# Functional and topological characteristics of mammalian regulatory domains

Orsolya Symmons,<sup>1</sup> Veli Vural Uslu,<sup>1</sup> Taro Tsujimura,<sup>1</sup> Sandra Ruf,<sup>1</sup> Sonya Nassari,<sup>1</sup> Wibke Schwarzer,<sup>1</sup> Laurence Ettwiller,<sup>2,3</sup> and François Spitz<sup>1,4</sup>

<sup>1</sup>Developmental Biology Unit, European Molecular Biology Laboratory, 69117 Heidelberg, Germany; <sup>2</sup>Centre for Organismal Studies, University of Heidelberg, 69111 Heidelberg, Germany

Long-range regulatory interactions play an important role in shaping gene-expression programs. However, the genomic features that organize these activities are still poorly characterized. We conducted a large operational analysis to chart the distribution of gene regulatory activities along the mouse genome, using hundreds of insertions of a regulatory sensor. We found that enhancers distribute their activities along broad regions and not in a gene-centric manner, defining large regulatory domains. Remarkably, these domains correlate strongly with the recently described TADs, which partition the genome into distinct self-interacting blocks. Different features, including specific repeats and CTCF-binding sites, correlate with the transition zones separating regulatory domains, and may help to further organize promiscuously distributed regulatory influences within large domains. These findings support a model of genomic organization where TADs confine regulatory activities to specific but large regulatory domains, contributing to the establishment of specific gene expression profiles.

#### [Supplemental material is available for this article.]

Specificity of gene expression is key to tissue function and identity, and is in great part determined at the transcriptional level. Promoters, located proximally to transcriptional start sites, play an essential role in initiating gene expression, but their activity greatly depends on the action of more distal regulatory elements. Among these, enhancers remain the best characterized (Bulger and Groudine 2011; Ong and Corces 2011). They often show a specific chromatin signature, which has helped the annotation of enhancers active in human or mouse cell lines and tissues (Heintzman et al. 2009; Creyghton et al. 2010; Rada-Iglesias et al. 2011; The ENCODE Project Consortium 2012; Shen et al. 2012). Remarkably, a substantial fraction of these putative enhancers was found hundreds of kilobases from the nearest gene. Yet, how their activity is allocated to their target gene(s) across such large distances and sometimes several genes, remains unclear.

The evidence to date indicates that enhancer-promoter communication can be influenced by specific regulatory elements, which may contribute to enhancer-promoter interactions positively (promoter architecture [Ohtsuki et al. 1998], tethering elements [Calhoun and Levine 2003]), or negatively (e.g., insulators/ enhancer blockers [for review, see Gaszner and Felsenfeld 2006]). Also, the regulatory interactions between promoters and associated remote enhancers are usually associated with physical proximity (Bulger and Groudine 2011; for review, see Bickmore and van Steensel 2013). Several protein complexes, including CTCF, cohesin, and Mediator, have been proposed to have a role in organizing these regulatory contacts (Parelho et al. 2008; Wendt et al. 2008; Hadjur et al. 2009; Kagey et al. 2010). Furthermore, recent studies suggest that the genome is organized into relatively cell-type invariant topological domains (TADs) characterized by preferential self-contacts (Dixon et al. 2012; Hou et al. 2012; Nora et al. 2012;

# <sup>3</sup>Present address: New England Biolabs, Ipswich, MA 01938, USA <sup>4</sup>Corresponding author E-mail spitz@embl.de

Article published online before print. Article, supplemental material, and publication date are at http://www.genome.org/cgi/doi/10.1101/gr.163519.113. Sexton et al. 2012). Genes located in the same TAD show greater expression correlation than genes located in distinct ones (Nora et al. 2012), and TAD borders are enriched for factors implicated as insulator elements (Dixon et al. 2012; Hou et al. 2012). These observations suggested that TADs may form a backbone for tissuespecific regulatory interactions (Bickmore and van Steensel 2013; Gibcus and Dekker 2013; Nora et al. 2013).

However, despite a growing body of knowledge on individual regulatory elements and the three-dimensional organization of the genome, the "rules of engagement" (Splinter and de Laat 2011) that determine the activity of remote *cis*-acting elements on the surrounding genes remain elusive, in part because a direct assessment of how such activities are distributed in the genomic environment is still lacking. Scrutinizing endogenous gene activity offers only a partial view of this distribution, since expression of endogenous genes also depends on their different promoters, and distinct post-transcriptional regulation. Furthermore, it does not provide information about the gene deserts that constitute ~25% of mammalian genomes and where many functionally important regulatory elements lie (Nobrega et al. 2003; Ovcharenko et al. 2004).

We have recently developed an efficient in vivo transposition system that allows the rapid production of mice with a single-copy insertion of a regulatory sensor (Ruf et al. 2011). This regulatory sensor consists of a *lacZ* reporter gene driven by a weak promoter, and has minimal effect—if any—on the expression of the surrounding genes. The sensor is carried in a *Sleeping Beauty* transposon that integrates almost randomly in the genome (Horie et al. 2003; Liu et al. 2005). Importantly, the sensor is "naïve": It has not been subjected to the evolutionary selection that has shaped

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By analyzing the activity of the regulatory sensor in mouse embryos at several hundred insertion sites, we identified multiple factors that impinge on the action of *cis*-regulatory elements. We confirm circumstantial evidence that the influence of enhancers extends over several hundred kilobases, irrespective of the position of the normal target genes. By comparing adjacent insertions, we identified large regulatory domains as well as transition zones. We find that regulatory domains are included within TADs, and accordingly depleted in features associated with insulating activities, whereas TAD boundaries correspond to regulatory transitions. Thus, the influence of enhancers appears limited to the topological domain they are part of, providing direct support to the notion that TADs correspond to a functional subdivision of the genome into regions where otherwise promiscuous regulatory influences are confined.

## Results

Our initial analysis of about 160 insertions of a regulatory sensor had suggested that tissue-specific regulatory activities are pervasively present throughout the genome (Ruf et al. 2011). From this, we inferred that with an increased number and density of insertions, one could map—in the natural genomic context—the intervals that were responsive to enhancers, and thus investigate the regulatory architecture of the mouse genome in greater detail (Fig. 1A).

For this purpose, we generated and mapped more than 1000 new integration sites using extensive transposition from singlecopy starting points. We characterized a substantial subset of these



**Figure 1.** Mapping the distribution of regulatory activities along the genome. (*A*) Insertion of a regulatory sensor (drawing of sensor) at different distances from an enhancer (blue oval) and different positions relative to its target gene (blue arrow) can report on the domains of action of the enhancer. (*B*) A total of 734 insertions were characterized for expression in mid-gestation mouse embryos. About 55% of these reported regulatory activities. Insertions with (blue) or without (white) expression were broadly distributed around endogenous gene transcriptional start sites (distribution made with GREAT) (McLean et al. 2010). (C) Examples of the diverse expression justerns obtained in E11.5 embryonic forelimb (*left*) or forebrain (*right*). Numbers refer to insertion identifiers used in the TRACER database (Chen et al. 2013).

(747 insertions) for *lacZ* expression (Supplemental Table 1; online TRACER database) (Chen et al. 2013). These insertions are distributed throughout the genome, with a subset clustered in domains of up to 1 Mb, owing to the local hopping of the Sleeping Beauty transposon (Horie et al. 2003). Overall, most insertions are localized far from endogenous transcriptional start sites (TSS) (Fig. 1B), with two-thirds more than 50 kb from the nearest TSS. Consistent with our first analysis (Ruf et al. 2011), ~55% of all insertions showed expression, with the vast majority expressed in a tissue-restricted manner (Symmons and Spitz 2013). Insertions far from TSS are more likely to show expression than the ones next to TSS (Fig. 1B), probably due to local competition with endogenous gene promoters (Ruf et al. 2011). Importantly, independent insertions show extremely diverse expression patterns (Fig. 1C): Only nearby insertions or insertions around paralogous genes yielded expression patterns that were not clearly distinguishable.

# Binding of the enhancer-associated protein EP300 is predictive of tissue-specific expression over large distances

We first sought to obtain deeper insight into the range of action of enhancers. For this, we investigated whether insertions were more likely to show expression in a given tissue if located in the proximity of an active enhancer, and—if so—how far we could detect such enrichment. We compiled a list of 670 nonredundant viewpoints, annotated with a simple controlled vocabulary (Chen et al. 2013; Supplemental Table 2). Since binding of the coactivator protein EP300 has been reported to demarcate active enhancers in a given tissue (Visel et al. 2009), we used EP300 binding sites from E11.5 mouse embryonic forebrain, midbrain, limb, and heart (Visel et al. 2009; Blow et al. 2010). We identified pairs of sensor insertion/EP300 binding sites, and classified them as concordant, when the sensor was expressed in the tissue where the EP300

> binding site was detected. By calculating the enrichment of such concordant pairs compared with a set of random insertion/ EP300 site pairs, we revealed significant enrichment for concordant pairs up to a distance of 200 kb (Fig. 2A; Supplemental Fig. 1). When EP300 binding sites were paired with silent insertions or insertions expressed in different tissues, enrichment was weaker or not found. This analysis implied that enhancers-represented here by EP300-bound regions-exert their effects across large distances. A similar typical range has been observed when comparing the distance between enhancer regions and the TSS of endogenous genes (Blow et al. 2010; Chepelev et al. 2012; Li et al. 2012).

# Enhancer activity is not gene-centric and broadly distributed

To refine this analysis, we made use of enhancers for which transgenic activity had been previously documented in E11.5 embryos, mostly from the Vista Enhancer database (Visel et al. 2007). We carefully compared the spatial activity of these enhancers with that of the regulatory sensor



**Figure 2.** Expression of the regulator sensor is correlated with surrounding enhancers up to large distances. (*A*) Enrichment of insertions showing LacZ activity in a given tissue relative to limb EP300 binding sites. Enrichment for insertions with expression in the limb (green) compared with random insertions is calculated at increasing distances from the nearest EP300 site (*x*-axis). Error bars represent one standard deviation from the mean. Enrichments of insertions with activity in other tissues but not in the limb (heart: purple; forebrain: blue; midbrain: red) or with no LacZ activity (gray) are also displayed. Results for EP300 sites from other tissues are shown in Supplemental Figure 1. (*B*) Comparison of enhancer and sensor activity. Different groups were considered, according to the relative distance between the insertions and the enhancers (number of insertion–enhancer pairs indicated *above* each bar). The two random data sets are described in the Methods section. The proportions of concordant enhancer–insertion pairs in different groups were compared using Fisher's exact test. (*C*-*E*) Examples of concordant enhancer–insertion pairs. The different loci are schematized (enhancer: blue oval with VISTA reference; sensor: drawing of transposon; endogenous genes: arrows or gray bars with black exons), with putative target genes of enhancers indicated by labeling the gene the same color as the enhancer. Photos of representative embryos of the in vivo enhancer assays are from the Vista Enhancer Browser (Visel et al. 2007). (C) The sensor reported the activity of an intronic diencephalon/midbrain enhancer, which likely contributes to the regulation of the distant *Lhx2* gene (Gray et al. 2004). (*D*) Heart-specific expression of the sensor when inserted adjacent to a heart-specific enhancer, possibly regulating the adjacent *Myocd* gene (Wang et al. 2001). The eye expression shown on the representative transgenic embryo (\*) is ectopic. (*E*) The sensor, inserted next to *Znf503* (McClinn et al. 2008)

when inserted in their vicinity (Fig. 2B–E). Given the highly specific patterns of the different insertions and enhancers, this offered much greater confidence in a direct relationship between matching patterns. We defined pairs as "concordant" when the domain of enhancer activity was included in the one shown by the regulatory sensor; as "divergent" or "inactive," when the regulatory sensor was expressed elsewhere or not expressed (Supplemental Table 3). As shown in Figure 2B, all insertions positioned in close proximity (<10 kb) of an enhancer showed expression patterns highly concordant with its autonomous activity. This ratio decreased with increasing distance, but even at 100–200 kb, a third of insertions–enhancer pairs were concordant, compared with <7% for random associations of enhancers and insertions (Fig. 2B) (Fisher's exact test,  $P < 3.305 \times 10^{-6}$ ). Expression of our regulatory sensor was not dependent on its orientation (Supplemental Fig. 2).

We then examined whether enhancers distributed their activities preferentially toward their endogenous target gene. For this purpose, we considered a subset of enhancers for which we could confidently assign a—putative—target gene, using in situ gene expression data from the literature. We categorized these 107 enhancer-gene-insertion triplets (corresponding to 33 genomic loci) based on the relative order of their components, and determined whether the inserted sensor showed complete, partial, or no overlapping expression with the enhancer/gene pair (Fig. 3; Supplemental Table 4). Insertions located between an enhancer and its target gene showed overlap with the reported enhancer activity more frequently than insertions located either opposite to the gene or beyond the transcriptional start site (Fisher's exact test, P =0.001457 and P = 0.04238, respectively). However, about half of the insertions located opposite the target gene or beyond its



**Figure 3.** Non-gene-centric enhancer activities are detected across large distances. (*A*) The number of insertions that correctly (blue) or partially (light blue) reported the activity of a neighboring tissue-specific enhancer, or showed a different activity (orange). Insertions were grouped depending on their position relative to the enhancer/target gene, as shown *below* the chart. (*B–D*) Examples of expression detected with the regulatory sensor (photos) in non-gene-centered situations. Gene (arrows) and enhancer (ovals) activities are color-coded and shown on the embryo outline. (*B*) An insertion between the *En2* and *Cnpy1* genes matches their expression at the mid/hindbrain junction (Jukkola et al. 2006), as well as the activity of an enhancer on the far side of *En2* (see also Supplemental Fig. 8) (*C*) *Barhl2* expression in the midbrain and diencephalon requires remote enhancer (Saba et al. 2005), and a diencephalon enhancer (hs612) (Visel et al. 2007) is present upstream of this gene. Enhancer activity extends to a downstream insertion. (*D*) *Sall1* gene expression is controlled by multiple enhancers spread in the two surrounding gene deserts, and insertions flanking the gene display overlapping expression patterns.

promoter showed an expression pattern corresponding to the associated enhancer (Fig. 3), indicating that enhancer activity is not exclusive to the promoter of their target gene. Furthermore, given the relatively small sample size, it is possible that the weak directionality of enhancer activity that we observed is due to the different distances associated with the different categories.

Overall, these multiple analyses show that enhancers act on their environment broadly and in a largely undiscriminating manner.

#### Extended enhancer activity results in large regulatory domains

Consistent with these findings, we noted that adjacent insertions frequently had similar expression, even when hundreds of kilobases apart. Large domains of co-regulation have been reported before, usually around developmental genes (Spitz et al. 2003; Zuniga et al. 2004; Kikuta et al. 2007), but with our extensive survey of the mouse genome, we could expand this list and refine the extent of such domains. Importantly, since the regulatory sensor is driven by the same promoter at each insertion site, changes in expression are not due to differently responsive promoters, but really outline the limits of enhancer action. We cannot formally exclude that co-expression of the sensor at different posi-

tions results from activation by different enhancers with overlapping activities, like those discovered around developmental genes (Carvajal et al. 2001; Uchikawa et al. 2003; Hong et al. 2008; Marinić et al. 2013; Visel et al. 2013). But individual enhancers usually have distinct spatial activities (Visel et al. 2007) and therefore, the spatially restricted expression patterns that we use to define co-expression provide confidence for genuine co-regulation. Supporting this further, insertions into loci where enhancers have been meticulously mapped allowed us to assess that they respond to the same enhancer(s) at multiple distant positions (e.g., *Shh*, Supplemental Fig. 3; *Hoxd*, Supplemental Fig. 4).

We found 311 chromosomal intervals defined by the presence of two or more insertions within <2 Mb. We assigned these intervals to different categories, depending on the expression patterns of the insertions (Fig. 4A–C; Supplemental Figs. 3–5; Supplemental Table 5). We defined *regulatory domains* (RDs; 46 regions; size from 3.8 kb to 2.1 Mb; median size: 359 kb; total combined length: 22.3 Mb) as the largest possible interval containing multiple insertions with shared expression. This is a conservative definition, and RDs likely extend beyond the insertions that define them. RDs are found on all chromosomes, except possibly Y, where we did not obtain insertions with expression. Within RDs, we occasionally saw quantitative variation or no detectable expression of the sensor at different positions. However, we never observed



**Figure 4.** Extended domains of co-regulation correlate with the subdivision of the genome into topological domains. (*A*–C) Outlines of loci, with genes displayed as arrows and insertions as drawings of the transposon. Regulatory domains and transition zones are labeled, TADs (identified by Hi-C in mouse ES cells) are indicated by green and brown bars and unstructured regions by dashed lines. Hi-C interaction frequencies are represented as a two-dimensional heat map (from Dixon et al. 2012). (*A*) Multiple insertions in the chr3:7.3–8.3M interval outlined an extended regulatory domain characterized by shared expression in the facial and trunk mesenchyme, and in neural crest derivatives. This domain extends into the adjacent unstructured region (insertion 201179e9), but two telomeric insertions, located in a different TAD, showed different patterns (the proximal limb expression of 181912bc-133 is anterior, whereas insertions in the flanking RD have a more medial expression), defining two transition zones. (*B*) Multiple insertions in the ear (\*) is due to another insertion also present in 177175-emb7. The regulatory domain defined by the gene and the insertions is contained within a single TAD. A more detailed version of this panel is shown in Supplemental Figure SE. (C) Two insertions in a gene desert between *Kcnt2* and *Cdc73* are divergently expressed in the limb bud (lb) and forebrain (fb), delineating a transition zone. This coincides with the respective insertions being located in different TADs. (*D*) Size distribution (*y*-axis) and relationship with TADs (color-coded) of functionally defined intervals. Only a single regulatory domain (RD) overlaps a TAD boundary. Random permutation of regions 200 kb–1 Mb in length (boxed area), where the size distribution of the different functional categories is not statistically different (Kolmogorov-Smirnov test, *P* = 0.8560 for RD vs. TZ + A + B; *P* = 0.9240 for RD vs. TZ), showed that RDs are significantly underrepresented in the "separated TADs" category. (*E* 

insertions with divergent expression between two insertions with the same activity.

Next, we defined *transition zones* (TZs) as regions separating insertions with distinct expression profiles (66 intervals, ranging from 14 kb to 1.9 Mb; median size: 734 kb; total combined length: 53.8 Mb). Other types of intervals (class A: two or more insertions without detectable expression; class B: one insertion with expression and one with no expression) were also annotated. B regions may be a type of transition, but we considered them separately,

since several local features may lead to inactivity of the sensor (Ruf et al. 2011).

# Regulatory domains are included in topologically associating domains

Next, we looked to see whether our operational subdivision of the genome matched any structural features of the genome. To this end, we compared RDs with the largely cell-invariant self-associating

"topologically associating domains" (TADs) identified by Hi-C (Dixon et al. 2012). We found that the great majority of RDs (78%) were contained within one TAD (Fig. 4D). The remaining ones extended into flanking unstructured regions (for which Hi-C data did not highlight specific compartmentalization). In only one case, two adjacent insertions with partially overlapping-but not identical-patterns of expression were found in distinct adjacent TADs. Similarly, insertions mirroring the expression patterns of adjacent but remote genes were also found within the same TAD (Supplemental Fig. 5C-E). In contrast, one third of TZs, including relatively short regions in gene-deserts, were located in different TADs (Fig. 4C; Supplemental Fig. 6). Since, on average, RDs were smaller than TZs, we performed several analyses to verify the significance of this different distribution relative to TADs. We randomly permutated RD, TZ, A, and B regions, and found it was significant (P < 0.05) that RDs almost never overlapped two or more TADs, unless we considered only extreme size ranges (below 400 kb; above 1.5 Mb). In particular, for intervals between 200 kb and 1 Mb, a range where the size distribution of the RD, TZ, A, and B regions was not significantly different, the altered distribution of RDs and TZs relative to TADs was highly significant (Fig. 4D). We also calculated the density of TAD ends for the real intervals and for 1000 random distributions of intervals of the same size in the genome, and found that RDs, but not the other categories, showed a significant depletion of TAD ends (Fig. 4E).

## Distribution of regulatory activities and insulators

Next, we compared how insulators or elements with enhancerblocking activities are distributed between the four types of operationally defined domains. We considered CTCF sites, SINE B1 and SINE B2 elements, as well as the TSS of protein-coding and noncoding genes from the RefSeq collection (Supplemental Table 6). All four elements were clearly depleted from RDs (Fig. 5A; Supplemental Fig. 7), whereas type A and B control regions never showed depletion, and TZs showed no, or far less depletion. This depletion was largely maintained, even when we considered potentially confounding factors, such as the clustering of CTCF sites and TSS (Kim et al. 2007; Shen et al. 2012) or the overlap of CTCF sites and SINE B2 elements (Schmidt et al. 2012) (Supplemental Fig. 7). The proportion of RDs containing at least one cell-invariant CTCF site was also lower than for TZ (39% compared with 70%), consistent with the proposed contribution of this protein in organizing regulatory interactions (Phillips and Corces 2009). However, since TAD boundaries are enriched for the different elements associated with insulator activity (Dixon et al. 2012), we also restricted our analysis to intervals located within TADs. In this case, we still found lower than expected density of TSS and SINE B2. However, the depletion of CTCF binding sites and SINE B1 X35S in RDs was no longer statistically significant (Fig. 5B; Supplemental Fig. 7), suggesting that it correlated with the topological, rather than regulatory subdivision of the genome.

As CTCF (Phillips and Corces 2009), together with cohesin complexes (Hadjur et al. 2009; Kagey et al. 2010; Merkenschlager and Odom 2013), has been proposed to have a major role in organizing regulatory interactions we examined their relationship to RDs in more detail, using available data sets (Supplemental Table 6). In agreement with previous analyses, we found a high degree of overlap between CTCF and cohesin-binding across mouse cell-types and tissues (Parelho et al. 2008; Wendt et al. 2008; Remeseiro et al. 2012). In several cases, cohesin and CTCF-bound regions lie within regulatory TZs, often also corresponding to TAD boundaries, an observation consistent with their proposed role (Phillips-Cremins et al. 2013).

However, even if slightly less abundant in RDs (Fig. 5C), many CTCF and cohesin binding sites are also interspersed within RDs, between co-expressed genes, insertions, and their associated enhancers (Fig. 6; Supplemental Figs. 4, 5, 8). We noted that RDs can also be further subdivided in more "specialized" ones, each characterized by expression specificities additional to the ones defining the RD (Fig. 5C; Supplemental Fig. 5A,E). Some of these subdivisions, with the current resolution offered by the available insertions, might be outlined by the presence of CTCF/cohesin sites (Fig. 5). However, by and large, it appeared difficult to match the mere distribution of the CTCF/cohesin sites with the distribution of regulatory activities, highlighting that, within TADs, the binding of cohesin/CTCF, defined by chromatin-immunoprecipitation, may not be a sufficient indicator of regulatory boundaries.

## Discussion

In vertebrates, since many critical enhancers lie at considerable distances from the genes they influence, ensuring and controlling proper interactions between regulatory elements and promoters is



**Figure 5.** Depletion of CTCF and cohesin in regulatory domains reflects the topological segmentation of the genome. Tissue-invariant CTCF sites (bound in more than nine tissues) are significantly depleted (but not absent) in regulatory domains when considering all functionally defined regions (*A*). However, the statistical significance of this depletion becomes marginal if the comparison is limited to intervals included in TADs, and not compared against the overall genome (*B*). Constitutive cohesin complex-binding sites (data not shown), or CTCF/cohesin co-occupied regions (*C*) also showed only a slightly reduced density in RDs, when compared with the part of the genome that is included in TADs.



**Figure 6.** CTCF and cohesin sites are interspersed in regulatory domains. Schematic representation of the large topological/regulatory domain on chr7:75.5M–77.8M. The two genes (*Arrdc4* and *Nr2f2*) are represented as arrows. The corresponding TAD is represented by a two-dimensional heat map (Dixon et al. 2012). Several constitutive CTCF sites (red lollipop, color intensity proportional to cell invariance), largely co-bound by cohesin (purple rings), are interspersed in this interval. Insertions spread across almost 2 Mb showed highly overlapping patterns in the proximal limb (blue arrow, *top*), face (blue arrow, *middle*), and at the midbrain/diencephalon boundary (blue arrow, *bottom*), forming a large regulatory domain. This large domain can be subdivided into smaller tissue-specific landscapes (green, purple, and brown) based on expression patterns. These different regulatory influences overlap with *Nr2f2* expression, detected by whole-mount in situ hybridization. In situ hybridization with *Arrdc4* probes did not reveal specific expression in E11.5 embryos. Embryos 183036-emb4 and 176069-emb50 were described previously (Ruf et al. 2011).

essential (Bulger and Groudine 2011; Splinter and de Laat 2011; Williamson et al. 2011). Our mechanistic understanding of this process is derived from studies carried out on a limited number of model loci (Gaszner and Felsenfeld 2006; Simonis et al. 2006; Vernimmen et al. 2007; Jhunjhunwala et al. 2008; Amano et al. 2009; Montavon et al. 2011; Tena et al. 2011; Marinić et al. 2013). Here, we expanded these by probing the regulatory landscape of the entire mouse genome with a transposable, naïve *lacZ* sensor. This large-scale exploration of the genomic regulatory architecture identified—in an operational manner—the widespread presence of large regulatory domains (RDs), within which the sensor displayed highly similar expression patterns at multiple distant positions. These RDs largely overlapped with TADs, the sub-megabase-sized

self-interacting intervals defined by chromosomal conformation capture analysis (Dixon et al. 2012; Nora et al. 2012). Insertions located in adjacent TADs almost systematically reported distinct regulatory activities, providing direct support for the suggested role of TADs as the basic building blocs of genomic regulatory architecture (Dixon et al. 2012; Hou et al. 2012; Nora et al. 2012; Gibcus and Dekker 2013). RDs are usually gene poor and shared several other features with TADs, such as a depletion of various elements (SINE B2 repeats, constitutive CTCFbinding sites).

Our operational approach also shed some light on the organization of these domains. Firstly, despite the overall inclusion of regulatory domains into TADs, the positions of the relative transitions were not always exactly superimposed, similarly to chromatin domains, which do not exactly correlate with TADs (Hou et al. 2012). These differences may partially arise from the low resolution of Hi-C, which can locate topological transitions with only limited precision (20 kb). Furthermore, topological transitions may not necessarily constitute absolute barriers, but act more like dampers (Andrey et al. 2013).

Within TADs, we found that enhancer activities are broadly distributed. and not targeted to specific regions (i.e., proximity of gene promoters or TAD borders). From their discovery, enhancers have been shown to act irrespectively of their orientation (Banerji et al. 1981). Genome-wide studies have further shown that enhancers can be found both 3' and 5' of their endogenous target genes (The ENCODE Project Consortium 2012; Li et al. 2012; Shen et al. 2012). Our data further stresses that orientation independence is an intrinsic property of enhancers in their normal context: in addition to controlling of target genes, enhancers generally act pervasively throughout their regulatory domains.

This broad distribution of enhancer activities along large domains has several implications. It may account for the transcriptional "ripple" effect observed upon growth-factor stimulation (Ebisuya et al. 2008) and bystander gene activation (Spitz et al. 2003; Cajiao et al. 2004; Zuniga et al. 2004). Furthermore, widespread enhancer activities may have been evolutionarily advantageous for genes brought into a new neighborhood by chromosomal rearrangements (Cande et al. 2009; De et al. 2009) or retrotransposition (Vinckenbosch et al. 2006), or for emerging lncRNAs (Ponting et al. 2009; Kutter et al. 2012), facilitating the acquisition of new expression domains.

Constraining enhancer activity through the formation of distinct topological compartments may contribute to gene regu-

lation in two important ways. It may help specify gene-enhancer interactions, and restrict ectopic "enhancer adoption" to accidental disruption of existing topologies, for example, through chromosomal rearrangements (Kokubu et al. 2003; Spitz et al. 2003; Niedermaier et al. 2005; Gostissa et al. 2009; Kantaputra et al. 2010; Marinić et al. 2013). Alternatively, it may help integrate the activity of multiple regulatory elements spread along large intervals (Carvajal et al. 2001; Uchikawa et al. 2003; Montavon et al. 2011; Li et al. 2012; Sanyal et al. 2012; Shen et al. 2012; Delpretti et al. 2013; Marinić et al. 2013; Visel et al. 2013) into the coherent regulatory units that have been described as regulatory archipelagos, holo-enhancers or chromatin-hubs (Palstra et al. 2003; Montavon et al. 2011; Marinić et al. 2013). Our observation that in intact endogenous loci, randomly inserted sensors report the complete integrated output provided to normal target genes (and do not decompose it into the individual activities of the closest enhancers) (e.g., Hoxd, Supplemental Fig. 4; Foxg1, Fig. 4B; Twist1, Ruf et al. 2011; Birnbaum et al. 2012), further supports the notion that regulatory control is exerted by coordinated, and not individual action, of enhancers. It indicates that this integration is orchestrated at the level of the regulatory domain, not at endogenous gene promoters (Marinić et al. 2013).

Along a given regulatory domain, the expression level detected by the sensor can vary quantitatively and reveal subdomains, including few positions apparently refractory to activation. We speculate that these "cold-spots" within otherwise permissive domains may shield genes from the influence of surrounding enhancers (Marinić et al. 2013), adding to other mechanisms of specificity such as core promoter sequences (Ohtsuki et al. 1998). This fine-scale organization of activities may be due to the positions of the corresponding enhancers, to local epigenetic modifications (Akhtar et al. 2013; Arnold et al. 2013), or to different tissue-specific three-dimensional organization within domains. Indeed, restructuring of physical interactions within TADs has been shown during lineage-commitment (Phillips-Cremins et al. 2013) and in different cell-types (Dixon et al. 2012). We suggest that such changes in local interactions may correlate with the different subregulatory domains that we observed within TADs. It will be an exciting avenue to explore what proteins (Hadjur et al. 2009; Kagey et al. 2010; Remeseiro et al. 2012; Aragon et al. 2013; Merkenschlager and Odom 2013) and nuclear substructures (Bickmore and van Steensel 2013; Gibcus and Dekker 2013; Meuleman et al. 2013) may be involved. Even though we found frequent co-occurrence of CTCF-bound regions with transitions between regulatory domains, our current analyses also highlight the difficulty of inferring the structure of these domains from the mere presence of CTCF/cohesin binding sites detected by chromatin-immunoprecipitation. This reinforces the growing perception that CTCF may have a versatile role (Handoko et al. 2011; Sanyal et al. 2012; Phillips-Cremins et al. 2013), perhaps different within topological domains and at their borders, and that analyzing these activities in the context of a 3D genome will be essential (Handoko et al. 2011; DeMare et al. 2013). Our data also emphasizes the need of direct functional approaches, as the one developed here, to map regulatory interactions and to compare them with physical conformations or chromatin maps. The confrontation of these approaches will be essential to understand the mechanistic basis and principles underlying the organization of the genome. Ultimately, such an understanding will be crucial to predict whether and how variants (such as structural variants) affect topological and regulatory organization, and consequently influence gene expression and phenotype (Weischenfeldt et al. 2013).

## Methods

#### Mouse lines and embryos with SBlac insertions

Insertion sites of the transposon were generated, mapped, and *lacZ* expression analysis was performed as described previously (Ruf et al. 2011). Information on insertions is found in Supplemental Table 1 and details are on the TRACER website (tracerdatabase.embl.de) (Chen et al. 2013). Whole-mount in situ hybridizations were carried out as described in Ruf et al. (2011). Templates for mRNA antisense probe synthesis were described in Spitz et al. (2003) (*Hoxd13*), Ruf et al. (2011) (*Sall1*), or produced by PCR on mouse cDNA using the following primers (Foxg1\_2F: GCCAAGCTGGCCTTTAAGC; Foxg1\_2R: ATTCTCCCACATTGC ACCTC; Nr2f2\_3Sp6: CATTTAGGTGACACTATAGCCACATGGG CTACATCAGAC; Nr2f2\_5: GGGCGGAGGAACCTGAGCTACAC; Hand2\_F: AGGACTCAGAGCATCAACAGC; Hand2-R: AGCGGATG CTCAAAGGTG).

Mouse experiments were conducted in accordance with the principles and guidelines in place at the European Molecular Biology Laboratory, as defined and overseen by its Institutional Animal Care and Use Committee, in accordance with the European Convention 18/3/1986 and Directives 86/609/EEC and 2010/63/EU.

#### Genomic data sets and resources used for the study

Genomic data sets were obtained from public sources and are summarized in Supplemental Table 6.

#### SB-EP300 comparison

We considered SB insertions with expression in heart, limb, midbrain, or forebrain, where EP300 binding data has been generated (Visel et al. 2009; Blow et al. 2010), and merged insertions with the same expression pattern <5 kb apart into one (Supplemental Table 2). In a given tissue, we calculated the enrichment of insertions with expression in that tissue, compared with the same number of randomly selected insertions with no expression in that tissue (200 randomizations). Enrichment was calculated for increasing distances (with steps of 5000 bp from 0 to 1 Mb, provided more than five insertions were found within). To test whether this enrichment was specific to the relevant tissue, we repeated the analysis using insertions expressed in other tissues or not expressed. For this, we removed EP300 sites if they were within 10 kb of a EP300 site in the first tissue considered, to reduce the possible confounding signal arising from clusters of different tissue-specific EP300 sites (Visel et al. 2009).

# Comparison of insertions, autonomous enhancer activity, and endogenous genes

We complemented the set of enhancers active at E11.5 from the VISTA Enhancer Browser (Visel et al. 2007) with additional enhancers from the literature. We visually inspected expression overlap between enhancer–insertion pairs (where insertions were within 200 kb of these enhancers; Supplemental Table 3), and annotated pairs as concordant, discordant, or inactive. As random control data sets (Supplemental Table 3), we first performed random permutation of the positions of the insertions and enhancers, and extracted new adjacent pairs. Second, we randomly drew insertion—enhancer pairs and compared their expression patterns.

For the enhancer-regulatory sensor-gene triplets, we compared the expression of characterized enhancers to reported in situ hybridization gene expression of either the first two flanking genes or genes <2 Mb away using gene expression resources (Supplemental Table 6). We considered sensor insertions lying within the same genomic interval, and annotated their expression as concordant, partially overlapping or divergent (no or different expression) with the enhancer-gene pair. These enhancer-gene-insertion triplets were then categorized depending on the position of the insertion relative to the enhancer and gene TSS. For neighboring insertions that showed the same annotation/relative position for a given enhancer, we kept only one triplet. Informative triplets are listed in Supplemental Table 4.

#### Defining regions with characteristic expression patterns

We functionally defined different classes of genomic intervals by comparing the expression of adjacent insertions <2 Mb apart. Intervals where both insertions showed no expression were annotated as class A. Class B was formed by pairs where one insertion was expressed whereas the other was not. Intervals defined by two insertions with discordant (nonoverlapping) expression patterns formed transition zones. Intervals where insertions showed overlapping *lacZ* expression were grouped as regulatory domains (RDs). To define a RD, the observed spatial pattern had to be identical, not only correspond to the same broad anatomical domain: Patterns as in Figure 1B are considered divergent. In cases where multiple insertions within a 2-Mb window were concordantly expressed, we considered the most centromeric and telomeric insertions as the boundaries of the RD, but only if no insertions in between were discordantly expressed. Thus, we merged adjacent RDs with the same activity together, sometimes including small B regions. Similarly, we determined the minimal extent of transition zones as the smallest region flanked by divergently expressed insertions (i.e., where none of the expression patterns overlapped), even if they were disrupted by insertions with no expression. We did not further group insertion pairs, where one or both insertions are not expressed, except when three or more consecutive insertions showed no expression, and one pair was <10 kb apart. The extended annotated regions are listed in Supplemental Table 5.

## Comparison to topological domains and other genomic landmarks

We compared the occurrence of certain features in our functionally defined regions to a random model. To estimate the significance (*P*-value) of the frequency of certain features (e.g., topological boundaries, CTCF sites, SINE B2 elements) for each category (i.e., all A, B, RD, and TZ domains, or those found intra-TAD only), we applied bootstrap following recommendations from Phipson and Smyth (2010). For details, see Supplemental Note 1. For TAD boundaries we also performed random permutation, as described in the text and Supplemental Note 1. For the analysis we used the data sets described in Supplemental Table 6.

## Data access

All insertions generated for this study are documented in the TRACER database (Chen et al. 2013) (http://tracerdatabase.embl.de; or http://www.ebi.ac.uk/panda-srv/tracer/index.php).

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