Integration of RNA processing and expression level control modulates the function of the *Drosophila* Hox gene *Ultrabithorax* during adult development

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SUMMARY

Although most metazoan genes undergo alternative splicing, the functional relevance of the majority of alternative splicing products is still unknown. Here we explore this problem in the *Drosophila* Hox gene *Ultrabithorax* (Ubx). Ubx produces a family of six protein isoforms through alternative splicing. To investigate the functional specificity of the Ubx isoforms, we studied their role during the formation of the *Drosophila* halteres, small dorsal appendages that are essential for normal flight. Our work shows that isoform Ia, which is encoded by all Ubx exons, is more efficient than isoform IVa, which lacks the amino acids coded by two small exons, in controlling haltere development and regulating Ubx downstream targets. However, our experiments also demonstrate that the functional differences among the Ubx isoforms can be compensated for by increasing the expression levels of the less efficient form. The analysis of the DNA-binding profiles of Ubx isoforms to a natural target, shows no major differences in isoform DNA-binding activities, suggesting that alternative splicing might primarily affect the regulatory capacity of the isoforms rather than their DNA-binding patterns. Our results suggest that to obtain distinct functional outputs during normal development genes must integrate the generation of qualitative differences by alternative splicing to quantitative processes affecting isoform protein expression levels.

KEY WORDS: *Drosophila*, *Ultrabithorax*, Hox, Alternative splicing, *spalt*

INTRODUCTION

Just over three decades ago, the discovery of splicing brought a revolution to molecular biology (Gilbert, 1978). It was then realized that alternative splicing could provide a higher coding capacity to a single gene by generating distinct proteins (Black, 2003). Although it is known that at least 43% of the genes of *Drosophila melanogaster*, and about two-thirds of mouse and human genes, encode alternatively spliced mRNA sequences (Johnson et al., 2003; Stole et al., 2004; Park and Gravely, 2007; Ben-Dov et al., 2008), the functional relevance of the majority of alternative splicing events remains to be established.

In *Drosophila*, an extreme case of diversity obtained through alternative splicing is observed in the *Dscam* gene, which may potentially code for more than 38,000 splicing variants (Graveley, 2005). Alternative splicing in the *double sex* gene provides a clear example of functional divergence of protein isoforms, as it determines the specific development of males or females (Schütt and Nöthiger, 2000). Within the Hox genes, which encode homeodomain-containing transcription factors that specify structures along the anteroposterior axis of bilaterians (Pearson et al., 2005; Foronda et al., 2009) there are several examples of alternative splicing in mouse, human and *Drosophila* (Mlodzik et al., 1988; Bermingham and Scott, 1988; Celniker et al., 1989; Zavortink and Sakonju, 1989; deLorenzi and Bienz, 1990; Shen et al., 1991; Cribbs et al., 1992; Sham et al., 1992; Fujimoto et al., 1998; Patel et al., 1999). However, the relevance of alternative splicing for Hox gene function has been little explored – except, perhaps, in the *Drosophila* Hox gene *Ultrabithorax* (Ubx).

The *Drosophila* Ubx gene specifies the development of adult derivatives such as the third leg, the haltere and the first abdominal segment (Lewis, 1963; Morata and Kerridge, 1981). Wings and halteres are serially homologous, and different, thoracic dorsal appendages. *Ubx* is expressed in the haltere discs but not in the wing discs – the primordia of halteres and wings, respectively (White and Wilcox, 1984; Beachy et al., 1985). Mutations that eliminate *Ubx* expression transform halteres into wings, whereas ectopic expression of *Ubx* transforms wings into halteres (Lewis 1963; Lewis, 1978; Cabrera et al., 1985; White and Wilcox, 1985; White and Akam, 1985). *Ubx* mutations that slightly change either *Ubx* expression pattern or levels in the haltere disc lead to milder, yet easily visible, transformations of haltere, revealing a highly sensitive response to the levels of *Ubx* input. Several specific targets regulated by *Ubx* in the haltere imaginal disc have been identified (Weatherbee et al., 1998; Mohit et al., 2006; Hersh et al., 2007) (T. Pavlopoulos and M.A., unpublished). Thus, the haltere system provides an excellent framework for the qualitative and quantitative assessment of Ubx function at both developmental and molecular levels.

*Ubx* alternative splicing leads to the formation of six different *Ubx* mRNAs according to the use of alternative splice donor sites at the start of the first exon added to the optional inclusion of two 51 bp microexons located between the 5′ and 3′ common exons (Fig. 1A). The resulting Ubx proteins are named I, II or IV depending on whether they are encoded by: all the exons (I); the 5′ exon, the second microexon, and the 3′ exon (II); or just the 5′ and 3′ exons...
(IV). Each Ubx protein variant (i.e. I, II or IV) is named as ‘a’ or ‘b’ depending on: (a) the lack of, or (b) the inclusion of a small nine amino acid segment encoded between the two donor splicing sites at the 3’ end of the first exon (O’Connor et al., 1988; Kornfeld et al., 1989) (Fig. 1A). Previous studies have revealed that each isoform has distinct expression in the embryo, that class ‘b’ isoforms are much less abundant than class ‘a’ proteins, and that isoform Ia is the predominant one in most stages (O’Connor et al., 1988; Kornfeld et al., 1989; López and Hogness, 1991; Arteto et al., 1992; Bonmze and López, 1994; López et al., 1996).

The extent to which the different isoforms possess different functions remains unclear. A pioneer analysis was done with the Ubx\textsuperscript{MXT} mutation, an inversion with breakpoints to the left and right of the second microexon, and which forms only isoforms IVa and IVb (Busturia et al., 1990; Subramaniam et al., 1994). Homozygous Ubx\textsuperscript{MXT} adults present small changes in phenotype, most obviously a partial transformation of haltere to wing. This effect, however, was attributed to a reduction in Ubx protein expression in the haltere disc and not to changes in the distribution of the different Ubx proteins, thus suggesting all Ubx isoforms are functionally equivalent (Busturia et al., 1990).

Other experiments, however, showed that changes in the activities of the Ia and IVa isoforms affected the peripheral nervous system (Mann and Hogness, 1990; Subramaniam et al., 1994) or the Klin’s organs, particular sensory structures (Gebelen et al., 2002), differently. Further work on the Ubx\textsuperscript{MXT} mutants showed defects in flight and behaviour (Subramaniam et al., 1994). Besides, no reduction in expression was observed in haltere discs of the mutant larvae, suggesting functional differences between Ubx isoforms (Subramaniam et al., 1994). Regardless of this evidence, and perhaps because of the discrepancies between the different studies, most recent work has assumed that alternative splicing is irrelevant to Ubx function, and accordingly, a single Ubx isoform (i.e. Ubxla) is normally considered to be representative of the function of all Ubx proteins.

Here we re-examine the role of alternative splicing on Ubx function, focusing on haltere development. Our experimental design has allowed us to assess, with high sensitivity, the abilities of the different Ubx isoforms to control haltere development or the regulation of target genes in haltere discs, and to compare these results with those obtained in Ubx\textsuperscript{MXT} mutants. Finally, we test the DNA-binding profiles of different Ubx isoforms to one of these molecular targets, the gene spalt. We find that Ubx isoform IVa is not able to specify the normal development of halteres or to regulate Ubx target genes as efficiently as isoform Ia when expressed at comparable levels in the haltere disc. However, higher levels of isoform IVa can compensate for its lower activity and form normal halteres. The differential activity of Ia and IVa proteins does not seem to depend on major differential DNA-binding profiles. Our results indicate that alternative splicing significantly modulates Ubx function during Drosophila adult development, and also reveal that low-performance splicing isoforms can improve their function if levels of expression are increased. We suggest that developmental genes must coordinate the formation of different proteins by alternative splicing with the regulation of their levels of expression in different tissues.

**MATERIALS AND METHODS**

**Genetics**

We used the following stocks: Ubx-Gal4\textsuperscript{LDN} (de Navas et al., 2006), scalloped-Gal4 (Garaulet et al., 2008), C-765-Gal4 (Guillén et al., 1995), MS1096-Gal4 (Capdevila and Guerrero, 1994), UAS-Ubxla (Castelli-Gair et al., 1994), UAS-GFP (Drysdale et al., 2005), abx bcl\textsuperscript{pxb} and Ubx\textsuperscript{cl} (Lewis, 1982). In most experiments we have used a recombinant Ube-Gal4\textsuperscript{LDN} UAS-y\textsuperscript{+} chromosome, and grown the larvae at 17°C (except those with the UAS-1va2 line, grown at 25°C or 29°C), to reduce the amount of Gal4 protein (normally in excess) that activates each UAS construct. The sal\textsuperscript{E/Py} enhancer was described previously (Barrio and de Celis, 2004).

**Immunostaining**

Immunostaining was performed according to standard protocols (Wolff, 2000). The antibodies used were mouse and rabbit anti-β-galactosidase (Cappel), mouse anti-Ubx (White and Wilcox, 1984), rabbit and rat anti-Sal (Barrio et al., 1999), mouse anti-Wg (Brook and Cohen, 1996), rat anti-Ara (Díez del Corral et al., 1999) and rat anti-Tub (YL/2) (Serelab). The intensity of the fluorescence signal in the haltere pouch was measured with the Measure tool of the MetaMorph program (Universal Imaging Corporation) or the Measure tool of ImageJ. Larvae of the different genotypes were put in tiny baskets, and all the baskets placed together in a wide vial in which all the fixation, incubation with antibodies and staining were carried out, allowing the free movement of the liquids, so that discs of all the genotypes received the same conditions. We took three measurements in each of five different discs and measured the average fluorescence.

**RT-PCR studies**

RNA was isolated using illustra RNAspin Mini (GE Healthcare) from 50 haltere discs of either wild-type or Ubx\textsuperscript{MXT} homozygous larvae. cDNA synthesis was performed with the M-MLV reverse transcriptase (Invitrogen) using the Ubx 3A1 (5’-GCCGGTCAGATAATGGTTCTG-3’), which hybridizes to nucleotides 78 to 98 bp downstream to the 3’ splice site of the 3’ exon. The cDNA obtained was amplified by PCR using the Ubx.5S1 (5’-TGGATGCTGCAAATGACCATC-3’), which hybridizes to the Ubx 5’ exon at nucleotides −133 to −113 bp upstream to the ‘b’ 5’ splice site, and Ubx.3A1 primers. The analysis of the PCR product was made in 2% agarose gels.

**Construction of UAS-Ubx transgenic lines**

This has been previously described (Reed et al., 2010).

**Western blots**

Western blotting was performed as described (Ausubel et al., 2004). We used biotinilated anti-rat (1:10,000) and anti-mouse antibodies (1:200). Measurement of the signal was carried out with a GS-710 Calibrated Image Densitometer (BioRad) for the digitalization of the film and the Quantity One (BioRad) software for the analysis of the data.

**DNA-binding experiments**

A series of DNA elements derived from the sal cis-regulatory region were analysed by electrophoretic mobility shift assay (EMSA) (the sequence of each element is provided on request). In brief, double-stranded oligonucleotides (33-38mer) centred on Ubx binding sites sal3, sal4 and sal5/6 (Galant et al., 2002; Walsh and Carroll, 2007) bearing 5’ protruding T-overhangs (2/3x) were radioactively labelled with \(^{32}P\) by end-filling with the Klenow fragment of the DNA polymerase I (New England Biolabs). Proteins were produced by in vitro translation using the TnT system (Promega, Madison, USA) in the presence of small amounts of \(^{35}S\)-methionine (MP Biomedicals). Protein quality and size were examined in 12% PAGE gels, protein bands were quantified in a Typhoon Trio Scanner (Molecular Dynamics) overnight. Images were collected as tiff files using ImageQuant software (Amersham Biosciences).
Adult cuticle analysis
This was done as previously described (de Navas et al., 2006).

RESULTS
We analysed the abilities of the Ubx splicing isoforms (Fig. 1A) to specify haltere development by two methods: first, we combined the use of Ubx mutants and a specific Ubx-Gal4 driver line to examine Ubx functions during haltere development; second, we re-examined the phenotype and Ubx protein expression in Ubx

Rescue of the Ubx mutant phenotype by different Ubx isoforms
We first studied which Ubx isoforms are present in the wild-type haltere disc during late larval stages by RT-PCR. As shown in Fig. 1B, isoforms I and II are abundant in these discs, and a small amount of isoform IV can also be detected. This agrees with results showing that cDNAs encoding isoforms IVA and IVB are much less abundant than other Ubx cDNAs in third instar larvae (O'Connor et al., 1988; Kornfeld et al., 1989). We cannot resolve whether isoforms a and b (Ia and Ib, IIa and IIb) are equally represented, but previous data indicate that the 'a' isoforms are much more abundant than the 'b' ones (O'Connor et al., 1988; Kornfeld et al., 1989; Subramaniam et al., 1994; Bomze et al., 1994).

To carry out a functional analysis of the different isoforms in forming a haltere we used the GAL4/UAS method (Brand and Perrimon, 1993) with the Ubx-Gal4Ldn insertion. This Gal4 line is inserted in the Ubx gene, drives expression mainly in the haltere pouch and is mutant for Ubx (de Navas et al., 2006). Ubx-Gal4Ldn UAS-y+/Ubx bx+ pbx adults (we refer to this mutant combination as the Ubx mutant background, or UMB), show a transformation of the distal part of the haltere (the capitellum) into the wing (de Navas et al., 2006); haltieres are bigger and bear bristles in their margins and veins in their surface (Fig. 1E, the wild-type haltere and wing are shown in Fig. 1D and 1C, respectively). As shown in Fig. 1B, this mutant combination does not alter the isoform distribution in haltere discs. We introduced different Ubx isoforms (UAS-Ubx constructs) into this mutant background and checked whether the mutant phenotype was rescued. Because the 'b' forms are much less abundant than the 'a' ones throughout development, we concentrated on studying the 'a' variants. As shown in Fig. 1F,G, the expression of isoforms Ia and IIa in the mutant background rescues much of the mutant phenotype, reducing the size of the haltere and suppressing the development of margin bristles and veins. We found, however, that a line that expresses the IVA protein (line UAS-IVA2) only partially rescued the mutant phenotype (Fig. 1H): the size of the transformed haltere was much bigger than that of the wild type and there were marginal bristles and veins. Although the initial experiments with this line were done at 25°C, the rescue was not much better at 29°C (not shown). Similar results were observed when expressing these lines in the wing disc with scaled-Gal4, MS1096-Gal4 or C-765-Gal4 lines: the flies expressing the IVA protein variably transformed wings into haltieres, even though the crosses are made at 17°C (Fig. 1J,L; data not shown), whereas adults expressing the IVA2 line only showed mild effects in wing development, even at 25°C (Fig. 1K; data not shown). We decided to study this line further and compare it with our standard Ia line (encoding the most abundant Ubx isoform).
The UbxIVa isoform is not as efficient as the UbxIa isoform in promoting a wild-type haltere

The partial rescue of the haltere mutant phenotype obtained with the Iva2 line could be due to its expressing very low Ubx levels. To check this possibility we compared, by measuring fluorescent staining and in western blots, the levels of Ubx products in the haltere discs of the wild-type and Ubx mutant background larvae with those observed when different UAS-Ubx constructs were expressed in the same mutant background. In all the cases we used an anti-Ubx antibody that recognizes all Ubx proteins (White and Wilcox, 1984).

We examined the fluorescent staining levels of haltere pouches of the following genotypes: wild type, Ubx1+/+, Ubx-Gal4LDN UAS-y+/abx bx+/ pbx (UMB), Ubx-Gal4LDN UAS-y+/abx bx+/ pbx UAS-Ubx1a (SG1 line) (UAS-Ubx1a/+; UMB) and UAS-UbxIaVa2/++; Ubx-Gal4LDN UAS-y+/abx bx+/ pbx (UAS-UbxIVa2/+; UMB; this cross was made at 29°C) (Fig. 2A-E; the pattern driven by the Ubx-Gal4LDN line is shown in Fig. 2F). Data showing fluorescence intensities in the pouches are summarized in Table 1. We found that the average intensity in the haltere pouch of UAS-UbxIVa2/++; UMB larvae is 60.2±2.4% of the wild type, a figure not statistically significantly different (P<0.05) from that observed in Ubx1/+ larvae (66.6±2.3%); the latter percentage is higher than 50% due to Ubx negatively regulating its own transcription (Irvine et al., 1993; Garaulet et al., 2008; Crickmore et al., 2009). Measurements in western blots of haltere discs gave comparable results: 62±6.7% for the UAS-UbxIVa2/++; UMB discs and 65.6±2.3% for the Ubx+/+ genotype (Fig. 2I and Table 1), the two measurements not being statistically significantly different (P<0.05). Therefore, the total amount of Ubx protein in the haltere pouch of UAS-UbxIVa2/++; UMB or Ubx+/+ larvae did not differ much. However, the phenotype in the halteres of the corresponding adult flies was different: Ubx+/+ heterozygous flies only showed a slight enlargement of the capitellum and, occasionally, one or two bristles, whereas UAS-UbxIVa2/++; UMB adults showed a partial transformation into wing (compare Fig. 2G,H). We note that in the western blots we compared the amount of protein in the whole disc, whereas our immunofluorescence measurements were concentrated in the pouch. However, the similar results obtained with the two methods support the conclusion that isoform IVa is not as efficient as isoform Ia in making a wild-type haltere when similar levels of expression are achieved.

Two results indicate, nevertheless, that isoform IVa is not completely unable to direct haltere development. First, when we expressed this protein at higher levels in the mutant background, by using the UAS-UbxIVa1 or UAS-UbxIVa33 lines (Fig. 3B, compare with a UAS-UbxIVa2/++; UMB disc in Fig. 3A), the halteres were smaller than those of wild-type flies and slightly malformed, but showed no wing characteristics (Fig. 3D, arrow, compare with a UAS-UbxIVa2/++; UMB haltere in Fig. 3C; data not shown); second, when we introduced two UAS-UbxIaVa2 insertions in our mutant background we obtained an almost complete rescue of the mutant phenotype (Fig. 3E). Our conclusion is supported by the results obtained when we used the UAS-UbxIVa1 and UAS-UbxIVa33 constructs to express the UbxIVa protein in the wing pouch with the MS1096-Gal4 or the scalledoped-Gal4 lines at 17°C: in these cases we obtained a transformation of wings into halteres (Fig. 3F,G, compare with Fig. 1K and 1I, respectively; data not shown).

In our experiments the Ia isoform rescued the Ubx mutant phenotype but at levels higher than those of the wild type or those obtained with the Iva2 line (see Table 1). To ascertain if lower levels of this protein in the mutant background were sufficient to make a normal haltere, we reduced the amount of UbxIa protein by increasing the number of UAS ‘neutral’ constructs to titrate the Ubx protein. In UAS-GFP/UAS-GFP; UbxIa/UMB haltere

Table 1. Relative abundance of Ubx protein in the haltere disc in different mutant backgrounds when compared with the wild type (100%)

<table>
<thead>
<tr>
<th>Measurement in discs (n=15)</th>
<th>Measurement in westerns (n=3)</th>
<th>Isoform Iva in haltere disc</th>
<th>Transformation to wing</th>
</tr>
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<tbody>
<tr>
<td>Wild type</td>
<td>100%</td>
<td>100%</td>
<td>–</td>
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<tr>
<td>UMB</td>
<td>31.4±2.0%</td>
<td>35.5±0.5%</td>
<td>–</td>
</tr>
<tr>
<td>Ubx+/+</td>
<td>66.6±2.3%</td>
<td>65.6±2.3%</td>
<td>–</td>
</tr>
<tr>
<td>UMB Ia</td>
<td>151.0±3.9%</td>
<td>136.6±13.5%</td>
<td>–</td>
</tr>
<tr>
<td>UMB Iva</td>
<td>60.2±2.4%</td>
<td>62.0±6.7%</td>
<td>+</td>
</tr>
<tr>
<td>UbxMX17</td>
<td>72.0±1.9%</td>
<td>76.3±9.6%</td>
<td>+</td>
</tr>
<tr>
<td>2xUASGFP UMB Ia</td>
<td>55.2±2.4%</td>
<td>n.d.</td>
<td>–</td>
</tr>
</tbody>
</table>

*Only high levels considered.

n.d., not determined.
The gene *wingless* (*wg*) is required for the formation of the wing margin and is expressed at the dorsoventral boundary of wing discs (Phillips and Whittle, 1993; Couso et al., 1994) (Fig. 4A). By contrast, it is only present in the anterior compartment of the haltere disc dorsoventral boundary (Weatherbee et al., 1998; Shashidara et al., 1999) (Fig. 4B, arrow). The *aranca* (*ara*) gene is required for the development of vein L3 and associated sensilla campaniformia (Gómez-Skarmeta et al., 1996a). *ara* is expressed in two patches of anterior cells close to the anteroposterior boundary of the wing pouch (Fig. 4I, arrows) (Gómez-Skarmeta et al., 1996a; Gómez-Skarmeta et al., 1996b), but it is not present in the haltere pouch (Fig. 4J). Finally, the gene *spalt* (*sal*) is needed for the positioning of veins L2 and L5 (reviewed in de Celis and Barrio, 2009) and is expressed in the central domain of the wing pouch (de Celis et al., 1996; Lecuit et al., 1996; Nellen et al., 1996) (Fig. 4Q) but not in the haltere pouch (Weatherbee et al., 1998; Barrio et al., 1999) (Fig. 4R).

In *Ubx<sup>+/+</sup>* haltere discs, the *wg*, *ara* or *sal* expression patterns remain as in the wild type (Fig. 4C,K,S), but they change to the wing pattern in the UMB haltere discs (Fig. 4D,L,T). The wild-type haltere pattern for any of these three genes is restored if the constructs express the Ia (Fig. 4F,N,V) or IVa (line IVa1) (Fig. 4G,O,W) proteins are introduced in the mutant background (except that the IVa1 construct also eliminates Wg expression in the anterior pouch), but not if we introduce the Iva2 line in the same mutant background (Fig. 4E,M,U). These results indicate that: (1) *wg*, *ara* and *sal* are repressed by just one dose of *Ubx* in the haltere pouch; (2) isoform Ia is able to repress the expression of these targets in the mutant background that transform haltieres into wings; and (3) isoform IVa is able to achieve a similar repression only when expressed at high levels but not when present at comparable levels to those of *Ubx<sup>+/+</sup>* haltieres.

These results are paralleled by those obtained when expressing different isoforms in the wing disc, where no endogenous protein is present. In *sd-Gal4<sup>+/+</sup>*; UAS-UbxIVa2/+ wing discs, the expression of *Sal* was only barely changed (Fig. 4Y), whereas expression of the Ia isoform under the same driver eliminated *Sal* expression (Fig. 4Z).

**A functional analysis of the *Ubx<sup>MX17</sup>* mutation in haltere development**

The *Ubx<sup>MX17</sup>* mutation is an inversion between the second and third introns of the *Ubx* gene that prevents the correct splicing between microexons I and II so that only isoforms IVa and IVb are made in *Ubx<sup>MX17</sup>* homozygotes (Fig. 5A,B) (Busturia et al., 1990; Subramaniam et al., 1994). The distribution of the IVa (and IVb) proteins in *Ubx<sup>MX17</sup>* embryos follows that of the compound wild-type *Ubx* pattern, and not the particular one of the IV isoforms in the wild type (Busturia et al., 1990; Subramaniam et al., 1994). Similarly, the expression of the IV variants in *Ubx<sup>MX17</sup>* third leg and haltere discs (Fig. 5E,F) closely resembled the *Ubx* wild-type pattern (Fig. 5C,D), although the levels of expression were slightly reduced (Table 1).

*Ubx<sup>MX17</sup>* homozygous haltieres are partially transformed into wings (Busturia et al., 1990) (Fig. 5H, compare with the wild type in Fig. 5G) and the number of bristles in the first abdominal segment is reduced compared with that of the wild type (Fig. 5J, compare with Fig. 5I) (Busturia et al., 1990). We have also found small changes in the pattern of the metathoracic legs: in 83% of the mutant legs (*n=24*) there was a small apical bristle in the anterior compartment (Fig. 5K, arrow), similar to that present in the wild-type mesothoracic leg (Fig. 5L, arrow) but absent in the wild-type
anterior metathoracic leg (Fig. 5M, n=15). There is also a reduction in the number of big bristles present in the posterior compartment of the metathoracic basitarsus of UbxMX17 flies (1.2 bristles; Fig. 5N, arrow) compared with the wild type (2.6 bristles; Fig. 5P, arrow), indicating a partial transformation into the basitarsus of the wild-type mesothoracic leg (which bears no bristles; Fig. 5O). All these characteristics point to a partial anteriorwards transformation of the third thoracic and first abdominal segments; that is, a partial reduction of Ubx activity.

In accordance with these transformations, in UbxMX17 haltere pouches there was weak expression of wg in the posterior compartment (Fig. 4H, compare with the wild-type wing and haltere disc expression in Fig. 4A,B, respectively) and of sal in the central region of the pouch (Fig. 4X; the wild-type wing disc pattern in the mutant background, independently of whether or not there is expression of the Ia line). The expression of sal in the haltere disc, as functional Ubx-binding sites are present in Galant et al. (Galant et al., 2002) and Walsh and Carroll (Walsh and Carroll, 2007). To this end, in vitro translated UbxIa and UbxIVa proteins were resolved by PAGE, quantified (see Materials and Methods) and used in EMSA applied to the cis-regulatory sequences of Ubx target genes; (2) distinct regulatory capabilities of the Ubx proteins independently of their binding profiles; or (3) a combination of both. To investigate this issue, we measured the abilities of Ubx isoforms Ia and IVa to form DNA-protein complexes on Ubx-binding sites present in the sal regulatory region contained in the salE/Pv fragment [named sal1.1 in Galant et al. (Galant et al., 2002) and Walsh and Carroll (Walsh and Carroll, 2007)]. To this end, in vitro translated Ubx proteins were resolved by PAGE, quantified (see Materials and Methods) and used in EMSA applied to the sal3, sal4 and sal5/6 DNA elements (Galant et al., 2002) (Fig. 6A).

Our results indicate that UbxIa and UbxIVa proteins possess very similar DNA-binding abilities to sal4 and sal5/6 DNA elements (Fig. 6B), forming binary DNA-protein complexes in comparable proportions in the presence of excess DNA target. By contrast, isoform UbxIa seems to bind the sal3 element in a slightly more stable manner than UbxIVa (assayed with proteins at equal molar proportions) (Fig. 6B). Mutation of Ubx-binding sites leads to no significant binding of either Ia or IVa proteins (Fig. 6B), showing that these proteins bind specifically to these sites (see Galant et al., 2002).
DISCUSSION

Although the prevalence of alternative splicing in metazoans is indisputable, the functional relevance of most alternative splicing events is still unknown (Bienzowe, 2006; Hartmann and Valcárcel, 2009). Here we examined the impact of alternative splicing on the functions of the Drosophila Hox gene Ubx.

Specificity of Ubx isoforms, levels of Ubx protein and morphology

Previous experiments resulted in conflicting views regarding specificity of Ubx isoforms: some claim that UbxIa and UbxIVA perform different roles in some contexts (Mann and Hogness, 1990; Subramaniam et al., 1994; Gebelein et al., 2002; Reed et al., 2010) whereas in the original work describing the UbxMX17 mutation it was concluded that the activity of all Ubx isoforms was basically the same (Busturia et al., 1990). However, some of these studies did not take into account the amount of Ubx protein obtained in each experiment. Furthermore it has been recently demonstrated that relatively small differences in the amount of Ubx protein cause major changes in development (Tour et al., 2005).

Our study has re-examined this controversy, focusing on the development of the haltere, and measuring the amount of Ubx protein in different mutant combinations. We have not found a major difference in Ubx protein staining in UbxMX17 haltere (or third leg) discs, which would account for the mutant phenotype (Busturia et al., 1990). By contrast, the analysis of UbxMX17 animals, and of flies expressing different isoforms in a Ubx mutant background, suggests that isoform UbxIVA is not able to form a normal haltere at the levels that the Ubx isoform Ia does, suggesting different activity and helping to explain the UbxMX17 mutant phenotype. Although we have based some of these conclusions on experiments expressing different isoforms in a non-null Ubx mutant background, which allowed the precise comparison of Ubx protein levels in the haltere disc, we have also obtained similar results when expressing different isoforms in the wing disc, where no Ubx is present, except in the peripodial membrane.

We have also found that if a higher amount of the Iva protein is achieved (using the Iva33 or Iva1 insertions, or with two doses of the Iva2 construct) the transformation of haltere into wing in the mutant background is suppressed. Accordingly, some Ubx targets (wg, sal, ara), silenced in the wild-type haltere pouch, are derepressed when the UbxIVA isoform is expressed at low levels in the mutant background, but suppressed if the levels are high. Therefore, higher levels of expression can compensate for the functional differences of different Ubx isoforms caused by alternative splicing. During normal development, genes must integrate the generation of qualitative differences by alternative splicing with the regulation of the levels of expression of each isoform; the molecular coupling of transcriptional regulation to alternative splicing via modulations in transcriptional elongation rate may provide a mechanism to achieve this molecular coordination during gene expression (Cramer et al., 1997; de la Mata et al., 2003).

Binding efficiency and activity of Ubx proteins

Most Hox proteins include a hexapeptide motif that is required to interact with the extradenticle protein, the main Hox protein cofactor in Drosophila (Mann and Chan, 1996). The Ia and Iva proteins differ in the length of a region located between the hexapeptide motif and the homeodomain (the linker region, longer in the Ia variant). It has been shown that this linker region is necessary for Ubx repression of embryonic Dil but not for the binding of Ubx to the relevant Dil binding region, and that the UbxIa and UbxIVA proteins have different transcriptional regulatory properties (Gebelein et al., 2002). Similarly, we have shown that the Ia and Iva proteins do not show major differences in binding in vitro to a relevant region of the sal regulatory domain (Galant et al., 2002) but do not repress the sal gene with equal
efficiency. The repression of Dll by Ubx, however, requires the co-factors Extradenticle (Exd) and Homothorax (Hth) (Gebelein et al., 2002). By contrast, Exd is cytoplasmic and inactive in the haltere pouch (González-Crespo and Morata, 1995; Aspland and White, 1997) and Hth is not expressed and probably not required in the haltere pouch (as it is not in the wing pouch) (Pai et al., 1998; Casares and Mann, 2000; Azpiazu and Morata, 2000). Therefore, a different interaction with Exd or Hth cannot account for the different activity of the two splicing variants in the distal haltere disc. It may be that other, yet undiscovered, co-factors may differentially interact with Ubx Ia and IVa proteins in the haltere disc. Alternatively, the different activity may rely on slight differences in binding affinity, or in the transcription-regulating properties of the bound proteins. We have found slight differences between the two isoforms of Ubx in binding to one element containing three Ubx core consensus binding sites within the sal regulatory domain, but no differences in Ubx binding to other sites within this same region. The Ubx-binding sites within the sal regulatory domain contribute additively to the repression of sal by Ubx (Galant et al., 2002). It is possible, therefore, that even small differences in the binding of Ubx to each of these sites may accumulate and result in appreciable differences in gene expression. We speculate that higher amount of isoform IV may compensate for its lower binding or activity by increasing the probability of physical interactions with the basal transcription machinery. Perhaps the functional differences observed among Ubx isoforms may explain, to some degree, the evolutionary conservation in the spatial distribution of Ubx isoforms within distantly related Drosophila species.

**Regulation of Ubx targets by different Ubx isoforms**

Based on our data and on previous reports we think that there may be three different responses when either the Ia or IVa proteins are expressed: (1) some Ubx targets, such as decapentaplegic in the visceral mesoderm (at least as to its anterior repression) and those needed to specify the embryonic cuticle, probably respond similarly to the different Ubx proteins (Busturia et al., 1990; Mann and Hogness, 1990; Subramaniam et al., 1994; Gebelein et al., 2002; Reed et al., 2010); (2) other targets, such as sal, wg and ara in the haltere disc, as well as others needed to repress wing development and promote haltere development, are more efficiently regulated by isoform Ia than IVa, but if the levels of the latter are increased, a similar regulation is achieved; and (3) the expression of dpp in the posterior visceral mesoderm (Reed et al., 2010) or of targets needed to specify the segmental pattern in the embryonic peripheral nervous system (Mann and Hogness, 1990; Subramaniam et al., 1994), are differently controlled by proteins Ia and IVa, independently of their levels of expression.

Other Ubx targets can be assigned to one of these three classes with less certainty. Thus, Dll is repressed in the embryo by the Ia protein, and not by the IVa isoform (Gebelein et al., 2002), and may be included in the second group. Although Dll expression in Ubx<sup>MXT</sup> animals, or in embryos with high levels of the IVa isoform, has not been detailed, high levels of this protein repress the formation of Keilin’s organs, which is driven by Dll expression (Mann and Hogness, 1990). We note that Ubx<sup>MXT</sup> adults have abnormalities in metathoracic legs and the A1 (Busturia et al., 1990; and this report), in the development of A1 and A2 larval abdominal muscles (Reed et al., 2010) and are not as healthy as wild-type flies (Subramaniam et al., 1994), suggesting that isoform IVa may be less efficient than isoform Ia in controlling many Ubx targets, not just those making a haltere. In conclusion, we think that the particular architecture of cis-regulatory regions in each Ubx target may account for the inclusion of each gene in one of these three categories, so a specific research may be needed for each case.

Our studies also point to the importance of measuring protein levels when trying to correlate isoform activity and morphology. Several studies in different systems have also shown that the different levels of Ubx (Tour et al., 2005) or other Hox proteins may also end up in distinct phenotypes (Cribbs et al., 1995; Greer et al., 2000; Trvdik and Capecchi, 2006). In sum, we conclude that alternative splicing modulates Ubx function during *Drosophila* adult development, providing a good example of the functional relevance of alternative splicing events and protein level control within the physiological context of animal development.

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Supplementary material

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