# REVIEW

# *Distal-less* homeobox genes of insects and spiders: genomic organization, function, regulation and evolution

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**Abstract** The *Distal-less (Dll)* genes are homeodomain transcription factors that are present in most Metazoa and in representatives of all investigated arthropod groups. In *Drosophila*, the best studied insect, *Dll* plays an essential role in forming the proximodistal axis of the legs, antennae and analia, and in specifying antennal identity. The initiation of *Dll* expression in clusters of cells in mid-lateral regions of the *Drosophila* embryo represents the earliest genetic marker of limbs. *Dll* genes are involved in the development of the peripheral nervous system and sensitive organs, and they also function as master regulators of black pigmentation in some insect lineages. Here we analyze the complete genomes of six insects, the nematode *Caenorhabditis elegans* and *Homo sapiens*, as well as multiple *Dll* sequences available in databases in order to examine the structure and protein features of these genes. We also review the function, expression, regulation and evolution of arthropod *Dll* genes with emphasis on insects and spiders.

Key words *Distal-less*; evolution; function; genomic organization; regulation

# Introduction

The Distal-less (Dll) gene is a homeodomain transcription factor, named Dlx in vertebrates and Dll in all other metazoans (Zerucha & Ekker, 2000). Dll is expressed in representatives of onychophorans and many arthropod groups including chelicerates, myriapods, crustaceans and insects (Williams & Nagy, 1996; Panganiban *et al.*, 1997; Popadic *et al.*, 1998; Scholtz *et al.*, 1998; Thomas & Telford, 1999; Mittmann & Scholtz, 2001; Pechmann *et al.*, 2010). Dll has its earliest origin in metazoan anteroposterior head axis patterning, and subsequently it was likely co-opted for proximodistal patterning of body appendages in arthropods, including serial homologous and non-homologous appendages (Lemons *et al.*, 2010).

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It is in this role of appendage development that Dll is best known – a role found to occur in virtually all arthropod species where *Dll* expression is known (Robledo et al., 2002). That the evolution of appendages is central to the diversification and success of arthropods means that Dll likely has played a key role in the success of this large group (Pechmann et al., 2010). Such appendages include the legs, antennae, wing, analia and mouthparts in insects (Casci, 2002; Lin et al., 2014), coxapophyses and ocularia in harvestmen (Sharma et al., 2013), mouthparts, legs and spinnerets in spiders (Abzhanov et al., 1999; Prpic & Damen, 2004; Pechmann & Prpic, 2009), nasi in termites (Toga et al., 2012), and horns in beetles (Moczek & Nagy, 2005). Dll has also been involved in the development of the peripheral nervous system and sensitive organs (Casci, 2002; Plavicki et al., 2012), in butterfly eyespots (Carroll et al., 1994; Brakefield et al., 1996; Beldade et al., 2002), and in melanin synthesis in the wings of both flies and butterflies (Arnoult et al., 2013; Monteiro et al., 2013). Vertebrate Dlx genes are involved in a variety of other developmental processes ranging from neurogenesis to hematopoiesis (Depew et al., 1999;

Shimamoto *et al.*, 2000). Earlier reviews of vertebrate *Dlx* genes and *Drosophila melanogaster Dll* (Panganiban, 2000; Zerucha & Ekker, 2000; Panganiban & Rubenstein, 2002) did not include much information from other protostomes.

Here we review information pertaining to *Dll* across all available protostome systems with emphasis on insects and spiders, including recent *Drosophila* work. We start by analyzing the complete genomes of seven protostome species and a deuterostome outgroup (humans), and many *Dll* sequences available in the databases in order to examine evolution of copy number and gene sequence. We also describe the *Dll* gene structure and its main sequence features, and summarize the function, expression and regulation of protostome *Dll* genes, mainly in insects and spiders.

# Genomic organization and gene structure

We investigated the complete genome of seven protostome species (six insects and one nematode) as well as a deuterostome outgroup, humans, and determined that there is only one *Dll* copy in protostomes (Table 1). For comparison, the basal chordate amphioxus also has a single Dlx gene (Holland et al., 1996), whereas the more derived urochordate, Ciona intestinalis, and the vertebrates, mouse and humans, have two and six *Dlx* genes, respectively (Digregorio et al., 1995; Zerucha & Ekker, 2000). The six vertebrate Dlx genes can be grouped into two clades (Dlx1/4/6 and Dlx2/3/5) based on sequence similarity, and are organized as three gene pairs (Dlx 1/2,Dlx3/4 and Dlx5/6) with each pair closely linked on the genome (see Table 1, [Zerucha & Ekker, 2000]). Two additional Dlx genes have been identified in zebrafish, which are not linked to each other and do not appear to exist in other vertebrates such as mammals (Stock et al., 1996).

Primary messenger RNA (mRNA) transcript sizes (including introns) of insect *Dll* genes (19 773–72 999 bp) are much larger than that of human *Dlx* (3313–5396 bp) and *Caenorhabditis elegans Dll* (3303 bp) genes (Table 1). Correspondingly, the exon number (5–7 exons), amino acid length (312–356 amino acid [aa]) and molecular mass (31.74–38.36 kDa) of insect *Dll* genes are also larger than those of human *Dlx* (3 exons, 240–328 aa and 26.26–34.24 kDa) and *C. elegans Dll* (5 exons, 273 aa and 30.16 kDa) genes. Interestingly, one of the central introns of insect *Dll* genes spans more than half of the gene's primary transcript size and is largely responsible for the observed difference in gene size relative to the

human and *C. elegans* homologues (Fig. 1). This large intron is the third intron in *D. melanogaster*, *Anopheles gambiae*, *Aedes aegypti* and *Apis mellifera* and the second intron in *Bombyx mori* and *Tribolium castaneum*.

Two possible *Dll* splicing patterns, RA and RB, were detected in the genomes of the Diptera, Lepidoptera and Hymenoptera species investigated, whereas a single splice variant was found for T. castaneum and C. elegans (Table 1, Fig. 1). The RA splicing pattern is a modification of the RB splice variant where an extra exon encoding a conserved VWPAV motif is added to the N-terminal end of the RA protein. These two splice variants match all Dll mRNA sequences available in the databases for these insect orders except that of the lepidopteran Precis coenia (AF404110.1), which has 27 unique residues at the Nterminal end instead of the common VWPAV motif. All the alternatively spliced protein sequences share a common NPS(G)LL(V)T motif at their C-terminus, whereas T. castaneum and C. elegans lack this motif. By comparison, vertebrate *Dlx* genes produce multiple transcripts by alternative transcription initiation (e.g. Dlx1) (McGuinness et al., 1996) as well as alternative splicing (e.g. Dlx4 and *Dlx5*) (Liu *et al.*, 1997).

The *Dll/Dlx* genes in the eight genomes investigated have different sized exons and introns; however, the homeodomain is commonly split between codons 43 and 44 by the largest intron in insects, C. elegans (1414 bp), and humans (431-2019 bp) (Fig. 1). This shared splice site suggests the presence of an ancient homologous intron across all these species. The two homeodomain sections encode 43 and 17 aa in their anterior and posterior exons, respectively, with the exception of C. elegans whose anterior 43 aa homeodomain section is additionally interrupted by a short intron and split into 19 and 24 codons. The first two exons of D. melanogaster, An. gambiae, Ae. aegypti and A. mellifera Dll genes, and the unique first exon of B. mori and T. castaneum encode the NM-1 and NM-2 regions of Dll (Fig. 2A). These regions are conserved regions, which will be described more fully later. The third exon of D. melanogaster, An. gambiae, Ae. aegypti and A. mellifera and the second exon of B. mori and T. castaneum are homologous and display the highest degree of sequence similarity across these species, perhaps because these exons include the largest portion of the homeodomain. Hereafter exons are quite varied in number and sequence. The last exon of the Dll-RB splice variant in all insect species but T. castaneum, encodes the conserved motif NPS(G)LL(V)T at the C-terminal end, whereas the Dll-RA variant has an additional exon encoding the conserved motif VWPAV in all insects examined (Fig. 2A).

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			Accessi	on number			
Gene/splicing	mRNA	Amino acid.	Chromosomal location $^{\dagger}$	Exon no.‡	Amino acid (length)	Mass (kDa)	Genome size (bp) <sup>¶</sup>
Drosophila melanogaster							
DII-RA	NM_079133.1	NP_523857.1	2R+ 60E2	7(7)	327	35.260	20334
DII-RB	NM_166689.1	NP_726486.1	2R+ 60E2	(9)9	322	34.71	19773
Anopheles gambiae							
DII-RA	XM_308706.1	XP_308706.1	2L-	7(7)	300	32.29	48680
DII-RB	Newly predicted	Newly predicted	2L-	(9)9	295	31.74	48506
Aedes aegypti							
DII-RA	Newly predicted	Newly predicted	N/A	7(7)	301	32.43	72999
Dll-RB	Newly predicted	Newly predicted	N/A	6(6)	296	31.88	72825
Bombyx mori							
DII-RA	Newly predicted	Newly predicted	N/A	7(7)	356	38.36	43839
DII-RB	Newly predicted	Newly predicted	N/A	(9)9	351	37.81	43539
Apis mellifera	4 6	4		к И			
DII-RA	Newly predicted	Newly predicted	Group 13 +	(9)9	349	36.28	69484
DII-RB	Newly predicted	Newly predicted	Group13 +	5(5)	344	35.73	68848
Tribolium castaneum							
DII	AF317551.1	AAG39634.1	Contig1484-, Contig1541+	5(6)	312	34.40	61104
Caenorhabditis elegans			1				
Dll (Ceh-43)	NM_065503.2	NP_497904.1	3-	5(5)	273	30.16	3303
Homo sapiens							
DlxI	NM_178120.4	NP_835221.2	2+q31.1	3(3)	255	27.32	4195
Dlx2	NM_004405.3	NP_004396.1	2-q31.1	3(3)	328	34.24	3313
Dlx3	NM_005220.2	NP_005211.1	17 - q21.33	3(3)	287	31.74	5220
Dlx4	NM_138281.1	NP_612138.1	17 + q21.33	3(3)	240	26.26	5396
Dlx5	NM_005221.5	NP_005212.1	7-q21.3	3(3)	289	31.54	4442
Dlx6	NM_005222.2	NP_005213.2	7+ q21.3	3(3)	265	29.39	5063
<sup>†</sup> +, forward strand; –, rev unavailability of the chrom	erse strand; both pr tosomal information.	eceded by chromoso	omal name or sequence group/	contig, and fo	ollowed by location in t	he cytological	map. N/A indicates

<sup>‡</sup>Total number of exons in coding regions. The numbers in brackets include exons in untranslated regions.

<sup>¶</sup>The size corresponds to the primary mRNA transcript including introns.

Table 1 Dll genes present on the genomes of eight species.



**Fig. 1** Scaled intron/exon structures of *Dll/Dlx* genes. The filled rectangles depict protein-coding sequences, and unfilled rectangles represent untranslated regions. Introns are the open spaces between these rectangles with intron numbers partially marked.

#### **Protein sequence features**

The *Dll* homeodomain is invariable across all arthropod sequences available in the databases and only 20 of the 60 residues vary across all available animal sequences (Fig. 2A). The homeodomain codes for a homeobox that binds DNA through a helix-turn-helix structure (HTH, Fig. 2A), usually in the regulatory region of *Dll/Dlx* target genes. There are four additional motifs that were identified in this study to be relatively conserved across arthropods, NM-1 and NM-2 located at the N-terminus, and HD-5' and HD-3' connected to the homeodomain at its 5'- and 3'-ends, respectively. NM-1 is 22 aa long in D. melanogaster, with eight identical aa throughout Insecta. NM-2 is 36 aa long, with three and 20 identical aa throughout Arthropoda and Insecta, respectively. HD-5' is approximately 13 aa long, with three and five identical aa throughout Metazoa and Arthropoda, respectively. HD-3' is 12 aa long, with three identical aa throughout Arthropoda. The functions of these motifs are still unknown, but likely to mediate interactions with other proteins.

All Dll/Dlx proteins possess two conserved tryptophan (Trp) residues that are C-terminal to the homeodomain. The first (not shown) is typically followed by an aspartic acid, whereas the second (Fig. 2A) is followed by a tyrosine (Panganiban & Rubenstein, 2002). Dll/Dlx proteins lack Trp N-terminal of their homeodomains.

#### **DII** expression and function

#### Anteroposterior head axis patterning

Early *Dll* expression in cnidarians (Ryan *et al.*, 2007), and in particular the head regions of mollusks (Lee & Jacobs, 1999), hemichordates (Lowe *et al.*, 2003) and spiders (Pechmann *et al.*, 2011) suggests that an anteroposterior axis patterning role existed ancestral to arthropods and that this role was later co-opted into a proximodistal patterning role for the serial homologous appendages (Lemons *et al.*, 2010). Remarkably, RNA interference (RNAi) suppression of early *Dll* expression in spiders shows a novel gap gene-like function with the loss of the first or the first and second walking leg body segments, and corresponding legs (Pechmann *et al.*, 2011) (Figure 4).

# Legs

*Dll* is activated in the thoracic limb primordia of the fly embryo soon after gastrulation, and is one of the earliest known markers of these primordia (Cohen *et al.*, 1989; Cohen, 1990). *Dll* is first expressed in all six thoracic primordia at late embryonic stage 10 and is expressed in the entire limb thoracic primordium at stage 11 (Co-



**Fig. 2** Analysis of the *Dll* sequences of representative insects. (A) Alignments of conserved motifs. NM-1 and NM-2 are two N-terminal motifs separated by a number of residues not shown, and HD-5' and HD-3' are the motifs connected to the homeobox at its 5'-and 3'-end. The shaded residues are identical either throughout Metazoa (red), Arthropoda (blue) or Insecta (yellow). The conserved residue tryptophan and the motif NPS(G)LL(V)T in C-termini are marked in green. Dots and dashes represent gaps and unavailability in individual sequences. AME, *Apis mellifera*; BMO, *Bombyx mori*; CEL, *Caenorhabditis elegans*; CSA, *Cupiennius salei*; DME, *Drosophila melanogaster*; GMA, *Glomeris marginata*; HAS, *Homo sapiens*; OFA, *Oncopeltus fasciatus*; TCA, *Tribolium castaneum*. (B) The most parsimonious *Dll* gene tree inferred from conserved protein sequences with maximum parsimony phylogenetic analysis, which was conducted using PAUP\* v4.0b10 (Swofford, 2002). The bootstrap percentages of 1000 replicates are shown on the branches where they exceed 50%.

hen et al., 1989; Cohen, 1990). The second thoracic limb primordium gives rise to three structures: the leg, wing and a larval sensory organ called the Keilin's organ. At stage 12, *Dll* expression becomes restricted to the center of the primordia of the combined disc for wing, leg and Keilin's organ. The central cells of the combined disc were thought to contribute to the distal leg but it was later proposed that they give rise to the Keilin's organ (Fig. 3A) (Kubota et al., 2003; Bolinger & Boekhoff-Falk, 2005). The Keilin's organ precursor cells express Dll, Cut and Couch potato (Cpo) but lack the leg imaginal determinant Escargot (Esg). Distal leg disc cells are marked by the coexpression of *Dll* and *Esg*, whereas proximal cells express only Esg (Fig. 5B) (Bolinger & Boekhoff-Falk, 2005). Dll expression is lost from medial leg cells either before or during the second larval instar (Weigmann & Cohen, 1999). Late in development, during the third instar. Dll expression is activated in a proximal ring corresponding to the trochanter where it seems to play an important role in preventing mixing of proximal and medial cells (Wu & Cohen, 1999) and is expressed at low levels in the developing femur (Weigmann & Cohen, 1999). In the late pupal stage, *Dll* is expressed in the distal trochanter and tibia, and in all tarsal segments (Figs. 3B, 4) (Dong *et al.*, 2002; Panganiban & Rubenstein, 2002).

Across the insects, with the exception of dipterans that have limbless larvae, limb primordia established during embryogenesis give rise to well-formed larval limbs. Before limb primordia are morphologically discernible, *Dll* expression in *Tribolium* closely resembles that of *Drosophila*. Limb development in *Drosophila* is not visible from the outside through the larval stages until the pupal stage when a fly pupal leg resembles an embryonic leg in *Tribolium* (Beermann *et al.*, 2001). Similar patterns of *Dll* expression have been observed in several insect embryos, including lepidopterans (Zheng *et al.*, 1999), grasshoppers (Jockusch *et al.*, 2000) and crickets (Niwa *et al.*, 1997).



**Fig. 3** Expression patterns of domain genes *Dll*, *Dac* and *Hth* along the PD axis of appendages. (A) Modified from Inoue *et al.* (2002). *Dll* is expressed in the entire leg primordium in the first to third thoracic segments at stage 11 of *Drosophila* embryonic development, and restricted in the central domain at stage 15. In the late second or early third instar the leg disc is defined by three discrete domains of gene expressions: distal (*Dll*, red), medial (*Dac*, green) and proximal (*Hth*, dark blue). Following this, *Dll* and *Dac* expressions overlap in a broad domain (yellow). By the late third instar, *Dll* expression is reactivated in a proximal ring corresponding to the trochanter domain (grey), where all three proteins overlap (Postleth & Schneide, 1971; Dong *et al.*, 2001; Inoue *et al.*, 2002). (B) *Dll*, *Dac* and *Hth* expression in the antenna, leg, maxilla and labium at a late disc developmental stage. cx, coxa; tr, trochanter; fe, femur; ti, tibia; t1-t5, tarsi 1–5; cl, claw; a1–a5, antennal segments 1–5; ar, arista; cl, claw; p1–p5, palps 1–5.

Despite some discrepancies in the early leg bud, expression patterns of *Dll* in its later stages are basically consistent between dipterans, orthopterans (Jockusch *et al.*, 2000; Inoue *et al.*, 2002), hemipterans (Rogers *et al.*, 2002), coleopterans (Beermann *et al.*, 2001), myriapods (Prpic & Tautz, 2003), chelicerates (Panganiban *et al.*, 1995; Thomas & Telford, 1999; Abzhanov & Kaufman, 2000; Schoppmeier & Damen, 2001) and crustaceans (Panganiban *et al.*, 1995; Popadic *et al.*, 1998; Abzhanov & Kaufman, 2000). These data suggest a conserved role for *Dll* in patterning the distal part of all arthropod limbs (Fig. 4).

In the abdominal prolegs of the butterfly *Junonia coenia*, the expression of *Dll* is restricted to a proximal ring, and the more distal expression is lacking or highly reduced in levels (Panganiban *et al.*, 1994) (Fig. 4). However, *Dll* is not expressed in the prolegs of sawfly larvae *Neodiprion abietis*, *Diprion similis* and *Athalia rosae* (Suzuki & Palopoli, 2001; Oka *et al.*, 2010). In two crustaceans, *Triops* and *Nebalia*, *Dll* is not expressed prior to branching in all limbs, nor is it expressed in every limb branch, which postulates that *Dll* only plays a role in establishing a single proximal-distal (PD) patterning axis within these branched limbs (Williams, 1998). This suggests that sawfly prolegs may derive from these branch elements but not from the main PD axis (Oka *et al.*, 2010). In addition, the distinct *Dll* expression patterns suggest that larval prolegs have evolved independently in Hymenoptera and Lepidoptera (Suzuki & Palopoli, 2001).



**Fig. 4** Expression domains and function of *Dll* in a chimeric arthropod with insect and spider features. *Dll* functions as a gap gene in spiders. It is a positive regulator of horn size in beetles, nasus size in termites and eyespot size in butterflies. *Dll* is required for the development of the analia and all ventral appendages except mandibles (early in development), distal limbs (later in development), and it specifies antennal identity. *Dll* is required for ocularium development in harvestmen, and endites, book lungs, and spinnerets in spiders. *Dll* is also expressed in prolegs in Lepidoptera but it is still unclear whether it has a function in the development of this trait. *Dll* is required for wing margin development in flies, and for the development of multiple parts of the nervous system and sensory cells (not shown). Head appendage nomenclature used is that of insects.

In arthropods, investigation of *Dll* function has been largely focused on D. melanogaster. Dll is required for limb outgrowth and for differentiation of distal limb structures. Dll-null mutants die as embryos because they lack the rudimentary larval limbs (Cohen & Jurgens, 1989). Clones of homozygous Dll<sup>-</sup> cells are incapable of contributing to any structure but the leg coxal segment, whereas those of  $Dll^+$  develop normally (Lindsley & Zimm, 1992). Hypomorphic fly Dll alleles have been reported to result in various transformations, malformations, fusions and/or reductions of legs (in the ventral thoracic disc, trochanter, femur, tibia, tarsus and joint) (Dong et al., 2000, 2002; Beermann et al., 2001; Dworkin, 2005). Ectopic *Dll* expression in the proximal region of ventral appendages induces nonautonomous duplication of legs by the activation of *Wingless* (Wg) and *Decapentaplegic* (Dpp) (Gorfinkiel et al., 1997). The requirement for Dll in the fly femur and most of the tibia is lost by about the early third instar, and the distal tibia and the tarsus remain the only regions where Dll function is required late in development (Campbell & Tomlinson, 1998).

In the arachnid *Cupiennius*, the silencing of *Dll* by RNAi leads to missing distal portions of limbs but to normal proximal parts (Schoppmeier & Damen, 2001). However, *Dll* is also expressed in the spider's palpal coxae, and may be responsible for the endites that extend from

them (Pechmann & Prpic, 2009; Pechmann *et al.*, 2010), especially seeing as *Dll* silencing in harvestmen results in the loss of coxapophyses (Sharma *et al.*, 2013). In *Onthophagus* beetles, *Dll* down-regulation in the last larval instar led to loss or fusion of distal appendage regions in pupal and adult legs, as well as mouthparts and antennae (Moczek & Rose, 2009). In *T. castaneum*, disruptions of *Dll* function in earlier larval instars led to more severe leg disruptions, suggesting a role for this gene in maintaining the integrity of the whole larval leg (Suzuki *et al.*, 2009). In the crustacean *Parhyale hawaiensis*, *Dll* small short interfering RNA (siRNA) injections into embryos led to hatchings with truncated appendages (Liubicich *et al.*, 2009).

Phenotypic analysis indicates that distal leg (and antennal) structures are more sensitive to changes in *Dll* levels than are medial structures. Based on this, it was anticipated that *Dll* would be expressed in a graded manner along the developing PD axis, with the highest levels present distally (Cohen *et al.*, 1989). However, neither RNA *in situ* data nor antibody stainings have provided convincing support for this view. It may be that there is a shallow *Dll* gradient, not readily observed using standard techniques. Alternatively, there may be a gradient early in development that has disappeared before the third instar, which is the stage most commonly analyzed. Yet



Fig. 5 The role of *Dll* in the genetic pathway of imaginal disc differentiation and subsequent subdivision in *D. melanogaster*. A. Modified from Panganiban (2000). Dll is activated by Wg in the embryonic thorax, and repressed by Dpp dorsally and DER ventrally. Dll is not expressed in the abdomen because of repression by Ubx and AbdA. B. Modified from Bolinger and Boekhoff-Falk (2005). Wg and Dpp serve key roles in the subdivision of thoracic limb primordium through determinants such as Dll and Esg (Cohen, 1990b; Goto & Hayashi, 1997). The Keilin's organ (red) is specified by Dll and by the downstream acuaete-scute (ASC) complex. The specification involves both the activation of the neural genes Cut and Cpo and the repression of Est (Bolinger & Boekhoff-Falk, 2005). High Dpp activity (solid arrow) is required for specification of the wing disc and lower (dashed arrow) for specification of the leg disc (Raz & Shilo, 1993; Goto & Hayashi, 1997; Kubota et al., 2000). Imaginal components are patterned by Esg (leg disc) and Snail and Vg (wing disc, blue), and Dll is a determinant for distal leg (green) but not for proximal leg (purple) (Hayashi et al., 1993; Fuse et al., 1996; Bolinger & Boekhoff-Falk, 2005). C. Subdivision of eye-antenna imaginal disc by Eyeless (Ey) – Dll selectors. Dll and Ey are expressed throughout the combined eye-antenna disc early in fly development (Kumar & Moses, 2001). Beginning late in the second-larval stage, Dll expression becomes confined to the antenna portion (Diaz-Benjumea et al., 1994) and Ey to the eye portion of the disc (Halder et al., 1998; Quiring et al., 1994). Dpp controls the subdivision (Kenyon et al., 2003), and Der and Notch participate in this subdivision through delimiting *Dll* and *Ey* expression to the antenna and eye portions, respectively (Kumar & Moses, 2001). Embryos mutant for Btd, Ems and Otd lack Dll expression in the primordium of the antenna (Inoue et al., 2002), which indicates that these genes regulate Dll.

another possibility is that other factors present in a graded manner in the developing limbs differentially sensitize cells to homogeneous levels of Dll protein. For instance, it is possible that Dll cooperates with the graded Dpp and Wg signals to achieve differential target gene regulation along the limb (Panganiban, 2000).

# Wings

Although *Dll* is expressed early in the imaginal discs of *Drosophila* wings and halteres, its function is not required for their formation (Weihe *et al.*, 2004). Subsequently during embryogenesis *Dll* expression is lost from the wing and haltere discs (Kaphingst & Kunes, 1994), but is reactivated along the presumptive wing margin at some point before the third and final larval instar (Weigmann & Co-

hen, 1999) (Fig. 4). However, this expression is modified in halteres (Weatherbee *et al.*, 1999).  $Dll^{17}$  affects the wing margin (including hairs and bristles) and vein differentiation in the vicinity of the margin during the third larval stage (Gorfinkiel *et al.*, 1997; Campbell & Tomlinson, 1998). Ectopic expression of Dll in the third larval wing discs leads to the formation of distal leg elements. This effect is likely due, in part, to repression of the wing selector gene *vestigial* (*vg*) by ectopic Dll (Gorfinkiel *et al.*, 1997).

#### Wing eyespots and melanic spots

*Dll* is also expressed along the margin of both butterfly and moth wings (Carroll *et al.*, 1994; Kango-Singh *et al.*, 2001; Reed & Gilbert, 2004; Reed & Serfas, 2004; Monteiro et al., 2006). In derived lepidopteran lineages, Dll is additionally expressed along intervenous stripes that, in species with border eyespots, become enlarged at the proximal end and resolved into a circular pattern. This group of cells map to the center of the future border eyespots (Brakefield et al., 1996; Reed & Serfas, 2004) (Fig. 4). In saturniid moths with more centrally located discal-cell eyespots Dll is also expressed in the future eyespot centers, which later are intercepted by a cross-vein (Monteiro et al., 2006). This cross-vein expression of Dll is also seen in nymphalid butterflies that do not carry discal-cell eyespots (Reed & Gilbert, 2004). In the pupal stage of nymphalid butterflies, *Dll* expression extends from the center of the future eyespot pattern to a disc of cells that maps to one of the concentric rings of colored scales of the adult eyespots where black pigmented scales, likely containing melanin, later appear (Brakefield et al., 1996; Brunetti et al., 2001) (Fig. 4). Polymorphic markers at the Dll locus in a nymphalid butterfly were associated with the size of the eyespots, suggesting a role for this gene in the control of eyespot size (Beldade et al., 2002). This role was supported with functional studies where the over-expression of Dll during late larval development led to both larger eyespots as well as additional eyespots, and ectopic expression during the early pupal stage led to patches of black pigmentation (Monteiro et al., 2013). These experiments indicated that *Dll* is a positive regulator of eyespot size, as well as a master control gene promoting black pigmentation. The latter function of Dll is also present in *D. biarmipes* where *Dll* expression is associated with a black spot of melanization on the tip of the wing and where *Dll* down-regulation removes the spot and over-expression induces melanization across the whole wing (Arnoult et al., 2013) (Fig. 4).

#### Mouthparts

In the fly embryo, shortly after blastoderm formation (at stage 11), *Dll* is expressed in the precursors of the labrum, maxillae and labium (Cohen *et al.*, 1989; Cohen & Jurgens, 1989; Cohen, 1990), and *Dll* is required for the development of the mouthparts, for example, clypeolabrum, maxillary palps and labial palps (Cohen & Jurgens, 1989) (Fig. 4). *Dll* expression is lost from presumptive proximal cells in all these structures during either embryogenesis or the first larval instar (Panganiban, 2000).

*Dll* is also expressed in the labrum of representatives of Chelicerata, Myriapoda, Crustacea and Insecta (Panganiban *et al.*, 1995; Popadic *et al.*, 1998; Abzhanov *et al.*, 1999; Thomas & Telford, 1999; Beermann *et al.*, 2001; Schoppmeier & Damen, 2001; Urbach & Technau, 2003), and *Dll* RNAi embryos of a spider lack a labral structure (Schoppmeier & Damen, 2001) (Fig. 4). However, *Dll* appears to lack an AP axis patterning role in the mouthparts of myriapods (Prpic & Tautz, 2003). In the primitive mandibulate insect mouthparts, *Dll* is expressed and functionally required in the palps and the medial endites of maxillae and labium (Fig. 3B) (Abzhanov & Kaufman, 2000; Beermann *et al.*, 2001). No *Dll* expression was found in the mandibles of insects (Panganiban *et al.*, 1994) or other arthropods (Scholtz *et al.*, 1998) (Fig. 4). Moreover, levels of *Dll* expression can influence the length of the gnathal appendages during larval development in Hemiptera (Angelini & Kaufman, 2005).

#### Analia

*Dll* is expressed in the genital discs and is required in the formation of the analia (dorsal and ventral anal plates) (Gorfinkiel *et al.*, 1999) (Fig. 4). The lack of *Dll* function in the anal primordia transforms the anal tissue into hindgut by the extension of the *even-skipped* (*eve*) domain, whereas ectopic *Dll* represses *eve* expression and hindgut formation in the fly (Gorfinkiel *et al.*, 1999). *Dll* is required for the development of both anal plates in males but only for the dorsal anal plate in females, supporting the idea that the analia arise from two primordia (Gorfinkiel *et al.*, 1999). Fly *Dll*<sup>2</sup> somatic clones do not develop anal plates in males, or dorsal anal plates in females (Gorfinkiel *et al.*, 1999). So far there is no report for *Dll* involvement in genital and hindgut primordia development and their morphogenesis.

#### Antennae

Similar to its expression in the mouthparts, *Dll* is also expressed in the precursors of the antennae in the fly embryo at stage 11 (Cohen *et al.*, 1989; Cohen, 1990). In the late pupal stage, *Dll* is expressed in the antennae from distal segment 2 through to the tip, or arista (Figs. 3B, 4) (Panganiban & Rubenstein, 2002).

Ectopic *Dll* expression in the head-eye region leads to ectopic antennae (Gorfinkiel *et al.*, 1997), whereas clones of homozygous *Dll*<sup>-</sup> cells are incapable of contributing to any structure but the first antennal segment (Lindsley & Zimm, 1992). Hypomorphic fly *Dll* alleles result in various transformations, malformations, fusions and/or reductions of antennae (in segments 2 and 3, and arista) (Dong *et al.*, 2000, 2002; Dworkin, 2005).

*Dll* also plays an essential role in antennal identity. *Dll* and *Eyeless* (*Ey*), are initially co-expressed in the fly eyeantennal disc (Fig. 5C) (Curtiss *et al.*, 2002), but after the input of *Dpp* that appears first in eye and later in the antennal portion of the disc, *Ey* and *Dll* become restricted to either the eye or the antennal disc, respectively, and function as selector genes (Kenyon *et al.*, 2003). *Dll* can induce ectopic antennae in the eye part of the eye-antenna disc (Dong *et al.*, 2000). In addition, both *Dll* and *Homothorax* (*Hth*) function together in specifying antennae from legs. Hypomorphic alleles of either of these genes lead to antenna-to-leg transformation (Dong *et al.*, 2000), and ectopic co-expression of *Dll* and *Hth* can induce antennal differentiation in the leg, head and genital discs (Dong *et al.*, 2000).

#### Novelties in Chelicerates

Chelicerates have separately evolved the suppression of legs in the abdomen relative to insects and crustaceans (Khadjeh *et al.*, 2012), and early expression of *Dll* correlates with the development of book lungs in arachnids, endites on palps, and of spinnerets in spiders (Abzhanov *et al.*, 1999; Pechmann & Prpic, 2009; Pechmann *et al.*, 2010) (Fig. 4). Whether the spinnerets are homologous with the wings of insects or the gills of crustaceans (Damen *et al.*, 2002) or whether they are serial homologs with legs (Pechmann & Prpic, 2009) is not yet clear, but it is clear that they are novelties, since the ancestors to spiders lacked spinneret-like appendages.

#### Horns, nasus and ocularium

*Dll*, together with other appendage patterning genes, is also expressed in horn primordia of *Onthophagus* horned beetles suggesting that these novel structures may have co-opted the limb developmental network in a novel location on the head (Moczek & Nagy, 2005; Moczek *et al.*, 2006; Monteiro & Podlaha, 2009) (Fig. 4). Interestingly, Dll accumulates in distal regions of the male pupal horn, but not in corresponding regions of the female. *Dll* down-regulation in late larval instars led to shorter horns (Moczek & Rose, 2009).

*Dll* is also expressed in the primordia of the nasus, a novel horn-like frontal projection that is present in termite soldiers, and *Dll* down-regulation represses nasus growth (Toga *et al.*, 2012) (Fig. 4). *Dll* is expressed in the ocularium of harvestman and its down-regulation leads to the loss of this head protuberance beneath the eyes (Sharma *et al.*, 2013) (Fig. 4).

#### Nervous system

*Dll* is required for the formation of parts of the central and peripheral nervous systems. Dll is expressed in some brain precursor cells at stage 15 of the fly embryo, and in both the optic lobe neurons of the brain and in the glial cells of the ventral nerve cord at a late stage (Kaphingst & Kunes, 1994). The proximal ring of Dll expression in the fly leg correlates with the location of a group of campaniform sensillae in the adult, and fly embryos that are Dll null lack certain sensilla, including Keilin's organs and antennal, maxillary, labial and labral sense organs. All of these sense organs are thought to correspond to vestiges of the distal sensilla of rudimentary larval appendages (Cohen & Jurgens, 1989; Lindsley & Zimm, 1992). Various fly Dll alleles also affect the development of the mechanosensory bristles and macrochaeta over cuticle, the labial, maxillary and antennal sense organs of the head, and Keilin's organs and leg sensilla (bracts, sex combs) (Campbell & Tomlinson, 1998; Dworkin, 2005). Moreover, it has been shown that *Dll* expression along the wing margin is autonomously required for activation of the proneural gene achaete in ventral sensory bristles (Campbell & Tomlinson, 1998). Similarly, loss of Dll activity in the genital disc of the fly eliminates sensory bristles from the analia and genitalia (Gorfinkiel et al., 1999).

In basal insects such as the silverfish *Lepisma* (Mittmann & Scholtz, 2001), Dll protein accumulation correlates with sensory organs on the mouthparts and terminalia. In the mandibles of this species, specific sensory cells are the only nuclei that stain for Dll. Similarly, in the myriapod *Glomeris* embryonic expression of *Dll* appears in presumptive sensory organs of the maxilla and mandible (Prpic & Tautz, 2003). In the crustaceans *Thamnocephalus* and *Triops*, *Dll* expression appears in cells throughout the limbs at the base of bristle-like setae, which likely have a sensory function (Williams *et al.*, 2002). In the chelicerate *Limulus*, Dll protein correlates with developing mechanoreceptors and sensory neurons in the proximal legs, book gill opercula and dorsal body surface (Mittmann & Scholtz, 2001).

# Dll regulation and downstream targets

So far only a few *Dll* regulatory elements have been characterized in the fly. One of the best characterized is enhancer element 208. A 877 bp region within this element, known as Dll304, is sufficient to recapitulate the early expression pattern of *Dll* in the embryonic leg primordium, and is located approximately 12 kb upstream of the presumed *Dll* promoter region. The remainder of the 208 element directs *Dll* expression in the head (Vachon *et al.*, 1992). In addition to activation functions, the Dll304 region contains two Hox binding sites, Bx1 and Bx2, which repress *Dll* activity in the abdomen. Bx1 is 26 bp long and sufficient to fully repress *Dll*, while BX2 has only a small repressive effect (Vachon *et al.*, 1992; White *et al.*, 2000; Gebelein *et al.*, 2002).

The regulation of *Dll* expression is dynamic and tissue specific. In the fly embryo, *Dll* is activated in the thoracic imaginal primordia by Wg and repressed by Dpp dorsally and epidermal growth factor receptor (DER) ventrally (Fig. 5A) (Cohen et al., 1989; Cohen, 1990; Raz & Shilo, 1993; Goto & Hayashi, 1997). Once activated, maintenance of *Dll* expression in the embryo does not require continued Wg signaling (Cohen et al., 1993). The subdivision of the thoracic imaginal primordia is allocated in response to determinants such as Dll, Esg, Snail and Vg under the control of Wg and Dpp (Fig. 5B) (Bolinger & Boekhoff-Falk, 2005). The expression of buttonhead (btd) and Sp1 is necessary to allow expression of Dll in the leg and sufficient to induce leg identity (Estella et al., 2003). Dll is activated by Wg, expressed along the wing margin, and together these genes participate in the formation of the wing margin (Tabata & Takei, 2004). Dll is also activated by Wg signaling in the optic lobes of the brain (Kaphingst & Kunes, 1994). Expression of Dll in the embryonic head has been shown to depend partially on Wg and Engrailed (En) in the gnathal segments (Cohen, 1990), and on btd, empty spiracles and orthodenticle in the antenna (Cohen & Jurgens, 1990). The homeodomain of Ey is able to downregulate the expression of Dll, which is required during endogenous eye development (Fig. 5C) (Punzo et al., 2004). In addition, expression of Dll in the ventral maxillary segment is dependent upon Deformed (Dfd) (Ohara et al., 1993). Maintenance and refinement of several *Dll* expression patterns through the larval stages requires cooperative positive inputs from both Dpp and Wg, as well as autoregulatory inputs from Dll itself (Goto & Hayashi, 1997; Lecuit & Cohen, 1997). For example, the activation of *Dll* for analia development is dependent on the combined action of Wg and Dpp (Gorfinkiel et al., 1999). In the larval leg and antennal imaginal discs, both Wg and Dpp are required for the maintenance of Dll expression (Diaz-Benjumea et al., 1994; Lecuit & Cohen, 1997). Dpp is expressed dorsally along the anteriorposterior axis, whereas Wg is expressed ventrally along each segment's anterior-posterior compartment boundary (Blair, 1995; Held, 1995). Thus, only cells in the center of the disc are exposed to high levels of both Dpp and Wg. As the discs grow, cells at the periphery of the Dll expression domain may continue to be exposed to high Dpp or high

Wg, but not both, and therefore stop expressing *Dll*. At some point during the second larval instar, *Dll* expression becomes independent of *Dpp* and *Wg* (Lecuit & Cohen, 1997), probably due to autoregulation (Gorfinkiel *et al.*, 1997; Weigmann & Cohen, 1999). *Dll* itself is also able to induce Wg and Dpp signals, which in turn induce *Dll* expression nonautonomously (Gorfinkiel *et al.*, 1997).

Gradients of the morphogens Dpp and Wg initiate the PD organization of the fly leg by activating Dll and repressing dachshund (dac) and homothorax (hth) in the center of the disc, and by allowing the activation of dac while repressing hth medially (Lecuit & Cohen, 1997; Abu-Shaar & Mann, 1998). This creates three domains, distal (center of the disc), medial and proximal, that are specified respectively by the expression of Dll, dac and hth (Fig. 3A). Additionally, duplication of dac in spiders created a paralog, dac2, which is responsible for an additional leg segment, the patella (Turetzek et al., 2016). Mutually antagonistic interactions between the genes expressed in the proximal and medial and between medial and distal domains maintain the domain identity in the leg (Dong et al., 2001). Antennapedia (Antp) (together with *Dll*) represses distal expression of *Hth* in the leg, precluding the overlap of *Dll* and *Hth* and thereby preventing antennal differentiation (Dong et al., 2000). As in the leg, Dll and hth are required to specify the distal and proximal domains of the antenna (Dong et al., 2000). However, dac, required for the patterning of the medial leg (Mardon et al., 1994), has a different function in the antenna. These three genes extensively overlap in expression in the antenna and there is no mutual antagonism between Dll and hth (Dong et al., 2001). Instead of dac, spalt (sal) functions in the specification of the medial domain in antennae (Dong et al., 2000), while aristaless (al) and bric à brac (bab) are required for the patterning of the tarsus (Campbell & Tomlinson, 1998). Mutually repressive interactions are also required to separate domains along the PD axis of the fly wing (Dong et al., 2001), but these involve hth and vestigial (vg).

The role of *Dll* and that of other genes in the genetic network for the specification of limb axes has probably been extensively modified across arthropods. An indication of network evolution is the different requirements found for *Wg* in limb axis specification in *Drosophila*, *Tribolium*, *Oncopeltus* and *Gryllus* (Angelini & Kaufman, 2005). Obviously, more sampling of taxonomic groups and more functional studies with other members of the network will be necessary for a complete understanding of the evolutionary history of limb axis specification.

In the fly and the moth *Manduca*, *Dll* expression and limb formation are repressed in the abdomen by the Hox proteins Ultrabithorax (Ubx) and Abdominal A (AbdA)

(Fig. 5A) (Vachon et al., 1992; Zheng et al., 1999). Spiders also prevent limb formation in the abdomen, but have evolved this convergently: Antp, and both Antp and Ubx, in the first and second abdominal segments respectively, each suppressing *Dll* expression (Khadjeh et al., 2012). In insects, these bithorax-complex proteins are known to repress Dll expression by binding to a small number of specific sites in a minimal *cis*-regulatory enhancer element (Vachon et al., 1992). However, Dll repression is absent in the abdomens of species that carry appendages such as in myriapods and crustaceans (Averof & Cohen, 1997; Grenier et al., 1997), and in the first abdominal segments (A1) of the beetle Tribolium (Lewis et al., 2000) and the grasshopper Schistocerca (Lewis et al., 2000). In Tribolium, Abd-A and Ubx diverge in function relative to limb repression. Whereas Abd-A represses early expression of *Dll* in the embryonic abdomen, Ubx, expressed in A1, appears to allow A1 appendage growth (Lewis et al., 2000). Thus, the repression of Dll by one or more Hox genes was apparently acquired progressively within the arthropod lineage.

*Dll* expression in butterfly hindwings is down-regulated directly or indirectly by *Ubx*. Clones of cells lacking *Ubx* within an eyespot field lead to over-expression of *Dll* and subsequent alterations in the size of the hindwing eyespots relative to their forewing counterparts (Weatherbee *et al.*, 1999). Additionally, ectopic expression of *Ubx* on the pupal wings of *B. anynana* activates the black-scale associated genes *sal* and *Dll*, and leads to the differentiation of black wing scales (Tong *et al.*, 2014).

In butterfly larval wings, *Notch* upregulation is followed by *Dll* up-regulation in an intervenous line of cells, as well as in cells that map to the center of the eyespots (Reed & Serfas, 2004), but functional essays to test the hypothesis of direct regulation of *Dll* by *Notch* are still lacking.

*Dll* expression in both the anal and genital primordia in the third instar larvae of the fly is induced by the joint activities of caudal (cad) and the hedgehog pathways (Gorfinkiel *et al.*, 1999; Moreno & Morata, 1999).

All of Dll's putative targets in the fly embryo and/or larvae encode transcription factors. In the embryo, Dll activates *disconnected* (*disco*) and *D-Wnt5* in the thorax (Cohen *et al.*, 1991; Emerald *et al.*, 2003), *al* in the antennal and maxillary segments (Panganiban, 2000), and represses *hth* in the head and thorax (Panganiban, 2000). In the leg and/or antenna imaginal discs, Dll activates *al* (the most distal elements of the antenna and leg) (Campbell & Tomlinson, 1998), *bab* (fourth-fifth antennal segments) (Campbell & Tomlinson, 1998), *bar* (fourth through to arista antennal segments), *spineless* (*ss*) (tarsal segments, and second and arista antennal segments), *dac* (third antennal segment), *sal* and *atonal* (*ato*) (second antennal segment) (Dong et al., 2002), distal antenna (dan) and distal antenna-related (danr) (distal antennal segments) (Emerald et al., 2003), and hernandez (hern) and fernandez (fer) (third antennal segment) (Suzanne et al., 2003). Dll represses hth in the leg (Abu-Shaar & Mann, 1998; Wu & Cohen, 1999) but not in the antenna (Dong et al., 2000), and dac in the distal leg (Dong et al., 2002). Other genes regulated by Dll include BarH1/BarH2 (Kojima et al., 2000) and Notch ligand Serrate (Ser) (Rauskolb, 2001) in the leg disc. In the wing disc, Dll regulates expression of achaete (ac) (Campbell & Tomlinson, 1998) and bab (Panganiban, 2000). Dll has low DNA binding site specificity so, in order to activate specific target genes, Dll probably forms complexes with other transcription factors (Panganiban & Rubenstein, 2002). So far, from the four identified candidate cofactors, all are homeodomain proteins: two Hox proteins, Deformed (Dfd) and Antp, and two TALE proteins Exd and Hth (Panganiban, 2000). Dll cooperates with Dfd to establish ventral maxillary identity (Ohara et al., 1993), and may interact with Antp to specify leg identity (Struhl, 1981). Both hth and Dll are required to establish antennal identity (Casares & Mann, 1998), and are also needed to activate antenna-specific transcription of sal (Dong et al., 2000) and probably dac (Dong et al., 2002).

#### Molecular and functional evolution

Our Basic Local Alignment Search Tool (BLAST) analysis shows that Ceh-43 of C. elegans is the homolog of Dll and Dlx of arthopods and vertebrates, respectively. This C. elegans gene is the most similar to Dll/Dlx, having 31.3% and 27.7% identity with Dll-RA of flys and Dlx1 of humans, respectively. The presence of a single Dll-like gene in C. elegans appears to represent the more ancestral state in Metazoa. Ceh-43 is physically linked to the C. elegans Hox cluster and located on the side of the cluster corresponding to the most posteriorly expressed genes (Stock et al., 1996). However, in flies, Dll became separated from the Hox cluster via a translocation, as these genes do not map to the same chromosomes (Cohen et al., 1989). In the lineage leading to the Deuterostomes, Dll was duplicated multiple times to give rise to the six Dlx genes in mice and humans and eight Dlx genes in zebrafish. Fly and amphioxus *Dll* are most closely related to *Dlx1* (Holland et al., 1996; Stock et al., 1996), suggesting that Dlx1 may have retained most of the ancestral functions of the vertebrate *Dlx* family of genes. The very large genomic size of *Dll* in the insect genomes analyzed, due for the most part to the extremely long central intron, appears to be a derived feature of the insect lineage.

Evolution of the Dll protein sequence throughout the metazoans is mainly occurring in the C-terminus following the homeodomain motif (Fig. 2A). This part of the protein is mostly varying in length across arthropod lineages. For instance, the C-terminus length in outgroups *C. elegans* and humans, ranges from 95 aa to 65–118 aa (across the six *Dlx* copies), respectively. In the spider *Cupiennius salei*, this same region is 90 aa in length, whereas in the coleopteran, *T. castaneum*, Dll is 115 aa long, and in the more derived Holometabola, Dll is at least 125 aa long.

Using the few but representative full-length Dll sequences available from the National Center for Biotechnology Information (NCBI) we attempted to estimate the phylogeny of the represented taxa using this gene. Using only the conserved Dll protein-coding regions we observed that the clades containing Diptera, Lepidoptera, Coleoptera and Eumetabola are strongly supported monophylies with at least 87% bootstrap values (Fig. 2B). The phylogeny is congruent with the inference of insect relationships based on morphological and other molecular data (Wheeler et al., 2001), but lacks bootstrap support at some nodes. In addition, Myriapoda appears as the sister taxa of Chelicerata instead of Insecta, as expected, but this relationship does not have strong bootstrap support. This analysis suggests that *Dll* alone may be insufficient to resolve phylogenetic relationships among Arthropod lineages.

Dll is likely to have acquired its multiple functional roles in development in a gradual fashion. By analyzing the common roles of Dll and Dlx in protostome and deuterostome development it was proposed that the ancestral Dll gene may have functioned first in the developing nervous system of both invertebrates and vertebrates, acquiring roles in appendage development later in evolution, but still before the split of protostomes and deuterostomes (Panganiban & Rubenstein, 2002). According to this hypothesis, Dll would initially be involved in patterning structures of the peripheral nervous system when selection for sensory structures to protrude from the body wall to better sample the environment would have modified Dll's role into a gene that promotes outgrowth. Later, because of Dll's pre-existing association with the sensory protrusions, Dll would be a good candidate to co-opt into the PD axis patterning process of limbs. Thus, primitively Dll would have at least two main roles during development, one in the formation of peripheral sensory structures (such as setae) and another in PD axis formation (Williams et al., 2002). Within the insects, this second function appears to have been co-opted into horn development in beetles, whereas a new non-outgrowth function evolved in the patterning of moth discal-cell eyespots and butterfly border eyespots. The origin of the gap gene function of Dll is unclear because this function was discovered in a single spider species. More comparative work will be required here. Also, it is still unclear when Dll's function of regulating melanin synthesis genes evolved. It may have evolved independently in butterfly and fly lineages or in a common ancestor to both lineages. *Dll/Dlx* genes have also acquired functions in patterning other organs/tissues, including the mouthparts, the auditory and olfactory systems, the hematopoietic system, and skeleton and connective tissue systems. However, it is not yet known whether these roles were acquired independently in the protostomes and deuterostomes or whether they also predate the divergence of these animal lineages.

#### Conclusions

The study of genomic and individual sequences of *Distalless* showed that there is only one *Dll* copy in protostomes and two *Dll* splicing variants, RA and RB, in most species. The genome sizes of insect *Dll* are much larger than those of human and nematode homologues due, for the most part, to a large intron with a shared splicing site in protostomes and deuterostomes. In addition to the homeodomain, four additional motifs are identified to be relatively conserved across arthropods, NM-1, NM-2, HD-5' and HD-3. Evolution of the Dll protein sequence throughout metazoans is mainly occurring in the C-terminus following the homeodomain motif.

The data reviewed here indicate that Dll functions as a gap gene in spiders, and is required for limb (leg, antenna, mouthparts, annalia) outgrowth and for differentiation of distal limb structures. Dll participates in the differentiation of the wing margin (including hairs and bristles) and of veins in the vicinity of the margin, and is a major regulator of melanin synthesis in flies and butterflies. Dll also plays a role in eyespot development in butterflies, horn development in beetles, nasi development in termites, and ocularium development in harvestman, outgrowths that are novel traits in the respective lineages. Dll, functioning as a selector gene, also plays an essential role in antennal identity. In addition, Dll is required for the formation of parts of the central and peripheral nervous systems. Dll is activated by a variety of genes, depending on the species and the developmental context, and affects a variety of downstream targets. Its complex regulation is probably due to the gradual evolution of multiple enhancers in its cis-regulatory region. Its multiple numbers of targets, so far all transcription factors, are the result of the evolution of Dll binding sites in these genes' cis-regulatory domains. Dll acts on the target genes either as a repressor or activator of gene transcription. Target specificity appears to require the binding of Dll to additional co-factors such as the homeodomain proteins Dfd, Antp, Exd and Hth. The ancestral *Dll* may have functioned first in the developing nervous system, acquiring roles as a gap gene, and in head patterning and appendage development later in evolution.

With only a few exceptions, most of the functional work on *Dll* in arthropods has been performed on *Drosophila*. Further functional work on a variety of other arthropod lineages would be welcome to confirm the inferred ancestral functions for this gene, but mostly to determine the branches on the phylogeny where the new functions evolved. Additionally, as the regulatory code becomes better understood, and further arthropod genomes are sequenced, comparative sequence analyses alone may provide insights into *Dll*'s functional evolution both in the ontogeny of an organism and across the tree of life.

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#### Disclosure

The authors declare no conflict of interest.

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