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An ancient genomic regulatory block conserved across bilaterians and its dismantling in tetrapods by retrogene replacement

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Developmental genes are regulated by complex, distantly located *cis*-regulatory modules (CRMs), often forming genomic regulatory blocks (GRBs) that are conserved among vertebrates and among insects. We have investigated GRBs associated with *Iroquois* homeobox genes in 39 metazoans. Despite 600 million years of independent evolution, *Iroquois* genes are linked to ankyrin-repeat-containing *Sowah* genes in nearly all studied bilaterians. We show that *Iroquois*-specific CRMs populate the *Sowah* locus, suggesting that regulatory constraints underlie the maintenance of the *Iroquois*–*Sowah* syntenic block. Surprisingly, tetrapod *Sowah* orthologs are intronless and not associated with *Iroquois*; however, teleost and elephant shark data demonstrate that this is a derived feature, and that many *Iroquois*–CRMs were ancestrally located within *Sowah* introns. Retroposition, gene, and genome duplication have allowed selective elimination of *Sowah* exons from the *Iroquois* regulatory landscape while keeping associated CRMs, resulting in large associated gene deserts. These results highlight the importance of CRMs in imposing constraints to genome architecture, even across large phylogenetic distances, and of gene duplication-mediated genetic redundancy to disentangle these constraints, increasing genomic plasticity.

[Supplemental material is available for this article.]

The high complexity of transcriptional control in animals is known to have a significant impact on how their genomes are shaped through evolution. In particular, key developmental genes show an exceptionally complex regulation, with very specific and intricate spatio-temporal expression patterns. Numerous *cis*-regulatory modules (CRMs) create vast regulatory landscapes around these loci (Nelson et al. 2004) that often extend to neighboring genes. This imposes constraints on genomic restructuring, creating “solid” regions, recognizable as conserved regulatory blocks within phyla, with very low rates of rearrangement (Becker and Lenhard 2007; Engstrom et al. 2007; Kikuta et al. 2007). Nevertheless, it is unclear whether long-range *cis*-regulatory interactions may have had an impact on genome architecture over deeper evolutionary distances (i.e., across different phyla) (Koonin 2009).

Iroquois (*Iro/Irx*) genes encode highly conserved homeobox transcription factors (TFs) of the TALE class with multiple and fundamental roles in animal development (Cavodeassi et al. 2001). In addition, they provide arguably one of the most paradigmatic examples of the distinct evolutionary fate followed by genomic

regions surrounding major developmental regulators. First, *Irx* genes are flanked by large genomic regions devoid of genes (gene deserts) (Nobrega et al. 2003; Ovcharenko et al. 2005) in all studied bilaterians (Irimia et al. 2008). In vertebrates, as well as in other lineages, Conserved Noncoding Regions (CNRs) (Aparicio et al. 1995; Bejerano et al. 2004; Woolfe et al. 2005; Pennacchio et al. 2006) are highly enriched in these unusually long intergenic distances (Sandelin et al. 2004; de la Calle-Mustienes et al. 2005; Irimia et al. 2008; Tena et al. 2011), and they often act as tissue-specific enhancers (de la Calle-Mustienes et al. 2005; Tena et al. 2011). Second, *Irx* genes have independently evolved a cluster organization in at least four bilaterian groups (Irimia et al. 2008; Takatori et al. 2008; Kerner et al. 2009). Finally, *Irx* genes show the most widespread and ancient conserved linkage to a phylogenetically unrelated gene known to date: *Iroquois* and the ankyrin-repeat-containing *Sosondowah* (*Sowah*) genes are tightly associated in the genome in nearly all studied bilaterians, including fast-evolving species such as flies and nematodes, but with the intriguing exception of tetrapods (Supplemental Fig. S1; Irimia et al. 2008; Kerner et al. 2009).

Each of these features might be explained by the presence of strong regulatory constraints, such as shared enhancers or global control regions (de la Calle-Mustienes et al. 2005; Irimia et al. 2008). Consistently, functional studies in vertebrates and flies have shown that some regulatory elements are shared between clustered *Irx* genes (Gomez-Skarmeta et al. 1996; Tena et al. 2011), providing an explanation for the preservation of the cluster organization in these lineages, although functional data is still missing for other groups. In addition, no studies have yet provided an

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explanation for the striking linkage of *Irx* genes to *Sowah* in invertebrates and their dissociation in tetrapods. The strong conservation of synteny in invertebrates suggests that the *Sowah* loci could be part of the *Irx* regulatory landscape. Therefore, a full understanding of *Irx* regulation and evolution might not be complete without addressing this issue.

Here, we have studied the *Irx-Sowah* regulatory landscape in different animal groups. First, we show that *Sowah* genes are widely conserved, showing several deeply conserved intron positions and relative lengths, with high density of CNRs. Second, we demonstrate that noncoding sequences within the *Sowah* loci from amphioxus and *Drosophila* can act as transcriptional regulators, and are crucial for the proper expression of *Irx* genes, at least in *Drosophila*. Third, we show that despite the general absence of *Irx*-associated *Sowah* in vertebrates, several key *Irx* regulators in this lineage were originally located within a *Sowah* gene that specifically lost its coding sequence aided by the genetic redundancy produced by an early retroposition event. Finally, we discuss the impact of *Sowah* linkage on *Irx* regulation and evolution, showing how the recurrent remodeling of the *Sowah* loci has shaped the *Irx* regulatory landscape in different bilaterians.

Results

Irx-Sowah linkage and *Sowah* gene structure are highly conserved in metazoans

To investigate the extent of conservation of the *Irx-Sowah* linkage during bilaterian evolution, we identified and characterized *Sowah* genes in 39 genomes (Supplemental Figs. S1, S2; see Methods). *Irx* complexes are linked to *Sowah* genes in nearly all bilaterian lineages, with the exception of tetrapods, tunicates, and the leech *Helobdella robusta*. Alignment of intron/exon structures of metazoan *Sowah* genes revealed that several intron positions (ancestral introns 5–9, Supplemental Fig. S3) are conserved across all *Irx*-linked genes, including nematodes (except for intron 5) and insects, which are known to have dramatically divergent intron/exon structures (Rogozin et al. 2003; Coulombe-Huntington and Majewski 2007). In addition, the lengths of these introns are often exceptionally large, up to 70 times the species average (Supplemental Table S1). A possible explanation for the extraordinary conservation of these intron positions and relative lengths may be the presence of regulatory elements within them, which would prevent intron loss in diverse lineages (Irimia et al. 2011). In addition to a potential regulatory function, we found two groups of highly conserved microexons (one of 12 and four of three nucleotides) within the ankyrin-repeat domain (introns 6 and 7, Supplemental Fig. S3) in species with enough available expression data (including cnidarians, ecdysozoans, lophotrochozoans, and chordates).

Sowah loci are populated by CNRs that are likely associated with *Irx*

As potential indicators of regulatory sequences, we analyzed CNRs within *Sowah* loci in different groups. We performed interspecies comparisons within flies and nematodes, lineages with availability of an appropriate taxon sampling (i.e., within the phylogenetic ranges of CNR detection) (Boffelli et al. 2004). We first determined CNR density in the *Irx-Sowah* genomic region using Ancora (Engström et al. 2008). Major peaks of CNR density often overlap with developmental genes and can be used to delimit the regulatory landscapes associated with these genes (Engstrom et al. 2007,

2008). In both nematodes and flies, *Irx* genes were located in regions with very high CNR density. Importantly, the neighboring *Sowah* genes were fully embedded within these *Irx*-associated CNR-dense peaks, indicating that *Sowah* intronic sequences are enriched in CNRs as sequences immediately flanking *Irx* (Fig. 1A,B). Indeed, VISTA analyses of the *Sowah* regions using stringent parameters revealed dozens of CNRs within the long *Sowah* introns and in the intergenic regions between *Irx* and *Sowah*, consistent with previous analyses in nematodes (Vavouri et al. 2007) and with the high number of evolutionarily constrained elements detected by phastCons (Fig. 1A,B; Siepel et al. 2005). These results further emphasize the potential functional importance of *Sowah* intronic sequences.

In the case of the cephalochordate amphioxus, there are currently no sequenced genomes within the appropriate phylogenetic range to perform informative interspecies comparisons (Pascual-Anaya et al. 2008), and therefore Ancora and phastCons tools cannot be used in this lineage. However, VISTA analysis between paralogous regions have been successfully applied before in this species (Jiménez-Delgado et al. 2006; Irimia et al. 2008). We had previously identified several highly conserved noncoding sequences by crossed VISTA alignments between the regions surrounding the three *Irx* genes in the amphioxus cluster (Irimia et al. 2008). Here, we have expanded these analyses to include *Sowah* (Fig. 2). We divided the amphioxus cluster plus *Sowah* into four regions. Three contained one of the *Irx* genes (*IrxA*, *IrxB*, or *IrxC*), their surrounding noncoding sequences and, in the case of *IrxA*, the entire *Sowah* locus, which is immediately next to it. We also defined a fourth region corresponding to a putative “*IrxD*” locus that was lost during amphioxus evolution, as suggested by CNR complements after preliminary analyses (see below and Methods for further details). Crossed VISTA comparisons of these regions revealed many repeated blocks with high-sequence similarity that were not detected in our previous survey (Irimia et al. 2008). Importantly, these novel conserved repeated blocks have one of their copies lying either within *Sowah* introns or in the intergenic region between *Sowah* and *IrxA* (Fig. 2; Supplemental Fig. S4; Supplemental Data S1). Like the previously identified CNRs, most conserved elements are present in two copies, and some in three, indicating differential losses after the gene duplications that gave rise to the cluster (note that potentially functional single-copy sequences cannot be detected with this analysis). In addition, some CNRs were present in four copies, despite the fact that there are only three *Irx* genes. The implications of these findings are threefold.

First, it was not the ancestral *Irx* alone, but the pair *Irx-Sowah* that duplicated in tandem to generate the amphioxus cluster. In fact, remains of three exons from pseudogenized *Sowah* copies are still present next to *IrxB* and *IrxC*, as can be noticed in the VISTA plots (Fig. 2; Supplemental Fig. S4; Supplemental Data S1).

Second, the four-copy CNRs and the respective order and orientation of both genes and CNRs suggest that, in addition to the three *Irx* genes and their respective *Sowah* genes and pseudogenes, a fourth duplicate of the tandem *Irx-Sowah* may have been present in the cluster during amphioxus evolution. Despite the loss of an *IrxD* gene, this “*D*” paralogous region is clearly identifiable as a CNR array of sequences that were originally in the intronic and downstream regions of a *SowahD* locus (Fig. 2; Supplemental Fig. S4). Since CRMs located in gene clusters may target multiple paralogous genes (Tena et al. 2011), the CNRs may have been conserved due to interactions with other *Irx* genes, despite the loss of their original target gene. These results, together with data from phylogenetic analysis (Irimia et al. 2008) and CNR similarity (Supplemental Fig. S4)

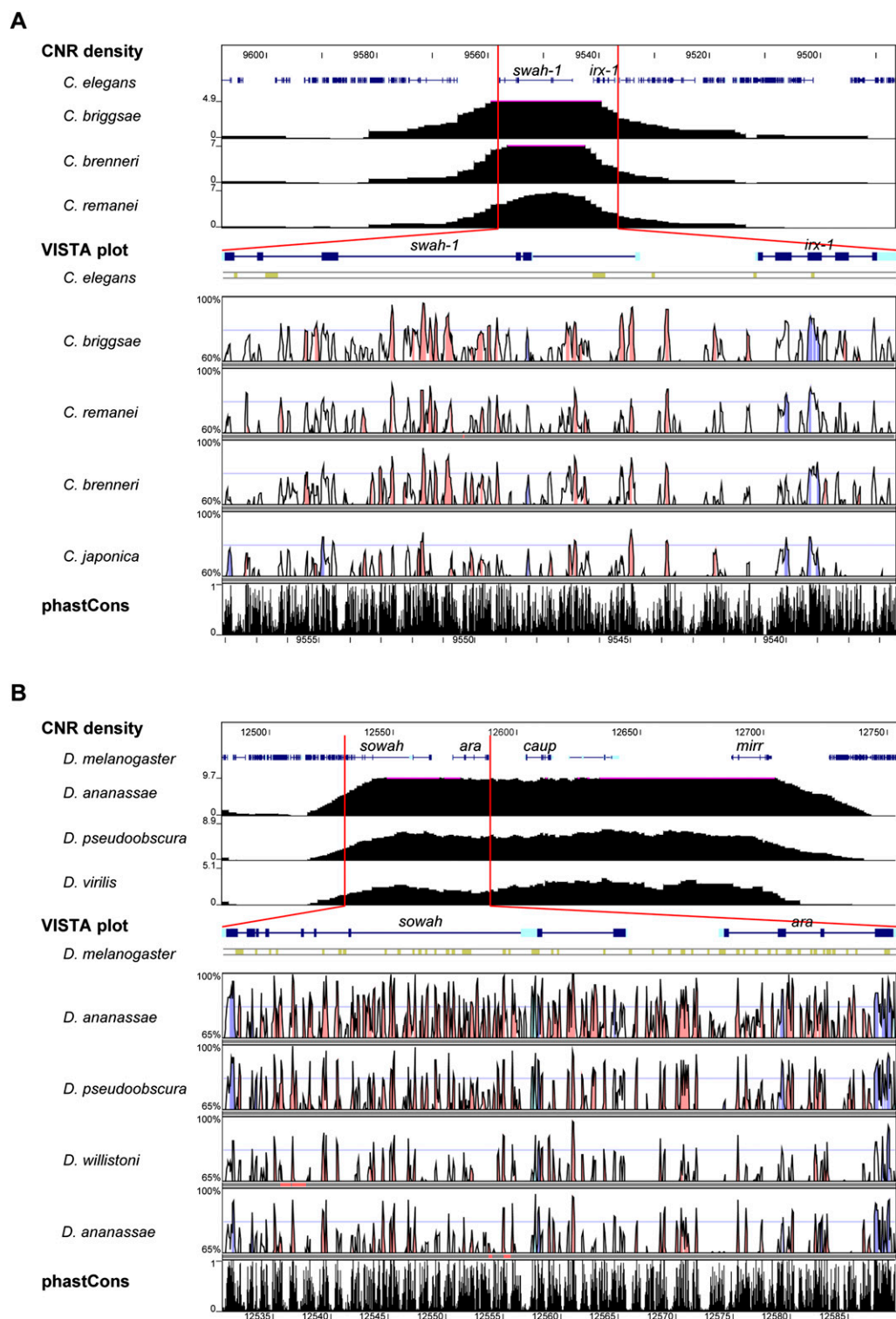


Figure 1. CNRs in nematode and drosophilid *Sowah* genes. Ancora plots of CNR density (top), VISTA plots (middle), and phastCons tracks (bottom) of the *Irx-Sowah* region of nematodes (A) and flies (B). (Red bars) Region depicted in Ancora plots zoomed in on VISTA and phastCons tracks. VISTA colored peaks (blue, coding; turquoise, UTR; pink, noncoding) indicate regions of at least 50 bp and $\geq 90\%$ similarity ($\geq 85\%$ in the case of *Caenorhabditis japonica*) in nematodes and 60 bp and $\geq 90\%$ similarity in flies. Only gene symbols corresponding to *Sowah* (*swah-1* in nematodes) and *Irx* (*ara*, *caup*, and *mirr* in *Drosophila*) are indicated. Numbers at the left correspond to the percentage of base pairs covered by CNRs in Ancora plots, percentage identity in VISTA analyses, and conservation scores in phastCons.

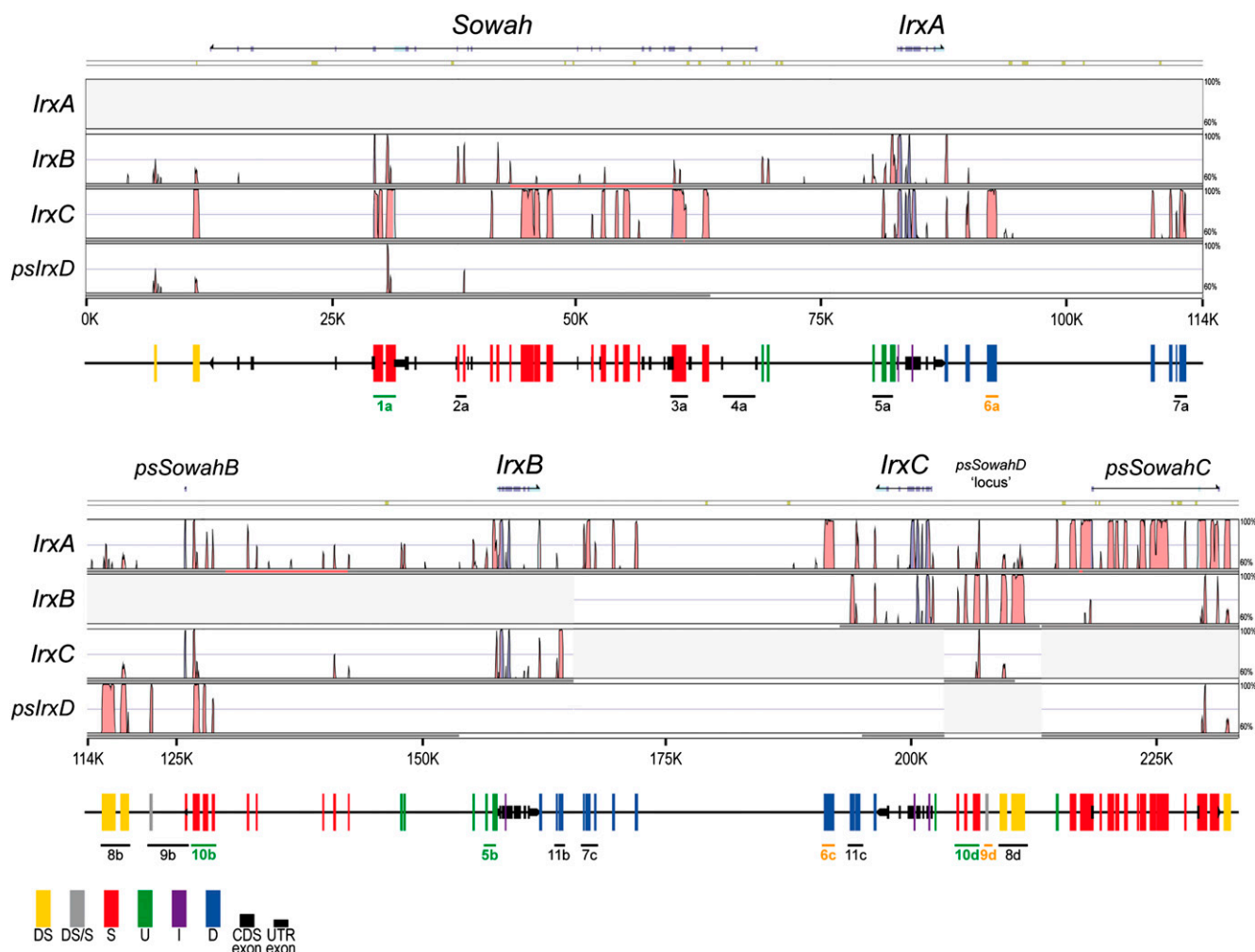


Figure 2. Internal organization of the *Irx-Sowah* complex in *B. floridae*. VISTA plot of the alignments between each of the three *Irx* genes (plus a fourth region corresponding to a putative *IrxD* locus lost during amphioxus evolution) and their respective surrounding noncoding regions, including *Sowah* in the case of *IrxA*. Colored peaks (blue, coding; turquoise, UTR; pink, noncoding) indicate regions of at least 100 bp and $\geq 70\%$ similarity. High-copy number elements (such as repeats and mobile elements) are masked and their presence is indicated by khaki segments above the VISTA plot. Vertical bars of different colors below the VISTA plot represent the different conserved blocks, indicating their respective location to *Sowah* and *Irx* loci (DS, Downstream *Sowah*; S, within *Sowah*; U, Upstream *Irx*; I, Introns of *Irx*; D, Downstream *Irx*. DS/S bars indicate elements of uncertain identity [DS or S]). Black rectangles and arrows indicate the exon sequences of *Irx*, *Sowah*, or the remains of *Sowah* duplicates. The blocks tested for transcriptional enhancer activity are indicated and named with a number and letter code. The letter refers to the *Irx* locus with which they are associated; color indicates whether they are tissue-specific enhancers (green), unspecific enhancers (yellow), or negative elements (black).

indicating that A duplicate is more closely related to C, and that B is more similar to D, provide a more detailed picture of the evolutionary origin of the amphioxus *Irx* cluster (Supplemental Fig. S5; Supplemental Discussion S1).

Third, the selective loss of *Sowah* exons versus their intronic conserved elements suggests that the CNRs in the *Irx-Sowah* genomic region are most likely associated with *Irx*, not with *Sowah*, providing a potential explanation for the conservation of the *Iroquois-Sowah* syntenic block. Adding to this idea, the expression pattern of amphioxus *Sowah* differs widely from those of *Irx* genes (Fig. 3A–D; Kaltenbach et al. 2009; Irimia et al. 2010), suggesting distinct regulations, and thus, that *Sowah* and *Irx* genes are not likely to share major transcriptional enhancers. A comparable situation is found in *Drosophila*, where the expression pattern of *Sowah* is not similar to that of *Irx* complex genes, *araucan* (*ara*), *caupolican* (*caup*), and *mirror* (*mirr*) (Gomez-Skarmeta et al. 1996; McNeill et al. 1997), indicating

that *Irx* and *Sowah* genes are probably not coregulated (Fig. 3E–H; data not shown).

Conserved sequences from the amphioxus *Irx-Sowah* locus drive expression consistent with that of *Irx* in zebrafish stable transgenic lines

Next, we tested the functionality of the conserved sequences within the amphioxus *Irx-Sowah* locus. We selected 18 amphioxus CNRs (Supplemental Table S2) from different locations: (1) between two *Irx* genes, (2) in the intergenic region between *Irx* and *Sowah*, and (3) within *Sowah* introns (Fig. 2). The potential regulatory activity of each of the 18 CNRs was assayed by generating stable transgenic zebrafish lines. To this end, we used the ZED vector (Bessa et al. 2009), which contains the *gata2a* minimal promoter with a GFP reporter gene, and RFP driven by the cardiac

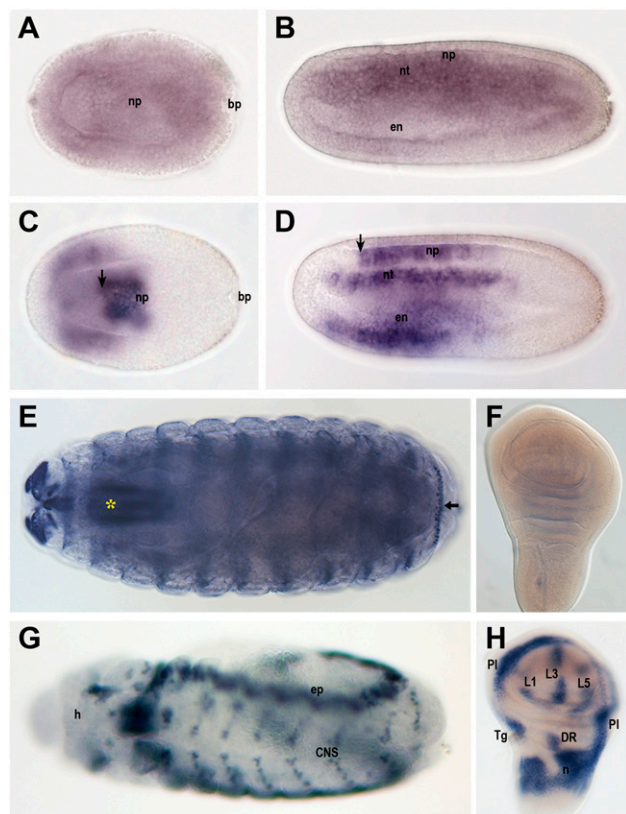


Figure 3. Comparison of the expression patterns of *Sowah* and *Iroquois* genes in amphioxus and fly. In situ hybridization of *B. lanceolatum* *Sowah* (A,B) and *IrxB* (C,D) genes in 15-h early neurulas (A,C) and 21-h neurulas (B,D) in dorsal and lateral views, respectively. *Sowah* transcripts were detected almost ubiquitously, with stronger expression in the dorsal half of the embryos. In contrast, *IrxB* showed a very defined and restricted pattern in the endoderm (en), notochord (nt), and neural plate (np). The anterior limit of expression in the neural plate, which is conserved in evolution (Irimia et al. 2010), is indicated by an arrow. The expression of *IrxA* and *IrxC* was similar at these stages (data not shown). (bp) Blastopore. In situ hybridization of *sowah* (E,F) and *caup* (G,H) in *D. melanogaster* stage 17 (E) and stage 12 (G) embryos (dorsal views) and third instar larvae wing imaginal discs (F,H), anterior is to the left). (E,F) *sowah* is expressed in the pharynx (*) and cephalic nervous system of late embryos (arrow in E points to nonspecific staining of the cuticular denticle belts), but is undetectable in the wing imaginal discs. (G,H) During embryonic development, *ara* and *caup* display coincident dynamic expression patterns in the epidermis (ep), central nervous system (CNS) and mesoderm, as well as in the head (h). In the wing disc, *caup* is expressed in the prospective regions of the 1, 3, and 5 longitudinal veins (L1, L3, L5), pleura (Pl); tegula (Tg); dorsal radius (DR); alula and lateral notum (n).

actin promoter as positive control for successful transgenesis. Four of the amphioxus CNRs (1a, 5b, 10b, and 10d [letters indicate the associated *Ir*x loci]) drove reproducible tissue-specific GFP expression in zebrafish embryos in at least three independent founders (Fig. 4; Supplemental Figs. S6, S7). Three other elements (6a, 6c, and 9d) drove GFP expression in a ubiquitous or variable way, suggesting that they may be less-specific enhancers (Supplemental Fig. S6; data not shown; Komisarczuk et al. 2009). No GFP expression could be detected for the other 11 tested elements, for which cardiac and muscle expression of the control RFP was detected in several founders studied for each element (data not shown). These transcriptionally inactive CNRs may not have been recognized by zebrafish TFs or they could be active in developmental stages or

tissues not surveyed in this work. Alternatively, they may be involved in negative transcriptional regulation or other regulatory functions that could not be detected in this assay.

Of the four tissue-specific enhancers, three are located within the introns of *Sowah* or its pseudogenized copies (two of them, 10b and 10d, are copies of a duplicated element), whereas the other (5b) lies between *IrxB* and its respective pseudogenized *Sowah* gene (Fig. 2). The duplicated elements, 10b and 10d, drove similar expression in the central nervous system (CNS) (Fig. 4), in a pattern consistent with *Ir*x genes in both amphioxus (Fig. 3D; Kaltenbach et al. 2009) and zebrafish (Lecaudey et al. 2001; Feijóo et al. 2004). Strikingly, this expression is reminiscent of previously characterized *Ir*x cis-regulatory elements in vertebrates (de la Calle-Mustienes et al. 2005; Visel et al. 2007; Tena et al. 2011), which are also active in wide CNS domains, including anterior regions normally devoid of *Ir*x expression. The CNR 5b drove GFP expression to the spinal cord and the telencephalon (Fig. 4). Again, while the first tissue is a major domain of *Ir*x expression in amphioxus, the second one shows no expression of these genes (Kaltenbach et al. 2009). As it has been proposed in vertebrates (de la Calle-Mustienes et al. 2005), it is likely that other negative cis-regulatory elements contribute to the down-regulation of *Ir*x expression in the telencephalon. Finally, the intronic element 1a was consistently active in a highly restricted domain, the blood islands (Supplemental Fig. S6), a “tissue” with no obvious known counterpart in amphioxus. To date, *Ir*x genes have not been implicated in hematopoiesis in vertebrates (Ferrell et al. 2005), and therefore, it is not clear whether the regulatory activity

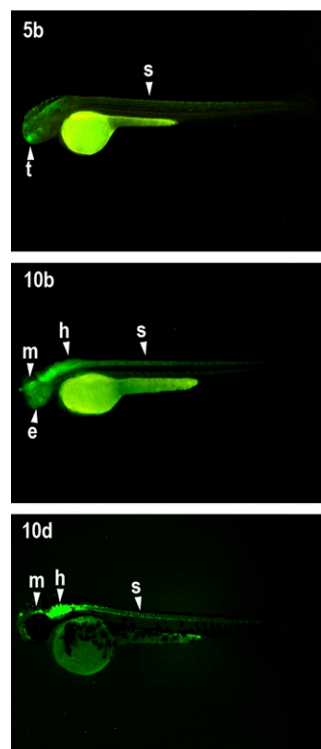


Figure 4. Transcriptional enhancer activity of *B. floridae* sequences from the *Ir*x-*Sowah* complex. Lateral views of 48-hpf zebrafish showing GFP expression driven by the 5b, 10b, and 10d CNRs. The 5b-driven expression is detected in the spinal cord and in the telencephalon; 10b and 10d consistently drove expression throughout the CNS (midbrain, hindbrain, and spinal cord) and in the eye. Anterior is to the right. (e) eye; (h) hindbrain; (m) midbrain; (s) spinal cord; (t) telencephalon.

of element 1a is compatible with either the expression of amphioxus *Irx* or *Sowah* genes, or whether this sequence is being read by different regulatory states in vertebrates and amphioxus. This last alternative illustrates the complexity of interpreting trans-species reporter assays at deep evolutionary distances, where the possibility of obtaining reproducible but biologically meaningless results cannot be completely ruled out. Nevertheless, very deep conservation of regulatory states does exist, even across phyla (Royo et al. 2011), and consistent results such as those observed for elements 10b, 10d, and 5b are thus highly suggestive of functional conservation.

Sowah contains regulatory elements essential for proper Iroquois expression in flies

The tissue-specific expression driven by some amphioxus *Sowah*-derived CNRs in zebrafish embryos, which encompasses *Irx* expression domains, suggests that these elements may be regulating *Irx* transcription. To obtain direct evidence that *Sowah* noncoding sequences were responsible for part of the transcriptional regulation of *Irx* genes, we then resorted to *Drosophila*, where the appropriate genetic tools are available. The expression of *Drosophila* members of the *Iro* family has been extensively characterized. *ara* and *caup* show identical expression patterns, including specific wing and leg imaginal disc regions and several embryonic domains (Figs. 3G,H, 5B,E,H,H',J; Gomez-Skarmeta et al. 1996; Diez del Corral et al. 1999; Letizia et al. 2007; Carrasco-Rando et al. 2011), suggesting that *ara* and *caup* share specific CRMs (Gomez-Skarmeta et al. 1996; Letizia et al. 2007). In contrast, *mirr* expression pattern is slightly different (McNeill et al. 1997; Kehl et al. 1998).

CRMs responsible for *ara/caup* expression in the prospective notum region of the wing disc are located between *caup* and *mirr* (Letizia et al. 2007). On the contrary, no CRMs accounting for the remaining expression domains of the wing disc have been characterized so far. Interestingly, genetic data suggested the presence of L3 vein-specific CRMs upstream of *ara* (Gomez-Skarmeta et al. 1996), consistent with a putative location within the *sowah* locus. To find these putative wing CRMs we generated three molecularly defined deletions (*sowah*^{EGP1}, *sowah*^{EGP2}, and *sowah*^{EGP3}) (Fig. 5A) and monitored the expression of *ara*, *caup*, and *mirr* in the wing discs by in situ hybridization (Fig. 5B–E). In every mutant condition examined, *ara* and *caup* expressions were undistinguishable from each other, and thus, only *caup* expression is shown; *mirr* expression was not affected by any of the deletions (data not shown).

sowah^{EGP2} harbors a deletion spanning the central area of *sowah*, including its longest introns (ancestral introns 5 and 6) (Fig. 5A). In these flies, *ara/caup* expression patterns were dramatically affected,

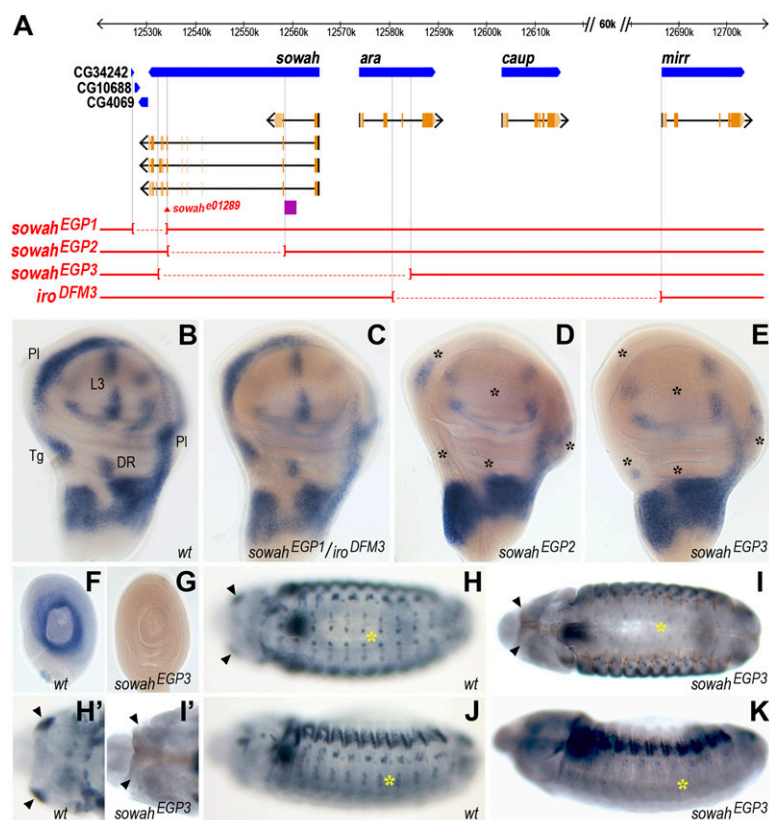


Figure 5. Expression of *caup* in *sowah*^{EGP} imaginal discs and embryos. (A) Physical map of the Iro-C locus. Genomic DNA is shown as a thick black bar with a 60-kb gap delimited by // . Transcripts are shown as black arrows below the genes (blue). Exons are shown in orange, with protein-coding regions colored darker. Red broken lines within brackets represent deleted regions. The purple box represents a region bound by several transcription factors as determined by ChIP-on-chip assays. (B–E) In situ hybridization with a *caup* probe of wing (B–E) and leg (F,G) discs of the indicated genotype. *caup* expression is not affected in the *trans*-heterozygous combination of *sowah*^{EGP1} and *iro*^{DFM3}, used to rescue the early embryonic lethality of *sowah*^{EGP1}. In *sowah*^{EGP2} (D) and *sowah*^{EGP3} (E) discs, *caup* expression is absent in L3 and DR domains and strongly reduced in PI and Tg regions (marked with black asterisks). (G) *sowah*^{EGP3} leg disc, in which *caup* wild-type expression in a ring-like pattern (F) is lost. (H–K) Anti-Caup staining of wild-type (H,H',J) and *sowah*^{EGP3} (I,I',K) late-stage 13 embryos. Yellow asterisks mark the expression of *ara/caup* in the nervous system; arrowheads point to the head mandibular segment. The brown signal in I corresponds to a nonspecific staining of the tracheal (respiratory) system. (H,I) ventral, (J,K) lateral views, (H',I') enlarged views of the head.

lacking the expression from the L3 vein, pleura (PI), tegula (Tg), and dorsal radius (DR) domains of the wing disc (black asterisks in Fig. 5D). This suggests that CRM(s) driving *ara/caup* expression in these territories are located within this region, most likely within the long introns. Consistently, the *sowah*^{DSO10} mutation—associated with the insertion of an insulator-containing transposon in the second intron of *sowah*—shows a similarly altered expression of *ara/caup* in the wing discs (Supplemental Fig. S8). The presence of this insulator is likely to prevent the interaction of CRM(s) located upstream of the insertion point with the downstream *ara* and *caup* promoters. Importantly, *sowah* transcripts are undetectable in wild-type imaginal discs (Fig. 3F), suggesting that the loss of *ara/caup* expression is not due to *sowah* loss of function. This notion is further supported by the fact that *sowah* exons are unaffected in *sowah*^{DSO10} flies; however, we cannot completely rule out that the insertion of a transposon within the intronic sequence could have mildly affected *sowah* activity.

sowah^{EGP3} has a larger deletion than *sowah*^{EGP2}. It encompasses the region deleted in *sowah*^{EGP2} and spans from the ancestral

ninth intron of *sowah* to the third intron of *ara* (Fig. 5A). Remarkably, in addition to the domains of *ara/caup* expression lost in *sowah*^{EGP2} and *sowah*^{EGP3} (black asterisks in Fig. 5E), the *sowah*^{EGP3} deficiency also altered the expression of *Iro* genes in leg discs (Fig. 5G), and in some domains of the embryonic head (arrowheads in Fig. 5I,I') and nervous system (yellow asterisks in Fig. 5I,K). This suggests that CRM(s) responsible for these embryonic expression domains and for the "ring" pattern of the leg discs are located somewhere in the intergenic region between *ara* and *sowah* and/or in the first two introns of *sowah*. Noteworthy, ChIP-on-chip experiments in embryos (<http://genome.ucsc.edu/>) show that several TFs bind to a region in *Drosophila sowah* intron 2 (ancestral intron 3, purple box in Fig. 5A), suggesting that this intron could contain the CRM(s) that drive *ara/caup* expression in the head and nervous system.

In *sowah*^{EGP1}, which lacks most of the *sowah* coding region (but not the longest introns) and the three CGs immediately distal to *sowah* (CG34242, CG10688, and CG4069) (Fig. 5A), and in *sowah*^{e01289} flies, which harbor a transposon inserted in the *sowah* exon 7 (Fig. 5A), *ara/caup* expression was unmodified (Fig. 5C; data not shown). This further reinforces the idea that the changes in the expression of *ara/caup* are not related to the absence of *sowah*

function. Thus, our results show conclusively that a substantial part of the complement of CRMs of the *Iro* family lies within the *sowah* locus, suggesting that long-range *cis*-regulatory interactions are the major constraint preventing the disruption of the *Iroquois*–*Sowah* genomic block, at least in flies.

Sowah gene complements, intron–exon structures, and synteny in vertebrate genomes

Despite the strong linkage constraint and intron position conservation in other bilaterian lineages, the four *Sowah* orthologs in the human genome (*SOWAHA*, *SOWAHB*, *SOWAHC*, *SOWAHD*, also known as *ANKRD43*, *ANKRD56*, *ANKRD57*, and *ANKRD58*) are not associated with *Irx* complexes and are intronless. Instead, they are all linked to *Septin* genes (*SEPT8*, *SEPT11*, *SEPT10*, and *SEPT6*, respectively), in a head-to-head orientation (Fig. 6A). This suggests that the human *Sowah* genes originated by a single retroposition event into a new genomic location that occurred before the two ancient rounds of whole-genome duplication (WGD) (Dehal and Boore 2005; Putnam et al. 2008). Therefore, we collectively refer to them as r-*Sowah* genes hereafter (the letters A–D indicate their retroposition origin, in contrast to the canonical intron-containing

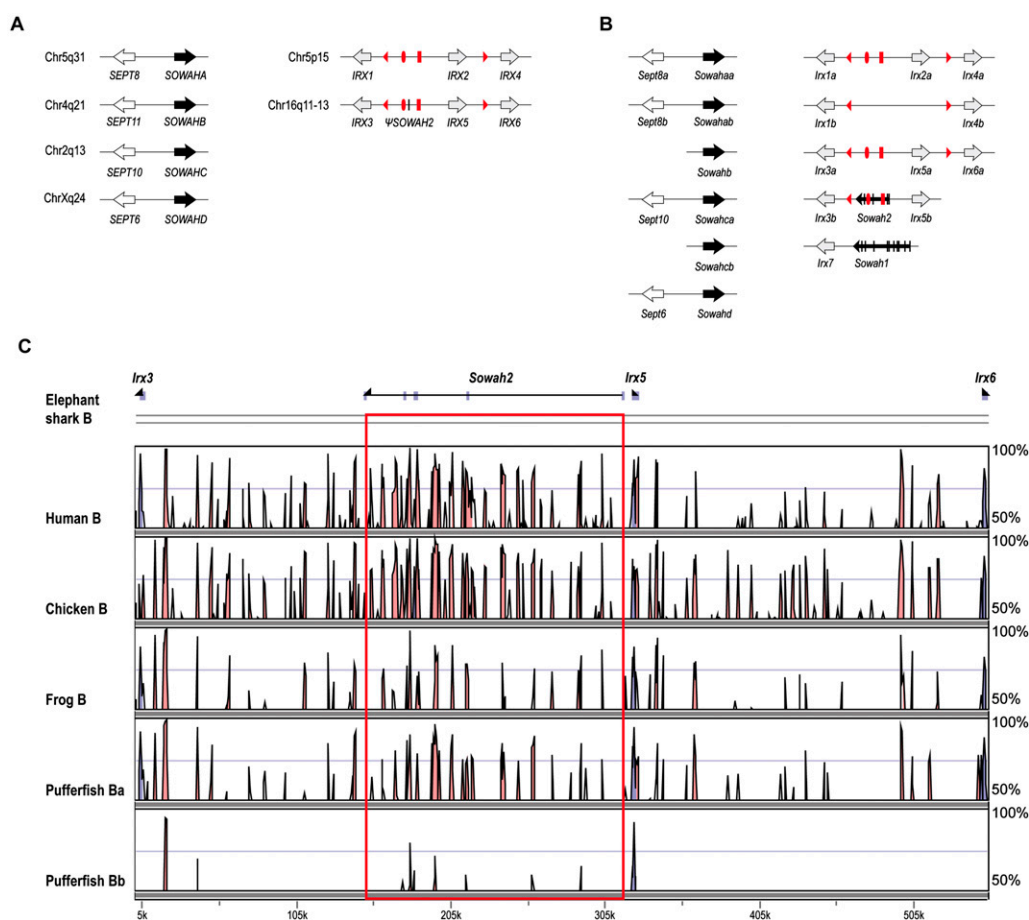


Figure 6. Genomic organization of *Sowah* and *Irx* genes in vertebrates. Schematic representation of the genomic organization of r-*Sowah* and *Sowah* (black arrows), *Septin* (white arrows), and *Irx* (gray arrows) genes in humans (A) and a generalized teleost (B). Red geometrical shapes represent CNRs within *Irx* clusters: triangles represent the UltraConserved Regions (UCRs) (de la Calle-Mustienes et al. 2005), and ellipses and rectangles the only two CNRs present within *Sowah2* in teleosts. For simplicity, only schematic intron–exon structures are indicated for *Sowah1* and *Sowah2*. (C) VISTA plot of the alignments between *IrxB* clusters of different vertebrate species, using elephant shark as a reference sequence for the comparison. Colored peaks (blue, coding; turquoise, UTR; pink, noncoding) indicate regions of at least 100 bp and $\geq 70\%$ similarity. *Sowah2*-CNRs are demarcated by a red rectangle.

sowah1-2 genes in fish, see below). The same pattern was found in all tetrapod species examined (data not shown). However, in the teleost fish fugu and zebrafish, in addition to six *Septin*-associated intronless copies, two “canonical”, intron-containing *Sowah* genes are still linked to *Irx* (Fig. 6B; Supplemental Fig. S9). They are linked to *Irx7* (see Supplemental Discussion S2 for the potential implications on the debated *Irx7* orthology) (Lecaudey et al. 2001, 2005; Itoh et al. 2002; Dildrop and Rütther 2004; Feijóo et al. 2004) and *Irx5b*, and we termed them, respectively, *Sowah1* (LOC100331692-LOC100148636 in zebrafish) and *Sowah2* (previously unannotated and very divergent, since the first four to five and the last two to three ancestral exons could not be detected in any available teleost genome, neither by sequence conservation nor using expression data). Thus, teleosts have partially retained the *Irx*–*Sowah* linkage typical of invertebrate animal lineages.

To further explore the early evolution of the *Irx*–*Sowah* linkage in vertebrates, we searched the available genomic contigs of a basal jawed vertebrate, the elephant shark (*Callorhynchus milii*), a species with a slowly evolving genome (Venkatesh et al. 2006; Ravi et al. 2009). We found three intronless r-*Sowah* genes (Supplemental Fig. S9), but as in the case of teleost fish, we also identified fragments of at least two intron-containing *Sowah* genes. However, the highly fragmented nature of the available assembly impeded further synteny analyses and gene annotation. We thus screened an elephant shark genomic BAC library and obtained the full sequence of the *IrxB* cluster, which contains a *Sowah2* gene, as in teleosts. The elephant shark *IrxB* cluster locus contains the three *Irx* genes (*Irx3*, *Irx5*, and *Irx6*) and their immediate upstream and downstream flanking genes (*Fto* and *Mmp2*) (Supplemental Fig. S10), with a total cluster length of 553 kb (similar to zebrafish *IrxBa* locus and shorter than those of tetrapods, Supplemental Fig. S11). The synteny of *Irx* and *Fto* and *Mmp2* genes is conserved in the zebrafish *IrxBa* locus and all sequenced tetrapod *IrxB* loci (Supplemental Fig. S11). As predicted, we found a *Sowah2* ortholog in the intergenic region between *Irx3* and *Irx5*, in the same orientation as teleosts, and corresponding to some of the fragments previously identified in the blast searches. In addition to exons 5–9 of *Sowah2* gene, we could also identify the first exon, indicating that the elephant shark *Sowah2* gene is more conserved than its teleost counterparts. Nevertheless, as in teleosts, the remaining ancestral exons were either lost or too divergent, raising the question about the functional status of *Sowah2* genes. First, *Sowah2* genes were transcriptionally active in at least zebrafish and medaka, as evidenced by the presence of Expressed Sequence Tags (five and two ESTs, respectively), and RNAseq data from different zebrafish tissues (see Methods). Second, if still active as protein-coding genes, *Sowah2* exonic sequences are expected to reflect the action of selective pressures. Thus, as an indicator of ongoing selection, we estimated the ratio between non-synonymous (d_N) and synonymous (d_S) nucleotide substitution rates (d_N/d_S or ω) for those exons that could be retrieved in at least four different species (exons 5, 7, 8, and 9). In all cases, ω values were very low ($\omega < 0.12$), indicating active purifying selection (Supplemental Fig. S12). These data suggest that, although divergent, *Sowah2* is likely a functional protein-coding gene in teleosts and elephant shark.

Finally, we characterized the expression of *Sowah* genes in zebrafish. As in invertebrate lineages, zebrafish *sowah1* and r-*sowah* genes showed ubiquitous and weak expression that was, in some cases, indistinguishable from the in situ hybridization background (data not shown), although expression of all of the genes was detectable by qRT–PCR (Supplemental Fig. S13). Thus, zebrafish *sowah* expression also differ extensively from the complex expression

patterns of zebrafish *Irx* genes (Feijóo et al. 2004; Lecaudey et al. 2005), suggesting that, regardless of their genomic location, vertebrate *Sowah* genes are not coregulated with *Irx*.

Elephant shark and teleost *Irx*-linked *Sowah* gene CNRs include *cis*-regulators of *Irx* in tetrapods

The identification of exons of *Sowah2* gene in elephant shark and teleosts suggests that *Sowah2* coding sequences in tetrapods have undergone a process of pseudogenization/loss. VISTA analysis with elephant shark as a reference allowed us to confirm this hypothesis: Highly divergent, pseudogenized exonic remnants are detectable in the chicken, anole lizard, and several mammalian genomes, including human (blue circles in Supplemental Fig. S12). The presence of *Sowah2* “pseudoexons” across deeply diverged tetrapod lineages was surprising, since nonfunctional pseudogenetic sequences are expected to decay relatively rapidly, and could be suggestive of (ancestral) recruitment of these exons into a *cis*-regulatory role as CNRs (Dong et al. 2010; Eichenlaub and Ettwiller 2011). However, the degree of degeneration of these pseudogenized remnants is very variable across lineages, with a patchy distribution of pseudoexon presence. For example, three exons (7, 8, and 9) are still clearly identifiable in the chicken genome, and the only evidence of pseudogenization is the lack of splice sites, indicating a very recent inactivation. In contrast, no traces are detectable in the amphibian *Xenopus tropicalis*, and only one exon is present in lizard (exon 9) and placental mammals (exon 7), in both cases containing frameshift and splice-site mutations. Marsupials present an intermediate situation, with two identifiable exons (7 and 9) in the genome of *Monodelphis domestica*. We then checked for the presence of epigenetic marks indicative of regulatory enhancer activity in species with available data. Deposition of histone marks was not consistent with enhancer activity in any species, including human cell lines and mouse embryos, as well as *sowah2* protein-coding exons in zebrafish embryos (Supplemental Fig. S12; data not shown). In support of this idea, we found no evidence for evolutionary constraint when comparing pseudoexon sequences across closely related species (e.g., within primates, Supplemental Fig. S12). Taken together, these results suggest that the presence of recognizable orthologous pseudogenetic exons in deeply diverged tetrapod lineages is probably due to independent and recent *Sowah2* inactivation events, and not due to functional conservation. Nevertheless, we cannot rule out that *Sowah2* remnants could have *cis*-regulatory roles in some lineages or developmental stages.

On the contrary, the scenario for noncoding elements within *Sowah* was completely different: The majority of the CNRs from the *IrxB* cluster were ancestrally located within the regions orthologous to elephant shark *Sowah2* introns (Fig. 6C). The elephant shark *Sowah2* locus extends over 167 kb, accounting for >50% of the “gene desert” that separates *Irx3* and *Irx5* in vertebrates (de la Calle-Mustienes et al. 2005), and almost one-third of the total cluster length (553 kb). This *Sowah2* region contains 58% (47 out of 81) of all *IrxB* cluster CNRs (between elephant shark and human, Supplemental Table S3), and 71% of those between *Irx3* and *Irx5*. In addition, four *Sowah2* CNRs are present in both *IrxA* and *IrxB* paralogous clusters (McEwen et al. 2006), suggesting that *Sowah* CNRs predate the WGD event that generated the two vertebrate *Irx* clusters, and that a *Sowah* gene was also originally present in the *IrxA* complex.

Interestingly, CNRs from both *IrxA* and *IrxB* clusters have been extensively studied and functionally characterized (de la Calle-Mustienes et al. 2005; Tena et al. 2011), including some of

the elements that are now recognized as originally located within *Sowah* loci. Nine of these *Sowah*-CNRs (three of them duplicated elements present in both clusters) (McEwen et al. 2006) show transcriptional enhancer activity when assayed in both *Xenopus* and zebrafish (de la Calle-Mustienes et al. 2005; Tena et al. 2011). These nine CNRs drove consistent tissue-specific expression in subdomains of the endogenous *Irx* expression patterns (Fig. 7; de la Calle-Mustienes et al. 2005; Tena et al. 2011), and 3C experiments showed that at least one of these elements specifically interacts with the promoters of both *Irx1* and *Irx2* in the *IrxA* cluster (element 3240) (Tena et al. 2011). These results demonstrate that *cis*-regulatory elements that were originally located within a *Sowah* intron are unequivocally *Irx* transcriptional regulators in vertebrates and have been maintained despite the loss of the associated *Sowah* coding regions.

Discussion

CRMs controlling developmental genes extend along enormous distances, often populating unrelated neighboring transcriptional units (bystander genes) (Engstrom et al. 2007; Kikuta et al. 2007). These intricate and intermingled genomic domains containing a developmental gene and its CRM array encompassing one or several bystander genes are typically known as genomic regulatory blocks (GRBs) (Becker and Lenhard 2007; Kikuta et al. 2007) and create evolutionarily “solid” regions of conserved microsynteny. The presence of conserved GRBs had been previously described only within vertebrates or within insects (Engstrom et al. 2007;

Kikuta et al. 2007), but none of them was shown to be conserved between the two lineages. Here, we describe the most ancient and widely conserved GRB, present since the origin of bilaterians. We show that regulatory elements present in the introns of the bystander gene *Sowah* are likely to regulate *Irx* in flies, amphioxus, and vertebrates, providing a general explanation for the maintenance of this GRB over 600 MY of independent evolution in several bilaterian lineages. Although we could not find any detectable sequence similarity between the *Sowah* noncoding regions and CRMs of different phyla (an extremely rare phenomenon) (Royo et al. 2011), most validated *Sowah* enhancers were located in a common set of conserved and exceptionally long introns (introns 5–9), suggesting the possibility of an ancestral regulatory system for this GRB in all bilaterians. In addition, these results show that GRBs, as a particular type of functional genomic organization, originated in a common ancestor of bilaterians, and that the evolution of complex *cis*-regulatory systems could have been determining genome remodeling processes since the origin of metazoans. Although the *Irx*-*Sowah* syntenic block constitutes the first example, this may well be the tip of the iceberg, and global approaches may reveal a wealth of ancient GRBs.

Intriguingly, despite this striking syntenic conservation, a few lineages evolved ways to disentangle the regulatory constraints and break *Irx* and *Sowah* apart. We found three different evolutionary paths, all of them involving the creation of genetic redundancy, resulting in the recurrent specific loss of the coding region of the bystander gene, while keeping the intervening CRMs (Fig. 8) (a loss process that could be a general mechanism for the formation of gene

deserts, whose origins remain largely mysterious). First, in early vertebrates, a *Sowah* retroposition event facilitated the disruption of the *Irx*-*Sowah* linkage by allowing the specific loss of the parental, *Irx*-linked coding sequence, and maintenance of the functional noncoding elements. This is, to our knowledge, the first example in which a retroposition event can be related to the genomic restructuring and loss of its parental locus and to the dismantlement of functionally constrained genomic linkages. Second, in the teleost lineage, the extra *Irx* clusters provide an example of the most classical and commonly reported case of disruption of synteny and GRBs after WGDs (Kikuta et al. 2007). Finally, the amphioxus case illustrates a third mechanism by which the coding regions of bystander genes containing CRMs can be lost: tandem duplication. Interestingly, the only other lineages in which *Sowah* is not linked to *Irx* genes—the tunicates and the leech *Helobdella robusta*—have multiple and genomically disperse *Irx* duplicates (Supplemental Fig. S1), suggesting that similar processes of gene duplication helped to disrupt the GRBs in these species.

These processes illustrate the high degree of mutual interdependence between the extremely complex transcriptional regulation of development and genome architecture through bilaterian evolution.

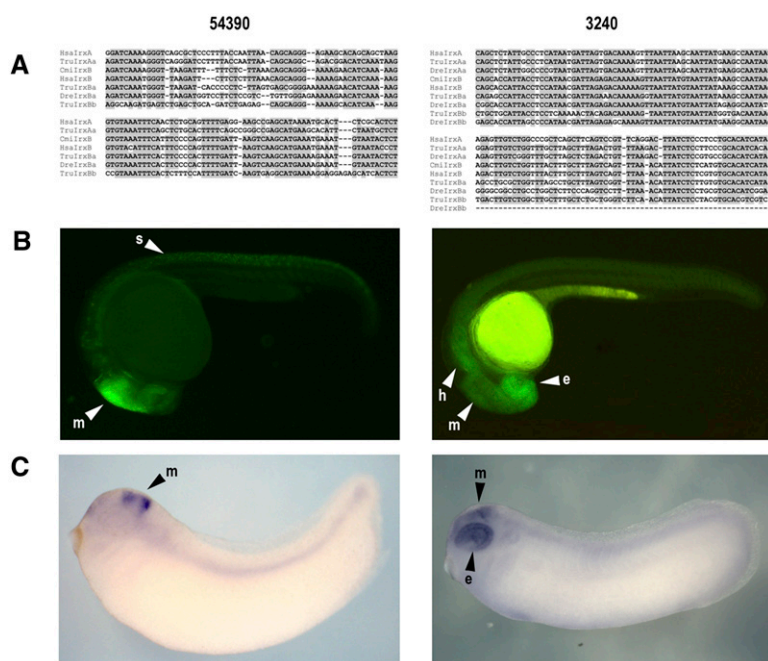


Figure 7. *Sowah* intronic CNRs. Two of the *Sowah2* pre-WGD CNRs that function as tissue-specific enhancers in reporter assays (represented by red ovals and rectangles in Fig. 6). (Left) CNR 54390 (*Sowah2* intron 7); (right) CNR 3240 (*Sowah2* intron 8). (A) Sequence alignment of the 54390 and 3240 *Sowah*-CNRs in different *Irx* complexes of several species. Shadowed nucleotides correspond to >60% sequence conservation. (B,C) GFP expression driven by the elements 54390 and 3240. Paralogous sequences of the element 54390 in complexes *IrxA* and *IrxB* drove similar expression patterns in *Xenopus* embryos (C) and zebrafish (B) transgenic lines. In the case of the CNR 3240, only that present in the *IrxA* complex was found to be positive in transgenesis studies. (e) Eye; (m) midbrain; (h) hindbrain; (s) spinal cord.

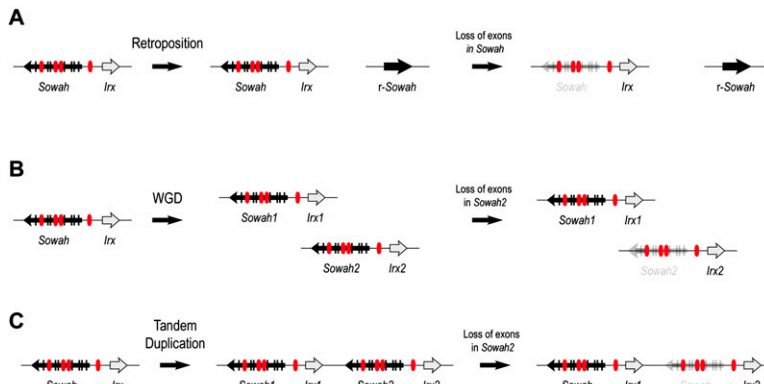


Figure 8. Evolutionary scenarios for the convergent loss of *Sowah* coding exons near *Irx* genes in chordates. (A) A retroposition event to other parts of the genome (indicated as a solid black arrow, r-*Sowah*) allows the original, *Irx*-linked *Sowah* to lose the coding sequences (black bars), while retaining the functional noncoding regions (red). A similar event occurred at the base of the vertebrates. (B) A polyploidization (a WGD) creates redundancy of *Sowah* genes. Therefore, some *Sowah* genes can lose their coding sequences. This event was observed in teleosts, after the third round of WGD. (C) Gene redundancy is acquired by tandem duplication of *Irx* and *Sowah*, as reported for amphioxus. Subsequently, one of the *Sowah* copies loses its coding sequences, whereas the functional noncoding regions are maintained.

Methods

Search for *Sowah* genes in metazoan genomes

In species where *Sowah* genes were not previously described or available gene predictions were fragmentary or poorly annotated, we built new manually curated predictions as described before (D’Aniello et al. 2008). We used the following databases: v1.0 of the genomes of *Trichoplax adhaerens*, *Nematostella vectensis*, *Branchiostoma floridae*, *Lottia gigantea*, *Capitella teleta*, and *H. robusta*, v1.0 and v2.0 of *Ciona intestinalis*, v4.1 of *Xenopus tropicalis*, and v4.0 of *Takifugu rubripes* from JGI (http://genome.jgi-psf.org/euk_home.html), *Homo sapiens* Build37.1, *Gallus gallus* Build2.1, *Danio rerio* Zv8 and Zv7, *Strongylocentrotus purpuratus* Build2.1, *Anopheles gambiae* AgamP3.3, *Apis mellifera* Amel4.5, *Nasonia vitripennis* Build1.1, and *Tribolium castaneum* Build2.1 from NCBI (<http://www.ncbi.nlm.nih.gov/blast/Blast.cgi>), *Saccoglossus kowalevskii* 2008-Dec-09 scaffolds from HGSC Baylor College of Medicine (<http://blast.hgsc.bcm.tmc.edu/blast.hgsc?organism=20>), *Gasterosteus aculeatus* 1.0, *Oryzias latipes* 1.0, *Anolis carolinensis* 2.0, *Ornithorhynchus anatinus* 1.2, *Monodelphis domestica* 2.2, *Loxodonta africana* 3.0, *Canis familiaris* 2.0, *Mus musculus* 37, and *Pan troglodytes* 2.1 from UCSC (<http://genome.ucsc.edu/>), *Drosophila melanogaster*, *D. ananassae*, *D. pseudoobscura*, *D. willistonii*, *D. virilis* from FlyBase (<http://flybase.org>), *Caenorhabditis elegans*, *C. briggsae*, *C. brenneri*, *C. remanei*, and *C. japonica* from WormBase (<http://www.wormbase.org>), and *Oikopleura dioica* v3 from Genoscope (<http://www.genoscope.cns.fr>). *C. milii* contigs were searched at the NCBI genomic BLAST webpage for unfinished eukaryotic genomes (http://www.ncbi.nlm.nih.gov/sutils/genom_table.cgi?organism=eukaryotes). Manually curated *Sowah* mRNA sequences from selected metazoan species are included as Supplemental Data S2. All other final gene annotations, including the correspondent genomic sequences and alternative splice variants are available upon request. In the study of the syntenic and regulatory interactions between *Sowah* and *Irx* loci, we applied a strict definition of GRB, requiring the presence of a bystander gene (Engstrom et al. 2007; Kikuta et al. 2007).

Phylogenetic analyses

We aligned *Sowah* protein sequences from multiple species using MAFFT (Katoh et al. 2002, 2005) as implemented in Jalview 2.4

(Waterhouse et al. 2009), and manually curated the alignments (available upon request) using information on intron positions (Irimia and Roy 2008). We performed two different phylogenetic analyses. First, to establish orthology of all studied *Sowah* genes, we used an alignment containing only the highly conserved ankyrin-repeat domain (Supplemental Fig. S2B). As there is no published information on closely related genes that could be used as outgroups, we performed BLASTP searches against the “Non-redundant protein sequences” database at NCBI. We selected BLAST hits with the highest score after applying two filters: (1) The gene is not one of our previously identified putative *Sowah* members, (2) the gene must have the same number of ankyrin-repeat domains as *Sowah* (four repeats, two highly conserved central domains, and two more divergent flanking ones spanning ancestral exons 5–9, Supplemental Fig. S3). Through this procedure we selected *Ilk* genes as outgroups. Orthology was further supported by the presence of a highly conserved domain in the N terminus of *Sowah* proteins that could not be identified in any other protein family (see the three first ancestral exons in Supplemental Fig. S3). This domain was clearly identifiable in all surveyed *Sowah* sequences except in *C. elegans*, teleost *Sowah2*, and vertebrate *SowahD*, whose orthology was otherwise well supported by the Bayesian trees and synteny data.

Second, to allow confident assignment of r-*Sowah* genes as in-groups within chordate *Sowah* genes, we increased the number of informative alignment positions, adding the aforementioned N-terminal domain and excluding the *Ilk* outgroups, the genes without the conserved N-terminal domain and divergent sequences from fast-evolving species (Supplemental Fig. S2B). We generated gene trees with MrBayes 3.1.2 (Huelsenbeck and Ronquist 2001; Ronquist and Huelsenbeck 2003) using two independent runs (each with four chains). Model selection using ProtTest (Drummond and Strimmer 2001; Guindon and Gascuel 2003; Abascal et al. 2005), convergence determination, burn-in, and consensus tree calculations were done as previously described (D’Aniello et al. 2008).

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Analysis of CNRs in *Sowah*–*Irx* loci

For amphioxus, we downloaded the *B. floridae* *Irx* cluster from JGI (<http://genome.jgipsf.org/Braf11/Braf11.home.html>). Two different haplotypes are normally present in the v1.0 assembly (Putnam et al. 2008). In the case of the *Irx*–*Sowah* cluster, one is located in scaffold-90, whereas the other is scrambled into several scaffolds: 632, 2884, 229, 975, and 90 (downstream from the other haplotype). We generated a consensus sequence of both haplotypes using scaffold-90 as default. We used the other scaffolds to remove gaps, correct assembly errors, and remove polymorphic repetitive elements (Supplemental Data S1). Preliminary blast searches allowed us to identify the presence (or traces of) of four tandem *Irx*–*Sowah* loci. Accordingly, we divided the nucleotide sequence of *B. floridae* cluster (i.e., from the end of the gene model preceding *Sowah* [*Rpgrip*] to the start of the gene model immediately after *IrxC*, the *CAVIII* gene) in four regions, each containing one of the *Irx* genes and surrounding noncoding sequences, plus a fourth region that contained the segment corresponding to the ancient

IrxD–*SowahD* locus, that has been partially lost. In the case of *IrxA*, we also included the entire *Sowah* locus. With these four sequences, we performed crossed VISTA analysis using default parameters (Frazer et al. 2004). It should be noted that regulatory blocks that are not repeated cannot be detected with this analysis. We also searched repeated sequences using blast-2-sequence (bl2seq) software at <http://www.ncbi.nlm.nih.gov/blast/bl2seq/wblast2.cgi>. Evidence for loss of *IrxD* was thoroughly verified using data from the assembly of both haplotypes to discard possible artifacts due to gaps or assembly errors.

For nematodes and flies, genome-wide VISTA analyses have not been done before. Therefore, we chose appropriate parameters for these species based on the length and percentage identity of CNRs previously identified in these lineages. In nematodes we used *C. elegans* as a reference sequence, using a window size of 50 bp and 90% minimum identity (85% in *C. japonica*), based on previously published data on worm CNRs (Vavouri et al. 2007) with slight modifications to take into account the inclusion of more divergent species. For flies, following previously reported CNR definitions (100% identity over 50 bp) (Glazov et al. 2005), we used a less-stringent identity criterion to account for the inclusion of *D. virilis* (90% identity) and a larger minimum window size (60 bp) following works reporting longer CNRs in the vicinity of developmental genes such as *Hox* genes (Negre et al. 2005). In vertebrates, VISTA analyses were performed using default parameters ($\geq 70\%$ identity over 100 bp), except for the detection of *Sowah2* pseudogenetic sequences (see below). In the case of teleosts *IrxBb* clusters, we complemented the VISTA analysis with bl2seq and ClustalW.

Ancora data (<http://ancora.genereg.net/>) were visualized as custom tracks in the UCSC Genome Browser (<http://genome.ucsc.edu/>). All available Ancora precomputed CNR density tracks (based on different CNR definitions) were checked, yielding equivalent results. For representation in Figure 1, we used the “96% identity over 30 column” density track for nematodes and the “98% identity over 50 columns” for flies.

Conservation tracks by phastCons and PhyloP of *Sowah* regions were downloaded from the UCSC Genome Browser (<http://genome.ucsc.edu/>).

Sequencing of *IrxB* locus in elephant shark

We searched the elephant shark *C. milli* 1.4 \times assembly (Venkatesh et al. 2007) using TBLASTN with human IRX3 and IRX5 protein sequences. We identified two scaffolds (AAVX01172485.1 and AAVX01359425.1) that contained exons for *Irx3* and *Irx5* genes, respectively. We cloned different DNA probes for each scaffold using PCR and used them to screen pooled DNA of an elephant shark BAC library (see Ravi et al. 2009). We selected one representative BAC each for *Irx3* and *Irx5*. After sequencing these BACs completely, we identified overlapping BACs by PCR. Altogether, we sequenced six BAC clones (Supplemental Fig. S10) to obtain the complete sequence of elephant shark *IrxB* locus (GenBank accession number JN228895). BAC sequencing was done using a standard shotgun sequencing technique, with the BigDye Terminator Cycle Sequencing Kit (Applied Biosystems). Sequences were processed and assembled using *phred-phrap* and *Consed* (<http://www.phrap.org/phredphrapconsed.html>). Elephant shark genes were predicted based on their homology with known genes in other vertebrates and as indicated above.

Functional status of *Sowah2* genes

Estimation of ω was performed with the maximum-likelihood (ML) method as implemented in CODEML in PAML v.4.2 (Yang 2007)

using the general model and WAG + Γ . Each *Sowah2* exon was analyzed independently, since not all of them could be identified in all studied species. *Sowah2* pseudogenized exonic sequences in tetrapods were detected with VISTA. Most of them could be retrieved using elephant shark as a reference sequence and low-stringency parameters (60% identity in 60 bp), although in some particular cases we used even a lower stringency (50% in 50 bp for pseudoxon 9 in *M. domestica*) and a different species as a reference sequence (human in the case of *M. musculus* pseudoxon 7). These analyses were complemented with bl2seq and ClustalW, and all pseudoxons were aligned and translated to highlight their mutations (data not shown). Selected tetrapod *Sowah2* pseudogenetic sequences are included in Supplemental Data S3. Expression data for teleost *Sowah2* genes were obtained by blasting nucleotide sequences from all teleost species against the teleost EST database at the NCBI blast webpage (<http://blast.ncbi.nlm.nih.gov/Blast.cgi>), and by mapping available short RNAseq reads from zebrafish head, ovary, and embryos against zebrafish *sowah2* mRNA using bowtie (Langmead et al. 2009) with default parameters. Raw ChIP-seq data for H3K4me1 and H3K4me3 in whole zebrafish embryos at 24 hpf was obtained from Aday et al. (2011). Highly enriched regions (peaks) of histone methylation were obtained by the MACS (v.1.3.3) algorithm (Zhang et al. 2008) using standard settings with one modification (mfold = 20). These results were uploaded to the UCSC browser (<http://genome.ucsc.edu/>) as custom tracks for visualization in *Danio rerio* Zv8. *Sowah2* exon 8 (located in an unmapped contig <2.7 kb long) data were not represented in Supplemental Figure S12 due to the lack of genomic context. Equivalent data from human and mouse were checked in roadmap (<http://www.roadmapepigenomics.org/>) and the UCSC Genome Browser, respectively.

Cloning of *Sowah* genes from amphioxus, *D. melanogaster*, and zebrafish, and of amphioxus CNRs

We designed primer pairs to span partial coding sequences of *D. melanogaster* and *B. floridae* *Sowah* and zebrafish *sowah* - r-*sowah* complement. We screened aliquots of cDNA libraries from different developmental stages of both *B. floridae* and *Branchiostoma lanceolatum* by PCR using the *B. floridae* *Sowah* primers. *D. melanogaster* *Sowah* was amplified by PCR using cDNA from larvae. Zebrafish genomic DNA and cDNA was used as a template to PCR amplify non-*Irx* linked r-*sowah* genes and *sowah1*, respectively. We cloned all genes in TA Cloning pCRII vectors (Invitrogen) and sequenced them using standard M13F and M13R primers. *B. lanceolatum* *Sowah* sequence has been submitted to GenBank (accession number JN609218).

For each amphioxus CNR, we designed primers to span the whole conserved sequence, plus a padding of ~ 100 nt on each side. We performed PCRs on genomic DNA using iProof High-Fidelity DNA Polymerase (Bio-Rad). We cloned amplicons in pCR8GW/TOPO vector (Invitrogen) according to the manufacturer's instructions. We then transferred sequence-verified clones with the Gateway recombination system (Invitrogen) to the ZED vector (Bessa et al. 2009). We purified the final transgenic constructs using phenolchloroform and normalized at 50 ng/mL in DEPC water prior to microinjection.

Sequences of all primers are provided in Supplemental Table S2.

sowah^{EGP} deficiency generation and fly stocks

The FLP–FRT recombination method and the FRT-bearing *piggyBac* insertion lines from the Exelixis collection were used to generate *w*⁺ *sowah*^{EGP} deficiencies, which were confirmed by PCR analyses (Parks et al. 2004; Thibault et al. 2004). We combined e03723 and e01289 starting insertions to generate *sowah*^{EGP1}, e01289, and

f05010 for *sowah*^{EGP2}, and f01127 and e02801b for *sowah*^{EGP3}. We confirmed the localization of the *piggyBac* insertion site by PCR in all lines, matching their previously described localization (Thibault et al. 2004). Note that there is an error in FlyBase: f01127 insertion site is reported >16 kb downstream from its actual location. *sowah*^{EGP1} is embryonic lethal, while a few LIII escapers are found in *sowah*^{EGP2} and *sowah*^{EGP3}.

iro^{DFM3} is a deficiency obtained by imprecise excision of the P[LacZ] element of *iro*^{FE209}, which removes *ara*, *caup*, and the promoter of *mirr* (Gomez-Skarmeta et al. 1996). *sowah*^{e01289} is a putative null allele of *sowah*: due to the transposon insertion at the beginning of the region encoding the highly conserved ankyrin repeat domain (Fig. 5A), this allele could encode a Sowah protein deficient in this domain.

In situ hybridization and immunohistochemistry in different species

In situ hybridization of whole mounts of *Drosophila* imaginal discs, amphioxus specimens of the European species *B. lanceolatum*, and zebrafish embryos with digoxigenin-labeled antisense RNA probes and immunocytochemistry of *Drosophila* embryos, were performed as previously described (Cubas et al. 1991; Tena et al. 2007; Yu and Holland 2009; Irimia et al. 2010; Carrasco-Rando et al. 2011). The following primary antibodies were used: rat anti-Caup, an antibody that recognizes both Ara and Caup 1:50 (Diez del Corral et al. 1999), and rabbit anti- β -galactosidase (Cappel; 1:5000).

Zebrafish microinjections and husbandry

We injected 50–100 pg of each ZED-CNR vector into one-cell stage embryos together with 50–100 pg of Tol2 mRNA. We observed injected fish at 24 and 48 hpf for GFP expression. As internal injection quality control, we determined muscle RFP expression at 72 hpf. Embryos were selected and raised to sexual maturity. Three or more independent stable transgenic lines were generated for each construct. We cloned PCR fragments in pCR8/TOPO and generated RNA probes using T7 polymerase and standard procedures. Embryos were reared at 28°C in standard E3 medium.

Real time-qPCRs (RT-qPCRs)

We studied *expression dynamics of Sowah* paralogs by RT-qPCRs. We isolated total RNA from 20 embryos each from 24 and 48 hpf stages, and 30 embryos from 80% of epiboly stage. cDNA was synthesized from total RNA by reverse transcription, and the relative amounts of different gene products were measured by RT-qPCR. We normalized all data using the gene *elongation factor 1-alpha*. We took the 24-hpf stage sample as reference.

Data access

The sequence data generated for this study have been submitted to GenBank (accession numbers JN228895 and JN609218).

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Authors' contributions: I.M. and M.I. performed the bioinformatic analyses, the amphioxus analyses, and cloning of CNRs. J.J.T. with the collaboration of I.M. carried out the zebrafish experiments. E.G.-P. performed the *Drosophila* experiments. D.T. and V.R. cloned and sequenced the elephant shark cluster. D.T., V.R., B.V., I.M., and M.I. analyzed the elephant shark data. All authors participated in the experimental design. I.M. and M.I. wrote the first draft of the manuscript and J.J.T., E.G.-P., B.V., S.C., J.L.G.-S., and J.G.-F. helped with the writing. I.M. and M.I. conceived the work.

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