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An evolutionarily conserved three-dimensional structure in the vertebrate *lrx* clusters facilitates enhancer sharing and coregulation

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Developmental gene clusters are paradigms for the study of gene regulation; however, the mechanisms that mediate phenomena such as coregulation and enhancer sharing remain largely elusive. Here we address this issue by analysing the vertebrate *lrx* clusters. We first present a deep enhancer screen of a 2-Mbp span covering the *lrxA* cluster. Using chromosome conformation capture, we show that enhancer sharing is widespread within the cluster, explaining its evolutionarily conserved organization. We also identify a three-dimensional architecture, probably formed through interactions with CCCTC-binding factor, which is present within both *lrx* clusters of mouse, *Xenopus* and zebrafish. This architecture brings the promoters of the first two genes together in the same chromatin landscape. We propose that this unique and evolutionarily conserved genomic architecture of the vertebrate *lrx* clusters is essential for the coregulation of the first two genes and simultaneously maintains the third gene in a partially independent regulatory landscape.

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The *Iroquois* (*Irx/iro*) genes, present in all metazoans, encode TALE class homeoproteins that exert multiple functions during animal development (reviewed in ref. 1). One of the striking characteristics of the members of this gene family is their organization in large genomic clusters, which have arisen independently across the animal kingdom^{2–6}. In these clusters, the *Irx* genes are separated by large intergenic regions, denoted gene deserts. Most vertebrates contain six *Irx* genes grouped in two paralog clusters of three genes each^{2,3}. The *IrxA* cluster contains *Irx1*, *Irx2* and *Irx4*, and the *IrxB* cluster contains *Irx3*, *Irx5* and *Irx6*. In all vertebrates analysed, the *Irx1/Irx2* and *Irx3/Irx5* pairs have very similar expression patterns^{3,7–14}. The expression of the third gene in each cluster, *Irx4* or *Irx6*, is usually more divergent. However, in some tissues, all the genes of a cluster, or even of both clusters, are identically expressed^{3,7,10,14}.

Why these genes tend to be organized in such large clusters remains a mystery. One possibility is that the different genes within a cluster share conserved intergenic regulatory elements. Consistent with this idea, we have identified many highly conserved non-coding regions (HCNRs) that function as *cis*-regulatory elements and are distributed throughout the gene deserts of the *IrxB* cluster³. These regulatory sequences drive the expression in sub-domains of the territories expressing the *IrxB* genes. Many of these sub-domains show expression of more than one *IrxB* gene, pointing again to the presence of shared enhancers. However, to date there has been no reported genetic or molecular evidence that enhancer sharing exists within vertebrate *Irx* clusters.

Here we present an extensive enhancer screen of the *IrxA* cluster. We demonstrate the presence of multiple *cis*-regulatory elements throughout this genomic region, which promote expression in sub-domains of the *IrxA*-expressing territories. Moreover, using the chromosome conformation capture (3C) technology, a PCR-based method that allows determining the physical *in vivo* interaction between any chromatin segments (reviewed in ref. 15), we examined the interaction of four representative enhancers with the promoters of the three *IrxA* genes. We show that enhancers are shared among the *Irx* promoters, although they preferentially interact with the first two genes of the cluster. This provides direct physical evidence that enhancers are shared in the cluster, probably placing a strong evolutionary constraint that maintains the linear association of these genes. Most interestingly, we also identify an evolutionarily conserved three-dimensional (3D) architecture, present in both *Irx* clusters of all vertebrates, that brings the *Irx1/3* and *Irx2/5* promoters into physical contact. This loop might explain the preferential interaction of the various enhancers with the first two genes of the *IrxA* cluster and the coregulation of these genes in both clusters. We finally show that the formation of this evolutionarily conserved loop in vertebrates is probably mediated by CCCTC-binding factor (CTCF). Our work not only provides the first demonstration that enhancers are shared within the *Irx* clusters, but also identifies a deeply conserved three-dimensional architecture that predates cluster duplication. This architecture would help to generate different regulatory landscapes for linearly arranged genes.

Results

***Cis*-regulatory landscape of the *IrxA* cluster.** Many reports have shown that HCNRs are enriched in *cis*-regulatory elements (see for example refs 3, 16, 17). To gain insight into the regulatory landscape of the *IrxA* cluster in vertebrates, we used an *in vivo* transgenic assay to examine the activity of 88 HCNRs distributed throughout the cluster and present in all tetrapods. These regions were amplified from *Xenopus tropicalis* genomic DNA and cloned in an appropriate vector for *Xenopus* transgenesis (see Methods). Of the 88 HCNRs, 16 (18%) activated expression in a robust and reproducible manner at the tail-bud stage (Fig. 1), a proportion in line with our previous analysis of HCNRs from the *IrxB* cluster³. These HCNRs drove

reporter expression within sub-domains of the endogenous pattern of *IrxA* genes at this stage¹⁴, such as the midbrain, hindbrain, spinal cord, kidney and otic vesicle (Fig. 1). These patterns are discrete and restricted to specific territories, although the domains of several elements partially overlap.

Eight of the *IrxA* HCNRs are partially conserved in the *IrxB* cluster in largely equivalent genomic locations¹⁸ indicating an origin that predates the duplication of the clusters. Interestingly, of these eight paralogous sequences, five were positive in our *Xenopus* transgenic assay (2,165, 2,695, 3,173, 3,240 and 3,565) and all *IrxA* and *IrxB* paralogues show similar regulatory activities, with the exception of the paralogue region of 3,240 which showed no enhancer activity (Supplementary Fig. S1 and see ref. 3).

To test the evolutionary conservation of enhancer function in vertebrates, we next examined the activity of four of these HCNRs in mouse and zebrafish transgenic assays. In most cases, these orthologous regions activated the expression of reporter genes in territories largely equivalent to those promoted by the *Xenopus* HCNRs (Fig. 1).

***IrxA* enhancers interact with multiple *Irx* promoters.** Most enhancers identified in this study drive expression in territories expressing more than one *Irx* gene, making it likely that their regulatory activity is shared by different genes of the cluster. To explore this, we examined physical interactions *in vivo* between different regulatory elements and the three *Irx* promoters by means of 3C¹⁹. For these experiments, we selected four enhancers well distributed along the cluster, two located between the *Irx1* and *Irx2* genes (3,565 and 3,240, Figs 1 and 2a) and the other two located in the *Irx2/Irx4* intergenic region (2,260 and 2,165, Figs 1 and 2a). Of these four enhancers, two are conserved in both *Irx* clusters (3,565 and 2,165) and the other two are specific for the *IrxA* cluster (3,240 and 2,260). Moreover, as for our 3C experiments we used tissues dissected from embryos, to increase homogeneity, we selected enhancers driving strong expression in as many cells as possible. The enhancers were taken as fixed positions, and interactions were tested with each promoter and two flanking regions ~30 kbp upstream and downstream. As we observed a clear inverse relationship between distance and interaction level, we defined the average of the interactions of each enhancer with its two promoter-flanking regions as the background level of interaction (see Fig. 2b). These enhancer–promoter interactions were examined in midbrain–hindbrain and limb tissues. As all four enhancers are predominantly active in the neuroectoderm (Figs 1 and 2a), we expected that any enhancer–promoter interactions would be stronger in neural tissues than in limb tissues. This was indeed the case, and all enhancers seem to interact preferentially with *Irx* promoters rather than with the neighbouring chromatin region, this interaction being stronger in neural tissue (Fig. 2b). Several enhancers show significant interaction with more than one promoter (3,565 with *Irx1* and *Irx4*; 3,240 with *Irx1* and *Irx2*), and these enhancer–promoter interactions can occur even over unprecedentedly long distances of 1.5 Mbp (3,565 with *Irx4* and 2,260 with *Irx1*). It is also noteworthy that most enhancers interact preferentially with the first two genes of the cluster, which raises the question of what molecular mechanism underlies such preferential interaction.

An evolutionarily conserved 3D architecture of *Irx* clusters. We next evaluated the existence of enhancer–enhancer and promoter–promoter interactions. No interactions were observed between enhancers (Supplementary Fig. S2a), but a very strong and significant interaction was detected between the *Irx1* and *Irx2* promoters (Fig. 3a). In contrast, no interactions were found between either of these promoters and the *Irx4* promoter (Fig. 3a and Supplementary Fig. S2b). Moreover, in contrast to the enhancer–promoter interactions, which were tissue specific, the *Irx1*–*Irx2* interaction

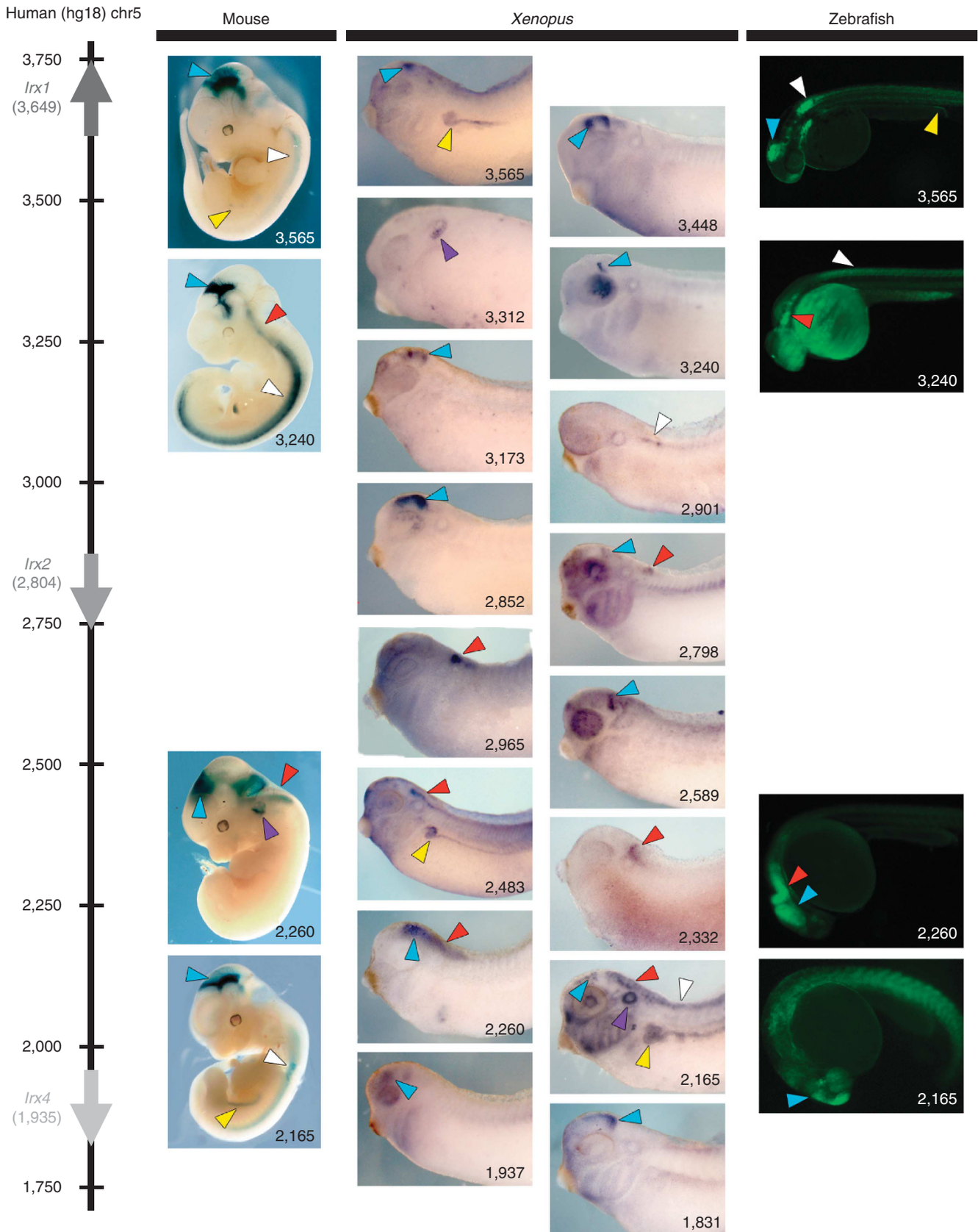


Figure 1 | Multiple enhancers are distributed along the the *IrxA* cluster. The black vertical line at the left depicts the *IrxA* cluster, showing coordinates (kbp) of human chromosome 5. Columns show enhancer activity of different HCNRs in embryos of mouse (stage E11.5), *Xenopus* (stage 30) and zebrafish (24–36 h.p.f.). Enhancers are named according to the coordinates (in kilobases) of their human HCNr counterparts. Orthologous regions promote expression in largely equivalent territories. Arrowheads mark tissues by colour as follows: blue, midbrain; red, hindbrain; white, spinal cord; yellow, kidney; and purple, otic vesicle.

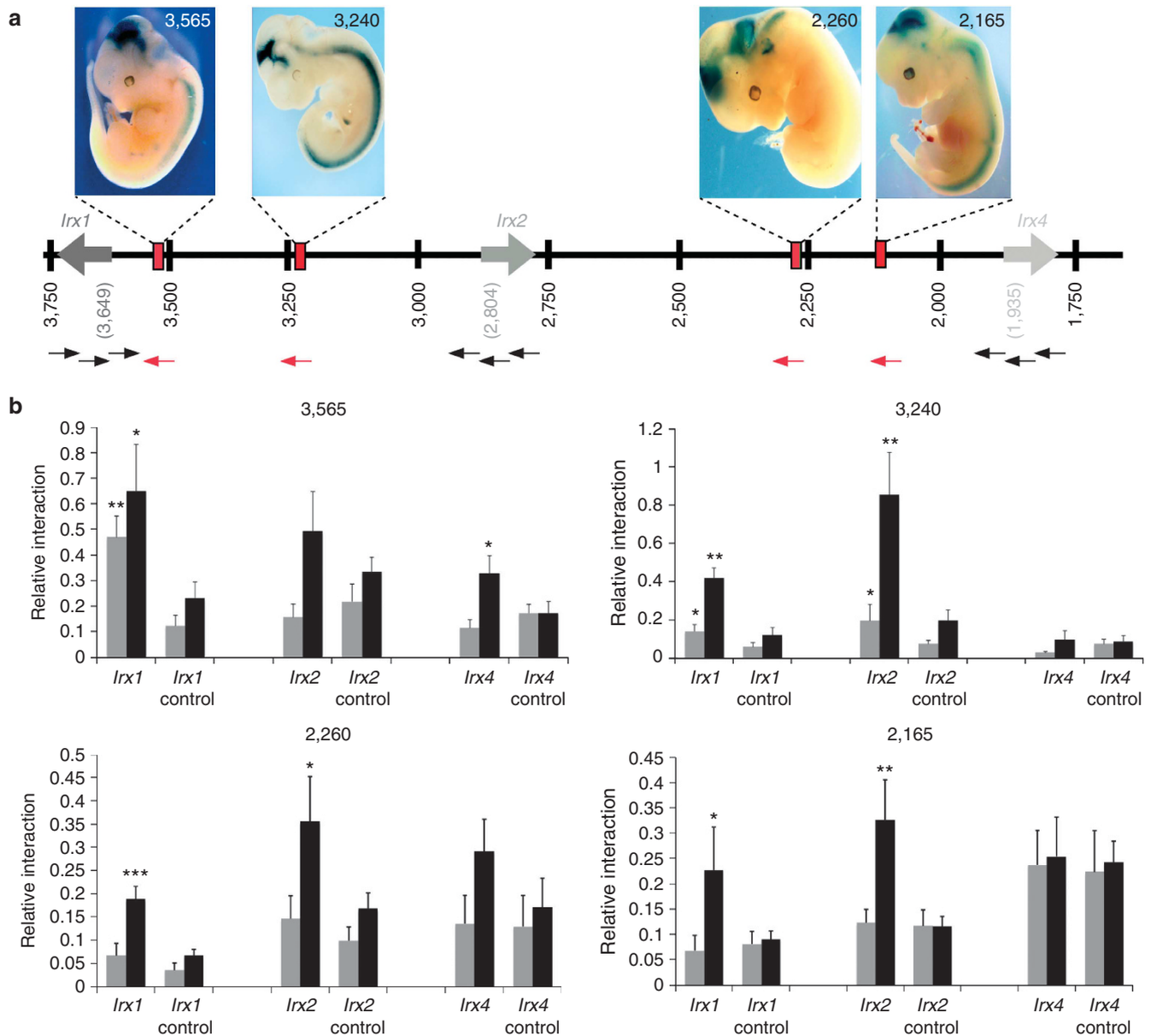


Figure 2 | Enhancers physically interact with multiple *Irx* promoters. (a) Expression patterns of the enhancers tested in mouse and their respective positions on human chromosome 5. Arrows mark primer positions used for 3C studies; fixed positions are coloured in red and variable positions in black. (b) Graphical representation of interactions between each enhancer with the different *Irx* promoters, as determined by 3C in two tissues: midbrain/hindbrain (black bars) and limbs (grey bars). The background interaction was calculated as the average of the interactions observed between each fixed position and two regions flanking the promoters. Graphs show means of at least three independent experiments for each enhancer. Asterisks indicate enhancer-promoter interactions that differ significantly from enhancer-control interactions: * $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$; Student's *t*-test. Error bars indicate s.e.m. Exact *P*-values are listed as follows: 2,260 versus *Irx1* (brain), $P = 0.0003$; 2,260 versus *Irx2* (brain), $P = 0.020$; 3,240 versus *Irx1* (limb), $P = 0.043$; 3,240 versus *Irx1* (brain), $P = 0.001$; 3,240 versus *Irx2* (limb), $P = 0.047$; 3,240 versus *Irx2* (brain), $P = 0.003$; 3,565 versus *Irx1* (limb), $P = 0.002$; 3,565 versus *Irx1* (brain), $P = 0.014$; 3,565 versus *Irx4* (brain), $P = 0.049$; 2,165 versus *Irx1* (brain), $P = 0.024$; 2,165 versus *Irx2* (brain), $P = 0.002$.

was found in all tissues examined, independently of whether they express these genes (midbrain-hindbrain and E15 limbs, Fig. 3) or not (telencephalon and E10 limbs, Supplementary Fig. S3). We therefore conclude that this 3D architecture may enable positive or negative coregulation of *Irx1* and *Irx2* by facilitating equal access of different *IrxA* enhancers or repressors to their promoters.

The two vertebrate *Irx* clusters arose from a genome duplication event that took place at the base of the vertebrate lineage. As a consequence, the overall genomic structure of both clusters is very similar and has been maintained throughout vertebrate evolution. Thus, in all vertebrates, both *Irx* clusters are very large, the relative distance between genes is similar and there is a conserved orienta-

tion of transcripts and distribution of paralogous HCNRs. Moreover, in both clusters, the first two genes are largely coregulated, whereas the third gene shows a more divergent expression pattern (reviewed in ref. 1). To test whether the 3D architecture described here, which brings the first two genes of the *IrxA* cluster into close proximity, is evolutionarily conserved and also forms at the *IrxB* cluster, we examined the putative interaction between *Irx3* and *Irx5*, and found that their promoters strongly and significantly interact in both mouse and in *Xenopus* (Fig. 3b,c). Similar to what occurs in the *IrxA* cluster, no interaction was found between the *Irx3* and the *Irx6* promoter (Supplementary Fig. S2c). By bringing the first two genes of both *Irx* clusters together, this evolutionarily conserved 3D

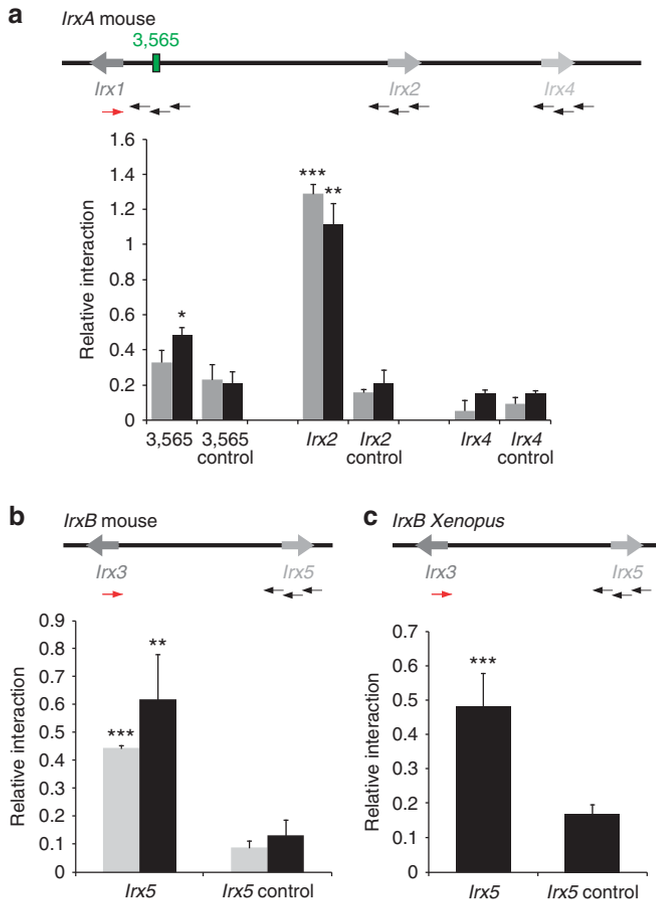


Figure 3 | An evolutionarily conserved 3D architecture is formed between the first two genes of each *Irx* cluster. Arrows show the fixed (red) and variable (black) primer positions used in 3C assays. (a) In mouse tissues, the *Irx1* promoter strongly and significantly interacts with the 3,565 enhancer and, even more frequently, with the *Irx2* promoter, but not with *Irx4*. (b) An equivalent interaction between *Irx3* and *Irx5* promoters was observed in mouse and *Xenopus* embryos. In these experiments, the fixed primers are situated at the *Irx3* promoters. Background interaction was calculated, for each species, as the average of the interactions between each fixed position and two flanking regions at a distance of 30 kbp from the *Irx5* promoters. Graphs show means from at least three independent experiments. Asterisks indicate enhancer–promoter interactions that differ significantly from enhancer–control interactions: * $P < 0.05$; ** $P < 0.01$; *** $P < 0.001$; Student's *t*-test. Error bars indicate s.e.m. Exact *P*-values are listed as follows: *mIrx1* versus 3,565 (brain), $P = 0.023$; mouse *Irx1* versus *Irx2* (brain), $P = 0.001$; mouse *Irx1* versus *Irx2* (limb), $P = 3.9 \times 10^{-6}$; mouse *Irx3* versus *Irx5* (brain), $P = 0.004$; mouse *Irx3* versus *Irx5* (limb), $P = 9.2 \times 10^{-6}$; *Xenopus Irx3* versus *Irx5*, $P = 2.4 \times 10^{-4}$.

architecture places them in the same regulatory landscape, probably facilitating similar access by all enhancers.

CTCF likely helps form the 3D *Irx* architecture. CTCF is a DNA-binding factor known to facilitate the formation of chromatin loops²⁰. To gain insight into the possible implication of CTCF in the physical interaction between *Irx* promoters, we explored the ENCODE database²¹, available via the UCSC browser²², to examine CTCF distribution in different human cell lines. In most human cell lines (Fig. 4a), CTCF is bound to the promoters of the *Irx* genes, preferentially to those of the first two genes of each complex (*Irx1/Irx2* and *Irx3/Irx5*). A similar situation occurs in mouse ES and fibroblast cells²³. This may therefore be an evolutionarily conserved feature

of the *Irx* genes. Accordingly, using the Jaspar and CTCF-binding site databases^{24,25}, we also found potential CTCF-binding sites in the vicinity of the *Xenopus* and zebrafish *Irx* genes (Supplementary Fig. S4). CTCF is therefore a good candidate mediator of the formation of the conserved *Irx* 3D architecture. To test this possibility, we knocked down CTCF function in zebrafish using two specific morpholinos (see Methods and Supplementary Fig. S5a–c for details). Examination of the contact between the *irx3a* and *irx5a* promoters in non-injected (non-morphant) zebrafish embryos revealed a clear and significant interaction (Fig. 4b), indicating that the 3D architecture that associates these promoters is also conserved in the fish genome. Reduction of CTCF function in morphant embryos of both types significantly reduced the contact between these promoters (Fig. 4b), suggesting that CTCF likely participates in the formation of this 3D architecture in all vertebrates. However, we cannot discard the possibility that the reduced contact between the *irx3a* and *irx5a* promoters in the CTCF morphant embryos is an indirect consequence of reduced expression of downstream CTCF targets.

The 3D architecture of the *Irx* cluster, likely facilitated by CTCF, may serve two functions. First, by bringing together the promoters of the first two genes of the clusters, it might make both genes similarly accessible to most *cis*-regulatory elements, thus contributing to their coregulation. Second, it might generate a more divergent regulatory landscape for the third gene of the cluster, by reducing the ability of some *cis*-regulatory elements within the loop to function on the promoter of the third gene. We therefore expect that CTCF impairment should have an impact on the expression patterns of all three genes in the cluster, and this is what we saw in the zebrafish *IrxBa* cluster in embryos injected with either of the two CTCF-specific morpholinos (Fig. 4c). Both *irx3a* and *irx5a* were overexpressed all along the neural tube, especially in the spinal cord, and *irx6a* was downregulated in the midbrain and hindbrain, whereas its expression increased in the otic vesicle and in the lateral epidermis (Fig. 4c). Interestingly, these two last domains correspond to tissues expressing one or both of the other *irx* genes, suggesting that, in the morphants, some enhancers that normally preferentially function on these other genes are now able to efficiently activate *irx6a*. Our results are therefore compatible with the 3D architecture shown in Figure 4d.

Remodelling of the *IrxA* architecture during teleost evolution.

If the above model is correct, any disruption of the *Irx* complexes should have a major impact on the expression of the first two genes, which are coregulated, whereas the third gene, which is present in a more independent regulatory landscape and is controlled by fewer *cis*-regulatory elements, should be less affected. To test this prediction, we took advantage of a system in which the highly conserved structure of the vertebrate cluster has not been maintained. The genome duplication that occurred at the base of the teleost lineage has provided extra copies of *IrxA* genes. This permitted divergent evolution^{26,27}, enabling comparison of the expression of the *IrxA* genes in clusters with different architectures. In medaka, the *IrxAa* cluster contains the full set of *Irx* genes, organized in a manner similar to that of other vertebrates. The expression patterns of medaka *IrxAa* genes are also similar to those of tetrapod *IrxA* genes, such as mouse or *Xenopus* (Figs 2a and 5a,b). For example, in medaka and *Xenopus*, *irx1/irx1a* and *irx2/irx2a* are largely coexpressed in broad areas of the neuroectoderm, whereas *irx4/irx4a* expression is limited to a restricted area of the hindbrain and to the heart (Fig. 5a,b).

In contrast to medaka, the zebrafish *IrxAa* cluster has split into three (Fig. 5c). *irx1a* remains associated with most of the original *Irx1/Irx2* intergenic region (200 kbp) and contains many HCNRS, including those corresponding to the 3,565 and 3,240 enhancers. *irx2a* and *irx4a*, on the other hand, retain only 25 and 45 kbp, respectively, from the original cluster sequences, with very few

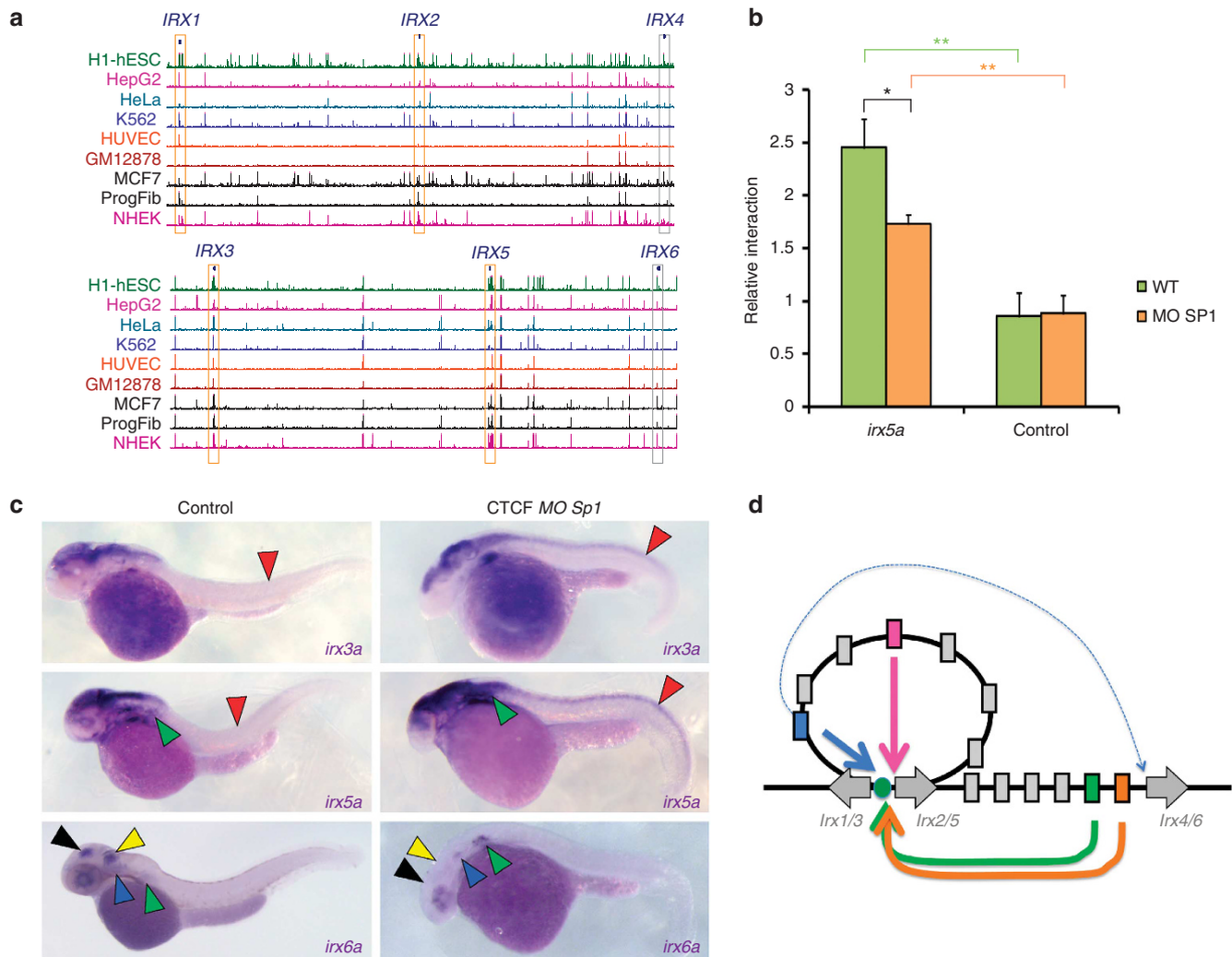


Figure 4 | The three-dimensional *lrx* architecture depends on CTCF. (a) Distribution of CTCF binding within the *lrx* clusters as determined by ChIP-seq in different cell types in the ENCODE project. In most cell types, CTCF is bound to the *lrx* promoters (coloured boxes). Note that the first two genes of each complex (orange boxes) are bound by CTCF in more cell types than the promoter of the third gene (grey boxes). (b) 3C assays to detect *lrx3* and *lrx5* interactions in zebrafish embryos. The fixed primer is located at the *lrx3* promoter and the background interaction was calculated as the average of the interactions between the fixed position and two flanking regions at a distance of 30 kbp from the *lrx5* promoter. A clear interaction is detected between the *lrx3* and *lrx5* promoters (green bars), which is significantly reduced in CTCF morphant embryos (orange bars). Graphs show means from at least three independent experiments: * $P < 0.05$; ** $P < 0.00$; Student's *t*-test. Error bars indicate s.e.m. (c) Expression of zebrafish *lrxAa* genes in control and CTCF morphant embryos. Note that the levels of *lrx3a* and *lrx5a* expression are increased, especially in spinal cord (red arrowheads), whereas *lrx6a* is downregulated in midbrain and hindbrain (black and yellow arrowheads) and upregulated in the otic vesicle and lateral epidermis (blue and green arrowheads, respectively). (d) Model of the 3D architecture of the *lrx* clusters. The proximity of the promoters of the first two genes is probably facilitated by CTCF (green circle). Similar access of enhancers to these genes is shown with coloured thick arrows. Restricted access of an enhancer to the third gene of the cluster is shown by the discontinuous thin blue arrow. Coloured boxes: relative positions of the enhancers analysed by 3C. Grey boxes: relative positions of other enhancers identified in this study.

associated HCNRs. The expression of zebrafish *lrx1a* is more restricted than that of the medaka *lrx1a* or *lrx2a* genes (Fig. 5b,c). Specifically, zebrafish *lrx1a* is only weakly expressed in the posterior hindbrain and spinal cord. This is in agreement with the reduced number of HCNRs associated with *lrx1a* in zebrafish and with the fact that most hindbrain and spinal cord enhancers identified in this study are located in the intergenic territory between *lrx2* and *lrx4*. Because of the chromosomal rearrangement in zebrafish, this territory is no longer associated with these genes. The expression of zebrafish *lrx2a* is restricted to only a few domains (Fig. 5c), again correlating with the low number of HCNRs associated with this gene. *lrx4a* is expressed in the hindbrain and in the heart (Fig. 5b,c), in a manner similar to medaka *lrx4a*, which would indicate that most *lrx4a* enhancers remain associated with this gene in zebrafish.

The expression patterns of zebrafish *lrxAa* genes are therefore compatible with our predictions based on the architectural features of the *lrx* clusters, namely, that the disruption of the *lrxAa* cluster in zebrafish has had a major impact on *lrx1a* and *lrx2a* expression, but much less of an effect on *lrx4a*.

Discussion

Developmental genes are expressed in a variety of tissues and at different embryonic stages, fulfilling multiple functions in different processes. To achieve this, their rate of transcription is controlled by multiple *cis*-regulatory elements, some of them functioning over very long distances^{28,29}. Our analysis has identified a large number of *cis*-regulatory elements, located in HCNRs and distributed throughout the 2-Mbp *lrxA* cluster, and that altogether appear to recapitulate

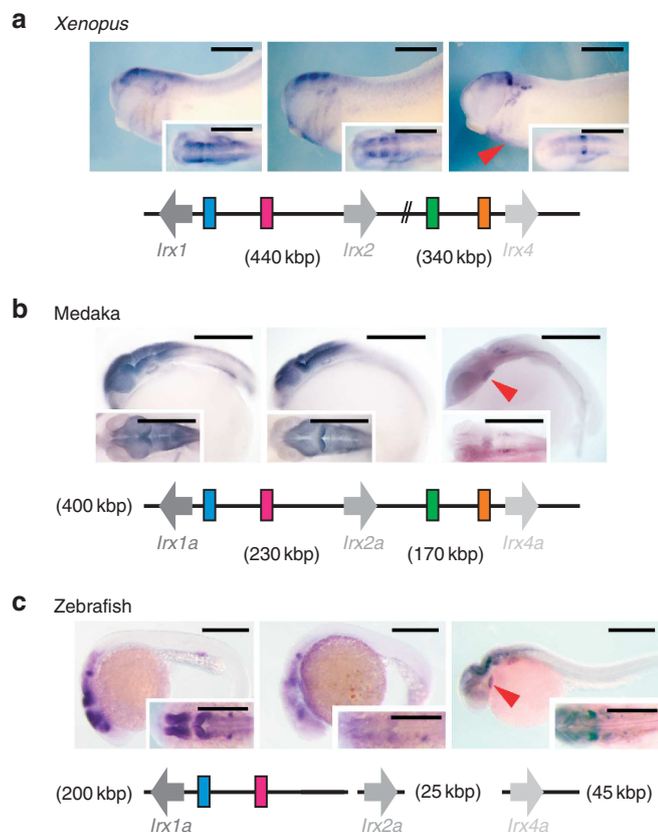


Figure 5 | Disassembly of the *IrxA* cluster alters the expression patterns of *IrxA* genes. Genome architecture and expression patterns of *IrxA* genes in (a) *Xenopus*, (b) medaka and (c) zebrafish. Expression patterns at neurula stages are shown in lateral and dorsal (insets) views. In a, b, numbers in parentheses indicate the intergenic distances, and in c it represents the size of orthologous *IrxAa* DNA associated with each gene. Red arrowheads point at *Irx4* expression in the heart. Coloured rectangles: enhancers analysed in this study by 3C. Some of these enhancers, as well as others, were lost upon disaggregation of zebrafish *IrxAa* cluster. Scale bars, 0.5 mm.

the expression patterns of *IrxA* genes. Most enhancers drive expression in territories expressing more than one *Irx* gene, making it likely that different genes of the cluster are regulated in concert. Sharing of *cis*-regulatory elements between neighbouring genes has often been suggested, but only demonstrated in a handful of cases^{30–34}. In all these examples, a single enhancer or a locus control region interacts with multiple genes. Using 3C (ref. 19), we show that *cis*-regulatory elements located throughout the *IrxA* cluster physically interact with more than one *Irx* promoter, even over distances of 1.6 Mbp (which, to the best of our knowledge, is the most distant physical interaction between an enhancer and a promoter so far reported). Interestingly, the interactions of each enhancer with the promoters seem to occur one by one, as we could not detect interactions between different enhancers. We propose that the presence of shared enhancers is probably the major evolutionary constraint that maintains the association of the *Irx* genes in clusters.

Despite the sharing of enhancers, the expression patterns of all the *Irx* genes do not completely overlap; the first two genes of both *Irx* clusters are coexpressed in most tissues, but the third gene shows a more limited and divergent expression pattern^{3,7,11–14,35–37}. Accordingly, we find that enhancers preferentially interact with the two first genes of the *IrxA* cluster. This raises the important question of how this differential regulation takes place. We demonstrate the existence of a conserved 3D architecture in the *Irx* clusters, prob-

ably dependent on CTCF, which helps to explain this differential regulation. This 3D conformation, which seems to be independent of transcription, brings the promoters of the first two genes of each cluster into close proximity. This would place these genes in the same regulatory landscape, thereby probably facilitating their coregulation, both positive and negative. Moreover, the same 3D architecture would be likely to trap *cis*-regulatory elements and thus reduce their access to the promoter of the third gene of the complex (Fig. 4d), generating a rather different regulatory landscape for this gene. If this model is correct, disruption of the cluster would more strongly affect the expression of the first two genes. This prediction is borne out by the naturally occurred genomic dispersion of the *IrxAa* cluster in zebrafish. However, definitive demonstration of this model would require the targeted rearrangement of the existing *Irx* clusters in a model system such as mouse.

In several aspects, the *Irx* genes resemble the *Hox* genes. Both types of genes encode homeoproteins that are essential for animal development, both are organized in clusters, genes within these clusters share regulatory elements, and the clusters form complex 3D architectures that are probably CTCF dependent^{1,38–40}. However, despite these similarities, there are also important differences. During evolution, the original *Hox* cluster appears to have arisen before the arthropod–vertebrate split³⁸. In contrast, *Irx* clusters seem to have arisen independently several times during evolution^{2,6}. Therefore, in contrast to *Hox* clusters, clustering of *Irx* genes in different lineages are not related. Once generated, both these clusters are probably maintained by the use of shared regulatory elements. In *Hox* clusters, shared regulation is mainly mediated by global control regions that lie outside the clusters³⁹. In the *Irx* clusters, we find many and diverse shared enhancers within the clusters. Nevertheless, we cannot exclude the possibility that other shared *Irx* enhancers lie outside these clusters. Finally, in *Hox* clusters, multiple loops, probably dependent on CTCF, are present in silent clusters. These loops are released upon cluster activation, suggesting that they may be associated with the process of cluster silencing. This contrasts with our results, which suggest that contact between promoters in the *Irx* clusters occurs in a similar manner in tissues independently of whether they express *Irx* genes, probably facilitating either their coactivation or their corepression. Moreover, our data show that the architecture of the two paralogous clusters is similar, whereas the architecture of the various human paralogous *HOX* clusters appears to be specific in each case⁴⁰. Finally, it remains to be shown whether, as we show for the *Irx* clusters, the 3D architecture of orthologous *Hox* cluster is conserved.

On the basis of our findings, we propose that the formation of internal loops in gene clusters, which may be regulated in time and space, would facilitate the formation of specific regulatory landscapes for certain genes of the cluster in a tissue- and stage-dependent manner. This may favour the further sub-functionalization of these independently regulated genes. Thus, the presence of loops may have evolved in these clusters as a new dimension to satisfy a requirement for differential expression of otherwise coregulated genes organized in a linear arrangement in the genome.

Methods

Animal transgenesis. All HCNRs were amplified by PCR from mouse, *Xenopus tropicalis* or zebrafish genomes using the primers listed in Supplementary Table S1. The PCR fragments were subcloned in PCR8/GW/TOPO vector and, using Gateway technology, transferred to the corresponding destination vectors for transgenesis in mouse, *Xenopus tropicalis* or zebrafish. For the generation of transgenic mice, the genomic fragments were transferred into a vector containing the human minimal β -globin promoter, *lacZ* and a SV40 polyadenylation signal⁴¹. Vectors were subsequently linearized, the vector backbone removed and the construct microinjected into one-cell mouse embryos. F0 embryos at stages 11.5–13 d.p.c. were collected and stained for *lacZ* activity. An enhancer was considered positive when three or more independent transgenic embryos showed the same expression pattern. For the generation of transgenic *Xenopus*, the fragments were transferred into a vector containing the *Xenopus* 0.6-kbp *Gata2* minimal promoter driving

green fluorescent protein (GFP)⁴². Transgenesis was performed using the I-SceI method following the protocol already described⁴³. Briefly, after *in vitro* fertilization of the oocytes, one-cell-stage *Xenopus* embryos were injected with 5 nl of the injection mix, composed of the construct DNA (final concentration 5 ng μ l⁻¹), 5 U of I-SceI enzyme (New England Biolabs) and 1X I-SceI buffer. *In situ* hybridization to detect GFP mRNA was used to identify the transgenic embryos. An enhancer was considered positive when ten or more independent transgenic embryos showed the same expression pattern. For the generation of transgenic zebrafish, the fragments were transferred into the ZED vector⁴⁴. Zebrafish transgenic embryos were generated using the Tol2 transposon/transposase method⁴⁵, with minor modifications. One-cell embryos were injected with a 2-nl volume containing 25 ng μ l⁻¹ of transposase mRNA, 20-ng μ l⁻¹ phenol/chloroform, purified ZED constructs and 0.05% phenol red. For zebrafish, three or more independent stable transgenic lines were generated for each construct.

All animal experiments were conducted following guidelines established and approved by local Government and our Institutional Animal Care and Use Committee, and in accordance with best practices outlined by the European Union.

Chromosome conformation capture assays. Midbrain, hindbrain and spinal cord (M–H samples), limbs and heart were dissected from E15 mouse embryos and processed to obtain single-cell preparations. For *Xenopus* and zebrafish, single-cell preparations were prepared from total embryos. A total of 10⁷ isolated cells were fixed and lysed, and nuclei were then digested with *Hind*III endonuclease (Roche). DNA was then treated with low concentrations of T4 DNA ligase (Promega) to favour intramolecular ligations. A set of locus-specific primers (Supplementary Table S2) was designed with the online program Primer3 v. 0.4.0 (ref. 46), each primer being close to a *Hind*III site flanking a chromosomal site of interest. For mouse and *Xenopus* 3C experiments, we measured the relative enrichment of each ligation product by semi-quantitative PCR, as described in ref. 20. Each DNA sample was calibrated so that the PCR products were always in the linear range. Primers next to each enhancer were considered fixed primers, and different interactions were tested using primers close to the promoters. For each interaction, two negative-control primers were designed to target sites ~30 kbp upstream and downstream of the region of interest. PCR products were run on an agarose gel and measured with a Typhoon scanner. For the zebrafish 3C experiments, which required precise quantifications, we followed the quantitative PCR protocol indicated in ref. 47. In all cases, product values were related to a control composed of bacterial artificial chromosomes (BACs) that encompass all our regions of interest (RP23-127L21, RP23-347L10, RP23-93E17, RP23-52B16 and RP23-131B17 for mouse genome; OAAA043I21 and OAAA098B15 for *Xenopus*; and DKEY-103G18 and CH211-25M11 for zebrafish). To compare data from different tissues, PCR values were normalized by means of control primers targeting the *Erc3* gene locus in each species.

Statistical analysis. Assuming that data is normally distributed, one-tailed *t*-test was performed to test significance of differences among sample averages. In all tests, we adopted an alpha level of 0.05. Differences were considered significant or highly significant when *P*-values were situated below 0.05 or 0.01, respectively.

Zebrafish morpholino injections. Two zebrafish morpholinos to knockdown CTCF function were used. *MOatg* (5'-CGGCCTCAGTCGGTCCCTCCAT-3') was designed to target the mRNA region spanning the first ATG, to block its translation. *MOSP1* was designed to bind to the acceptor splice site between intron 2 and exon 3 (5'-AGCAAATATCACACTCACCTTTC-3'). A total of 4 or 15 ng of the *MOatg* or *MOSP1* morpholinos, respectively, were injected into one-cell-stage embryos. Morpholino specificity was evaluated by detecting the levels of CTCF protein in the morphant embryos (Supplementary Fig. S5a,b). CTCF was detected with the chick CTCF antibody⁴⁸, which recognizes zebrafish CTCF⁴⁹. Western blots were performed as reported in ref. 49 using the anti-CTCF antibody diluted 1:500. To further examine the inhibitory action of *MOSP1* on CTCF mRNA splicing, we designed primers targeting exons 2 and 5 (5'-GGAAGAAGAAATGGCTGAACC-3' and 5'-GGCATAACTGCACAGACTGC-3'). These primers should amplify a 726-bp band only if the morpholino inhibits correct removal of intron 2; retention of this intron in the mRNA introduces several precocious stop codons (Supplementary Fig. S5c). For reverse transcription-PCR, total RNA was extracted at 48 h.p.f. from 25 morphants and control embryos, and amplification was carried out for 30 cycles.

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Author contributions

J.J.T. and E.d.I.C.-M. performed the *Xenopus* and zebrafish experiments. M.E.A. carried out the mouse transgenic assays. J.J.T. with the collaboration of E.S. and W.d.L. performed the 3C assays. J.L.G.-S. and M.M. designed the experiments and wrote the manuscript, and J.L.G.-S. conceived the work.

Additional information

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