM values.

*P<0.05 relative to Hfla1a+/+ cells.

Found at: doi:10.1371/journal.pgen.1000079.s007 (0.35 MB TIF)

Figure S7 Active Hnf1α-dependent and independent loci preferentially localize in Hnf1α-rich domains. (A) Quantitative analysis of Hnf1α immunofluorescence signal intensity at the Hnf1α-target locus Cyp2j5, at a non Hnf1α-target active control gene (Actb), and at an inactive locus (Lys9) in wild-type hepatocytes. Immunofluorescence signals were normalized as described in Figure 3. (B) Percentage of alleles located in RNA polymerase II-rich domains that are also located in Hnf1α enriched domains in wild-type hepatocytes. Immunofluorescence signals were normalized and categorized essentially as described in Figure 3. *P<0.05 and **P<0.01 relative to Lys9.

Found at: doi:10.1371/journal.pgen.1000079.s008 (0.22 MB TIF)

Figure S8 Hnf1α-deficiency causes altered positioning of Cyp2j5 relative to its chromosomal territory. (A–C) Representative confocal images of Cyp2j5 DNA FISH (red) and chromosome 4 paint (green) in hepatocytes counterstained with TO-PRO-3 (blue) showing alleles that are classified as being either in, out or in contact of the chromosome territory. More details on the criteria for classification are described in methods. Higher magnifications of Cyp2j5 FISH signals are shown in the upper right panels. (D–E) Percentage of Cyp2j5 (D) and 114C9 (E) alleles that are located in, out, or in contact with the chromosome 4 territory in Hfla1α+/− (black bars) and Hfla1α−/− (white bars) hepatocytes. Significance values for the comparison of allele distributions between Hfla1α+/+ and Hfla1α−/− hepatocytes were obtained by Fisher’s exact test.

Found at: doi:10.1371/journal.pgen.1000079.s009 (0.24 MB TIF)

Table S1 Oligonucleotide sequences used in chromatin immunoprecipitation analysis.

Found at: doi:10.1371/journal.pgen.1000079.s010 (0.03 MB DOC)

Table S2 Bacterial artificial chromosomes used in this study.

* Gene expression is represented as −, +, or ++ based on qualitative judgment of expression (not expressed, relative low expression, or relative high expression, respectively). ND: not determined. For genes marked in bold blue this information has been obtained experimentally by reverse transcription PCR or microarray expression analysis, while for genes marked in black this information has been collected from the Unigene EST Expression profile viewer (www.ncbi.nlm.nih.gov/entrez).

*Cyp2j6 locus is immediately adjacent but not included in the BAC clone 263F12. Note that Hnf1β mRNA is mildly increased in Hfla1α−/− hepatocytes.

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Author Contributions

Conceived and designed the experiments: RL MM NS DZ JF. Performed the experiments: RL MM NS DZ JF. Analyzed the data: RL MM NS DZ JF. Wrote the paper: RL DZ JF.

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