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Architectural Protein Subclasses Shape 3D Organization of Genomes during Lineage Commitment

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SUMMARY

Understanding the topological configurations of chromatin may reveal valuable insights into how the genome and epigenome act in concert to control cell fate during development. Here, we generate high-resolution architecture maps across seven genomic loci in embryonic stem cells and neural progenitor cells. We observe a hierarchy of 3D interactions that undergo marked reorganization at the submegabase scale during differentiation. Distinct combinations of CCCTC-binding factor (CTCF), Mediator, and cohesin show widespread enrichment in chromatin interactions at different length scales. CTCF/cohesin anchor long-range constitutive interactions that might form the topological basis for invariant subdomains. Conversely, Mediator/cohesin bridge short-range enhancer-promoter interactions within and between larger subdomains. Knockdown of Smc1 or Med12 in embryonic stem cells results in disruption of spatial architecture and downregulation of genes found in cohesin-mediated interactions. We conclude that cell-type-specific chromatin organization occurs at the submegabase scale and that architectural proteins shape the genome in hierarchical length scales.

INTRODUCTION

Genomes are organized at multiple length scales into sophisticated higher-order architectures (Misteli, 2007). Individual chromosomes occupy distinct spatial territories with respect to each other in interphase nuclei (Cremer and Cremer, 2001). Within each territory, at intermediate length scales of ~1–10 Mb, compartments of transcriptionally active euchromatin tend to group together, and independent from, compartments of inactive heterochromatin (Lieberman-Aiden et al., 2009). At the subcompartment level, chromatin is further organized into megabasesized topologically associating domains (TADs) that represent spatial neighborhoods of high-frequency chromatin interactions (Dixon et al., 2012; Hou et al., 2012; Nora et al., 2012; Sexton et al., 2012). Within TADs, however, the precise features of chromatin folding at the submegabase scale remain poorly understood.

Emerging evidence suggests that nuclear architecture is critically important for cellular function. Seminal microscopy studies have linked the spatial positioning of specific genomic loci to gene expression (Fraser and Bickmore, 2007; Kosak and Groudine, 2004; Lanctôt et al., 2007), replication (Gilbert et al., 2010), X chromosome inactivation (Erwin and Lee, 2008; Nora and Heard, 2010), DNA repair (Misteli and Soutoglou, 2009), and chromosome translocations (Roix et al., 2003). Moreover, molecular methods based on proximity ligation, such as chromosome conformation capture (3C) or circularized-3C (4C), have been used to detect functional long-range interactions between two specific genomic loci in a population of cells (Dekker et al., 2002; Simonis et al., 2006; Zhao et al., 2006). Principles from these studies have been difficult to generalize, however, because most previous reports focus on interrogation of 3D interactions between specific preselected fragments (Kurukuti et al., 2006; Noordermeer et al., 2011; Schoenfelder et al., 2010; Vakoc et al., 2005). More recently, technologies for genome-wide mapping of chromatin architecture have been described, but comprehensive detection comes at the expense of resolution for mammalian genomes (Hi-C) (Dixon et al., 2012; Lieberman-Aiden et al., 2009) or is restricted to only interactions mediated by a preselected protein of interest (ChIA-PET) (Handoko et al., 2011; Li et al., 2012). Thus, there is a great need to elucidate principles of genome folding at the submegabase



Figure 1. High-Resolution Mapping Reveals a Hierarchy of Architectural Subdomains within Larger Topological Domains (A–F) 5C and Hi-C interaction frequencies represented as normalized 2D heat maps. (A), (B), (D), and (E) Hi-C data (adapted from Dixon et al., 2012) displayed for (A) and (D) 10 Mb and (B) and (E) 1 Mb regions around (A) and (B) Sox2 and (D) and (E) Olig1-Olig2 for mouse E14 ES cells (top) and mouse cortex (bottom). TADs reported in (Dixon et al., 2012) are represented as tracks for domain calls (blue bars) and a directionality index (downstream bias, green; upstream bias, red). (C) (legend continued on next page)

scale by mapping 3D chromatin interactions in an unbiased manner at high resolution.

Megabase-scale TADs appear to be constant between mammalian cell types and conserved across species (Dixon et al., 2012; Nora et al., 2012). Therefore, we hypothesized that genome organization at the submegabase scale, e.g., within TADs, plays a critical role in the establishment and/or maintenance of cellular state. To test this hypothesis, we present an unbiased, large-scale, and high-resolution analysis of 3D chromatin architecture in a continuous developmental system. We employed chromosome conformation capture carbon copy (5C) (Dostie et al., 2006) in combination with highthroughput sequencing to map higher-order chromatin organization during differentiation of pluripotent mouse embryonic stem cells (ES) cells along the neuroectoderm lineage. An alternating 5C primer design was applied to guery long-range chromatin interactions in a massively parallel manner across six 1-2 Mb-sized genomic regions around key developmentally regulated genes (Oct4, Nanog, Nestin, Sox2, Klf4, and Olig1-Olig2). Our analyses reveal that distinct combinations of architectural proteins shape the 3D organization of mammalian genomes at different length scales for unique functional purposes during lineage commitment.

RESULTS

Generation of High-Resolution Chromatin Interaction Maps

To investigate cell-type-specific patterns of higher-order chromatin organization, we first derived homogeneous populations of multipotent neural progenitor cells (NPCs) from mouse ES cells using a well-established, four-stage procedure (Mikkelsen et al., 2007; Okabe et al., 1996). qRT-PCR analysis and confocal microscopy coupled with immunofluorescence staining confirmed a >95% pure population of Nestin/Sox2-postitive, Oct4/Nanog-negative cells displaying morphological characteristic of NPCs (Figure S1 available online).

We then employed 5C in combination with high-throughput sequencing to generate high-resolution long-range interaction maps for two biological replicates of ES cells and ES-derived NPCs (Dostie et al., 2006). 5C, a high-throughput derivative of 3C, involves the selective amplification of chromatin interactions within specific genomic loci of interest. By preselecting regions to be queried, we were able to obtain insight into chromatin architecture at the resolution of single interrogated fragments (~4 kb), which is not yet feasible in a cost-effective manner with genome-wide Hi-C technologies in mammalian systems. Forward and reverse 5C primers were designed in an alternating scheme using tools from the publicly available my5C suite (Lajoie et al., 2009) (Figure S2E). The tiled, alternating design queried ~90,000 cis and

 \sim 500,000 *trans* interactions in parallel across seven genomic loci surrounding developmentally regulated genes (*Oct4*, *Nanog*, *Sox2*, *Klf4*, *Nestin*, *Olig1-Olig2*, and gene-desert control) (Table S1).

We first evaluated the quality of our raw 5C data by (1) assessing consistency between biological replicates and (2) comparing our high-resolution 5C data to Hi-C data recently reported at 40 kb resolution for E14 ES cells and primary cells isolated from mouse cortex (Dixon et al., 2012). Raw 5C counts were highly correlated between replicates (ES 1 versus ES 2, Pearson correlation coefficient: 0.91; NPC 1 versus NPC 2, Pearson correlation coefficient: 0.89) and more correlated than between ES cells and NPCs (Pearson correlation coefficient: 0.70) when considering the log of counts for all fragment combinations with >100 reads. Furthermore, heatmaps of raw 5C data showed high similarity between biological replicates at each individual locus (Figure S2F), suggesting that regionspecific primers amplified each region in a robust and consistent manner. Importantly, we also observe a striking similarity between global topological features in 5C and Hi-C data for all regions queried, albeit with marked differences in resolution (Figure S3). These results indicate that our 5C libraries are high quality, consistent between replicates, and achieve notable similarity to data generated with an independent method (Hi-C) in an independent study with similar cellular phenotypes.

Unique Hierarchy of Topological Subdomains at Each Genomic Locus

We next examined large-scale architectural features by visualizing heatmaps of 5C counts in ES cells and NPCs (Figures 1C and 1F). Comparison of high-resolution 5C maps to Hi-C maps at each region revealed a complex hierarchy of chromatin organization. TADs are readily detected in both Hi-C and 5C data sets. Importantly, the higher resolution of 5C data revealed that TADs previously defined with Hi-C are further subdivided into smaller subtopologies (sub-TADs). We systematically identified sub-TADs in 5C data with a Hidden Markov Model-based approach (Extended Experimental Procedures). Using this method, we uncovered numerous distinct subtopologies arranged in a hierarchy within the larger TAD organization. Indeed, >60 invariant and cell-type-specific sub-TAD boundaries were identified with our 5C data at the submegabase scale, whereas only 7 TAD boundaries were called in our regions of interest with a previous Hi-C analysis (Dixon et al., 2012) (Figures 1G and 1H). Topological features were unique to each region (Figure S3), suggesting that each genomic locus has an architectural signature that may reflect the functional activity of that region. Taken together, these data demonstrate that 5C achieves a marked increase in resolution compared to Hi-C, which enables mapping of finer-scale architectural features within TADs.

and (F) 5C data displayed for 1 Mb regions around (C) Sox2 and (F) Olig1-Olig2 for mouse V6.5 ES cells (top) and ES-derived NPCs (bottom). Constitutive and celltype-specific subdomains called with a Hidden Markov Model (see Extended Experimental Procedures) are represented as black lines overlaid on 5C heatmaps and a directionality index displayed as a hierarchy of black wiggle tracks.

⁽G and H) Overlap between cell types for (G) TAD boundaries called from Hi-C data in (Dixon et al., 2012) and (H) subdomain boundaries called in the present work from 5C data.

See also Figures S1, S2 and S3.



Figure 2. Genome Architecture Undergoes Marked Reorganization at the Submegabase Scale upon Differentiation

(A) Scatterplot comparison of interaction scores between ES cells and NPCs. Thresholds for constitutive and cell-type-specific looping interactions are represented as colored boxes (brown, constitutive; red, ES specific; orange, NPC specific; gray, background).

(B) Scatterplot comparison of interaction scores after randomly permuting replicates as described in the Extended Experimental Procedures. (C) Interactions called significant in ES cells and NPCs.

(D–F) Chromatin interactions and epigenetic modifications at specific genomic loci in ES cells and NPCs. ChIP-seq reads are displayed for CTCF, Med12, Smc1, Sox2, Oct4, Nanog, H3K27Ac, H3K4me1, and H3K4me3 in ES cells (above gene track) and CTCF, Smc1, H3K27Ac, H3K4me1, and H3K4me3 in NPCs (below gene track). 3D interactions are represented as mirror image arcplots for ES cells (above gene track) and NPCs (below gene track), with constitutive and cell-type-specific interactions displayed in black and red, respectively. Black bars represent HindIII restriction fragments queried by the alternating 5C primer design scheme. Forward 5C primers, upper track. Reverse 5C primers, lower track. (D) ES-specific interactions between *Sox2* and a putative enhancer. (E) ES-specific interactions between *Oct4* and a putative enhancer. (F) Constitutive interactions around *Nanog and Slc2a3*. See also Figure S4.

Constitutive and Cell-Type-Specific Features of 3D Chromatin Organization

Our observation that submegabase-scale architectural features undergo marked changes between cell types prompted us to systematically identify constitutive and cell-type-specific looping interactions within and between larger-scale TADs. To account for bias intrinsic to all 3C-based methods, as well as to 5C specifically, we developed a probabilistic model that simultaneously captures the distance-dependent background level of nonspecific chromatin interactions and the nonbiological contribution from each primer (Imakaev et al., 2012; Yaffe and Tanay, 2011) (Extended Experimental Procedures). Our model produces an interaction score that is comparable within and between experiments and allows for robust detection of fragment-to-fragment looping interactions that are significant above the expected background signal (Figure S4).

3D contacts with interaction scores greater than stringent, pre-established thresholds in both biological replicates were subjected to further analysis (Figure 2A). To rigorously minimize false positives, thresholds were selected so that the large majority of cell-type-specific interactions were lost when randomly permuting data (Figure 2B). By applying these stringent thresholds, we identified 83 ES-cell-specific interactions that are lost upon differentiation, 260 constitutive interactions that are constant between cell types, and 165 NPC-specific interactions that are absent in ES cells and acquired upon differentiation

We next integrated 5C data with other epigenomic data sets. We observed that a significant proportion of fragments engaged in 3D interactions were occupied by specific histone modifications. For example, a series of ~80- to 120-kb-sized looping interactions connect the Sox2 gene with a putative downstream enhancer in ES cells marked by H3K4me1, H3K27ac, and low levels of H3K4me3 (Creyghton et al., 2010; Heintzman et al., 2009; Rada-Iglesias et al., 2011) (Figure 2D). Loss of enhancer marks in NPCs occured in parallel with loss of ES-specific looping interactions, suggesting that this particular chromatin conformation has important functional significance. Similarly, an ES-specific interaction connects the Pou5f1/Oct4 gene to a putative enhancer ~ 25 kb upstream marked by H3K4me1, H3K27ac, and low levels of H3K4me3 (Figure 2E). By contrast, we detected a hierarchy of constitutive interactions (constant between cell types) around the pluripotent genes Nanog and Slc2a3 despite changes in gene activity during differentiation (Figure 2F). These examples provide evidence that a notable proportion of looping interactions identified in this study may be involved in genome function.

Candidate Architectural Protein Subclasses

To gain more insight into organizing principles governing genome folding, we integrated 5C data with genome-wide maps of protein occupancy. We first examined factors that have been reported as both essential for cellular functions and correlated with a specific looping interactions using 3C technology. The top three candidates fulfilling these criteria were CCCTC-binding factor (CTCF), cohesin, and Mediator (Hadjur et al., 2009; Handoko et al., 2011; Kagey et al., 2010; Kurukuti et al., 2006; Splinter et al., 2006). Genome-wide binding sites for CTCF, mediator subunit Med12, and cohesin subunit Smc1 have been previously identified in ES cells by ChIP-seq (Kagey et al., 2010; Stadler et al., 2011). We first used the ChIP-seq data to quantify unique and overlapping occupied sites in our regions of interest. As previously reported, high-confidence Smc1 binding sites significantly overlapped high-confidence CTCF and Med12 binding sites (Kagey et al., 2010). However, in addition to Med12+Smc1 and CTCF+Smc1 co-occupied sites, we also found notable subclasses of CTCF alone and Med12 alone both genome wide and in our regions of interest in ES cells (Figures 3A and 3B). Noteworthy, Med12 rarely overlaps CTCF in the absence of cohesin, but a subclass with occupancy of all three proteins (i.e., Med12+Smc1+CTCF) does emerge as significant.

Architectural Proteins Organize the Genome at Different Length Scales

We next examined the enrichment of CTCF, Med12, and Smc1 in 5C looping interactions. Unsupervised cluster analysis demonstrated that >80% of significant interactions were anchored by some combination of CTCF, Med12, or Smc1 in ES cells, which is significantly higher than the enrichment of these proteins in all queried background interactions (Figure 3C). By contrast, only ~40% of interactions were occupied by some combination of Oct4, Nanog, and/or Sox2, which is not significant compared to the expected background enrichment of these proteins (Figure 3D). The widespread occupancy of CTCF, Med12, and Smc1 in 3D interactions led us to hypothesize that these three proteins might have important architectural roles in shaping 3D genome organization.

We next explored the specific role for each candidate architectural protein subclass in genome organization. We observed a striking pattern in which multiple adjacent binding sites for the same architectural protein subclass were often found at the base of significant interactions. Indeed, enrichment for a particular architectural subclass in 3D interactions showed a strong correlation with the number of binding sites (Figure 3E). Therefore, to explore only high-confidence interactions, we focused our analysis on only loops anchored by >2 or >3 co-occupied sites (Figure 3F). CTCF+Smc1 and CTCF alone subclasses were highly overrepresented at the base of constitutive interactions compared to background nonloops (Figure 3F). By contrast, Med12+Smc1 and Med12 alone subclasses were predominantly enriched in only ES-specific looping interactions. Intriguingly, sites co-occupied by Med12+CTCF+Smc1 showed enrichment in both constitutive and ES-cell-specific interactions.

We also noticed that interactions mediated by each candidate architectural subclass displayed markedly different size distributions (Figure 4A). Med12+Smc1 co-occupied sites were predominately enriched at the smallest <100 kb length scale (Figure 4B), whereas Med12 alone sites, independent from cohesin, were enriched at intermediate length scales of 600– 1,000 kb (Figure 4D). The subclass with all three proteins (Med12+CTCF+Smc1) also displayed a loop size distribution shifted toward small to intermediate (<300 kb) length scales (Figure 4C). By contrast, loops connected by CTCF+Smc1 and CTCF Alone subclasses were significantly biased toward interactions greater than 1 Mb in size (Figures 4E and 4F). Together, these results support our hypothesis that architectural protein subclasses function at different length scales to fulfill distinct roles in genome organization.

CTCF and Cohesin Anchor Constitutive Interactions

To further explore the molecular mechanisms regulating constitutive chromatin interactions, we mapped CTCF and Smc1 occupancy in NPCs using ChIP-seq. Genomic loci co-occupied by CTCF and Smc1 in both ES cells and NPCs represented the largest architectural subclass genome wide and in our regions of interest (n = 159) (Figures 5A and 5B). Moreover, the sites with constant occupancy of CTCF+Smc1 between cell types were highly enriched in constitutive interactions compared to background (Figure 5C). This result is illustrated with a series of loops around Nanog and SIc2a3 (Figure 2F) and Olig1 and Olig2 (Figure 5D). At both genomic loci, fragments anchoring the base of constitutive interactions contain CTCF+Smc1 cooccupied sites that remain constant between ES cells and NPCs. Thus, constitutive CTCF occupancy may be a critical mechanism regulating the establishment and/or maintenance of constitutive chromatin architecture.

We next used high-resolution 3D fluorescent in situ hybridization (FISH) to assess the importance of CTCF/cohesin in configuring chromatin architecture. Two 10 kb probes (Figure 5D)

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Figure 3. Architectural Protein Subclasses Have Distinct Roles in Genome Organization

(A) Heatmap representation of ChIP-seq signal for distinct architectural protein subclasses genome wide.

(B) Venn diagram comparing binding patterns for high-confidence ($p < 1 \times 10^{-8}$) CTCF, Med12, and Smc1 occupied sites in 5C regions.

(C and D) Unsupervised hierarchical clustering for significant interactions in ES cells enriched for (C) CTCF, Med12, and Smc1 or (D) Oct4, Nanog, and Sox2. (E) Fold enrichment of architectural protein subclasses in looping interactions versus the number of occupied sites per anchoring fragments.

(F) Fraction of constitutive or ES-specific looping interactions enriched with architectural protein occupied sites compared to the expected enrichment in background. Fisher's Exact test, * $p \le 0.05$.

corresponding to fragments anchoring the base of a predicted constitutive looping interaction around *Olig1-Olig2* produced virtually superimposable FISH signals in the majority of wild-type ES cells (Figures 5E and 5F). Having validated this looping interaction with an independent assay, we then directly tested the role for CTCF and cohesin by knocking down these proteins in V6.5 ES cells. CTCF and Smc1 mRNAs were markedly depleted to <20% of their wild-type expression levels after trans-

duction of ES cells with lentiviral shRNA constructs and subsequent puromycin selection (Figure S5). By contrast to observations in wild-type nuclei, FISH probes were no longer colocalized in CTCF- and Smc1-KD ES cells (Figures 5E and 5F). These data indicate disruption of chromatin organization and provide strong evidence that 3D contacts identified by 5C represent bona fide chromatin interactions. We conclude that both CTCF and Smc1 are essential for maintaining this particular



Figure 4. Architectural Protein Subclasses Function at Different Length Scales

(A) Size distributions of chromatin interactions anchored by distinct subclasses of architectural proteins.

(B–F) Histograms binned by loop size displaying fold enrichment of chromatin interactions connected by (B) Med12+Smc1 (navy), (C) Med12+CTCF+Smc1 (light blue), (D) Med12 alone (green), (E) CTCF+Smc1 (red), or (F) CTCF alone (orange) compared to background interactions depleted of the occupied sites in ES cells (gray). Fisher's Exact test, *p \leq 0.05.

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Figure 5. Constitutive Looping Interactions Are Anchored by Constitutive Binding of CTCF and Cohesin

(A) Heatmap representation of distinct subclasses of architectural protein occupancy between cell types genome wide.

(B) Venn diagram representing unique and overlapping high-confidence ($p < 1 \times 10^{-8}$) CTCF and Smc1 occupied sites in ES cells and ES-derived NPCs in 5C regions.

(C) Fraction of constitutive or cell-type-specific looping interactions enriched with constitutive occupancy of CTCF+Smc1 compared to the expected enrichment in background noninteractions. Fisher's Exact test, * $p \le 0.05$.

(D–F) DNA FISH analysis of chromatin interactions connected by sites constitutively bound by CTCF+Smc1. (D) Arcplot of constitutive interactions anchored by constitutive CTCF+Smc1 occupied sites (black) and cell-type-specific interactions anchored by ES-specific Smc1 occupied sites (red) compared to epigenetic marks around *Olig1* and *Olig2* genes. Shaded gray vertical bars highlight genomic fragments constitutively bound by dual CTCF+Smc1 sites anchoring the base of a series of constitutive looping interactions. Black bars represent HindIII restriction fragments queried by the alternating 5C primer design scheme. Upper track, fragments represented by forward 5C primers were windowed around adjacent fragments represented by reverse 5C primers. Lower track, fragments represented by reverse 5C primers were windowed around adjacent fragments represented by forward 5C primers. Lower track, fragments represented by forward 5C primers were windowed around adjacent fragments represented by forward 5C primers. Lower track, fragments represented by reverse 5C primers. Lower track, fragments represented by reverse 5C primers were windowed around adjacent fragments represented by forward 5C primers. Lower track, fragment A (green) and fragment B (red) were used to perform DNA FISH in wild-type V6.5 ES cells and ES cells treated with lentiviral shRNA for CTCF or Smc1. Scale bar, 1 μ m. (F) Quantification of spatial distances separating FISH probes (mean \pm SD). Wild-type V6.5 ES cells (0.144 \pm 0.05 μ m, n = 126), CTCF knockdown (0.421 \pm 0.21 μ m, n = 130), and Smc1 knockdown (0.385 \pm 0.13 μ m, n = 113).

See also Figures S5 and S6.

constitutive interaction and propose that similar mechanisms will apply to constitutive interactions genome wide.

Mediator and Cohesin Bridge Proximal Enhancer-Promoter Interactions

To understand the organizing principles regulating celltype-specific chromatin architecture, we first examined CTCF-independent Smc1 sites that were occupied only in ES cells and then lost upon differentiation (n = 123) (Figures 5B and 6A). Loss of ES-specific Smc1 in NPCs occurred in parallel with abrogation of ES-specific interactions (Figure 6B), supporting the idea that Smc1 can function in a CTCF-independent manner as an architectural protein essential for cell-type-specific chromatin interactions.

We then set out to identify cofactors that partner with cohesin to bridge cell-type-specific interactions. Genome-wide analysis of CTCF-independent, ES-specific Smc1 binding sites revealed a strong colocalization with Mediator and pluripotent transcription factors Oct4, Sox2, and Nanog (OSN) (Soufi et al., 2012) (Figure 6A). Indeed, >95% of all ES-specific Smc1 binding sites colocalize with Med12, whereas only ~55% of these sites colocalize with OSN in our regions of interest (Figure 6C). Importantly, ES-specific Smc1 binding sites were enriched in ES-specific interactions in both cases where these sites colocalize with OSN and also in cases where these sites did not colocalize with OSN (Figure 6D). These observations suggest that cohesin does not require OSN transcription factors to serve an architectural role in the establishment and/or maintenance ES-specific chromatin interactions.

We further investigated the mechanistic link between pluripotent transcription factors and chromatin architecture by parsing OSN subclasses genome wide and in our regions of interest (Figures 6E and S6D). We noticed a partial overlap between OSN occupied sites genome wide and architectural proteins in ES cells. Indeed, in our regions of interest, ~50% of OSN-binding sites colocalized with Med12+Smc1, whereas ~25% did not colocalize with any architectural protein subclass (Figure 6F). Importantly, OSN occupied sites were only enriched in ES-specific looping interactions in cases where these proteins colocalized with architectural proteins (Figure 6G). Together, these data suggest that OSN transcription factors do not have a specific role in chromatin organization independent from architectural proteins.

To validate the roles for Mediator and cohesin in ES-cell-specific looping interactions, we carried out high-resolution 3D-FISH in wild-type and Med12- or Smc1-knockdown V6.5 ES cells. For this analysis, we chose an interaction between *Olig1* and a putative downstream ES-cell-specific enhancer (Figures S6B and S6C). Probes generated from these interacting regions (Figure S6C) colocalized in WT ES nuclei, but not in Med12- or Smc1-KD cells (Figures 6H and 6I). We conclude that Mediator and cohesin are essential for formation of an ES-cell-specific loop at *Olig1* and propose that similar mechanisms will apply to other ES-specific chromatin interactions.

Cohesin-Mediated Interactions Are Functionally Linked to Gene Expression

To further test the hypothesis that Med12+Smc1-mediated interactions have functional significance during lineage commitment, we examined the expression of genes anchoring loops connected by these proteins. Analysis of microarray data generated in ES cells and NPCs (Creyghton et al., 2010) demonstrated that ES-specific, Smc1-mediated interactions are biased toward connecting genes that are highly expressed in ES cells and turned off in NPCs (Figure 6J). Gene ontology analysis confirmed an overrepresentation of developmentally regulated pluripotent genes (e.g., *Pou5f1/Oct4, Sox2,* and *Notch4*) in ES-specific interactions connected by ES-specific Smc1 compared to all genes in ES-specific interactions (Figure S6E). By contrast, the expression distribution of genes colocalized with CTCF+Smc1 in constitutive looping interactions was not significantly different from the expression distribution of all genes in constitutive looping interactions (Figure 6K).

We then examined the effect of knocking down Med12 and Smc1 on gene expression. After siRNA knockdown of either Smc1 or Med12 in ES cells (Kagey et al., 2010), expression of genes anchoring the base of cohesin-mediated interactions was markedly reduced compared to expression of all genes found in ES-specific interactions (Figure 6L). Noteworthy, the reduction in gene expression after siRNA treatment was more severe for cohesin-colocalized genes anchoring the base of looping interactions versus nonlooping background interactions. These results expand upon previous reports at specific genomic loci (Kagey et al., 2010) by suggesting that the architectural roles for Mediator and cohesin might be a widespread mechanism linking gene expression and chromatin organization genome wide.

Overall, data are consistent with a model in which Mediator/ cohesin connect ES-cell-specific looping interactions between proximal regulatory elements and promoters of developmentally regulated pluripotent genes. This idea is illustrated at the Sox2 locus, where a series of ~80- to 120-kb-sized looping interactions connect the Sox2 TSS to a putative active enhancer (Figure 2E). OSN transcription factors and the Med12+Smc1 architectural subclass colocalize at the fragments anchoring these loops. Loss of architectural protein binding and ES-specific looping interactions in NPCs occurs in parallel with loss of pluripotent gene expression, suggesting that chromatin structure and function are intricately linked.

Architectural Proteins Facilitate Looping of Cell-Type-Specific Enhancers

The involvement of Mediator and cohesin in relatively shortrange enhancer-promoter interactions prompted us to look more broadly at the relationship between architectural proteins and distal-cell-type-specific regulatory elements. Putative ESspecific enhancers were parsed as genomic loci at least 2 kb distal from TSSs with high-confidence signal for H3K4me1 and H3K27ac in ES cells and loss of these chromatin marks in NPCs (Figure 7A). Similarly, putative NPC-specific enhancers were parsed as genomic loci at least 2 kb distal from TSSs that do not display H3K27ac signal in ES cells, but acquire high-confidence signal for H3K4me1 and H3K27ac in NPCs (Figure 7F). We noticed that putative ES-specific and NPC-specific enhancers could be sorted from high to low intensity of H3K4me3 signal (Figures 7A and 7F). A strong correlation was observed between ES-specific enhancers displaying H3K4me3

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signal and RNA Pol II occupancy, suggesting that enhancers marked by H3K4me3 might have active transcription of eRNAs (Kim et al., 2010) (Figure 7A).

To further explore the link between enhancers and chromatin architecture, we parsed ES-specific and NPC-specific enhancers in our regions of interest into high, intermediate, and low levels of H3K4me3 (Figures 7B and 7G). Noteworthy, only ES-specific enhancers with high levels of H3K4me3 were enriched in ES-specific looping interactions, whereas ES-specific enhancers with low levels of H3K4me3 were enriched only in constitutive interactions compared to background (Figure 7C). Similarly, NPC-specific enhancers with high and intermediate levels of H3K4me3 were enriched in NPC-specific looping interactions, whereas NPC-specific enhancers with low levels of H3K4me3 were not enriched in chromatin interactions compared to background (Figure 7H). Altogether these observations provide support for the idea that eRNA transcription correlates with enhancer activity and subsequent activity-dependent looping of cell-type-specific enhancers (Sanyal et al., 2012).

As a final step, we queried the potential colocalization of enhancers with architectural proteins. We noticed that the majority of ES-specific enhancers (~95%) colocalized with architectural proteins in ES cells (Figure 7D). Consistent with our previous analyses, ES-specific enhancers that colocalized with Smc1 alone were enriched in ES-specific looping interactions, whereas ESspecific enhancers colocalized with CTCF+Smc1 were enriched predominantly in constitutive interactions (Figure 7E). Intriguingly, only $\sim 25\%$ of NPC-specific enhancers colocalized with architectural proteins (Figure 7I). NPC-specific enhancers that colocalized with CTCF+Smc1 were enriched in NPC-specific looping interactions, whereas NPC-specific enhancers that did not colocalize with architectural proteins were not enriched in looping interactions (Figure 7J). Because the CTCF+Smc1 architectural subclass is markedly enriched in constitutive interactions and also displays a slight enrichment in NPC-specific interactions (Figure 5C), we hypothesize that one functional purpose for constitutive subdomains is to premark specific locations in

the genome that will acquire or lose enhancer activity during development.

An example of architectural proteins cooperating with enhancers is shown at the Sox2 locus, where the Sox2 gene is highly expressed in both ES cells and NPCs (Figure 7K). Data indicate that this developmentally regulated gene is controlled by different regulatory elements even though expression levels remain high as cells switch fate. In ES cells, the Sox2 TSS is connected to a proximal enhancer through a series of ~100kb-sized interactions, whereas in NPCs these smaller looping interactions break apart, and a larger (~450 kb) subdomain is present between the Sox2 TSS and a more distal NPC enhancer. Mediator/cohesin mark the proximal ES-specific enhancer, whereas CTCF/cohesin premark the sub-TAD boundary in ES cells that ultimately acquires a distal NPC-specific enhancer upon differentiation. These results support a prevalent role for architectural proteins in spatially connecting proximal and distal enhancer elements to the genes that they regulate. Although enhancers and insulators are traditionally thought to serve distinct mechanistic functions in gene regulation, our data suggest that enhancers and architectural proteins may work in collaboration to organize the genome much more than previously realized.

DISCUSSION

Analyses presented here provide an important step toward understanding the link between higher-order chromatin architecture, epigenetic modifications, and cell-type-specific gene expression. By analyzing the genome in 3D, we now discover that three proteins thought to play more traditional roles in transcriptional activation and insulation might belong to a class of architectural proteins with primarily "chromatin organizing" function. It was originally suggested that vertebrate CTCF is an insulator protein based on transgene studies demonstrating that this protein blocks communication between adjacent regulatory elements in a position-dependent manner. However, data presented here are more consistent with recent reports

Figure 6. Mediator and Cohesin Bridge ES-Specific Enhancer-Promoter Interactions

(A) Heatmap representation of all ES-specific Smc1 occupied sites compared to Med12, Oct4, Sox2, and Nanog occupied sites genome wide.

(B) Fraction of constitutive or cell-type-specific looping interactions enriched with ES-specific Smc1 occupied sites compared to the expected background

enrichment. Fisher's Exact test, *p \leq 0.05.

(C) Fraction of ES-specific Smc1 occupied sites colocalized with Med12 or Oct4/Sox2/Nanog in 5C regions.

(D) Fraction of constitutive or cell-type-specific looping interactions enriched with ES-specific Smc1 occupied sites with or without Oct4/Sox2/Nanog compared to the expected enrichment in background. Fisher's Exact test, * $p \le 0.05$.

(E) Heatmap representation of all Oct4/Sox2/Nanog subclasses compared to architectural proteins sorted by Med12 occupancy genome wide.

(F) Pie chart showing percentages of Oct4/Sox2/Nanog occupied sites colocalized with architectural proteins in 5C regions (n = 102).

(G) Fraction of constitutive or cell-type-specific looping interactions enriched with Oct/Sox2/Nanog with or without architectural proteins compared to the expected enrichment in background. Fisher's Exact test, $*p \le 0.05$.

(H) Probes specific for fragment A (green) and fragment B (red) anchoring an ES-specific looping interaction connected by an ES-specific cohesin site were used to perform DNA FISH in wild-type V6.5 ES cells and ES cells treated with shRNA for Med12 or Smc1. Scale bar, 1 μ m.

(I) Quantification of spatial distances separating FISH probes (mean \pm SD). Wild-type V6.5 ES cells (0.139 \pm 0.04 μ m, n = 114), Med12 knockdown (0.390 \pm 0.16 μ m, n = 123), and Smc1 knockdown (0.462 \pm 0.21 μ m, n = 123).

(J and K) Gene expression ratio between ES cells and NPCs for (J) genes in ES-specific interactions colocalized with ES-specific Smc1 compared to all genes in ES-specific interactions or (K) genes in constitutive interactions colocalized with constitutive CTCF+cohesin compared to all genes in constitutive interactions. Kolmogorov-Smirnov test: *, $p \leq 0.05$.

(L) Gene expression ratio between siRNA treatment for Med12 or Smc1 and wild-type ES cells for genes in ES-specific interactions colocalized with ES-specific Smc1 compared to either all genes in ES-specific interactions or all genes colocalized with Smc1 in background noninteractions. Kolmogorov-Smirnov test, * $p \le 0.05$.

See also Figures S5 and S6.

Cell



Figure 7. Architectural Proteins Cooperate with Cell-Type-Specific Enhancers to Form Cell-Type-Specific Interactions

(A) Heatmap representation of chromatin modifications demarcating putative ES-specific enhancers genome wide.

(B) Fraction of ES-specific enhancers colocalized with high, intermediate, or low levels of H3K4me3 in 5C regions.

(C) Fraction of constitutive or cell-type-specific looping interactions enriched with ES-specific enhancers with high, intermediate, or low levels of H3K4me3 compared to the expected background enrichment. Fisher's Exact test, * $p \le 0.05$.

(D) Fraction of ES-specific enhancers colocalized with architectural proteins in 5C regions.

(E) Fraction of constitutive or cell-type-specific looping interactions enriched with ES-specific enhancers colocalized with Smc1 Alone or CTCF+Smc1 compared to the expected enrichment in background. Fisher's Exact test, * $p \le 0.05$.

(F) Heatmap representation of chromatin modifications demarcating putative NPC-specific enhancers genome wide.

(G) Fraction of NPC-specific enhancers colocalized with high, intermediate, or low levels of H3K4me3 in 5C regions.

(H) Fraction of constitutive or cell-type-specific looping interactions enriched with NPC-specific enhancers colocalized with high, intermediate, or low levels of H3K4me3 compared to the expected background enrichment. Fisher's Exact test, * $p \le 0.05$.

(I) Fraction of NPC-specific enhancers colocalized with architectural proteins in 5C regions.

suggesting that enhancer blocking or barrier insulation may only occur in rare, context-dependent cases as a consequence of CTCF's primary role in connecting long-range interactions (Handoko et al., 2011; Phillips and Corces, 2009; Sanyal et al., 2012). Although traditionally considered an adaptor protein with multiple subunits essential for transcriptional activation (Kornberg, 2005; Malik and Roeder, 2000), we suggest that the widespread enrichment in 3D interactions predicts a similar architectural role for Mediator. In the case of cohesin, a protein with a well-known architectural function during mitosis, recent reports have suggested a critical mechanistic role in insulation based on the

finding that CTCF and cohesin colocalize across the genome at thousands of loci (Parelho et al., 2008; Wendt et al., 2008). Our data predict that cohesin functionally affects gene expression through its architectural role during interphase instead of through classic insulator mechanisms (Hadjur et al., 2009).

The molecular mechanisms governing chromatin folding at the submegabase scale remain critical unanswered questions in nuclear biology. CTCF is present at most boundaries between conserved megabase-sized TADs (Dixon et al., 2012; Nora et al., 2012). However, this protein cannot be considered the sole determinant of topological organization because the majority of CTCF sites are found within TADs. Here, we expand our understanding of chromosome organization at the submegabase scale by combining high-resolution 5C data and a detailed probabilistic model to computationally resolve individual fragmentto-fragment looping interactions within TADs. Our analyses reveal that larger, invariant TADs are hierarchically organized into constitutive and cell-type-specific subtopologies. By integrating 5C architecture maps with genome-wide maps of epigenetic modifications, we observed that a large proportion of subdomains coincide with specific looping interactions between architectural proteins and other regulatory sequences. Thus, we favor the idea that architectural protein-binding sites found within larger TADs could be responsible for connecting interactions that form the topological basis for subdomains. For example, the megabase-sized region around the Olig1 and Olig2 genes spans a boundary between two larger megabasesized TADs. Interactions identified in the present work with 5C reveal a further nested hierarchy of constitutive interactions anchored by CTCF+Smc1 and ES-specific interactions within and between subdomains connected by Med12+Smc1 (Figure S7).

We suggest a refined model for genome organization in which architectural protein subclasses function at different length scales to fulfill distinct roles in genome organization (Figure 7L). Data presented here predict that megabase-sized TADs are constant throughout development and are demarcated by constitutive occupancy of CTCF/cohesin at their boundaries. Within TADs, at intermediate length scales of 100 kb–1 Mb, CTCF/ cohesin co-occupied sites create subdomains by anchoring constitutive interactions around developmentally regulated or repressed tissue-specific genes. One possible functional role for CTCF/cohesin-mediated interactions at this intermediate length scale is to cooperate with distal regulatory elements to connect long-range enhancer-promoter interactions. Finally, at the smallest length scale (<100 kb), Mediator and cohesin cooperate to bridge ES-specific interactions between enhancers and core promoters of developmentally regulated genes. We also note that a small but significant proportion of CTCF/Mediator/ cohesin binding sites are not involved in chromatin interactions. It is possible that these sites are involved in interactions outside of the 5C regions queried in this study. Moreover, it is also possible that architectural proteins require additional layers of regulation that we do not yet understand, such as posttranslational modifications or additional binding partners that regulate protein-mediated chromatin organization in a spatiotemporal manner (Phillips and Corces, 2009).

Overall, this work provides insights into the organizing principles governing higher-order chromatin architecture and is significant toward understanding how the genome and the epigenome act in concert to regulate the formation of a diverse array of cell types during development. Powerful insights into the link between the hierarchical organization of 3D genomes and cellular function could be realized by future studies combining high-resolution analyses of chromatin architecture with genetic experiments in developmentally relevant model systems.

EXPERIMENTAL PROCEDURES

Extended Experimental Procedures are included in the Supplemental Information.

ES Cell Expansion and Differentiation

Murine V6.5 ES cells were expanded on Mitomycin-C-inactivated embryonic fibroblasts under standard pluripotent conditions. After initial expansion, ES cells were passaged 1–2 times on tissue culture plates coated with 0.1% gelatin to remove contaminating feeder cells. V6.5 ES cells were differentiated into NPCs using established procedures detailed in the Extended Experimental Procedures (Mikkelsen et al., 2007; Okabe et al., 1996).

Generation and Analysis of 5C Libraries

3C templates and 5C libraries were generated for ES cells and ES-derived NPCs according to standard procedures with some modifications (Dekker et al., 2002; van Berkum and Dekker, 2009). Regions queried with 5C primers are summarized in Table S1. All libraries were prepared for Illumina sequencing as described in the Extended Experimental Procedures. Reads were aligned to a pseudogenome consisting of all 5C primers (Tables S2, and S3) using Bowtie version 0.12.2 (Langmead et al., 2009). To account for poor quality reads, sequences were required to have only one unique alignment and 5 and 3 bases were trimmed from the 5' and 3' ends of the read, respectively. A summary of sequencing details for each biological replicate is provided in Table S5. Interactions were counted when both paired end reads could be uniquely mapped to the 5C primer pseudogenome. Only interactions between forward-reverse primer pairs were tallied as a true count. Primers showing counts >100,000 or <100 total reads were deemed outliers and removed from subsequent an alyses (Table S4).

⁽J) Fraction of constitutive or cell-type-specific looping interactions enriched with NPC-specific enhancers colocalized with CTCF+Smc1 or without architectural proteins compared to the expected enrichment in background. Fisher's Exact test, *p \leq 0.05.

⁽K) 5C interaction frequencies and epigenetic modifications at the Sox2 locus in ES cells (above gene track) and NPCs (below gene track).

⁽L) Architectural length-scale model for developmentally regulated chromatin organization.

5C data were corrected and analyzed with a probabilistic model detailed in the Extended Experimental Procedures. TADs were systematically identified with a Hidden Markov Model (HMM)-based approach also detailed in the Extended Experimental Procedures. Algorithms for the 5C peak-calling pipeline and HMM can be found at: https://bitbucket.org/bxlab/ phillips-cremins_cell_2013.

Generation and Analysis of ChIP-Seq Libraries

ChIP was performed on NPC pellets as previously described with minor modifications (Kagey et al., 2010). ChIP-seq libraries for Smc1a and CTCF were generated using Bethyl Laboratories (A300-055A) and Upstate (07-729) polyclonal antibodies, respectively.

A summary of ChIP-seq libraries analyzed in this study is provided in Table S6. All ChIP-seq data sets were aligned to build version MM9 of the mouse genome using default parameters (-v1 -m1) in Bowtie version 0.12.2 (Lang-mead et al., 2009). Only sequences that mapped uniquely to the genome were used for further analysis. The model-based Analysis for ChIP-sequencing peak finding algorithm (MACS, version 1.4.1) was used to identify regions of ChIP-seq enrichment over background (Zhang et al., 2008). Peak-called data were analyzed as detailed in the Extended Experimental Procedures.

Lentiviral Transductions

Lentiviral shRNA plasmids cloned into the pLKO.1 vector were purchased from Open Biosystems (Thermo Scientific). Specific clones were screened for robust knockdown of CTCF, Med12, and Smc1 (Figure S5). Lentiviral particles were produced using the TransLenti Viral Packaging Mix (TLP4615, Open Biosystems) and the Arrest-In Transfection Reagent in H293T cells as described in the kit manual. Murine V6.5 ES cells were transduced with lentiviral particles and 6 μ g/ml hexadimethrine bromide (polybrene, Sigma) as detailed in the Extended Experimental Procedures. To select for stable integration of shRNA constructs, cells were treated with 2–3.5 μ g/ml puromycin in ES cell media for 3–4 days starting at 48 hr postinfection.

DNA FISH

Three-color DNA FISH was carried out according to procedures described previously (Guo et al., 2011) in wild-type V6.5 ES cells, ES cells after lentiviral shRNA for CTCF, ES cells after lentiviral shRNA for Med12, and ES cells after lentiviral shRNA for Smc1. Position-specific 10 kb probes were amplified by long-range PCR using BAC templates with primers listed in Table S7. Ten kilobase FISH probes for fragment A and fragment B anchoring the base of a specific looping interaction were labeled with Alexa Fluor 488 (green) and 594 (red), respectively. BACs were used as anchors and labeled with Alexa Fluor 697 (blue). All probes and BACs were hybridized to ES cell slides prepped as described in the Extended Experimental Procedures. Distances between red and green probes were visualized and measured according to procedures also detailed in the Extended Experimental Procedures.

ACCESSION NUMBERS

The Data have been deposited in GEO under accession number GSE36203.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Extended Experimental Procedures, seven figures, and seven tables and can be found with this article online at http://dx. doi.org/10.1016/j.cell.2013.04.053.

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