Interactome maps of mouse gene regulatory domains reveal basic principles of transcriptional regulation

Kyong-Rim Kieffer Kwon,1,*, Zhonghui Tang,2,*, Ewy Mathe,1,*, Jason Qian,1,*, Myong-Hee Sung,3,*, Guoliang Li,2, Wolfgang Resch,1, Songjoon Baek,3, Nathanael Pruett,1, Lars Grøntved,3, Laura Vian,1, Steevenson Nelson,1, Hossein Zare,4, Ofir Hakim,5, Deepak Reyon,6,7, Arito Yamane,1, Hirotaka Nakashashi,1, Alexander L. Kovalchuk,8, Jizhong Zou,9, J. Keith Joung,6,7, Vittorio Sartorelli,4, Chia-Lin Wei,10, Xiaoan Ruan,2, Gordon L. Hager,3,*, Yijun Ruan,2,*, and Rafael Casellas,1,11,*

1Genomics & Immunity, NIAMS, National Institutes of Health, Bethesda, MD 20892, USA
2The Jackson Laboratory for Genomic Medicine, and Department of Genetic and Development Biology, University of Connecticut, 400 Farmington, CT 06030, USA
3Laboratory of Receptor Biology and Gene Expression, NCI, National Institutes of Health, Bethesda, MD 20892, USA
4Laboratory of Muscle Stem Cells and Gene Regulation, NIAMS, National Institutes of Health, Bethesda, MD 20892, USA
5Bar-Ilan University, Ramat-Gan, 5290002, Israel
6Molecular Pathology Unit, Center for Computational and Integrative Biology, and Center for Cancer Research, Massachusetts General Hospital, Charlestown, MA 02129, USA
7Department of Pathology, Harvard Medical School, Boston, MA 02115 USA
8Laboratory of Immunogenetics, National Institute of Allergy and Infectious Diseases, National Institutes of Health, Rockville, MD 20852, USA
9Laboratory of Stem Cell Biology, NIAMS, National Institutes of Health, Bethesda, MD 20892, USA
10DOE Joint Genome Institute, 2800 Mitchell Drive, Walnut Creek, CA 94598
11Center of Cancer Research, NCI, National Institutes of Health, Bethesda, MD 20892, USA

Correspondence: rafael.casellas@nih.gov.
*These authors contributed equally to this work

ACCESSION NUMBERS
All accession numbers for raw data are provided in Table S1. All bed files used in the text are also available upon request.

SUPPLEMENTAL INFORMATION
Supplemental information includes extended experimental procedures, 7 figures, and 3 tables.

CONFLICT OF INTEREST STATEMENT
J.K.J. has a financial interest in Transposagen Biopharmaceuticals. J.K.J.’s interests were reviewed and are managed by Massachusetts General Hospital and Partners HealthCare in accordance with their conflict of interest policies.

Publisher’s Disclaimer: This is a PDF file of an unedited manuscript that has been accepted for publication. As a service to our customers we are providing this early version of the manuscript. The manuscript will undergo copyediting, typesetting, and review of the resulting proof before it is published in its final citable form. Please note that during the production process errors may be discovered which could affect the content, and all legal disclaimers that apply to the journal pertain.
Abstract

A key finding of the ENCODE project is that the enhancer landscape of mammalian cells undergoes marked alterations during ontogeny. However, the nature and extent of these changes are unclear. As part of the NIH Mouse Regulome Project, we here combined DNaseI hypersensitivity, ChIP-Seq, and ChIA-PET technologies to map the promoter-enhancer interactomes of pluripotent ES cells and differentiated B lymphocytes. We confirm that enhancer usage varies widely across tissues. Unexpectedly, we find that this feature extends to broadly-transcribed genes, including *Myc* and *Pim1* cell cycle regulators, which associate with an entirely different set of enhancers in ES and B cells. By means of high-resolution CpG methylomes, genome editing, and digital footprinting we show that these enhancers recruit lineage-determining factors. Furthermore, we demonstrate that the turning on and off of enhancers during development correlates with promoter activity. We propose that organisms rely on a dynamic enhancer landscape to control basic cellular functions in a tissue-specific manner.

INTRODUCTION

Gene expression during development is orchestrated by promoter sequences and a variety of distal *cis*-regulatory elements. Key among these are enhancers, which associate with promoters to increase the transcriptional output of target genes in a tissue-specific manner (Visel et al., 2009). Enhancers are typically distinguished from non-regulatory DNA by their hypersensitivity to DNaseI digestion (Sabo et al., 2006), and binding of chromatin modifiers. The CBP/p300 acetyltransferase for instance mediates H3K27 acetylation of chromatin at active enhancers (Creyghton et al., 2010). In addition, enhancers display high levels of H3K4 monomethylation (H3K4me1, (Buecker and Wysocka, 2012)), and a relative depletion of H3K4me3 (Heintzman et al., 2007) and the histone variant H2AZ (Kouzine et al., 2013). Based on these parameters, ~400,000 genomic sites displaying enhancer-like features were recently discovered, spanning nearly 10% of the human genome (Consortium et al., 2012).

Enhancers control lineage identity by recruiting transcription factors, cofactors, and RNA Polymerase II (PolII) to target genes. They physically interact with promoters resulting in looping out of intervening sequences (Krivega and Dean, 2012), which in some instances can span over 1Mb of DNA (Nobrega et al., 2003). In contrast to promoters and insulators, which vary little across cell types, the enhancer landscape changes considerably during development (Thurman et al., 2012). This feature predicts that functional connectivity in mammalian cells i) must display a high degree of tissue specificity and ii) should closely reflect transcriptome changes during cell differentiation. However, these ideas have not been fully explored because of the difficulty of mapping promoter-enhancer connections during development.

In the absence of direct approaches, enhancers have been typically assigned to “cognate” promoters based on linear proximity or shared chromatin states. This strategy has limitations because enhancers do not always regulate nor share chromatin profiles with the nearest promoter. Alternatively, chromosome conformation capture techniques have been used to explore regulatory interactions at predefined genomic loci. However, the coverage of 3C-
based techniques alone is insufficient to map promoter-enhancer connectivity in entire genomes (Xie and Ren, 2013). To overcome this challenge, the ChIA-PET protocol was recently developed (Fullwood et al., 2009). ChIA-PET is a ChIP-based method that captures long-range chromatin interactions involving or mediated by a protein of interest such as estrogen receptor alpha in adenocarcinoma cells (Fullwood et al., 2009) or RNA PolII in human cell lines (Li et al., 2012).

We here introduce the NIH Mouse Regulome Project, an initiative that seeks to define the 3D interplay of gene regulatory domains in developing mouse primary cells. In this first report we compare pluripotent embryonic stem (ES) cells and differentiated B lymphocytes. By combining ChIA-PET, CpG methylomes, DNaseI hypersensitivity, transcriptomes, digital footprinting, and TALEN-mediated genome editing, our studies reveal the dynamics of the mouse regulome during ontogeny.

RESULTS

A comprehensive map of regulatory domains and their interactions in mouse primary cells

To characterize the mouse regulome in primary cells we first applied DNaseI hypersensitivity (DHS) followed by deep-sequencing to CD43− B lymphocytes activated in the presence of lipopolysaccharide and interleukin-4 (LPS+IL-4). From two independent experiments (268 million aligned reads) we identified 90,015 high-confidence DNaseI domains in B cells. As expected, DNaseI-Seq profiles were highly reproducible between biological replicates (Spearman’s ρ = 0.99, Figure S1A).

To identify DHS sites associated with gene regulatory domains, we next mapped Nipbl, Med12, and p300 by ChIP-Seq (Figure 1A). We chose the cohesin-loading factor Nipbl and the Med12 component of mediator because they demarcate enhancers tethered with core promoters (Kagey et al., 2010). Likewise, the transcription regulator p300 occupies a subset of active promoters and enhancers (Chen et al., 2008). There was a considerable, although incomplete, overlap in the recruitment of these factors in B cells (Figure 1A, Figure S1B). Active promoters were thus identified as Nipbl+, Med12+, or p300+ DHS sites that overlapped with ENSEMBL-annotated transcription start sites (TSSs). Based on this strategy we identified 17,004 DHS promoter elements associated with 16,931 genes, 49,763 enhancer elements, and an additional 23,248 that did not overlap with either (Figure 1B). Promoters were in general H3K4me1 low H3K4me3 high H2AZ high, whereas the opposite signature demarcated enhancer domains (Figure 1A, (Kouzine et al., 2013)). DHS analysis of mouse ES cells uncovered a similar number of DHS promoters (16,771) and an increased number of enhancers (62,766, Figure 1B).

To directly map the promoter-enhancer interactome, we applied chromatin interaction analysis by paired-end-tag sequencing (ChIA-PET, (Fullwood et al., 2009; Zhang et al., 2012)), which combines PolII ChIP with 3C technology (Figure 1C, Figure S2A). We generated two independent B cell ChIA-PET libraries, from which ~15 million reads were uniquely aligned and classified into two separate datasets: 5.7 million reads of PolII chromatin occupancy, and 9.2 million reads clustered into 14,247 high-confidence PolII
long-range cis interactions or PETs (Figure 1D, and Table S1). Both datasets were correlated between replicates (Spearman’s ρ > 0.83, Figure S2B–C).

Attesting to the specificity of ChIA-PET, most PolII long-range interactions (13,070, 92%) were linked to at least one gene regulatory domain (Figure S1C). Furthermore, of 16,931 B cell promoters associated with DHS domains, 6,890 were involved in PolII long-range interactions. In general, these genes were transcribed 2-fold higher (P < 2e-16, Figure S1D) and recruited more PolII (P < 2e-16, Figure S1E) than non-anchored ones. We also detected 6,813 DHS enhancer domains involved in PolII interactions. Of these, 71% were active (H3K27Ac+), whereas up to 60% of non-anchored ones were poised (H3K27Ac−, Figure S1F). In general, the number of ChIA-PET interactions per regulatory site was proportional to the extent of DNaseI digestion (Figure S1G). Thus, ChIA-PET preferentially detects PolII long-range interactions involving H3K27Ac+ enhancers and transcriptionally active promoters.

As previously shown (Li et al., 2012), PolII interactions fell into four distinct groups: i) intragenic, connecting promoters to gene bodies; ii) extragenic, connecting promoters to distal regulatory elements; iii) intergenic, tethering promoters from different genes; and iv) enhancer-enhancer interactions (Figure S1H). Examples of these are provided in Figure 1D for the Atp5j-Mir155 gene locus. Consistent with high expression of Mir155 in activated B cells (Kuchen et al., 2010), its promoter was associated with 83 long-range interaction tags (Figure 1D, upper panel). Of these, 70 were extragenic, involving 5 upstream enhancer domains, while 13 were intragenic, connecting the promoter to downstream sequences. An additional 23 PolII long-range interactions interconnected the 5 enhancers upstream of Mir155. In contrast to B cells, ES cells actively transcribe Mrpl39, Jam2, and Atp5j but express little Mir155 mRNA (Figure 1D, lower panel). Consistent with this, we identified 30 intergenic connections between Mrpl39, Jam2, and Atp5j promoters in ES cells, whereas few connections involved Mir155 (Figure 1D). As in previous ChIA-PET studies, both direct and indirect interactions were considered in our analysis (Figure S1I).

**TALEN-mediated validation of promoter-enhancer connectivity**

ChIA-PET confirmed established connections between genes regulatory domains. For instance, the pluripotent gene Sox2 was associated in ES cells with a series of enhancers recently described by 5C studies (Figure S3A, (Phillips-Cremins et al., 2013)). Likewise, the immunoglobulin heavy chain (Igh) 3′Eα enhancer was found in spatial proximity to transcribed Igμ, Igγ1, and Igε in B cells only (Figure S3B). We also found evidence of Igγ-Igα associations, representing either synapses between the recombining genes (Wuerffel et al., 2007), or fully recombined DNA. At the Igκ locus, the VJκ-5′Eκ domain was connected to the previously characterized 3′Eκ and Ed enhancers (Figure 2A, (Liu et al., 2002; Meyer and Neuberger, 1989)). Unexpectedly, the analysis uncovered two additional enhancers located 8kb (E4) and 15kb (E5) downstream of Ed (Figure 2A).

We found additional enhancers (E1–E2) associated with the activation induced deaminase (AID) gene Aicda (Figure 2B). The three enhancers previously shown to regulate AID transcription in vivo were also linked by PolII long-range interactions in the analysis (E3–E5, Figure 2B, (Crouch et al., 2007; Huong le et al., 2013; Sayegh et al., 2003)).
Apobec1 promoter and a sixth enhancer located in Apobec1 intron 2 were also clustered (Figure 2B). To validate ChIA-PET associations, we deleted E1 and E2 in CH12 mouse lymphoma cells. We chose this B cell line because upon activation it transcribes high levels of AID and undergoes efficient Igμ-Igα recombination (Nakamura et al., 1996). To facilitate homozygous gene targeting, knockout constructs were co-transfected with enhancer-specific transcription activator-like effector nucleases (TALENs), assembled via a solid-phase high-throughput system ((Reyon et al., 2012), Figure S4A). Upon activation, wild type CH12 cells increased AID mRNA expression ~5 fold and recombined to IgA (15%, Figure 2C and 2D). Deletion of E1 or E2 however markedly reduced AID transcription and IgA expression (Figure 2C and 2D), consistent with the notion that the extent of switching is proportional to AID expression (Takizawa et al., 2008). Transcription of Apobec1 was also impaired in the mutant cells, whereas noninteracting Ezh2 and Cd83 genes were unaffected (Figure 2C).

Importantly, E1−/− and E2−/− cells displayed an overall reduction in Nipbl and PolII occupancy at all regulatory domains within the Aicda locus, including the Apobec1 promoter (Figure 2E and S4B). In contrast, this effect was not observed at the Foxj2-Necap1 locus, ~190kb downstream of Apobec1 (Figure 2E and S4B). Thus, E1 and E2 regulate AID and Apobec1 transcription by controlling local recruitment of PolII.

To further validate the ChIA-PET results we targeted additional regulatory elements associated with the Pou2af1 (OCA-B), and Cd79a genes. We uncovered an intronic enhancer (E3) ~15kb downstream of OCA-B TSS required for transcriptional upregulation upon B cell activation, but dispensable for basal transcription in non-stimulated cells (Figure S4C). This activity is consistent with the reported dynamics and signaling requirements of OCA-B expression during B cell differentiation (Casellas et al., 2002; Qin et al., 1998). A similar analysis confirmed the presence of enhancer elements that augment basal Cd79a transcription (Figure S4D). Additional gene targeting experiments within the Pim1 oncogene locus are discussed below (Figure 5). Taken together, these results demonstrate that at least a fraction of PolII long-range interactions, as defined by ChIA-PET, represent functional promoter-enhancer connections in B lymphocytes.

**Single- and higher-order gene clusters in primary mouse cells**

Up to 54% of genes recruiting PolIII in activated B cells were associated with long-range interactions (6,890 of 12,652). Of these, 1,231 (18%) represented single promoters tethered to at least one enhancer (Figure 3A). These clusters created complex architectures and spanned an average of 78kb of genomic DNA (Figure S5A). The most elaborate of this group was the Gpr183 promoter, which was connected either directly or indirectly to 12 enhancers via 76 long-range interactions (Figure 3A). Another example was Cd83, which displayed 158 PolIII intragenic and extragenic connections (Figure 3B).

Among single promoter gene clusters we found examples of the recently dubbed super-enhancer domains (Whyte et al., 2013), which were defined based on clustering of gene regulatory domains: e.g. mir290/295 and Sox2 loci (Figure S5B and S3A). However, the vast majority of anchored genes (5,606, 81%) formed higher-order multigene complexes (1,481 B cell clusters, Figure 3C), which could not be easily deduced as interacting based on visual inspection of DHS island distribution. The average span of these clusters was 179kb.
(Figure S5A). Its prime example in B cells was the *Rela* cluster in chromosome 19, which was composed of 66 genes and 398 long-range interactions (Figure 3C). Promoters linked by intergenic connections displayed higher PolII density and mRNA synthesis relative to genes from single-promoter clusters or not anchored to other domains (Figure S5C). Furthermore, families of genes co-expressed during ontogeny were overrepresented in the multiple-promoter gene group (see Methods). Among these we found the major histocompatibility complex *H2-Mβ* cluster (Fisher exact test $P = 1.3e-14$), the *Hist1h* histone family ($P = 7.7e-82$), and the lymphoid signaling *Gimap* cluster ($P = 7.6e-14$, Figure S5D).

In contrast to promoters, which readily formed higher-order complexes, the vast majority of enhancers (~90%) were linked to a single promoter, and less than 2% of all enhancers were linked to more than 2 promoters (Table S1). One exception was an enhancer downstream of *Gmap6*, which was directly linked to 7 promoters (Figure S5D).

**Transcriptional correlation between IncRNA and associated coding genes**

Long non-coding RNAs (IncRNA) are a new class of RNAs believed to play regulatory functions (Batista and Chang, 2013). ChIA-PET identified hundreds of associations between protein-coding and IncRNA genes in multiple-promoter clusters. For instance, IncRNA E*E*ENSMUSG85930 is extensively associated with *Clec2d* and to a lesser extent to *Cd69* (Figure 3D). Other examples involving genes key for B cell development included *Ptprc*-*cap-E90702*, *Cd81*-*E59277*, and *Bcl11a*-*E123592* pairs (Figure S5E). IncRNAs are believed to modulate transcription of neighboring genes by promoting local topological changes in chromatin (Ponting et al., 2009). Consequently, transcription of IncRNAs and their targeted genes is often coordinated (Guil and Esteller, 2012). To test this idea across the genome we measured expression of IncRNAs and their interacting protein-coding genes as defined by ChIA-PET. We found that genes associated with highly abundant IncRNAs were transcribed at higher levels than those associated with IncRNAs detected at low or trace levels ($P < 0.05$, Figure 3E). These findings are consistent with the proposal that transcription of IncRNAs and their targets can be coordinated. Whether the same scenario applies to promoter-promoter clusters not involving IncRNA remains to be determined.

**Broadly expressed genes are linked to cell-type specific enhancers**

As expected, genes differentially expressed in B and ES cells were linked to tissue-specific regulatory elements. The pluripotent gene *Sox2* for instance was associated with ES cell-specific enhancers (Figure S3A). Conversely, the B cell specific *Cd79b* gene was only anchored to enhancers in the B cell compartment (Figure S6A).

We next turned our attention to genes transcribed in both cell types. Of 6,890 promoters anchored by ChIA-PET in B cells, 4,854 (70%) were also anchored in ES cells (Figure 4A, Venn diagram). As an example, the *Hexim1-2* genes were linked to the same downstream enhancer (E1) in B and ES cells (Figure 4B). Surprisingly, most anchored promoters in the two cell types (4,430, 94%) were associated with at least one additional tissue-specific enhancer (Figure 4A, lower pie chart). A striking example was the *Myc* proto-oncogene, which displayed a completely different enhancer landscape in ES and B cells (Figure 4C). In B cells *Myc* was linked to 10 enhancers (E5–E14) located near or downstream of exon 3 of
the lncRNA Pvt1, whereas in ES cells all enhancers associated with Myc (E1–E4) were found upstream of this site (Figure 4C). Other examples included Tgf1, Smad7, and Malat1, which were preferentially linked to ES cell-specific enhancers, whereas Swap70, Env5, and Pim1 were tethered to a greater number of enhancers in B lymphocytes (Figure 4D).

To explore whether changes in the enhancer landscape impacts transcription of this gene group, we measured their expression by calculating mRNA copy numbers per cell. Genes that turned on or off a single tissue-specific enhancer displayed little or no changes in transcription levels in the two cell types (Figure 4E). However, as genes interacted with 2 or more additional enhancers their expression was significantly different ($P < 9e-10$, Figure 4E). Myc for instance was transcribed ~4 times higher in B cells than in ES cells (Table S2). This observation is consistent with the notion that, in general, transcription levels of a give promoter are commensurate with the number of regulatory domains it is regulated by (Li et al., 2012). On the basis of these findings we conclude that i) broadly-expressed genes can be regulated by cell type-specific enhancers and ii) the turning on and off of enhancers during ontogeny impacts transcription levels.

**Dynamic CpG methylation of cell-type specific regulatory domains**

Cellular differentiation is accompanied by changes in DNA methylation at promoters and distal regulatory domains (Shen et al., 2013; Song et al., 2013; Stadler et al., 2011; Ziller et al., 2013). To explore whether the dynamics of DNA methylation correlate with differential enhancer usage we applied bisulphite sequencing (Bis-Seq) and generated methylome libraries at single nucleotide resolution. Bis-Seq of activated B cells provided a total of 148 billion mappable methylome bases. We complemented this dataset with Bis-Seq libraries from mouse ES cells (Stadler et al., 2011) and calculated the percentage of CpG methylation at gene regulatory domains. With few exceptions, activated B cell-specific enhancers were highly methylated in ES cells (>80% of CpGs), whereas enhancers common to both cell types displayed a broad range of CpG methylation levels (Figure 5A). High methylation was also observed at ES cell-specific enhancers in activated lymphocytes whereas B cell enhancers displayed low methylation levels (Figure 5A). Importantly, the level of CpG methylation was lower at active than at poised enhancers ($P < 3e-16$, Figure 5A). As expected, promoters of silent genes displayed on average higher CpG methylation than active ones ($P < 2.2e-16$, Figure 5A). Methylation levels were also inversely proportional to the extent of ChIA-PET signals (Figure S6B). Thus, enhancers are highly methylated when inactive, but become demethylated during development concomitant with the presence of tissue-specific PolII interactions.

We explore in Figure 5B transcriptional regulation of the Pim1 oncogene, whose promoter is tethered to an entirely different set of enhancers in activated B cells and ES cells. The analysis shows a direct correlation between CpG demethylation and enhancer usage. For instance, B cell enhancers E2 and E5 and ES cell enhancers E6 and E7 display nearly complete CpG demethylation in a cell type specific manner (Figure 5B). To confirm that these enhancers truly promote Pim1 transcription, targeted E2 and E6 in CH12 B cells and ES cells respectively. As measured by qPCR, we found a significant decrease in Pim1 mRNA levels in the targeted cells, whereas expression of Brd2, Mtx1, and Gapdh was

*Cell. Author manuscript; available in PMC 2014 December 19.*
unaffected (Figure 5C). Attempts to delete E2 and E6 in the cell type where they are inactive were unsuccessful (not shown), likely due to the inability of TALENs to target methylated DNA (Bultmann et al., 2012).

In ES cells, B cell-specific enhancers, including those linked to broadly expressed genes, were hypermethylated (Figure S6C). To examine at which stage during B lymphopoiesis these regulatory elements become demethylated, we generated methylome libraries from bone marrow hematopoietic stem cells (KSL), B lymphoid precursors (CLP), and peripheral G0 resting B cells. Of 1,518 B cell-specific enhancers that were linked by long-range interactions and hypomethylated during activation, only 82 (5%) were also hypomethylated in ES cells (Figure 5D). However, in KSL precursors nearly half (714, 47%) of activated B cell enhancers were already demethylated (Figure 5D). This group included Pim1 enhancers E2, E3, and E5, which displayed nearly identical CpG methylation levels in KSL, CLP, G0 resting, and cycling B cells (Figure S6D). As KSLs develop into CLP B cell precursors, demethylation was observed in 61% (926) of B cell enhancers. In resting G0 B cells, this number increased to 80% (1,207, Figure 5D). At this stage of development, the overall mean methylation was comparable between activated B cell enhancers and those functional both in B and ES cells (Figure S6C). Thus, most activated B cell enhancers, including those associated with broadly expressed genes, are demethylated by the time naive lymphocytes migrate from the bone marrow to the periphery. This finding is consistent with the notion that the genome of G0 lymphocytes is primed for activation, and that most genes expressed during the humoral immune response are transcribed at basal levels in the naive compartment (Kouzine et al., 2013; Nie et al., 2012). At the same time, it is important to point out that ~20% of activated B cell enhancers do not become fully demethylated until activation occurs. Among these we find Pim1 E4 and Aicda E3 and E4 (Figure S6D and S6E).

**Digital genomic footprinting characterizes TF binding in the mouse genome**

The observation that cell-type specific enhancers can promote transcription of broadly expressed genes implies that factors driving lineage specification are involved in this regulation. To explore this idea we sought to comprehensively catalogue transcription factor occupancy in mouse B lymphocytes and ES cells. To this end, we took advantage of the fact that transcription factors protect their binding sites from DNaseI cleavage, leaving nucleotide-resolution footprints within DHS islands (Neph et al., 2012). Figure 6A for instance shows four DNaseI footprints at the Pold4 gene promoter in G0 and cycling B cells. Importantly, these footprints overlap with recognizable binding motifs for transcription factors PU.1, Ebf1, Egr1, and Sp1 (Figure 6A). By applying an established algorithm (Baek et al., 2012), we detected 706,669 high-confidence (FDR <5%) footprints within 75,917 B cell DHS domains (70% of total DHS). To link these footprints to known transcription factor recognition sequences, we examined all empirically defined DNA binding motifs, compiled by HOMER, UniPROBE, JASPAR, and similar databases. We found a significant enrichment of transcription factor binding motifs within DHS footprints ($P < 1e-06$, Figure S7A). Altogether, we linked 247 distinct transcription factor DNA motifs to 122,505 footprints in B cells (Table S1). In addition, *de novo* motif discovery yielded 18 new binding
sites that did not match known recognition sequences (Table S1). A similar analysis linked 306 DNA motifs to 346,284 footprints in ES cells (Table S1).

Figure 6B shows examples of cleavage profiles for transcription factors Irf8, Sp1, Nrf1, PU.1, and CTCF (an extended view of footprints is provided in Figure S7B). Importantly, ChIP-Seq analysis showed a correlation between PU.1 and CTCF occupancy, their DNA binding motifs, and cognate footprints (Figure 6C). Similar results were obtained with Irf8 and Ebf1 (not shown). To confirm that footprinting profiles truly reflect the morphology of DNA-protein interactions, we turned to available transcription factor:DNA cocrystal structures. For instance, consistent with the published Ebf1:DNA structure (Treiber et al., 2010), backbone phosphates in direct contact with Ebf1 amino acids via hydrogen bounds (in red in Figure 6D) were the most protected from DNaseI cleavage, whereas more exposed residues (in cyan) displayed by comparison greater susceptibility to digestion. Similarly, for the cRel (NF-κB) factor DNaseI cleavage paralleled the topology of the protein-DNA interphase with a marked depression at DNA binding motif (Figure 6E, (Huang et al., 2001)). We conclude that DNaseI cleavage profiles can recapitulate the structural features of transcription factor:DNA interactions and thus they reflect the occupancy of gene regulatory proteins across the genome.

**Lineage-determining factors regulate transcription of broadly expressed genes**

Having validated the genomic footprinting approach we next turned to the question of whether lineage-specific factors associate with enhancers controlling transcription of broadly expressed genes. To this end we classified B and ES cells enhancers into three groups: i) those shared between the two cell types and linked to the same promoter as determined by ChIA-PET, ii) cell-type specific enhancers bound to promoters active both in B and ES cells, and iii) cell-type specific enhancers linked to cell type-specific promoters (Figure 7A). Within the B cell compartment, all transcription factors analyzed were associated with the three enhancer groups. Ebf1, Oct2, and E2A footprints for instance showed no significant differences in their distribution regardless of enhancer specificity (Figure 7B). Other factors, such as E2f2 and Foxo1 displayed statistical significant biases for shared and cell type-specific enhancers respectively (Figure 7B). However, even in these cases footprints were not excluded from any enhancer group. A similar distribution was observed in ES cells (Figure S7C). Thus, lineage specification factors associate both with tissue-specific and broadly active enhancers. Confirming this finding, ChIP-Seq analysis of the pluripotent factors Nanog, Oct4, and Sox2 showed occupancy of B and ES cell shared enhancers at the *Ube2g1* locus as well as binding to ES cell specific enhancers at the *Myc* locus (Figure 7C). A global analysis of Irf4, Irf8, Pu.1, Ebf1, and Stat6 ChIP-Seq from B cells also corroborated the results (not shown). We conclude that transcription factors driving lineage specification associate with regulatory elements of broadly-expressed genes. We point out that the functional significance of TF recruitment as defined by digital footprinting remains to be empirically determined. At the same time, our findings are consistent with the notion that broadly-expressed genes can be regulated in a tissue-specific manner.

*Cell. Author manuscript; available in PMC 2014 December 19.*
We have here characterized the first interactomes of gene regulatory domains in primary cells. The data provide a wealth of information from the mouse genome with thousands of promoter-enhancer pairs. Even for loci that have been extensively studied, the analysis uncovered new connectivities. At the Igκ locus for instance we identified novel enhancers linked to 5′Eκ, the regulatory domain from which NF-κB was originally isolated 27 years ago (Sen and Baltimore, 1986). ChIA-PET also uncovered three additional enhancers tethered at a long distance (up to 50kb) to the AID gene promoter; the previously characterized AID enhancers are all located within 15kb of the TSS. Thus, one clear advantage of ChIA-PET lies in its ability to identify long-range interactions, even when they leapfrog non-interacting genes, as is the case for 65% of all enhancers (Table S1). A striking example is a pair of giant enhancers linked to Pax5 by skipping over ~250Kb of DNA containing the Zcchc7 gene (Figure S7D).

Like other conformation capturing techniques, ChIA-PET does not directly address functionality of chromatin interactions. This can only be determined empirically by other means. Typically, enhancer activity is defined by luciferase-based plasmids or LacZ transgenes. However, these experimental approaches only provides an incomplete view of transcriptional regulation because they either lack proper chromatin structure or the influence of neighboring enhancers, insulators, and silencers is not taken into account. Conversely, genome editing provides a means to measure the impact of enhancer deletion in the physiological context. In most cases, we found that cognate promoter activity was partially reduced following enhancer ablation, supporting the model that the contribution of individual enhancers to gene expression is additive in nature. Examples of this category were enhancers linked to the Cd79a, Pou2af1, and Pim1. On the other hand, the targeting of AID 5′ enhancers E1 or E2 nearly entirely abolished AID expression and activity, a result that is reminiscent of those obtained upon deletion of AID enhancers E3 and E5 in BAC transgenic mice (Crouch et al., 2007; Huong le et al., 2013), or by interference with E4 activity in primary B cells (Sayegh et al., 2003). Thus, rather than working in additive fashion, AID gene regulatory elements seem to synergize or act as a cooperative unit. Conceivably, the local topology of the Aicda-Apobec1 locus requires activation of all enhancers for optimal transcription to occur. This strategy may be useful for genes that require tight regulation as AID, whose expression is strictly limited to activated B cells to minimize its well known tumorigenic activity (Casellas et al., 2009). Consistent with this model, our methylome analysis indicates that the AID locus is not completely demethylated until B cells are activated.

A direct comparison between the B and ES cell interactomes revealed that up to 95% of genes anchored in both cell types are associated with at least one tissue-specific enhancer. Transcription modularity, the mechanism whereby genes accumulate regulatory elements during ontogeny, is a well-described phenomenon that controls the spatiotemporal expression of developmental genes (Davidson, 2001; Visel et al., 2007). For instance, the cardiac homebox gene Nkx2-5 is targeted to specific sub-regions of the developing heart by turning on additional cis-regulatory domains over time (Schwartz and Olson, 1999). Similarly, expression of the human apolipoprotein E gene is triggered in hepatocytes and...
astrocytes by enhancers only active in those tissues (Grehan et al., 2001). Our studies demonstrate that the turning on and off of enhancers is not a singularity of developmental gene loci. Instead, it is a widespread mechanism that regulates broadly expressed genes involved in basic cellular functions, such as cell cycle initiation (Myc), signal transduction (β-catenin), and cellular motility (Malat1).

Mechanistically, we show that the changing enhancer landscape in mammalian development results from the unbiased recruitment of lineage-determining factors, which associate with enhancers anchored not only to tissue-specific promoters but also to constitutively active ones. Based on these observations we propose a “relay race” model of transcriptional regulation, whereby broadly active genes make use of tissue-specific cis-regulatory elements and transcription factors as cells progress through development (Figure 7D). For genes that only replace a subset of their regulatory domains, transcription is roughly maintained at comparable levels in different cell types. However, as the number of connected enhancers fluctuates considerably, promoter activity can be significantly altered.

We can think of at least two reasons why higher organisms modulate the enhancer landscape of broadly expressed genes. First, as aforementioned, it enables fine-tuning of protein output, which in turns controls protein activity. Second, it places basic cellular functions under the control of tissue-specific factors. In the B cell compartment, these strategies are perhaps best illustrated during the immune response to invading pathogens. In this microenvironment B lymphocytes move rapidly from a G0, quiescent state to one of the fastest proliferative rates among eukaryotic cells (Liu et al., 1991). Key in this process is Myc which, along with TFIIH, triggers a ~10-fold amplification of the lymphocyte transcriptome concomitant with cell cycle entry (Kouzine et al., 2013; Nie et al., 2012). Thus, vis-à-vis proliferation B cells differ substantially from continuously dividing ES cells, in that they must rapidly engage a burst of Myc expression and activity during the immune response to cope with fast dividing pathogens. Our studies imply that this unique response is mediated, at least in part, by the large number of cis- and trans-responsive elements that associate with the Myc promoter in the B cell compartment.

**EXPERIMENTAL PROCEDURES**

**Cell Isolation and Culture**

Hematopoietic stem cells were isolated from the bone marrow of 25 six-week old C57BL6 mice (Jackson Laboratory). Cells were purified following Ema et al.’s protocol (PMID: 17406558). KSL (KIT+, SCA1+, Lin−, IL-7R−) and CLP (KIT+, SCA1+, Lin−, IL-7R+) cells were sorted using MoFlo Legacy (Beckman Coulter) and BD FACSaria III. Resting splenic B cells were isolated from 6–8 week-old wild-type C57BL6/J mice with anti-CD43 Microbeads (anti-Ly48; Miltenyi Biotech) and were activated for 48–60 h with LPS (50 μg/ml; Sigma), IL-4 (5 ng/ml; Sigma) and 0.5 μg/ml of anti-CD180 (RP105) antibody (RP/14, BD Pharmingen). E14 tg2A mouse embryonic stem cells were maintained as described in (PMID:18555785). Switchable IgM+→IgA+ murine CH12-F3 Ly-1+ B cell lymphoma line was maintained and passaged every 2 days in RPMI 1640 supplemented with 10% FBS (ATCC), 1% penicillin/streptomycin (Invitrogen), 55 uM 2-β mercaptoethanol (Invitrogen). All cells were maintained at 37°C and 5% CO2 in a humidified incubator.
Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

We thank Kefei Yu for CH12 cells and protocols; J. Simone and J. Lay for cell sorting; G. Gutierrez for technical assistance with Illumina analyzer. This work was supported by the Intramural Research Program of NIAMS and NCI, and internal Jackson Laboratory fund JAX19020120 to Y.R.. J.K.J. was supported by NIH grants DP1 GM105378 and P50 HG005550, the Defense Advanced Research Projects Agency grant W911NF-11-2-0056 and The Jim and Ann Orr Massachusetts General Hospital Research Scholar Award. J.Q. was supported by the NIH UGSP program. All animal experiments were performed according to NIH guidelines. High-performance computation was performed using NIH Helix Systems (http://helix.nih.gov).

References


Meyer KB, Neuberger MS. The immunoglobulin kappa locus contains a second, stronger B-cell-specific enhancer which is located downstream of the constant region. EMBO J. 1989; 8:1959–1964. [PubMed: 25073121]


ChIA-PET is used to characterize the regulome of primary mouse cells. Broadly-expressed genes change their enhancer landscape during development. The dynamics of CpG methylation mirrors changes in the enhancer landscape. Most transcription factors bind to cell-specific and constitutively active enhancers.
Figure 1. Characterization of regulatory domains and their interactions in mouse primary B and ES cells

(A) Pax5 locus in activated B cells displaying DNaseI hypersensitivity (DHS); recruitment of Nipbl, Med12, and p300; and chromatin marks H2AZ, H3K4me1, and H3K4me3. (B) Bar graphs showing the number of DHS islands in B and ES cells overlapping with promoters (TSS+, white), enhancers (TSS−, Nipbl+, or Med12+, or p300+, red), or non-overlapping (blue). (C) The ChIA-PET protocol combines PolII ChIP with conformation capturing techniques to map the interaction of active promoters with gene regulatory domains. (D) Examples of ChIA-PET clusters at the Mir155 locus in activated B cells (red connectors) or ES cells (blue connectors). Each connector links 2 or more long-range interactions (PETs) separated by <500 bps (Figure S2A). ChIP-Seq data are represented as reads per Kb per million sequences (RPKM). Promoters (P) and enhancers (E) are boxed and the number of total PETs is provided in parenthesis. Interactions between enhancers and Mir155 are represented by semi-circle connectors. mRNA expression is provided for B and ES cells as RPKM values (+ strand transcription in green, − strand in blue).
Figure 2. In vivo validation of ChIA-PET by genome editing

(A) ChIA-PET at the Igκ locus identifies previously characterized 5′Ex, 3′Ex, and Ed enhancers, as well as new enhancers E4 and E5. Number of PETs associated with each regulatory domain (boxed) are provided in parenthesis. The DHS activated B cell track is also provided (black). (B) Regulatory map of the Aicda-Apobec1 locus in activated B cells. Deletion of selected enhancers (E1 and E2) was carried out in CH12 B cells using knockout targeting cassettes (cyan) and TALEN endonucleases. (C) qPCR analysis of Aicda, Apobec1, Ezh2, and Cd83 expression in wild type (WT), and E1 or E2 deleted (Δ) CH12 cells. Data are represented as the mean +/− SEM (n = 6). P values were < 0.0001 (Aicda), and = 0.008 (Apobec1). (D) Flow cytometry analysis of recombination to IgA in activated WT, ΔE1, or ΔE2 cells. (E) Nipbl (black) and PolII (red) occupancy at the Aicda-Apobec1 and Foxj2-Necap1 loci in WT or ΔE2 cells. The two loci are separated on chromosome 6 by 188kb.
Figure 3. Gene clusters identified by ChIA-PET

(A) Single-promoter clusters in activated B cells connecting 1,231 gene promoters to at least one enhancer. Right: e.g. the Gpr183 promoter (blue circle) is linked to 12 enhancers (red circles) via 76 interactions. PETs anchored outside enhancers are represented with grey circles. Circles are sized according to the absolute number of anchored PETs. (B) Interactions at the Cdl83 single gene cluster. (C) Multi-promoter clusters (n = 1,481) identified in B cells. The Rela cluster display 398 interactions involving 66 genes. (D) PolII connections between lncRNA E85930, Clec2d, and Cdl69. Promoters and enhancers are boxed and number of PETs are provided in parenthesis. (E) Transcription levels of genes
associated with trace (no detectable FPKM), low (<0.9 FPKM), or highly transcribed IncRNAs (≥0.9 FPKM).
Figure 4. Tissue-specific enhancers contribute to the transcriptional regulation of broadly expressed genes

(A) Venn diagram showing the number of ChIA-PET anchored promoters in B cells (left), ES cells (right), or in both cell types (middle). For the latter group, the pie chart below shows the number of promoters linked to the same (yellow) or to at least one cell-type specific enhancer (grey). (B) Acbd4 and Hexim1/2 gene promoters associate with the same downstream enhancer (E1) in B and ES cells. (C) The Myc oncogene is linked to an entirely different set of enhancers in ES (blue) and B (red) cells. Enhancers are numbered from 1–10 based on proximity to the Myc promoter (P1). (D) Genes classified by the total number of tissue specific enhancers they interact with. Individual examples are highlighted and the number of B cell-specific (red) and ES cell-specific (blue) enhancers is provided in parenthesis. (E) Box plot providing changes in mRNA copy number for genes that are anchored both in B and ES cells and that associate with ≤2 ES cell-specific enhancers (blue), ≥2 B cell-specific enhancers (red), or genes in between (grey).
Figure 5. DNA demethylation demarcates differential enhancer usage

(A) CpG methylation at regulatory elements in B and ES cells. (B) DNA demethylation at the Pim1 mouse locus in B (upper) and ES (lower) cells. Demethylation was calculated by subtracting normalized ES cell methylation values from the activated B cell Bis-Seq and vice versa. (C) Bar graph showing qPCR analysis of Brd2, Mtc1, and Pim1 expression in WT (black) or ΔE2 (red) CH12 cells. Same analysis for WT or ΔE6 (blue) ES cells. Data are represented as the mean ± SEM (n = 6). Pim1 P < 0.005 in B cells and P < 0.003 in ES cells. (D) Bar graph represents the fraction (%) of 1,518 activated B cell enhancers that are CpG demethylated in ES, KSL, CLP, and G0 resting (r) B cells. Absolute numbers of demethylated enhancers are provided on top of each bar.
Figure 6. Digital genomic footprinting

(A) Characterization of footprints at the Pold4 promoter in primary resting or cycling CH12 B cells. Transcription factor binding motifs overlapping with each footprint are shown below the graph (red rectangles). Tracks were configured to display the maximum (light grey) and one standard deviation above the mean (dark grey). (B) Examples of footprints overlapping with Irf8, Sp1, Nrf1, PU.1, and CTCF DNA recognition motifs. (C) Composite of PU.1 and CTCF ChIP-Seq (blue data points, upper graphs) and cumulative footprinting (lower graphs) associated with cognate binding motifs (middle logos) in B cells. The absolute number of motif occurrences is provided. Grey data points represent ChIP-seq signals at footprints not associated with PU.1 or CTCF motifs. (D) The co-crystal structure of Ebf1 bound to its DNA ligand is compared to its cognate footprint profile. Motif nucleotides least sensitive to cleavage are depicted in red; most sensitive residues are depicted in cyan. (E) Similar analysis as in panel D for the NFκB-cRel dimer.
Figure 7. Distribution of transcription factor binding across the B and ES cell genome

(A) Classification of B and ES cell enhancers into those present in both cell types and associated with shared promoters (light blue), cell-type specific enhancers associated with broadly active promoters (grey), and cell-specific enhancers associated with cell-specific promoters (blue). (B) Fraction of each enhancer type harboring a particular motif in a footprint. (C) ChIP-Seq analysis of pluripotent factors Nanog, Oct4, and Sox2 at the Ube2g1 and Myc gene loci. (D) Transcription of broadly-expressed genes is driven during development by a relay-race type of regulation, where the landscape of active enhancers varies in different cell types concomitant with the dynamics of CpG methylation.