

Conservation of regulatory sequences and gene expression patterns in the disintegrating *Drosophila Hox* gene complex

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Homeotic (*Hox*) genes are usually clustered and arranged in the same order as they are expressed along the anteroposterior body axis of metazoans. The mechanistic explanation for this colinearity has been elusive, and it may well be that a single and universal cause does not exist. The *Hox*-gene complex (HOM-C) has been rearranged differently in several *Drosophila* species, producing a striking diversity of *Hox* gene organizations. We investigated the genomic and functional consequences of the two HOM-C splits present in *Drosophila buzzatii*. Firstly, we sequenced two regions of the *D. buzzatii* genome, one containing the genes *labial* and *abdominal A*, and another one including *proboscipedia*, and compared their organization with that of *D. melanogaster* and *D. pseudoobscura* in order to map precisely the two splits. Then, a plethora of conserved noncoding sequences, which are putative enhancers, were identified around the three *Hox* genes closer to the splits. The position and order of these enhancers are conserved, with minor exceptions, between the three *Drosophila* species. Finally, we analyzed the expression patterns of the same three genes in embryos and imaginal discs of four *Drosophila* species with different *Hox*-gene organizations. The results show that their expression patterns are conserved despite the HOM-C splits. We conclude that, in *Drosophila*, *Hox*-gene clustering is not an absolute requirement for proper function. Rather, the organization of *Hox* genes is modular, and their clustering seems the result of phylogenetic inertia more than functional necessity.

[Supplemental material is available online at www.genome.org. The sequence data from this study have been submitted to GenBank under accession nos. AY900631–AY900632 and AY897430–AY897434.]

Homeotic (*Hox*) genes were discovered in *Drosophila melanogaster* as mutations that transform one body part into another. Lewis (1978) and Kaufman et al. (1980) found that these genes are clustered and arranged in the chromosome in the same order as their domains of action in the body of flies. Homologous *Hox* genes were subsequently found in many other animals and their arrangement in complexes (HOM-C) shown to be the general rule (McGinnis and Krumlauf 1992; Ruddle et al. 1994). *Hox* genes encode transcription factors involved in the determination of segment identity along the anteroposterior body axis, and thus, play a fundamental role in animal development. The conserved colinearity between *Hox* gene chromosomal arrangement and expression domain is a basic notion of developmental biology, yet this is an enigmatic phenomenon for which no single satisfactory explanation exists (Kmita and Duboule 2003). Furthermore, HOM-C splits have been observed in *Drosophila* (Von Allmen et al. 1996; Lewis et al. 2003; Negre et al. 2003), *Bombix* (Yasukochi et al. 2004), nematodes (Aboobaker and Blaxter 2003), and tunicates (Ikuta et al. 2004; Seo et al. 2004).

Ten genes arranged in a single complex comprised the an-

cestral HOM-C of arthropods (Cook et al. 2001; Hughes and Kaufman 2002; Hughes et al. 2004). In winged insects, including *Drosophila*, the genes *Hox3* and *fushi tarazu* (*ftz*) lost their homeotic function, and thus, only eight truly homeotic genes remain. Three different splits of the ancestral HOM-C have been found so far in the *Drosophila* genus (Fig. 1A). In *D. melanogaster*, the complex is split between the genes *Antennapedia* (*Antp*) and *Ultrathorax* (*Ubx*), leaving two separate gene clusters as follows: the Antennapedia complex, ANT-C (Kaufman et al. 1990) that specifies the identity of the mouth parts and anterior thorax, and the Bithorax complex, BX-C (Duncan 1987; Martin et al. 1995) involved in the development of the posterior thorax and abdomen. In *D. pseudoobscura*, the HOM-C is also similarly divided in the ANT-C and BX-C complexes (Lewis et al. 2003). A different split between *Ubx* and *abdominal A* (*abdA*) occurs in *D. virilis* (Von Allmen et al. 1996), *D. repleta* (Ranz et al. 2001), *D. buzzatii*, and other species of the *Drosophila* subgenus (Negre et al. 2003; Fig. 1B). Finally, an additional split, between *labial* (*lab*) and *proboscipedia* (*pb*), is present in *D. buzzatii* and other species of the *repleta* group (Negre et al. 2003). This third split separated the gene *lab* far from *pb* and the anterior genes of the *Hox* complex and relocated it near the posterior genes *abdA* and *Abdominal B* (*AbdB*) in a flagrant violation of the colinearity rule. The functional consequences of these splits are unknown.

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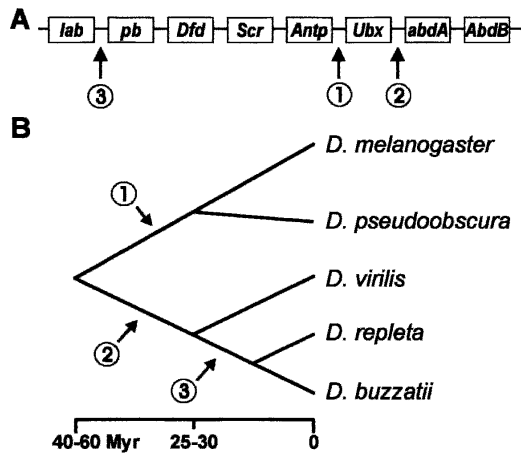


Figure 1. Genomic (A) and phylogenetic (B) localization of the three *Hox* gene complex splits observed in the *Drosophila* genus. (A) Ancestral arrangement of the eight *Hox* genes within the insects is as follows: *labial* (*lab*), *proboscipedia* (*pb*), *Deformed* (*Dfd*), *Sex combs reduced* (*Scr*), *Antennapedia* (*Antp*), *Ultrabithorax* (*Ubx*), *abdominal A* (*abdA*), and *Abdominal B* (*AbdB*). (B) Phylogenetic relationships and divergence times for the five *Drosophila* species included in this study. *D. melanogaster* and *D. pseudoobscura* belong to the *Sophophora* subgenus. *D. repleta* and *D. buzzatii* (both in the *repleta* species group) and *D. virilis* (*virilis* species group) belong to the *Drosophila* subgenus (see Negre et al. 2003 for details).

In order to ascertain the consequences of *Drosophila* HOM-C splits, we have carried out a genomic and functional characterization of the two splits present in *D. buzzatii*. We isolated and sequenced two BAC clones containing the *lab-abdA* and *pb* chromosomal regions of *D. buzzatii*. The gene organization in these regions is compared with that of the homologous regions in *D. melanogaster* and *D. pseudoobscura* to map the precise site of the two splits. None of the two splits has altered the coding regions of *Hox* genes. We then searched for Conserved Noncoding Sequences (CNS), which are putative regulatory sequences, around the genes *lab*, *pb*, and *abdA*, to find out whether the splits removed or altered any *Hox*-gene enhancer. The position of CNS around *Hox* genes is compared with experimentally identified *Hox*-gene enhancers, and the arrangement of CNS is compared between *Hox* and non-*Hox* genes. Finally, we analyzed the expression patterns of three *Hox* genes, *lab*, *pb*, and *abdA*, in four *Drosophila* species with different *Hox*-gene organizations (with and without the splits) in whole-mount embryos and imaginal discs. The results show that, in *Drosophila* species, *Hox* genes, as well as their regulatory regions and expression patterns, are conserved, despite the *Hox* complex breaks. Thus, the functional significance of the *Hox*-gene clustering in *Drosophila* is questionable.

Results

Molecular characterization of *Hox*-gene complex breakpoints

To characterize the two HOM-C splits present in *D. buzzatii*, we isolated and sequenced two BAC clones, one (5H14, 124,024 bp) containing the *lab-abdA* region, and another (40C11, 132,938 bp) including the *pb* region (see Methods). The organization of the two regions of *D. buzzatii* chromosome 2 is shown in Figure 2 along with the homologous regions of *D. melanogaster* and *D. pseudoobscura* for comparison. *D. melanogaster* and *D. pseudoobscura* are homosequential in the analyzed regions, except where indicated. The sequenced *pb* region (Fig. 2A) contains 16 ORFs including *Dbuz\pb*, *Dbuz\zerknüllt* (*Dbuz\zen*), *Dbuz\zerknüllt-related* (*Dbuz\zen2*), and *Dbuz\bicoid* (*Dbuz\bcd*). These four genes are present in the ANT-C of *D. melanogaster* and also in the homologous region of *D. pseudoobscura* (Fig. 2B,C). The orientation of *Dbuz\zen2* is the same as that of *Dpse\zen2*, but inverted with regard to *Dmel\zen2*. The remaining 12 genes in this region are orthologous to *D. melanogaster* genes from four different regions (84D1–2, 89D2, 84E5, and 91D4–5) of chromosomal arm 3R. One of the genes, *CG14609*, is represented by six copies, in contrast to the single copy present in *D. melanogaster* or *D. pseudoobscura*. A total of four breakpoints are fixed in this region between *D. buzzatii* and *D. melanogaster* beside the *zen2* microrearrangement. That corresponding to the *lab-pb* split is located in the ~3-kb intergenic segment between *Dbuz\pb* and *Dbuz\CG17836* (Fig. 2A).

The sequenced *lab-abdA* region contains 11 ORFs, including *Dbuz\lab*, the cuticular cluster genes (*Dbuz\Ccp*), and *Dbuz\abdA*

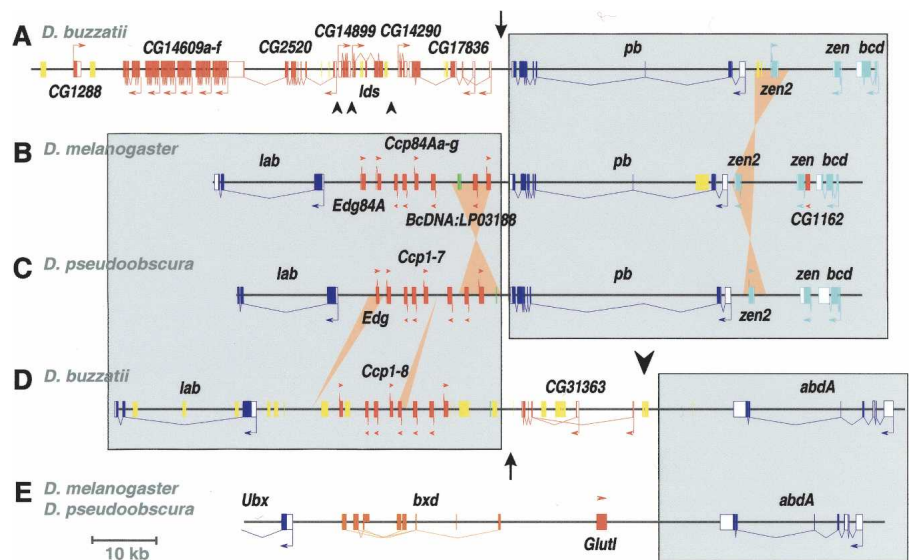


Figure 2. Gene organization of the *lab-abdA* and *pb* genomic regions of *D. buzzatii* compared with the homologous regions of *D. melanogaster* and *D. pseudoobscura*. The localization of the *lab-pb* split (arrow) and the *Ubx-abdA* split (large arrowhead) are indicated. (A) Sequence of *D. buzzatii* BAC 40C11 containing the *pb* region. (B) Organization of the *lab-pb* region in *D. melanogaster*. (C) *Idem* in *D. pseudoobscura*. (D) Sequence of *D. buzzatii* BAC 5H14 containing the *lab-abdA* region. (E) Organization of the *abdA* region in *D. melanogaster* and *D. pseudoobscura*. Genes are represented as open (UTRs) and filled boxes (coding sequences) with arrows indicating the sense of transcription. *Hox* genes are colored in dark blue, *Hox*-derived genes in light blue, non-*Hox* genes in red, noncoding RNA genes in orange, and the BcDNA:LP03188 and orthologous sequences in green. Transposable element insertions (usually *ISBu* elements, see Negre et al. 2003) are shown as yellow boxes. Large shaded rectangles include homologous *Hox*-gene regions in different species. Ochre triangles denote small inversions and insertions or deletions. Small arrowheads show breakpoints between *D. buzzatii* and *D. melanogaster* in non-*Hox* regions.

(Fig. 2D). The number of *Ccp* copies (including the gene *Edg*) is eight in the three species, but there is a small inversion encompassing two copies (plus the cDNA *BcDNA:LP03188*) in *D. melanogaster* in comparison to *D. buzzatii* or *D. pseudoobscura*, as well as one gain and one loss (Fig. 2B–D). These 11 genes come from three different regions (84A2–5, 86E11–13, and 89E2) of *D. melanogaster* chromosomal arm 3R, which means two fixed breakpoints between *D. buzzatii* and *D. melanogaster*, beside the small inversion of *Ccp* genes. One breakpoint corresponds to the *lab-pb* split and is found ~40 kb upstream of *Dbuz\lab*, in the 5-kb between the sequence similar to *BcDNA:LP03188* and the gene *Dbuz\CG31363*. The second breakpoint is that of the *Ubx-abdA* split and is located between 11 and 15 kb downstream of *Dbuz\abdA*. The two breakpoints are separated by a DNA segment of only ~22 kb encoding a single gene, *Dbuz\CG31363* (Fig. 2D).

Conserved noncoding sequences in *Hox* gene regions

We analyzed the conservation of noncoding sequences around the three *Hox* genes *lab*, *pb*, and *abdA* by comparing the sequences of the three species *D. buzzatii*, *D. melanogaster*, and *D. pseudoobscura* as done previously by other authors (Bergman and Kreitman 2001; Bergman et al. 2002) (see Methods). Figure 3 shows the VISTA graph, where the conservation between the aligned sequences is plotted (when higher than 50%) and the regions that meet the selected criteria (75% identity in a 25-bp window) are highlighted for both coding and noncoding sequences. A preliminary analysis showed no differences between intergenic and intronic regions, in agreement with previous studies (Bergman and Kreitman 2001). Thus, CNS are defined as intergenic (excluding UTRs) or intronic sequences that meet the above criteria. The characteristics of observed CNS are given in

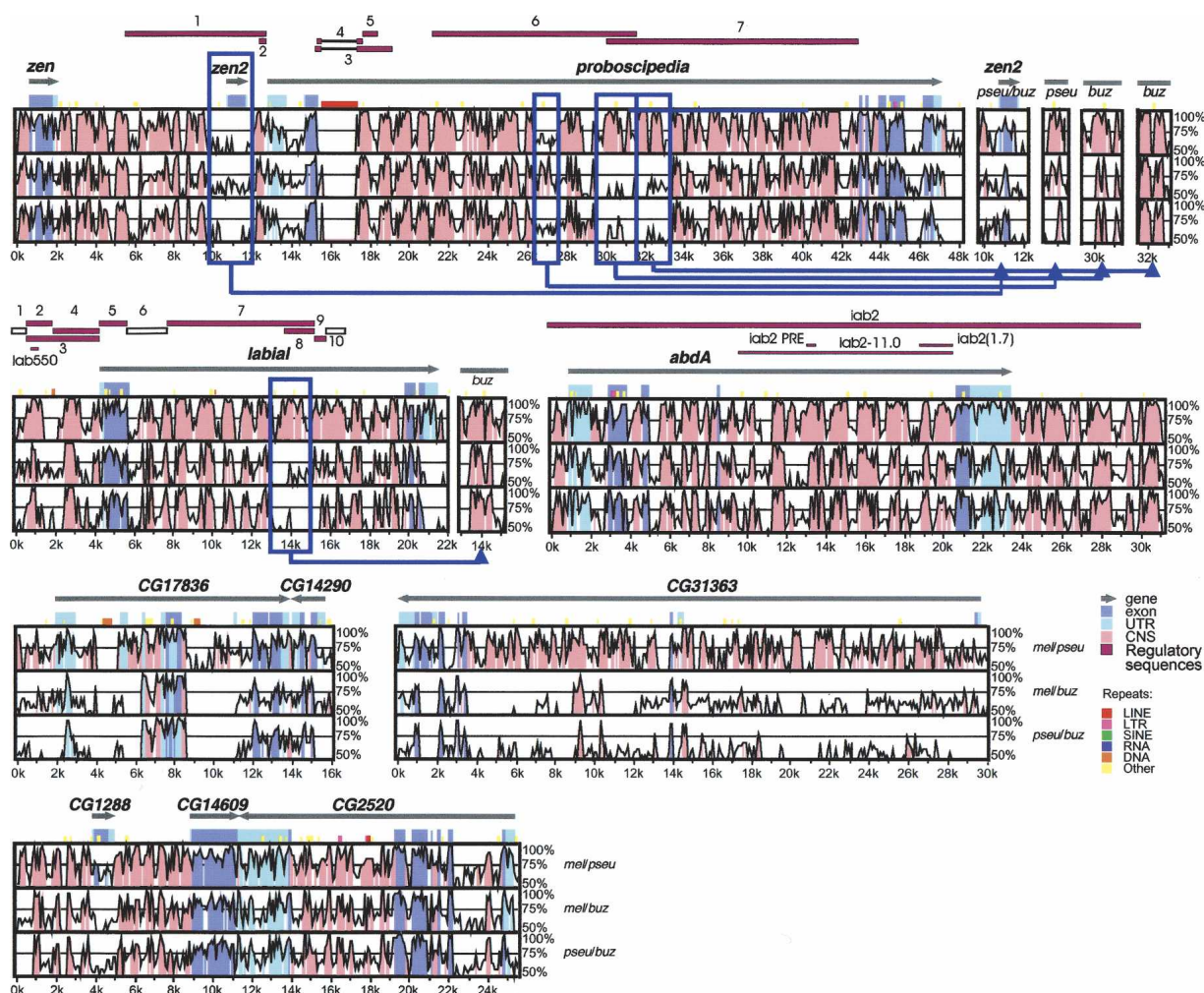


Figure 3. Nucleotide sequence conservation in the *lab-abdA* and *pb* regions between *Drosophila* species. The three panels in each VISTA plot represent pairwise comparisons between *D. melanogaster* and *D. pseudoobscura* (*mel/pse*), *D. melanogaster* and *D. buzzatii* (*mel/buz*) and *D. pseudoobscura* and *D. buzzatii* (*pse/buz*). The *x*-axis represents *D. melanogaster* coordinates, and *y*-axis sequence identity (50%–100%). Gray arrows show the location and orientation of genes. Conservation in exons and UTRs is shown in dark and light blue, respectively. Pink regions represent CNS. Experimentally identified regulatory sequences (solid purple bars) or segments with negative results (empty bars) are indicated on top of each plot. Five microinversions detected in the *lab* or *pb* regions are enclosed in blue frames, and the VISTA graphs generated with the inverted sequences shown to the right of the main plots. VISTA plots for the CG17836–CG14290, CG31363, and CG1288–CG14609–CG2520 regions (adjacent to *Hox* genes) are shown at the bottom of the figure for comparison.

Tables 1 and 2, and the results of statistical analysis are shown in Supplemental Tables S1 and S2.

When *D. buzzatii* is compared with *D. melanogaster* or *D. pseudoobscura*, 395 and 440 CNS are found, respectively, around the three *Hox* genes (Table 1). This gives a density of 4.5 and 5 CNS per kilobase, respectively. These conserved blocks show a mean size of 44 bp with 86.5% nucleotide identity and represent 20%–22% of the analyzed noncoding sequence. When *D. melanogaster* and *D. pseudoobscura* are compared, 563 CNS are detected (6.5/kb) with a mean size of 55 bp and an average identity of 87.4%. In this comparison, the sequence in CNS represents 36% of noncoding sequence. In all three comparisons, the three regions around the *Hox* genes *lab*, *pb*, and *abdA* are homogeneous with little variation either in CNS density, size, or nucleotide identity (Supplemental Table S1). It is worth noting that CNS are coincident in all three comparisons (Fig. 3), which means that all CNS detected when comparing *D. buzzatii* with either *D. melanogaster* or *D. pseudoobscura* are also found in the comparison between the latter two species. Although most CNS keep colinearity (relative position and orientation), we could identify four microinversions, around 1–2 kb in size. One is located within the large intron of *lab* and the other three in introns 2 and 3 of *pb* (Fig. 3).

D. buzzatii is equally distant phylogenetically from either *D. melanogaster* or *D. pseudoobscura* (Fig. 1). The latter two species belong to the same subgenus and are phylogenetically closer. We compared the characteristics of the CNS found in the three pairwise comparisons. As expected, there are no statistical differences between the CNS found when comparing *D. buzzatii* with either *D. melanogaster* or *D. pseudoobscura* (Supplemental Table S2). The CNS density and the proportion of sequence in CNS are significantly higher when comparing the phylogenetically closer species *D. melanogaster* and *D. pseudoobscura*. Increasing divergence time does not seem to affect the nucleotide identity of the CNS, although the size of the CNS detected in the *Hox*-gene regions shows a significant decrease (Supplemental Table S2).

Conserved noncoding sequences in non-*Hox* gene regions

To find out whether the observed pattern of CNS is a particular feature of *Hox* genes, we also analyzed the presence of CNS in regions of the sequenced BACs adjacent, but unrelated, to *Hox* genes. We used the three microsyntenic regions between *D. buzzatii*, *D. melanogaster*, and *D. pseudoobscura* longer than 10 kb, i.e.,

the *CG31363* gene region, between *lab* and *abdA*, and the *CG17836-CG14290* and *CG1288-CG2520* regions, near *pb* (Fig. 2). These regions include one, two, and three genes, respectively. The pattern of CNS detected is shown in Figure 3 and summarized in Table 2. In the comparisons with *D. buzzatii*, we found around 100 CNS (~2/kb), which represents <8% of noncoding sequence. Thus, in these non-*Hox* regions, a much smaller number of CNS is observed and the proportion of sequence in CNS is also significantly lower than in *Hox*-gene regions (Supplemental Table S1). In the *D. melanogaster*–*D. pseudoobscura* comparison, there are 326 CNS (5.7/kb) which represents a 23% of noncoding sequence. Thus, in this case, the density is similar between *Hox* and non-*Hox*-gene regions, but the size of CNS is significantly smaller in the latter regions (Supplemental Table S1). Consequently, the proportion of sequence in CNS is also significantly lower in the non-*Hox*-gene regions. It should be noted that non-*Hox* regions show a significant variation for CNS density and also for the proportion of sequence in CNS that is not observed in *Hox*-gene regions (Supplemental Table S1). The higher variation observed between non-*Hox* regions is probably due to the heterogeneity of the sample from a functional point of view. There is little information available on the function and expression pattern of the six non-*Hox* genes analyzed, which probably represent a mixture of genes with different regulatory needs and number of enhancers.

Conservation of known regulatory sequences

Regulatory sequences of the genes *lab*, *pb*, and *abdA* have been experimentally identified in *D. melanogaster* (Karch et al. 1985; Chouinard and Kaufman 1991; Kapoun and Kaufman 1995; Martin et al. 1995). We compared their position with the pattern of CNS found around *Hox* genes. As shown in Figure 3, the regulatory sequences identified in *D. melanogaster* generally contain or correspond to CNS in *D. buzzatii*. For instance, CNS are found in the sites corresponding to the *iab2 PRE* and *iab2(1.7)* enhancers of *abdA* (Shimell et al. 1994, 2000). Similarly, a prominent conservation peak is observed at the site of the *lab550* enhancer, which directs the expression of *lab* in the embryo midgut (Marty et al. 2001). Also, the inverted segment found in the large intron of the *lab* gene roughly corresponds to the segment responsible for *lab* expression in the posterior midgut. Sequence details of the *lab550* and *iab2(1.7)* enhancer and binding site conservation are shown in Supplemental Figure S1. The Homeotic Response Ele-

Table 1. Characteristics of conserved noncoding sequences (CNS) detected with mVISTA in comparisons of *Hox* gene regions between *D. melanogaster* (mel), *D. pseudoobscura* (pse), and *D. buzzatii* (buz)

Region	Noncoding nucleotides	Species pair	Number of CNS	Density ^a (SD)	Mean size (nt) (SD)	Mean nucleotide identity (%)	Sequence in CNS (%)
<i>lab</i>	19,227	mel/pse	129	6.71 (0.59)	53.20 (34.03)	87.69	35.69
		mel/buz	73	3.80 (0.44)	46.70 (24.35)	87.33	17.73
		pse/buz	84	4.37 (0.48)	44.54 (27.05)	88.08	19.46
<i>pb</i>	42,056	mel/pse	265	6.30 (0.39)	55.04 (35.79)	87.29	34.68
		mel/buz	196	4.66 (0.33)	41.88 (22.38)	87.09	19.52
		pse/buz	215	5.11 (0.35)	42.92 (24.65)	86.38	21.94
<i>abdA</i>	26,043	mel/pse	169	6.49 (0.50)	59.29 (38.43)	87.44	38.45
		mel/buz	126	4.84 (0.43)	45.98 (26.15)	86.18	22.25
		pse/buz	141	5.41 (0.46)	46.11 (24.97)	85.44	24.97
Total Hox gene regions	87,326	mel/pse	563	6.45 (0.27)	55.89 (36.23)	87.42	36.03
		mel/buz	395	4.52 (0.23)	44.08 (24.04)	86.83	19.94
		pse/buz	440	5.04 (0.24)	44.25 (25.53)	86.39	22.30

^aDensity = number of CNS per kilobase.

Table 2. Characteristics of conserved noncoding sequences (CNS) detected with mVISTA in comparisons of non-*Hox* gene regions between *D. melanogaster* (mel), *D. pseudoobscura* (pse), and *D. buzzatii* (buz)

Region	Noncoding nucleotides	Species pair	Number of CNS	Density ^a (SD)	Mean size (nt) (SD)	Mean nucleotide identity (%)	Sequence in CNS (%)
CG1288-CG2520	18,333	mel/pse	127	6.93 (0.61)	43.48 (28.16)	86.31	30.12
		mel/buz	65	3.55 (0.44)	45.26 (31.51)	86.30	16.05
		pse/buz	67	3.65 (0.45)	42.44 (29.40)	86.81	15.59
CG17836-CG14290	10,921	mel/pse	46	4.21 (0.62)	45.02 (33.27)	82.67	18.96
		mel/buz	18	1.65 (0.39)	39.61 (22.67)	82.88	6.53
		pse/buz	22	2.01 (0.43)	42.09 (24.34)	84.34	8.48
CG31363	27,510	mel/pse	153	5.56 (0.45)	35.51 (14.95)	87.17	19.75
		mel/buz	22	0.80 (0.17)	26.09 (3.94)	82.93	2.09
		pse/buz	23	0.84 (0.17)	28.78 (7.70)	83.23	2.41
Total non- <i>Hox</i> gene regions	56,764	mel/pse	326	5.74 (0.32)	39.96 (24.15)	86.09	22.95
		mel/buz	105	1.84 (0.18)	40.28 (27.50)	85.27	7.45
		pse/buz	112	1.97 (0.19)	39.59 (25.88)	85.76	7.83

^aDensity = number of CNS per kilobase.

ment (HOMRE) of the *lab550* enhancer contains four binding sites; all of them are conserved in the three species. In the *iab2(1.7)* enhancer, there are five Hunchback (HB)-binding sites, three of which are conserved in the three species, whereas the other two vary in position between species. This enhancer also contains a unique Krüppel (KR)-binding site, where point mutations in *D. melanogaster* cause gain-of-expression mutants (Hab1 and Hab2) (Shimell et al. 1994). This binding site is conserved in all three species (Supplemental Fig. S1). The conservation between *D. melanogaster* and *D. buzzatii* around *abdA* ends 9 kb (in *D. melanogaster*) and 11 kb (in *D. buzzatii*) downstream of this gene (Fig. 3). This boundary lies between the *iab2* and *pbx* regulatory sequences, which control the expression of *abdA* and *Ubx*, respectively (Karch et al. 1985). We have shown that the *iab2* region downstream of *abdA* is conserved in *D. buzzatii*. We have not sequenced the *Ubx* region in *D. buzzatii*, but we assume that the *pbx* regulatory sequence will conserve its position upstream of *Ubx*, i.e., there are no rearrangements between *Ubx* and its regulatory sequences (see below).

It is worth noting though, that CNS were also found in fragments not experimentally tested or described as with no effect on expression (Fig. 3). This observation suggests that the regulation of these genes may be even more complex than currently envisaged, and that more regulatory modules may be operative in nature than those experimentally identified in the laboratory.

Hox gene expression patterns

The conservation of regulatory sequences suggests that splits of the HOM-C had no consequences on *Hox*-gene expression. To test this prediction, we compared the expression patterns of the *Hox* genes *lab*, *pb*, and *abdA* between *D. melanogaster*, *D. virilis*, *D. buzzatii*, and *D. repleta*. These four *Drosophila* species represent three different *Hox*-gene organizations (Figs. 1,2). *D. melanogaster* possess the *Antp-Ubx* split only, whereas *D. virilis* has the *Ubx-abdA* split instead. Both *D. buzzatii* and *D. repleta* present the *Ubx-abdA* and *lab-pb* splits. We used in situ hybridization and antibody staining to whole-mount embryos and to imaginal discs from third instar larvae and prepupae (see Methods). Detailed results are given in Figure 4 and Supplemental Figures S2–S6. The expression patterns of the four species closely follow those described for *D. melanogaster* (for review, see Hughes and Kaufman 2002). Interspecific variation was detected only in the *pb* gene, which in *D. virilis* presents an extra domain in the embryo mesoderm (Fig. 4). As this expression domain is not shared by *D. melanogaster*, it is seemingly not related with the *lab-pb* split. Although our analysis is qualitative, and slightly quantitative changes or domain changes of a few cells may remain undetected, it shows that the reorganization of the HOM-C caused no major alterations of the expression patterns of the three *Hox* genes adjacent to the splits, in good agreement with the conservation of regulatory sequences (see above). Likewise, Bomze and

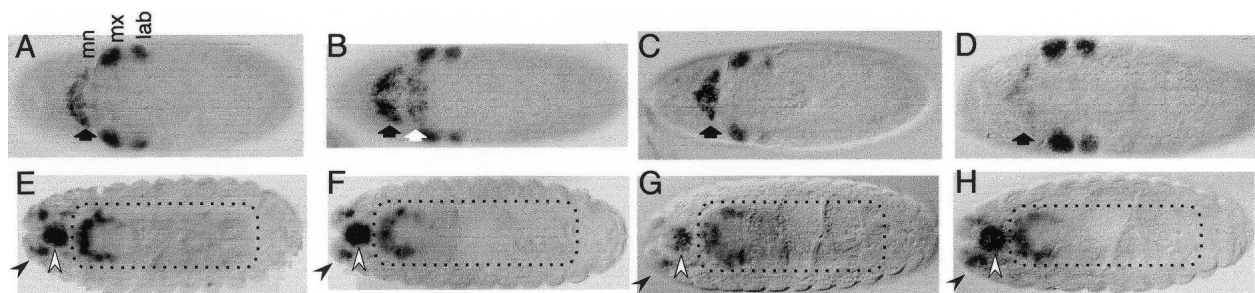


Figure 4. Expression pattern of *pb* in embryos. (A–D) stage 11 embryos, (E–H) stage 17 embryos. (A,E) *D. melanogaster*, (B,F) *D. virilis*, (C,G) *D. buzzatii*, and (D,H) *D. repleta*. (A–D) Expression on the ectoderm of the maxillary and labial lobes. Later in development (E–H) *pb* is detected in the derivatives of the maxillary (white arrowhead) and labial (black arrowhead) lobes, and in the ventral nervous system (boxed area). (A–D) *pb* expression is detected in the mesodermal layer of the mandibular segment (black arrow) in all four species. In *D. virilis* only (B), *pb* is also expressed in the mesodermal layer of the maxillary segment (white arrow). The mandibular (mn), maxillary (mx), and labial (lab) segments are shown in A.

López (1994) found that the expression pattern of *Ubx* in embryos is conserved between *D. melanogaster*, *D. pseudoobscura*, *D. virilis*, and *D. hydei* (a species of the *repleta* group), despite their different *Hox*-gene organization (Figure 1).

Discussion

zen2 predates the *Drosophila* radiation

The *zen* and *bcd* genes come from a duplication of *Hox3* in the ancestor of Cyclorhaphan flies (Stauber et al. 2002). A second duplication of *zen* gave birth to *zen2*, which was thought to be a recent event in *D. melanogaster* (Randazzo et al. 1993), where it has no discernible function. However, the existence of *Dpse\zen2* and *Dbuz\zen2* shows that the *zen-zen2* duplication must predate the divergence of the *Sophophora* and *Drosophila* subgenus, and that this gene has been kept during at least 40–60 Myr of evolution. Whether this gene is also present in other flies outside of the *Drosophila* genus is still unknown.

Patterns of conserved noncoding sequence evolution

Cis-Regulatory Modules (CRM) are transcription regulatory DNA segments (from a few hundred base pair to 1 kb in size) that control gene expression in higher eukaryotes (Wray et al. 2003). CRM have a complex structure still not fully understood. They contain one or several binding sites for different transcription factors, which act cooperatively to activate or repress transcription of the target gene. As CRM are functionally constrained to maintain the expression of the target gene, they evolve slower than nonfunctional sequences. Therefore, the conservation of noncoding sequences between phylogenetically distant species may be used as a guide for identification of regulatory sequences. Several recent studies (Bergman and Kreitman 2001; Bergman et al. 2002; Cooper and Sidow 2003; Nobrega et al. 2003; Santini et al. 2003) support the use of comparative sequence analysis and characterization of CNS as a useful approach to detect putative CRM in *Drosophila* and other organisms. The clustering of previously characterized transcription-factor binding sites may be also used for detection of CRM (Berman et al. 2004). However, the absence of high-quality binding data for most *Drosophila* transcription factors represent a great current limitation in the widespread application of this method.

We exhaustively searched for CNS around *lab*, *pb*, and *abdA* and around adjacent non-*Hox* genes by comparing three species pairs. A plethora of highly conserved blocks was found surrounding the three *Hox* genes in the comparison between the phylogenetically distant species *D. buzzatii* and *D. melanogaster* or *D. pseudoobscura* (Fig. 1). The proportion of noncoding sequence included in CNS was 20%–22%. In most cases, these CNS keep their relative position and colinearity, although a few microrearrangements were found. The interpretation of these CNS as regulatory sequences is supported by the high neutral substitution rate (Moriyama and Gojobori 1992) and intrinsic rate of DNA loss (Petrov et al. 1996; Singh and Petrov 2004) in *Drosophila*. Noncoding sequences are not expected to be conserved between such distantly related species unless they are functionally constrained. The coincidence between CNS and known enhancers such as *iab2 PRE* or *lab550* (Supplemental Fig. S1) further supports this interpretation.

A lower CNS density was observed around non-*Hox* genes. This result fits well with previous observations showing that genes with complex developmentally regulated expression show

a higher degree of conservation in noncoding regions than more simple genes with metabolic or housekeeping functions (Bergman and Kreitman 2001; Bergman et al. 2002; Halligan et al. 2004). Moreover, *Hox* genes are associated with larger noncoding regions. *Hox* genes harbor some of the longest introns of any *Drosophila* gene (Moriyama et al. 1998) and mean intron size is significantly greater in the *Hox* than in the non-*Hox* genes analyzed here ($F = 4.69$, $df = 1$, $P < 0.05$). This observation also fits with the notion that the amount of noncoding DNA must be larger in those genes with complex developmental functions in order to harbor the required CRM (Nelson et al. 2004).

HOM-C evolution in *Drosophila*

In *Drosophila*, *Hox* genes are arranged in the same 5'→3' orientation (with only one exception, the *Deformed* gene in *D. melanogaster*). Their regulatory sequences are usually located upstream of each gene and in the introns. If we look at the three HOM-C splits known in *Drosophila*, a common pattern arises. As can be seen in Figure 2, the *lab-pb* split took place close to the 3' end of *pb* and far from the *lab* 5' end. Likewise, the split between the genes *Ubx* and *abdA* took place near the *abdA* 3' end and far from the *Ubx* 5' end, in the short space between their respective regulatory sequences *pbx* and *iab2*. This is approximately the same position where an experimental break that does not affect development has been observed (Struhl 1984), although the deficiencies used in the complementation tests both carry a fraction of the *pbx* and *iab2* regions. Finally, sequence comparison between *D. melanogaster* and *D. virilis* (Lewis et al. 2003) show that both the insertion of the *CG31217* gene and the *Antp-Ubx* split took place close to the *Ubx* 3' end, and far from the *Antp* 5' end (results not shown). Thus, all three splits seem to have occurred far from the 5' end of one gene and much closer to the 3' end of the next one, in such a way as to keep in place the regulatory sequences of both genes. In this way, rearrangements did not alter any of the known regulatory sequences of these *Hox* genes; this would explain the absence of gene expression changes.

In the *repleta* group species, the anterior gene *lab* is located near the posterior genes *abdA* and *AbdB*. The sequence analysis shows that *lab* and *abdA* are only 75 kb apart and show the same orientation. The breakpoint of the *lab-pb* split occurred at ~22 kb from that of the *Ubx-abdA* split. None of those splits seem to have affected the regulatory regions of the *Hox* genes, because the expression patterns of *lab* and *abdA* are unaffected. Although it is intriguing, the proximity between these genes in the *D. buzzatii* genome seems purely accidental and lacking any functional significance.

The most likely mechanisms for the generation of the HOM-C splits are paracentric inversions (Ranz et al. 2001; Gonzalez et al. 2002). A plausible reconstruction of HOM-C evolution in the *Drosophila* subgenus that accounts for the current organization of *Hox* genes in *D. buzzatii* is shown in Figure 5. In lower Dipterans, such as *Anopheles gambiae*, the eight *Hox* genes, plus *Hox3* and *ftz*, are arranged as a single cluster (Powers et al. 2000). Before the radiation of the *Drosophila* genus, two transpositions occurred as follows: the *Ccp* gene cluster between *lab* and *pb*, and the gene *CG31217* between *Antp* and *Ubx* (Lewis et al. 2003). Also, *zen*, *zen2*, and *bcd* evolved from the *Hox3* gene (see above). In the lineage of the *Drosophila* subgenus, an inversion took place with one breakpoint between *Ubx* and *abdA* (split 2 in Fig. 1) and the other one between *CG31363* and an unknown ORF (*X*). This HOM-C structure is now present in species of the *Drosophila* sub-

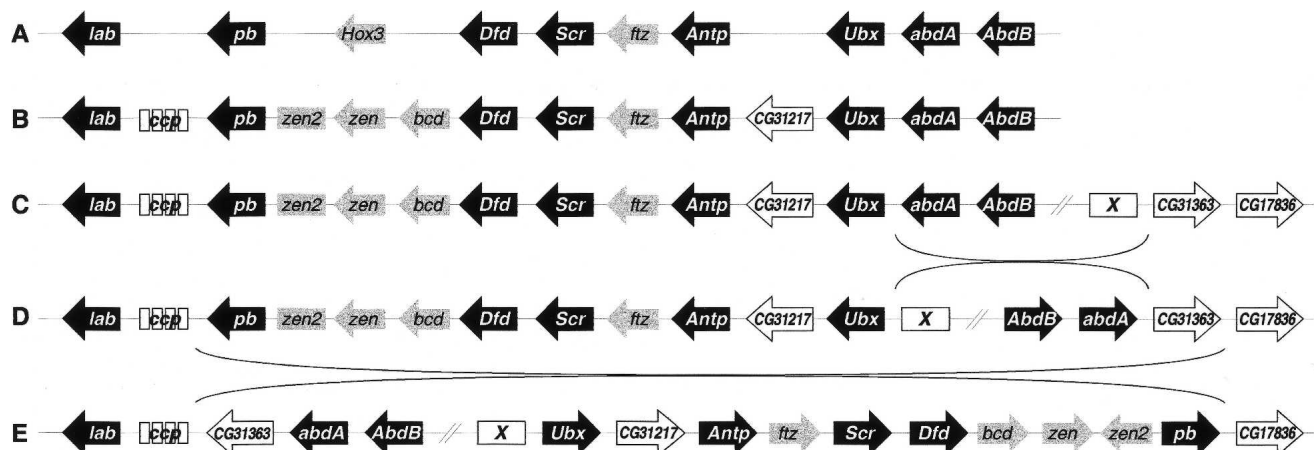


Figure 5. Reconstruction of the *Hox* gene complex evolution in the *Drosophila* subgenus. Genes are shown as arrows when the orientation 5'→3' is known, and as rectangles otherwise. *Hox* genes are in black, *Hox*-related genes in gray, and non-*Hox* genes in white. (A) Lower Dipterans. (B) Before the radiation of the *Drosophila* genus. (C) *Drosophila* subgenus after its separation from that of the *Sophophora* subgenus. (D) Ancestor of the *repleta* group. (E) Present arrangement of *Hox* genes in *Drosophila buzzatii* (cf. Fig. 2A,D).

genus outside the *repleta* group, such as *D. virilis* (see Fig. 1). A second inversion, in the ancestor of the *repleta* group, split the HOM-C between *lab* and *pb* (split 3 in Fig. 1). This inversion, which relocated *lab* close to *abdA*, had one breakpoint between *pb* and the *Ccp* cluster genes and the second breakpoint between *CG31363* and *CG17836*. These two genes are not adjacent in the *D. melanogaster* genome, but we infer that they were so in the ancestor of the *Drosophila* subgenus.

Do flies have a *Hox* gene complex?

Despite the striking conservation of *Hox*-gene clustering in metazoans, if we compare two of the most deeply studied organisms, *Drosophila* and vertebrates, important differences arise (Ferrier and Minguillon 2003; Santini et al. 2003; Wagner et al. 2003). *Drosophila* *Hox*-gene regions (1) are much larger than those of vertebrates, e.g., the human *HoxA* cluster is only 110 kb long, whereas the *D. melanogaster* HOM-C spans 665 kb; (2) contain transposable element insertions, which are remarkably absent in those of vertebrates; (3) contain also non-*Hox* genes that are inserted between the *Hox* genes, and tandem duplications within the complex, such as those of the *zen*-related genes; (4) allow for small inversions of *Hox* genes, such as *Dfd* (Randazzo et al. 1993), and non-*Hox* genes, such as *zen2* (Fig. 2); and (5) are split in three ways in different lineages, apparently without consequences on gene expression. These observations suggest a highly dynamic evolution in *Drosophila* that contrasts with the compact structure seen in vertebrates. Thus, the splits of HOM-C in *Drosophila* indicate a release of functional requirements present in other metazoan.

Moreover, *Drosophila* is not the only organism known to have a split HOM-C. Split *Hox*-gene complexes were also known in nematodes, and recently have been described in *Bombyx* and tunicates. What do those organisms have in common in addition to the split HOM-C? Vertebrate development follows a rostral-to-caudal temporal progression, and the colinearity of *Hox* genes is not only spatial, but also temporal (the *Hox* clock) (Kmita and Duboule 2003). In the tunicate *Oikopleura*, *Hox* gene expression still evokes spatial colinearity but not temporal (Seo et al. 2004), which favors the argument that the constraining force of HOM-C structure conservation is temporal colinearity (Ferrier and Min-

guillon 2003). In nematodes, the pattern of *Hox*-gene evolution seems indicative of the move to a deterministic developmental mode (Aboobaker and Blaxter 2003). *Bombyx* embryogenesis, which is difficult to assign to a short or a long germ insect, is characterized by a quick development (Davis and Patel 2002). *Drosophila* is a long germ insect, where all *Hox* genes are activated almost simultaneously during the cellular blastoderm stage. Thus, none of these organisms seems to show temporal colinearity. A common feature between all organisms shown so far to have a split *Hox* complex seems to be a derived mode of embryogenesis characterized by a fast early development.

The loss of temporal progression in the activation of *Hox* genes in a very rapid mode of embryogenesis could be the ultimate cause for the modular organization of those *Hox* "clusters," where modules can be taken apart without loss of function. Given the high rate of chromosomal rearrangement in the genus *Drosophila* (Ranz et al. 2001; Gonzalez et al. 2002), we anticipate that an even greater variety of *Hox*-gene organizations will be discovered when more species are investigated. It is ironic that *Hox*-gene colinearity was discovered in *Drosophila*, an organism with a partially disassembled complex, which may be the by-product of phylogenetic inertia more than that of functional necessity.

Methods

Flies

D. buzzatii stock st-1 was used for construction of a genomic BAC library (González et al. 2005). The following species and stocks were used for gene expression experiments: *D. buzzatii* (j19), *D. repleta* (1611.2), *D. virilis* (Tokyo-Japan), and *D. melanogaster* (Canton S and Oregon R).

BAC sequencing

The genomic BAC library was screened with probes from the *lab*, *pb*, and *abdA* genes (González et al. 2005). Positive clones were used to build physical maps for the *lab-abdA* and *pb* chromosomal regions, and one BAC clone from each region was chosen for sequencing. Shotgun sublibraries were constructed for each BAC using the vector TOPO, and enough plasmid clones were se-

quenced by both ends to reach an $\sim 6\times$ redundancy. Reads were assembled with the PHRED-PHRAD-CONSED software (Ewing and Green 1998; Ewing et al. 1998; Gordon et al. 1998) and sequences finished with one round of AUTOFINISH (Gordon et al. 2001), followed by PCR to bridge the remaining gaps. A continuous high-quality sequence (PHRED score >40) was obtained for BAC clones 5H14 (124,024 bp), and 40C11 (132,938 bp). Statistical details of the sequencing process are given in Supplemental Table S3.

Sequence annotation

Nucleotide sequences were annotated with the aid of GENESCRIP (Hudek et al. 2003) and ARTEMIS (Berriman and Rutherford 2003). Predicted ORFs were corroborated with GOFigure (Khan et al. 2003) for automatic Gene Ontology (Harris et al. 2004) annotation, and BLAST (McGinnis and Madden 2004) for similarity searches. *D. buzzatii* sequences were compared with those of homologous regions in *D. melanogaster* (Celniker et al. 2002) and *D. pseudoobscura* (Richards et al. 2005) genomes. *D. melanogaster* sequences used were as follows: AE001572 (ANT-C), DMU31961 (BX-C), and AE003692, AE003672, AE003713, AE003676, and AE003724 (other regions). *D. pseudoobscura* contigs AADE01000437 (*lab*), AADE01000149 (*pb*), AADE01000036 (*abdA*), and AADE01000014, AADE000175, AADE01002495, AADE01000322 (non-*Hox* genes) were identified with Genome VISTA (Dubchak et al. 2000) and the regions of interest annotated.

Analysis of regulatory sequences

Pairwise alignments of six homologous genomic regions between *D. buzzatii*, *D. melanogaster*, and *D. pseudoobscura* were performed with the AVID global-alignment tool using default parameters (Bray et al. 2003). CNS were identified in the alignments with mVISTA (Mayor et al. 2000) using a window size of 25 bp and a minimum identity of 75%. Statistical tests were carried out to compare the characteristics of the CNS found in the different regions. Comparisons of CNS size distributions, which depart significantly from normality, were conducted using the G-test (Sokal and Rohlf 1995). The number of CNS and the proportion of sequence within CNS was scored for 1-kb windows along the analyzed regions (masking out exons). The resulting variables (density and percent sequence in CNS) as well as the nucleotide identity (per CNS) were tested using ANOVA (Sokal and Rohlf 1995). A complete list of CNS detected is provided in Supplemental Table S5.

Gene-expression experiments

In situ hybridizations and antibody staining were performed to whole-mount embryos and to imaginal discs from third-instar larvae and prepupae as described (Alonso and Akam 2003; Suzanne et al. 2003). cDNA clones were obtained for *lab* from the four species, *pb* from *D. buzzatii* and *D. melanogaster* and *abdA* from *D. buzzatii*, *D. repleta*, and *D. virilis* as described (Negre et al. 2003) (for primers see Supplemental Table S4). Sense and anti-sense RNA probes were produced as described (Suzanne et al. 2003). When no species-specific probe was available, at least two different ones were used in independent experiments, and the results were always consistent. Specific antibodies against the protein were used for *abdA* (Macias et al. 1990).

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