Large-Scale Functional Organization of Long-Range Chromatin Interaction Networks

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SUMMARY

Chromatin interactions play important roles in transcription regulation. To better understand the underlying evolutionary and functional constraints of these interactions, we implemented a systems approach to examine RNA polymerase-II-associated chromatin interactions in human cells. We found that 40% of the total genomic elements involved in chromatin interactions converged to a giant, scale-free-like, hierarchical network organized into chromatin communities. The communities were enriched in specific functions and were syntenic through evolution. Disease-associated SNPs from genome-wide association studies were enriched among the nodes with fewer interactions, implying their selection against deleterious interactions by limiting the total number of interactions, a model that we further reconciled using somatic and germline cancer mutation data. The hubs lacked disease-associated SNPs, constituted a nonrandomly interconnected core of key cellular functions, and exhibited lethality in mouse mutants, supporting an evolutionary selection that favored the nonrandom spatial clustering of the least-evolving key genomic domains against random genetic or transcriptional errors in the genome. Altogether, our analyses reveal a systems-level evolutionary framework that shapes functionally compartmentalized and error-tolerant transcriptional regulation of human genome in three dimensions.

INTRODUCTION

Long-range chromatin interactions are pervasive in the human genome and serve to regulate gene expression (Göndör and Ohlsson, 2009; Schoenfelder et al., 2010). Proximity ligation in combination with next-generation sequencing has recently enabled us to explore genome-wide spatial crosstalk in the chromatin (Fullwood et al., 2009; Lieberman-Aiden et al., 2009). By implementing Chromatin Interaction Analysis using Paired End Tags (ChIA-PET) (Fullwood et al., 2009), we recently mapped all-to-all chromatin interactions associated with RNA polymerase II (RNAPII) at base-pair resolution. In addition to widespread promoter-enhancer chromatin interactions, our analysis revealed a range of distinct types of chromatin crosswirings, including promoter-enhancer, enhancer-enhancer, promoter-terminator, and, intriguingly, promoter-promoter interactions. These interactions constitute a basic topological template for transcriptional coordination (Li et al., 2012). The observation of most interest was that interacting promoters not only correlate with gene coexpression, but can also regulate each other’s transcriptional states, which blurs the traditional definitions of gene-regulatory elements in the genome. These observations support the notion of a chromatin interactome encompassing a dense repertoire of regulatory elements for transcriptional regulation.

Whole-genome chromatin interaction data sets are too complex to be analyzed by conventional approaches. To gain a better understanding of these interactions, we performed a complex network analysis by integrating chromatin interactions and several other genomic data sets (Table S1). Network analysis has emerged as a powerful tool for obtaining novel insights into complex systems. The nonrandom topological properties of most real-world networks are strongly associated with their robustness and functional organization (Albert et al., 2000; Barabási and Albert, 1999; Barabási and Oltvai, 2004), which has motivated molecular biologists to explore cellular regulation using a systems approach. Although most cellular networks, such as gene-regulatory, metabolic, protein-protein interaction, and signaling networks, are being widely studied, the extensive communications among regulatory elements in the genome have not been viewed in a complex-network context (Singh Sandhu et al., 2011).
were called at FDR sequencing using the Illumina GAIIx platform. Unique PETs were mapped DNA material was subjected to PET extraction and next-generation in the diluted conditions. After the chromatin complexes were decrosslinked, (blue). Specific linkers were added to the open ends and subsequently ligated plexes bound with RNAPII (green) were pulled down using 8WG16 antibody.

Figure 1. Simplified Illustration of the ChIA-PET Technique and the Network Construction.
(A) ChIA-PET technique. Formaldehyde crosslinked chromatin was sonicated and chromatin complexes bound with RNAPII (green) were pulled down using 8WG16 antibody (blue). Specific linkers were added to the open ends and subsequently ligated in the diluted conditions. After the chromatin complexes were decrosslinked, DNA material was subjected to PET extraction and next-generation sequencing using the Illumina GAIIx platform. Unique PETs were mapped back to the reference genome (Hg19) and statistically significant interactions were called at FDR < 0.05 using the ChIA-PET tool (Li et al., 2010). To construct the network, the redundancy in the data were removed by merging the overlapping interaction sites. The cutoff taken to merge the overlapping sites is shown in Figure S1A.

We show that a large proportion of the human genome converges to a complex hierarchical network to orchestrate transcription in functionally compartmentalized and evolutionarily constrained chromatin communities. We demonstrate that the hubs (i.e., nodes with a disproportionately high number of interactions) and spokes (i.e., nodes with fewer interactions) of the network exhibit distinct functional and etiological properties. Together, our findings present a chromatin-level explanation for how disease-associated mutations are tolerated during development and how the key cellular genes maintain their consistent and error-free expression.

RESULTS

Transcription-Associated Chromatin Interactions Form a Complex Hierarchical Network

ChIA-PET is a logical extension of proximity-ligation-based techniques such as chromosomal conformation capture (3C) and circularized 3C (4C). In brief, the chromatin is crosslinked using 1% paraformaldehyde and sonicated, and chromatin complexes are pulled down using a specific antibody against a particular protein factor (in this case, 8WG16 antibody against RNAPII). Specific linkers are added to the open ends and the complexes are ligated in the diluted conditions. The ligated material is then subjected to PET extraction and next-generation sequencing (Figure 1A). Using K562 and/or MCF7 ChIA-PET data sets (Li et al., 2012), we constructed an RNAPII-associated chromatin interaction network (ChIN) by denoting the distinct genomic sites as vertices (nodes) and statistically significant (false discovery rate [FDR] < 0.05; Extended Experimental Procedures) chromatin interactions among those sites as edges (links) (Figures 1B and S1A; Extended Experimental Procedures). To remove redundancy from the ChIA-PET data, we merged the neighboring overlapping sites as illustrated in Figures 1B (left panel) and S1A. Several randomly selected intra- (cis) and interchromosomal (trans) interactions had been validated with Chromosomal Conformation Capture (3C) and DNA fluorescence in situ hybridization (FISH) assays in our earlier study (Li et al., 2012). The topological and functional properties of the ChIN presented in this study were also scrutinized against the artifacts of genomic rearrangements in the MCF7 and K562 cell lines (Figures S1C, S1D, S3C, S4C, S5C, and S6A).

The strategy elaborated in Figure 1 and Extended Experimental Procedures yielded a comprehensive network map of chromatin interactions with ~10,000 connected network components. Surprisingly, however, ~40% of the total nodes formed a giant network component of 36,748 nodes sharing 55,039 links among them (Figure S1E; Tables S2 and S3), suggesting that the vast majority of the transcriptionally active genome displays widespread communication, implying an unprecedented level of regulatory influence among genes and their associated genetic elements. This could lead to common pleiotropic gene effects. It is obvious that most of these interactions would not occur at the same time in the same cell due to spatial constraints, but rather represent highly dynamic interactions across a large population of cells (Sandhu et al., 2009).

Except for some of the properties concerning the topology of the network, which were determined using the giant network component, most analyses presented in this study were performed on the whole network map, including the smaller components. Functional analyses were performed on the K562 and/or MCF7 data sets, depending on the availability of other related genomic data sets, although the overall properties were coherent between the two cell lines as shown in Figure S1.

The giant network component of ChIN followed a scale-free-like degree distribution, according to which very few nodes would have a disproportionately large number of interactions, and most others would be weakly connected (Barabási and Albert, 1999) (Figure 2A, top panel). We confirmed this observation for the complete networks for both the cell lines, as well as for the publicly available Hi-C data set (Lieberman-Aiden et al., 2009) (Figure S1F). Most real-world networks exhibit scale-free-like behavior (Albert et al., 2000), and the property ascribes error tolerance against random malfunctions, indicating that the ChINs allow for robust systems.

Furthermore, the ChIN displayed a hallmark of hierarchical network topology, characterized by a strong negative correlation between the degree (i.e., number of interactions per locus) and the clustering coefficient (i.e., the tendency of a node to form triangles; Pearson correlation coefficient [PCC] = −0.81; Figure 2A, lower panel). A hierarchical network exhibits high modularity in addition to scale-freeness and is an inherent property of biological networks that governs functional organization.
We further illustrated the hierarchical nature of the ChIN in an example in which the long-range (<200 Kb) interaction clusters, namely the HIST1 gene clusters, further converge via super-long-range (>500 Kb) interactions in a hierarchical manner (Figure 2B). The convergence of these three HIST1 clusters correlates significantly with their coexpression (Li et al., 2012), suggesting that super-long-range interactions are important for the global coordination of distant gene clusters. Indeed, we observed that the super-long-range and trans chromatin interactions are critical for maintaining the overall ChIN topology, despite having a lower frequency of interactions and accounting for a smaller proportion of all chromatin interactions (Figure S1G). Abolishment of these interactions would break the ChIN into smaller disconnected components and consequently alter the global coordination among distant genes, as in the case of HIST1 clusters.

**Strong Regulatory Marks Govern the Modular Topology of the ChIN**

To further investigate the modular nature of the ChIN, we used the ModuLand algorithm (Kovács et al., 2010) and mapped the network modules, which we refer to as chromatin communities.

**Figure 2. Topological Properties of Transcription-Associated ChIN**

(A) Top panel: Log-log plot of the node degree distribution for ChIN constructed from the K562 ChIA-PET dataset (plots for other data sets are given in Figure S1F). The plot shows heavy tailed distribution atypical of scale-free-like networks. Bottom panel: Log-log scatter plot of node clustering coefficients and degree. The strong inverse correlation is atypical of scale-free networks.

(B) An example of hierarchical chromatin organization on chromosome 6. Three distant HIST1 gene clusters (HC1, HC2, and HC3) converge in a hierarchical manner, as shown in the heat-map representation of ChIA-PET data.

(C) Hierarchical topology of K562 ChIN. The color gradient represents the hierarchical organization of chromatin communities (#1173); red and blue indicate the most central and most peripheral chromatin communities, respectively, as identified by the ModuLand algorithm. Other network properties for K562 and MCF7 ChINs are given in Figures S1 and S2.

(D) Community centralities of nodes having distinct chromHMM profiles (Ernst et al., 2011) in the K562 cell line. Red, active/weak promoter; magenta, poised promoter; orange, strong enhancer; yellow, weak enhancer; blue, insulator; green, transcribed region; gray, repressed region; white, all. Asterisks indicate the chromatin types for which the community centralities were significantly greater when compared with all nodes. The Mann-Whitney U test was used to calculate the p values (<2.2e-16 for each asterisk). See also Figure S2J.

(E) Public and private nature of enhancers. Shown is the bar plot for the proportion of strong and weak enhancers in private and public enhancers. The p value was calculated using Fisher’s exact test. A box-plot representation of overall promoter interactions of strong and weak enhancers is given in Figure S2K.

(F) High-resolution example of physical interactions among distinct chromatin types. Shown are the tracks for UCSC known genes, RNA-Seq, RNAPII enrichment, ChIA-PET, and chromatin types (chromHMM) in the K562 cell line. The network constructed from this locus is shown in the right panel. Nodes are colored according to their chromatin types and their size is scaled to their degree.
A chromatin community can include loci from different parts of the genome and, therefore, represents an extension of the multigene complex (Li et al., 2012) that was methodically restricted in the genomic range of 1 Mb. The ModuLand algorithm identified 1,173 communities in the giant component of the K562 ChIN (Figure 2C). Most of the intracommunity interactions were enriched with cis interactions, whereas intercommunity interactions were mostly mediated by super-long-range or trans chromatin interactions (Figure S1H), adhering to the fractal nature of chromatin folding (Lieberman-Aiden et al., 2009; Sexton et al., 2012). We then asked whether distinct genomic elements and chromatin types, as identified by Ernst et al. (2011), could contribute distinctly to the modular topology of the ChIN (Figures 2D and S2). We calculated a centrality score, called the community centrality score, which is a cumulative measure of the influence of the entire network to the given node, and is maximal at the central core of the network modules (Kovács et al., 2010). The active/weak promoter and the strong enhancer elements showed significantly greater community centrality scores than the other categories, suggesting that the modular structure of the ChIN is primarily shaped around these genomic elements (Figures 2D and S2J). Therefore, a single promoter can have multiple enhancers, and a single enhancer can have multiple target promoters, mounting the regulatory complexity of the genome. We further classified the enhancer nodes as private or public enhancers based on their attainment by one or multiple (≥2) gene promoters, respectively. Interestingly, >70% of the public enhancers were also strong enhancers (p < 2.2e-16), whereas private enhancers were equally represented by strong and weak enhancers (Figures 2E and S2K). We scrutinized and confirmed this observation against the possibility of differential enrichment of RNAPII at strong and weak enhancer sites by restricting the analysis to sites of similar levels of RNAPII enrichment (Figure S2L). A specific example is shown in Figure 2F. Three active promoters, three strong enhancers, and three weak enhancers converged to a network complex. Here again, the active promoters b, d, and g were central to the network segment, undergoing three, five, and five interactions, respectively. Similarly, the strong enhancer c interacted with all three active promoters in the locus, whereas the other strong enhancer, h, interacted with two of the three active promoters (Figure 2F). On the other hand, the weak enhancers f, i, and j were peripheral to the chromatin communities connecting to individual genes (Figure 2F). Therefore, strong enhancers not only have a greater enhancing effect on transcription, which is the original definition of strong enhancers (Ernst et al., 2011), but also have the potential to regulate multiple genes (pleiotropic regulation).

We then sought a possible explanation for the greater centrality of strong/public enhancers. We assessed the correlations with individual histone modifications (Figure S2E). Hyperacetylation of nodes was associated with higher degree, which is in line with the supposition that hyperacetylation endows greater chromatin mobility (Brown et al., 2008; Krajewski and Becker, 1998). We previously showed that the abundance of chromatin interactions correlates with genomic descriptors such as SINE and LINE densities (Li et al., 2012). Therefore, we controlled our present analyses for these genomic correlates. The partial correlations controlled for SINE and LINE densities clearly suggested that the correlations between degree and SINE/LINE densities do not account for the correlations observed between node degree and enrichment of chromatin marks (Figures S2H and S2I). Furthermore, the elements bound with chromatin remodeling factors such as BRG1 and IN1 were more interactive than the rest (Figure S2M). We hypothesize a prominent role for chromatin-remodeling factors in determining the ChIN topology, which is also in line with earlier reports on individual loci (Kim et al., 2009; Ni et al., 2008; Zhang et al., 2006). Surprisingly, contrary to the prevailing view on the role of CTCF in chromatin architecture, we did not observe strong association between CTCF binding and the number of RNAPII-associated chromatin interactions (Figures 2D and S2E), suggesting that CTCF orchestrates the genome architecture in a manner reasonably distinct from that of RNAPII, possibly by enclosing the chromatin communities in large chromatin compartments and thus ascribing a basic chromatin skeleton for transcription-associated complex connectivity (G.L. et al., unpublished data). Therefore, the notion that CTCF is the “master-weaver of the genome” needs to be reconciled by taking into account the role of other factors in three-dimensional genome organization.

Chromatin Communities Organize Functional Compartmentalization

The modular nature of the ChIN raises the possibility of functional compartmentalization of chromatin in the nucleus. To assess the functional enrichment in chromatin communities using network-based ontology tools (see Experimental Procedures), we focused only on promoter-promoter interactions. This resulted in the decomposition of the giant network into several smaller network components (Figures S3A and S3B). We analyzed the enrichment of gene ontology (GO) terms among the top 30 network components, containing at least 20 genes each (Extended Experimental Procedures; Table S4). Out of 30 such subnetworks, we observed the enrichment (FDR < 0.01) of one or more functions in 18 (60%; Figure 3A, left panel). Using an example of a network component, we further showed that the enrichments of multiple functions were localized in distinct chromatin communities within a network component (Figure 3A, right panel; Table S4). Figure S3C illustrates that the observed functional organization is not an artifact of genomic rearrangements. We further validated two interesting examples using DNA FISH experiments: (1) A common enhancer interacted with two brain-related proteases, both expressed in MCF7, in cis and trans. Interestingly, the enhancer locus was specifically conserved among primates, hinting at the possibility of primate-specific gene expression regulated via long-range chromatin interactions (Figure S3D). (2) Two small nuclear noncoding RNA loci were found to be interacting in trans (Figure S3E). DNA FISH experiments confirmed the significant interaction frequencies among the loci involved (p = 6.8e-07 and 2.2e-16, respectively; binomial test; Figures S3D and S3E).

Often, not all of the genes in a community served the same function, suggesting that the chromatin communities were not absolutely dedicated to a particular function and often incorporated overlapping secondary functions, which might be indirectly...
Chromatin Communities Are Evolutionarily Constrained

We speculated that evolutionary constraints may have shaped the functional compartmentalization of chromatin. To test this, we analyzed the density of interactions (i.e., the number of interactions per Mb) within genomic blocks that were syntenic or nonsyntenic to chimp and mouse genomes. We used a moderate level of coarse graining to map syntenic blocks using the Centery algorithm (Sinha and Meller, 2007) (Table S1), which revealed human-chimp and human-mouse syntenic blocks covering 48% and 45% of the human genome, respectively. Subsequent analysis revealed a nonrandomly higher density of chromatin interactions within the syntenic blocks than in the nonsyntenic blocks (Figures 4A–4C). Moreover, the frequency of loops connecting syntenic and nonsyntenic blocks was also very low (four and seven loops per megabase for human genomic blocks syntenic to chimp and mouse genomes, respectively). Because syntenic regions are expected to show higher gene density and expression, we performed some control analyses by selecting syntenic regions are expected to show higher gene density and expression, we performed some control analyses by selecting syntenic to chimp and mouse genomes, respectively). Because syntenic regions are expected to show higher gene density and expression, we performed some control analyses by selecting syntenic and nonsyntenic regions of similar gene density and expression. The analysis consistently showed a significantly higher density of chromatin interactions in the syntenic blocks (Figure S4A). Furthermore, a brief analysis on some of our unpublished ChIA-PET data for mouse embryonic fibroblasts (MEFs) suggests that RNAPII-mediated chromatin interactions tend to accumulate in mouse regions that are syntenic to the human genome (Figure S4B), reconciling the evolutionarily constrained nature of chromatin communities. The observed synteny of chromatin communities was also robust against the possibility of artifacts due to genomic rearrangements in human cancer cell lines (Figure S4C). The above observations were also supported by our analyses of human-mouse orthologous genes, conservation of genomic neighborhood, asynonymous sequence divergence between human-chimp genomes, and mammalian phastCons conservation scores (Figures S4D–S4G).

To obtain further details, we plotted the genomic distance between interacting loci of human genome against that of corresponding sites in the mouse genome (UCSC’s liftOver; 95% sequence similarity; Figure 4B). We had three key

related to the primary function. For instance, a protease (SIPA1), a protease inhibitor (CST6), and a DNA repair-related factor (MUS81) were embedded in a community significantly enriched in defense/immunity-related genes (FDR < 0.01; Figure 3B). We hypothesize that such interactions might orchestrate a coordinated response to external stimuli. The overlapping functional enrichments in chromatin communities could also help in efficient reconfiguration of community function in response to external signals, as proposed earlier (Mihalik and Csermely, 2011; Pál et al., 2006).

Importantly, the community structures were largely conserved between MCF7 and K562 cell lines. Out of 1,783 total gene communities with at least three genes each, 1,279 (71%) showed >75% overlap in MCF7 and K562 cell lines (Figure S3F). Upon closer examination, we observed that the nodes with K562- and MCF7-specific interactions were often embedded in the communities enriched with genes common to both cell lines (Figure S3G). Therefore, the cell-line specificity is defined either by individual long-range transient interactions that do not converge to the same community or by fine level differences in chromatin looping within communities. Such fine differences in chromatin architectures have also been observed by others in different contexts (Filion and van Steensel, 2010; Lienert et al., 2011; Peric-Hupkes et al., 2010).
observations: (1) Paired coordinates for 32% and 38% of total interactions in K562 and MCF7 cell lines, respectively, could be directly mapped to the mouse genome as compared with 13% of randomly selected coordinates with the same span distribution (p < 2.2e-16). (2) The correlation between locus distances in human and mouse genomes was significantly higher for interacting loci than for randomly selected pairs of loci of the same span distribution, supporting the higher conservation of synteny for interacting pairs (PCC = 0.90, FDR = 0.004; Figures 4B and S4H). (3) There were very few instances in which proximal mouse genomic sites were rearranged to distant sites in the human genome (Figure 4B), and in contrast, there was nonrandomly higher representation (FDR = 0.009; Figure S4H) of sites that were distant in the mouse genome but were rearranged to proximal domains in the human genome. This suggests two possibilities: (1) The physical interactions, if any, among distant genomic sites in the ancestor genome may have served as an evolutionary mechanism to translocate the interacting loci to proximal regions in the human genome (Figure 4B). Indeed, spatial proximity has been shown to mediate genome rearrangements associated with cancer genomes (Lin et al., 2009). Therefore, the evolution of gene clusters may have been mediated by long-range chromatin interactions. (2) If the distant loci had no interaction in the ancestor genome, then the newly acquired linear proximity of loci through the process of translocation may have been the sole driving force behind chromatin interactions. A detailed analysis of RNAPII-associated chromatin interaction data from other lower species would allow further examination of such observations in the future.

We previously demonstrated a nonrandomly higher correlation among expression profiles of interacting genes across several gene-expression data sets (Li et al., 2012). Along similar lines, we now asked whether the expression of genes with promoter-promoter interactions is evolutionarily more conserved than the rest. To address this issue, we obtained an expression-divergence data set for multiple human and chimpanzee tissues from the literature (Khaitovich et al., 2005). Indeed, the genes that had promoter-promoter interactions showed a significantly lower divergence of gene expression and sequence during the evolutionary split of chimpanzees and humans (Figures 4D and S4F). These observations highlight the strong evolutionary selection of advantageous chromatin communities for functional coordination of related genes.

Disease-Associated Genetic Errors Are Enriched Among Spokes

Genetic errors in distal noncoding elements could influence the expressivity of the genome (Freedman et al., 2011; Mu et al., 2011). One way in which genetic errors could influence gene activity and, consequently, the phenotype is via long-range chromatin interactions (Ahmadiyeh et al., 2010; Ferrai and Pombo, 2009; Sandhu et al., 2009; Steidl et al., 2007; Visel et al., 2009). Therefore, we focused on disease-associated SNP data obtained from the genome-wide association studies (GWAS) catalog (Hindorff et al., 2009). The representation of GWAS SNPs among genic and intergenic sites did not differ from that of the overall representation of genic and intergenic sites in the giant ChIN (Figures S5A and S5B), suggesting that
disease-associated SNPs are equally probable for genic or intergenic regions in the ChIN. Although the target genes of intergenic GWAS SNPs determined by ChIA-PET showed good correspondence (70%) with the targets reported in the GWAS catalogue, there were SNPs that had different or additional targets other than with the known ones (Figures S5D and S5E), suggesting that the chromatin interaction data can help one determine the precise targets of noncoding SNPs. Because >95% of total genes engaged in RNAPII-associated long-range interactions were related to the transcriptionally active state of the genes (Li et al., 2012), we reason that if the gene is expressed in the tissue for which the GWAS study was performed, most likely it will also have the corresponding chromatin looping to regulatory elements. This is also supported by the observation that 79% of total interactions involving genes commonly expressed in the MCF7 and K562 cell lines were conserved among cell lines. Therefore, the cell-lineage discrepancy of integrated GWAS data and the ChIA-PET data sets might not be entirely incoherent in this context.

By mapping the GWAS SNPs onto the ChIN, we further showed that genomic elements with at least one disease-associated SNP were enriched to a lower degree (3–6; spokes) and that the ChIN hubs were devoid of such SNPs (FDR = 0.001; Figure 5A; for examples, see Figures 5C, 5D, SS, and SSE, and Li et al. [2012]). This observation was also true for distinct types of promoters or enhancer loci (Figure S5F). A relatively weaker second dip in the FDR curve in Figure 5A appeared to be due to a locus having an abnormally high copy number. Therefore, we assessed the representation of all of the nodes with GWAS SNPs in the normal- and abnormal-copy-number regions. More than 90% of the disease-associated SNPs were found to be in the normal-copy-number range (Figure S5C). Thus, we argue that our observation is not an artifact of the genomic abnormalities of cancer genomes. Furthermore, 80% of the chromatin interactions mediated by the nodes having GWAS SNP were generally restricted within the chromatin community (p = 1.07e-07, Fisher’s exact test), suggesting that in general, the chromatin interactions of disease-linked SNPs associate with the spatially localized dysregulation of a limited number of genes.

Similar observations were reported for most disease genes from the morbid entries in the Online Mendelian Inheritance in Man (OMIM) database (Hamosh et al., 2000), where hubs were mostly devoid of genes associated with disease phenotypes (Figure S5G). Therefore, we asked whether disease-associated
regulatory loci were selected against the possibility of erroneous interactions, like those reported elsewhere (De and Michor, 2011; Lin et al., 2009) or otherwise, by restricting themselves to fewer interactions. To test this, we compared the chromatin interactions of loci associated with germline and somatic cancer mutations (Futreal et al., 2004). Comparison revealed that the loci harboring cancer-associated germline mutations are less interactive than the ones with somatic mutations, despite the least difference in RNAPII enrichment between the two types of loci (Figures 5B and S5H). The data suggest that disease-associated regulatory elements generally function locally and have a rather limited repertoire of interactions. We propose that the possibility of erroneous genomic interactions and consequently functional dysregulation is minimized by means of restricting the total number of interactions of loci that are important for normal organism development. The germline transmission of genetic lesions having fewer interactions can thus be better tolerated. By contrast, somatic mutations are not under any selection pressure and thus could have a relatively wider exposure to the regulatory cross-wirings in the chromatin. Therefore, based on our analysis, we propose that disease-causing SNPs may generally be trapped in local chromatin communities that affect rather limited phenotypic traits, such as those shown in Figures 5C, 5D, S5D, and S5E.

Hubs Conform to a “Rich-Club” Organization of Key Cellular Functions

Given that the hubs had different characteristics as compared to spokes, we focused on the hubs (degree $\geq 60$) to determine whether the hubs showed any particular behavior. Interestingly, we found that the top hubs had a preferential link structure, i.e., a rich-club, among themselves (Figure 6A and 6B; total 385 hubs).
promoters and 2,386 other elements centered on 25 hubs). By analyzing our in-house data sets of genomic rearrangements uncovered by the genomic DNA paired end tag (G-PET or DNA-PET) sequencing approach (Hillmer et al., 2011), we ensured that the rich-club is not an artifact of genomic rearrangements (Figure S6A).

Rich-clubs in several real-world networks were previously reported and were proposed to contribute to the greater robustness of the network against random hub failures (Colizza et al., 2006; Shi and Mondragon, 2004) and to enhance global cooperation in several biological systems (Bastolla et al., 2009; Saavedra et al., 2011). To test this hypothesis, we disrupted rich-club connectivity and performed a network-resilience analysis of the ChIN. We observed that the ChIN with an intact rich-club had greater topological robustness than the one with a disrupted rich-club (p = 0.004; Figures S6B and S6C).

Functionally, the rich-club genes (n = 385) were enriched in essential cellular functions, including chromatin assembly (e.g., HIST1 genes, TTF2, MTA2, TAF6, and BRD2), cellular organization (e.g., ACTB, ACTG1, CIT, KIF1B, KIF2C, and TRIP6), and primary metabolic processes (e.g., SLC17A7, SIC3A2, ITPA, ATP1A2, DXH29, MAP4K4, EEF1A2, and PLEC1), when compared to spokes (degree 1–6) or the target genes of GWAS SNPs, which were relatively enriched in development-related functions (FDR < 0.05; Figures 6C, S6D, and S7A). More importantly, 62% of the rich-club genes, and the genes interacting with them, which could be mapped to mouse phenotype information available from the Mouse Genome Informatics (MGI) database (Shaw, 2009), had “lethal,” “death,” or “mortality” phenotypes in mouse (red-colored nodes in Figure 6D and Table S5), whereas only ~23% of the mouse genes had shown these phenotypes (Shaw, 2009; p = 6.34e-08). A specific example of rich-club organization across an ~14 Mb region on chr6 is shown in Figure 6E. The hubs centered on JARID2 (chromatin modifying), E2F3 (transcription factor), and c6orf62 and HIST1 (chromatin assembly) genes converge via super-long-range interactions. These observations indicate that the hubs collectively perform essential cellular functions by conforming to a rich-club. Nonrandom spatial clustering of essential genomic loci might also relate to an evolutionary strategy to reduce expression noise by locating the essential loci to site(s) of abundant transcription, as discussed below.

**DISCUSSION**

In brief, we have demonstrated that chromatin interactions form a giant, interconnected network organized into three key interrelated structures: communities, hubs, and peripheral spokes. Communities are primarily centered on hyperacetylated, strong regulatory marks and organize the genome into distinct functional compartments. Hubs conform to an interconnected core, or rich-club, of key cellular functions, whereas spokes are relatively enriched in development-related and lineage-specific genes (Figure 7). This peculiar nonrandom functional organization of hubs and spokes might have evolved to keep the interactome healthy and robust against random deleterious genetic or transcriptional errors in the genome.

Functional enrichment of chromatin communities could be a potent constraint that ties together transcription-associated chromatin in the nucleus. Our observation strongly supports the notion of specialized transcription factories (Pombo et al., 2000; Xu and Cook, 2008), wherein promoters with common properties, such as binding sites for a particular transcription factor, share the same transcription factory. We reason that the enrichment of secondary functions in the same community might relate to (1) the limitation of available GO knowledge, (2) the transcription of genes that need to be expressed as a coordinated response to external stimuli, or (3) neutral coexpression clusters, which were previously proposed to be a result of neutral coevolution (Sémon and Duret, 2006). Interestingly, the chromatin communities are weakly interconnected with each other through trans or super-long-range cis interactions, which are generally transient in nature, suggesting that the trans chromatin interactions might be critical for cross-functional communication of genes to enable a coordinated response to external signals and allow the genome to easily reconfigure under environmental changes.

Evolutionary conservation of genomic neighborhood, sequence, and gene expression clearly supports the evolutionary constraints of transcription-associated chromatin proximity. This is also in line with a recent Hi-C study on human and mouse embryonic stem cells (Dixon et al., 2012), which appeared when this work was in communication. Interestingly, we observed a population of interacting loci that were distant in the mouse genome but proximal in the human genome, hinting at the possibility of evolution of gene clusters by means of long-range chromatin interactions. Loci that are to be transcribed in a cooperative manner may have been located at genomically
distant but spatially proximal sites in the ancestor genome, and may have translocated to proximal genomic sites in higher primates by a similar mechanism that mediates the genomic rearrangements in cancer genomes (Lin et al., 2009). Such hypotheses can be tested further when high-resolution RNAPII-associated chromatin interaction data become available for other mammalian and vertebrate species in the future.

The nonrandom enrichment of disease SNPs and germline mutations among the spokes hints at their selection against diverse chromatin interactions. This might be important for the fine-level regulation of development-related genes. Highly diverse interactions at these loci might increase their susceptibility to erroneous interactions and eventually to transcriptional dysregulation. Moreover, it was previously shown that chromatin interactions can mediate mutations (De and Michor, 2011; Lin et al., 2009). Therefore, the genomic loci with disease-associated mutations would survive through development only if their interactions were limited. The mutations that occur at the hub loci would be lethal and would not be observed in the population.

Nevertheless, the ChIN had hubs that were enriched in key cellular functions. How do these hubs escape random malfunctions? Based on our analysis, we reason that (1) the ChIN follows a scale-free-like distribution of node degrees, which means that the number of hubs would be very low in the network and hence the probability of an error hitting a hub would also be very low; and (2) hubs are not randomly distributed in the ChIN and instead are arranged nonrandomly into an interconnected core or rich-club, which further reduces the probability of being hit by random malfunctions.

A rich-club of key cellular functions implies two things: (1) In addition to the partial or complete loss of the known protein function, which may or may not explain the lethality, genetic or epigenetic errors in the top hub loci in the ChIN may have other consequences, such as alteration of transcription of other genes through promoter-promoter interactions (Li et al., 2012), followed by a cascading dysregulation of the downstream gene regulatory network, and eventually contributing to lethality. Because the top hubs are directly interconnected through promoter-promoter interactions, we hypothesize that it is this core, rather than a single gene, that gets transcriptionally dysregulated to cause lethality. (2) Nonrandom rich-club organization in the chromatin interactome may have evolved to shield the genes with key biological functions from random malfunctioning and also ensure their robust, high, and synchronized transcription through promoter-promoter interactions (Li et al., 2012; Figure S2G, degree correlation with gene expression).

Indeed, nonrandom linear genomic clustering of essential genes was previously proposed to be associated with lower expression noise (Batada and Hurst, 2007). Along similar lines, nonrandom three-dimensional (3D) clustering of essential cellular genes at nuclear sites of abundant transcription may regulate their lower expression noise. This clustering can be attributed to 3D organization of gene-dense regions (~70% of hubs were located in regions with >20 genes per Mb) in the nuclear core, which was previously shown to be evolutionarily conserved (Neusser et al., 2007; Tanabe et al., 2002). Therefore, an interconnected core of housekeeping genes might suggest a selection mechanism that evolved to reduce the variation in gene expression at essential gene loci associated with core cellular functions. In contrast, such variations in expression in the peripheral, nonhub nodes associated with lineage-specific and developmental functions may have been relatively tolerant in the context of cell survival. Nonetheless, certain type of genetic errors and dysregulated expression levels at these loci could be lethal in the context of organism survival.

Interestingly, the rich-club remains intact after the genomically rearranged regions are removed from the network, hinting at two possibilities: (1) either the loci in the rich-club are protected against DNA breaks, possibly via efficient DNA repair and protection mechanisms, or (2) the genomic rearrangements at these loci are deleterious in the cell-survival context and hence not observed in the cancer cell lines. The former possibility is also supported by the fact that these loci are hyperacetylated, which also allows for efficient DNA repair (Ikura et al., 2007), and the hub loci generally locate to early replicating domains (Figure S2N), which are less susceptible to genetic errors (Stamatoyannopoulos et al., 2009) due to lower accumulation of single-stranded DNA.

Taken together, our results obtained via a network approach uncover evolutionary and functional constraints, which might have shaped the 3D organization of the human genome. We propose that the human genome exhibits a robust systems organization of chromatin interactions to regulate transcription by compartmentalizing biological functions into distinct chromatin communities, and by ensuring the robust and consistent transcription of key essential genes in the interconnected dense core. The modular topology of the chromatin interactome may also guide GWAS studies to prioritize the SNPs for genotype-phenotype associations.

This work also suggests several future perspectives. First, by integrating the gene regulatory circuitry into the ChIN, investigators would be able to study and predict the erroneous waves a genetic or epigenetic lesion might radiate in a diseased genome. Second, for a full exploration of the emergent properties of chromatin interactome networks that arise over time, the dynamics of chromatin interactions during normal cell-lineage specification and evolution will need to be examined.

EXPERIMENTAL PROCEDURES

Data Sets

We used our previously published RNAPII ChIA-PET data sets (Li et al., 2012) to perform the comprehensive network analysis. These data sets are available from the Gene Expression Omnibus (GEO, GSE33664; Edgar et al., 2002) and from our in-house server. Other genomic data sets were taken from the resources listed in Table S1.

Network Construction

The ChIN was constructed using nonoverlapping distant genomic sites present in our RNAPII ChIA-PET libraries. The detailed strategy for network construction is elaborated in the Extended Experimental Procedures. Nodes were then demarcated as TSS, TES, GBD, and IGN, and as distinct chromatin types using genome annotations from the University of California Santa Cruz (UCSC) and ENCODE (Ernst et al., 2011).

Network Analysis

We used the igraph library on the R platform to analyze topological descriptors, such as the node degree (k), average degree of nearest neighbors (knn),
average path length, clustering coefficient, and various node or edge centralities of ChIN. To assign communities and their centralities, we used the ModuLand algorithm (Kovács et al., 2010). We detected the rich-club using the recently proposed rich-club coefficient (Colizza et al., 2006). We performed a network resilience analysis by progressively deleting random nodes from the ChIN and measuring the network destruction as a function of the average path length or the number of disconnected network components (Albert et al., 2000).

**GO Analysis**
We used network ontology analysis (NOA: Wang et al., 2011), BINGO (Maere et al., 2005), and PANTHER (Mi et al., 2010) to assess the enrichment of specific functions in chromatin communities.

**Visualization**
Networks were visualized on Cytoscape (Kohli et al., 2011) and Gephi. Spring-embedded layouts were used throughout the analysis. Chromatin loops and associated genomic features were browsed and analyzed on an advance genomic browser developed in-house (F.H.M. et al., unpublished data). Most of the plots were made on the R platform.

**Statistics**
Statistical tests of significance (i.e., Wilcoxon’s rank sum test, Fisher’s exact test, and binomial tests) were performed on the R platform. The FDR, when applicable, was calculated by randomizing the data sets several thousand times.

**SUPPLEMENTAL INFORMATION**
Supplemental Information includes Extended Experimental Procedures, seven figures, and six tables and can be found with this article online at [http://dx.doi.org/10.1016/j.celrep.2012.09.022](http://dx.doi.org/10.1016/j.celrep.2012.09.022).

**LICENSING INFORMATION**
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Supplemental Information

EXTENDED EXPERIMENTAL PROCEDURES

ChIA-PET Data Analysis
ChIA-PET reads were analyzed using the ChIA-PET tool (Li et al., 2010) and the strategy elaborated elsewhere (Li et al., 2012). Statistically significant chromatin interactions were identified using a random null model, which assumes that the probability of a ChIP enriched fragment to get ligated to another ChIP enriched region is random. Under this model, the probability of finding a particular cross-ligated product would follow a hypergeometric distribution. This probability was used to assign a p value to each chromatin interaction. FDR was calculated by adjusting p values using Benjamini-Hochberg correction method of multiple comparisons. The raw data are available from ChIA-PET and from GEO. The detailed properties of the data sets are elaborated elsewhere (Li et al., 2012). Numbers of uniquely mapped PETs are as follows:

| ChIA-PET library # unique PETs |
|-------------------------------|
| K562 replicate1 14,177,547     |
| K562 replicate2 14,365,592     |
| MCF7 replicate1 15,283,270     |
| MCF7 replicate2 15,622,720     |

Construction of the RNAP II-Associated ChIN
A large proportion (~50%) of interaction sites or anchors (average size = 500bp) was redundant, i.e., closely overlapping, in the original dataset (red peak in Figure S1A). We observed a clear separation of overlapping (redundant) and nonoverlapping interaction sites present in our ChIA-PET dataset (Figure S1A). We, therefore, merged the neighboring overlapping anchors and the maximum PET signal in the region was taken as the anchor-center. For further data-mining purpose, we extended each anchor +/- 2.5kb upstream and downstream from the anchor center to acquire nodes of 5kb size. Nodes were then connected as per their connectivity in the ChIA-PET dataset. Pairs of interacting nodes were imported to igraph library, available for R-package, to construct an undirected graph of chromatin interactions. The graph consisted of several small network components and one giant network, which covered up to 40% of total nodes present in the network. Though we used the complete graph for genomic, epigenomic and functional characterization of the chromatin interactome, only the giant network component was used to decipher topological properties of RNAPII associated chromatin interaction network (ChIN). Copy number variations in cancer genomes might contribute to bias in number of interactions of involved loci. To address this, we performed some control analyses as mentioned in the section 1.12.

Analysis of Network, Node, and Edge Descriptors
Several network descriptors like scale-freeness, network hierarchy, disassortativity were assessed using log-log plots of node descriptors. We performed Kolmogorov-Smirnov test in combination with maximum-likelihood for goodness-of-fit for power-law distribution \( f(k) = \frac{C}{k^\beta} \) of node degrees \( k \). The test was positive for non-TSS sites in ChIN, while it failed for TSS sites (Table S6). Several real world networks do not have a strict scale-free degree distribution but possess a scale-free like, heavy tailed degree distribution, which is sufficient for the explanation of our data as explained in detail elsewhere (Clauset et al., 2009). Therefore, we use the term “scale-free-like network” throughout this work. Network modularity was analyzed using ModuLand algorithm (Kovács et al., 2010). The existence of rich club was identified using the rich-club coefficient proposed by others (Colizza et al., 2006). Node descriptors namely degree, betweenness (number of shortest paths passing through a node), closeness (number of shortest paths required to reach any other node in the network), eigen vector centrality, transitivity (clustering coefficient), knn (average degree of nearest neighbors) etc were calculated using igraph functions on R platform. Community centrality scores were assigned using the ModuLand algorithm (Kovács et al., 2010). Edge betweenness was calculated using igraph to assign centralities to cis and trans chromatin interactions. Global topological properties of MCF7 and K562 ChIN were compared using log-log plots among the node descriptors. The two networks appear to be very similar topologically (Figure S1). Detailed properties of MCF7 and K562 ChIN nodes are given in Tables S2 and S3.

Detection of Rich-Club
We detected rich-club by calculating the following rich-club coefficient as proposed by Colizza et al. (2006):

\[
\sigma(k) = \frac{2E_{\geq k}}{N_{> k}(N_{> k} - 1)}
\]

where
- \( E_{\geq k} \) = number of observed edges at degree cut-off \( k \)
- \( N_{> k} \) = number of nodes at degree cut-off \( k \)
- \( N_{> k}(N_{> k} - 1) \) = number of expected edges at degree cut-off \( k \).
The coefficient, $c(k)$, was calculated for the original network and for $10^3$ randomly rewired networks. The average ratio of original versus random $c(k)/c_{ran}(k)$ was then plotted against degree cut-off. The 95% confidence interval was based on the distribution of $c_{ran}(k)$ values.

**Analysis of Genomic and Epigenomic Descriptors**

Genomic and epigenomic descriptors such as local SINE & LINE density, H3K4me3, H3K4me1, H3K9me3, H3K27me3, H3K9ac, H3K14ac, FAIRE, CTCF, replication timing, chromatin types (chromHMM) were taken from databases and published literature (Hansen et al., 2010; Joseph et al., 2010; Li et al., 2012; http://genome.ucsc.edu; http://www.genome.gov/10005107). Enrichment of each of the descriptors was calculated in 5kb bins across the whole genome. Two kinds of analyses were performed: 1) the correlation of node centrality (degree or community centrality) with enrichment of genomic/epigenomic marks to identify which mark associates best with number of interactions, 2) Pair-wise correlation analysis for interacting nodes to analyze the marks associated with chromatin interactions. Analysis of chromatin types (K562) is presented in Figures 2D and 2F, while plots for individual genomic and epigenomic marks are shown in Figure S2. Gene expression data sets used in the study are our in-house RNA-Seq and time course microarray data sets (Fullwood et al., 2009; Li et al., 2012). We analyzed MCF7 and/or K562 ChIN based on the availability of other epigenomic and functional data sets, including chromatin types, replication timing, and time-course microarray data.

**Expression Breadth Analysis**

RNAseq data for GM12878, HCT116, HeLa, K562 and MCF7 cell-line were downloaded from ENCODE. Expression breadth was calculated as the number of cell-lines a particular gene is expressed in (RPKM > 0). Expression breadth of 1 means that the gene is highly cell-line specific, while the value 5 signifies house-keeping gene.

**GO Analysis**

Most of the GO analyses shown in the article were performed using hypergeometric test available in BiNGO, a plugin for Cytoscape software. To confirm our observations by edge based ontology methods, we used Network Ontology Analysis (NOA). PANTHER was used occasionally to validate the enrichments. In each of methods, p values were corrected using standard Benjamini-Hochberg method.

**Analysis of Evolutionary Conservation**

Genomic blocks syntenic between human and chimp, mouse and zebra fish genome were downloaded from Cinteny database. The number of chromatin interactions per Mb were calculated for syntenic and nonsyntenic blocks. The interactions between syntenic and nonsyntenic blocks were negligible. These observations were also confirmed using data from ECRbase (not shown). Mouse genomic coordinates for corresponding interacting genomic sites in human genome, were lifted-over using the liftOver utility from UCSC browser. The FDRs were calculated using the approach given in the section 1.15. Conservation of genomic neighborhood (CGN) scores were taken from De et al. (2009). Sequence and expression divergence data were taken from Khaitovich et al. (2005). Human-mouse orthologs were downloaded from MGI.

**Analysis of Disease-Associated Genetic Mutations**

A list of disease-associated SNPs identified by genome-wide association studies (GWAS) was downloaded from GWAS catalog (Hindorff et al., 2009). SNPs were mapped to all types of nodes (TSS, TES, GBD, IGN and different chromatin types) of uniform size as mentioned in the previous section. FDR at each degree cut-off was calculated using $10^3$ randomizations of GWAS SNP locations in the ChIN. OMIM (Hamosh et al., 2000) data were used to analyze other disease-associated mutations. Cancer associated somatic and germline mutations were downloaded from the cancer census (Futreal et al., 2004). Only the TSS nodes were considered for genic mutations.

**Mouse Phenotype Analysis**

Mouse phenotype information was taken from MGI database. The phenotypes showing the terms “lethal,” “death” or “morality” were considered as lethal phenotypes. MGI showed a total of 10,692 genotypes with these terms out of total 46,178 mouse genotypes available.

**Network Resilience Analysis**

To assess whether the rich club of critical biological functions could indeed make the ChIN relatively more robust, we performed a network resilience analysis of ChIN. We gradually deleted the random nodes (mutation) from the ChIN with intact rich club and with disrupted rich-club. The expansion of average path length (average of all shortest path lengths in the network) and the increase in number of network components were taken as measures to calculate network disruption. From Figures S6B and S6C, it is clear that ChIN with intact rich club is relatively more tolerant to node deletions when compared to ChIN with disrupted (disconnected) rich-club suggesting that the rich club structure is indeed important for the robustness of ChIN topology against random errors.
Control Analyses for Genomic Rearrangements

Structural abnormalities of cancer genomes can alter the network and functional analyses and interpretations presented in this study. To scrutinize our observations, we performed some control analysis of genomic rearrangements in MCF7 and K562 genomes. We used our in-house G-PET data on several different kinds of structural variations (Hillmer et al., 2011). The plots shown in Figure S1C suggest that the copy numbers 1.25 to 4 are the most common in the MCF7 and K562 genomes. We, therefore, filtered out genomic regions having less than 1.25 or more than 4 copies in the MCF7/K562 genomes. We, then, removed the other abnormalities like translocation, inversion, insertion, deletions, which mapped proximal to chromatin interaction sites. We then re-plotted important topological descriptors, like degree, centrality and clustering coefficient as well the key observations of the study before and after filtering of structural abnormalities. We did not observe any significant bias which could influence the overall conclusions presented in the article (Figure S1D).

Control Analyses for Genomic Correlates

Genomic descriptors like SINE, LINE & gene densities and GC content strongly correlate with gene expressivity. In our earlier analyses, we showed that these descriptors also correlate with density of chromatin interactions. Therefore, to assess the extent these correlations account for the degree correlations shown in Figure S2E, we controlled our analyses using partial correlations by eliminating the effects of two such descriptors, namely SINE and LINE densities, which are anticorrelated with each other. The comparisons shown in Figure S2I, clearly suggest that these genomic variables do not account for the strong degree correlations observed in Figure S2E.

Data Visualization

Spring embedded layouts of ChIN were visualized using Cytoscape (Kohl et al., 2011) and Gephi software. Genomic tracks for genes, chromatin interactions and other epigenomic marks were visualized using an in-house genome browser. Most plots were made on R platform.

Statistics

Wilcoxon’s rank sum test was used to test the significance between distributions shown in the article. FDRs in Figure 5A were calculated using 10^4 randomizations of GWAS SNPs in the ChIN. FDR values in Figure 4B were calculated from 10^6 random data sets of paired coordinates of same genomic span distribution as the original interacting pairs. Following equation, which represents the upper bound of p value, was used to calculate the FDR:

\[ P_{upper} = \frac{b+1}{m} \]

Where \( b \) = number of values \( \geq \) observed value, and \( m \) = number of permutations. All statistical tests were performed on R.

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Figure S1. Detailed Topological Description of ChIN, Related to Figures 1 and 2

(A) Distribution of genomic distances between neighboring interaction sites (anchors) in the RNAII ChIA-PET dataset (MCF7).

(B) Global comparison of network descriptors of MCF7 (red) and K562 (blue) ChINs. k: degree; b: betweenness centrality (i.e., the number of shortest paths passing through a node); c: closeness centrality (i.e., the average number of shortest paths needed to reach from a node to all other nodes in the network); e: eigenvector centrality; t: transitivity or clustering coefficient (number of observed triangles/number of expected triangles) incident on a node; knn: average degree of nearest neighbors. Overall, the two networks had similar topological properties.

(C) Copy-number variations in MCF7 and K562 cell lines. Regions with high or low copy numbers were discarded for control analyses.

(D) Topological descriptors were compared before and after removal of genomic abnormalities (high/low copy number and genomic rearrangements revealed by G-PET analyses). The statistically significant difference was assessed via the Mann-Whitney U test. None of the descriptors showed a significant p value.

(E) Giant network components of ChINs constructed from MCF7 ChIA-PET (left, 5 kb resolution) and K562 HiC (right, 1 Mb resolution) datasets. The colors of the nodes depict distinct chromosomes.

(F) Scale-free-like degree distribution of ChINs constructed from MCF7/K562 ChIA-PET (left) and K562 HiC (right) data.

(G) Top panel: Distribution of edge centralities (edge betweenness) for cis (intra-chr) and trans (inter-chr) chromatin interactions in MCF7 and K562 cell lines. Bottom panel: Correlation plots between edge centralities and genomic span of chromatin interactions in MCF7 and K562 cell lines. Edge betweenness represents the number of shortest paths passing through a particular edge. The figure demonstrates the greater importance of superlong-range and trans chromatin interactions in maintaining the ChIN topology. Abolishment of these superlong-range and trans interactions would break the ChIN topology into smaller fractions.

(H) Left panel: Genomic span distribution of intra- and intercommunity interactions. Right Panel: Fraction of interchromosomal interactions in intra- and inter-community interactions.
Figure S2. Detailed Genomic and Epigenomic Characterization of ChIN Topology, Related to Figure 2

(A) Pie chart representing different types of nodes. (B) Boxplots for the distribution of degree ($k$) and the average degree of neighbors ($k_{nn}$) of distinct types of nodes. TSS, transcription start site; TES, transcription end site; GBD, gene BoDy; IGN, intergenic sites.

(D) Log-log plot of degree ($k$) and average degree of neighbors ($k_{nn}$). Inverse correlation suggests that ChIN is disassortative, i.e., highly connected nodes preferably interact with weakly connected nodes. Disassortativity reflects a functional segregation of TSS (black), which have more neighbors and are mostly connected to distal regulatory elements (non-TSS; gray) with relatively fewer neighbors. Note that the rich-club phenomenon, as reported in this article, and the assortative/disassortative mixing properties of a network are not trivially related or derived from each other, as elaborated elsewhere (Colizza et al., 2006).

(F) Correlations of degree with distinct chromatin modifications (per 5 kb). The strongest correlation of degree was seen with H3K9 acetylation. These observations were true for all types of nodes (All, TSS, TES, GBD, and IGN). Numbers on the bottom are the PCC values for the correlation between degree and enrichment of corresponding chromatin mark for all of the nodes in the network.

(F) Pairwise correlations between chromatin modifications at interacting sites. The X and Y axes are interacting partners.

(G) Correlation of degree with gene expression (RNA-Seq).

(H) Correlations of degree with SINE and LINE densities (per 50 Kb).

(I) Partial correlations of epigenomic marks with degree when controlled for SINE and LINE densities.

(J) Detailed distribution of community centrality scores among distinct chromatin types. Asterisks indicate the chromatin types for which community centralities were significantly greater when compared with all nodes. The Mann Whitney U test was used to calculate the p values (≤2.2e-16 for each asterisk).

(K) Distribution of the number of promoter interactions of nodes with strong and weak enhancer marks. The p value was calculated using the Mann Whitney U test.

(M) Degree distribution of all active promoters and strong enhancers, and those marked with chromatin-remodeling factors BRG1 or INI1. Active promoters and strong enhancers marked with BRG1/INI1 were significantly more interactive.

(N) Degree distribution for early- and late-replicating sites (per 5 kb).

(O) Correlation of degree with the expression breadth (i.e., the number of cell lines in which a gene is expressed) of genes.

(A)–(I) refer to MCF7, and (J)–(O) refer to K562 ChIN. Numbers shown on the scatter plots are PCCs.
Figure S3. In Situ Validations and Other Details about the Functional Organization of ChIN, Related to Figure 3
(A) Component size analysis for ChIN containing only TSS sites; ~600 nodes converge to a giant complex. (B) Mapping of chromatin communities using the ModuLand algorithm. A total of 1,783 communities spread across all network components were identified. Only components with at least 20 genes were considered for the GO analysis presented in Figure 3A. (C) Functional organization of the giant network component before (+G-PET) and after (−G-PET) the removal of genomically rearranged regions. Loss of links does not alter overall functional organization (related to Figure 3). (D and E) DNA FISH validations of interchromosomal interactions between functionally related genes. (D) A common enhancer interacts with PRSS22 (a brain-specific serine protease) in cis and with CASP2 (cysteine-aspartic acid protease, involved in neurodegenerative disorders) in trans. Interestingly, the enhancer locus appears to be primate specific. Both genes are expressed in the MCF7 cell line. Tracks for regulatory marks (FAIRE, H3K14ac, H3K9ac, and CTCF), and RNA-Seq, and conservation across primates, mammals, and vertebrates are shown. Primate-specific conservation of enhancer and brain-associated gene functions prompted us to test the underlying interactions by single-cell experiment. (E) Small nuclear RNAs on chr3 and chr10 converge via trans chromatin interaction. This interaction was selected for FISH validation to illustrate the physical association of noncoding gene loci. The p values were calculated via a binomial test. (F) Proportion of cell-line-specific (MCF7, for illustration) communities and the communities common (>75% overlap) to MCF7 and K562 cell lines. (G) Example of MCF7-specific (red) and K562-specific (blue) nodes embedded in housekeeping (common, yellow) chromatin communities.
Figure S4. Control Analyses for Evolutionary Constraints in ChIN, Related to Figure 4

(A) Synteny analysis on genomic sites of similar gene density and expression. Left and middle panels: Distribution plots for gene density and expression values for control syntenic and nonsyntenic blocks (human-mouse). Right panel: Density of K562 chromatin interactions in the syntenic and nonsyntenic regions.

(B) Density of MEF chromatin interactions in the mouse genomic blocks that are syntenic to the human genome.

(C) Synteny analysis before (+G-PET) and after (−G-PET) filtering the genomic rearrangements in the MCF7 cell line. The difference in interaction densities between the syntenic and nonsyntenic regions remains significant.

(D) Distribution of human-mouse orthologous genes in the K562 chromatin communities and random chromatin communities of the same number of nodes.

(E) Conservation of the genomic neighborhood (CGN).

(F) Asynonymous sequence divergence of genes with and without promoter-promoter interactions, in human and chimp genomes.

(G) Correlation of degree (k) with mammalian sequence conservation (46way phastCons elements) for distinct types of nodes.

(H) Left and middle panels: Nonrandom covariance and PCC between genomic spans of chromatin loops in the human genome (K562) and corresponding sites in the mouse genome. Right panel: Nonrandom enrichment of data points in the top-left quadrant (distant in the mouse genome and proximal in the human genome) of the plot in Figure 4B. The histograms are the distributions of values for random genomic spans of the same length as the original.
Figure S5. Detailed Characterization of Nodes with GWAS SNPs, Related to Figure 5

(A and B) Distribution of genic and intergenic sites in (A) all nodes and (B) GWAS SNP-containing nodes in MCF7 ChIN. There is no observable difference in the representation of genic and intergenic nodes in ChIN with and without GWAS SNPs.

(C) Distribution of GWAS SNP-containing nodes with normal and abnormal copy number in the MCF7 genome.

(D and E) Individual examples of long-range chromatin interactions between promoters (highlighted in gray) and intergenic nodes with GWAS SNP (highlighted in orange). The phenotype and function of the linked gene are noted at the bottom. Functional correspondence between the phenotype and the gene function is apparent.

(F) Degree distribution for distinct types of nodes with and without GWAS SNP. 1: active promoter; 2: weak promoter; 4–5: strong enhancer (class 1, 2); 6–7: weak enhancer (class 1, 2).

(G) Degree distribution for nodes mapping to nondisease genes (NDG), single disease genes (SDG), and multidisease genes (MDG) as obtained from OMIM.

(H) RNAPII enrichment on promoters of genes having somatic or germline cancer mutations. There was no remarkable difference in RNAPII enrichment between the two data sets.
Figure S6. Topological and Functional Properties of Rich-Club Genes, Related Figure 6

(A) Rich-club analysis after removal of genomically rearranged regions. The rich-club remains intact.

(B and C) Network resilience analysis of ChIN (K562).

(B) Fold change (from the initial value) of the average path length after gradual deletion of random nodes from ChIN when the intraconnectivity of the rich-club was preserved (blue) and when edges (#48) connecting the hubs were disconnected and randomly rewired to other non-rich-club nodes (orange).

(C) The same analysis as in (B), except that the total number of network components was considered as a measure of network destruction. The expansion of average path length and the increase in the number of network components represent the network destruction. The analyses consistently show slightly greater resilience of ChIN with intact rich-club. After ~1,500 nodes were deleted (highlighted in gray), the curves show a statistically significant difference.

(D) Functional enrichment among the rich-club genes versus the genes having disease SNPs at genic (promoter, terminator, exon, or intron) regions.
Figure S7. Topological and Functional Properties of Rich-Club Genes—Continuation of Figure S6, Related to Figures 6 and 7

(A) Functional enrichment among the rich-club genes versus the genes having disease SNPs at intergenic regions. For intergenic SNPs, target genes were determined by exploiting their chromatin interactions.

(B and C) Comparative analyses of evolutionary divergence in (B) sequence and (C) gene expression of spoke and rich-club genes.

**Figure S7.** Topological and Functional Properties of Rich-Club Genes—Continuation of Figure S6, Related to Figures 6 and 7

(A) Functional enrichment among the rich-club genes versus the genes having disease SNPs at intergenic regions. For intergenic SNPs, target genes were determined by exploiting their chromatin interactions.

(B and C) Comparative analyses of evolutionary divergence in (B) sequence and (C) gene expression of spoke and rich-club genes.