Boundaries support specific long-distance interactions between enhancers and promoters in *Drosophila Bithorax* complex

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Abstract

*Drosophila* bithorax complex (BX-C) is one of the best model systems for studying the role of boundaries (insulators) in gene regulation. Expression of three homeotic genes, *Ubx, abd-A*, and *Abd-B*, is orchestrated by nine parasegment-specific regulatory domains. These domains are flanked by boundary elements, which function to block crosstalk between adjacent domains, ensuring that they can act autonomously. Paradoxically, seven of the BX-C regulatory domains are separated from their gene target by at least one boundary, and must “jump over” the intervening boundaries. To understand the jumping mechanism, the *Mcp* boundary was replaced with *Fab-7* and *Fab-8*. *Mcp* is located between the *iab-4* and *iab-5* domains, and defines the border between the set of regulatory domains controlling *abd-A* and *Abd-B*. When *Mcp* is replaced by *Fab-7* or *Fab-8*, they direct the *iab-4* domain (which regulates *abd-A*) to inappropriately activate *Abd-B* in abdominal segment A4. For the *Fab-8* replacement, ectopic induction was only observed when it was inserted in the same orientation as the endogenous *Fab-8* boundary. A similar orientation dependence for bypass activity was observed when *Fab-7* was replaced by *Fab-8*. Thus, boundaries perform two opposite functions in the context of BX-C – they block crosstalk between neighboring regulatory domains, but at the same time actively facilitate long distance communication between the regulatory domains and their respective target genes.
**Author Summary**

*Drosophila* bithorax complex (BX-C) is one of a few examples demonstrating *in vivo* role of boundary/insulator elements in organization of independent chromatin domains. BX-C contains three *HOX* genes, whose parasegment-specific pattern is controlled by *cis*-regulatory domains flanked by boundary/insulator elements. Since the boundaries ensure autonomy of adjacent domains, the presence of these elements poses a paradox: how do the domains bypass the intervening boundaries and contact their proper regulatory targets? According to the textbook model, BX-C regulatory domains are able to bypass boundaries because they harbor special promoter targeting sequences. However, contrary to this model, we show here that the boundaries themselves play an active role in directing regulatory domains to their appropriate *HOX* gene promoter.

**Introduction**

The three homeotic (*HOX*) genes in the *Drosophila* Bithorax complex (BX-C), *Ultrabithorax* (*Ubx*), *abdominal-A* (*abd-A*) and *Abdominal-B* (*Abd-B*), are responsible for specifying cell identity in parasegments (PS) 5-14, which form the posterior half of the thorax and all of the abdominal segments of the adult fly [1–3]. Parasegment identity is determined by the precise expression pattern of the relevant *HOX* gene and this depends upon a large *cis*-regulatory region that spans 300 kb and is subdivided into nine PS domains that are aligned in the same order as the body segments in which they operate [4–6]. *Ubx* expression in PS5 and PS6 is directed by *abx/bx* and *bxd/pbx*, while *abd-A* expression in PS7, PS8, and PS9 is controlled by *iab-2*, *iab-3*, and *iab-4* [7–10]. *Abd-B* is regulated by four domains, *iab-5*, *iab-6*,...
iab-7 and iab-8, which control expression in PS10, PS11, PS12 and PS13 respectively [6,11,12].

Each regulatory domain contains an initiator element, a set of tissue-specific enhancers and Polycomb Responsible Elements (PREs) and is flanked by boundary/insulator elements (Fig 1A; Maeda and Karch 2006). BX-C regulation is divided into two phases, initiation and maintenance [15,16]. During the initiation phase, a combination of gap and pair-rule proteins interact with initiation elements in each regulatory domain, setting the domain in the on or off state. In PS10, for example, the iab-5 domain, which regulates Abd-B, is activated by its initiator element, while the more distal Abd-B domains, iab-6 to iab-8 are set in the off state (Fig 1B). In PS11, the iab-6 initiator activates the domain, while the adjacent iab-7 and iab-8 domains are set in the off state. Once the gap and pair-rule gene proteins disappear during gastrulation, the on and off states of the regulatory domains are maintained by Trithorax (Trx) and Polycomb (PcG) group proteins, respectively [17,18].

In order to select and then maintain their activity states independent of outside influence, adjacent regulatory domains are separated from each other by boundary elements or insulators [19–25]. Mutations that impair boundary function permit crosstalk between positive and negative regulatory elements in adjacent domains and this leads to the misspecification of parasegment identity. This has been observed for deletions that remove five of the BX-C boundaries (Front-ultraabdominal (Fub), Miscadestral pigmentation (Mcp), Frontadominal-6 (Fab-6), Frontadominal-7 (Fab-7), and Frontadominal-8 (Fab-8)) [6,18,20,21,23,24,26,27].

While these findings indicate that boundaries are needed to ensure the functional autonomy of the regulatory domains, their presence also poses a paradox [14,28]. Seven of the nine BX-C regulatory domains are separated from their target HOX gene by at least one
intervening boundary element. For example, the *iab-6* regulatory domain must “jump over” or “bypass” *Fab-7* and *Fab-8* in order to interact with the *Abd-B* promoter (Fig 1A). That the blocking function of boundaries could pose a significant problem has been demonstrated by experiments in which *Fab-7* is replaced by heterologous elements such as *scs*, *gypsy* or multimerized binding sites for the architectural proteins dCTCF, Pita or Su(Hw) [26,29–31]. In these replacements, the heterologous boundary blocked crosstalk between *iab-6* and *iab-7* just like the endogenous boundary, *Fab-7*. However, the boundaries were not permissive for bypass, preventing *iab-6* from regulating *Abd-B*.

A number of models have been proposed to account for this paradox. One is that BX-C boundaries must have unique properties that distinguish them from generic fly boundaries. Since they function to block crosstalk between enhancers and silencers in adjacent domains, an appealing idea is that they would be permissive for enhancer/silencer interactions with promoters (Fig 1B). However, several findings argue against this notion. For one, BX-C boundaries resemble those elsewhere in the genome in that they contain binding sites for architectural proteins such as Pita, dCTCF, and Su(Hw) [24,31–35]. Consistent with their utilization of these generic architectural proteins, when placed between enhancers (or silencers) and a reporter gene, BX-C boundaries block regulatory interactions just like boundaries from elsewhere in the genome [20,36–42]. Similarly, there is no indication in these transgene assays that the blocking activity of BX-C boundaries are subject to parasegmental regulation. Also arguing against the idea that BX-C boundaries have unique properties, the *Mcp* boundary, which is located between *iab-4* and *iab-5*, is unable to replace *Fab-7* [31]. Like the heterologous boundaries, it blocks crosstalk, but it is not permissive for bypass. A second model is that there are special sequences, called promoter targeting
sequence (PTS), located in each regulatory domain that actively mediate bypass (Zhou and Levine 1999; Chen et al. 2005; Lin et al. 2003). While the PTS sequences that have been identified in iab-6 and iab-7 enable enhancers to “jump over” an intervening boundary in transgene assays, they do not have a similar function in the context of BX-C and are completely dispensable for Abd-B regulation [19,30].

A third model (Fig 1C) is suggested by transgene “insulator bypass” assays [46,47]. In one version of this assay, two boundaries instead of one are placed in between an enhancer and the reporter. When the two boundaries pair with each other, the enhancer is brought in close proximity to the reporter, thereby activating rather than blocking expression. Consistent with a possible role in BX-C bypass, these pairing interactions can occur over large distances and even skip over many intervening boundaries [48–51]. The transgene assays point to two important features of boundary pairing interactions that are likely to be relevant in BX-C. First, pairing interactions are specific. For this reason boundaries must be properly matched with their neighborhood in order to function appropriately. A requirement for matching is illustrated in transgene bypass experiments in which multimerized binding sites for specific architectural proteins are paired with themselves or with each other[52]. Bypass was observed when multimerized dCTCF, Zw5 or Su(Hw) binding sites were paired with themselves; however, heterologous combinations (e.g. dCTCF sites with Su(Hw) sites) did not support bypass.

The fact that both blocking and bypass activities are intrinsic properties of fly boundaries suggests that the BX-C boundaries themselves may facilitate contacts between the regulatory domains and their target genes (Fig 1C). Moreover, the non-autonomy of both blocking and bypass activity could potentially explain why heterologous Fab-7 replacements
like gyspy and Mcp behave anomalously while Fab-8 functions appropriately. Several observations fit with this idea. There is an extensive region upstream of the Abd-B promoter that has been implicated in tethering the Abd-B regulatory domains to the promoter [53–56] and this region could play an important role in mediating bypass by boundaries associated with the distal Abd-B regulatory domains (iab-5, iab-6, iab-7). Included in this region is a promoter tethering element (PTE) that facilitates interactions between iab enhancers and the Abd-B promoter in transgene assays [57,58]. Just beyond the PTE is a boundary element, AB-I. In transgene assays AB-I mediates bypass when combined with either Fab-7 or Fab-8. In contrast, a combination between AB-I and Mcp fails to support bypass [59,60]. The ability of both Fab-7 and Fab-8 to pair with AB-I is recapitulated in Fab-7 replacement experiments.

Unlike Mcp, Fab-8 has both blocking and bypass activity when inserted into Fab-7 [30]. Moreover, its’ bypass but not blocking activity is orientation-dependent. When inserted in the same orientation as the endogenous Fab-8 boundary, it mediates blocking and bypass, while it does not support bypass when inserted in the opposite orientation.

In the studies reported here we have tested this model by replacing the endogenous Mcp boundary with heterologous boundaries. Mcp defines the border between the set of regulatory domains that control abd-A and those that control Abd-B expression (Fig 1A). Unlike the boundaries that are within the Abd-B regulatory domain (e.g. Fab-7 or Fab-8), Mcp is not located between a regulatory domain and its target gene. Instead, it defines the boundary between regulatory domains that target abd-A and those that target Abd-B. For this reason, we expected that it does not need bypass activity. Consistent with this expectation, we find that multimerized dCTCF binding sites fully substitute for Mcp. A different result is obtained for the Abd-B-associated boundaries, Fab-7 and Fab-8. Both boundaries are (for the
most part) able to block crosstalk between the *abd-A* regulatory domain *iab-4*, which specifies A4 (PS9) and the *Abd-B* regulatory domain *iab-5*, which specifies A5 (PS10). Their blocking activity is orientation independent. However, in spite of blocking crosstalk, these replacements still inappropriately induce *Abd-B* expression in A4 (PS9), causing the misspecification of this segment. For the *Fab-7* replacements, this occurred in both orientations, while for the *Fab-8* replacement ectopic induction was only observed when it was inserted in the same orientation as the endogenous *Fab-8* boundary. We present evidence showing that the boundary replacements activate the *Abd-B* gene in A4 (PS9) by inappropriately targeting the *iab-4* domain to the *Abd-B* promoter. In addition to altering the specification of A4 (PS9), the *Fab-7* replacements induce novel transformations of A5 and A6. These findings indicate that when *Fab-7* is inserted into the BX-C in place of *Mcp*, it perturbs the function not only of *iab-4*, but also *iab-5* and *iab-6*.

**Results & Discussion**

**Substitution of *Mcp* by an *attP* integration site in the BX-C**

The *Mcp* boundary is defined by 340 bp core sequence that has enhancer blocking activity in transgene assays [36] and blocks crosstalk between *iab-6* and *iab-7* when substituted for *Fab-7* [31]. Located just distal to the boundary is a PRE that negatively regulates the activity of the *iab-5* enhancers [61]. We used CRISPR to delete a 789 bp DNA segment including the *Mcp* boundary and the PRE and replace it with an *eGFP* reporter flanked by two *attP* sites (*Mcp^attP*) (S1 Fig). The presence of two *attP* sites in opposite
orientation allows integration of different DNA fragments by recombination mediated cassette exchange (RMCE; Bateman et al) using the phiC31 integration system [62].

**Multimerized dCTCF sites substitute for Mcp**

The Mcp boundary marks the division between the set of regulatory domains that control the abd-A and Abd-B genes (Fig 1A). The iab-4 domain is just proximal to Mcp, and it directs abd-A expression in PS9. The iab-5 domain is on the distal side and it regulates Abd-B in PS10. A boundary in this position would be needed to block crosstalk between iab-4 and iab-5; however, neither of these domains would require the intervening boundary to have bypass activity. On the proximal side, iab-4 must bypass the putative Fab-3 and Fab-4 in order to activate the abd-A promoter, while on the distal side, iab-5 must bypass Fab-6, Fab-7 and Fab-8 in order to activate Abd-B. If this expectation is correct, a generic boundary that has blocking activity but is unable to direct iab-4 to the abd-A promoter or iab-5 to the Abd-B promoter should be able to substitute for Mcp. To test this prediction (Fig 2), we introduced either the iab-5 PRE itself (Mcp\textsuperscript{PRE}) or the PRE in combination with four dCTCF sites (Mcp\textsuperscript{CTCF}). In Fab-7 replacement experiments four dCTCF sites in combination with the iab-7 PRE block crosstalk between the iab-6 and iab-7 domains, but do not allow the iab-6 domain to regulate Abd-B expression in PS11 [30].

The Abd-B protein is master regulator of pigmentation in the male abdominal A5 and A6 segments due to the regulation of genes involved in melanin synthesis [63–65]. Flies carrying the null y\textsuperscript{I} allele lack black melanin but still have brown melanin that is also regulated by the Abd-B protein [65,66]. In order to be able to recover recombinants and also
to monitor the blocking activity of the replacement sequence and the on/off state of the iab-5 domain, we included a minimal yellow (mini-y) reporter in our Mcp replacement construct. The mini-y reporter consists of the cDNA and the 340 bp yellow promoter and lacks the wing, body and bristle enhancers of the endogenous yellow gene. As a result, activity of the mini-y reporter depends upon proximity to nearby enhancers. Expression of the mini-y reporter was examined in the y\textsuperscript{1} background.

Based on previous studies [5,22,67], we assume that the activity of this reporter will be determined by the activity state of the iab-5 domain. When iab-5 is off in PS9 and more anterior parasegments, the mini-y reporter will also be off. When iab-5 is on in PS10 and more posterior parasegments, the mini-y reporter will be expressed. This parasegment-specific regulation of the reporter activity will be reflected in the segmental pattern of black melanin pigmentation in the adult cuticle. In replacements in which blocking activity is compromised, mini-y will be expressed in PS9 in adults the A4 tergite will be black, just like the A5 and A6 tergites.

When we replaced the Mcp deletion by the iab-5 PRE alone (Mcp\textsuperscript{PRE}) the mini-y reporter was active not only in A5 (PS10) and more posterior segments, but also in A4 (PS9). As shown in Fig 2, the pigmentation in A4 is black like that in A5 indicating that the reporter is expressed in both segments (Fig 2). This finding shows that, similar to classical Mcp deletions, the Mcp\textsuperscript{PRE} replacement does not have blocking activity. In these Mcp deletions iab-5 is ectopically activated in PS9 by the iab-4 initiator and as a consequence there is a gain-of-function transformation in parasegment identity from PS9 to PS10. We used two approaches to test whether this was true for the Mcp\textsuperscript{PRE} replacement. In the first, we excised the mini-y reporter and introduced an y\textsuperscript{+} X chromosome. Since Abd-B directly regulates
endogenous yellow expression in abdomen [64,66], a transformation of PS9 into PS10 should be accompanied by a PS10-like pattern of pigmentation. Fig 2 shows that this is indeed the case. We also examined the pattern of Abd-B protein expression in the embryonic CNS. In wild type embryos Abd-B is not expressed in PS9, while it is expressed at low levels in PS10. As shown in Fig 3A, Abd-B protein is detected in both PS9 and PS10 at similarly high levels in the Mcp\textsuperscript{PRE} replacement.

As predicted, a quite different result is obtained when we combined the iab-5 PRE with multimerized dCTCF sites. Expression of the mini-y reporter in the Mcp\textsuperscript{CTCF} replacement was restricted to A5 (PS10) and A6 (PS11) as would be expected if the multimerized dCTCF sites block crosstalk between the iab-4 and iab-5 domains (Fig 2). The same pigmentation pattern is observed for the endogenous yellow in the Δmini-y derivative of Mcp\textsuperscript{CTCF}, indicating that Abd-B is not turned on ectopically in PS9. This conclusion is confirmed by antibody staining experiments (Fig 3A). Thus, unlike replacements of Fab-7, a generic boundary can fully substitute for Mcp.

**Substitution of Fab-7 for Mcp disrupts Abd-B regulation in parasegments PS9, PS10 and PS11**

We next tested whether the Fab-7 boundary can substitute for Mcp. The Fab-7 region consists of a minor (HS*) and three major (HS1, HS2 and HS3) nuclease hypersensitive sequences [18,22,23,41,42]. Unlike Mcp or other known or suspected boundaries in BX-C, dCTCF does not bind to Fab-7 [33,68]. Instead, Fab-7 boundary function depends upon two BEN domain protein complexes, Elba and Insensitive, the C\textsubscript{2}H\textsubscript{2} zinc finger protein Pita, and a
large multiprotein complex, called the LBC (Fedotova et al. 2017; Kyrchanova et al. 2017; Wolle et al. 2015; Cleard et al. 2017; Kyrchanova et al. 2018). In addition to a boundary function, the HS3 sequence can also function as a PRE (iab-7 PRE; Mishra et al. 2001; Kyrchanova et al. 2018). In previous studies, we found that a combination of HS1+HS2+HS3 can functionally substitute for the complete Fab-7 boundary in vivo and we used this sequence (named for simplicity F7) for the Mcp replacements (Fig 4). Although Fab-7 has only limited orientation dependence in its endogenous context (Kyrchanova et al. 2016; Kyrchanova et al. 2018), we inserted the HS1+HS2+HS3 sequence in both the forward (same as endogenous Fab-7) and reverse orientations in the Mcp replacement platform. The phenotypic effects of the Fab-7 replacement inserted in the forward orientation, McpF7, are considered first.

Like the McpCTCF replacement, the mini-y reporter is turned on in A5 (PS10) and A6 (PS11) in McpF7 males, and the tergites in both of these segments are black. However, McpF7 differs in two important respects from McpCTCF. First, there are one or two small patches of darkly pigmented cuticle in the A4 tergite (marked by the arrow). These patches are variable and appear to be clonal in origin. This finding indicates that the blocking activity of McpF7 is incomplete, and the mini-y reporter is ectopically activated by the iab-4 domain in a small number of PS9 cells. Second, instead of a stripe of light yellow-brown pigmentation along the posterior margin, nearly the entire A4 tergite is covered in yellow-brown pigmentation. This pattern of pigmentation is not observed in A4 in y1 males carrying the McpCTCF replacement (Fig 2) and the mini-y reporter or for that matter in wild type y1 males (see Fig 4). The presence of yellow-brown pigmentation throughout most of the A4 tergite suggests that the cells in this segment (PS9) are not properly specified. This is the case. When the mini-y
reporter was excised and replaced by the endogenous X-linked \( y^+ \) gene, the A4 tergite has a black pigmentation like A5 and A6 (Fig 4). Since expression of the \textit{yellow} gene is controlled by \textit{Abd-B}, this observation indicates that \textit{Abd-B} must be ectopically activated throughout A4. Antibody staining experiments of the CNS in \textit{Mcp}\(^{F7}\) embryos indicate that this inference is correct (Fig 3B).

A simple interpretation of these findings is that \textit{Mcp}\(^{F7}\) is unable to block crosstalk between \textit{iab-4} and \textit{iab-5} and, as a result, \textit{iab-5} is ectopically activated in all PS9 cells. However, such interpretation is inconsistent with the expression pattern of the \textit{mini-y} reporter; it is only activated in small clones in the A4 tergite and not in the entire A4 tergite. By way of comparison, the black pigmentation generated by the reporter in \textit{Mcp}\(^{PRE}\), which has no boundary activity, was clearly quite different from the light yellow-brown pigmentation observed for the reporter in \textit{Mcp}\(^{F7}\).

There are other reasons to think that this simple interpretation is incorrect and that \textit{Mcp}\(^{F7}\) replacement has a much more complicated effect on the operation of \textit{iab-4} and of the regulatory domains that control \textit{Abd-B} expression. In wild type males, the A6 sternite has a banana shape and no bristles, while the A5 and A4 sternites resemble isosceles trapezoids and are covered with bristles. As can be seen in Fig 4, the A4 and A5 sternites in \textit{Mcp}\(^{F7}\) males are split into two connected lobes and resemble the banana shape of the A6 sternite. These morphological changes are indicative of a GOF transformation of both A4 (PS9) and A5 (PS10) toward an A6 (PS11) identity. This type of transformation is not observed in \textit{Mcp} boundary deletions, nor is it observed in the \textit{Mcp}\(^{PRE}\) replacement.

Further evidence of a novel A4/A5\(\rightarrow\)A6 transformation can be seen in the pattern of trichome hairs in the tergites. In wild type flies, the A4 and A5 tergites are covered with
trichomes, while trichomes are only found along the anterior and ventral margins of the A6 tergite (see darkfield image in Fig 4). In the $Mcp^{F7}$ replacement, there are large regions of the A4 and A5 tergite that are devoid of trichomes. There are even anomalies in A6: the band of trichomes along the anterior margin is absent. Similar alterations in cuticular phenotypes are observed in $Mcp^{F7}$ females (S2 Fig). These findings indicate that the normal regulation of $Abd-B$ is disrupted in several parasegments when $Mcp$ is replaced by the $Fab-7$ boundary.

In its endogenous context, the functioning of $Fab-7$ is weakly orientation dependent. For this reason, we anticipated that the reverse $Mcp$ replacement, $Mcp^{F7R}$, would alter the $Abd-B$ expression pattern in several parasegments and give a similar though perhaps milder spectrum of phenotypic effects. Fig 4 shows that this is the case. In $y^+$ background, large regions of the A4 tergite have a black pigmentation like A5 and A6. The ectopic activation appears to be weaker than in the $Mcp^{F7}$ replacement as there are regions in A4 in which the endogenous $yellow$ gene is not turned on. Also, and unlike $Mcp^{F7}$, there are no bald patches in the A4 trichomes, while the sternite appears to have a normal isosceles trapezoid shape. However, the novel transformations seen in $Mcp^{F7}$ in the more posterior segments A5 (PS10) and A6 (PS11) are still evident. The A5 tergite is not completely covered with trichomes, while the trichomes along the anterior margin of A6 are absent. The A5 sternite is also misshapen. Thus, like $Mcp^{F7}$, introducing a reversed $Fab-7$ boundary in place of $Mcp$ disrupts $Abd-B$ regulation across several parasegments.
The *Fab-8* boundary displays orientation-dependent effects on ectopic activation of *Abd-B* in the A4 abdominal segment

In previous *Fab-7* replacement experiments we found that a 337 bp fragment (*F8*\textsuperscript{337}) spanning the *Fab-8* boundary nuclease hypersensitive site is sufficient to fully rescue a *Fab-7* boundary deletion (Kyrchanova et al. 2016). In the direct (forward) orientation this fragment not only blocks crosstalk but also supports bypass. However, when the orientation of the *Fab-8* boundary is reversed, bypass activity is lost, while blocking is unaffected. Since *F8*\textsuperscript{337} appears to be fully functioning, we inserted this fragment in both orientations next to the *iab-5 PRE* in the *Mcp* deletion (*Mcp*\textsuperscript{F8} and *Mcp*\textsuperscript{F8R}).

The effects of the *Fab-8* replacement in the reverse orientation, *Mcp*\textsuperscript{F8R}, will be considered first. Like the *Mcp*\textsuperscript{CTCF} replacement, *Mcp*\textsuperscript{F8R} blocks crosstalk between *iab-4* and *iab-5* and the *mini-y* reporter is off in A4 (Fig 5). After the deletion of the *mini-y* reporter and introducing a wild type *y*\textsuperscript{+} allele, the pigmentation in the adult male abdomen is equivalent to that in wild type flies. The morphological features of *Mcp*\textsuperscript{F8R} tegites and sternites also resemble those in wild type flies or the *Mcp*\textsuperscript{CTCF} replacement and there is no indication of the other abdominal transformations seen in the *Fab-7* replacements. Consistent with the phenotype of the adult cuticle, the pattern of *Abd-B* expression in the embryonic CNS resembles wild type (Fig 3B). Thus, the *Mcp*\textsuperscript{F8R} replacement fully substitutes for the endogenous *Mcp* boundary.

A different result is obtained when the *F8*\textsuperscript{337} sequence is inserted in its normal forward orientation. Like the reverse orientation *Mcp*\textsuperscript{F8R}, *Mcp*\textsuperscript{F8} efficiently blocks crosstalk between *iab-4* and *iab-5* and the *mini-y* reporter is not activated in A4 (PS9). On the other hand, like
the Fab-7 replacements (Mcp\textsuperscript{F7} and Mcp\textsuperscript{F7R}) most of the A4 tergite is covered in a light yellow-brown pigmentation instead of the normal stripe of yellow-brown pigmentation along the posterior margin of the tergite that is seen in y\textsuperscript{1} males. Moreover, when the reporter is excised and the y\textsuperscript{1} allele replaced by the wild type y\textsuperscript{+} gene, nearly the entire A4 tergite is black. Consistent with the induction of y\textsuperscript{+} expression in A4, Abd-B is active in PS9 in the embryonic CNS (Fig 3B). The GOF transformation of A4 (PS9)\(\rightarrow\)A5 (PS10) is not the only anomaly in Mcp\textsuperscript{F8} flies. While there does not seem to be any misspecification of the tergite or sternites in A5 (PS10), the line of trichomes along the anterior margin of the A6 tergite is disrupted or absent altogether indicating that there are abnormalities in the temporal and/or special pattern of Abd-B expression in PS11.

**Ectopic expression of Abd-B in A4 (PS9) requires a functional iab-4 domain**

In the Fab-7 replacement experiments, the relative orientation of the Fab-8 boundary was thought to be important because it determined whether the chromatin loops formed between the replacement boundary and the AB-I element and/or the PTE sequence upstream of the Abd-B transcription start site were circle loops or stem loops [30,74]. In the forward orientation circle loops are expected to be formed and in this configuration, the downstream iab-5 regulatory domain is brought into close proximity with the Abd-B promoter. In the reverse orientation, iab-6 and iab-7 are predicted to form stem loops, and this configuration would tend to isolate the iab-5 regulatory domain from the Abd-B promoter.

It seemed possible that a similar mechanism might be in play in the Fab-8 replacements of Mcp. In the forward orientation (Mcp\textsuperscript{F8}), the iab-4 regulatory domain would
be brought into close proximity to the *Abd-B* gene, activating its ectopic expression in A4 (PS9). In the opposite orientation, the spatial relationship between the *iab-4* domain and the *Abd-B* promoter would not be conducive for activation. In this case, *Abd-B* would be off in A4 (PS9). A strong prediction of this model is that the inappropriate activation of *Abd-B* in PS9 in the *Mcp*$_{F8}$ replacement should depend on a functional *iab-4* domain.

To test this prediction, we used CRISPR (see S3 Fig) to delete a 4,401 bp sequence (*iab-4*$_{Δ}$) that spans the putative *iab-4* initiation element in flies carrying the *Mcp*$_{F8}$ replacement. The *iab-4*$_{Δ}$ sequence was selected based on the clustering of multiple binding sites for embryonic gap and pair-rule gene proteins [75]. Fig 5 shows that the ectopic activation of $y^+$ in A4 in *Mcp*$_{F8}$ flies was eliminated by the *iab-4*$_{Δ}$ deletion. Moreover, and as predicted, *Abd-B* was not activated in A4 (PS9) in the embryonic CNS of *iab-4*$_{Δ}$ *Mcp*$_{F8}$ embryos (Fig 3B). Interestingly, the loss of trichomes along the anterior margin of the A6 tergite in *Mcp*$_{F8}$ also seemed to depend on a functional *iab-4* domain. As can be seen in Fig 5, the trichome pattern in the A6 tergite of *iab-4*$_{Δ}$ *Mcp*$_{F8}$ flies resembled that of wild type.

**Conclusion**

*Mcp* defines the boundary between the regulatory domains that control expression of *abd-A* and *Abd-B*. In this location, it is required to block crosstalk between the flanking domains *iab-4* and *iab-5*, but it does not need to mediate bypass. In this respect, it differs from the boundaries that are located within the set of regulatory domains that control either *abd-A* or *Abd-B*, as these boundaries must have both activities. Consistent with this limited role, we found that *Mcp* can be replaced by multimerized binding sites for the dCTCF protein. Quite
different results are obtained when *Mcp* is replaced by *Fab-7* or *Fab-8*. Although *Fab-7* is able to block crosstalk between *iab-4* and *iab-5*, its blocking activity is incomplete and there are small clones of cells in which the *mini-y* reporter is activated in A4. In contrast, the *mini-y* reporter is off throughout A4 in the *Fab-8* boundary replacements, indicating that it efficiently blocks crosstalk between *iab-4* and *iab-5*. One plausible reason for this difference is that *Mcp* and the boundaries flanking *Mcp* (*Fab-4* and *Fab-6*) utilize dCTCF as does *Fab-8*, while this architectural protein does not bind to *Fab-7* [33].

In spite of their normal (or near normal) blocking activity, both boundaries still perturb *Abd-B* regulation. In the case of *Fab-8*, the misregulation of *Abd-B* is orientation dependent just like the bypass activity of this boundary when it is used to replace *Fab-7* [30]. When inserted in the reverse orientation, *Fab-8* behaves like multimerized dCTCF sites and it fully rescues the *Mcp* deletion. In contrast, when inserted in the forward orientation, *Fab-8* induces the expression of *Abd-B* in A4 (PS9), and the misspecification of this parasegment. Our results, taken together with our previous studies [30,59,60], support a model in which the chromatin loops formed by *Fab-8* inserted at *Mcp* in the forward orientation brings the enhancers in the *iab-4* regulatory domain in close proximity to the *Abd-B* promoter, leading to the activation of *Abd-B* in A4 (PS9). In contrast, when inserted in the opposite orientation, the chromatin loops formed by the ectopic *Fab-8* boundary are not permissive for interactions between *iab-4* and the *Abd-B* promoter. Importantly, the ectopic activation of *Abd-B* in A4 when *Fab-8* is inserted in the forward orientation suggests that the bypass activity has a predisposed preference, namely it is targeted for interactions with the *Abd-B* gene. From this perspective, it would appear that boundary bypass for the regulatory domains that control *Abd-B* expression is not a passive process in which the boundaries are simply permissive for
interactions between the regulatory domains and the *Abd-B* promoter. Instead, it appears to be an active process in which the boundaries are responsible for bringing the regulatory domains into contact with the *Abd-B* gene. It clearly will be of interest to test the out of context functional properties of the boundaries associated with the *abd-B* and *Ubx* genes to see if they behave like *Fab-7* and *Fab-8*.

While similar conclusions can be drawn from the induction of *Abd-B* expression in A4 (PS9) when *Fab-7* is inserted in place of *Mcp*, this boundary causes even more profound disruptions in the normal pattern of *Abd-B* regulation. In the forward orientation, A4 and A5 are transformed towards an A6 identity, while A6 is also misspecified. Similar though somewhat less severe effects are observed when *Fab-7* is inserted in the reverse orientation. Although the mechanisms responsible for these novel phenotypic effects are uncertain, a plausible idea is that pairing interactions between the *Fab-7* insert and the endogenous *Fab-7* boundary disrupt the normal topological organization of the regulatory domains in a manner similar to that seen in boundary competition transgene assays [76]. Further studies will be required to test this idea.

**Materials and Methods**

**Generation of *Mcp*_{attP} by CRISPR/Cas9-induced homologous recombination**

The backbone of the recombination plasmid was designed *in silico* and contains several genetic elements in the following order: [MCS5]-[attP]-[3xP3-EGFP-SV40polyA]-[attP]-[FRT]-[MCS3]. This DNA fragment was synthesized and cloned into pUC57 by Genewiz. The two multiple cloning sites MCS5 and MCS3 were used to clone homology
arms into this plasmid. The orientations of two the \textit{attP} sites are inverted relative to each other and serve as targets for \textit{ф}C31-mediated recombination mediated cassette exchange [77]. The \textit{3x3P-EGFP} reporter [78] was introduced as a means to isolate positive recombination events. The \textit{Flp}-recombinase target \textit{FRT} [79] were included for the deletion of the selectable \textit{mini-yellow} marker after recombination mediated cassette exchange.

Homology arms were PCR-amplified from \textit{yw} genomic DNA using the following primers: CCTGCCGACTGAACGAATGC and ACGCCCTGATCCCGATACACATAC for the proximal arm (\textit{iab}-4 side; 3967 bp fragment), and GCGTTTTGTGTTAGTAAATGTATC and AAAGGCCAACAAAGAAGACATGGACG for the distal arm (\textit{iab}-5 side; 4323 bp fragment). A successful homologous recombination event will generate a 789 bp deletion within the \textit{Mcp} region (Genome Release R6.22: 3R:16’868’830 – 16’869’619; or complete sequence of BX-C (Martin et al. 1995): 113821 - 114610).

The recombination plasmid was injected into \textit{yw} \textit{vas-Cas9} embryos together with two gRNAs containing the following guides: GCTGGCTTTTACAGCATTTC and GCTTTGTTACCCCTGAAAAT. Concentrations were as described in Gratz et al.[80]. The injected embryos were grown to adulthood and crossed with \textit{yw} partners. Among the few fertile crosses, one produced many larvae with a clear GFP-signal in the posterior part of their abdomens. This observation suggested that these animals had successfully integrated the recombination plasmid and that the \textit{3x3P-EGFP} reporter acts as an enhancer trap for \textit{Abd-B} specific enhancers. GFP positive larvae were isolated and grown to adulthood. Emerging males showed the expected \textit{Mcp} phenotype. Also, and as expected for a reporter located in the BX-C, no fluorescence signal could be detected in their eyes, indicating that the \textit{3x3P-EGFP} reporter is silenced in eye cells where the \textit{3x3P} promoter is usually active. The planned
homologous recombination event could later be verified by PCR and sequencing. We will refer to it as $Mcp^{attP}$.

12 EGFP- and Mcp-positive candidate males were individually crossed with $y\ w$ virgins. Only 2 were fertile. The sterility of others may be caused by presence of off-targets as a frequent non-specific result of CRISPR/Cas9 editing. Starting from the two fertile crosses, 2 independent balanced stocks could be obtained according to established crossing schemes. One of them was used to obtain a $y\ w\ M\{\text{vas-integrase}\}zh-2A\ ;\ Mcp^{attP}/TM3,Sb$ stock for recombination mediated cassette exchange. Because of poor survival rates in injection experiments, the $Mcp^{attP}$ chromosome was also temporarily combined with $Dp(3R)P9, Sb$ ($y\ w\ M\{\text{vas-integrase}\}zh-2A\ ;\ Mcp^{attP}/Dp(3R)P9, Sb$). By selection we obtained homozygous $Mcp^{attP}$ line that was subsequently used for fly injections.

**Generation of $iab-4^{A}$ by CRISPR/Cas9-induced homologous recombination**

For generating dsDNA donors for homology-directed repair we used $pHD-DsRed$ vector that was a gift from Kate O'Connor-Giles (Addgene plasmid # 51434). The final plasmid contains genetic elements in the following order: [$iab-4$ proximal arm]-[attP]-[lox]-[$3xP3$-dsRed-SV40polyA]-[lox]-[$iab-4$ distal arm]. Homology arms were PCR-amplified from $yw$ genomic DNA using the following primers:

TTTGAAATTTCTTCCAGACACGCATCGGG and

AAACATATGCTTGCTATCGACCCTCCTC for the proximal arm (846 bp fragment), and

AATACTAGTCCTCGGAAAGGAAGAAAGTTTC and

TACTCGAGCCGCTAAAAGGAGGTCTTCG for the distal arm (835 bp fragment). A successful homologous recombination event will generate a 4401 bp deletion within the $iab-4$
region (Genome Release R6.22: 3R:16,861,368..16,869,768; or complete sequence of BX-C [4]: 120073-115673).

Targets for Cas9 were selected using “CRISPR optimal target finder” – the program from O'Connor-Giles Lab. The recombination plasmid was injected into McpF8 vasa-Cas9 embryos together with two gRNAs containing the following guides:

ATAGCAAGTAGGAGTGGAGT and GAACCTCTTCCCTTTCCGAGCGG.

Concentrations were as described in Gratz et al. (2014). Injectees were grown to adulthood and crossed with y w; TM6/MKRS partners. Flies with clear dsRed-signal in eyes and the posterior part of their abdomens were selected into a new separate line. The successfully integration of the recombination plasmid was verified by PCR.

Cuticle preparations

Adult abdominal cuticles of homozygous enclosed 3-4 day old flies were prepared essentially as described in (Kyrchanova et al. 2017) and mounted in 100% glycerol.

Photographs in the bright or dark field were taken on the Nikon SMZ18 stereomicroscope using Nikon DS-Ri2 digital camera, processed with ImageJ 1.50c4 and Fiji bundle 2.0.0-rc-46.

Embryo immunostaining

Primary antibodies were mouse monoclonal anti-Abd-B at 1:100 dilution (1A2E9, generated by S.Celniker, deposited to the Developmental Studies Hybridoma Bank) and polyclonal rabbit anti-Engrailed at 1:1000 dilution (a kind gift from Judith Kassis). Secondary antibodies were goat anti-mouse Alexa Fluor 488 and anti-rabbit Alexa Fluor 647 (Molecular
Probes) at 1:2000 dilution. Stained embryos were mounted in the following solution: 23% glycerol, 10% Mowiol 4-88, 0.1M Tris-HCl pH 8.3. Images were acquired on Leica TCS SP-2 confocal microscope and processed using GIMP 2.8.16, ImageJ 1.50c4, Fiji bundle 2.0.0-rc-46.

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Figure captions

Fig 1. Models of an enhancer – promoter interactions in BX-C. (A) Regulatory region of the distal part of the BX-C. Horizontal arrows represent transcripts for abd-A (blue) and Abd-B (green). iab enhancers are shown as ovals color-coded with respect to the gene they control (darker shades of color indicate higher expression levels). The arrow arches are a graphical illustration of the targeting of each cis-regulatory domain to the abd-A or Abd-Bm promoter. Vertical lines mark boundaries (Fab, Fab-3, Fab-4, Mcp, Fab-6, Fab-7, and Fab-8) of regulatory iab domains which are delimited by brackets behind the map. There is also a boundary-like element AB-I upstream of the Abd-B promoter that has communicator activity in bypass assays. (B) and (C) Schematic representation of the models explaining interaction of the iab enhancers with the Abd-B promoter.

Fig 2. The CTCF sites block crosstalk between the iab-4 and iab-5 domains. (A) Molecular maps of the Mcp boundary. The coordinates of the Mcp\textsuperscript{amP} deletion and Mcp\textsuperscript{PRE} Mcp\textsuperscript{CTCF} replacement fragments according to complete sequence of BX-C in SEQ89E numbering [4] are shown below. DNAse hypersensitive site is shown as a light gray box above the coordinate bar. Binding sites for GAF, Pita and dCTCF are indicated by blue, red and green ovals, respectively. PRE element from iab-5 is marked as a blue stripe. Replacement fragments are shown below. (B) The cuticle preparations of wt, Mcp\textsuperscript{PRE} and Mcp\textsuperscript{CTCF} males. The morphology of the 2th to 6th abdominal segments is shown. Abnormalities in segment phenotype are shown by the red arrows. The localization of trichomes on the 4th to 6th abdominal tergites are shown in dark field.

Fig 3. Expression of Abd-B in Mcp replacement embryos. (A) Abd-B expression in wt, Mcp\textsuperscript{PRE} and Mcp\textsuperscript{CTCF} embryos. (B) Abd-B expression in Mcp\textsuperscript{F7}, Mcp\textsuperscript{F8}, Mcp\textsuperscript{F8R} and iab-4\textsuperscript{M} Mcp\textsuperscript{F8} embryos. Each panel shows an image of the embryonic CNS of stage 14 embryos stained with antibodies to ABD-B (red) and Engrailed (En, green). En is used to mark
parasegments, which are numbered from 9 to 14 on the right side of the panels; approximate positions of segments are shown on the left side of the wild type (wt) panel and marked A4 to A8. The wild type expression pattern of Abd-B in the embryonic CNS is characterized by a stepwise gradient of increasing protein level from PS10 to PS14. The $Mcp^{F8}$ or $Mcp^{F7}$ embryos have similar low Abd-B expression in PS9 and PS10. The Abd-B expression in PS9 is absent in $iab-4^A Mcp^{F8}$ and $Mcp^{F8R}$ embryos.

**Fig 4.** $Mcp^{F7}$ and $Mcp^{F7R}$ support Abd-B activation in the A4 segment. (A) Schematic representation of the $Fab-7$ boundary. The 1.1 kb $Fab-7$ replacement consists of HS1, HS2 and HS3 ($iab-7$ PRE) regions (shown as gray boxes). (B) Morphology of the 2th to 6th abdominal segments in $Mcp^{F7}$ and $Mcp^{F7R}$ males. Other designations are as in Fig 2.

**Fig 5.** Activation of Abd-B by the $iab-4$ enhancer depends on the orientation of the $Fab-8$ insulator in $Mcp^{F8}$ and $Mcp^{F8R}$ mutants. (A) Molecular maps of the $Fab-8$ boundary and $F8^{337}$. The $Fab-8$ insulator is shown as a horizontal bar. The proximal and distal deficiency endpoints of the $Fab-8$ deletions are shown below. For other designations see Fig 2. (B) Morphology of the 2nd to 6th abdominal segments in insulator in $Mcp^{F8}$, $Mcp^{F8R}$ and $iab-4^A Mcp^{F8}$ males. Other designations are as in Fig 2.

**Supporting information captions**

**S1 Fig.** The strategy to create Mcp replacement lines. On the top: schematic representation of regulatory region of the $abd-A$ and Abd-B genes (blue and green, respectively). The 789 bp Mcp region that was deleted (coordinates according to complete sequence of BX-C in SEQ89E numbering) and replaced by two attP sites for the integration of the tested constructs. $3xP3$-eGFP was used as a marker gene. $frt$ site was used for excision of yellow
maker gene. The plasmid that was injected into Mcp<sup>attP</sup> line, contains two attB site for integration, iab-5 PRE for restoring functional integrity of the iab-5 domain, the frt site for excision of yellow gene, lox sites for excision of testing element. Testing elements were inserted just in front of iab-5 PRE.

**S2 Fig. The abdominal cuticles of wt, Mcp<sup>F8</sup>, Mcp<sup>F8R</sup>, Mcp<sup>F7</sup> and Mcp<sup>F7R</sup> females.**

Morphology of the 2<sup>th</sup> to 6<sup>th</sup> abdominal segments in wt, Mcp<sup>F8</sup>, Mcp<sup>F8R</sup>, Mcp<sup>F7</sup> and Mcp<sup>F7R</sup> females. The expression of mini-γ (black pigment) is shown on the upper panel. Localization of trichomes on tergites is shown lower.

**S3 Fig. The strategy to create iab-4 deletion.** The scheme of the regulatory region in the distal part of the BX-C. Horizontal arrows represent transcripts for abd-A (blue) and Abd-B (green). The iab-4 region was selected using FlyBase, based on the clustering of multiple binding sites for embryonic gap and pair-rule gene proteins. The screenshot show localization of the 4401 bp of iab-4 deletion with R6 genome release coordinates. The coordinates of iab-4 deletion according to complete sequence of BX-C (in SEQ89E numbering ) are 120073-115673 (shown lower). The deletion was made using CRISPR/Cas9 strategy. Targets for Cas9 were selected using “CRISPR optimal target finder” – program from O’Connor-Giles Lab. Vector for generating dsDNA donors for homology-directed repair contains the visible marker 3xP3-DsRed. pH-DsRed was a gift from Kate O’Connor-Giles (Addgene plasmid # 51434). dsRed gene was using for selection of flies with iab-4 deletion.
