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# Global histone acetylation induces functional genomic reorganization at mammalian nuclear pore complexes

Christopher R. Brown,<sup>1,3</sup> Caleb J. Kennedy,<sup>1,3</sup> Valerie A. Delmar,<sup>2</sup> Douglass J. Forbes,<sup>2</sup> and Pamela A. Silver<sup>1,4</sup>

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The nuclear localization of genes is intimately tied to their transcriptional status in *Saccharomyces cerevisiae*, with populations of both active and silent genes interacting with components of the nuclear envelope. We investigated the relationship between the mammalian nuclear pore and the human genome by generating high-resolution, chromosome-wide binding maps of human nucleoporin 93 (Nup93) in the presence and absence of a potent histone deacetylase inhibitor (HDACI). Here, we report extensive genomic reorganization with respect to the nuclear pore following HDACI treatment, including the recruitment of promoter regions, euchromatin-rich domains, and differentially expressed genes. In addition to biochemical mapping, we visually demonstrate the physical relocalization of several genomic loci with respect to the nuclear pore following HDACS leads to significant changes in genomic organization, recruiting regions of transcriptional regulation to mammalian nuclear pore complexes.

[Keywords: NPC; nucleoporin; nuclear organization; Nup93; ChIP-chip]

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The nucleus is a structurally and functionally complex organelle with a nonuniform interior consisting of distinct chromatin domains and several proteinaceous subcompartments. Chromosomes occupy nonrandom intranuclear positions with respect to each other and the nuclear periphery (Croft et al. 1999; Parada and Misteli 2002; Parada et al. 2002, 2004a,b; Tanabe et al. 2002a,b; Cremer et al. 2006). Chromosome positioning is believed to expose genomic loci to functionally distinct regions in the nucleus, generating transcriptional regulatory domains favoring either activation or repression. Distinct subnuclear regions also direct specific transcriptional programs by organizing genomic loci around specialized protein hubs. The nucleolus is one such region, mediating the localized transcription of ribosomal RNA genes encoded on multiple chromosomes. In the budding yeast, Saccharomyces cerevisiae, a nucleolar-proximal region has also been shown to cluster several tRNA genes (Thompson et al. 2003).

Components of the nuclear envelope have been shown to assert a repressive role in transcription. In yeast, the silent mating-type loci and telomeres are regulated by components of the nuclear periphery, including the nuclear pore complex (NPC) (Stavenhagen and Zakian 1994; Thompson et al. 1994; Maillet et al. 1996; Marcand et al. 1996; Andrulis et al. 1998; Feuerbach et al. 2002). However, several recent studies have reported NPCproximal transcriptional activation, with the concomitant recruitment of induced genes from the nuclear interior to the periphery (Brickner and Walter 2004; Casolari et al. 2004, 2005; Menon et al. 2005; Cabal et al. 2006; Dieppois et al. 2006; Drubin et al. 2006; Schmid et al. 2006; Taddei et al. 2006; Brickner et al. 2007; Luthra et al. 2007; Sarma et al. 2007). NPC association can increase the efficiency of mRNA processing and export through associations with the SAGA and TREX complexes, regulate the absolute levels of gene expression, and establish an epigenetic state that confers transcriptional memory and rapid reactivation of genes (Cabal et al. 2006; Taddei et al. 2006; Brickner et al. 2007). Furthermore, several components of the nuclear transport machinery possess boundary activity, potentially facilitating the presence of both repressive and activating domains at the nuclear periphery (Ishii et al. 2002).

Elements of the nuclear periphery are also involved in transcriptional regulation in *Drosophila melanogaster*. In *Drosophila*, the dosage compensation complex (DCC),

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required for the twofold increase in gene expression on the peripherally localized male X chromosome, interacts with two nuclear pore components (Mendjan et al. 2006). The deletion of these pore proteins eliminates the hypertranscription of the male X chromosome, implicating *Drosophila* nuclear pores in transcriptional activation. In support of this, the *Drosophila* proteins E(y)2 and Xmas-2 were recently shown to regulate mRNA expression, export, and the subnuclear positioning of the *hsp70* gene cluster (Kurshakova et al. 2007). E(y)2 and Xmas-2 are components of the *Drosophila* SAGA and TREX complexes, whose homologs in yeast play a role in NPCassociated transcriptional activation (Cabal et al. 2006; Dieppois et al. 2006; Taddei et al. 2006).

Interestingly, a recent genome-wide study in Drosophila probing interactions between the nuclear lamina and the genome uncovered a repressive role for the nuclear periphery (Pickersgill et al. 2006). Lamins are integral components of a protein network that lines the inner surface of the nuclear envelope between NPCs and have been shown to bind chromatin in vitro. These results suggest that distinct peripheral components, NPCs, and lamins, may have divergent roles in transcriptional activation and repression, respectively. In addition, a lamin-associated protein, LAP2B, binds to histone deacetylase 3 (HDAC3), a member of a large family of proteins that removes acetyl modifications from histones (Somech et al. 2005). Highly acetylated histones are found in the promoters of transcriptionally active genes, suggesting that the lamin-mediated enrichment of HDAC3 at the nuclear periphery could aid in the maintenance of a transcriptionally repressive environment (Fukuda et al. 2006).

Observations in both mouse and human cells indicate the presence of an equally diverse transcriptional regulatory domain at the nuclear periphery. For example, gene-poor chromosomes are reproducibly located near the nuclear periphery in human lymphocytes and fibroblasts (Croft et al. 1999; Boyle et al. 2001; Tanabe et al. 2002b). However, a study of the murine  $\beta$ -globin locus during erythroid maturation showed that significant transcriptional activity occurred at the nuclear periphery prior to the locus's transit to the interior (Ragoczy et al. 2006). While NPCs have yet to be implicated in  $\beta$ -globin activation, evidence has emerged that a murine nucleoporin, Nup96, is involved in interferon gene regulation (Faria et al. 2006).

The mammalian NPC is thought to contain as many as 30 unique proteins that are present in multiple copies due to the eightfold symmetry of the complex (Cronshaw et al. 2002). Nucleoporin 93 (Nup93) is a mammalian nucleoporin centrally located in the nuclear pore (Rout et al. 2000; Krull et al. 2004). siRNA-mediated depletion of Nup93 in HeLa cells resulted in misshapen nuclei lacking some, but not all nucleoporins, while depletion of both Nup93 and the transmembrane nucleoporin Ndc1 led to a complete disruption of NPCs (Mansfeld et al. 2006). Nup93, as well as importin- $\alpha$  and CAS (an exportin), have been shown to interact with the histone acetyltransferase (HAT), CREB-binding protein (CBP) (Ryan et al. 2006). This interaction is similar to one seen in *S. cerevisiae*, where association of a HAT with the NPC is mediated by interactions with SAGA, a multimeric complex required for the expression of numerous yeast genes (Green et al. 2003; Rodriguez-Navarro et al. 2004). As mentioned previously, NPC–SAGA interactions are thought to play a major role in mediating gene recruitment to the NPC in *S. cerevisiae* (Cabal et al. 2006; Dieppois et al. 2006; Taddei et al. 2006).

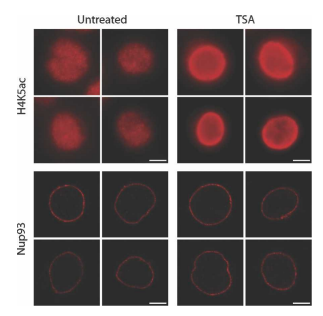
HDAC inhibitors (HDACIs) globally elevate levels of histone acetylation in the nucleus by inhibiting Class I and II HDACs (Drummond et al. 2005). HDACI treatment also leads to the enrichment of acetylated histones at the nuclear periphery (Taddei et al. 2001; Gilchrist et al. 2004; Drummond et al. 2005). Local changes in genomic organization have also been reported following treatment with HDACIs (Taddei et al. 2001; Zink et al. 2004; Pickersgill et al. 2006). For example, repressed genes are no longer associated with lamins in Drosophila after treating cells with the HDACI trichostatin A (TSA) for 24 h (Pickersgill et al. 2006). In addition, the human CFTR gene moves away from the nuclear periphery upon treatment with TSA for 10 h (Zink et al. 2004). Importantly, several HDACIs are in clinical trials for various forms of cancer due to their ability to induce the expression of repressed genes that lead to growth arrest, differentiation, and apoptosis in transformed cells (Drummond et al. 2005; Glaser 2007).

Disparate observations of both transcriptional activation and repression at the nuclear periphery in S. cerevisiae and Drosophila led us to investigate the presence of a similar regulatory domain in human nuclei. Using genomic location analysis (Ren et al. 2000), we report Nup93 interactions with human chromosomes 5, 7, and 16. To investigate the effects of global histone acetylation on NPC-chromatin interactions, we treated cells with TSA, a reversible HDACI that does not perturb gross nuclear structure (Taddei et al. 2001; Gilchrist et al. 2004; Pickersgill et al. 2006). Upon treatment with TSA, Nup93 distribution was significantly altered across all three chromosomes, indicative of a large-scale nuclear reorganization event with associated effects on gene expression. Analysis of Nup93-binding sites identified enrichments for several genomic features involved in transcriptional regulation. Together, these features define regions of functional interaction between the nuclear pore and the human genome.

#### Results

#### Identification of Nup93-binding sites

We investigated Nup93–genomic association in the presence and absence of TSA, a potent HDACI. We also performed chromatin immunoprecipitation (ChIP) on several other nucleoporins including Nup62, Nup107, and Nup205, isolating significant amounts of DNA (data not shown). Nup93 was chosen for further study due to the robustness of Nup93 ChIPs and the strictly maintained subnuclear localization of Nup93 at the nuclear enve-



**Figure 1.** TSA treatment of HeLa S3 cells. Histone acetylation was monitored with an anti-acetylated H4K5 antibody (H4K5ac) in untreated and TSA-treated HeLa S3 cells. Histone acetylation was enriched at the nuclear periphery after 12 h of exposure to 40 ng/mL TSA. H3K9 acetylation is also enriched, although to a lesser extent (Supplemental Fig. 1). The peripheral localization of Nup93 was unchanged following TSA treatment. Bars, 5 µm.

lope (Fig. 1). Immunofluorescence microscopy of HeLa S3 cells following a 12-h, 40-ng/mL TSA treatment revealed a marked increase in histone H4K5 acetylation at the nuclear periphery (Fig. 1), a finding supported by other studies (Taddei et al. 2001; Gilchrist et al. 2004). H3K9 acetylation was also enriched at the nuclear periphery (Supplemental Fig. 1), although to a lesser extent, consistent with previous observations (Gilchrist et al. 2004). Additionally, we found that Nup93 localization was unchanged following TSA treatment (Fig. 1). The subnuclear localization of two other nuclear envelope proteins, lamin B and LAP2, are also unaffected by TSA treatment (Taddei et al. 2001).

Next, we performed genomic location analysis by coupling ChIPs and microarray analysis (ChIP-chip) to map interactions between Nup93 and the human genome. This technique has been used extensively in S. cerevisiae and is emerging as a powerful tool in higher eukaryotes (Ren et al. 2002; Carroll et al. 2005, 2006; Kim et al. 2005). Asynchronously growing HeLa S3 cells were cross-linked, stabilizing both protein-protein and protein-DNA interactions. We then performed ChIPs of Nup93 and purified, amplified, and hybridized the associated DNA to Affymetrix tiled arrays (ChipE, also referred to as chip5) containing all nonrepetitive regions of human chromosomes 5, 7, and 16. This subset of chromosomes was examined based on preliminary data obtained from Nup62, Nup107, and Nup205 genomic association studies that showed significant interactions with chromosome 7 on ENCODE arrays, including binding at the *HOXA* locus and *CFTR* (data not shown). In addition, chromosome 7 has been shown to preferentially reside near the nuclear periphery in human cells (Boyle et al. 2001).

Nup93 was mapped to 207 sites on chromosomes 5, 7, and 16 in untreated cells and 170 sites in TSA-treated cells (Fig. 2A; Supplemental Table 1). The 18% reduction in total Nup93-binding sites was predominantly due to large-scale changes in binding to chromosome 5, where Nup93 association was reduced from 75 sites in untreated cells to 15 sites following TSA treatment (Fig. 2A; Supplemental Table 1). Significant binding sites were classified as either high confidence (143 untreated and 86 TSA sites,  $10^{-12} < P \le 10^{-7}$ ) or very high confidence (64 untreated and 84 TSA sites,  $P < 10^{-12}$ ) based upon our statistical analysis (see the Supplemental Material).

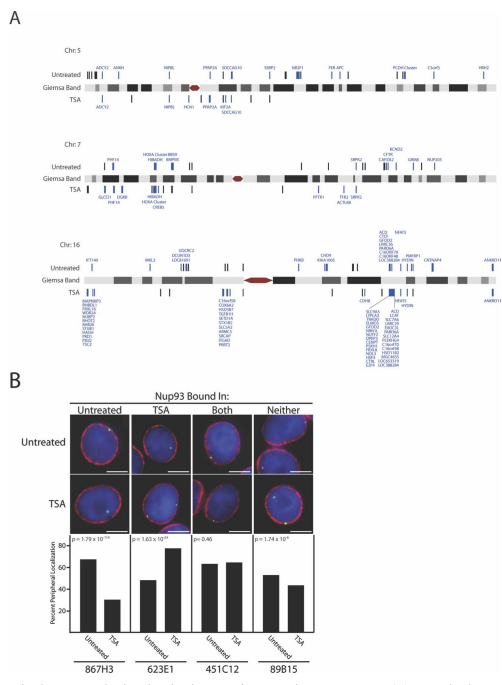
Nup93-binding sites were mapped to their nearest neighboring gene, resulting in the assignment of 86 untreated and 90 TSA-treated genes (some genes contained multiple Nup93-binding sites) (Supplemental Table 2). The untreated gene set did not contain any enriched gene ontology groups as reported by GOstat (Beissbarth and Speed 2004). However, the TSA-treated gene set contained two enriched ontology groups: DNA-binding activity (P = 0.002) and transcription factor activity (P = 0.02). These ontology groups include several genes that are members of the *HOXA* transcription factor cluster on chromosome 7, a gene-dense locus with extensive Nup93 association (Fig. 2A).

Mapping of Nup93 sites in untreated and TSA-treated cells across chromosomes 5, 7, and 16 revealed large changes in nuclear pore association (Fig. 2A). Significant reorganization of the q-arm of chromosome 5 occurred with the loss of statistically significant Nup93 binding over a 98-Mb region following TSA treatment (Fig. 2A). A similar change was seen on the q-arm of chromosome 7 where Nup93 association was lost over a 35-Mb region following TSA treatment (Fig. 2A). However, changes of this kind were not observed on chromosome 16, where many Nup93-binding sites, while mostly intergenic in untreated cells, overlapped with coding regions following TSA treatment (Fig. 2A). Interestingly, the subnuclear localization of chromosome territories has been shown to be unaffected by TSA, while local changes in chromatin condensation have been observed (Croft et al. 1999; Santos et al. 2002).

#### Visualization of Nup93-associated loci

To verify that Nup93-binding sites were located at the nuclear periphery, we performed FISH combined with confocal microscopy to visualize the intranuclear localization of several genomic loci. Four loci with distinct Nup93 association patterns were chosen (Fig. 2B). One locus, associated in untreated cells and absent in TSA-treated cells, was found at the nuclear periphery 66% of the time in untreated cells versus just 30% in TSA-treated cells (probe 867H3,  $P = 1.79 \times 10^{-114}$ ) (Fig. 2B). Another locus, associated in TSA-treated cells and absent in untreated cells and absent in untreated cells and absent in the time in untreated cells (probe 867H3,  $P = 1.79 \times 10^{-114}$ ) (Fig. 2B).

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**Figure 2.** Nup93 binding map and subnuclear localization of associated genomic regions. (*A*) Nup93-binding sites identified by ChIP-chip are plotted on schematic representations of chromosomes 5, 7, and 16. Nup93-binding sites are denoted by black bars *above* or *below* each chromosome for untreated and TSA-treated cells, respectively. Nup93-binding sites directly overlapping coding regions are labeled blue with the associated gene displayed. Chromosomes are not to scale. (*B*) FISH analysis of Nup93 targets. Four genomic loci were visually mapped in untreated and TSA-treated HeLa S3 cells. From *left* to *right*, these loci interact with Nup93 in untreated cells, TSA-treated cells, both conditions, and neither condition. The *top* images are examples from untreated cells, while the *bottom* row are examples from TSA-treated cells. In each image, the FISH signal is green; DNA stained with DAPI is blue; and lamin B, a marker of the nuclear periphery, is red. The bar graphs report percent peripheral localization of each locus in untreated and TSA-treated cells. *P*-values comparing the difference in percent peripheral localization for each locus were calculated using the  $\chi^2$  test. BAC probes used to study each genomic region are identified at the *bottom* of each column. Bars, 5 µm.

ery just 48% of the time in untreated cells versus 77% in TSA-treated cells (probe 623E1,  $P = 1.63 \times 10^{-63}$ ) (Fig. 2B). A third locus, associated in both conditions, was

found at the nuclear periphery 63% of the time in untreated cells and 64% of the time in TSA-treated cells (probe 451C12, P = 0.46) (Fig. 2B). Finally, a genomic re-

gion that was not bound in our ChIP-chip analysis was found at the nuclear periphery 52% and 43% of the time in untreated and TSA-treated cells, respectively (probe 89B15,  $P = 1.74 \times 10^{-6}$ ) (Fig. 2B). The high percentage of peripheral localization observed for the control probe (89B15) in both conditions and probe 623E1 in untreated cells may reflect the peripheral positioning of chromosome 7 in mammalian cells (Boyle et al. 2001). Interestingly, probe 867H3, located on chromosome 5, shows only 30% peripheral association in TSA-treated cells, likely reflecting the preferentially internal positioning of chromosome 5. Associated P-values test the significance of the difference in the percent peripheral localization for each locus in untreated and TSA-treated cells. These results verify that ChIP-chip identifies Nup93-binding sites enriched at the nuclear periphery and is capable of distinguishing dynamic changes in localization that are influenced by TSA treatment.

Differential Nup93 binding was also observed at the *CFTR* locus on the q-arm of chromosome 7. Specifically, ChIP–chip showed that Nup93 was associated with the *CFTR* locus in untreated cells, but was not associated with the locus in TSA-treated cells (Fig. 2A). This result is consistent with previously reported visual evidence showing that *CFTR* is peripherally localized in mammalian cells and moves to the nuclear interior upon TSA treatment (Zink et al. 2004).

## Comparison of Nup93 association in untreated and TSA-treated cells

We investigated the magnitude of TSA-induced genomic reorganization and found that only 34 binding sites and 25 genes were shared between untreated and TSAtreated Nup93-binding sites, indicating a significant shift in peripheral association (Fig. 3A). Comparison of untreated and TSA-treated Nup93-binding sites with chip5 distributions reveals significant changes in association across five genomic regions: exons, introns, promoters, downstream elements, and intergenic regions  $(P = 0.001; \chi^2)$ . Untreated binding sites are random relative to all categories except for introns when compared with chip5 (P = 0.002). However, when compared with chip5 distributions, TSA-binding sites are significantly enriched for introns and promoters (P = 0.005 and) $P = 1.2 \times 10^{-14}$ , respectively) and depleted for exons and intergenic regions ( $P = 5.7 \times 10^{-4}$  and  $P = 5.4 \times 10^{-5}$ , respectively) (Fig. 3B). In agreement with this observation, we mapped the distance from the center of all untreated and TSA-treated Nup93-binding sites to the nearest transcription start site (Fig. 3C). Untreated binding sites show a slight bias toward the nearest transcription start site (P = 0.002), while TSA-binding sites are significantly closer to their nearest transcription start site  $(P = 6.8 \times 10^{-6})$ . Additionally, TSA-binding sites are significantly closer to the nearest transcription start site compared with untreated sites (P = 0.02).

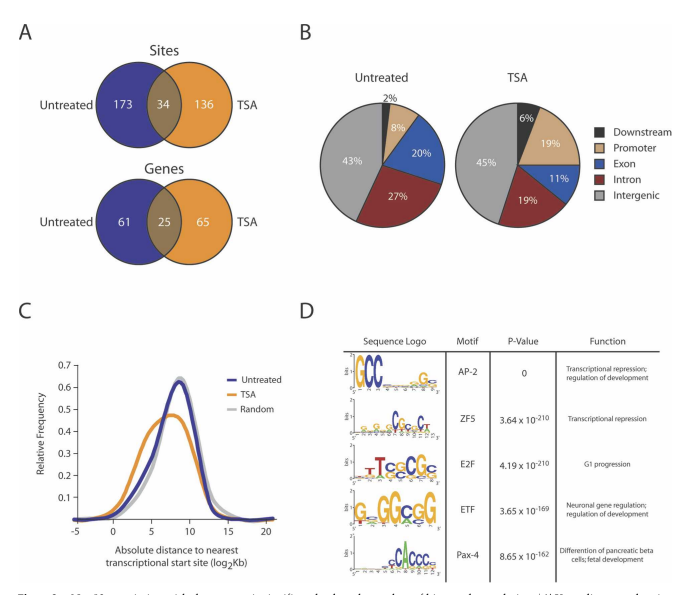
We identified several enriched transcription factorbinding motifs in our TSA-treated data set by analyzing our sites with CEAS (*Cis*-regulatory Element Annotation System, http://ceas.cbi.pku.edu.cn), a Web-based ChIPchip annotation system (Ji et al. 2006). TSA-treated Nup93-binding sites contained 106 enriched transcription factor-binding motifs; the top five are shown in Figure 3D with a complete list included in Supplemental Table 3. However, we identified only one enriched transcription factor-binding motif in untreated Nup93-binding sites, E2F ( $P = 8.71 \times 10^{-7}$ ). Importantly, multiple iterations of randomly generated data sets failed to identify any significant motifs using CEAS. The increase in the number of transcription factor-binding motifs in the TSA-treated data set likely reflects the shift in Nup93 binding to promoter regions and transcription start sites. The identified motifs are recognized by a wide array of transcription factors whose diverse functions include transcriptional repression, transcriptional activation, G1 progression, and neuronal development.

## Nup93 binding is proximal to differentially expressed genes

We performed RNA expression analysis that revealed patterns of expression upon TSA treatment: 753 genes were induced and 828 were repressed (Supplemental Table 4). Of these genes, 91 mapped to chromosome 5 (34 induced, 57 repressed), 70 to chromosome 7 (33 induced, 37 repressed), and 29 to chromosome 16 (17 induced, 12 repressed). However, there was no obvious overlap between Nup93-binding sites and transcriptionally induced or repressed genes. To probe the relationship further, we mapped all untreated and TSA-treated Nup93binding sites to the nearest differentially expressed gene (induced and repressed). While no distance-to-gene relationship was uncovered for untreated Nup93-binding sites (Fig. 4A), we did uncover a strong correlation in TSA-treated Nup93-binding sites (Fig. 4B). One example of the proximal positioning of Nup93-binding sites to differentially expressed genes following TSA treatment is shown in Figure 4C. In untreated cells, Nup93-binding sites were identified in the intergenic region between two genes, RPA3 and GLCCI1, on chromosome 7. In TSA-treated cells, Nup93 sites were also found in both promoter regions, consistent with the promoter enrichment discussed earlier (Fig. 3B). Interestingly, while GLCCI1 expression was induced following TSA treatment, RPA3 expression was unaffected. Observations such as these show that differentially expressed genes may localize to the nuclear periphery, but that proximal positioning to the NPC is not always coupled with a transcriptional change.

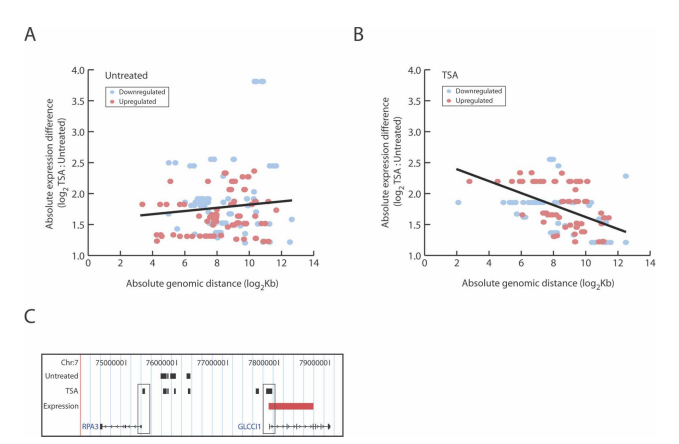
### *Nup93-binding sites are enriched for specific histone methylations*

Untreated and TSA-treated Nup93-binding sites were enriched for histone methylations associated with transcriptionally silent and active regions, respectively (Table 1). A recent study mapped 20 unique histone methylations, the histone variant H2A.Z, the insulating factor CTCF, and RNA Polymerase II (PoIII) over the



**Figure 3.** Nup93 association with the genome is significantly altered upon loss of histone deacetylation. (*A*) Venn diagrams showing the overlap between untreated and TSA-treated Nup93-binding sites and nearest genes. (*B*) Comparison of untreated and TSA-treated Nup93-binding sites with chip5 distributions reveals significant changes in association across five genomic regions: exons, introns, promoters, downstream elements, and intergenic regions (P = 0.001;  $\chi^2$ ). Untreated sites are enriched for introns when compared with chip5 (P = 0.002). TSA-binding sites are significantly enriched for introns and promoters (P = 0.005 and  $P = 1.2 \times 10^{-14}$ , respectively) and depleted for exons and intergenic regions ( $P = 5.7 \times 10^{-4}$  and  $P = 5.4 \times 10^{-5}$ , respectively) when compared with chip5 distributions. (*C*) Distribution of distances from the center of each untreated Nup93-binding site, TSA-treated Nup93-binding sites to the nearest transcription start site. *P*-values for C were calculated by comparing untreated and TSA-treated Nup93-binding sites to randomly distributed sites using the two-tailed Mann-Whitney *U*-test. (*D*) Sequence logos for the five most significantly enriched transcription factor-binding motifs identified in TSA-treated Nup93-binding sites. One-hundred-six motifs were identified in TSA-treated binding sites by CEAS (Supplemental Table 3).

whole human genome (Barski et al. 2007). We compared the reported distributions of histone methylations from Barski et al. (2007) with our untreated and TSA-treated Nup93-binding sites and found several significant overlaps (Table 1). Nine of the histone modifications are associated with active genes, five with silent genes, and six showed no bias. Untreated Nup93 sites were enriched for three specific histone modifications associated with silent genes, depleted for two modifications associated with active genes, and depleted for RNA PolII-binding sites. The opposite was found with TSA-treated Nup93 sites, which were enriched for seven histone modifications associated with active genes, depleted for three modifications associated with silent genes, and enriched in RNA PolII-, H2A.Z-, and CTCF-binding sites. These results indicate that regions associated with increased transcriptional activity become associated with Nup93 upon treatment with TSA.



**Figure 4.** Nup93-binding sites are proximal to differentially expressed genes following TSA treatment. Untreated and TSA-treated Nup93-binding sites were mapped to the nearest differentially expressed gene. The  $log_2$ -transformed distance-to-gene value (*X*-axis) was plotted against the magnitude of differential expression (*Y*-axis). Spearman rank order correlation was used to assess statistical significance. Down-regulated genes (blue) and up-regulated genes (red) are shown in each plot. The black trend lines were calculated by least squares best fit. (*A*) There was no correlation between distance-to-gene and magnitude of differential expression for untreated Nup93-binding sites (*P* = 0.0241; *P* = 0.7198). (*B*) Significant correlation was observed when comparing distance-to-gene and magnitude of differential expression for TSA-treated Nup93-binding sites (*P* = 0.5056; *P* = 6.28 × 10<sup>-11</sup>). (*C*) Nup93-binding sites near two genes on chromosome 7, *RPA3* and *GLCCI1*. Untreated and TSA-treated sites overlap in the intergenic space while TSA treatment induces Nup93 binding to the promoter region of both genes, denoted by thin black rectangles. *GLCCI1* expression is induced upon TSA treatment, illustrated by the red rectangle in the Expression track.

#### Nup93 association with Giemsa (G) bands

To investigate chromosome-wide Nup93-binding patterns we compared the distribution of Nup93-binding sites with the G bands of chromosomes 5, 7, and 16. Staining with G dye reveals chromosome-wide banding patterns that are divided into five classes based on their staining intensity (Furey and Haussler 2003). The G-negative (Gneg) class consists of nonstaining bands, while the Gpos25, Gpos50, and Gpos75 classes contain progressively darker G-positive bands. The Gpos100 class consists of the darkest staining bands. Areas of heavy G staining represent condensed regions of the genome that have been shown to be gene poor, transcriptionally inactive, AT-rich, and replicated late in S phase. Of all the classes of G bands, Gneg bands have the highest density of genes, the largest average number of CpG islands, and are considered to represent euchromatin.

We identified nonrandom distributions of Nup93 across G bands in both untreated and TSA-treated

data sets when compared with chip5 distributions (Fig. 5A). Untreated binding sites were enriched in Gpos50 bands (27% compared with 15% on chip5), while depleted in Gneg/Gpos25 bands (44% compared with 52% on chip5), suggesting a bias toward heterochromatin ( $P < 1.4 \times 10^{-32}$ ;  $\chi^2$ ). Conversely, TSA-induced binding sites were enriched in Gneg/Gpos25 bands (60% compared with 52% on chip5), while showing significant depletion in both Gpos75 and Gpos100 bands (9% and 8% compared with 16% and 13% on chip5, respectively), indicating a shift to regions of the genome enriched in euchromatin ( $P < 4.6 \times 10^{-33}$ ;  $\chi^2$ ) (Fig. 5A).

Next, we investigated the distribution of Nup93-binding sites within G bands. All G bands on chromosomes 5, 7, and 16 were normalized to the same length and then partitioned into 10 sections (bins). We assigned each Nup93 site a bin (0–9) based on its relative position between its nearest G-band boundaries. For example, a Nup93-binding site at the extreme 5' G-band boundary would be placed in bin 0, while a binding site at the

Table 1.	Histone	methylation	enrichments
in Nup93	-binding	sites	

Gene bias	Data set	Untreated	TSA
Active	H3K4me1		+++
	H3K4me2		+++
	H3K4me3		+
	H3K9me1		++
	H3K27me1		+
	H3K36me1		
	H3K36me3		
	H4K20me1	-	+++
	H2BK5me1	-	+++
	H2A.Z		+
	RNA PolII	-	+++
Silent	H3K9me2		-
	H3K9me3	++	
	H3K27me2		-
	H3K27me3	++	-
	H3K79me3	++	
None	H3K79me1		
	H3K79me2		
	H3R2		
	H3R2me2(as)		
	H4K20me3	++	
	H4R3me2		
N/A	CTCF		++
,			

Genomic locations of several histone methylations as well as the histone variant H2A.Z, RNA PolII, and the insulating factor CTCF were mapped over the whole genome using the ChIP-Seq method (Barski et al. 2007). Untreated, TSA-treated, and random intervals were mapped to sequence tags specific to each factor by direct overlap. Each histone modification, as well as H2A.Z, RNA PolII, and CTCF, was associated with a discrete tag frequency distribution for untreated and TSA-treated Nup93binding sites, which was compared individually with an expected distribution (random sites) using a  $\chi^2$  goodness-of-fit statistical test. Gene biases were reported in Barski et al. (2007). Tag biases for each factor are reported as either highly enriched (+++;  $P \le 1 \times 10^{-10}$ , moderately enriched (++;  $1 \times 10^{-10} \le P \le 1 \times 10^{-4}$ ), slightly enriched (+;  $1 \times 10^{-4} \le P \le 0.05$ ), or depleted (-;  $P \le 0.05$ ) in untreated or TSA-treated Nup93-binding sites. Blank cells indicate the lack of a statistically significant bias (P > 0.05).

extreme 3' G-band boundary would be placed in bin 9. As a result, each bin shown in Figure 5B represents the relative position of several Nup93-binding sites (between 10 and 40 sites per bin). A randomized data set failed to produce any enrichment across the normalized G bands, as did mapping of RefSeq genes (Fig. 5B). However, Nup93 had distinct enrichment patterns across G bands in both untreated and TSA-treated data sets, with two very strong peaks of enrichment in the untreated data set flanking a large region of enrichment found solely in the TSA-treated data set (Fig. 5B). These observations are indicative of a nuclear reorganization event induced by the inhibition of histone deacetylation.

#### Discussion

Our results show that the mammalian nuclear pore interacts with the human genome. To probe the functionality of this association we increased global levels of histone acetylation by treating cells with the HDAC inhibitor, TSA. Upon drug treatment, interactions between the nuclear pore and chromatin changed significantly. We observed changes in localized regions of transcriptional importance in addition to chromosome-wide changes in NPC association (Fig. 6).

#### Local chromatin changes alter NPC association

TSA treatment induces the global hyperacetylation of histones by reversibly inhibiting Class I and II HDACs. The resulting histone acetylation, which is enriched at the nuclear periphery in TSA-treated cells, decreases

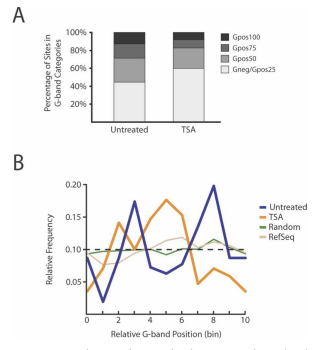
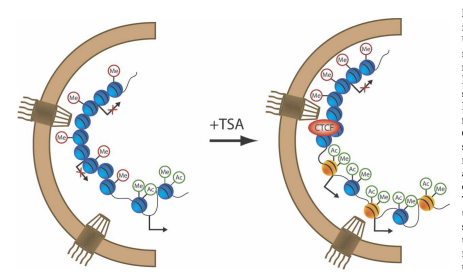


Figure 5. Distribution of Nup93-binding sites within G bands. (A) Nup93 is enriched within Gpos50 bands and depleted in Gneg/Gpos25 bands in untreated conditions when compared with chip5 ( $P = 1.4 \times 10^{-32}$ ;  $\chi^2$ ). However, a similar comparison with chip5 distributions shows that TSA-treated Nup93-binding sites are enriched within Gneg/Gpos25 bands and depleted within Gpos75 and Gpos100 bands ( $P = 4.6 \times 10^{-33}$ ;  $\chi^2$ ). *P*-values were calculated by comparing the observed G-band distribution to chip5 G-band distributions using the  $\chi^2$  test. (B) Binding distribution across G bands. All G bands were normalized by dividing each into 10 equal sections (bins). We assigned each Nup93 site a bin (0–9) based on its relative position between its nearest G-band boundaries. Each bin shown represents the relative position of several Nup93-binding sites (between 10 and 40 sites per bin). Intervals from randomized data sets were uniformly distributed across G bands, which reflected the probe distribution of chip5. Refseq genes failed to show significant positional enrichments (P = 0.97;  $\chi^2$ ). However, both untreated and TSA-treated Nup93-binding sites were enriched in specific regions as shown by the peaks in blue and orange, respectively  $(P = 3.1 \times 10^{-5} \text{ and } P = 2.1 \times 10^{-6}, \text{ respectively; } \chi^2)$ . Furthermore, untreated and TSA-treated sites were positioned nonrandomly relative to each other, with significant TSA enrichment between two peaks of untreated enrichment ( $P = 5.4 \times 10^{-23}$ ;  $\chi^2$ ).



chromosome condensation through the recruitment of chromatin remodeling factors such as the SWI/SNF complex (Agalioti et al. 2002). Several other proteins, including CBP and the general transcription initiation factor, TFIID, are also recruited to histones through interactions with acetyl-lysine residues (Agalioti et al. 2002; Fukuda et al. 2006). Acetylated histones are typically enriched in the promoter regions of active genes and we found that Nup93 was highly enriched in these regions following TSA treatment. In addition, we identified >100 transcription factor-binding motifs in the TSA-treated Nup93-binding sites, a finding indicative of the functional repositioning of chromatin. Further evidence for a role of the mammalian nuclear transport machinery in transcriptional regulation comes from the transport protein CAS, the mammalian homolog of the yeast exportin, Csel. A recent study has shown that CAS associates with p53 at the promoter regions of several proapoptotic genes and contributes to their transcriptional activation (Tanaka et al. 2007).

Histone methylation also plays an important role in transcriptional regulation. Several H3K4 methylations are associated with actively expressed genes, while the H3K27me3 modification is associated with silent genes (Barski et al. 2007). Accordingly, Nup93-binding sites in TSA-treated cells are significantly enriched in H3K4me1–3 and depleted in H3K27me3 (Table 1). The opposite histone methylation biases are observed for untreated Nup93-binding sites, which are random with respect to H3K4me1–3 and enriched in H3K27me3.

#### Transcriptional regulation at the NPC

TSA-treated Nup93-binding sites were enriched in promoter regions and were closer to transcription start sites than their untreated counterparts. We also show that following TSA treatment, Nup93-binding sites are enriched in RNA PolII-associated regions and are closer to genes exhibiting differential expression. These regions include genes that are both highly expressed and strongly Figure 6. Model summarizing the factors involved in Nup93-genomic interactions. Untreated Nup93-binding sites contain a mix of active and inactive genes at the nuclear periphery, with enrichment for silent histone methylation marks ("Me" inside red circles). Following TSA treatment, Nup93-binding sites are enriched in several factors including the insulator protein CTCF, the histone variant H2A.Z (yellow spheres), and active histone methylation marks ("Me" inside green circles). Histone acetylation ("Ac" inside green circles) was enriched at the nuclear periphery following TSA treatment. One of the two NPCs on the *right* is proximal to a transcription start site, representing the promoter-binding and transcription factor-binding motif enrichment observed in Nup93-binding sites upon TSA treatment.

repressed following drug treatment. The presence of both active and repressed genes at nuclear pores has been observed previously in S. cerevisiae (Casolari et al. 2004). The role of the NPC as a boundary element may help explain these observations (Ishii et al. 2002). Interestingly, TSA-treated Nup93-binding sites are enriched in CTCF-associated regions. CTCF, a protein that recognizes cis-acting transcriptional insulator elements and marks histone modification boundaries (Splinter et al. 2006; Barski et al. 2007), copurifies with the histone variant H2A.Z, as well as two components of the nuclear envelope, lamin A/C and importin  $\alpha 3/\alpha 1$  (Yusufzai et al. 2004). These associations suggest that CTCF interacts with the nuclear periphery and that regions of CTCFmediated regulation are repositioned proximal to nuclear pores following TSA treatment.

The balance of HAT and HDAC activity is essential for the maintenance of boundaries between active and repressed chromatin (Kimura et al. 2002; Suka et al. 2002). Indeed, boundary elements recruit both HATs and HDACs in order to prevent spreading of silent and active chromatin, respectively (West et al. 2004; Yusufzai et al. 2004). Mutations in HATs and HDACs can significantly shift the borders of functional regions in the genome. For example, deletion of a subunit of the Rpd3 HDAC complex in yeast leads to aberrant transcriptional initiation (Carrozza et al. 2005). The mammalian nuclear periphery contains at least one HAT and HDAC, CBP and HDAC3, respectively. CBP interacts with the NPC, while HDAC3 associates with the lamin-binding protein LAP2<sub>β</sub>. These proteins, along with CTCF, could be important components of boundary elements at the mammalian nuclear periphery.

Evidence from work in *S. cerevisiae* has shown that recently activated genes remain at the nuclear periphery for hours after they have been repressed (Brickner et al. 2007). This localization differs from their preactivated positioning, which appears to be random with respect to the nuclear periphery (Drubin et al. 2006). This novel form of transcriptional memory is functional; it was

shown that the recently repressed *GAL1* gene was activated faster than a long-term repressed form. In yeast, Nup2p and the histone variant H2A.Z were involved in this process. Interestingly, we found that TSA-treated Nup93-binding sites were enriched in H2A.Z-binding sites. The presence of H2A.Z at sites of NPC-chromatin interaction could mean a similar form of transcriptional memory is present in human nuclei. Alternatively, H2A.Z site enrichment could reflect the role of this histone variant in peripherally localized boundary activity as H2A.Z has been shown to copurify with CTCF in HeLa cells and block the spread of telomeric heterochromatin in yeast (Meneghini et al. 2003; Yusufzai et al. 2004).

#### Chromosome-wide changes in nuclear organization

Large-scale changes in NPC-chromosome interactions were observed after treatment with TSA. Stretches of chromosomes 5 and 7 were no longer associated, while chromosome 16 retained the majority of its associated regions (Fig. 2A). These changes appear to be part of an extensive nuclear reorganization event in response to the global increase in histone acetylation.

The NPC has been implicated in boundary activity in *S. cerevisiae* and chromosome looping is thought to occur at the nuclear periphery in *Drosophila* (Gerasimova et al. 2000; Ishii et al. 2002). In addition, evidence has emerged that two distally localized genes on murine chromosome 7 can be localized to the same nuclear compartment upon activation (Osborne et al. 2004). It is possible that CTCF-mediated insulator activity near mammalian NPCs could generate similar loops that functionally isolate regions of activation or repression at the nuclear periphery.

Chromosome G-band patterns represent regions of genomic enrichment in either heterochromatin or euchromatin. We show that Nup93-binding sites are more likely found in heterochomatin in untreated cells, while favoring euchromatin following TSA treatment. The significant and opposing histone modifications and G-band enrichments found in Nup93-binding sites reflect a global reorganization event that exchanges silent for active chromatin at NPCs.

In summary, we show that the human genome interacts with Nup93, a component of the mammalian nuclear pore. Nup93 is associated with regions of transcriptional repression and enriched heterochromatin content in HeLa cells. Upon global histone acetylation mediated by the HDAC inhibitor, TSA, Nup93 associates with regions important for transcriptional regulation. Our results hold significance for both the general mechanisms of gene expression in mammalian cells and the global effects of histone modifications on nuclear organization.

#### Materials and methods

#### Cell culture and drug treatment

HeLa S3 cells were grown in DMEM supplemented with 10% FBS. Exponentially growing cells were incubated in the presence

of 40 ng/mL TSA (Sigma) for 12 h. Cells were grown to near confluency before fixation and harvesting for ChIP-chip, indirect immunofluorescence, FISH, and expression profiling.

#### Immunofluorescence

HeLa S3 cells were grown to near confluency and adhered to polylysine-coated coverslips. Cells were prepared for labeling as described previously (Brown et al. 1996). Primary antibodies used include anti-lamin B diluted 1:50 (Santa Cruz Biotechnology), anti-Nup93 diluted 1:200 (D. Forbes), anti-acetylated H4K5 diluted 1:100 (Abcam), and anti-acetylated H3K9 diluted 1:100 (Upstate Biotechnology). Other antibodies used for preliminary ENCODE studies include anti-Nup62 (BD Transduction Laboratories), anti-Nup107, and anti-Nup205 (generous gifts from V. Cordes). The secondary antibody used to detect polyclonal primaries was goat anti-rabbit Alexa594 diluted 1:1000 (Molecular Probes). Images were acquired on a Nikon E800 epifluorescence microscope equipped with a Radiance 2000 confocal laser scanning system (Bio-Rad).

#### FISH

Bacterial artificial chromosomes (BACs) were purified using a Large-Construct Kit (Qiagen). Digoxigenin-dUTP was incorporated into purified BACs using a DIG Nick Translation Kit (Roche). For each sample, 40–160 ng of DIG-dUTP-labeled probe were combined with 40 µg of unlabeled COT1 DNA (Roche) and 20 µg of salmon sperm DNA (Ambion), a portion of which was used for hybridization. Cells were prepared and probes hybridized as described previously (Solovei et al. 2002). The probe signal was detected using an antibody enhancer kit (Roche). Vectashield containing DAPI (Vector Laboratories) was added to the slides and the coverslips were sealed with nail polish. More than 200 cells were counted for each experimental condition. FISH signal was counted as peripheral if it either overlapped with or was immediately adjacent to the nuclear periphery signal.

#### ChIP-chip

ChIP was performed essentially as described (Brodsky et al. 2005) with the following differences. Chromatin was prepared from four independently grown batches of cells-two untreated and two treated with 40 ng/mL TSA for 12 h before fixation to generate replicates for each condition. Cleared chromatin was incubated overnight with polyclonal Nup93 antibody at 4°C. After incubation, a fresh batch of 50:50 protein A/G sepharose beads were added to the chromatin/antibody solution and incubated overnight. Beads were washed five times for 10 min each, followed by DNA elution overnight, proteinase K treatment, phenol extraction, and RNase treatment. Isolated DNA was amplified isothermally using random nonamers and Klenow polymerase (Invitrogen) for 2 h, yielding ~4 µg of DNA per ChIP. DNA was prepared and hybridized to Affymetrix GeneChip Human Tiling 2.0R ChipE (chip5) arrays using fragmentation, hybridization, staining, and scanning procedures described previously (Kennedy et al. 2003). A sample of chromatin was set aside before immunoprecipitation and used to represent the input DNA.

#### Expression profiling

Total RNA was extracted with Trizol and purified with an RNeasy Mini Kit (Qiagen) with an added DNase step (RNasefree DNase; Qiagen). Fifteen micrograms of RNA from each

sample were sent to the DFCI Microarray facility for labeling and hybridization to Affymetrix Human Genome U133 Plus2.0 expression arrays. Raw hybridization intensity (CEL) files from three biological replicates were processed using the RMA method (Bolstad et al. 2003; Irizarry et al. 2003a,b). Significant differentially expressed genes were considered those with average log<sub>2</sub>-transformed fold-difference ratios (TSA:Untreated)  $\geq$ 1.2 or less than or equal to -1.2 and *P* < 0.01 (two-tailed Student's *t*-test).

#### Analysis of genomic binding data

All genomic track data (sequence conservation scores, SNPs, Refseq genes, and Giemsa ideograms) were downloaded July 2006 from the University of Californai at Santa Cruz genome bioinformatics Web site (ftp://hgdownload.cse.ucsc.edu/ goldenPath) for the May 2004 assembly. To maintain consistency in the track variables, these originally downloaded data were used for all subsequent analyses. Duplicate Nup93 ChIP samples were submitted to the Dana-Farber Cancer Institute Microarray Core facility for hybridization. Enrichments in ChIPs compared with input were calculated from raw intensity (CEL) files using a nonparametric statistical method implemented in Affymetrix Tiling Analysis Software (TAS) (Ghosh et al. 2006). Binding significances (expressed as probe P-values) were used to detect statistical dependencies between Nup93binding enrichments and sequence conservation (see the Supplemental Material for a technical description of the method, including statistical significance assessment and false discovery).

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