# Evolution of the entire arthropod *Hox* gene set predated the origin and radiation of the onychophoran/arthropod clade

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**Background:** Dramatic changes in body size and pattern occurred during the radiation of many taxa in the Cambrian, and these changes are best documented for the arthropods. The sudden appearance of such diverse body plans raises the fundamental question of when the genes and the developmental control systems that regulate these designs evolved. As *Hox* genes regulate arthropod body patterns, the evolution of these genes may have played a role in the origin and diversification of the arthropod body plan from a homonomous ancestor. To trace the origin of arthropod *Hox* genes, we examined their distribution in a myriapod and in the Onychophora, a sister group to the arthropods.

**Results:** Despite the limited segmental diversity within myriapods and Onychophora, all insect *Hox* genes are present in both taxa, including the trunk *Hox* genes *Ultrabithorax* and *abdominal-A* as well as an ortholog of the *fushi tarazu* gene. Comparative analysis of *Hox* gene deployment revealed that the anterior boundary of expression of trunk *Hox* genes has shifted dramatically along the anteroposterior axis between Onychophora and different arthropod classes. Furthermore, we found that repression of expression of the *Hox* target gene *Distal-less* is unique to the insect lineage.

**Conclusions:** A complete arthropod *Hox* gene family existed in the ancestor of the onychophoran/arthropod clade. No new *Hox* genes were therefore required to catalyze the arthropod radiation; instead, arthropod body-plan diversity arose through changes in the regulation of *Hox* genes and their downstream targets.

# Background

The sudden appearance and remarkable diversity of complex animals in the Cambrian has prompted extensive paleontological [1,2], comparative [3,4] and molecular systematic [5,6] studies of animal relationships and the origin and evolution of metazoan body plans. Current debate is focused on three central issues: the origin of taxa (cladogenesis), the variety of basic designs (morphological disparity) and the evolution of the developmental control systems that regulate these designs [7–9]. Comparisons of evidence from fossils and molecular studies may help to reveal whether the Cambrian marks the true origin of many higher taxa or whether major lineages diverged well before the Cambrian [7-10]. In either of these scenarios, it appears that dramatic changes in body size, pattern and diversity occurred during the radiation of many taxa in the Cambrian, raising a fundamental question: which new developmental mechanisms [11] and genetic information were responsible for the evolution and diversification of larger and more complex animals?

The analysis of arthropods and their relatives may present the best opportunity for an integrated approach to the Addresses: \*Howard Hughes Medical Institute and Laboratory of Molecular Biology, University of Wisconsin, 1525 Linden Drive, Madison, Wisconsin 53706, USA. <sup>†</sup>Department of Zoology, University of New England, Armidale, New South Wales 2351, Australia.

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problem of the evolution of body plans: the fossil record of arthropods is relatively abundant; arthropods are the most speciose and morphologically diverse taxa; and many recent advances in developmental genetics have emerged from the study of one arthropod, *Drosophila melanogaster*. Paleontological, comparative and molecular systematic evidence suggests that arthropods are monophyletic [3,12,13] and descended from a lobopodian ancestor [14]. The most striking trend in the evolution of the lobopodian/arthropod clade has been the diversification of segment types, from the simplicity of some Cambrian lobopodians, with four types of unjointed appendages and a uniformly patterned (homonomous) trunk [14–16], to the complexity of some extant crustaceans and insects, with as many as ten distinguishable appendage types and a highly diversified trunk.

As segmental diversity in highly derived insects such as *Drosophila* is regulated by eight *Hox* genes, it has been postulated that primitive arthropods possessed a more limited set of *Hox* genes which expanded during the course of arthropod and insect evolution [17,18]. Comparative studies of *Hox* genes in various metazoans indicate that many *Hox* genes predate the origin of the insects

[19,20]. The common ancestor of arthropods and vertebrates possessed five or six Hox genes [21-23], and most of the insect Hox genes were present in the annelid/arthropod ancestor [24-29]. Importantly, arthropods have two unique Hox genes, Ultrabithorax (Ubx) and abdominal-A (abd-A), which are not found in vertebrates [21] or annelids [24,26,27]. The Ubx and abd-A genes are present in both crustaceans [30] and insects [19,20], the two arthropod classes that have the greatest segmental diversity. It is possible that the evolution of these two Hox genes facilitated the diversification of trunk segments during the evolution of diverse arthropods from an ancestor with homonomous trunk segments, a process that included the subdivision of the insect body plan into thorax and abdomen (tagmosis). In this study, we ask the following questions: when did all of the arthropod Hox genes arise, and was this before or during the arthropod radiation? And, if Ubx and abd-A are present in animals with homonomous trunks, how are these genes deployed?

# **Results and discussion** Myriapod and onychophoran *Hox* genes

# To address these questions, we first examined the *Hox* genes from a myriapod, the scolopendromorph centipede *Ethmostigmus rubripes*. In order to identify *Hox* orthologs unambiguously, gene sequences obtained from an initial PCR survey were further extended using vector ligation and degenerate PCR. Characteristic residues within and flanking the homeodomain enabled the identification of centipede orthologs of the *labial (lab)*, *Deformed (Dfd)*, *Sex combs reduced (Scr)*, *Antennapedia (Antp)*, *Ubx* and *abd-A* genes (Figure 1a). The presence of the trunk genes *Ubx* and *abd-A* in the homonomously organized centipede demonstrates that these genes are not unique to tagmatized arthropods, and suggests that they might have been present in even more primitive arthropods.

In order to trace the origin of the Ubx and abd-A genes further back through evolution, we cloned the Hox genes from the Australian onychophoran Acanthokara kaputensis. As the Onychophora are considered to be a sister group to the arthropods [2,3] with a fossil record extending back to the late Early Cambrian, the complement of Hox genes shared between Onychophora and arthropods would reflect the condition prior to the origin and radiation of the arthropods and perhaps the entire lobopodian/arthropod clade. A PCR survey and the analysis of larger genomic clones revealed that this onychophoran possesses candidate orthologs of all of the Hox genes found in Drosophila. Sequence alignments have allowed the identification of onychophoran lab, proboscipedia (pb), Dfd, Scr, Ubx, abd-A, Abd-B, and an additional Hox gene which, by virtue of its sequence and by process of elimination, is presumably the Antp gene (Figure 1a). The onychophoran Hox genes are more similar (in deduced amino-acid sequence) to their arthropod orthologs than to annelid Hox genes (Figure 1a),

a finding that is consistent with a closer affinity between Onychophora and arthropods than between either group and annelids [2,3]. In particular, the conserved regions of Onychophora Ubx (O-Ubx) and Onychophora abd-A (O-abd-A) extend into the sequence downstream of the homeodomain, thus identifying these genes to be orthologs of the arthropod Ubx and abd-A genes.

# Onychophoran Ubx gene structure

Drosophila Hox gene structure is unusually complex when compared with vertebrates and annelids. Because the analysis of onychophoran Hox genes could shed light on the nature of Hox gene structure before the arthropod radiation and the evolution of insects, we cloned and characterized the entire reading frame of O-Ubx (Figure 1b). The O-Ubx protein is encoded by two exons; the first exon contains three short regions of amino-acid similarity with the first exon of Drosophila Ubx, but is about 120 amino acid residues shorter than the Drosophila equivalent (Figure 1b). A single small intron is located in a conserved position just upstream of the homeodomain-containing exon (Figure 1b). Thus, the O-Ubx gene is much more compact (~2.5 kb) than the equivalent region of the Ubxgene in Drosophila (73 kb) [31], and is more similar in size and organization to chordate Hox genes.

# Independent Hox gene duplication events in the annelid and onychophoran/arthropod lineages

The presence of Ubx and abd-A orthologs in Onychophora and their absence in annelids suggests that these two genes have arisen more recently than other Hox genes. To analyze the origins of Ubx and abd-A, we aligned the amino-acid sequences of the homeodomain and flanking regions of onychophoran and arthropod Ubx and abd-A proteins and the annelid Lox2 [28,29] and Lox4 proteins [26] to construct a gene tree. Both parsimony and distance analyses suggest that a single Hox gene precursor existed in the common ancestor of both annelids and the onychophoran/arthropod clade and that this gene duplicated and diverged independently in each lineage to create the gene pairs Ubx/abd-A and Lox2/Lox4 ([26,27]; Figure 2a). The presence of Ubx and abd-A in the onychophoran shows that the duplication event that created these genes occurred before the separation of the onychophoran and arthropod lineages and the diversification of arthropod body plans.

# The ftz gene predates the arthropods

The centipede and the onychophoran each possess an additional *Antp*-class gene that is not obviously orthologous to any *Drosophila Hox* gene. Alignment of these centipede and onychophoran genes with the *Drosophila fushi tarazu* (*ftz*) gene [32], other insect *ftz* orthologs [33,34], and the crustacean Hx1 gene [30] along with gene-tree analysis suggests that the centipede and onychophoran genes are most closely related to each other and to the *ftz*-related genes and

# Figure 1

Identification of centipede and onychophoran Hox gene orthologs. (a) Alignment of the homeodomain and flanking sequences of arthropod [30], onychophoran and annelid [26--29,44] Hox genes. Orthologous genes are grouped except for the Ubx/abd-A/Lox2/Lox4 genes which are shown together to facilitate direct comparison. Sequences that extend beyond the homeodomain and flanking regions of homology are truncated. Within the homeodomain. residues identical to the Drosophila Antp gene are indicated with dashes. The onychophoran Antp gene was identified on the basis of its sequence homology to insect Antp genes and is ruled out as an Scr candidate because a large unambiguous Scr clone was obtained. In the regions flanking the homeodomain, boxed residues indicate identity among a majority of arthropod sequences (Lox2/Lox4 residues are boxed when they match either the arthropod Ubx or abd-A majority). The epitope recognized by the FP6.87 monoclonal antibody is marked by asterisks. Gaps in the alignment are indicated by a full stop. Intron/exon boundaries are inferred from consensus splice sequences. Af = Artemia franciscana; Cen = centipede; Dm = Drosophila melanogaster; Hm = Hirudomedicinalis; Hro = Helobdella robusta; Htr = Helobdella triserialis; Ony = Onychophora. (b) Comparison of the genomic structure and conserved motifs of the Drosophila and onychophoran Ubx genes. The coding regions of the Drosophila and onychophoran Ubx exons are shown as open boxes separated by introns of the indicated sizes. The 3' exon of each gene contains the homeodomain (light grey region), and the 5' exon encodes three motifs that are shared between the Drosophila and onychophoran genes: MxSxFExA, NGYKxxW, and FYPWM (dark grey boxes; exact sequences for each motif shown in alignment). The onychophoran Ubx intron does not appear to contain any microexons similar to those in Drosophila Ubx.



thus we suggest that they are *ftz* orthologs (Figure 2b,c). This group of genes shows significantly more amino-acid variation within the homeodomain than typical *Hox* genes. However, all of these genes except *Drosophila ftz* have the YPWM tetrapeptide motif, suggesting that they are related to *Hox* genes. As no *ftz* orthologs have yet been identified outside of Onychophora and arthropods, *ftz* may represent a rapidly evolving *Hox* gene unique to this clade.

# Evolution of *Hox* gene regulation in animals with homonomous trunks

The complement of myriapod and onychophoran *Hox* genes demonstrates that the segmental diversity of arthropods evolved without an increase in *Hox* gene number. The evolution of arthropod segmental diversity

must therefore have involved regulatory changes in Hox genes and/or their targets. More specifically, as some of the major differences between arthropods involve the tagmosis and segmental diversity of the trunk, it is possible that the regulation of Ubx and abd-A could differ significantly between different arthropod classes. To address this possibility, we examined the expression domains of Ubx and abd-A in the centipede embryo. Using the monoclonal antibody FP6.87 [35], which recognizes an epitope found in both the centipede Ubx and abd-A proteins (Figure 1a), we observed antigen expression over almost the entire trunk of the embryo (Figure 3a,b). The initial anterior boundary of Ubx/abd-A expression in the body wall is in the T2 segment and appears parasegmental (Figure 3a inset), and in later stages of development the boundary shifts





The *Ubx*, *abd-A*, and *ftz* genes are unique to the onychophoran/arthropod clade. (a) Phylogenetic gene tree indicating the relationships of *Ubx*, *abd-A*, *Lox2* and *Lox4*. The orthology relationships of the onychophoran *Ubx* and *abd-A* genes are highly supported, as indicated by the monophyly of the *Ubx* and *abd-A* genes of onychophora/arthropods. This and other phylogenetic analyses [26,27] suggest that the gene pairs *Ubx/abd-A* and *Lox2/Lox4* arose from independent duplications in the Onychophora/arthropod lineage and the annelid lineage, respectively. *Antp* orthologs from both arthropods and annelids are used as the outgroup. (b) Monophyly of *ftz* orthologs. The orthology of onychophoran and arthropod *ftz* genes [30,32–34] is indicated on a phylogenetic gene tree with the *Antp*-class *Drosophila* 

forward to the anterior of the T2 segment (Figure 3b). The *Ubx/abd-A* boundary correlates with the transition in appendage morphology from the poison claw (T1 segment) to the first walking leg (T2 segment; Figure 3b).

The onychophoran body plan and appendage organization are simpler than that of arthropods, with only three distinct head segments followed by a series of homonomous trunk segments, and may reflect a more primitive condition. Given the expression of *Ubx/abd-A* across the trunk of myriapods and crustaceans [36,37], one might expect that these *Hox* genes would be expressed across the lobopod-bearing trunk. We again used the monoclonal antibody FP6.87 to determine the deployment of *Ubx/abd-A* in onychophoran embryos. Although there is a substitution in the presumed epitope recognized by this antibody in O-Ubx (Figure 1a), the antibody recognizes both the O-Ubx and O-abd-A proteins (see Materials and methods). Surprisingly, expression homeodomain sequences used as an outgroup. For all trees, branch lengths are proportional to the changes per amino-acid position (scale shown) and bootstrap values calculated by least-squares distance methods are shown at each node. Af = Artemia franciscana; Cen = centipede; Dm = Drosophila melanogaster; Hro = Helobdellarobusta; Hm = Hirudo medicinalis; Ony = onychophoran; Sg = Schistocerca gregaria; Tc = Tribolium castaneda. (c) Alignmentof ftz orthologs. Amino-acid sequences are aligned with <math>DrosophilaAntp from the YPWM tetrapeptide motif through the homeodomain. The locations of the introns are indicated by (>) and gaps are indicated by a full stop. The presence of the YPWM motif in the centipede gene is implied by the primer used to clone this sequence (see Materials and methods).

of *Ubx/abd-A* is restricted to the very posterior end of the onychophoran embryo, in the last pair of lobopods and in the terminus (Figure 3c). This *Ubx/abd-A* boundary may correlate with a cryptic transition in segmental identity in this species, as the last lobopod of some fossil lobopodians and extant onychophorans is truncated or vestigial [15,38].

The marked difference in the expression patterns of *Ubx/abd-A* in the centipede and the onychophoran indicates that their homonomous trunks are not regulated by the same *Hox* genes. Although the deployment of *Ubx/abd-A* in the common ancestor of Onychophora and arthropods cannot be determined, it is clear that the anterior boundary of *Ubx/abd-A* expression must have shifted along the anteroposterior axis in one or both lineages. This shift might have been achieved by changes in the regulation of *Hox* genes and/or by the addition or loss of segments. Importantly, the boundary of *Ubx/abd-A* expression consistently correlates

# Figure 3

Regulation of Ubx/abd-A expression in embryos with homonomous trunks. Doublelabel antibody stains of Ubx/abd-A (red) and Dll (green) expression in centipede and onychophoran embryos. (a) Stage 4 centipede embryo [45] shown with anterior to the left. Dll is expressed in all body outgrowths, including the antenna (Ant), mouthparts (Md, mandible; Mx1, Maxillary segment 1; Mx2, Maxillary segment 2), maxilliped (T1) and walking legs (T2-T22) [45]. The anterior boundary of Ubx/abd-A expression lies in T2, where staining appears to be parasegmental in the body wall but the entire T2 limb stains. (Inset: the arrows indicate expression around the base of the limb bud and the anterior boundary in the body wall.) The overlap of Ubx/abd-A and Dll expression in the trunk limbs is shown in yellow. (b) The right side of a slightly older centipede embryo is shown with anterior at the top left. The larger antenna obscures the mandibular and maxillary segments. The anterior boundary of Ubx/abd-A has moved to the anterior of the T2 segment in the body wall, but staining in the nervous system (ns) is apparently parasegmental. Ubx/abd-A expression continues to the posterior of the embryo at this stage. (c) Onychophoran embryo shown with anterior at the top left. The antenna (Ant), jaws (J), slime papillae (SP), lobopods (L1-L15) and some cells in the body wall express Dll. Ubx/abd-A are expressed in the last pair of lobopods (L15) and in the terminus. A higher magnification view of the posterior segments (inset) shows



that the *Ubx/abd-A* boundary lies within the segment bearing the last lobopods (arrow) and that the antigens are localized, as expected, to cell nuclei. *Dll* and *Ubx/abd-A* 

are co-expressed in the last lobopod and in some cells in the terminus; this overlap of *Ubx/abd-A* and *Dll* expression is shown in yellow.

with a transition in appendage morphology in the trunk of each animal ([36,37] and summarized in Figure 4b). Our data suggest that the expression domains of *Ubx/abd-A* do not demarcate homologous body regions in these taxa, but instead, evolutionary changes in *Ubx/abd-A* deployment underlie the diversification of arthropod body plans that are evident in the Cambrian fossil record.

# Hox regulation of specific target genes differs between taxa

The *Hox* genes control segmental identity by regulating the expression of downstream target genes. In insects, for example, the *Ubx* and *abd-A* gene products suppress limb formation in the abdomen via repression of the *Distal-less* (*DII*) gene [39]; however, *DII* is not repressed in the *Ubx/abd-A* domains in centipede (Figure 3a,b) or onychophoran trunks (Figure 3c) at the stages we surveyed, nor in crustacean trunks [37,39], providing evidence that *Hox*-mediated repression of *DII* evolved in the insect lineage to sculpt the distinctive limbless insect abdomen. Thus, the evolution of *Hox* regulation of target genes is a second developmental mechanism underlying the diversification of arthropod body plans.

# Conclusions

Our results demonstrate that the entire set of arthropod Hox genes was present in the onychophoran/arthropod ancestor, and that no new Hox genes were required to catalyze the radiation of the arthropods. Instead, changes in the regulation of trunk Hox genes along the anteroposterior axis and in Hox regulation of major downstream target genes appear to have enabled the morphological diversification of arthropod body patterns. Indeed, if certain phylogenies that depict Onychophora as a primitive lobopodian are correct [14], the diversification of arthropod Hox genes would predate the lobopodian radiation as well. The fossil record of Onychophora, other lobopodians and arthropods extends to the Late Early Cambrian (~530 million years ago) [14,15] and the appearance of these taxa is contemporaneous with the 'explosion' of other taxa. Hox gene diversification must then have predated this period and could extend to the base of the Cambrian (~543 million years ago) or earlier.

Three new *Hox* genes did arise early in the onychophoran/arthropod clade before the divergence of the Onychophora and the arthropod radiation (Figure 4a). The *Ubx*, *abd-A* and *ftz* genes may represent synapomorphies

# Figure 4

Hox gene evolution and the regulation of Ubx/abd-A expression in arthropods and onvchophorans. (a) The Hox genes expanded in both the annelids and the onychophoran/arthropod clade by independent duplication events in each lineage. The Ubx, abd-A and ftz genes are unique to the Onvchophora and arthropods and did not exist in the ancient protostome ancestor of annelids and Onychophora/arthropods. Some annelid Lox genes are not shown. (b) The expression domains of Ubx (red) and abd-A (purple) are indicated on the body plans of several arthropods and an onychophoran. Where the expression domains of Ubx and abd-A overlap (Drosophila, Artemia) or are not yet distinguishable (centipede and onychophoran), both red and purple are shown. Transient or low level patterns of Ubx expression (for example, in Drosophila T2 segment) [46] are not shown. The anterior boundary of Ubx/abd-A differs between the arthropod classes and the Onychophora, and correlates with transitions in appendage morphology along the anteroposterior axis in each animal: in insects (Drosophila), the boundary falls in T3, between the thorax (walking legs) and abdomen (no



appendages), and the wing and haltere; in crustacea (*Artemia*), the boundary is between gnathal and thoracic segments that bear distinct limbs; in centipedes, the boundary is located between the poison claw (T1 segment) and the first walking leg (T2 segment); and in Onychophora, the boundary lies between the penultimate and terminal lobopod. The differences in the *Ubx/abd-A* expression domains among arthropods and Onychophora suggest that the evolution of arthropod segmental diversity arose through changes in the regulation of *Hox* genes.

(shared derived characters) for the onychophoran/arthropod clade, as no clear orthologs have yet been identified in other taxa. This raises the exciting possibility that the presence of these genes could be used as phylogenetic tools to resolve the relationship of arthropods to other taxa. The traditional view of annelids and arthropods as sister taxa has been challenged [4] and most protostome relationships are not resolved [2]. Thorough characterization of the *Hox* genes from other protostomes might help to identify the sister taxa of the lobopodian/arthropod clade.

# Materials and methods

# Cloning of Hox genes

Centipede and onychophoran adult DNA was purified from carcasses after the gut contents were removed to prevent potential contamination with prey items. Hox gene fragments were amplified from centipede and onychophoran genomic DNA using the degenerate Hox primers A, B, E and F [40,41]. Onychophoran abd-B was amplified using the Hox F primer and two nested abd-B specific primers (RNK-3 and RNK-4) [36]. PCR products were ligated into the TA Cloning vector pCR2.1 (Invitrogen). Multiple copies of all PCR products were sequenced, however, only a single clone of Dfd was recovered from each species. Two different nucleic acid sequences encoding the same O-abd-B amino-acid sequence were obtained. A caudal ortholog from Onychophora and an engrailed ortholog from the centipede were also identified (data not shown). No orthologs of Hox3 were identified, perhaps because of substitutions in the Hox degenerate primer binding sites. Similarly, no centipede pb or Abd-B orthologs were cloned, but their presence in Onychophora and chordates demonstrates their existence before the evolution of arthropods. DNA flanking the initial centipede and onychophoran Hox clones was isolated by vector-ligation PCR. Genomic DNA was digested with restriction enzymes and ligated and into

pBluescript (Stratagene) or pCR2.1. DNA was then amplified using nested gene-specific primers designed from previously isolated PCR products and vector-specific primers. *Hox* gene products were detected by Southern blotting analysis using <sup>32</sup>P-labelled DNA encoding the *Drosophila*, centipede or onychophoran homeodomains as probes. The centipede *ftz* gene fragment was amplified using a degenerate primer for the POIYPWM (single-letter amino-acid code) sequence (dGAGGATC-CCCNCARATHTAYCCNTGGATG) and the F primer. The centipede *Scr* sequence was extended in the 5' direction using a degenerate primer recognizing ETKRQRT (dGACACNAARMGNCARMGNAC) in nested PCR with the *Hox* B and F primers and a gene-specific primer.Onychophora *Ubx* (2648 bp) and *Scr* (1296 bp) genomic clones were isolated from two subgenomic lambda gt10 libraries after identifying bands of interest on Southern blots of *Eco*RI-digested genomic DNA.

### GenBank accession numbers

Centipede gene sequences have accession numbers AFO10172--AFO10179. Onychophoran gene sequences have accession numbers AFO11272-AFO11282.

# Gene trees

Gene trees were constructed with PHYLIP (Phylogeny Inference Package v3.572c). For the *Ubx/abd-A/Lox2/Lox4* tree, up to 83 amino acids containing the homeodomains and flanking conserved sequences were aligned as shown in Figure 1a. The *ftz* alignment consisted of homeodomain sequences only (60 amino acids). The Seqboot program resampled each data set 100 times for bootstrap analysis. The Protdist program calculated distances using a PAM–Dayhoff distance matrix [42] and the Fitch program used the Fitch–Margoliash least-squares model to search for the best trees. Maximum parsimony trees were constructed with the Protpars program. Bootstrap values were calculated with the Consense program.

Immunohistochemistry and analysis of antibody reactivity Immunohistochemical staining with the FP6.87 and *DI*/ antibodies was performed as previously described [37,43]. In order to examine the reactivity of O-Ubx with the FP6.87 antibody, the entire exons encoding the homeodomains of O-Ubx and O-abd-A were cloned in-frame into the pET28a protein expression vector (Novagen) and expressed in *E. coli* strain BL21 DE3. Western blots of the induced lysates containing onychophoran homeodomain fusions were then assayed with the monoclonal antibody FP6.87 [35]. The putative epitope for this antibody has been mapped to six amino acids that are shared between Ubx and abd-A but are not present in Antp [35,36]. There is a single amino-acid replacement in one of these six residues in O-Ubx (Figure 1a). The O-Ubx and O-abd-A proteins were detected equally well by the FP6.87 antibody by western blotting analysis, indicating that the observed *in situ* immunoreactivity includes both proteins.

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### References

- Conway Morris S: The fossil record and the early evolution of the Metazoa. Nature 1993, 361:219-225.
- Valentine J, Erwin D, Jablonski D: Developmental evolution of metazoan bodyplans: The fossil evidence. Review. Dev Biol 1996, 173:373-381.
- 3. Wheeler W, Cartwright P, Hayashi C: Arthropod phylogeny: a combined approach. *Cladistics* 1993, **9**:1-39.
- Eernisse D, Albert J, Anderson F: Annelida and arthropoda are not sister taxa: a phlyogenetic analysis of spiralian metazoan morphology. Syst Biol 1992, 41:305-330.
- Raff R, Marshall C, Turbeville J: Using DNA sequences to unravel the Cambrian radiation of the animal phyla. *Annu Rev Ecol Syst* 1994, 25:351-375.
- Lake J: Origin of the Metazoa. Proc Natl Acad Sci USA 1990, 87:763-766.
- Fortey R: The Cambrian evolutionary explosion decoupling cladogenesis from morphological disparity. *Biol J Linn Soc* 1996, 57:13-33.
- Erwin D, Valentine J, Jablonski K: The origin of animal body plans. Am Sci 1997, 85:126-137.
- 9. Conway Morris S: Molecular clocks: defusing the Cambrian 'explosion'? Curr Biol 1997, 7:R71-R74.
- Wray G, Levinton J, Shapiro L: Molecular evidence for deep precambrian divergences among metazoan phyla. Science 1996, 274:568-573.
- Davidson E, Peterson K, Cameron R: Origin of bilateran body plans: evolution of developmental regulatory mechanisms. *Science* 1995, 270:1319-1325.
- 12. Boore J, Collins T, Stanton D, Daehler L, Brown W: Deducing the pattern of arthropod phylogeny from mitochondrial DNA rearrangements. *Nature* 1995, **376**:163-165.
- Friedrich M, Tautz D: Ribosomal DNA phylogeny of the major extant arthropod classes and the evolution of myriapods. *Nature* 1995, 376:165-167.
- 14. Budd G: The morphology of *Opabinia regalis* and the reconstruction of the arthropod stem-group. *Lethaia* 1996, **29:**1-14.
- 15. Hou X, Bergström J: Cambrian lobopodians ancestors of extant onychophorans? Zoo J Linn Soc 1995, 114:3-19.
- Ramsköld L, Xianguang H: New early Cambrian animal and onychophoran affinities of enigmatic metazoans. *Nature* 1991, 351:225-228.
- Lewis EB: A gene complex controlling segmentation in Drosophila. Nature 1978, 276:565-570.
- Akam M: The molecular basis for metameric pattern in the Drosophila embryo. Development 1987, 101:1-22.
- Akam M, Averof M, Castelli-Gair J, Dawes R, Falciani F, Ferrier D: The evolving role of Hox genes in arthropods. *Development* 1994, Supplement:209-215.
- 20. Carroll S: Homeotic genes and the evolution of arthropods and chordates (Review). *Nature* 1995, **376**:479-485.
- 21. Krumlauf R: *Hox* genes in vertebrate development. *Cell* 1994, **78:**191-201.

- 22. Schubert F, Nieselt-Struwe K, Gruss P: **The** *Antennapedia*-type homeobox genes have evolved from three precursors separated early in metazoan evolution. *Proc Natl Acad Sci USA* 1993, **90:**143-147.
- Garcia-Fernández J, Holland P: Archetypal organization of the amphioxus Hox gene cluster. Nature 1994, 370:563-566.
  Invine S, Martindale M: Cellular and molecular mechanisms of
- Irvine S, Martindale M: Cellular and molecular mechanisms of segmentation in annelids. Cell Dev Biol 1996, 7:593-604.
- Dick M, Buss L: A PCR-based survey of homeobox genes in Ctenodrilus serratus (Annelida: Polychaeta). Mol Phylogen Evol 1994, 3:146-158.
- Wong V, Aisemberg G, Wen-Biao G, Macagno E: The leech homeobox gene Lox4 may determine segmental differentiation of identified neurons. J Neurosci 1995, 15:5551-5559.
- Kourakis M, Master VA, Lokhorst DK, Nardelli-Haefliger D, Wedeen CJ, Martindale MQ, Shankland M: Conserved anterior boundaries of *Hox* gene expression in the central nervous system of the leach Helobdella. *Dev Biol* 1997, in press.
- Wysocka-Diller JW, Aisemberg GO, Baumgarten M, Levine M, Macagno ER: Characterization of a homologue of bithorax-complex genes in the leech *Hirudo medicinalis*. *Nature* 1989, 341:760-763.
  Nardelli-Haefliger D, Shankland M: Lox2, a putative leech segment
- Nardelli-Haefliger D, Shankland M: Lox2, a putative leech segment identity gene, is expressed in the same segmental domain in different stem cell lineages. Development 1992, 116:697-710.
- Averof M, Akam M: HOM/Hox genes of Artemia: implications for the origin of insect and crustacean body plans. Curr Biol 1993, 3:73-78.
- Hogness DS, Lipshitz HD, Beachy PA, Peattie DA, Saint RB, Goldschmidt-Clermont M, et al.: Regulation and products of the Ubx domain of the bithorax complex. Cold Spring Harbor Symp Quant Biol 1985, 50:181-194.
- Laughon A, Scott MP: Sequence of a Drosophila segmentation gene: protein structure homology with DNA-binding proteins. Nature 1984, 310:25-31.
- Dawes R, Dawson I, Falciani F, Tear G, Akam M: Dax, a locust Hox gene related to fushi-tarazu but showing no pair-rule expression. Development 1994, 120:1561-1572.
- Brown S, Hilgenfeld R, Denell R: The beetle Tribolium castaneum has a fushi tarazu homolog expressed in stripes during segmentation. Proc Natl Acad Sci USA 1994, 91:12922-12926.
- Kelsh R, Weinzierl R, White R, Akam M: Homeotic gene expression in the locust Schistocerca: an antibody that detects conserved epitopes in Ultrabithorax and abdominal-A genes. Dev Genet 1994, 15:19-31.
- Averof M, Akam M: Hox genes and the diversification of insectcrustacean body plans. Nature 1995, 376:420-423.
- Panganiban G, Sebring A, Nagy L, Carroll S: The development of crustacean limbs and the evolution of arthropods. *Science* 1995, 270:1363-1366.
- Ruhberg H: Die Peripatopsidae (Onychophora): Systematik, ökologie, Chorologie und phylogenetische Aspekte. Zoologica 1985, 46:1-184.
- Vachon G, Cohen B, Pfeifle C, McGuffin M, Botas J, Cohen S: Homeotic genes of the *Bithorax* complex repress limb development in the abdomen of the *Drosophila* embryo through the target gene. *Cell* 1992, **71**:437-450.
- Murtha M, Leckman J, Ruddle F: Detection of homeobox genes in development and evolution. Proc Natl Acad Sci USA 1991, 88:10711-10715.
- Pendleton J, Nagai B, Murtha M, Ruddle F: Expansion of the Hox gene family and the evolution of chordates. Proc Natl Acad Sci USA 1993, 90:6300-6304.
- 42. Dayhoff M, Schwartz R, Orcutt B: *Atlas of Protein Sequence Structure.* Silver Spring: National Biomedical Research Foundation; 1978.
- Warren R, Nagy L, Selegue J, Gates J, Carroll S: Evolution of homeotic gene regulation and function in flies and butterflies. *Nature* 1994, 372:458-461.
- Aisemberg G, Macagno E: Lox1, an Antennapedia-class homeobox gene, is expressed during leech gangliogenesis in both transient and stable central neurons. Dev Biol 1994, 161:455-465.
- Whitington P, Meier T, King P: Segmentation, neurogenesis and formation of early axonal pathways in the centipede, *Ethmostigmus rubripes* (Brandt). *Roux's Arch Dev Biol* 1991, 199:349-363.
- Brower DL: Ultrabithorax gene expression in Drosophila imaginal discs and larval nervous system. Development 1987, 101:83-92.

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