The Genomic Landscape of the Somatic Linker Histone Subtypes H1.1 to H1.5 in Human Cells

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SUMMARY

Human cells contain five canonical, replication-dependent somatic histone H1 subtypes (H1.1, H1.2, H1.3, H1.4, and H1.5). Although they are key chromatin components, the genomic distribution of the H1 subtypes is still unknown, and their role in chromatin processes has thus far remained elusive. Here, we map the genomic localization of all somatic replication-dependent H1 subtypes in human lung fibroblasts using an integrative DNA adenine methyltransferase identification (DamID) analysis. We find in general that H1.2 to H1.5 are depleted from CpG-dense regions and active regulatory regions. H1.1 shows a DamID binding profile distinct from the other subtypes, suggesting a unique function. H1 subtypes can mark specific domains and repressive regions, pointing toward a role for H1 in three-dimensional genome organization. Our work integrates H1 subtypes into the epigenome maps of human cells and provides a valuable resource to refine our understanding of the significance of H1 and its heterogeneity in the control of genome function.

INTRODUCTION

The linker histone H1 is a key structural component of chromatin. It binds the DNA entering and exiting the nucleosome, sealing two turns of DNA around the histone octamers (Bednar et al., 1998; Thoma et al., 1979). Because of its capacity to promote and stabilize the folding of chromatin into higher-order structures in vitro, H1 might limit the access of transcription factors to DNA and thus has been proposed to act primarily as a transcriptional repressor (Schlissel and Brown, 1984). However, this model is difficult to reconcile with knockout studies in several organisms in which only a few changes in gene expression (including both up- and downregulation) occurred (Shen and Gorovsky, 1996; Takami et al., 2000; Fan et al., 2001; Alami et al., 2003). Moreover, it was recently demonstrated that in vitro, chromatin-containing H1 is not completely inaccessible and can still be remodeled by ATPase-containing complexes (Maier et al., 2008; Clausell et al., 2009).

An additional level of complexity is added by the existence of a variable number of closely related H1 proteins in higher eukaryotes, often referred to as subtypes (Izzo et al., 2008). In humans, 11 H1 subtypes with cell-type-specific and different temporal expression have been identified, including five canonical, replication-dependent somatic subtypes (H1.1–H1.5; for a review, see Happel and Doenecke, 2009). Despite their abundance and their potential specialization, gene ablation of individual somatic H1 subtypes in mice failed to reveal an essential role, leading to the assumption that H1 subtypes are redundant (Fan et al., 2001). However, inactivation of three somatic H1 subtypes in mice demonstrated that correct H1 levels are critical for proper development and that ~50% reduction of H1 is early-embryonic lethal (Fan et al., 2005). Moreover, several biochemical studies pointed toward more individual functions for H1 subtypes; for example, H1 subtypes differ in their capacity to compact nucleosomal arrays in vitro, and bind chromatin with different affinities in vivo (Liao and Cole, 1981; Talasz et al., 1998; Orrego et al., 2007; Clausell et al., 2009; Th’ng et al., 2005). Furthermore, H1 subtypes carry distinct posttranslational modifications and interact with different regulatory proteins (Wisniewski et al., 2007; Weiss et al., 2010; Lennox et al., 1982; Montes de Oca et al., 2005; Kamieniarz et al., 2012). Additionally, in microscopy and biochemical fractionation studies, individual subtypes showed discrete patterns of localization (Parseghian et al., 2000; Th’ng et al., 2005) and, importantly, displayed varied effects on gene regulation (Sancho et al., 2008; Gunjan and Brown, 1999; Bhan et al., 2008; Brown et al., 1996), arguing for H1-subtype-specific functions. Evolutionary analysis of the rates of nucleotide substitutions of H1 subtypes supports the notion of a functional differentiation of these subtypes (Ponte et al., 1998). Thus, it is possible that compensatory mechanisms activated to recapitulate a normal H1-to-nucleosome ratio could partially mask the functional importance of H1 heterogeneity. However, due to the lack of subtype-specific antibodies, the genomic distribution of H1 subtypes in vivo has not been addressed and the role of H1 subtypes in chromatin-related processes has thus far remained elusive.
The DNA adenine methyltransferase identification (DamID) technique is a method for identifying target sequences of chromatin-binding proteins. It is based on the expression of a protein of interest fused to Dam (van Steensel and Henikoff, 2000), a bacterial methyltransferase that specifically methylates adenines within GATC sequences (Greil et al., 2006). Because adenine methylation is absent in mammals, the methylation pattern can be used to infer the binding pattern of its fusion partner (van Steensel et al., 2001). DamID is now a well-established technology and has been used successfully to characterize the genomic binding of multiple chromatin components (Greil et al., 2006; Tolhuis et al., 2006; Nègre et al., 2006; Filion et al., 2010), including Drosophila H1 (Braunschweig et al., 2009).

Based on this, we employed the DamID technique as an antibody-independent approach to systematically map the genomic distribution of all replication-dependent somatic H1 subtypes in human cells. Our analysis revealed DNA and chromatin features that potentially regulate the recruitment of H1 subtypes to chromatin. In particular CpG-dense regions and regulatory regions associated with active transcription might be critical for H1 binding. Moreover, our data indicate a possible role of somatic H1 subtypes in the three-dimensional organization of the genome within the cell nucleus. Finally, our data point toward a special role for the H1.1 subtype in chromatin function.

RESULTS

Histone H1 Subtypes Display a Similar Genomic Distribution with Dips at Promoters

In order to map the genomic distribution of the five canonical, replication-dependent linker histone H1 subtypes, we employed the DamID technique in human fibroblast IMR90 cells. We verified correct and comparable expression of H1 subtypes by RT-PCR and confirmed that fusion to Dam has no detectable effect on chromatin binding (Figures S1A and S1B; data not shown). Each data set was generated in duplicate under standardized conditions and H1 binding was detected by hybridization on NimbleGen oligonucleotide arrays. The data were normalized as described in the Materials and Methods section, and H1 binding is presented throughout this work as the log2 ratio of Dam-H1 fusion to Dam enzyme alone: log2(Dam-H1/ Dam) (Braunschweig et al., 2009). A log2 ratio of zero can be interpreted as an average steady-state level of the given H1 subtype binding to chromatin, whereas negative and positive values correspond to a relative depletion and enrichment of the H1 subtype, respectively.

A comparison of the normalized DamID profiles showed an overall similar global distribution for the H1 subtypes H1.2–H1.5 (Figure S1C), with both dips and enrichments. According to our DamID data, H1.1 seems to have a different genomic localization compared with the other H1 subtypes. Using a simple running median-based algorithm, we identified specific regions that are either depleted or enriched for each H1 subtype in our data. The average size of the depleted and enriched regions was ~1,000 bp and 800 bp, respectively. Therefore, the H1-subtype enrichments identified are distinct and localized to particular genomic regions rather than to broad domains. Annotation of the identified regions to the human genome revealed a clear overrepresentation of depleted regions of all H1 subtypes at promoters (Figure 1A). In contrast to this, we found regions enriched for all H1 subtypes overrepresented at exons. Intergenic regions are preferentially bound by H1.2–H1.5, and a small fraction of H1.1-enriched regions correspond to promoters (Figure 1A). In addition to their similar distribution throughout the genome, a closer look at individual chromosomes in the DamID data revealed unique H1 subtype DamID patterns consisting of a combination of H1 subtypes with varying degrees of enrichment (Figure 1B). For example, chromosomes 14 and 15 are relatively depleted of H1, whereas chromosomes X, 18, and 20 show higher than average H1 binding (Figures 1B and S1D). Strikingly, H1.2 appears enriched at chromosome X, with no preference for any particular DNA feature analyzed (Figures 1B, 1C, and S1E), and H1.1 on the highly transcriptionally active chromosome 19 (Figures 1B, 1C, and S1D).

In order to identify distinct combinations of H1 subtypes that are recurrent throughout the genome, we performed segmentation using an unbiased hidden Markov model (HMM) approach. We found that a five-state model optimally described our DamID data (see Materials and Methods). In this model, state 3 is the most abundant one, covering 50% of all probed locations, and corresponds to an average steady-state level of all five H1 subtypes binding (Figure 1D). State 1 corresponds to regions strongly depleted of all H1 subtypes in the DamID data sets, and shows 3.6% genome coverage. States 2, 4, and 5 show a similar level of H1.1 binding and different levels of H1.2–H1.5 enrichment. State 2, which covers 19.3% of the genome, and state 4, with 20.5% coverage, show a slight depletion and enrichment, respectively, of H1.2–H1.5. State 5, covering 6.2% of probed locations, corresponds to high enrichments of H1.2–H1.5 and highly variable H1.1 binding. The fact that none of the identified states distinguished between the subtypes H1.2–H1.5 supports the similarity of their distribution. We further corroborated this observation by independently calculating the correlation coefficients among our data sets for all somatic H1 subtypes. Indeed, the DamID results suggest the existence of two distinct groups: one consisting of H1.1 and one comprised of H1.2–H1.5 (Figure 1E). To gain insight into the molecular function of genes whose transcription start site (TSS) is found within these five HMM states, we performed a Gene Ontology (GO) analysis (Table S1). Genes within states 1 and 2 have a wide range of molecular functions and are only weakly enriched in a wide range of GO categories (e.g., DNA or protein binding), whereas genes within states 3–5 belong preferentially to fewer specialized categories.

H1 Subtypes Are Depleted at Active Promoters and Regulatory Elements

Linker histone H1 has been associated with transcriptional regulation both in vivo and in vitro (Alami et al., 2003; Bhan et al., 2008; Koop et al., 2003; Talasz et al., 1998). In order to investigate whether the depletion of H1 subtypes at promoters is linked to the transcriptional status of the corresponding genes, we divided
Figure 1. Genomic DamID Profile of Somatic H1 Subtypes
(A) Genomic annotation of regions found to be enriched and depleted of individual H1 subtypes. The length of the colored bars represents the percentage of regions enriched or depleted of H1 subtypes (as defined in Materials and Methods) overlapping with the indicated genomic features. As a control, the genomic annotation of all array probes is shown. Promoters are defined as RefGene TSSs flanked by 2 kb upstream and downstream. The genomic coordinates of exons

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CpG Islands Are Depleted of Histone H1 Subtypes

In human cells, 70% of promoters are associated with CpG islands and therefore can be classified according to their CG content (Saxonev et al., 2006). High-CpG promoters (HCPs) and low-CpG promoters (LCPs) represent strong CpG islands and clearly non-CpG islands, respectively. Intermediate-CpG promoters (ICPs) represent “weak” CpG islands (Weber et al., 2007). When we analyzed the DamID binding values of H1 at these promoter classes, we observed that the CG content negatively correlates with H1 binding values (Figures 3A and S2A). H1 subtypes tend to be depleted from HCPs (with the strongest depletion for H1.4 and the least depletion for H1.1), but not from LCPs, and an intermediate situation is present at ICP (Figures 3A and S2A). However, when they were sorted according to their expression level, it became evident that H1.2–H1.5 are mainly excluded from active and poised promoters regardless of their CG richness (Figure 3B). This inverse correlation between H1 binding values and transcription does not apply to all inactive promoters, since HCPs seem to remain H1-free even when repressed (Figure 3B). It has been shown that HCPs have a “permissive” chromatin structure and are bound by RNA-Polymerase II even when the gene they regulate is repressed (Guenther and Young, 2010). Therefore, it is possible that the presence of the transcription machinery, rather than the functional status of a promoter, is sufficient to displace H1.

Despite their frequent association with promoters, CpG islands can also be intragenic or intergenic (also called “orphan” CpG islands). With the exception of H1.1, H1 subtypes tend to be depleted from all types of CpG islands (Figure 3C). However, all H1 subtypes seem to be less depleted from inter- and intragenic-CpG islands as compared with promoter-CpG islands, and H1.1 is enriched at intragenic-CpG islands as compared with its average binding (Figure S2B). Overall, our results indicate that the degree of H1 displacement from CpG islands can be dependent on their genomic location.

The Distribution of Histone H1 Subtypes at DNA-Methylated Regions Is Context Dependent

So far, our data argue that gene expression correlates inversely with H1 binding. However, transcription per se is not sufficient to explain H1 distribution, since inactive HCPs tend to be devoid of H1 (Figure 3B) and CpG islands can be differentially bound by H1 subtypes (Figures 3C and S2B). Promoter-CpG islands are typically in an unmethylated state, whereas intra- and intergenic-CpG islands are more often found DNA methylated (Illingworth et al., 2010; Maunakea et al., 2010). We therefore analyzed the DamID binding values of H1 subtypes at CpG islands according to their density of methylation, and observed a positive correlation with DNA methylation (Figures 4A and S3A; Table S2). This tendency was consistent for all CpG island types independently of their location (data not shown). Although in most cases DNA methylation at promoter-CpG islands leads to gene silencing, this is not the case for orphan-CpG islands (Zilberman et al., 2007). Interestingly, intragenic-CpG islands are especially prone to DNA methylation (Illingworth et al., 2010; Maunakea et al., 2010), and this is well reflected by the slight enrichment of H1 subtypes (Figure S2B). As expected, we also found a positive correlation between H1 binding and DNA methylation at HCPs and ICPs (Figures 4B and S3B; Table S1). However, at LCPs, the presence of
Figure 2. H1 Subtypes Are Depleted at Active TSS and Regulatory Regions

(A) H1 subtypes distribution observed around TSSs according to transcription levels. The averaged H1 DamID binding values (log₂Dam-H1/Dam) plotted 5 kb around the TSS of four categories of transcribed genes: highly transcribed (top 25%: high, light blue), medium highly transcribed (top middle 25%: medium high, medium light blue), medium lowly transcribed (bottom middle 25%: medium low, medium dark blue), and lowly/not transcribed (bottom 25%: low, dark blue). For comparison, the distribution of H3K4me3 reads around the TSS of the same transcribed genes is shown at the top.

(B) Enhancers and Active enhancers showing the log₂Dam-H1/Dam values around TSSs and CTCF peaks.

(C) Distribution of H1 DamID binding values around different regions: 5' UTR, Exons, Introns, and 3' UTR.

(D) Distribution of H3K4me1, H3K4me3, and H3K9ac around different regions, showing the log₂Dam-H1/Dam values.

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H1.2–H1.5 seems to correlate negatively with DNA methylation (Figures 4B and S3B; Table S2). In contrast to HCPs and ICPs, LCPs do not show any significant correlation between gene activity and abundance of CpG methylation (Weber et al., 2007). Therefore, it is possible that a low density of methylation is not sufficient for effective repression of LCP-associated genes, and that the preferential presence of H1 at LCPs might contribute to their silencing.

In contrast to what we observed for CpG islands and HCPs, the binding values of H1 subtypes to gene bodies seems to correlate negatively with DNA methylation (Figures 4C and S3C; Table S2), and H1 subtypes bind preferentially at unmethylated gene bodies (Figure S3C). Interestingly H1.1 seems to be enriched at highly methylated exons and its binding values decrease proportionally with the levels of DNA methylation (Figure S3D). In contrast to this, H1.2–H1.5 subtypes tend to be depleted at highly methylated exons.

The Spatial Distribution of H1 Subtypes Is Indicative of Functionally Distinct Chromatin Domains

The recent development of novel powerful techniques has provided insights into the three-dimensional organization of the genome, demonstrating the existence of many transient contacts between genomic loci that are up to several megabases apart (Fullwood and Ruan, 2009; van Steensel, 2011). These regions of highly frequent intra- and intermolecular interactions have been defined as “topological domains” and the shorter chromatin segments separating them as “topological boundaries” (Dixon et al., 2012). Because of its long-studied structural role in local chromatin compaction, linker histone H1 is a good candidate for regulating the spatial organization of the genome within the nucleus. Recently, Dixon et al. (2012) employed the high-throughput chromosome conformation capture (Hi-C) method to map chromatin interactions in IMR90 cells. We therefore compared their data set with our DamID profiles. The DamID binding values of H1 subtypes show a very similar distribution at both domains and boundaries, with a median binding centered around zero (Figure 5A). The higher variance at boundaries could be explained by their smaller size (Figure S4A). This suggests that H1 might not be involved in demarcating topological boundaries versus domains at a global level. However, it was previously suggested that repressive domains are governed by an intrinsic folding regime that is significantly different from active chromatin regions (Sexton et al., 2012), and differential H1 binding could mediate those differences. To investigate this notion, we performed clustering of topological domains according to the enrichment of known epigenetic marks and chromatin regulators as previously described in Drosophila (Sexton et al., 2012). We analyzed four major classes of domains with clear biological functions for the enrichment/depletion of H1 subtypes: active domains (characterized by enrichment of H3K36me3 and H3K4me3), PcG domains (H3K27me3 enriched, mainly transcriptionally repressed), HP1 domains (H3K9me3 enriched, mainly transcriptionally repressed), and null domains (not enriched for any mark). Based on our DamID mapping, we find that different combinations of H1 subtypes can associate with functionally distinct topological domains: H1.1 seems to be preferentially enriched in PcG domains and slightly enriched in active domains, whereas H1.2–H1.5 are enriched in HP1 and null domains and depleted from active domains (Figure 5B).

An additional mechanism that contributes to the spatial organization of the genome is the anchoring of chromatin segments to the nuclear periphery by lamina-associated domains (LADs) (Guelen et al., 2008; Peric-Hupkes et al., 2010). Interestingly, we observed a strong enrichment of H1 within LADs (Figures 5C and 5D), and that changes in H1.2–H1.5 distribution can demarcate LAD boundaries (Figure S4B). In support of this observation, we found that depletion of H1.4 can globally affect the localization of lamins at the nuclear periphery (Figure S4C).

Although the precise molecular basis for this association of H1 with specific functional domains remains to be elucidated, these data suggest that distinct H1 subtypes have a preference for certain genomic regions and might be implicated in the formation of repressive chromatin domains, clearly distinguishing them from active chromatin.

DISCUSSION

Here, we present a comprehensive analysis of the genomic distribution of all somatic, replication-dependent human linker histone H1 subtypes. Because currently the specificity of H1 subtype-specific antibodies in ChIP assays is questionable, we employed an unbiased method, DamID.

DamID as a Method for Studying H1 Subtypes

DamID is now a well-established technique and often is the only valid alternative to chromatin immunoprecipitation (ChIP) in genome-wide studies. Due to the lack of linker histone H1 subtype-specific antibodies, DamID is a method of choice for analyzing the genomic distribution of H1 subtypes. In addition, DamID has a number of other important advantages. First, DamID detects protein-DNA interactions as they occur in living cells. Moreover, the Dam fusion protein is expressed at very high levels.
low levels and thus is unlikely to interfere with the functions of the endogenous protein or its targets. Third, DamID is a suitable method for proteins with relatively short residence times (Gelbart et al., 2005; Schmiedeberg et al., 2009; Misteli et al., 2000) because the adenine methylation remains after the protein has dissociated. Overall, detailed comparisons have extensively shown that DamID identifies the same binding sites and profiles as ChIP for a variety of proteins in different species (Greil et al., 2006; Tolhuis et al., 2006; Nègre et al., 2006; Filion et al., 2010). Importantly, the DamID binding profile...
of *Drosophila* H1 (under conditions comparable to ours) has been directly compared twice with its genomic binding obtained by ChIP (Filion et al., 2010; Braunschweig et al., 2009), confirming that fusion to Dam does not alter the chromatin-binding properties of linker histone and that DamID is a valid method for studying H1 distribution. We also validated key features of the genomic distribution of H1 subtypes identified by DamID by quantitative ChIP-PCR in HEK293 stably expressing Flag-HA-H1.1 and Flag-HA-H1.4 (Figure S5), further strengthening our results.

However, the application of DamID has some potential limitations. It requires a relatively long time to express the fusion protein (Filion et al., 2010). However, this does not represent a limitation in our study, which focuses on unraveling the average chromatin-binding sites of linker histone H1 subtypes in a large population of unsynchronized cells. Furthermore, the DamID approach we used has a resolution of ~1 kb (Greil et al., 2006), which is comparable to the resolution of traditional ChIP with chromatin fragments of ~1 kb, but is 2–3 times lower than that achieved by recent ChIP-sequencing approaches. Although such resolution may not always be sufficient to determine the accurate binding position of every protein on a single-locus level, an average binding profile in a metagene-type analysis, such as the one obtained in our study, is highly informative and robust.

**Genomic Distribution of H1 Subtypes**

In our analysis, we found that globally human H1 subtypes (H1.2–H1.5) tend to have a very similar distribution, with both specific dips and enrichments. Although the enrichments/depletions we observe are relatively slight, we expect them to be biologically relevant, since a similar situation of small but functionally

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**Figure 4. DNA Methylation Can Affect H1 Binding in a Context-Dependent Manner**

A) Observed H1 subtypes distribution at CpG islands according to their density of methylation. DNA methylation density was defined as the proportion of methylated cytosines per basepair. CpG islands were scored for DNA methylation as described in Materials and Methods and then divided into three groups of decreasing DNA methylation density (high, medium, low). The distribution of the H1 DamID binding values (log2Dam-H1/Dam) is shown as a boxplot. The density of DNA methylation is color coded by decreasing intensities of red.

B) H1 subtypes distribution at promoter classes according to their density of methylation. Boxplots show distribution of the H1 DamID binding values for HCP, ICP, and LCP classes, with each further divided into three groups of decreasing DNA methylation density as described in Materials and Methods. Red, light red, and rose boxes correspond to high, medium, and low density of DNA methylation, respectively.

C) H1 subtypes distribution within gene bodies according to their density of methylation. DNA methylation density at the gene body, as defined in Materials and Methods, is scored as described for CpG islands, and the distributions of the H1 DamID binding values are shown as a boxplot. The levels of DNA methylation density are color coded as in (A) and (B).

See also Figures S3 and S5 and Table S2.
important enrichments on regulatory elements has also been reported for the core histone variant H3.3 (Goldberg et al., 2010).

In general, we found H1 subtypes to be depleted from active promoters as well as from regulatory elements controlling transcription, and enriched at regions carrying repressive histone marks such as H3K9me3 and H3K27me3. In particular, CpG-dense regions often seem to be devoid of H1.2–H1.5. We found that the depletion of H1 subtypes is generally less pronounced at intra- and intergenic (orphan) CpG islands than at promoter-CpG islands. Based on this, we speculate that CpG islands are differently bound by H1 depending on their location. Many orphan-CpG islands represent alternative promoters of annotated genes, whereas others are initiation sites of noncoding RNA transcription (Illingworth et al., 2010), suggesting that active transcription might trigger the loss of H1 at CpG-dense regions. Furthermore, we found that H1 subtypes are present within gene bodies, with less pronounced changes depending on the transcription levels (data not shown). This suggests that H1 binding might be more sensitive to initiation of transcription than to transcriptional elongation. This is in line with previous findings that other chromatin marks, such as DNA methylation and the methyl-CpG binding protein 2 (MeCP2), are associated with silencing when present at the TSS but are permissive for transcription when found within gene bodies (Jones, 1999).

Transcription does not seem to be the only determinant of the genomic distribution of H1, since, e.g., poised promoters and inactive HCPs can also be depleted of H1. A recent analysis indicated that promoter-CpG islands are kept in a nucleosome-free state through the stable binding of regulatory factors (Choi, 2010). Thus, H1 distribution might also be regulated by other chromatin proteins rather than simply being a consequence of transcription or an intrinsic limitation of H1 binding to CG-rich DNA. We therefore favor a model in which H1 binding is regulated at multiple levels and H1 subtypes act as fine-tuners of gene regulation.

The Link between DNA Methylation and H1 Binding

The functional link between DNA methylation and H1 binding has been under discussion for a long time, and partially contradictory

Figure 5. The Spatial Distribution of H1 Subtypes in the Nucleus Overlaps with Repressive Chromatin Domains
(A) H1 subtypes distribution at topological domains versus boundaries. Boxplots show the distribution of H1 DamID binding values (log2(Dam-H1/Dam)) within the topological domains and their boundaries, as defined using Hi-C in human IMR90 cells (Dixon et al., 2012). Violet boxes represent boundaries and light violet boxes indicate domains.

(B) H1 subtypes enrichment at topological domains assigned to four epigenetic classes. Heatmap shows enrichment (red) and depletion (blue) with respect to random expectations of specific histone marks, H1.1–H1.5, and LADs within each of four epigenetic classes of physical domains determined by Hi-C (Dixon et al., 2012) and defined according to Sexton et al. (2012). The epigenetic marks used for supervised clustering into polycomb (PcG), HP1, active, and null domains were H3K27me3, H3K9me3, H3K36me3, and H3K4me3, respectively.

(C and D) H1 subtypes enrichment at LADs.

(C) Black boxplots show the distributions of the H1 DamID binding values (log2(Dam-H1/Dam)) within LADs, as previously defined (Guelen et al., 2008).

(D) Genome browser snapshot of the H1 subtypes distribution at a subregion of chr16 (red box). The position of LADs is indicated by blue boxes at the bottom of H1 DamID profiles, and the highlighted light blue regions show the H1 distribution within LADs.

See also Figures S4 and S5.
results regarding the preferential recruitment of H1 on methylated versus unmethylated chromatin have been reported (Nightingale and Wolff, 1995; McArthur and Thomas, 1996; Ball et al., 1983; Campoy et al., 1995). There could be several reasons for these apparent discrepancies. In early in vitro experiments, investigators used total H1 purified from different sources without considering the variation in the relative amount of H1 subtypes (Helliger et al., 1992; Lennox and Cohen, 1983). Furthermore, the affinity of H1 subtypes for methylated DNA was often measured in vitro on naked DNA instead of nucleosomes (Strom et al., 1995; McArthur and Thomas, 1996). Purified H1 subtypes might also lack their original modifications, which are known to have important consequences for gene regulation (Kamieniarz et al., 2012; Weiss et al., 2010; Hergeth et al., 2011). Our data show that on a genomic scale in vivo, there seems to be no simple correlation between H1 binding and DNA methylation, and instead the presence of H1 at methylated regions could be dependent on the genomic location and therefore on the chromatin context.

**Role of H1 in Higher Order of Chromatin Structure**

It has now became evident that chromatin is organized within the nucleus according to specific biological functions (Duan and Blau, 2012). Three main molecular mechanisms have been proposed to explain how chromatin can acquire its three-dimensional configuration: (1) local chromatin compaction into irregular patches of aggregated nucleosomes, (2) inter- and intramolecular interactions between distant DNA elements into topological domains, and (3) anchoring of chromatin to nuclear scaffolds (van Steensel, 2011). Because of its ability to regulate chromatin compaction, it has been suggested that H1 could play a key role in the spatial organization of the genome. However, this interesting possibility has not been addressed thus far. Our mapping data and the analysis of H1 distribution within topological domains suggests that differences among H1 subtypes at the linear genomic level can be recapitulated within the tridimensional organization of nuclear space. In particular, H1 subtypes could take part in the global segmentation of the genome into active and inactive regions (domains) and be implicated in the establishment and/or maintenance of LADs. This suggestion is in agreement with previous findings that LADs contain mainly highly expressed genes (Guelen et al., 2008) and are mainly associated with HP1 and H1.1, H1.2–H1.5 (Figure 5B). Moreover, the insulator protein CTCF and CpG islands can delimit the borders of LADs (Guelen et al., 2008), regions which have found to be depleted of H1. In line with this, changes in H1.2–H1.5 distribution can demarcate LAD boundaries. Together, these results suggest a possible involvement of H1 in LAD confinement.

It has been proposed that in addition to gene regulation, attachment of genomic regions to the nuclear lamina contributes to the spatial organization of interphase chromosomes inside the nucleus (Guelen et al., 2008). In line with this, we noticed that different combinations of H1 subtypes can be enriched at individual chromosomes, and in particular chromosomes with the highest LAD coverage also have the strongest H1.2–H1.5 enrichment (Figures 1B and S1D). This specific H1 pattern might contribute to the compartmentalization of chromosomes into specific territories (Cremer and Cremer, 2001) through, e.g., direct local chromatin compaction, which could be dependent on the H1 subtypes present and/or the recruitment of H1 subtype-specific interactors, which in turn could tether H1-bound regions to different nuclear locations.

**H1.1 Is a Special H1 Subtype**

Our analysis suggests that, based on their DamID binding values, H1 subtypes can be divided into two groups: (1) H1.1 and (2) H1.2–H1.5. We found that the differences between the enrichment of H1.1 and that of H1.2–H1.5 seem to be more pronounced at regulatory regions marked by activating histone modifications such as promoters, enhancers, and CpG islands (Figures 2 and 3). For example, we observed in our DamID profiling that although H1.1 is not depleted at inactive and poised HCPs, H1.2–H1.5 enrichment levels remain high (Figure 3B). HCPs are often associated with developmentally regulated genes (Zhu et al., 2008), which need to be activated in response to external stimuli and therefore do not require a stable repressive chromatin organization. Moreover, although CG richness can affect H1 subtype binding to chromatin similarly (Figure S3A), H1.1 is the only subtype that displays a rather positive correlation with DNA methylation independently of its genomic location (Figure S3B; Table S1).

Based on these findings, we propose that H1.1 is somehow special among H1 subtypes. The chromatin structure promoted by H1.1 binding might support a level of compaction that facilitates rapid conversion into either an active or repressed state. Previous data also agree with this special role for H1.1. For example, it was shown that H1.1 had the least ability to condense nucleosomes in vitro (Khade and Rao, 1995), and had the lowest affinity to chromatin both in fluorescence recovery after photobleaching experiments in vivo and biochemical studies in vitro (Orrego et al., 2007; Th’ng et al., 2008). The dynamic binding of H1.1 might at least in part explain why chromatin-containing H1.1 does not necessarily impede gene expression. In line with this, Alami et al. (2003) showed that H1.1 has a unique role in activating a reporter transgene in mice. In view of its distinct distribution and its potential association with more active and open chromatin, it is not surprising that H1.1 expression is restricted to certain cells, thereby conferring some tissue-specific function (Piña et al., 1987). Interestingly, estimation of the degree of nucleotide substitutions during evolution also pointed toward a functional differentiation of H1.1 from the other H1 subtypes, further strengthening the uniqueness of H1.1 among the somatic H1 subtypes from an evolutionary perspective (Ponte et al., 1998).

In conclusion, we present here a genome-wide distribution of human linker histone H1 subtypes. Our comprehensive analysis represents a key step toward answering many open questions about the function of H1 and the significance of H1 heterogeneity in higher eukaryotes. Our work will also provide a resource for further studies to elucidate the molecular mechanisms that regulate H1 binding to chromatin, and the consequences of the functional diversity of H1 subtypes.

**EXPERIMENTAL PROCEDURES**

The DamID experiments were performed in IMR90 cells as previously described (Moorman et al., 2006). Normalization and analyses of DamID
data were performed using custom R scripts (http://www.r-project.org/), GenPlay software (Laugie and Bouhassira, 2011), and Galaxy (Blankenberg et al., 2010). For more details, see Extended Experimental Procedures. Coordinates for the human RefSeq genes, introns, exons, 5' UTRs, 3' UTRs, CpG islands, and LADs were retrieved from the UCSC Table Browser. Histone-modification maps, gene-expression data, and bisulfite sequencing data for IMR90 cells were obtained from the Gene Expression Omnibus (GEO; data set GSE16296). The CTCF map for IMR90 cells was retrieved from GEO (data set GSE5559). Table S1 summarizes the Gene Ontology (GO) analysis of genes present in the HMM states. Table S2 shows the correlations of H1 binding and DNA methylation, and Table S3 includes the list of primers used in this study.

ACCESSION NUMBERS

The data have been submitted to the NCBI GEO under accession number GSE40886.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Extended Experimental Procedures, five figures, and three tables and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2013.05.003.

LICENSING INFORMATION

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