

A feed-forward relay integrates the regulatory activities of Bicoid and Orthodenticle via sequential binding to suboptimal sites

Rhea R. Datta,¹ Jia Ling,¹ Jesse Kurland,^{2,3} Xiaotong Ren,¹ Zhe Xu,¹ Gozde Yucel,¹ Jackie Moore,¹ Leila Shokri,^{2,3} Isabel Baker,¹ Timothy Bishop,¹ Paolo Struffi,¹ Rimma Levina,¹ Martha L. Bulyk,^{2,3} Robert J. Johnston Jr.,⁴ and Stephen Small¹

¹Center for Developmental Genetics, Department of Biology, New York University, New York, New York 10003, USA; ²Division of Genetics, Department of Medicine, Brigham and Women's Hospital, Harvard Medical School, Boston, Massachusetts 02115, USA; ³Department of Pathology, Brigham and Women's Hospital, Harvard Medical School, Boston, Massachusetts 02115, USA; ⁴Department of Biology, Johns Hopkins University, Baltimore, Maryland 21218, USA

The K50 (lysine at amino acid position 50) homeodomain (HD) protein Orthodenticle (Otd) is critical for anterior patterning and brain and eye development in most metazoans. In *Drosophila melanogaster*, another K50HD protein, Bicoid (Bcd), has evolved to replace Otd's ancestral function in embryo patterning. Bcd is distributed as a long-range maternal gradient and activates transcription of a large number of target genes, including *otd*. Otd and Bcd bind similar DNA sequences *in vitro*, but how their transcriptional activities are integrated to pattern anterior regions of the embryo is unknown. Here we define three major classes of enhancers that are differentially sensitive to binding and transcriptional activation by Bcd and Otd. Class 1 enhancers are initially activated by Bcd, and activation is transferred to Otd via a feed-forward relay (FFR) that involves sequential binding of the two proteins to the same DNA motif. Class 2 enhancers are activated by Bcd and maintained by an Otd-independent mechanism. Class 3 enhancers are never bound by Bcd, but Otd binds and activates them in a second wave of zygotic transcription. The specific activities of enhancers in each class are mediated by DNA motif variants preferentially bound by Bcd or Otd and the presence or absence of sites for cofactors that interact with these proteins. Our results define specific patterning roles for Bcd and Otd and provide mechanisms for coordinating the precise timing of gene expression patterns during embryonic development.

[*Keywords:* Bicoid; Orthodenticle; enhancer; embryo; transcription factor binding; suboptimal binding]

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Animal body plans are established in large part by transcriptional networks that provide positional information during embryogenesis (Peter and Davidson 2015). The evolution of body plans is driven by alterations to these networks, including changes to *cis*-regulatory elements and neofunctionalization of transcription factors (TFs) after gene duplication (Thornton et al. 2003; Carroll 2008; Peter and Davidson 2011; Tautz and Domazet-Lošo 2011; Abascal et al. 2013).

Network interactions involve direct binding of TFs to binding sites in enhancers of target genes, and binding events are integrated into spatial and temporal patterns of expression along the major axes during development.

Understanding the mechanisms controlling the dynamics of gene expression remains a challenge, in part because TFs exist in families of closely related proteins and bind very similar sequence motifs *in vitro* (Noyes et al. 2008b). While possible mechanisms by which TF specificity is achieved (TF-preferred binding sequences, cofactor binding, etc.) have been proposed (Slattery et al. 2011), it is unclear how binding events are coordinated *in vivo* so that each protein can regulate specific gene targets.

The early *Drosophila melanogaster* embryo provides a unique system to study the spatiotemporal complexity of

Corresponding author: sjs1@nyu.edu

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TF–DNA interactions. *Drosophila* and other cyclorrhaphan Diptera develop via a “long germ” mode, in which all segments of the body plan are specified and positioned during the syncytial blastoderm stage (Lynch and Desplan 2003; Liu and Kaufman 2005; Peel 2008). In fruit flies, the maternal TF Bicoid (Bcd) sets up the first positional instructions for anterior patterning (Driever et al. 1990). *bcd* mRNA is sequestered at the anterior pole of the mature oocyte via sequences in its 3′ untranslated region (UTR) (Berleth et al. 1988). After fertilization, translation and diffusion establish an anterior gradient of Bcd protein (Fig. 1A,B; Driever and Nusslein-Volhard 1988; Little et al. 2011). Bcd contains a homeodomain (HD) and is a transcriptional activator of >50 target genes (Driever and Nusslein-Volhard 1989; Struhl et al. 1989; Finkelstein and Perrimon 1990; Stanojevic et al. 1991; Chen et al. 2012), which form a network of downstream regulatory interactions that precisely patterns the embryo. Embryos lacking Bcd fail to form any anterior structures, including all cephalic and thoracic segments, and show duplications of posterior structures (e.g., Filzkoerper [Fk]) in anterior regions (Frohnhofer and Nusslein-Volhard 1986). While the majority of its target genes is expressed during and after cellularization, the Bcd protein gradient is active only during the syncytial blastoderm stage, prior to cellularization.

Despite its critical functions in *Drosophila*, Bcd is not well conserved even within insects. Rather, Bcd arose after a recent gene duplication event and rapidly evolved to play an important role in embryonic patterning (Stauber et al. 1999; Casillas et al. 2006). In three species lacking Bcd (*Tribolium castaneum*, *Nasonia vitripennis*, and *Acyrtosiphon pisum*), at least some of the anterior

patterning functions of Bcd are performed by Orthodenticle (Otd) (Schroder 2003; Lynch et al. 2006; Huang et al. 2010). *otd* is maternally expressed in these species, and its disruption causes severe *bcd*-like defects in anterior patterning. In *Drosophila*, *otd* has evolved to become a zygotic target gene of Bcd-dependent activation (Finkelstein and Perrimon 1990). Loss of *Drosophila otd* causes embryonic lethality, but *otd* mutants show cephalic defects that are much less dramatic than the complete loss of anterior structures observed in *bcd* mutants (Finkelstein and Perrimon 1990; Finkelstein et al. 1990). Later in development, Otd is critical for central nervous system and eye development (Finkelstein et al. 1990)—roles that are conserved in vertebrates, including humans (Finkelstein and Boncinelli 1994).

Both Otd and Bcd contain a lysine at amino acid position 50 (K50) of their respective HDs and bind in vitro to the consensus sequence TAATCC (Treisman et al. 1989; Noyes et al. 2008a). The K50 residue and preference for TAATCC are conserved among all Otd homologs (Finkelstein and Boncinelli 1994) and thus are ancient, but the ancestral protein that gave rise to Bcd was a Hox3 protein with a glutamine (Q) at HD position 50 (Q50) (Stauber et al. 1999). This suggests that an important step in Bcd’s evolution was the conversion of Q50 in the ancestor to K50, which changed its DNA-binding preference and allowed it to usurp some of the anterior patterning roles played by Otd in ancestral insects. All known Bcd target gene enhancers contain multiple copies of TAATCC (Chen et al. 2012). We hypothesize that Otd regulated many of these enhancers in the ancestral species that gave rise to *Drosophila* and regulates a similar battery of

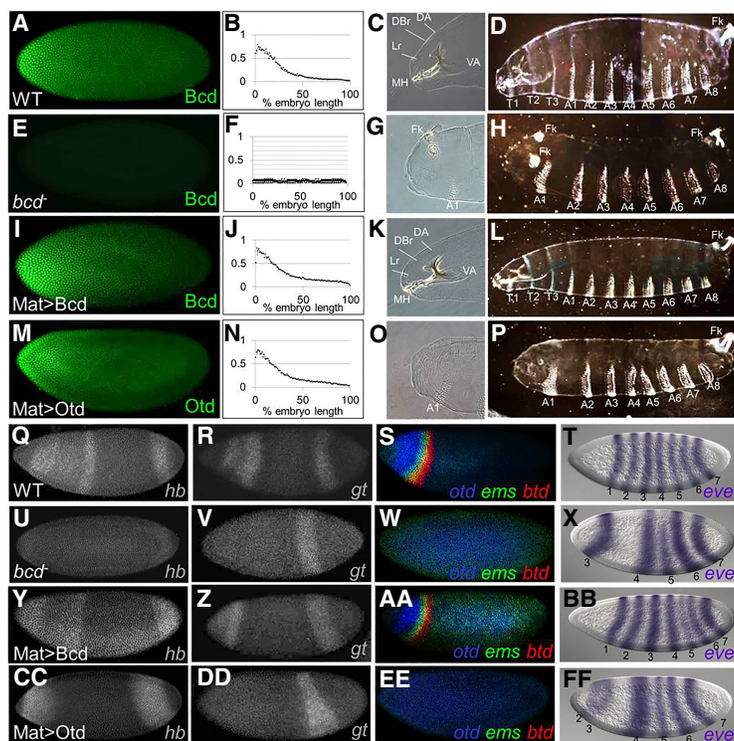


Figure 1. A maternal *Otd* gradient (Mat > *Otd*) cannot replace most *Bcd*-like functions. (A–P) Protein expression patterns (A,E,I,M), gradient quantifications (averaged from five embryos; B,F,J,N), anterior cuticle structures (C,G,K,O), and whole larva cuticles (D,H,L,P) are shown for wild type (A–D), *bcd* mutants (E–H), *bcd* mutants containing the Mat > *Bcd* transgene (I–L), and *bcd* mutants containing the Mat > *Otd* transgene (M–P). Labeled structures in anterior regions (C,G,K,O) include the dorsal arm (DA), dorsal bridge (DBr), labrum (Lr), mouthhooks (MH), ventral arm (VA), Filzkoerper (Fk), and first abdominal segment (A1). (D,H,L,P) Thoracic (T1–T3) and abdominal (A1–A8) segments are labeled. (Q–FF) mRNA expression patterns for six *Bcd* target genes [*hunchback* [*hb*], *giant* [*gt*], *otd*, *empty spiracles* [*ems*], *btd*, and *even-skipped* [*eve*]] in wild type (Q–T), *bcd* mutants (U–X), *bcd* mutants containing the Mat > *Bcd* transgene (Y–BB), and *bcd* mutants containing the Mat > *Otd* transgene (CC–FF). Assayed target genes are labeled in the bottom right corner of each panel. (T,X,BB,FF) Numbers correspond to *eve* stripes. All embryos in this study are oriented with anterior to the left.

enhancers in extant insect species that do not contain Bcd. If this is the case, perhaps Otd can replace many of Bcd's functions in *Drosophila* if it is maternally expressed and distributed in an anterior gradient.

We tested whether Otd can provide Bcd-like patterning functions through a transgenic gene replacement assay. We present here a comprehensive comparison of the *in vitro* binding preferences and *in vivo* binding profiles of Bcd and Otd. We also demonstrate that the two proteins bind sequentially to enhancers in feed-forward relays (FFRs) in which Bcd-binding initiates target gene activation, and Otd-binding maintains expression after the maternal Bcd gradient decays. Each protein also binds independently to distinct enhancers. We present evidence that Otd- and Bcd-specific binding activities *in vivo* are controlled by two mechanisms: subtle differences in binding preference and protein-specific interactions with cofactors. Our results define specific roles for Bcd and Otd in embryonic patterning in *Drosophila* and shed light on the molecular mechanisms that alter regulatory networks during evolution.

Results

A maternal gradient of Otd cannot replace Bcd in Drosophila

To assess the functional similarity between Bcd and Otd, we tested whether Otd could mediate Bcd-like activities using a gene replacement assay (Fig. 1). Coding regions for both Bcd and Otd were inserted into transgenes containing the *bcd* promoter (for maternal expression) and the *bcd* 3' UTR (for anterior mRNA localization) and were integrated into the same genomic position (Bateman et al. 2006). These constructs (designated Mat > Bcd and Mat > Otd) were crossed into *bcd*^{E1}-null mutant females, and embryos laid by those females were assayed for RNA and protein expression. mRNAs from the transgenes were maternally expressed and localized to the anterior pole region (Supplemental Fig. 1A). Antibody stains showed very similar gradients of Bcd and Otd in early embryos containing the transgenes, and the Bcd gradient from the rescue transgene was indistinguishable from the endogenous (wild-type) gradient (Fig. 1A,B,I,J,M,N).

Wild-type embryos develop cephalic structures and three thoracic segments in anterior regions during embryogenesis (Fig. 1C,D). All of these structures are missing in embryos laid by *bcd* mutant females, and posterior structures (Fk) are duplicated near the anterior pole (Fig. 1G,H). At the molecular level, *bcd* mutants fail to activate all Bcd target genes, including *hunchback* (*hb*), *giant* (*gt*), *otd*, *buttonhead*, (*btd*), *empty spiracles* (*ems*), and *even-skipped* (*eve*) stripes 1 and 2 (*eve-1* and *eve-2*) (Fig. 1U–X, cf. with Q–T), and fail to repress translation of *Cad* in anterior regions (Supplemental Fig. 1B).

As expected, when the control construct (Mat > Bcd) was crossed into embryos lacking Bcd, it completely rescued all morphological defects (Fig. 1K,L), and 95% of the embryos developed into fertile adults. This construct also activated the expression patterns of all six tested Bcd target

genes (Fig. 1Y–BB). In contrast, when the Mat > Otd construct was crossed into *bcd* mutants, none of the morphological structures missing in *bcd* mutants structures was rescued (Fig. 1O,P). However, we did detect the suppression of ectopic posterior Fk (Fig. 1O, cf. with G). Mat > Otd activated only two Bcd target genes (*hb* and *eve-2*) but failed to activate four others (*gt*, *otd*, *btd*, and *ems*) (Fig. 1CC–FF) and failed to translationally repress *caudal* (Supplemental Fig. 1B). We also tested whether Mat > Bcd and Mat > Otd could activate 24 other Bcd-dependent reporter genes (Chen et al. 2012). The Mat > Bcd construct activated expression of all 24 of these reporters, while Mat > Otd activated only two (Supplemental Fig. 2). Taken together, these results show that Otd cannot rescue most Bcd functions when provided maternally despite the fact that the two proteins bind very similar DNA sequences *in vitro*.

HDs mediate distinct in vivo activities of Otd and Bcd

The inability of Mat > Otd to activate most Bcd target genes or repress *Cad* is consistent with its failure to rescue anterior segments in embryos lacking Bcd. This result is not surprising in view of the fact that the evolved Bcd- and Otd-coding regions show very little sequence conservation (38% sequence identity within their HDs) (Fig. 2A) and no detectable homology outside the HD. To map regions of Bcd and Otd that mediate their distinct functions, we generated rescue constructs with chimeric proteins in which the DNA-binding HDs were precisely swapped (Mat > Bcd:OtdHD and Mat > Otd:BcdHD) (Fig. 2B,C,F,G). If the structural differences preventing Otd from rescuing *bcd* mutants lie inside its HD, inserting the Bcd HD into the Otd protein (Mat > Otd:BcdHD) should cause a strong rescue of the phenotype. If those differences lie outside the HD, inserting the Otd HD into the Bcd protein (Mat > Bcd:OtdHD) should result in a strong rescue.

Inserting the Bcd HD into the Otd protein (Mat > Otd:BcdHD) caused a dramatic rescue of anterior structures (Fig. 2D,E). Although rescue was incomplete (embryos died before hatching), all embryos containing the Mat > Otd:BcdHD formed three thoracic segments and at least some identifiable cephalic structures (Fig. 2D,E). Mat > Otd:BcdHD also activated all six tested Bcd target genes (Fig. 2J–M) and repressed *Cad* translation (Supplemental Fig. 1B). In contrast, the reciprocal swap (Mat > Bcd:OtdHD) caused very little rescue of the morphological defects of *bcd* mutants (Fig. 2H,I) and activated only *hb* and *eve2*, similar to rescue by Mat > Otd (Fig. 2N–Q, cf. with 1EE–HH). Together, these results indicate that important functional differences between the two proteins lie within their HDs, and regions outside their HDs are largely interchangeable.

In vivo and in vitro binding activities of Bcd and Otd

The result that the Bcd and Otd HDs are not interchangeable seems in conflict with the observation that Bcd and Otd bind the same TAATCC consensus sequence *in vitro* (Noyes et al. 2008b). However, it is possible that

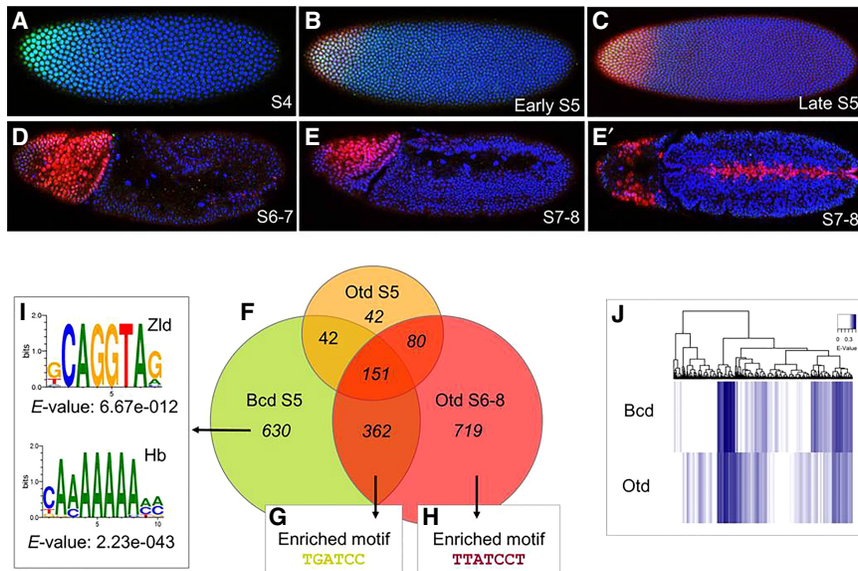


Figure 3. (A–E') A temporal comparison of Bcd and Otd expression. All embryos are stained for the Bcd protein (green), the Otd protein (red), and DAPI (blue), which marks individual nuclei (blue). The embryos shown represent a temporal series from A (youngest) to E (oldest). The ages of embryos are labeled by stage (S). (F–I) Genome-wide binding activities of Bcd and Otd. (F) A Venn diagram showing the number Bcd and Otd peaks in euchromatin (false discovery rate of <5%) in wild type embryos at the S5 and S6–S8 time points. (G–I) Motifs enriched in each data set are indicated. (J) Heat map comparisons of 9-mers bound by Bcd and Otd hierarchical clustering analysis of *E*-scores. All 9-mers bound by Bcd and Otd with an *E*-value of >0 are represented. Every position on the heat map is a single 9-mer.

early [S5], and L indicates bound late [S6–S8]) and use a consistent nomenclature here.

A FFR regulates enhancers bound by Bcd and Otd

To test the FFR hypothesis, we examined the activities of eight candidate relay enhancers that were bound by both Bcd and Otd, expressed in anterior regions, and continuously active during the period between S5 and S6–S8 (Fig. 4C–R; Supplemental Fig. 3A). In a previous study, we showed that removing Bcd function (which also removes Otd) completely abolished expression driven by all of these enhancers (Chen et al. 2012). If Otd is involved in maintaining the expression patterns driven by these enhancers, removing its function should cause a loss or reduction of expression at S6–S8. To ablate Otd function, we used CRISPR–Cas9 mutagenesis to delete the Otd HD in the endogenous locus (Materials and Methods). This mutation caused embryonic lethality but did not affect the shape of the Bcd gradient (Supplemental Fig. 4).

Expression patterns driven by the eight candidate relay enhancers were assayed in *otd* mutant embryos at three time points: early S5 (high Bcd and low Otd), late S5 (high Bcd and high Otd), and S6–S8 (no Bcd and high Otd). For six of the eight tested enhancers, removing Otd caused a substantial reduction in expression at late S5 and/or a complete loss of expression at S6–S8 but had little effect on early activation (Fig. 4C–H, K–P; Supplemental Fig. 3). One of these enhancers was the EHE enhancer from the *otd* gene itself (BcdOtd^{EL6}) (Supplemental Fig. 3), and the strong reduction of expression driven by that enhancer (Fig. 4M, N) suggests that maintenance of the normal *otd* pattern is mediated by a positive autoregulatory loop. The other two reporter genes (BcdOtd^{EL4} and BcdOtd^{EL8}) tested in *otd* mutants showed strong reductions even in early S5 (Fig. 4I, J, Q, R; Supplemental Fig. 3), suggesting that Otd is required for the initial activation of some Bcd target genes. We define these as “relay

enhancers”; they are bound and activated by Bcd prior to S5 and are then maintained by Otd in later development. Taken together, these results and the large number of shared peaks in the ChIP-seq experiments suggest that the FFR between Bcd and Otd is an important mechanism for extending the timing of expression of a subset of target genes in anterior regions of the embryo. In this relay mechanism, Otd’s primary role is the maintenance of gene expression.

The Bcd–Otd FFR involves sequential binding to a suboptimal site

One possibility is that FFR-regulated enhancers contain specific sequence motifs that facilitate sequential binding of Bcd and Otd. Using a discriminative motif search (see the Materials and Methods), we identified a single base variant of the K50 consensus, TGATCC, which is enriched in peaks bound by both proteins compared with peaks bound by Bcd or Otd alone (Fig. 3G). This site does not contain the TAAT core recognition site preferred by most HD-containing TFs (Treisman et al. 1989; Noyes et al. 2008b). This TGATCC motif appears in 62% of the overlapping peaks compared with 23% of Bcd S5-only peaks ($P=0$) and 38% of Otd S6–S8-only peaks ($P=9.05 \times 10^{-8}$). We also searched the data from our PBM experiments to see how this single base change affects the binding preferences of Bcd and Otd (Fig. 5). This search showed that TGATCC-containing 8-mers are suboptimal (less strongly bound by both proteins) compared with the TAATCC consensus (Fig. 5A). It also appears that Bcd prefers TGATCC-containing 8-mers more strongly than does Otd (Fig. 5B).

To test the *in vivo* function of the TGATCC suboptimal site, we mutated it in two different enhancers (BcdOtd^{EL4} and BcdOtd^{EL25}). Both are relay enhancers because they (1) are activated at both S5 and S6–S8 (Fig. 5C, E), (2) are bound by both Bcd and Otd, (3) lose expression in *bcd*

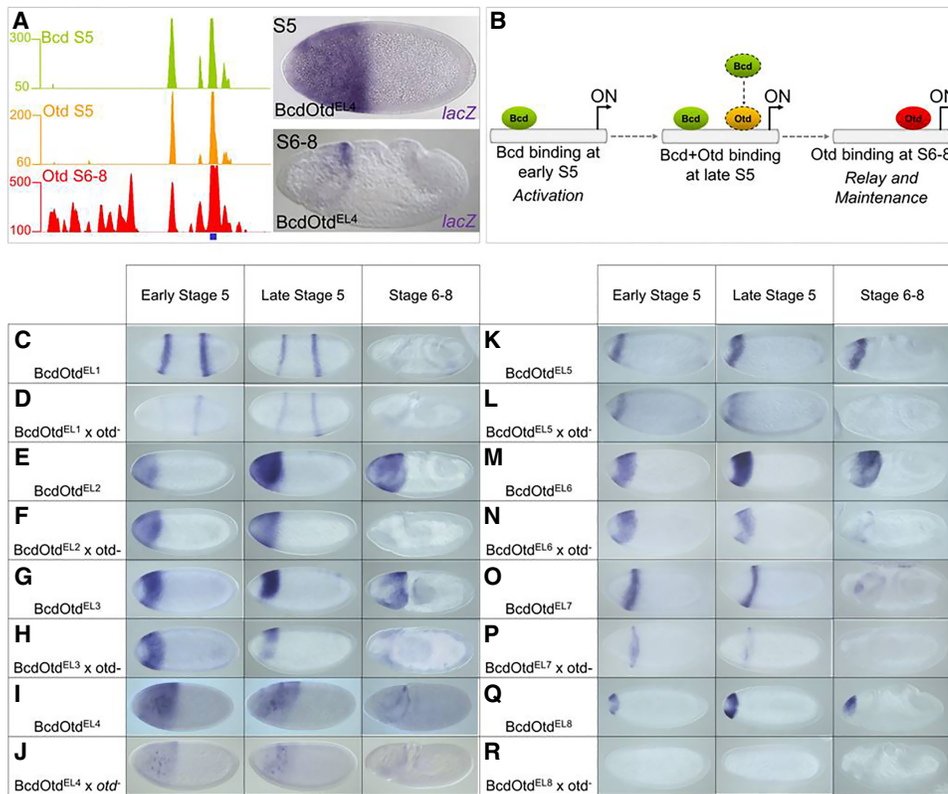


Figure 4. Temporal regulation of FFR enhancer activity by Bcd and Otd. (A) ChIP-seq peaks for Bcd and Otd around the *hb* locus. The position of a known Bcd-dependent enhancer is shown as a blue box. This enhancer drives expression early at S5 and later at S6–S8. (B) A model for a FFR coordinated by Bcd and Otd. (C–T) Testing FFR enhancers in *otd* mutant embryos. Reporter gene expression patterns in wild-type embryos (C, E, G, I, K, M, O, Q) and *otd* mutant embryos (D, F, H, J, L, N, P, R). Embryos are shown at three time points, as indicated at the top.

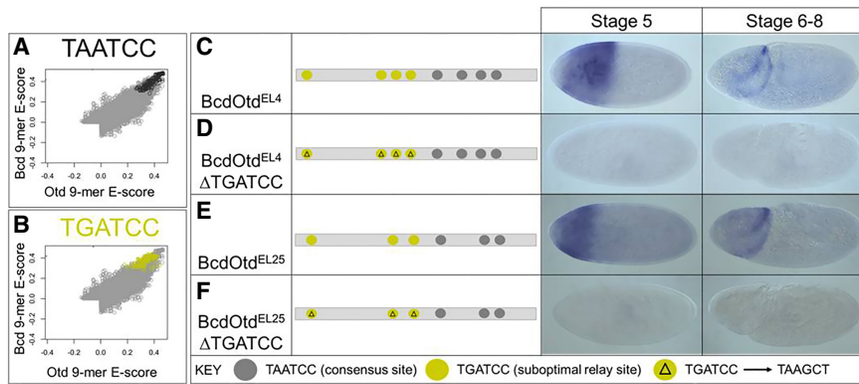
mutants, and (4) lose expression late in *otd* mutants (Fig. 4; Chen et al. 2012). The *BcdOtd^{EL4}* and *BcdOtd^{EL25}* enhancers contain three copies each of the TGATCC site and four and three copies, respectively, of the TAATCC consensus site. We converted the TGATCC sites in these enhancers to TAAGCT—a lower-affinity site for both proteins that is equally represented in all ChIP data sets. For both enhancers, these mutations caused a complete loss of expression (Fig. 5D, F), suggesting that this motif is important for both Bcd-mediated activation and Otd-dependent maintenance of expression and that the TGATCC-binding site is critical for the Bcd/Otd relay.

Anterior patterning by the integration of three classes of enhancers

In addition to the relay enhancers identified by overlapping peaks of Bcd and Otd binding, our ChIP-seq experiments yielded large numbers of unique binding peaks for each protein (Fig. 3). The 630 peaks unique to Bcd included 13 previously known Bcd-dependent enhancers (Chen et al. 2012). The expression patterns of eight of these enhancers perdured after the Bcd gradient decays (Supplemental Fig. 5) but were not bound by Otd, so we

hypothesize that other factors must be involved in the maintenance of their patterns (see the Discussion).

The ChIP experiments also identified many peaks bound only by Otd (122 peaks at S5 and 799 at S6–S8); 80 peaks were detected at both time points (Fig. 3F). We used *lacZ* reporter genes to test 19 of these peaks for enhancer activity. In our first experiments, we tested nine of the 719 fragments bound specifically by Otd only at the late time point (*Otd^L*) (Fig. 6B; Supplemental Fig. 5). As expected, all nine enhancers showed anterior expression patterns in S6–S8 embryos (Fig. 6B; Supplemental Fig. 5). We also tested 10 genomic fragments bound uniquely by Otd only at S5 (*Otd^E*; four fragments) or at both time points (*Otd^{EL}*; six fragments). Surprisingly, none of the 10 tested fragments directed any reporter gene expression at either stage of development (Fig. 6A; Supplemental Fig. 5). These results are consistent with the failure of Otd to rescue *bcd* mutant embryos (Fig. 1) and suggest that Otd binding to euchromatic regions at S5 does not lead to enhancer activation. The reason for Otd's failure to activate expression at S5 is not clear, but one possibility is that activation of anterior genes in the early embryo requires prior binding by Bcd. Alternatively, because activation by Bcd requires interactions with cofactors (Simpson-Brose et al. 1994; Crauk and Dostatni 2005;



Xu et al. 2014), Otd's inability to activate may be caused by a failure to make critical protein–protein contacts.

Taken together, our results define three classes of enhancers that mediate Bcd and/or Otd functions in vivo. Class 1 enhancers relay transcriptional initiation by Bcd to transcriptional maintenance by Otd, Class 2 enhancers are activated by Bcd and are maintained by an unknown factor. Class 3 enhancers are activated at a later time point by an Otd-dependent mechanism that is completely independent of Bcd.

Otd-dependent activation is mediated by another suboptimal site

We next searched for overrepresented motifs within the set of peaks bound specifically by Otd at S6–S8. This search identified an enriched K50 variant motif in Otd, TTATCCT, an extended variant of the canonical

TAATCC motif optimally preferred by Bcd and Otd (Fig. 3H). It appears in 26% of Otd S6–S8 peaks, 10% of Bcd + Otd peaks, and only 9% of Bcd S5 peaks ($P = 1.89 \times 10^{-15}$). An analysis of our PBM data showed that this suboptimal site is preferentially bound by Otd compared with Bcd (Fig. 6C), consistent with its overrepresentation in Otd-bound genomic fragments.

We tested the role of the TTATCCT motif in an enhancer (Otd^{L6}) that is bound by Otd and transcriptionally active only at S6–S8 in wild-type embryos and inactive in *otd* mutants (Fig. 6D; Supplemental Fig. 5). The Otd^{L6} enhancer contains two exact copies of the TTATCCT motif (Fig. 6D), one copy of the canonical K50HD-binding motif (TAATCC), and two copies of the TGATCC motif that is overrepresented in relay enhancers. We mutated the TTATCCT motifs, leaving the canonical and relay motifs intact, which caused a complete loss of expression (Fig. 6E). Based on our PBM data, ChIP data enrichment, and

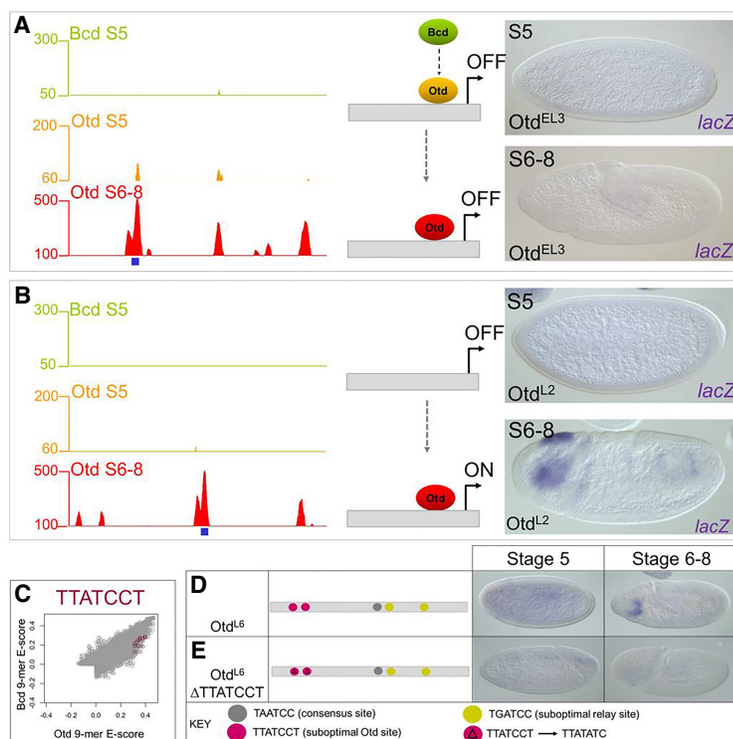


Figure 6. Reporter gene tests of genomic fragments bound only by Otd. (A) Otd^{EL3} is bound by Otd at S5 and S6–S8, but not by Bcd, and is inactive. (B) The Otd^{L2} enhancer is bound by Otd at S6–S8 and is active in the head. (C) Otd-preferred sequence TTATCCT that is enriched in the Otd S6–S8 ChIP data is preferred in vitro by Otd. (D,E) Mutating the Otd suboptimal site TTATCCT abolishes enhancer activity in a late acting Otd enhancer.

the enhancer site change data, we conclude that these suboptimal sites are necessary for activation by Otd at S6–S8.

Timing of enhancer activity is controlled by suboptimal motif preferences and cofactor interactions

Our motif searches identified suboptimal binding sites required for the activities of FFR enhancers (Fig. 5) and Otd^{L6} enhancers (Fig. 6), respectively. These results suggest that subtle binding preferences between the two proteins contribute to correctly timing enhancer activation in vivo. Alternatively, it is possible that the differential timing of enhancer activity is controlled by the presence or absence of binding sites for protein-specific cofactors. Previous studies identified two TFs, Hb and Zelda (Zld), as critical for Bcd-dependent activation (Small et al. 1992; Simpson-Brose et al. 1994; Porcher et al. 2010). Binding motifs for both of these proteins are enriched in Bcd-dependent enhancers (Schroeder et al. 2004; Ochoa-Espinosa et al. 2005; Nien et al. 2011; Xu et al. 2014). Consistent with this, a discriminative motif analysis revealed an enrichment of Hb and Zld motifs in regions bound in vivo by Bcd and Bcd + Otd compared with Otd only (Fig. 3I).

We hypothesized that early activation of enhancer activity by Bcd is controlled in part by cofactor interactions and that the addition of Hb and/or Zld sites might convert a late acting enhancer into a relay enhancer that is expressed earlier. We tested this hypothesis on Otd^{L6}, which is bound by Otd at S6–S8 and active only at this later stage.

This enhancer is not active early and is not bound by Bcd in vivo despite having two copies of the relay motif (TGATCC) (Fig. 7A). We added four high-affinity Hb sites to the Otd^{L6} enhancer, which resulted in early activation of expression as an anterior stripe (Fig. 7B). Adding four canonical Zld sites into this element also caused it to be activated early but in a complex pattern along the length of the embryo (Fig. 7C), and adding both Hb and Zld sites had an additive effect on the enhancer's activities (Fig. 7D). Otd^{L6} enhancers containing extra Hb and Zld sites were crossed into *bcd* mutants, which caused a complete loss of expression (data not shown), consistent with the hypothesis that the addition of Hb and Zld sites successfully converted the Otd^{L6} enhancer into an early acting Bcd-dependent enhancer.

Our experiments above suggest that the specific sequence variants TGATCC and TTATCCT are required for the activities of relay and Otd-activated enhancers, respectively (Figs. 3, 5, 6). The Otd^{L6} enhancer contains both sequence variants and thus provides an opportunity to test how each motif might interact with the Zld and Hb cofactors. Mutating the TTATCCT motifs in the Otd^{L6} enhancer causes a loss of expression even though there are two intact copies of the TGATCC sequence (Figs. 6E, 7E). We hypothesized that adding Hb and Zld sites to this inactivated enhancer might “rescue” regulatory activity. Adding Hb and Zld sites to Otd^{L6}ΔTTATCCT resulted in a stripe of expression that was expressed early and maintained in S6–S8 (Fig. 7F), a temporal pattern

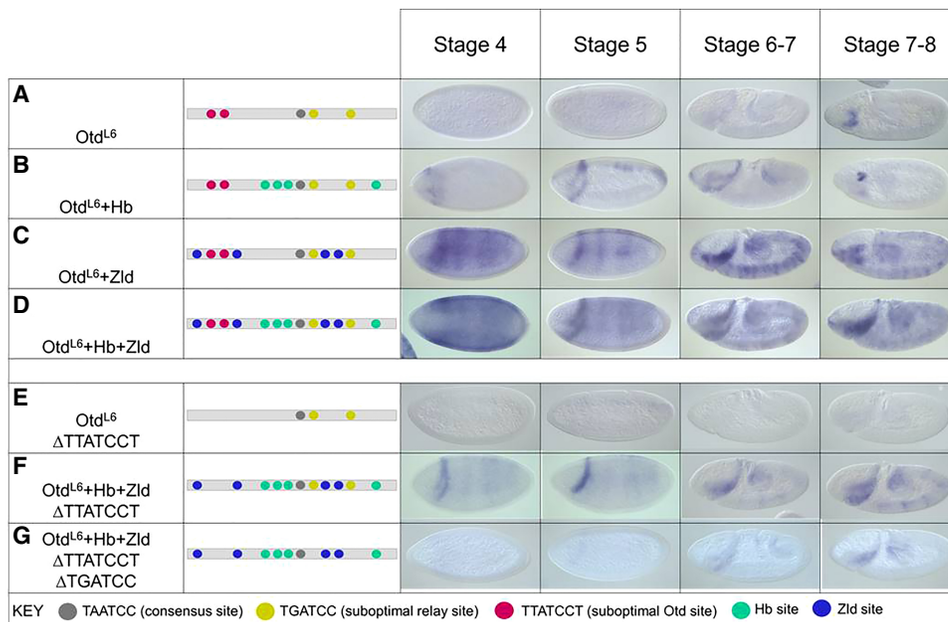


Figure 7. Binding site manipulations that change the timing of enhancer activity. (A–D) *lacZ* RNA expression patterns at four different time points driven by the wild-type Otd^{L6} enhancer (A) and the Otd^{L6} enhancer with four added Hb sites (B), four added Zld sites (C), and four extra Hb sites plus four extra Zld sites (D). (E) *lacZ* expression driven by the Otd^{L6} enhancer carrying mutations in three suboptimal Otd (TTATCCT) sites (compare with the patterns in A). (F) *lacZ* expression driven by the Otd^{L6} enhancer carrying mutations in three TTATCCT sites and four additional sites each for Zld and Hb. (G) *lacZ* expression driven by the Otd^{L6} enhancer carrying mutations in three TTATCCT sites, two TGATCC sites, and four additional sites each for Zld and Hb. See the text for a description of results and interpretations.

similar to that observed for other relay enhancers. This result suggests that the combination of only two TGATCC sites can mediate early activation if augmented with binding sites for Zld and Hb. A further mutation of the two TGATCC sites in the Otd^{L6} + Hb + Zld ΔTTATCCT enhancer resulted in a loss of expression, indicating the importance of those sites for early activation despite the presence of strong binding sites for Zld and Hb (Fig. 7G).

Taken together, our experiments suggest that the specific responses of enhancers to Bcd- and Otd-mediated activation are controlled in part by suboptimal motifs preferred by each protein and that early activation by Bcd requires interactions with the cofactors Hb and Zld.

Discussion

In this study, we compared the *in vivo* functions of two K50HD proteins (Bcd and Otd) within a framework of evolution. We showed that Otd and Bcd have evolved independent functions *in vivo*, and HD swaps between the two proteins indicated that the major structural differences mediating their distinct *in vivo* activities can be traced to their HDs. The two proteins bind to unique enhancers, and Otd also binds enhancers previously bound by Bcd via a FFR mechanism that extends the timing of the gene expression patterns that they regulate. We presented evidence that Otd binding does not lead to enhancer activation in the early embryo but is an effective activator after the first wave of zygotic target gene activation. Finally, we showed that enhancers respond in specific ways to Bcd and Otd through suboptimal binding sites for K50HD protein and binding sites for cofactors.

The Bcd–Otd FFR in Drosophila

Bcd has evolved rapidly to become a powerful morphogen in *Drosophila* but is not well conserved even within Diptera (Stauber et al. 1999). In *Drosophila*, *otd* has evolved to become a zygotic target gene of Bcd (Finkelstein and Perrimon 1990). *otd* RNA appears at S4, and its protein is detectable at early S5—at the same time as most other Bcd target genes. Despite having a lethal mutant phenotype, little is known about Otd's molecular functions in *Drosophila* embryogenesis. Otd binds to many enhancers that were initially activated by Bcd at S4 (Fig. 3). The reduced expression driven by these enhancers in *otd* mutants at S5 suggests that Otd protein binding is functional, and the loss of expression at S6–S8 demonstrates that Otd is required for maintaining their activity (Fig. 4). These results suggest strongly that Bcd and Otd participate in a relay in which Bcd initiates transcriptional activation of target genes, including *otd*, and Otd maintains its own expression and the expression of other target genes after the Bcd gradient decays. This relay is similar to the C1 feed-forward loop (FFL) with an “OR”-like input function described by Alon (2007) but is distinct in two respects. First, all target enhancers regulated by the Bcd–Otd FFR are initially activated by Bcd before Otd is present. Second, the relay involves the sequential binding of Bcd and Otd to the same binding motifs in the same target gene enhancers.

Because Bcd and Otd recognize similar sequence motifs *in vitro* and are involved in an FFR in *Drosophila*, one would predict that a maternal gradient of Otd (Mat:Otd) would activate many Bcd-dependent target genes when Bcd is genetically removed. On the contrary, we found that Mat:Otd cannot activate the great majority (more than 90%) of the tested Bcd-dependent transcriptional targets, including many that are regulated by the Bcd–Otd FFL. Also, our ChIP-seq experiments identified hundreds of peaks that bind Otd but not Bcd in early embryos. We tested 10 such fragments from euchromatic regions, but none showed any enhancer-like activity. Taken together, these results suggest that Otd is unable to efficiently activate transcription on its own in the early embryo.

The most likely explanation for Otd's inability to replace or function well without Bcd is that it is unable to interact efficiently with the Bcd cofactors Zld or Hb. Bcd activates the very first zygotic target genes as part of the maternal-to-zygotic transition (MZT), and a key factor in the MZT is Zld, which is thought to act as a pioneer that opens chromatin (Li et al. 2014; Sun et al. 2015). We showed previously that Zld facilitates Bcd binding to target gene enhancers (Xu et al. 2014). Similarly, most Bcd target genes require the activity of Hb to increase target gene sensitivity to Bcd-dependent activation (Simpson-Brose et al. 1994; Porcher et al. 2010). Interestingly, because insertion of the Bcd HD into the Otd protein confers on it many of Bcd's normal functions, these critical interactions may involve direct contacts with peptide motifs within Bcd's HD. We propose that similar motifs are not present in Otd's HD, which accounts for its inactivity at the early time point.

Mechanisms controlling different enhancer activities

The ChIP-seq experiments enabled us to classify genomic fragments based on their ability to bind Bcd and/or Otd and their temporal expression patterns (Fig. 8). Fragments in class 1 bind Bcd early and Otd late and mediate the FFR described above. Fifty-three of the 66 known Bcd-dependent enhancers belong to this class. Fragments in class 2 are bound early by Bcd and never bound by Otd. Only 13 of the 66 known Bcd-dependent enhancers are in this class. These enhancers are also active after the Bcd gradient degrades but do not bind Otd, and it is not clear how their expression is maintained. Fragments in class 3 (Otd^L) are never bound by Bcd but are bound by Otd at S6–S8. We tested nine Otd^L candidate enhancers, and all showed enhancer activity.

We have begun to understand the molecular mechanisms that distinguish the activities of the three enhancer classes. Motif searches identified two suboptimal motifs, TGATCC and TTATCCT, in class 1 enhancers and class 3 enhancers, respectively (Fig. 3G,H). Mutating these motifs abolished enhancer function, which is consistent with recent studies showing that suboptimal sites play critical roles in other systems (Lebrecht et al. 2005; Rowan et al. 2010; Ramos and Barolo 2013; Crocker et al. 2015; Farley et al. 2016). However, it is important to point out that almost all genomic regions bound by Bcd and Otd

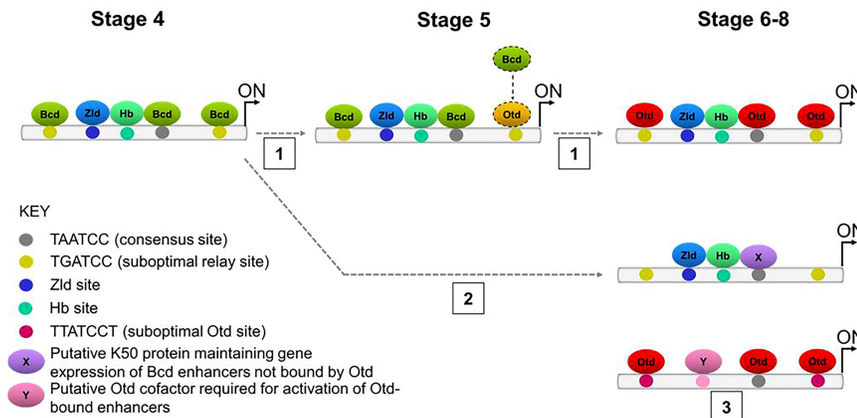


Figure 8. Three classes of Bcd- and Otd-dependent enhancers. Class 1: Bcd-Otd FFR enhancers are activated at S4 by Bcd with the cofactors Hb and Zld. Bcd protein binds to consensus (TAATCC) and suboptimal (TGATCC) sites required for gene activity. At S5, Otd protein binds to suboptimal sites, replacing Bcd. At S6–S8, Otd protein binds to all consensus and suboptimal sites to maintain gene activity. Class 2: These enhancers are activated by Bcd and maintained after the Bcd gradient decays but are never bound by Otd. These enhancers may be regulated by a FFL involving an unknown factor (X). Class 3: Other Otd activates gene targets in the late stage

through binding to both consensus and suboptimal (TTATCCT) sites. Otd enhancer activation at S6–S8 might require another cofactor (Y).

and all enhancers tested here also contain optimal TAATCC motifs. Previous studies have shown that optimal sites are critical for enhancer activation (Driever et al. 1989; Struhl et al. 1989; Stanojevic et al. 1991; Arnosti et al. 1996) and sufficient for *in vivo* activation when clustered (Driever et al. 1989; Schier and Gehring 1992; Hanes et al. 1994; Crauk and Dostatni 2005; Lebrecht et al. 2005; Barr et al. 2017). These studies and our results suggest that both optimal and suboptimal sites contribute binding events required for productive activation of the basal promoter. However, we propose that the presence or absence of a suboptimal site can bias an enhancer toward activation by a specific family member.

Sequence motifs for Zld and Hb were also overrepresented in ChIP-seq peaks bound by Bcd (Fig. 3), consistent with the combinatorial activation model mentioned above. Adding Zld- and Hb-binding sites to a class 3 enhancer normally activated later by Otd results in earlier activation and conversion of that enhancer into a Bcd-dependent class 1 enhancer (Fig. 7B–D,F). We hypothesize that class 3 enhancers contain binding sites for unknown Otd-specific cofactors, and identifying these sites and the proteins that bind them will be the focus of future studies. Taken together, our results indicate that both intrinsic DNA-binding preferences and interactions with cofactors control the distinct temporal and spatial patterns of expression driven by individual enhancers *in vivo*.

Convergent evolution and a robust core anterior patterning network

bcd and its paralog, *zen*, arose through duplication of an ancestral maternally expressed *Hox3*-like gene (Stauber et al. 1999, 2002). In insects lacking Bcd, Otd has Bcd-like properties (maternal expression and anterior mRNA localization and patterning) (Schroder 2003; Lynch et al. 2006). Since the two proteins do not share common ancestry (Otd is a paired class homeoprotein that diverged from the Hox cluster >800 million years ago), some functional convergence of these proteins must have occurred during insect evolution.

The evolution of Bcd likely involved the retention of the maternal promoter and the acquisition of three characteristics required for anterior embryonic patterning: (1) UTR sequences that control anterior mRNA localization, (2) protein motifs that mediate translational repression, and (3) amino acid substitutions that alter its DNA-binding preferences; namely, a Q50-to-K50 mutation in its HD. Since *otd* is zygotic in *Drosophila*, some of these characteristics described above must have been lost in the lineage that led to *Drosophila*. Such dramatic changes in these two genes may be attributed to reduced selective pressure on maternal genes (Barker et al. 2005; Demuth and Wade 2007), which permits the exploration of the evolutionary landscape and the acquisition of new functional roles.

What is striking is that Otd binds to at least half of the Bcd-bound target genes after they are activated by Bcd. Perhaps this set of target genes represents an ancestral core network that is well conserved in evolution. Many feed-forward enhancers are associated with the gap genes, which cross-regulate each other via repressive interactions. All of these enhancers contain DNA motifs that are recognized by a K50HD TF such as Bcd or Otd. Thus, it is possible that the *cis*-regulatory motifs (and consequently the enhancers) are functionally robust in the evolution of anterior embryo patterning, while *trans*-acting factors can accumulate mutations. This allows for a conserved set of targets that makes up a canalized anterior patterning network that allows the regulating TFs to evolve.

Materials and methods

D. melanogaster stocks

The following stocks were used in these experiments: *yw* (wild type), $\pm/Cyo\ bcd^+;bcd^{E1}/bcd^{E1}$, *Cyo bcd⁺/Sco;bcd^{E1}/bcd^{E1}*, and $\Phi C31\ (\gamma+);38F1\ (w+)$.

Maternal gene chimeras

The *bcd*- and *otd*-coding regions were amplified by PCR from pBS-SK⁺ cDNA clones. We cloned an injection plasmid (piattB40-Bcd) using traditional techniques from pBS-SK⁺ (Asp718/SacI)

and industrially synthesized oligonucleotides. This plasmid contained inverted Φ C31-specific recombination sequences (AgeI/HindIII), the fluorescent green eye marker Gmr-Gfp (HindIII/AscI), and a polylinker flanked by the *bcd* promoter and 3' UTR (AgeI/AscI). The amplification product was digested with RsrII and AscI and ligated into the plasmid fragment. HD swaps and residue changes were generated using standard cloning techniques and nested PCR. All transgenic lines were generated using the Φ C31 integration system, and constructs were integrated into the 38F1 landing site on the third chromosome (Bateman et al. 2006).

Embryo collections, cuticle preparations, and immunohistochemistry

Embryos were collected 2–3 and 3–5 h AEL. Embryos were dechorionated for 2 min in bleach, and a 2:1 mixture of methanol and heptane was used to remove the vitelline membrane. ISH and FISH were performed as described previously (Kosman et al. 2004) using digoxigenin-, fluorescein-, and biotin-labeled probes. Cuticle preparations were performed on embryos aged 20–24 h. For cuticle preparations, larvae were fixed overnight at 65°C in a 1:4 mixture of glycerol and acetic acid and mounted in a 1:1 mixture of Hoyer's medium and lactic acid. Rabbit anti-Bcd (1:400) and guinea pig anti-Otd (1:1000) were used for immunostaining (with rabbit FITC and 647 guinea pig as secondary antibodies at 1:400 each; Invitrogen). Guinea pig anti-Cad (1:400) and Alexa flour-conjugated 488 donkey anti-guinea pig (1:500; Invitrogen) were used to examine Cad protein expression. All antibodies were diluted in PBT. Data for immunostaining images were collected on a Leica TCS SP5 confocal microscope using the Leica confocal analysis software. Gradient quantifications were performed as described previously (Ochoa-Espinosa et al. 2009).

ChIP-seq

ChIP was performed using Bcd antibody (rabbit) and two Otd antibodies (guinea pig; GP-5 and GP-6, both provided by Tiffany Cook) on two biological replicates (two technical replicates per sample) of wild-type chromatin collected at 2–3 and 3–5 h AEL. Antibodies were protein A-purified using the protein A antibody purification kit (Sigma). The embryos from each collection were DAPI-stained to confirm embryonic stages, and 85% of the embryos were the correct age in each collection. Embryos were treated and fixed in 1.8% formaldehyde as described previously (Zeitlinger et al. 2007; Xu et al. 2014). Samples were sonicated for 10 min at setting 3 (30 sec on and 30 sec off) followed by 2.5 min at setting 4 (30 sec on and 30 sec off) using a Sonic Dismembrator model 550. Two-hundred microliters of embryonic lysate was used for each immunoprecipitation reaction. Two-hundred-forty microliters of buffer FA + PI (50 mM HEPES/KOH at pH 7.5, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate, 150 mM NaCl, protease inhibitor) was added to each sample to bring the total volume to 440 μ L. Five percent of each extract was removed as input. Ten microliters of each antibody was added to the sample (minus input) and incubated overnight at 4°C. Forty microliters of protein A sepharose bead slurry (Amersham Biosciences) was added per ChIP sample with 1 mL of FA buffer. The sample was centrifuged at 2500g for 1 min, and supernatant was discarded. One milliliter of FA buffer was added to each tube. Beads were suspended by inverting the tubes a few times and then centrifuging them again. This wash was repeated three times. After the final wash, the beads were suspended in 40 μ L of FA buffer. The ChIP sample was added to the bead slurry and rotated for 2 h at 4°C. Two microliters of 20 mg/mL RNase A was added to the inputs. The beads were washed at room temperature by adding

1 mL of each of the following buffers and incubating on a nutator (or a rotator): FA (twice for 5 min), FA–1 M NaCl (once for 5 min; 50 mM HEPES/KOH at pH 7.5, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate, 1 M NaCl), FA–500 mM NaCl (once for 10 min; 50 mM HEPES/KOH at pH 7.5, 1 mM EDTA, 1% Triton X-100, 0.1% sodium deoxycholate, 500 mM NaCl), TEL (once for 10 min; 0.25 M LiCl, 1% NP-40, 1% sodium deoxycholate, 1 mM EDTA, 10 mM Tris-HCl at pH 8.0), and Tris-EDTA (twice for 5 min). The beads were collected after each wash by centrifugation at 2500g for 1 min, and the supernatant was removed. To elute the immunocomplexes, we added 125 μ L of ChIP elution buffer and placed the tube for 15 min in a 65°C heat block. We spun down the beads at 6000g for 1 min and transferred the supernatant to a new tube. The elution was repeated, and supernatants were combined.

ChIP-seq library preparation and data processing

NEXTflex ChIP-seq kits from BIOO Scientific (5143-01) were used to prepare libraries, which were then barcoded using NEXTflex ChIP-seq barcodes (BIOO Scientific, 514120). We performed single-end HiSeq 2000 sequencing (Illumina) at the New York University Genome Center. Sequencing reads were mapped to the *D. melanogaster* genome release 5.3 using Bowtie, and duplicate reads were removed after combining data from each biological replicate. The MACS program was used to call peaks over input (Zhang et al. 2008; Liu et al. 2011). We removed all heterochromatic and uncharacterized chromatin from the analysis and further narrowed down relevant peaks by using only those that appeared in both antibodies (for Otd) and across all replicates. All ChIP-seq data processing was done on the Galaxy Cistrome platform. A false discovery rate cutoff of 5% was used for all further analysis on ChIP-seq peaks.

Motif enrichment analyses

Matlab was used for all motif analyses unless stated otherwise. Bcd and Otd peaks were compared, and peaks were considered shared if they overlapped by at least 200 base pairs. We compared all of our ChIP-seq data sets with prior DNase I, Hb ChIP-chip, and Zld ChIP-seq experiments using the same overlapping criterion. We used MEME-ChIP (Machanic and Bailey 2011) in the MEME suite (Bailey et al. 2009) to search for overrepresented motifs in each ChIP data set. We then used a discriminative de novo motif search to look for overrepresented 6-mers, 7-mers, 8-mers, and 9-mers in ChIP-seq peaks. Cofactor position-weight matrices used to search Bcd- and Otd-bound genomic regions were derived as follows: Bcd and Otd from PBMs done in this study and Zld and Hb from Fly Factor Survey (<http://mccb.umassmed.edu/ffs>).

GST-tagged proteins

HD-coding sequences (as well as 15-amino-acid flanking regions) for Bcd and Otd were PCR-amplified and cloned into a N-terminal GST fusion Gateway expression vector (pDEST15, Invitrogen), and the correct clones were confirmed by sequencing. Rosetta (DE3)-competent cells (BL21 derivatives from Novagen) were transformed with the GST-BcdHD or GST-OtdHD plasmids. For each transformation, a single colony was used to inoculate 0.5 L of LB + Amp¹⁰⁰ and shaken at 200 rpm at 37°C until the OD₆₀₀ reached 0.5–0.6. Proteins were induced for 3 h at 37°C by the addition of 0.5 mM IPTG. After induction, bacteria were harvested by centrifugation at 5000g for 15 min, and the cell pellets were stored at –80°C. Each frozen pellet was resuspended in 40 mL of lysis buffer (150 mM NaCl, 10 mM HEPES at pH 7.9,

Datta et al.

5 mM DTT, 10% glycerol, 0.5% Triton X-100, 0.5 mg/mL lysozyme, 1 mM MgCl₂, 1 mM benzamidine, 3 μM pepstatin A, 2 mM leupeptin, 1 mM PMSF) and sonicated using a Branson Sonifier 450 homogenizer equipped with a midi tip. The sonication parameters were six to eight strokes of 1 min with an output of 6, duty cycle of 50%, and a resting time of 2 min on ice between strokes. The crude lysate was centrifuged at 27,000g for 30 min at 4°C, and the supernatant was transferred to a new tube containing 0.5 mL of settled, prewashed, and pre-equilibrated glutathione sepharose 4B beads (GE Healthcare, 17075601) and rotated for 6 h. Elutions were performed with lysis buffer containing 5 mM (E1 and E2), 7.5 mM (E3), 10 mM (E4 and E5), 20 mM (E6 and E7) and 50 mM (E8 and E9) imidazole. A total of 10 mL of the peak eluates was dialyzed twice for 12 h each time against 1 L of storage buffer (150 mM NaCl, 5 mM HEPES at pH 7.9, 1 mM DTT, 10% glycerol, 1 mM PMSF). After dialysis, the samples were centrifuged at 21,000g for 5 min, aliquoted, flash-frozen in liquid nitrogen, and stored at -80°C.

PBMs

PBMs were performed as described previously (Berger et al. 2006) using a custom-designed “all 10-mer” universal array (Berger et al. 2008) in the 8x60K array format (Agilent Technologies, Inc., Agilent microarray design identifier [AMADID] 030236) (Nakagawa et al. 2013). Both proteins were tested at a concentration of 87 nM. Duplicate PBMs were performed for each protein. The array data were quantified as described previously (Berger et al. 2006), and 8-mer data were averaged over duplicate PBM experiments using the Universal PBM analysis suite (Berger and Bulyk 2009). Motifs were derived by the Seed-and-Wobble algorithm (Berger et al. 2006; Berger and Bulyk 2009), modified to use 90% of the foreground and background features (Busser et al. 2012). As described previously (Busser et al. 2012), each 9-mer was then assigned the *E*-score of the lower scoring of its two constituent 8-mers. In previous assays using PBM data for other *Drosophila* HDs, 9-mers with an *E*-score >0.31 were used with confidence to predict real binding events (Busser et al. 2012). 9-mer sequences bound preferentially by Bcd versus Otd were identified as described previously (Busser et al. 2012).

Generation of otd HD deletion by CRISPR

The CRISPR flies were generated as described (Gratz et al. 2013, 2014). gRNAs were inserted into the pU6-BbsI-chiRNA vector (gRNA1, CTTCGAAAAAAAAAACGAGTTAGC; gRNA2, CTTCGCATATATAAACATTATGTAC). Two homology arms flanking the cleavage sites were inserted into the multiple cloning sites of the pHD-DsRed-attP vector as donor template. The mixture of 500 ng/μL donor vector and two gRNA vectors (100 ng/μL each) was injected into embryos of y[1] M[vas-Cas9].RFP-ZH-2A w[1118]/FM7a. The F1 flies were crossed to YW flies, and F2s were screened for the dsRed⁺ transformants.

Synthetic enhancer constructs, transgenesis, and K50-dependent enhancer analysis

All genomic fragments were cloned into the piB-HC-lacZ vector described previously (Chen et al. 2012) using BglII and AscI. All constructs were inserted using ΦC31 integrase-mediated cassette exchange at 38F1 on chromosome II. We converted low-affinity Zld and Hb sites into high-affinity sites in the OtdL^A enhancer. Synthetic enhancer fragments for BcdOtd^{EL4} (HC_01), BcdOtd^{EL25} (HC_49), and OtdL⁶ were synthesized by Integrated DNA Technologies. Otd^{EL} and Otd^L enhancers were cloned

from genomic DNA. The dependency of enhancers on Otd, Bcd, and Mat > Otd was determined by crossing the enhancers to *otd* and *bcd* mutants and Mat > Otd virgin females, collecting progeny embryos, and checking to see how *lacZ* expression was affected by ISH.

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Rhea R. Datta, Jia Ling, Jesse Kurland, et al.

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