

Combinatorial Patterning of Chromatin Regulators Uncovered by Genome-wide Location Analysis in Human Cells

Oren Ram,^{1,2,3,4,7} Alon Goren,^{1,2,3,4,7} Ido Amit,^{1,2} Noam Shores,¹ Nir Yosef,^{1,2} Jason Ernst,^{1,5} Manolis Kellis,^{1,5} Melissa Gymrek,^{1,2,3,4} Robbyn Issner,¹ Michael Coyne,¹ Timothy Durham,¹ Xiaolan Zhang,¹ Julie Donaghey,¹ Charles B. Epstein,¹ Aviv Regev,^{1,2,6,*} and Bradley E. Bernstein^{1,2,3,4,*}

¹Broad Institute of MIT and Harvard, Cambridge, MA 02142, USA

²Howard Hughes Medical Institute, Chevy Chase, MD 20815, USA

³Department of Pathology, Massachusetts General Hospital and Harvard Medical School, Boston, MA 02114, USA

⁴Center for Systems Biology and Center for Cancer Research, Massachusetts General Hospital, Boston, MA 02114, USA

⁵MIT Computer Science and Artificial Intelligence Laboratory, Cambridge, MA 02139, USA

⁶Department of Biology, Massachusetts Institute of Technology, Cambridge MA 02140, USA

⁷These authors contributed equally to this work

*Correspondence: aregev@broad.mit.edu (A.R.), bernstein.bradley@mgh.harvard.edu (B.E.B.)

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SUMMARY

Hundreds of chromatin regulators (CRs) control chromatin structure and function by catalyzing and binding histone modifications, yet the rules governing these key processes remain obscure. Here, we present a systematic approach to infer CR function. We developed ChIP-string, a meso-scale assay that combines chromatin immunoprecipitation with a signature readout of 487 representative loci. We applied ChIP-string to screen 145 antibodies, thereby identifying effective reagents, which we used to map the genome-wide binding of 29 CRs in two cell types. We found that specific combinations of CRs colocalize in characteristic patterns at distinct chromatin environments, at genes of coherent functions, and at distal regulatory elements. When comparing between cell types, CRs redistribute to different loci but maintain their modular and combinatorial associations. Our work provides a multiplex method that substantially enhances the ability to monitor CR binding, presents a large resource of CR maps, and reveals common principles for combinatorial CR function.

INTRODUCTION

Gene regulation in eukaryotes relies on the functional packaging of DNA into chromatin, a higher-order structure composed of DNA, RNA, histones, and associated proteins. Chromatin structure and function are regulated by posttranslational modifications of the histones, including acetylation, methylation, and ubiquitylation (Kouzarides, 2007; Margueron and Reinberg, 2010; Ruthenburg et al., 2007).

Advances in genomic technologies—in particular, chromatin immunoprecipitation (ChIP) followed by sequencing

(ChIP-seq)—have enabled researchers to characterize chromatin structure genome-wide in different mammalian cells (Barski et al., 2007; Birney et al., 2007; Heintzman et al., 2007; Mikelsen et al., 2007; Zhang and Pugh, 2011; Zhou et al., 2011). The resulting maps have shown that distinct histone modifications often exist in well-defined combinations, corresponding to different genomic features (e.g., promoters, enhancers, gene bodies) or regulatory states (e.g., actively transcribed, silenced, poised). The number of chromatin types may, in fact, be relatively limited (Ernst and Kellis, 2010; Filion et al., 2010). For example, a study of chromatin landscapes across 9 different human cell types distinguished 15 dominant chromatin types, or “states,” based on their combinatorial histone modifications (Ernst et al., 2011). The chromatin state of each locus varies between cell types, reflecting lineage-specific gene expression, developmental programs, or disease processes.

It is compelling to hypothesize that combinatorial histone modification states are determined by different combinations of chromatin regulators (CRs). The human genome encodes hundreds of CRs that add (“write”), remove (“erase”), or bind (“read”) these modifications (Kouzarides, 2007; Ruthenburg et al., 2007). CRs are expressed in a tissue-specific manner and play important roles in normal physiology and disease (Ho and Crabtree, 2010). For example, cancer genome projects have unveiled prevalent mutations in CR genes, suggesting broad roles for these proteins in tumor biology (Elsässer et al., 2011). Despite their importance, the target loci and specific functions of most mammalian CRs remain unknown. In contrast to histone modifications that are readily mapped by ChIP-seq, systematic localization of CRs has proven challenging. Though recent studies in yeast (Venters et al., 2011) and fly (Filion et al., 2010) have profiled multiple CRs, few have been mapped in mammalian cells. Furthermore, the available profiles typically have lower signal-to-noise ratios than maps of histone modifications or transcription factors. This is likely due to the indirect associations between CRs and DNA, compounded by suboptimal antibody reagents and ChIP procedures. This severely

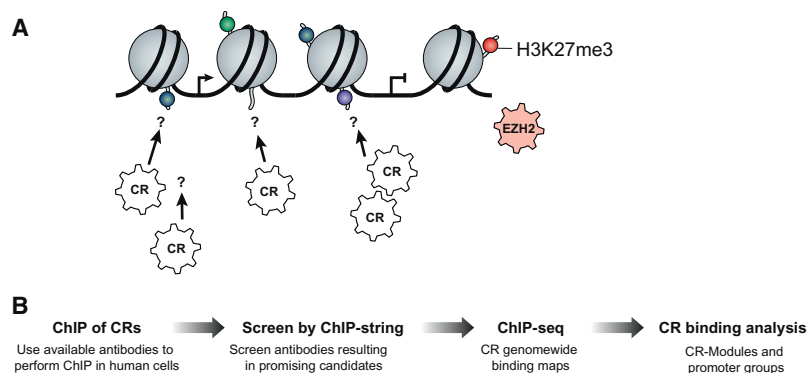


Figure 1. A Systematic Approach for Associating CRs to Genomic Loci and Chromatin States

(A) CR associations with chromatin modification states. (Right) The chromatin regulator EZH2 (pink gear) is associated (arrow) with a locus marked by the histone modification H3K27me3 (small red ball). (Left) For many other chromatin regulators (noncolored gears, CR), their target loci and associated modifications (different colored balls) remain unknown.

(B) Our process consists of four steps (left to right): (1) ChIP assays were performed with 145 different antibodies targeting 92 distinct CRs and 15 modifications; (2) ChIP samples were screened for enrichment across 487 signature loci by ChIP-string to identify promising CR antibodies; (3) These antibodies were applied in ChIP-seq to generate genome-wide maps for 29 CRs; and (4) We identified target loci and associated histone modifications for each CR. We found that the CRs partition into six modules with correlated binding patterns and also exhibit instances of combinatorial binding.

restricts our ability to identify the CRs that act at any given locus (Figure 1A), to determine how they impart distinct histone modifications, and to decipher how they affect the regulation of target loci in *cis*.

Here, we describe a general methodology for identifying effective procedures to map CRs in mammalian cells (Figure 1B) and demonstrate the usefulness of the approach by studying the localization of CRs in K562 cells and human embryonic stem (ES) cells. We first developed a meso-scale localization assay, ChIP-string, based on a signature readout of 487 loci representing diverse chromatin states. We then used ChIP-string to screen 145 antibodies and thereby identified effective reagents, which were used to generate genome-wide binding maps for 29 CRs by ChIP-seq.

The resulting data sets provide a comprehensive view of the associations between CRs and their relationships to histone modification states. We found that CRs bind in characteristic modular combinations, each associated with distinct modification patterns and genomic features, and often with different functional groups of genes. For example, HDAC1 and SAP30 cobind sharply over transcription start sites (TSS) of cell cycle-related genes, whereas SIRT6 and CHD7 cobind the proximal portions of highly active genes encoding ribosomal and chromatin architecture proteins. Other sets of CRs coassociate at distal elements or repressed loci. Remarkably, most modules combine CRs with opposing enzymatic activities that likely mediate homeostatic regulation of dynamic chromatin. When comparing different cell types, CRs often redistribute to different genomic regions yet maintain their characteristic modular associations. Our work provides a new experimental approach to use ChIP in high-throughput screens and identify effective antibodies; presents a valuable resource for studying CR location and function; and reveals key principles of chromatin organization.

RESULTS

ChIP-String: Meso-Scale Location Analysis for Chromatin Proteins

We developed a new method to determine, in multiplex, the enrichment of many CRs or histone modifications at hundreds

of representative loci. We reasoned that such a signature binding profile would be highly informative. First, querying several hundred regions is less biased than sampling a handful of loci as typically done by ChIP-PCR. Second, it yields a “signature” pattern that could help determine whether a CR is consistently associated with loci sharing a chromatin state. Third, a signature can be measured faster and at a much lower cost than a genome-wide profile and is thus appropriate for screening antibodies and ChIP conditions for difficult targets or for perturbation screens using RNA interference or small molecules.

As a signature readout, we assembled a panel of 487 genomic loci representing different types of chromatin environments. To choose the regions, we used genome-wide chromatin state annotations for human ES and K562 cells, derived from multiple histone modification maps (Ernst et al., 2011). We selected representative loci for each of the major states in the two cell types, e.g., active or repressed promoters, transcripts, distal elements, etc. (Experimental Procedures and Table S1 available online). We reasoned that individual CRs would localize to subsets of these representative loci and thus enable us to distinguish an effective CR ChIP assay.

To measure enriched binding at the signature loci, we developed the ChIP-string method. ChIP-string leverages the nCounter Analysis System platform (NanoString Technologies), originally developed for multiplex quantification of RNA molecules. We designed a probe set complementary to the signature loci and adapted the nCounter operating procedures for ChIP DNA (Experimental Procedures). We validated ChIP-string by analyzing histone modification ChIPs and comparing the measurements to ChIP-seq data (Figures 2A and S1A).

We evaluated the sensitivity of ChIP-string by conducting the assay with successively smaller quantities of ChIP DNA. We found that a minimum of ~5 ng of DNA is needed to maintain quantitative accuracy. Whereas histone modification ChIPs typically yield more than 5 ng of DNA, CR ChIPs yield much smaller quantities (<1 ng), even when millions of cells are used as starting material. We therefore implemented a rapid genomic amplification step to ChIP DNA prior to nCounter detection and confirmed that it faithfully maintains enrichment for a majority of the signature loci (Figure 2A).

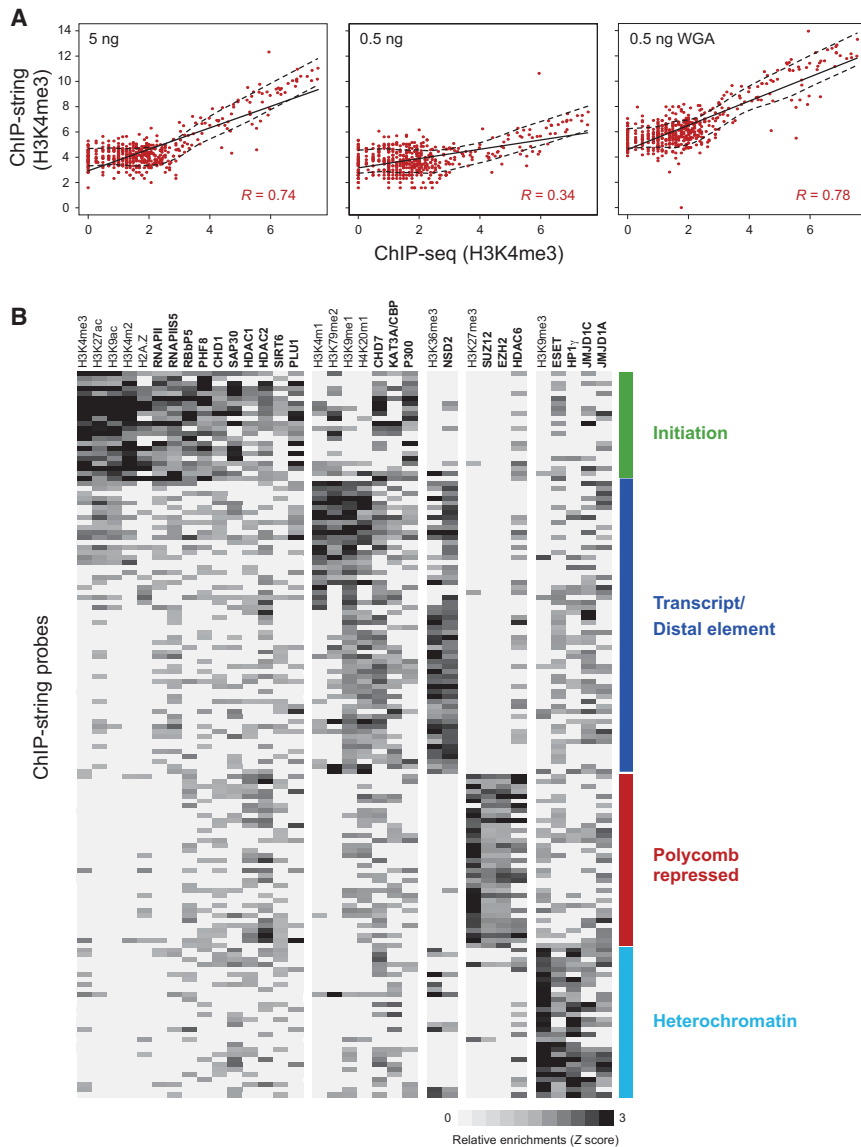


Figure 2. Screening CR Antibodies by ChIP-String

(A) Comparison of H3K4me3 ChIP-string to ChIP-seq data. Scatter plots compare ChIP-seq read density (x axis) against ChIP-string counts (y axis). The adjacent panels reflect ChIP-string experiments performed with 5 ng of DNA (left), 0.5 ng of DNA (middle), or after whole-genome amplification of 0.5 ng of DNA (right). Spearman correlations (R) are indicated at lower right.

(B) ChIP-string assays that scored positively associate with distinct histone modification “states.” Columns represent ChIP-string data for 21 CR antibodies (bold) and 12 histone modification antibodies (nonbold). Relative enrichments are indicated for the 200 most informative loci (rows); white indicates no enrichment, and black indicates high enrichment. The probes were clustered and then sorted by the “chromatin states” of the corresponding locus (initiation, green; elongation or enhancer, purple; Polycomb-repressed, red; and heterochromatin, light blue). The experiments (columns) were ordered by hierarchical clustering and then fine-tuned by visual inspection (Experimental Procedures). Supporting data for ChIP-string and antibody specificity are presented in Figure S1.

ficients between each pair of ChIP-string experiments and hierarchically clustered the data. A substantial majority of the CR binding signatures (~80) either clustered with IgG control antibodies or formed separate clusters with overall weak signals before standardization (Figure S1B). Furthermore, none of these experiments correlated well with any specific histone modification or chromatin state. Although we cannot rule out that they enrich regions not captured by our probe set, we designated these CR antibodies as “failed” in our screen.

The remaining CR ChIP-string experiments were clearly distinct from the IgG and input control experiments (Figure S1B), exhibiting both a larger number of enriched probes as well as higher enrichment values. In many cases, these CR experiments enriched subsets of loci in patterns reminiscent of individual histone modifications or chromatin states (Figure 2B). Regardless, we designated all of these remaining CR antibodies as “passed.” Notably, an alternative analysis procedure, which used different statistical methods in preprocessing and antibody assessment, led to highly similar results, supporting the robustness of the screen. This alternative procedure can be used even when only a few antibodies are tested (Experimental Procedures).

We carried out ChIP-seq for 39 CR antibodies that passed the screen and a sample of 9 failed CR antibodies. Of the 39 passed antibodies, 34 (~90%) yielded high-quality genome-wide profiles as reflected by robust enrichment of specific genomic

ChIP-String Screen Identifies Effective Reagents for Mapping CRs

We applied ChIP-string to 126 CR antibodies, 17 histone modification antibodies, and 2 IgG control antibodies (Table S2). We also analyzed 16 other control samples of unenriched chromatin input. We used chromatin from K562 cells, with the exception that ES cell-specific CRs were profiled using chromatin from ES cells. In 21 cases, more than one antibody was tested for the same target protein, allowing us to evaluate different epitopes. Overall, we screened ~150 samples. We normalized the data by sample and then by probe, using an approach analogous to methods applied to microarray data. We then standardized the measurements in each sample, creating a scale of relative probe enrichment that is comparable across samples (Experimental Procedures).

Next, we distinguished effective CR antibodies from those yielding nonspecific enrichment. We calculated correlation coef-

loci, whereas none of the failed antibodies yielded high-quality data. These results indicate that ChIP-string provides an independent and objective means to identify effective reagents for CR mapping.

A Compendium of Genome-wide CR Maps

We used 29 of the CR antibodies that passed our screen to generate 42 ChIP-seq data sets of the genome-wide distributions of 27 CRs in K562 cells and 15 CRs in ES cells (Figures 3A and S2 and Experimental Procedures). We confirmed the specificity of each of these antibodies by western blots (Figure S1C). We used two independent peak-calling procedures to collate enriched sites for each CR in each cell type (Table S3). The number of sites ranged from 1,680 for HP1 γ to 30,993 for RBBP5 to 39,180 for RNA polymerase II phosphorylated at serine 5 (RNAPIIS5P), with a median of 9,194 sites per CR. The vast majority of enriched regions were between 1 and 2 kb in size.

CR Binding Patterns Reveal a Modular Organization

Comparing the enrichment profiles of the CRs, we found that CRs bind in characteristic combinations. Specifically, we calculated correlations between each pair of CR binding profiles over all regions showing a significant peak in at least one data set (Experimental Procedures). This allowed us to not only compare the different bound locations, but also to consider the shapes of the binding peaks in cases in which the locations overlap. We then hierarchically clustered the CRs based on all pairwise correlations. The resulting dendrogram and correlation matrix reveal striking associations between groups of CRs. These are reflected in six major modules (Figure 3B), each containing between three and six CRs with similar binding profiles. The six modules encompass all of the CR profiles except RE1-binding protein (REST), whose profile is dissimilar to all others. Although REST is extensively implicated in CR recruitment, it is the only sequence-specific DNA binding protein in our compendium, which may explain its failure to conform to the modular organization seen for the other 28 CRs.

CR Modules Associate with Distinct Genomic Features and Chromatin Environments

We next studied the relationship of the CRs and CR modules to genomic features, including promoters, transcribed regions, and distal regulatory elements. CRs within each module exhibit remarkably similar patterns of association to genomic features and chromatin states (Figure 3C), which are distinct between modules. Each localization pattern is consistent with known biology while also providing insight into CR functions. We discuss each module below.

Module I (PHF8, RBBP5, PLU1, CHD1, HDAC1, and SAP30; promoters) is characterized by preferential binding at promoters (Figure 3C), with 65%–80% of binding sites overlapping transcriptional start sites (TSSs). The targets carry H3K4me3 and other modifications related to competent (i.e., nonrepressed) promoters but exhibit a wide range of transcriptional activity based on RNA-seq. The results are consistent with known biology: RBBP5 is a core component of MLL complexes that catalyze H3K4me3 (Smith and Shilatifard, 2010), and PHF8

and CHD1 both bind this modification (Flanagan et al., 2005; Kleine-Kohlbrecher et al., 2010; Sims et al., 2005). In addition, the module contains PLU1 (JARID1B), an H3K4me3 demethylase.

Module I also includes HDAC1 and SAP30, core members of the SIN3 histone deacetylase complex with exquisitely similar binding profiles ($R = 0.92$). Although deacetylases have generally been linked to repression, the robust occupancy of these factors at nonrepressed TSSs is consistent with a prior study that localized deacetylases to many active genes (Wang et al., 2009b). Importantly, such coassociation of CRs with “activating” and “repressive” characteristics in one module is also seen in other modules and nearly all classes of target loci (see below). The cobinding patterns likely reflect widespread roles for opposing CRs in fine-tuning chromatin structure at regulatory loci.

Module II (RNAPIIS5P, SIRT6, NSD2, and CHD7; transcribed regions) is characterized by binding to active promoters as well as proximal and distal transcripts (Figure 3C). In particular, the cobinding patterns suggest interplay between initiating RNAPII (Smith and Shilatifard, 2010) and SIRT6 ($R = 0.70$): 78% of SIRT6-enriched windows reside over the TSS or within the first 5 KB of an active gene (compare to 75% for RNAPIIS5P). Another member of the module, NSD2, also localizes to active transcripts but with greater preference for distal, elongating regions (49% of enriched intervals are within actively transcribed regions). This may reflect interplay between NSD2, a histone methyltransferase, and the elongation mark H3K36 methylation (Nimura et al., 2009). Finally, CHD7 binds promoters, transcribed regions, and some distal elements.

Module III (JARID1C, HDAC2, HDAC6, and ESET; promoters) comprises four CRs with catalytic activities typically associated with repression. These factors colocalize with active and competent promoters, similar to Module I, but also bind repressed targets. JARID1C (SMCX) is an H3K4 demethylase closely related to PLU1 (Module I). HDAC2 and HDAC6 complement HDAC1 (Module I) at active promoters but also associate with Polycomb-repressed targets. Finally, the H3K9 methyltransferase ESET expands the spectrum of known heterochromatic CRs at promoters. These binding patterns suggest prevalent roles for repressive CRs at sites of dynamic chromatin activity.

Module IV (P300, MI2, and LSD1; candidate enhancers) includes three CRs that preferentially bind distal regulatory elements, including sites with enhancer-like chromatin (Figure 3C). Consistent with prior reports (Heintzman et al., 2007), more than 70% of P300 sites are distal from TSSs, and ~50% of those distal regions are enriched for modifications that correlate with enhancer activity, such as H3K4me1 and H3K27Ac (Birney et al., 2007; Ernst et al., 2011; Heintzman et al., 2007). Moreover, ~55% of distal P300 sites coincide with highly conserved sequences (Lindblad-Toh et al., 2011). Module IV also contains two members of the NuRD repressor complex: MI-2 and LSD1 (Wang et al., 2009a). Both CRs bind distal elements, with ~30% overlap to P300 peaks. LSD1 is a demethylase that is specific for mono- and dimethylated H3K4 (Shi et al., 2004), two characteristic methylation states of enhancer chromatin (Birney et al., 2007; Heintzman et al., 2007). These

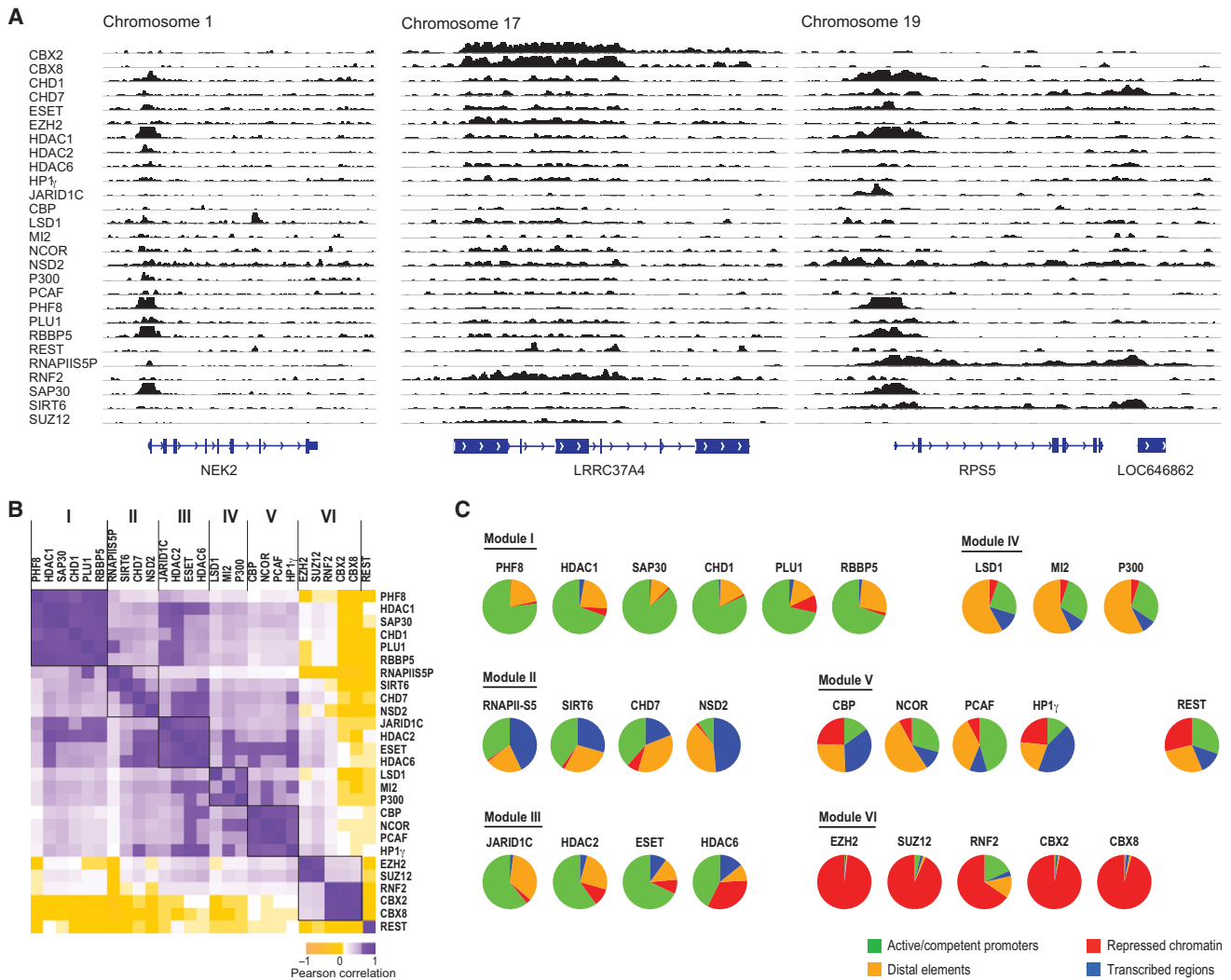


Figure 3. CR Binding Maps Reveal Modular Organization and Coherent Associations with Chromatin States

(A) Binding of CRs at representative genomic loci in K562 cells. ChIP-seq profiles for 27 CRs are shown across three loci (chromosome 1: 211,833,852-211,852,044; chromosome 17: 43,580,509-43,600,984; chromosome 19: 58,895,062-58,910,985). Examples of tracks derived from human ES cells (H1) are shown in Figure S2.

(B) CRs partition into modules with correlated binding profiles. Correlation matrix reflects pairwise correlations of binding peaks between CR data sets. (Purple) Positive correlation between CRs; (yellow) negative correlation; (white) no correlation. Correspondence is evident among CRs within each of the six CR modules (demarcated by black squares).

(C) For each CR, a pie chart indicates the proportion of binding sites that reside in regions with a given chromatin state annotation (green, active/competent promoter; gold, distal regulatory element/candidate enhancer; red, repressed chromatin; blue, transcribed region). CRs within a common module have similar distributions of binding.

associations support a model in which chromatin at distal regulatory elements is tightly regulated by opposing enzymatic activities, as observed for promoters above.

Module V (NCOR, PCAF, CBP, and HP1 γ ; candidate enhancers, other distal features) contains CRs that bind a more diverse set of elements. NCOR, PCAF, and CBP each bind thousands of distal elements, many with enhancer-like characteristics, such as P300 binding. Considering all distal P300 sites, 48% are cobound by CBP, 45% by NCOR, and 35% by PCAF ($p < 10^{-15}$ in all cases). Nevertheless, these three

CRs also bind many other loci, accounting for their overall lower correlation with P300 and separate module. CBP is closely related to P300 and has also been shown to bind enhancers (Kim et al., 2010). PCAF and NCOR are antagonistic regulators associated with nuclear hormone receptor activity and repression, respectively (Perissi et al., 2010). Although they are typically studied at promoters, their colocalization patterns suggest that they also act at enhancers. The partitioning of distal element CRs into separate modules suggests a high degree of specificity among enhancers and their regulators.

HP1 γ , a heterochromatin protein that physically interacts with H3K9me3, occupies diverse chromatin environments. Its association to this module reflects frequent cobinding of distal elements with CBP. However, HP1 γ also correlates with CRs in other modules and binds repetitive elements, Polycomb-repressed regions, and ZNF gene clusters (O'Geen et al., 2007).

Module VI (EZH2, SUZ12, CBX2, CBX8, and RNF2; Polycomb-repressed) comprises core components of Polycomb-repressive complexes 1 and 2 (PRC1 and PRC2). Binding occurs almost exclusively in regions enriched for H3K27me3, which typically correspond to transcriptionally inactive, GC-rich promoters (Ku et al., 2008; Lee et al., 2006; Simon and Kingston, 2009). However, RNF2 (RING1B), an E3 ubiquitin ligase also present in other protein complexes (Vidal, 2009), shows a limited extent of binding outside of H3K27me3-marked regions.

Together, these findings portray diverse regulatory functions for CRs and identify combinations of regulators that cobind and likely coregulate common genomic targets. In specific examples, coordination involves multiple CRs in the same protein complex. However, in most cases, CRs in a module show only partial, albeit significant, overlap, consistent with both shared and unique regulatory functions.

Fine-Scale Analysis of CR Binding Patterns across Promoters

To evaluate the extent and significance of the modular CR organization and whether it is also guided by combinatorial principles, we next systematically examined CR binding patterns at individual loci. We inspected all promoters bound by more than one regulator. We focused on promoters because roughly half of CR binding events occur within 3 kb of a TSS, and nearly all CRs show some binding across such regions. We clustered the 1,081 promoters that are highly enriched for at least two CRs (Experimental Procedures) by the combinatorial binding of the 18 CRs with substantial promoter occupancy. We also grouped the CRs based on their localization patterns across these loci. This promoter-focused grouping (CR groups) is largely consistent with the CR modules deduced from genome-wide correlations. However, this fine-scale analysis highlights differential associations of individual CRs with TSSs and flanking regions, as well as differential relations to gene activity.

A first group of CRs—PLU1, CHD1, SIRT6, and CHD7—exhibits binding profiles that are characteristic of RNAPII initiation (Figure 4A). Although enriched across all transcriptionally competent promoters, these CRs are most strongly bound at highly active promoters undergoing productive initiation and elongation, as indicated by the high expression levels of the corresponding genes. Their broad binding distributions over TSSs emulate RNAPIIS5P (Figures 4A and 4B). Fine binding patterns thus identify additional CRs with close connections—and possible direct physical interactions—with initiating RNAPII (Smith and Shilatifard, 2010).

A second larger CR group—ESET, HDAC6, JARID1C, HDAC2, HDAC1, SAP30, and RBBP5—binds active and competent promoters (Figure 4A) in sharp peaks that precisely coincide with TSSs (Figure 4C). In addition to facilitating RNAPII engagement, these CRs may help to maintain chromatin integrity around

the nucleosome-free TSSs (Jiang and Pugh, 2009) by fine-tuning modifications of the flanking -1 and $+1$ nucleosomes.

A third group of CRs—EZH2, SUZ12, RNF2, CBX2, and CBX8—includes core components of PRC2, which catalyzes H3K27me3, and PRC1, which binds H3K27me3 and mediates chromatin compaction (Margueron and Reinberg, 2010; Simon and Kingston, 2009) (Figure 4A). These CRs bind inactive promoters, many of which correspond to genes involved in development or signaling. Remarkably, PRC2 and PRC1 subunits exhibit distinct fine-scale binding profiles over the promoters (Figure 4D). PRC2 components (EZH2 and SUZ12) peak over TSSs, potentially reflecting interactions with DNA sequences in these nucleosome-depleted regions. In contrast, PRC1 components (CBX2 and CBX8) bind broadly across the same regions, likely promoted by physical interactions with flanking H3K27me3-marked nucleosomes (Figure 4D). Notably, Polycomb-repressed promoters are the only set of genomic elements in our study that are not subject to opposing chromatin regulatory activities, as they are bound exclusively by repressive CRs in K562 cells.

Fine-Scale Promoter Analysis Reveals Combinatorial Complexity of CR Associations

Although the promoter clustering largely corresponds to the modular organization discerned from genome-wide correlations, it also reveals several exceptions that may reflect combinatorial CR binding. In some cases, different CRs bind the same promoters but with distinct binding structures. For example, despite largely overlapping targets, CHD1 and PLU1 exhibit markedly different binding patterns. CHD1 peaks sharply over TSSs, whereas PLU1 extends well into transcribed regions (Figure 4E).

In other cases, a CR is associated with different CR groups under different promoter contexts (Figures 4A, 4F, and 4G). Particularly striking examples of such combinatorial partitioning involve deacetylase complexes (Yang and Seto, 2008). SIN3 complex members HDAC1, HDAC2, and SAP30 bind promoters of genes that oscillate during the cell cycle with an intensity that distinguishes them from all other targets (Figures 4A and 4G, cluster 5). In addition, HDAC1, HDAC2, and JARID1C (members of the CoREST complex [Tahiliani et al., 2007]) cobind along with HDAC6 to repressed PRC2 targets (Figure 4A, clusters 6–8). This association may reflect physical interactions between CoREST and Polycomb complexes (Ren and Kerppola, 2011; Tsai et al., 2010) and/or direct interactions between HDAC2 and PRC2 (van der Vlag and Otte, 1999). The fine binding patterns of these CRs vary dramatically based on the context of the target gene's activity or the cobinding CRs. For example, HDAC2 binds sharply over transcriptionally competent TSSs but distributes broadly over Polycomb-repressed promoters (Figures 4A, clusters 5 and 13, and 4F). This is consistent with a model in which histone deacetylases act as fine-tuners of accessible chromatin at competent TSSs but as enforcers of hypo-acetylated chromatin domains at Polycomb-repressed loci.

Combinatorial CR Binding Patterns Are Associated with Refined Functional Distinctions

We next explored whether individual CRs or CR combinations might be associated with specific cellular processes. The

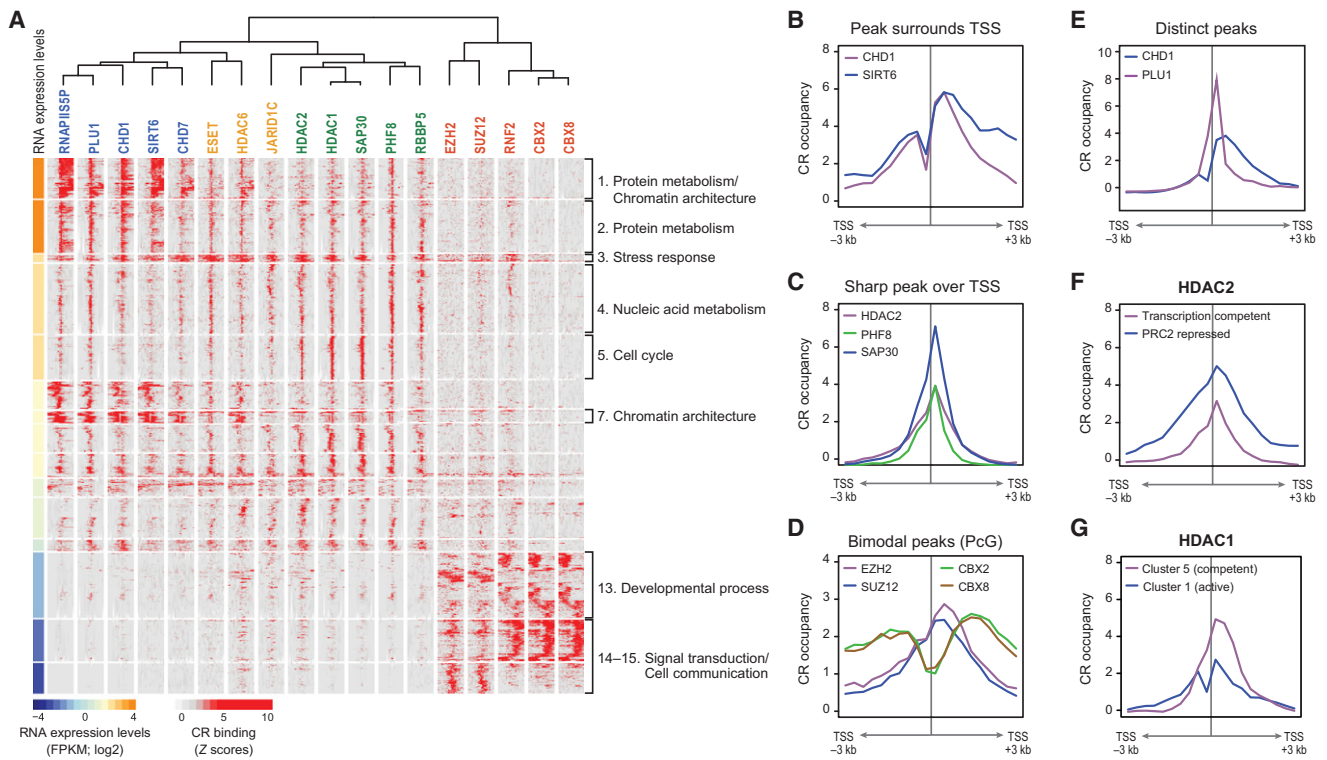


Figure 4. Fine-Scale CR Binding Profiles Distinguish Coherent Gene Sets

(A) Combinatorial binding patterns of CRs at individual promoters are associated with distinct expression and function. Fine-scale binding profiles are shown for CRs (vertical sections) across 1,081 target promoters (rows) after hierarchical clustering and reordering of major promoter clusters based on expression (original ordering shown in Figure S3). The profiles depict enrichment Z scores (red, high; white, low) for 300 bp windows within 3 kb of TSSs. Promoter clusters derived based on common CR binding profiles (indicated by thick white horizontal lines) share transcriptional status, and the corresponding genes often share coherent functions (labels on right, curated from enriched functional gene sets, listed in Table S4B). Left bar shows RNA expression levels (log₂(FPKM)) derived from RNA-seq data: orange, high; blue, low; white, median. CR labels are colored according to the dendrogram shown above.

(B–D) Composite profiles for CRs with similar binding patterns at shared promoters. Profiles reflect average binding of the indicated CRs (y axis) over cobound promoters, centered on TSSs (x axis).

(B) Peaks that surround TSSs but dip at the TSS itself for CHD1 (purple) and SIRT6 (blue).

(C) Sharp peaks over TSSs for HDAC2 (purple), PHF8 (green), and SAP30 (blue).

(D) Bimodal peaks of Polycomb CRs—diffuse peaks over TSSs for PRC2 members EZH2 (purple) and SUZ12 (blue) along with TSS-excluded peaks for PRC1 members CBX2 (green) and CBX8 (brown).

(E) Composite profiles for CRs with distinct binding patterns at shared promoters. PLU1 (purple) has a sharp peak over TSSs, whereas CHD1 (blue) has a broader peak that extends downstream. Note that the promoters in this composite differ from (B).

(F and G) Composite profiles show distinct patterns for the same CR at composite sets with different activity levels.

(F) HDAC2 at PRC2-repressed (blue) and transcriptionally competent promoters (purple).

(G) HDAC1 at transcriptionally competent (cluster 5, purple) and active (cluster 1, blue) promoters.

promoter-based analysis revealed 15 “combinatorial binding” gene clusters, each of which shares binding by a combination of CRs, as well as a fine CR location structure around their TSSs (Figures 4A and S3, horizontal blocks). The genes in many of these clusters are characterized by shared functional attributes (Figure 4A, labels on right, and Table S4). In particular, genes with similar expression levels but distinct biological functions are often bound by distinct combinations of CRs. For example, the “Protein Metabolism” cluster (Figure 4A, cluster 2; 110 genes) is comprised of highly expressed genes whose promoters are cobound by SIRT6, CHD1, PLU1, and RNAPIIS5P. A distinct cluster that consists of 84 genes with similarly high expression but whose promoters are cobound by these CRs along with CHD7 and HDAC6 is enriched

for genes involved in chromatin architecture (cluster 1). A separate binding cluster (cluster 5; 92 genes) enriched for cell-cycle gene promoters is unremarkable in terms of its intermediate expression levels but is prominently cobound by HDAC1, HDAC2, and SAP30. The physical association of these core SIN3 components with these promoters offers a mechanistic explanation for documented roles for this repressor complex and histone deacetylase activity in cell-cycle progression (David et al., 2008; Minucci and Pelicci, 2006). Interestingly, the promoters in the “stress response” cluster are cobound by most of the activating and repressive CRs in our panel, which may play important roles in the notable capacity of these genes to rapidly change their activity in response to stimuli.

CRs Occupy Different Loci in ES Cells but Maintain Their Modular Associations

We next explored whether the colocalization patterns and associations observed in K562 cells can be generalized to other cell types. We considered several layers of CR organization. First, we asked whether CRs distribute to different genomic locations, consistent with changed gene expression programs. Second, we asked whether the associations between individual CRs and chromatin modification states change between cell types. Third, we asked whether the modular relationships between CRs are maintained in different cell types.

To examine each of these possibilities, we generated ChIP-seq data for 15 CRs in human ES cells and analyzed their localization patterns (Figures 5A and 5B). We used the same computational methods as in K562 cells to identify regions of enrichment, which yielded similar overall statistics (Table S3).

The CRs differ substantially in their genomic location between the cell types, though the degree of overlap varies between CRs (Figure 5A, left). The patterns of relocalization are reminiscent of histone modifications (Figure 5A, right), which dynamically change between these cell types (Ernst et al., 2011), consistent with differential transcriptional programs.

Despite substantial differences in CR localization, the underlying CR organization is maintained between the cell types (Figure 5B). First, the degree of cobinding between pairs of CRs is conserved between the two cell types ($R = 0.64$). Similarly, the degree of correspondence between a given CR and a given histone modification is also well correlated ($R = 0.79$). Thus, CR-CR associations as well as CR-histone modification associations are globally preserved between cell types.

Furthermore, the relationships between individual CRs and genomic annotations remain largely unchanged. For most CRs, the distribution of binding between promoter, transcribed, distal, and repressed regions is highly concordant between K562 and ES cells (Figure 5C). Conservation of binding patterns is also evident when comparing the fine-scale promoter profiles of CRs in ES cells (Figure 5D) to those in K562 cells (Figure 4A). Consistent patterns of binding are evident for CRs associated with competent TSSs (e.g., PHF8, RBBP5, and SAP30), productive initiation (e.g., CHD1 and SIRT6) and Polycomb repression (e.g., EZH2 and SUZ12). Gene sets distinguished based on combinatorial binding profiles are also similar between the cell types (Figures 5D and S4 and Table S5).

Notably, when there are changes in CR localization, they tend to be shared by members of the same module and to relate to a fundamental difference in chromatin structure between cells (Figure 5C). For example, although Module I CRs (e.g., PHF8, CHD1, and RBBP5) are restricted to active and competent promoters in K562 cells, they also associate with Polycomb-repressed promoters in ES cells (Figures 5C and 5D). The presence of multiple activating CRs at these inactive targets is consistent with the enrichment of the underlying chromatin for opposing (bivalent) histone modifications. These CRs likely contribute to the poised character of the corresponding genes, many of which are induced during ES cell differentiation (Bernstein et al., 2006). In addition, P300 binds substantially fewer sites in ES cells than in K562 cells (Figures 5A and 5C), possibly

reflecting a lower prevalence of enhancer-like chromatin in ES cells (Ernst et al., 2011).

Overall, our analysis suggests that the modular and combinatorial structures of CRs, and their association with histone modification states, are constitutive features of the chromatin regulatory network. Thus, changes in CR binding tend to be coordinated at the level of modules and to correspond to changes in the underlying chromatin landscape.

DISCUSSION

Modular and Combinatorial Organization of the CR Network

Despite their large number and the importance of chromatin organization to gene regulation, the localization and function of individual CRs remains poorly understood. Studies of histone modification patterns have revealed a relatively limited number of chromatin configurations, or “states,” that distinguish different types of genome regulatory elements. It has been compelling to hypothesize that specific CRs contribute to the establishment and maintenance of these states in different cell types and that they work in a combinatorial fashion, akin to transcription factors, which are encoded in a comparable number in the genome. However, it has been difficult to develop detailed models of CR function given the limited availability of comprehensive measurements and the paucity of effective capture reagents.

Here, we presented a first systematic view of CR localization across the human genome in two cell types and a general methodology for studying the targeting and functions of such regulators. We reveal several major principles for the organization of the CR network in mammalian cells (Figure 6). (1) Coherent modules of CRs cobind to common target loci that share specific chromatin states; the modules often consist of modifying enzymes that catalyze activating and repressive modifications, offering a means for precise tuning of chromatin and gene regulation. (2) In addition to these global associations, the same CR may associate with different modules at different target loci, suggesting complex functional relationships, indicative of combinatorial regulation. (3) Specific combinations of CRs bind sets of genes with related functions, suggesting functional specificity. (4) When comparing different cell types, CRs distribute to different loci, often in conjunction with changes in chromatin states; however, (5) they largely retain their modular associations.

In many respects, this view is reminiscent of the organization of sequence-specific transcription factor networks. In particular, the association of CRs within modules—each related to different chromatin modification states, functional gene groups, and expression patterns—is consistent with the modular organization of transcription factor networks in organisms from yeast to human (Yosef and Regev, 2011). Nevertheless, we cannot rule out the possibility that other CRs, not tested in our study, might adopt different or possibly nonmodular organizations that do not conform to any CR modules defined here.

Due to their interconnected organization, changes in the expression of an individual CR may affect the function of one or more modules in which it participates, with potentially

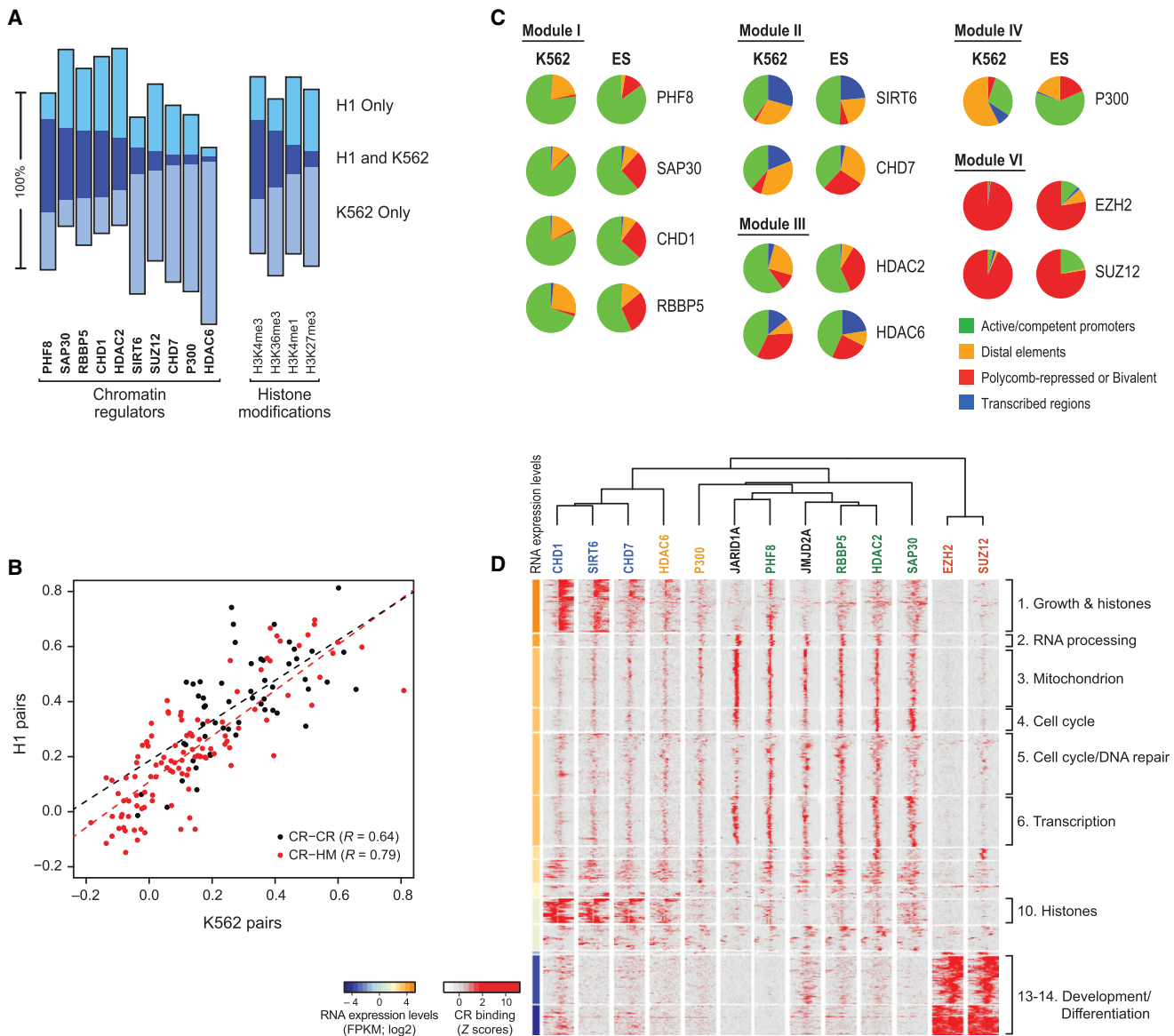


Figure 5. Comparisons of CR Binding and Modular Associations in K562 and ES Cells

(A) CRs distribute to distinct loci in ES and K562 cells. For each CR (left) or histone modification (right), bar graph indicates proportions of binding intervals that are ES cell specific (“H1 only,” light blue), K562 specific (“K562 only,” gray), or overlapping between cell types (“H1 and K562,” navy). The bars are vertically centered according to the overlap regions.

(B) CR-CR associations and CR-histone modification associations are largely preserved between K562 and ES cells. Scatter plot presents the correlations in localization profiles between every pair of CRs (black dots) and every CR-histone modification pair (red dots) in either K562 cells (x axis) or ES cells (y axis). Linear regression lines and correlation coefficients are indicated for each type of combination.

(C) CRs associate with similar chromatin states in ES and K562 cells with some distinctions. Pie charts indicate the proportion of CR binding sites that correspond to a given chromatin state annotation (green, active/competent promoter; gold, distal regulatory element/candidate enhancer; red, repressed chromatin [including bivalent state]; and blue, transcribed region). CRs are grouped according to the modules in Figure 3.

(D) Combinatorial binding profiles are shown for CRs in ES cells. Fine-scale binding profiles are shown for each CR across 1189 target promoters (rows) in ES cells, after hierarchical clustering and re-ordering of major promoter clusters as in Figure 4A (original ordering shown in Figure S4). Functional gene set annotations (right) curated from enriched sets (Table S5B). CR labels colored as in Figure 4A, with ES cell-specific CRs in black.

widespread consequences for gene expression and cellular phenotype. Such network properties could help explain how dynamic changes in CR expression guide differentiation processes and how genetic inactivation of CRs promotes tumor

progression. However, the binding modules derived here do not predict how the removal of specific components will affect other participants or downstream targets. Further study is therefore needed to derive more detailed functional models of the direct

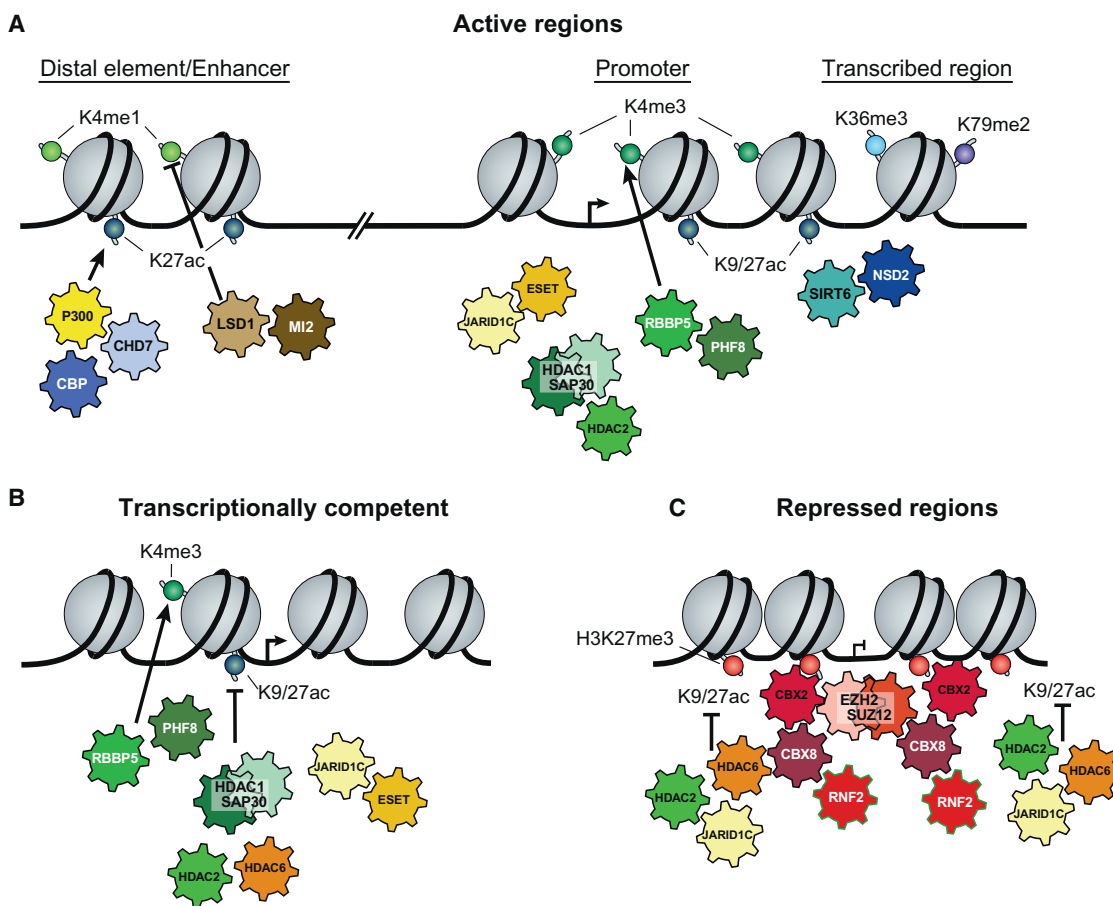


Figure 6. Principles of CR Organization

(A–C) Model of the association of CRs (gears) with histone modifications (little colored balls) across (A) actively transcribed genes, (B) transcriptionally competent TSSs, and (C) Polycomb-repressed regions. CRs colored as in Figures 4A and 5D.

physical interactions between CRs and the associated binding hierarchies.

The organizational principles also suggest how CR binding can tune expression programs. Each of the CR modules targeting transcriptionally active or competent (nonrepressed) promoters contain proteins with opposing activities—those that catalyze the addition of modifications associated with active/accessible chromatin and those that catalyze their removal. Such opposing activities in bifunctional modules may underlie homeostasis at active chromatin loci and allow precise tuning of gene expression. Because distal enhancers are also bound by activating and repressive CRs, they too may be subject to similar fine-tuning. Genomic loci targeted by Polycomb proteins in K562 cells are an outlier in this regard, as they appear to be exclusively subject to repressive histone modifiers.

Specific Hypotheses for CR Function

The network organization suggests many specific hypotheses regarding the functions or molecular mechanisms of individual CRs or CR complexes. For example, in both K562 and ES cells, components of the SIN3 repressor complex are strikingly enriched at cell-cycle gene promoters, providing a potential mech-

anistic explanation for known roles for deacetylases in cell-cycle progression. In another example, we find that several repressive CRs bind both to competent TSSs and to Polycomb-repressed targets. These repressors, which include histone deacetylases and an H3K4 demethylase, likely enforce the hypo-acetylated and H3K4 unmethylated state that is characteristic of these repressed loci.

A Meso-Scale Assay for CR-DNA Interactions

Our ChIP-string assay opens the way to further functional studies of these and other CRs in many cell systems. It allowed us to screen hundreds of antibody-condition combinations for CR localization, through which we identified a new set of effective reagents for mapping CRs. We expect this screening approach will help to overcome the current paucity of ChIP-seq grade antibodies for studying the several hundred CRs that control chromatin structure and function. The modest success rate of antibodies tested here suggests that this will also require substantial efforts to develop antibodies against different CR epitopes as well as new types of affinity reagents.

The multiplexed assay for CR binding and histone modifications also has the potential to greatly enhance functional

studies of chromatin. Its rapid turnaround and low cost will enable systematic studies of perturbations induced by small molecules or RNA interference, which have traditionally been restricted to downstream phenotypic readouts such as protein or RNA expression. In particular, this will help to assess the functional impact of the components and organization of the CR network.

A Systematic Resource of Genome-wide CR Binding Profiles

Our data set provides an important resource for studying CRs at an unprecedented scope. Prior studies of mammalian CR binding typically considered very few factors and used varied procedures and cell types, all of which precluded systematic comparisons. In contrast, our resource allows direct comparison of many CRs in the same cell and between cell types. It also provides a reference to which users may compare their CR or transcription factor profiles, with the potential to predict binding partners and cellular functions. It should therefore enable the large community of chromatin biologists to develop and test mechanistic hypotheses, ultimately leading to a more comprehensive understanding of chromatin organization and gene regulation.

EXPERIMENTAL PROCEDURES

Data Access

All raw data, mapped reads, and integrated profiles are available at <http://www.broadinstitute.org/software/crome/>. Data sets are also available at the ENCODE website (<http://genome.ucsc.edu/ENCODE>) and the Gene Expression Omnibus (GSE32509).

Chromatin Regulator Antibodies

We collated a list of 515 proteins with annotated functions related to histone modification, histone binding, or chromatin remodeling. We obtained a total of 128 antibodies to these proteins, which we tested in the ChIP-string assay. A list of all antibodies annotated by their performance in ChIP-string and ChIP-seq is provided in Table S2. The specificity of all antibodies used in ChIP-seq was confirmed by western blots (Figure S1C). Roughly 20 million K562 cells or H1 ES cells were used for each ChIP assay. Detailed procedures are in the Supplemental Information.

Representative Genomic Loci and nCounter Probe Design

We chose a set of genomic loci designed to be representative of diverse chromatin environments. We used a hidden Markov model (Ernst and Kellis, 2010) and ChIP-seq maps for ten chromatin marks in K562 and ES cells (Ernst et al., 2011) to identify ten major chromatin states and annotate the genome accordingly. For each state in each cell type, we randomly selected 20 loci and used the corresponding sequences for probe design (Table S1).

We modified the nCounter Analysis System platform (NanoString Technologies) to measure enriched genomic DNA from ChIP experiments (ChIP-string). Detailed descriptions of probe set design and ChIP-string procedures are in the Supplemental Information.

ChIP-String Data Analysis and Processing

We devised two alternative analysis methods for the ChIP-string screen. The first ("original") approach, optimal for large-scale screens, was used to score the screen and select ChIP experiments for sequencing. The second ("alternative") approach is suitable for both large- and small-scale studies, even for those testing just a few antibodies. The results of the two approaches on our screen data agree very closely. Detailed descriptions are in the Supplemental Information.

ChIP-Seq Analysis

ChIP-seq was performed as described (Mikkelsen et al., 2007), followed by identification of enriched intervals, which were correlated to genomic elements and chromatin states. Pearson correlations were calculated between every pair of CRs, based on signal distributions across enriched intervals, and were used to produce pairwise cluster maps. CR binding profiles were used to hierarchically cluster promoters. Expression values were derived from RNA-seq data. Full details are provided in the Supplemental Information.

ACCESSION NUMBERS

CR binding maps have been deposited at the Gene Expression Omnibus (GSE32509).

SUPPLEMENTAL INFORMATION

Supplemental Information includes Extended Experimental Procedures, four figures, and five tables and can be found with this article online at [doi:10.1016/j.cell.2011.09.057](https://doi.org/10.1016/j.cell.2011.09.057).

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REFERENCES

- Barski, A., Cuddapah, S., Cui, K., Roh, T.Y., Schones, D.E., Wang, Z., Wei, G., Chepelev, I., and Zhao, K. (2007). High-resolution profiling of histone methylations in the human genome. *Cell* 129, 823–837.
- Bernstein, B.E., Mikkelsen, T.S., Xie, X., Kamal, M., Huebert, D.J., Cuff, J., Fry, B., Meissner, A., Wernig, M., Plath, K., et al. (2006). A bivalent chromatin structure marks key developmental genes in embryonic stem cells. *Cell* 125, 315–326.
- Birney, E., Stamatoyannopoulos, J.A., Dutta, A., Guigó, R., Gingeras, T.R., Margulies, E.H., Weng, Z., Snyder, M., Dermitzakis, E.T., Thurman, R.E., et al; ENCODE Project Consortium; NISC Comparative Sequencing Program; Baylor College of Medicine Human Genome Sequencing Center; Washington University Genome Sequencing Center; Broad Institute; Children's Hospital Oakland Research Institute. (2007). Identification and analysis of functional elements in 1% of the human genome by the ENCODE pilot project. *Nature* 447, 799–816.
- David, G., Grandinetti, K.B., Finnerty, P.M., Simpson, N., Chu, G.C., and Depinho, R.A. (2008). Specific requirement of the chromatin modifier mSin3B in cell cycle exit and cellular differentiation. *Proc. Natl. Acad. Sci. USA* 105, 4168–4172.
- Eisässer, S.J., Allis, C.D., and Lewis, P.W. (2011). Cancer. New epigenetic drivers of cancers. *Science* 331, 1145–1146.
- Ernst, J., and Kellis, M. (2010). Discovery and characterization of chromatin states for systematic annotation of the human genome. *Nat. Biotechnol.* 28, 817–825.

- Ernst, J., Kheradpour, P., Mikkelsen, T.S., Shores, N., Ward, L.D., Epstein, C.B., Zhang, X., Wang, L., Issner, R., Coyne, M., et al. (2011). Mapping and analysis of chromatin state dynamics in nine human cell types. *Nature* 473, 43–49.
- Filion, G.J., van Bommel, J.G., Braunschweig, U., Talhout, W., Kind, J., Ward, L.D., Brugman, W., de Castro, I.J., Kerkhoven, R.M., Bussemaker, H.J., and van Steensel, B. (2010). Systematic protein location mapping reveals five principal chromatin types in *Drosophila* cells. *Cell* 143, 212–224.
- Flanagan, J.F., Mi, L.Z., Chruszcz, M., Cymborowski, M., Clines, K.L., Kim, Y., Minor, W., Rastinejad, F., and Khorasanizadeh, S. (2005). Double chromodomains cooperate to recognize the methylated histone H3 tail. *Nature* 438, 1181–1185.
- Heintzman, N.D., Stuart, R.K., Hon, G., Fu, Y., Ching, C.W., Hawkins, R.D., Barrera, L.O., Van Calcar, S., Qu, C., Ching, K.A., et al. (2007). Distinct and predictive chromatin signatures of transcriptional promoters and enhancers in the human genome. *Nat. Genet.* 39, 311–318.
- Ho, L., and Crabtree, G.R. (2010). Chromatin remodelling during development. *Nature* 463, 474–484.
- Jiang, C., and Pugh, B.F. (2009). Nucleosome positioning and gene regulation: advances through genomics. *Nat. Rev. Genet.* 10, 161–172.
- Kim, T.K., Hemberg, M., Gray, J.M., Costa, A.M., Bear, D.M., Wu, J., Harmin, D.A., Laptewicz, M., Barbara-Haley, K., Kuersten, S., et al. (2010). Widespread transcription at neuronal activity-regulated enhancers. *Nature* 465, 182–187.
- Kleine-Kohlbrecher, D., Christensen, J., Vandamme, J., Abarategui, I., Bak, M., Tommerup, N., Shi, X., Gozani, O., Rappsilber, J., Salcini, A.E., and Helin, K. (2010). A functional link between the histone demethylase PHF8 and the transcription factor ZNF711 in X-linked mental retardation. *Mol. Cell* 38, 165–178.
- Kouzarides, T. (2007). Chromatin modifications and their function. *Cell* 128, 693–705.
- Ku, M., Koche, R.P., Rheinbay, E., Mendenhall, E.M., Endoh, M., Mikkelsen, T.S., Presser, A., Nusbaum, C., Xie, X., Chi, A.S., et al. (2008). Genomewide analysis of PRC1 and PRC2 occupancy identifies two classes of bivalent domains. *PLoS Genet.* 4, e1000242.
- Lee, T.I., Jenner, R.G., Boyer, L.A., Guenther, M.G., Levine, S.S., Kumar, R.M., Chevalier, B., Johnstone, S.E., Cole, M.F., Isono, K., et al. (2006). Control of developmental regulators by Polycomb in human embryonic stem cells. *Cell* 125, 301–313.
- Lindblad-Toh, K., Garber, M., Zuk, O., Lin, M., Parker, B., Washietl, S., Kheradpour, P., Ernst, J., Jordan, G., Mauceli, E., et al. (2011). A high-resolution map of human evolutionary constraint using 29 mammals. *Nature*. Published online October 12, 2011. 10.1038/nature10530.
- Margueron, R., and Reinberg, D. (2010). Chromatin structure and the inheritance of epigenetic information. *Nat. Rev. Genet.* 11, 285–296.
- Mikkelsen, T.S., Ku, M., Jaffe, D.B., Issac, B., Lieberman, E., Giannoukos, G., Alvarez, P., Brockman, W., Kim, T.K., Koche, R.P., et al. (2007). Genome-wide maps of chromatin state in pluripotent and lineage-committed cells. *Nature* 448, 553–560.
- Minucci, S., and Pelicci, P.G. (2006). Histone deacetylase inhibitors and the promise of epigenetic (and more) treatments for cancer. *Nat. Rev. Cancer* 6, 38–51.
- Nimura, K., Ura, K., Shiratori, H., Ikawa, M., Okabe, M., Schwartz, R.J., and Kaneda, Y. (2009). A histone H3 lysine 36 trimethyltransferase links Nkx2-5 to Wolf-Hirschhorn syndrome. *Nature* 460, 287–291.
- O’Geen, H., Squazzo, S.L., Iyengar, S., Blahnik, K., Rinn, J.L., Chang, H.Y., Green, R., and Farnham, P.J. (2007). Genome-wide analysis of KAP1 binding suggests autoregulation of KRAB-ZNFs. *PLoS Genet.* 3, e89.
- Perissi, V., Jepsen, K., Glass, C.K., and Rosenfeld, M.G. (2010). Deconstructing repression: evolving models of co-repressor action. *Nat. Rev. Genet.* 11, 109–123.
- Ren, X., and Kerppola, T.K. (2011). REST interacts with Cbx proteins and regulates polycomb repressive complex 1 occupancy at RE1 elements. *Mol. Cell Biol.* 31, 2100–2110.
- Ruthenburg, A.J., Li, H., Patel, D.J., and Allis, C.D. (2007). Multivalent engagement of chromatin modifications by linked binding modules. *Nat. Rev. Mol. Cell Biol.* 8, 983–994.
- Shi, Y., Lan, F., Matson, C., Mulligan, P., Whetstone, J.R., Cole, P.A., Casero, R.A., and Shi, Y. (2004). Histone demethylation mediated by the nuclear amine oxidase homolog LSD1. *Cell* 119, 941–953.
- Simon, J.A., and Kingston, R.E. (2009). Mechanisms of polycomb gene silencing: knowns and unknowns. *Nat. Rev. Mol. Cell Biol.* 10, 697–708.
- Sims, R.J., III, Chen, C.F., Santos-Rosa, H., Kouzarides, T., Patel, S.S., and Reinberg, D. (2005). Human but not yeast CHD1 binds directly and selectively to histone H3 methylated at lysine 4 via its tandem chromodomains. *J. Biol. Chem.* 280, 41789–41792.
- Smith, E., and Shilatifard, A. (2010). The chromatin signaling pathway: diverse mechanisms of recruitment of histone-modifying enzymes and varied biological outcomes. *Mol. Cell* 40, 689–701.
- Tahiliani, M., Mei, P., Fang, R., Leonor, T., Rutenberg, M., Shimizu, F., Li, J., Rao, A., and Shi, Y. (2007). The histone H3K4 demethylase SMCX links REST target genes to X-linked mental retardation. *Nature* 447, 601–605.
- Tsai, M.C., Manor, O., Wan, Y., Mosammaparast, N., Wang, J.K., Lan, F., Shi, Y., Segal, E., and Chang, H.Y. (2010). Long noncoding RNA as modular scaffold of histone modification complexes. *Science* 329, 689–693.
- van der Vlag, J., and Otte, A.P. (1999). Transcriptional repression mediated by the human polycomb-group protein EED involves histone deacetylation. *Nat. Genet.* 23, 474–478.
- Venters, B.J., Wachi, S., Mavrich, T.N., Andersen, B.E., Jena, P., Sinnamon, A.J., Jain, P., Roller, N.S., Jiang, C., Hemeryck-Walsh, C., and Pugh, B.F. (2011). A comprehensive genomic binding map of gene and chromatin regulatory proteins in *Saccharomyces*. *Mol. Cell* 41, 480–492.
- Vidal, M. (2009). Role of polycomb proteins Ring1A and Ring1B in the epigenetic regulation of gene expression. *Int. J. Dev. Biol.* 53, 355–370.
- Wang, Y., Zhang, H., Chen, Y., Sun, Y., Yang, F., Yu, W., Liang, J., Sun, L., Yang, X., Shi, L., et al. (2009a). LSD1 is a subunit of the NuRD complex and targets the metastasis programs in breast cancer. *Cell* 138, 660–672.
- Wang, Z., Zang, C., Cui, K., Schones, D.E., Barski, A., Peng, W., and Zhao, K. (2009b). Genome-wide mapping of HATs and HDACs reveals distinct functions in active and inactive genes. *Cell* 138, 1019–1031.
- Yang, X.J., and Seto, E. (2008). The Rpd3/Hda1 family of lysine deacetylases: from bacteria and yeast to mice and men. *Nat. Rev. Mol. Cell Biol.* 9, 206–218.
- Yosef, N., and Regev, A. (2011). Impulse control: temporal dynamics in gene transcription. *Cell* 144, 886–896.
- Zhang, Z., and Pugh, B.F. (2011). High-resolution genome-wide mapping of the primary structure of chromatin. *Cell* 144, 175–186.
- Zhou, V.W., Goren, A., and Bernstein, B.E. (2011). Charting histone modifications and the functional organization of mammalian genomes. *Nat. Rev. Genet.* 12, 7–18.